# Homework 3: Binary and Count Regressions

## Solution Set

This homework set will cover problems concerning binary and count regression models. Point totals for specific problems are given, 10 points will be reserved for correct submission of the homework assignment.

**Problem 1** [10 points]: Manually write your own Fisher scoring algorithm which maximizes the logistic regression log likelihood for the CCSO example in the notes. Report  $\hat{\beta}$  and reproduce the summary table (up to convergence tolerance differences) without using the glm or summary commands. You can ignore deviance residuals.

#### Solution 1:

The log-likelihood is

$$l(\beta) = \sum_{i=1}^{n} y_i x_i^T \beta - \log(1 + exp(x_i^T \beta))$$
$$l'(\beta) = \sum_{i=1}^{n} \left( y_i x_i - \frac{x_i}{1 + exp(x_i^T \beta)} exp(x_i^T \beta) \right)$$
$$\implies l'(\beta) = X^T (Y - \pi)$$

where  $\pi_i = \frac{exp(x_i^T \beta)}{1 + exp(x_i^T \beta)}$ 

$$l''(\beta) = -\sum_{i=1}^{n} \left( \frac{x_i^2}{(1 + exp(x_i^T \beta))^2} exp(x_i^T \beta) \right)$$
$$\implies l''(\beta) = -X^T W X$$

where  $W = diag(\pi_i(1 - \pi_i))$ 

Thus the Fisher scoring algorithm:

$$\beta^{(t+1)} = \beta^{(t)} + (X^T W^{(t)} X)^{-1} X^T (Y - \pi)$$

# Reading in the data

library(data.table)

library(tidyverse)

```
## -- Attaching packages -----
                                          ----- tidyverse 1.3.2 --
## v ggplot2 3.4.0 v purrr 0.3.5
                       v dplyr 1.0.10
## v tibble 3.1.8
## v tidyr 1.2.1
                      v stringr 1.4.1
## v readr
           2.1.3
                       v forcats 0.5.2
## -- Conflicts ----- tidyverse conflicts() --
## x dplyr::between() masks data.table::between()
## x dplyr::filter()
                       masks stats::filter()
## x dplyr::first()
                       masks data.table::first()
## x dplyr::lag() masks stats::lag()
## x dplyr::last() masks data.table::last()
## x dplyr::lag()
                       masks stats::lag()
## x purrr::transpose() masks data.table::transpose()
library(MASS)
##
## Attaching package: 'MASS'
## The following object is masked from 'package:dplyr':
##
##
       select
CCSO = fread("https://uofi.box.com/shared/static/9elozjsg99bgcb7gb546wlfr3r2gc9b7.csv")
# Pre-Processing the data
CCSO_small <- CCSO %>% rename(Days = "Days in Jail", Age = "Age at Arrest",
Date = "BOOKING DATE", Sex = "SEX", Race = "RACE",
Crime = "CRIME CODE") %>%
mutate(atleastone = ifelse(Days > 0,1,0)) %>%
filter(Crime == "OTHER TRAFFIC OFFENSES") %>%
filter(Race %in% c("Asian/Pacific Islander", "Black", "White", "Hispanic")) %>%
filter(Sex %in% c("Female","Male")) %>%
dplyr::select(atleastone, Age, Sex, Date, Race) %>%
mutate(Race = fct_drop(Race), Sex = fct_drop(Sex))
CCSO_small <- CCSO_small[complete.cases(CCSO_small), ]</pre>
head(CCSO_small)
     atleastone Age
                                       Race
                       Sex
                               Date
         0 22
                      Male 1/1/2011
## 1:
                                      White
             0 26 Male 1/1/2011
## 2:
                                      White
## 3:
             0 32 Female 1/1/2011
                                    White
## 4:
             0 22 Male 1/2/2011
                                      White
              0 35
## 5:
                      Male 1/2/2011 Hispanic
## 6:
              0 35
                     Male 1/2/2011 Hispanic
#Creating the model matrix
X = model.matrix(atleastone ~ -1 + Race + Sex + Age, data = CCSO_small)
n = nrow(X)
p = ncol(X)
Y = CCSO small$atleastone
# Initializing the beta
beta = matrix(rep(0,6))
#Running the Fisher scoring iterations
for(t in 1:10)
```

```
pi = exp(X%*\%beta)/(1+exp(X%*\%beta))
  W = diag(c(pi*(1-pi)))
  beta = beta + solve(t(X) %*% W %*% X)%*%t(X)%*%(Y - pi)
}
# The final values
pi_CCSO = exp(X%*\%beta)/(1+exp(X%*\%beta))
W_{CCSO} = diag(c(pi_{CCSO}*(1-pi_{CCSO})))
var_matrix_CCSO = solve(t(X) %*% W_CCSO %*% X)
sd_beta = sqrt(diag(var_matrix_CCSO))
z_Val = beta/sd_beta
pvalue = 2*(1 - pnorm(abs(z_Val)))
#Deviance results
deviance\_res = -2*(t(Y)%*%X%*\%beta - sum(log(1+exp(X%*\%beta))))
beta_0 = 0
deviance_null = -2*(beta_0*sum(Y) - n*log(1+exp(beta_0)))
AIC = deviance_res + 2*p
The summary table:
tab = data.frame("Estimate" = beta, "Std.Error" = sd_beta, "z value" = z_Val, "Pvalue" = pvalue)
list("Coefficients" = tab, "Null deviation" = deviance_null, "Residual Deviance" = deviance_res, "Null
## $Coefficients
##
                                  Estimate
                                             Std.Error
                                                          z.value
                                                                         Pvalue
## RaceAsian/Pacific Islander -4.38986549 0.523613419 -8.383791 0.000000e+00
                              -1.87654964 0.144600944 -12.977437 0.000000e+00
## RaceBlack
## RaceHispanic
                              -2.80454861 0.173349092 -16.178617 0.000000e+00
## RaceWhite
                              -3.04322627 0.147160127 -20.679693 0.000000e+00
## SexMale
                               0.73983403 0.105379772 7.020646 2.208456e-12
                               0.00770504 0.003186262 2.418207 1.559721e-02
## Age
##
## $`Null deviation`
## [1] 8201.317
##
## $`Residual Deviance`
##
            [,1]
## [1,] 4668.728
##
## $`Null df`
## [1] 5916
## $`Residual df`
## [1] 5910
##
## $AIC
##
            [,1]
```

