# Assignment 2

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### Exercise 1

a) To investigate whether tree type influences total wood volume, we can perform a one-way ANOVA.

```
tree_df$type <- as.factor(tree_df$type)
tree_type_lm <- lm(volume~type, data=tree_df)
anova(tree_type_lm)

## Analysis of Variance Table
##
## Response: volume
## Df Sum Sq Mean Sq F value Pr(>F)
## type 1 380 380 1.9 0.17
## Residuals 57 11395 200
```

With p > 0.05, we can conclude that type does not have a significant effect on volume. Because the factor type has two levels, we can apply a two sample t-test.

```
mask <- tree_df$type == "beech"
t.test(tree_df$volume[mask], tree_df$volume[!mask])</pre>
```

This supports the result from the ANOVA test. The estimated volume is 30.2 for Beech trees and 35.2 for Oak trees.

### b)

To investigate this claim, we create two models, each including all three explanatory variables (type, diameter and height). In the first model, we also include the pairwise interaction between type and diameter.

```
tree_type_d_lm <- lm(volume~height+type*diameter, data=tree_df)
anova(tree_type_d_lm)</pre>
```

```
## Analysis of Variance Table
##
## Response: volume
##
                 Df Sum Sq Mean Sq F value Pr(>F)
## height
                  1
                       2188
                               2188 206.21 < 2e-16 ***
## type
                  1
                                      40.65 4.2e-08 ***
                        431
                                431
## diameter
                  1
                       8577
                               8577
                                     808.49 < 2e-16 ***
## type:diameter
                         6
                                  6
                                       0.52
                                               0.47
                  1
## Residuals
                 54
                       573
                                 11
## ---
## Signif. codes: 0 '*** 0.001 '** 0.01 '* 0.05 '.' 0.1 ' ' 1
```

In the second model, we include the pairwise interaction between type and height.

```
tree_type_h_lm <- lm(volume~diameter+type*height, data=tree_df)
anova(tree_type_h_lm)</pre>
```

```
## Analysis of Variance Table
##
## Response: volume
##
               Df Sum Sq Mean Sq F value Pr(>F)
## diameter
                   10827
                            10827 1045.97 < 2e-16 ***
## type
                                     4.37
                                            0.041 *
                1
                       45
                               45
## height
                1
                     324
                              324
                                    31.32 7.5e-07 ***
## type:height
               1
                      19
                               19
                                     1.88
                                            0.176
## Residuals
               54
                     559
                               10
## ---
## Signif. codes: 0 '*** 0.001 '** 0.01 '* 0.05 '.' 0.1 ' ' 1
```

We see that both pairwise interactions are not significant. Therefore, we can conclude that both height and diameter have the same influence on volume regardless of type.