**Problem 2** [10 points]: Manually write your own Fisher scoring algorithm which maximizes the Poisson regression log likelihood for the Galapagos example in the notes. Report  $\hat{\beta}$  and reproduce the summary

## [1,] 4680.728

table (up to convergence tolerance differences) without using the glm or summary commands. You can ignore deviance residuals.

#### Solution 2:

The log-likelihood is

$$l(\beta) = \sum_{i=1}^{n} (y_i x_i^T \beta - exp(x_i^T \beta))$$
$$l'(\beta) = \sum_{i=1}^{n} (y_i x_i - x_i exp(x_i^T \beta))$$
$$\implies l'(\beta) = X^T (Y - \pi)$$

where  $\pi_i = exp(x_i^T \beta)$ 

$$l''(\beta) = -\sum_{i=1}^{n} x_i^2 exp(x_i^T \beta)$$
$$\implies l''(\beta) = -X^T W X$$

where  $W = diag(\pi_i)$ 

Thus the Fisher scoring algorithm:

$$\beta^{(t+1)} = \beta^{(t)} + (X^T W^{(t)} X)^{-1} X^T (Y - \pi)$$

```
library(faraway)
```

```
#Pre-processing the data
data(gala)
gala <- gala %>%
mutate(Size = as.factor(1 + ifelse(Area > 1,1,0) + ifelse(Area > 25,1,0)))
head(gala)
```

```
##
                Species Endemics Area Elevation Nearest Scruz Adjacent Size
## Baltra
                              23 25.09
                                             346
                                                     0.6
                                                           0.6
                     31
                              21 1.24
                                             109
                                                                 572.33
                                                                           2
## Bartolome
                                                     0.6 26.3
## Caldwell
                     3
                               3 0.21
                                             114
                                                     2.8 58.7
                                                                   0.78
                     25
                                              46
## Champion
                               9 0.10
                                                     1.9 47.4
                                                                   0.18
                                                                           1
## Coamano
                      2
                               1 0.05
                                              77
                                                     1.9
                                                           1.9
                                                                 903.82
                                                                           1
## Daphne.Major
                     18
                              11 0.34
                                             119
                                                     8.0
                                                           8.0
                                                                   1.84
```

```
# Creating the model matrix
```

```
X = model.matrix(Species ~ Elevation + Nearest + Scruz + Adjacent + Size, data = gala)
n = nrow(X)
p = ncol(X)
Y = gala$Species
```