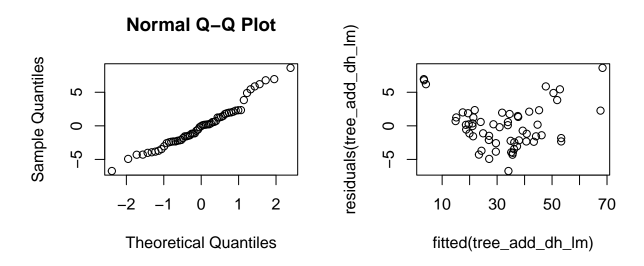
 $\mathbf{c}$ 

In (b), we saw that the interactions of *height* and *diameter* with *type* were not significant, and so we will investigate a purely additive model (assuming no interactions).

```
tree_add_all_lm <- lm(volume~diameter+height+type, data=tree_df)
drop1(tree_add_all_lm, test= "F")
## Single term deletions
## Model:
## volume ~ diameter + height + type
            Df Sum of Sq RSS AIC F value Pr(>F)
## <none>
                          578 143
## diameter 1
                    8577 9155 304 815.61 < 2e-16 ***
## height
             1
                    324 903 167
                                     30.82 8.4e-07 ***
## type
             1
                      23 602 143
                                      2.21
                                              0.14
## ---
## Signif. codes: 0 '*** 0.001 '** 0.01 '* 0.05 '.' 0.1 ' ' 1
We see that the effect of type is not significant in the additive model. Therefore we will investigate
an additive model that excludes type.
tree_add_dh_lm <- lm(volume~diameter+height, data=tree_df)</pre>
anova(tree_add_dh_lm)
## Analysis of Variance Table
##
## Response: volume
             Df Sum Sq Mean Sq F value Pr(>F)
## diameter
                10827
                         10827 1007.8 < 2e-16 ***
              1
## height
                   346
                           346
                                   32.2 5.1e-07 ***
              1
## Residuals 56
                   602
                            11
## ---
## Signif. codes: 0 '*** 0.001 '** 0.01 '* 0.05 '.' 0.1 ' 1
summary(tree_add_dh_lm)
##
## Call:
## lm(formula = volume ~ diameter + height, data = tree_df)
##
## Residuals:
      Min
              1Q Median
                            3Q
                                   Max
## -6.724 -2.278 -0.034 1.820 8.629
##
## Coefficients:
               Estimate Std. Error t value Pr(>|t|)
## (Intercept) -64.3697
                            5.5577 -11.58 < 2e-16 ***
## diameter
                 4.6325
                            0.1602
                                     28.92 < 2e-16 ***
```

```
## height 0.4289 0.0755 5.68 5.1e-07 ***
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
##
## Residual standard error: 3.28 on 56 degrees of freedom
## Multiple R-squared: 0.949, Adjusted R-squared: 0.947
## F-statistic: 520 on 2 and 56 DF, p-value: <2e-16</pre>
```

This model has a high R-squared value while using fewer variables, all of which are significant. Since simpler models are generally preferred, this is our model of choice to make predictions. As a final test, we need to check this model's assumptions to ensure that the conclusions we draw from it are valid:



While these plots are not perfect, we believe the model assumptions to be valid.

Therefore, the effects of type, diameter and height can be summarized as follows:

- The tree *type* does not affect volume significantly.
- Looking at the coefficients, we see that increasing both height and diameter result in an increase in volume, with diameter having a bigger impact (with a gradient of 4.63 compared to *height's* 0.43). This makes sense given that we know volume is proportional to the square of the diameter.

To predict the volume for a tree with the overall average diameter and height, we can use the following linear regression model:

$$volume = -64.37 + 4.63 * diameter + 0.43 * height$$

```
mean_d <- mean(tree_df$diameter)
mean_h <- mean(tree_df$height)
means <- data.frame(diameter=c(mean_d), height=c(mean_h))

predict(tree_add_dh_lm, means, interval = "confidence")</pre>
```

```
## fit lwr upr
## 1 32.6 31.7 33.4
```

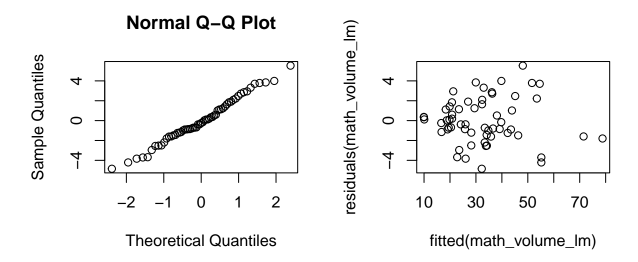
Therefore we expect the volume for such a tree to be 32.6.

d) Assuming that a tree is roughly cylindrical, we expect that *volume* would be proportional to the *height* multiplied by the square of *diameter*. We perform this transformation and add it as a new column in the data frame. We could apply the true transformation,  $V = h \times \pi (d/2)^2$ , but this would just add unnecessary constants which would already be captured in the regression coefficients. We also will not include *type* because it was not significant.

```
tree_df$math_volume <- tree_df$height * tree_df$diameter^2</pre>
math_volume_lm <- lm(volume~math_volume, data=tree_df)</pre>
anova(math_volume_lm)
## Analysis of Variance Table
##
## Response: volume
               Df Sum Sq Mean Sq F value Pr(>F)
                           11477
                                     2201 <2e-16 ***
## math_volume
               1
                   11477
## Residuals
               57
                     297
                               5
## ---
## Signif. codes: 0 '*** 0.001 '** 0.01 '* 0.05 '.' 0.1 ' 1
summary(math volume lm)
##
## Call:
## lm(formula = volume ~ math_volume, data = tree_df)
## Residuals:
##
              1Q Median
      Min
                            3Q
                                  Max
## -4.846 -1.343 -0.245 1.533 5.532
##
## Coefficients:
##
                Estimate Std. Error t value Pr(>|t|)
## (Intercept) -3.79e-01
                           7.63e-01
                                        -0.5
                                                 0.62
## math_volume 2.14e-03
                           4.57e-05
                                        46.9
                                               <2e-16 ***
## ---
                   0 '*** 0.001 '** 0.01 '* 0.05 '. ' 0.1 ' 1
## Signif. codes:
## Residual standard error: 2.28 on 57 degrees of freedom
## Multiple R-squared: 0.975, Adjusted R-squared: 0.974
## F-statistic: 2.2e+03 on 1 and 57 DF, p-value: <2e-16
```

We see that this transformation does indeed produce an explanatory value with a significant effect. We also see that the R-squared (0.975) and adjusted R-squared (0.974) values are higher than that

of the model chosen in (c) (tree\_add\_dh\_lm), indicating that it better explains the data. Finally, we check the assumptions of this model.

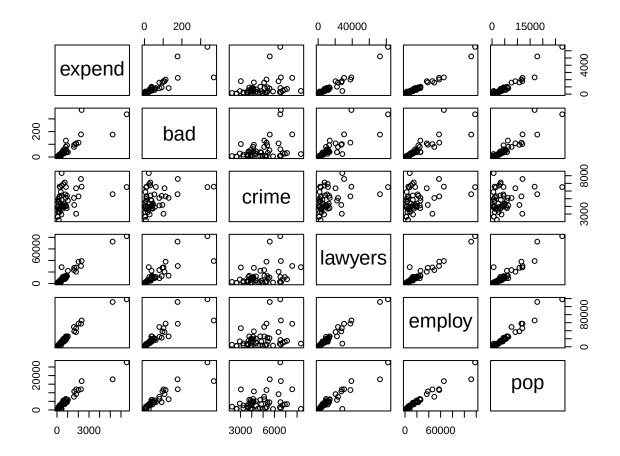


These plots are acceptable, meaning we can accept the model assumptions.

# Exercise 2

**a**)

To investigate the interactions between all the variables of interest, we can plot the pairwise scatter plots for all their combinations:



We see that *expend*, our response variable, appears to have a positive correlation with all the explanatory variables except for *crime*. There appear to be several outliers at the high end of the data which could skew the model. We can also see that collinearity exists between the explanatory variables *bad*, *lawyers*, *employ* and *pop*. This is a problem since the redundant information will make the regression coefficients difficult to estimate.

We can use Cook's distance to find the influence points (a distance greater than 1 indicates an outlier)

```
crime_lm <- lm(expend~bad+crime+lawyers+employ+pop, data=crime_df)
cooks.distance(crime_lm)[cooks.distance(crime_lm) > 1]
```

```
## 5 8 35 44
## 4.91 3.51 1.09 2.70
```

We can see that indices of 5, 8, 35 and 44 are outliers, which we can remove:

```
crime_df_upd <- crime_df[-c(5,8,35,44),]</pre>
```

To further investigate collinearity, we can examine the correlations between all the explanatory variables, which confirms strong correlations between bad, lawyers, employ and pop.

```
round(cor(crime_df[, c(exp_vars)]), 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
## pop
           0.92 0.28
                         0.93
                                0.97 1.00
```

We can also use the VIF to see which variables are collinear (VIF > 5 is cause for concern).

```
vif(lm(expend~bad+crime+lawyers+employ+pop, data=crime_df))
```

```
## bad crime lawyers employ pop
## 8.36 1.49 16.97 33.59 32.94
```

This further confirms that collinearity exists for the variables bad, lawyers, employ and pop.

From this point on, we will proceed without the influence points.

b)

The step-up process was carried out. The variables added in order were *employ*, *crime* and *pop*, after which no further added variables had significant p-values. Hence the final model is as follows:

```
step_up_lm <- lm(expend~employ+crime+pop, data=crime_df_upd)
summary(step_up_lm)</pre>
```

```
##
## Call:
## lm(formula = expend ~ employ + crime + pop, data = crime_df_upd)
##
## Residuals:
##
      Min
                1Q Median
                                3Q
                                       Max
## -179.99 -49.64
                      0.48
                             51.19
                                   266.63
##
## Coefficients:
                Estimate Std. Error t value Pr(>|t|)
## (Intercept) -2.47e+02
                           5.47e+01
                                      -4.52 4.8e-05 ***
## employ
                2.09e-02
                           3.95e-03
                                       5.30 3.7e-06 ***
## crime
                5.43e-02
                           1.13e-02
                                       4.82 1.8e-05 ***
                           1.79e-02
                                       4.00 0.00025 ***
## pop
               7.14e-02
## ---
                  0 '*** 0.001 '** 0.01 '* 0.05 '. ' 0.1 ' ' 1
## Signif. codes:
## Residual standard error: 91.4 on 43 degrees of freedom
## Multiple R-squared: 0.974, Adjusted R-squared: 0.973
## F-statistic: 547 on 3 and 43 DF, p-value: <2e-16
```

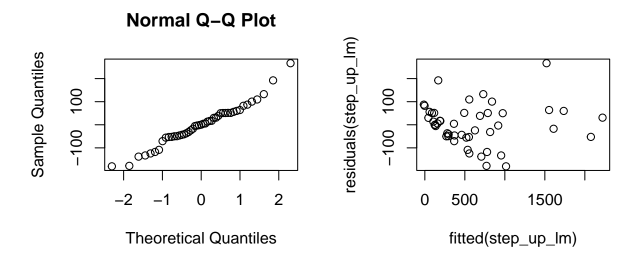
Final model: expend = -247 + 0.0209\*employ + 0.0543\*crime + 0.0714\*pop  $\pm$  error, with  $R^2 = 0.974$ . Using VIF we can

```
vif(step_up_lm)
```

```
## employ crime pop
## 18.08 1.14 17.51
```

We see that the step-up method naturally removes collinearity and produced a better model than was arrived upon using VIF in (a), which had an R-squared value of 0.957.

Finally, we check the model assumptions, which can be accepted based on the following plots:



**c**)

Using the step-up model found in (b), the 95% prediction interval for expend is given by:

```
new_data <- data.frame(bad=50, crime=5000, lawyers=5000, employ=5000, pop=5000)
predict(step_up_lm, new_data, interval="prediction", level=0.95)</pre>
```

```
## fit lwr upr
## 1 486 258 713
```

We cannot improve this interval since we have already removed the influence points from the data.

d)

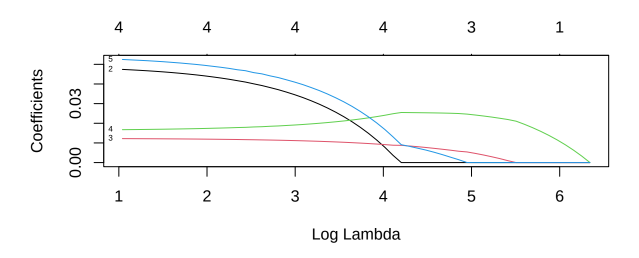
We can apply the LASSO method as follows:

```
x <- as.matrix(crime_df_upd[, exp_vars])
y <- as.matrix(crime_df_upd[, c(response)])</pre>
```

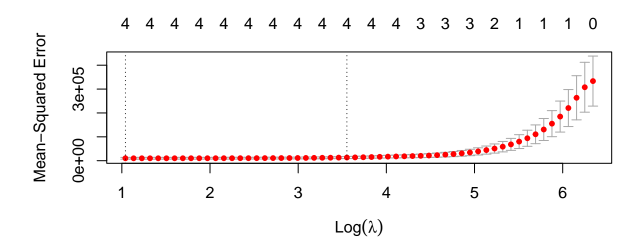
```
# train-test splitting
train <- (sample(1:nrow(x), 0.67*nrow(x))) # train by using 2/3 of the data
x.train <- x[train,]; y.train <- y[train]
x.test <- x[-train,]; y.test <- y[-train]

# fitting the model
lasso.mod <- glmnet(x.train, y.train, alpha=1)
cv.lasso <- cv.glmnet(x.train,y.train,alpha=1,type.measure='mse')</pre>
```

plot(lasso.mod, label=T, xvar="lambda") # have a look at the lasso path



plot(cv.lasso) # the best lambda by cross-validation



```
(lambda.1se <- cv.lasso$lambda.1se)
## [1] 34.9
\# https://glmnet.stanford.edu/articles/glmnet.html\#assessing-models-on-test-data-1
assess.glmnet(lasso.mod, newx = x.test, newy = y.test, s=cv.lasso$lambda.1se)
## $mse
##
      s1
## 17369
## attr(,"measure")
## [1] "Mean-Squared Error"
## $mae
##
## 94.1
## attr(,"measure")
## [1] "Mean Absolute Error"
Looking at lambda 1se
coef(lasso.mod, s=cv.lasso$lambda.1se) # beta's for lambda.1se
## 6 x 1 sparse Matrix of class "dgCMatrix"
##
## (Intercept) -44.2794
## bad
## crime
                 0.0233
## lawyers
                 0.0103
## employ
                 0.0212
                 0.0309
## pop
y.pred <- predict(lasso.mod, s=lambda.1se, newx=x.test) # predict for test
mse.lasso <- mean((y.test - y.pred)^2); mse.lasso # mse for the predicted test rows
## [1] 17369
To compare this to the step-up model in (b), we can find the MSE for this model.
new_data <- data.frame(x.test)</pre>
y.pred <- predict(step_up_lm, new_data, interval="confidence", level=0.95)</pre>
mse.step_up <- mean((y.test - y.pred)^2); mse.step_up # mse for the predicted test rows
## [1] 13030
```

We see that the step-up model outperforms the LASSO model, producing a smaller MSE. This could be because LASSO is better suited to situations with many more explanatory variables.

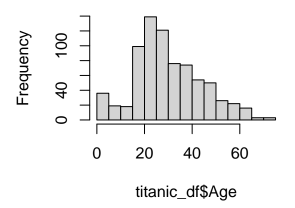
## Exercise 3

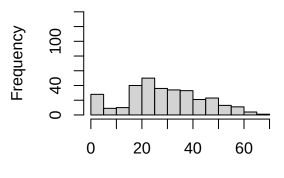
a)

```
titanic_df$PClass <- as.factor(titanic_df$PClass)</pre>
titanic_df$Sex <- as.factor(titanic_df$Sex)</pre>
summary(titanic_df)
##
                        PClass
                                                                  Survived
        Name
                                       Age
                                                     Sex
## Length:1313
                                  Min.
                        1st:322
                                         : 0
                                                 female:462
                                                              Min.
                                                                      :0.000
## Class :character
                        2nd:280
                                  1st Qu.:21
                                                 male :851
                                                              1st Qu.:0.000
## Mode :character
                        3rd:711
                                  Median:28
                                                              Median :0.000
##
                                  Mean
                                         :30
                                                              Mean
                                                                      :0.343
                                  3rd Qu.:39
                                                               3rd Qu.:1.000
##
                                         :71
##
                                  Max.
                                                              Max.
                                                                      :1.000
                                  NA's
##
                                          :557
tot_comb <- xtabs(~PClass+Sex, data=titanic_df)</pre>
tot_comb
         Sex
##
## PClass female male
      1st
             143 179
##
      2nd
             107 173
##
      3rd
             212 499
tot_comb.surv <- xtabs(Survived~PClass+Sex, data=titanic_df)</pre>
round(tot_comb.surv/tot_comb, 2)
##
         Sex
## PClass female male
            0.94 0.33
      1st
##
      2nd
            0.88 0.14
##
      3rd
          0.38 0.12
par(mfrow=c(1, 2))
hist(titanic_df$Age)
```

hist(titanic\_df\$Age[titanic\_df\$Survived == 1], ylim =c(0, 140))

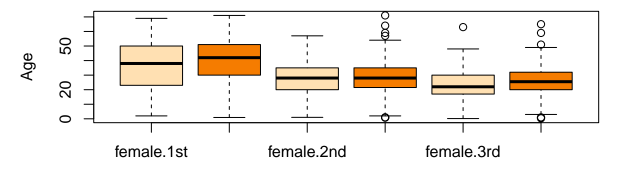
# Histogram of titanic\_df\$Age 'am of titanic\_df\$Age[titanic\_df\$Sul





titanic\_df\$Age[titanic\_df\$Survived == 1

boxplot(Age ~ Sex + PClass, data=titanic\_df, col = c("#FFE0B2", "#F57C00"))



Sex: PClass

Removing rows with missing ages:

```
titanic_df_upd <- na.omit(titanic_df)</pre>
```

Fitting the logistic regression model.

```
titanic_df_upd$PClass <- as.factor(titanic_df_upd$PClass)
titanic_df_upd$Sex <- as.factor(titanic_df_upd$Sex)
base_lm <- glm(Survived ~ Age+PClass+Sex, data = titanic_df_upd, family = binomial)</pre>
```

```
drop1(base_lm, test = "Chisq")
## Single term deletions
##
## Model:
## Survived ~ Age + PClass + Sex
         Df Deviance AIC
                          LRT Pr(>Chi)
##
## <none>
                 695 705
                724 732 28.5 9.6e-08 ***
## Age
         1
## PClass 2
                 796 802 100.4 < 2e-16 ***
                 910 918 214.8 < 2e-16 ***
## Sex
          1
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
summary(base lm)
##
## Call:
## glm(formula = Survived ~ Age + PClass + Sex, family = binomial,
      data = titanic_df_upd)
##
## 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|)
## (Intercept) 3.75966
                          0.39757
                                   9.46 < 2e-16 ***
             -0.03918
## Age
                          0.00762 -5.14 2.7e-07 ***
## PClass2nd -1.29196 0.26008 -4.97 6.8e-07 ***
## PClass3rd -2.52142
                          0.27666
                                   -9.11 < 2e-16 ***
## Sexmale
             -2.63136
                          0.20151 -13.06 < 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
## AIC: 705.1
##
## Number of Fisher Scoring iterations: 5
exp(coef(base_lm))
                           PClass2nd
                                       PClass3rd
                                                     Sexmale
## (Intercept)
                      Age
##
      42.9339
                   0.9616
                               0.2747
                                          0.0803
                                                      0.0720
```

TODO: add discussion of odds from the paper b) anova(glm(Survived ~ Age\*PClass, data = titanic\_df\_upd, family = binomial), test="Chisq") ## Analysis of Deviance Table ## ## Model: binomial, link: logit ## ## Response: Survived ## ## Terms added sequentially (first to last) ## ## Df Deviance Resid. Df Resid. Dev Pr(>Chi) ## ## NULL 1026 755 ## Age 1 2.8 754 1023 0.091 . 2 ## PClass 112.8 752 910 <2e-16 \*\*\* ## Age:PClass 1.2 750 909 0.558 2 ## ---0 '\*\*\* 0.001 '\*\* 0.01 '\* 0.05 '. ' 0.1 ' ' 1 ## Signif. codes: anova(glm(Survived ~ Age\*Sex, data = titanic\_df\_upd, family = binomial), test="Chisq") ## Analysis of Deviance Table ## ## Model: binomial, link: logit ## Response: Survived ## ## Terms added sequentially (first to last) ## ## Df Deviance Resid. Df Resid. Dev Pr(>Chi) ## NULL 755 1026 754 1023 ## Age 1 2.8 0.091 . ## Sex 1 227.1 753 796 < 2e-16 \*\*\* ## Age:Sex 1 25.0 752 771 5.6e-07 \*\*\*

Therefore we decided to keep following model as Age:Sex interaction is significant. PClass was significant by itself, so we include it in the final model.

0 '\*\*\* 0.001 '\*\* 0.01 '\* 0.05 '. ' 0.1 ' ' 1

««< INVESTIGATE VARIABLES USING anova(smaller model, bigger model) »»»

## Signif. codes:

```
final_lm <- glm(Survived ~ PClass+Age*Sex, data = titanic_df_upd, family = binomial)
summary(final_lm, test="Chisq")
##
## Call:
## glm(formula = Survived ~ PClass + Age * Sex, family = binomial,
       data = titanic_df_upd)
##
##
## Deviance Residuals:
##
      Min
               10 Median
                               3Q
                                       Max
## -2.435
          -0.656
                  -0.353
                            0.696
                                     2.728
##
## Coefficients:
##
               Estimate Std. Error z value Pr(>|z|)
## (Intercept) 2.75656
                           0.43764
                                       6.30 3.0e-10 ***
## PClass2nd
                                     -5.37 7.8e-08 ***
               -1.54337
                           0.28736
## PClass3rd
                           0.29142
                                      -9.11
                                             < 2e-16 ***
               -2.65398
## Age
                0.00244
                           0.01141
                                      0.21
                                                0.83
## Sexmale
               -0.50819
                           0.44251
                                      -1.15
                                                0.25
                                     -5.04 4.7e-07 ***
## Age:Sexmale -0.07559
                           0.01501
## ---
## 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: 667.08 on 750 degrees of freedom
## AIC: 679.1
##
## Number of Fisher Scoring iterations: 5
««< THIS MODEL MAY CHANGE »»>
newdata <- data.frame(Age=c(55, 55, 55, 55, 55, 55), PClass=c("1st", "1st", "2nd", "2nd", "3rd
predict(final_lm, newdata, type="response")
##
        1
                      3
                                     5
                                            6
## 0.9474 0.1450 0.7937 0.0350 0.5590 0.0118
For "female" all the probs > 0.5 and for the "male" probs are < 0.5.
c)
```

We can predict the survival status of the individuals by looking at the fitted values produced by the model, which gives the probability of survival. We can then set a threshold of 0.5. We can predict that those with a probability lower than this did not survive, and those with a probability higher than this did survive. To measure the quality of this prediction method, we can use tools such as a confusion matrix and log likelihood as quality measures.

d)

#### Survived vs SEX

For 2x2 tables we can obtain exact p-value using the Fisher test.

```
fisher.test(x=titanic_df_upd$Survived, y=titanic_df_upd$Sex)
```

```
##
## Fisher's Exact Test for Count Data
##
## data: titanic_df_upd$Survived and titanic_df_upd$Sex
## p-value <2e-16
## alternative hypothesis: true odds ratio is not equal to 1
## 95 percent confidence interval:
## 0.0586 0.1215
## sample estimates:
## odds ratio
## 0.0848</pre>
```

Therefore Sex and Survived are not independent.

```
chisq.test(x=titanic_df_upd$Survived, y=titanic_df_upd$PClass)
```

```
##
## Pearson's Chi-squared test
##
## data: titanic_df_upd$Survived and titanic_df_upd$PClass
## X-squared = 76, df = 2, p-value <2e-16</pre>
```

Therefore *PClass* and *Survived* are not independent.

**e**)

Logistic

A - Also tells you how and how much the response depends on the variable D - https://careerfoundry.com/en/blog/data-analytics/what-is-logistic-regression/

VS

Contingency

A - since only testing for effect, maybe more reliable outcome? D - Only tells you whether dependence exists... not the nature of it

 $_{
m VS}$ 

Fisher

A - exact p-value D - only works for 2x2

contingency table tells us only about the presence of effect and doesn't provide some quantitative characteristics

#### Exercise 4

```
coups_df$pollib <- as.factor(coups_df$pollib)</pre>
a)
poison_glm <- glm(miltcoup ~ oligarchy + parties + pctvote + popn + size + numelec + numregim +
drop1(poison_glm, test= "Chisq")
## Single term deletions
##
## Model:
## miltcoup ~ oligarchy + parties + pctvote + popn + size + numelec +
##
       numregim + pollib
             Df Deviance AIC LRT Pr(>Chi)
##
                    28.2 113
## <none>
## oligarchy 1
                    32.4 115 4.10 0.0428 *
## parties
              1
                    35.3 118 7.06 0.0079 **
## pctvote 1 30.6 113 2.32 0.1275
## popn 1 30.6 113 2.35 0.1252
## size 1 29.2 112 0.99 0.3202
## numelec 1 28.4 111 0.18 0.6705
## numregim 1 29.1 112 0.81 0.3681
              2 35.6 116 7.33
## pollib
                                     0.0256 *
## ---
## Signif. codes: 0 '*** 0.001 '** 0.01 '* 0.05 '.' 0.1 ' 1
summary(poison_glm)
##
## Call:
## glm(formula = miltcoup ~ oligarchy + parties + pctvote + popn +
       size + numelec + numregim + pollib, family = poisson, data = coups_df)
##
##
## Deviance Residuals:
##
      Min
               10 Median
                                3Q
                                       Max
## -1.508 -0.953 -0.310 0.486
                                     1.646
##
## Coefficients:
                Estimate Std. Error z value Pr(>|z|)
##
```

```
-0.23
## (Intercept) -0.233427
                            0.997611
                                               0.8150
## oligarchy
                0.072566
                            0.035346
                                        2.05
                                               0.0401 *
## parties
                0.031221
                                        2.80
                                               0.0052 **
                            0.011166
## pctvote
                0.015441
                            0.010103
                                        1.53
                                               0.1264
## popn
                0.010959
                            0.007149
                                        1.53
                                               0.1253
## size
               -0.000265
                            0.000269
                                       -0.99
                                               0.3244
## numelec
               -0.029619
                            0.069625
                                       -0.43
                                               0.6705
## numregim
                0.210943
                           0.233933
                                        0.90
                                               0.3672
## pollib1
                                       -1.68
               -1.103244
                            0.655811
                                               0.0925 .
## pollib2
               -1.690306
                           0.676650
                                       -2.50
                                               0.0125 *
## ---
                   0 '*** 0.001 '** 0.01 '* 0.05 '.' 0.1 ' ' 1
## Signif. codes:
## (Dispersion parameter for poisson family taken to be 1)
##
##
       Null deviance: 65.945
                                      degrees of freedom
                              on 35
## Residual deviance: 28.249
                              on 26
                                      degrees of freedom
## AIC: 113.1
##
## Number of Fisher Scoring iterations: 5
```

Through Poisson regression, and its summary, we can find that the variables that are significant in predicting number of successful military coups are: oliqarchy, pollib, and parties.

## b)

The step down method was applied, removing the variables: numelec, size, popn, and pctvote, respectively. After which, all remaining variables were significant, resulting in the model:

```
final_plm <- glm(miltcoup ~ oligarchy + parties + pollib, data = coups_df, family = poisson)
summary(final_plm, test="Chisq")</pre>
```

```
##
## glm(formula = miltcoup ~ oligarchy + parties + pollib, family = poisson,
       data = coups_df)
##
##
## Deviance Residuals:
##
      Min
               1Q Median
                                3Q
                                       Max
## -1.361 -1.041 -0.315
                             0.615
                                     1.754
##
## Coefficients:
##
               Estimate Std. Error z value Pr(>|z|)
## (Intercept)
                 0.2080
                             0.4457
                                       0.47
                                               0.641
## oligarchy
                 0.0915
                             0.0226
                                       4.05
                                               5e-05 ***
## parties
                                       2.46
                 0.0224
                             0.0091
                                               0.014 *
## pollib1
                -0.4954
                             0.4757
                                      -1.04
                                               0.298
## pollib2
                -1.1121
                             0.4595
                                      -2.42
                                               0.016 *
```

```
## Signif. codes: 0 '*** 0.001 '** 0.01 '* 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: 32.822 on 31 degrees of freedom
## AIC: 107.6
## Number of Fisher Scoring iterations: 5
In comparison with a) all the same variables are significant.
c)
coups_df$pollib <- as.factor(coups_df$pollib)</pre>
mean(coups_df$oligarchy); mean(coups_df$parties)
## [1] 5.22
## [1] 17.1
newdata <- data.frame(pollib=c("0", "1", "2"), oligarchy=c(5.22, 5.22, 5.22), parties=c(17.1,
predict(final_plm, newdata, type="response")
##
             2
       1
## 2.909 1.772 0.957
```

## ---

Our model predicts that there will be roughly 3 successful coups for pollib=0, roughly 2 successful coups for pollib=1 and 1 successful coup for pollib=2.