### # Initialising the beta

beta = c(0,apply(X,2,mean)[-1]/apply(X,2,var)[-1])

```
#Running the fisher scoring iterations
for(t in 1:10)
 pi = exp(X%*\%beta)
 W = diag(c(pi))
 beta = beta + solve(t(X) %*\% W %*\% X)%*\%t(X)%*\%(Y - pi)
}
# The final values
pi = exp(X%*\%beta)
W = diag(c(pi))
var_matrix = solve(t(X) %*% W %*% X)
sd_beta = sqrt(diag(var_matrix))
z_Val = beta/sd_beta
pvalue = 2*(1 - pnorm(abs(z_Val)))
#Deviance results
deviance_res = -2*(t(Y)%*%X%*\%beta - sum(pi)+ sum(Y - Y*log(Y)))
beta_0 = log(mean(Y))
deviance_null = -2*(beta_0*sum(Y) - n*exp(beta_0) + sum(Y - Y*log(Y)))
AIC = -2*sum(dpois(Y,pi,log = TRUE)) + 2*(p)
The summary table:
tab = data.frame("Estimate" = beta, "Std.Error" = sd_beta, "z value" = z_Val, "Pvalue" = pvalue)
list("Coefficients" = tab, "Null deviation" = deviance_null, "Residual Deviance" = deviance_res, "Null
## $Coefficients
                    Estimate
                                Std.Error
                                           z.value
## (Intercept) 2.7897964692 8.107802e-02 34.408787 0.0000000000
## Elevation 0.0009360990 5.402069e-05 17.328527 0.0000000000
## Nearest
              0.0064693041 1.747557e-03 3.701912 0.0002139805
               -0.0062664946 6.268336e-04 -9.997063 0.0000000000
## Scruz
              -0.0002857805 2.960795e-05 -9.652152 0.0000000000
## Adjacent
## Size2
              1.1276155415 9.535272e-02 11.825730 0.0000000000
               2.0586771315 9.419392e-02 21.855732 0.0000000000
## Size3
## $`Null deviation`
## [1] 3510.729
##
## $`Residual Deviance`
##
            [,1]
## [1,] 594.1753
##
## $`Null df`
## [1] 29
##
## $`Residual df`
## [1] 23
##
## $AIC
## [1] 769.0063
```

**Problem 3** [5 points]: Derive the log-odds ratio of x + 1 to x when Y = 1, and observe that the log-odds ratio does not depend on x. Comment on this finding.

#### Solution 3:

We know that in the logistic regression model we have

$$logit(\pi(x)) = x\beta$$
 where  $\pi(x) = E(Y|X=x) = P(Y=1|X=x)$ 

Thus the log of their ratio

$$\log\left(\frac{odds(Y=1|X=x+1)}{odds(Y=1|X=x)}\right) = \log\left(\frac{P(Y=1|X=x+1)/P(Y=0|X=x+1)}{P(Y=1|X=x+1)/P(Y=0|X=x+1))}\right)$$

$$= \log\left(\frac{\pi(x+1)/(1-\pi(x+1))}{\pi(x)/(1-\pi(x))}\right)$$

$$= \log\left(\pi(x+1)/(1-\pi(x+1))\right) - \log\left(\pi(x)/(1-\pi(x))\right)$$

$$= \log it(\pi(x+1)) - \log it(\pi(x))$$

$$= (x+1)\beta - x\beta$$

$$= \beta$$

Thus it does not depend on X. This shows that a unit increase in X leads to an increase of beta in the log-odds ratio i.e a unit increase in x leads to the ratio of their odds increasing by  $e^{\beta}$ 

**Problem 4** [10 points]: Complete the following parts:

- (a) Explain important findings and model information from the summary table produced by a call to summary(m1) in the CCSO example in the logistic regression notes. Keep in mind that we restricted attention to "other traffic offenses" in the CCSO example, and that this data is observational.
- (b) Explain important findings and model information from the summary table produced by a call to summary(m1) in the Galapagos islands example in the count regression notes.

### Solution 4:

```
# Creating the logistic regression model for the CCSO model
m1 <- glm(atleastone ~ -1 + Race + Sex + Age, data = CCSO_small,
family = "binomial", x = "TRUE")
summary(m1)
##
## Call:
## glm(formula = atleastone ~ -1 + Race + Sex + Age, family = "binomial",
##
       data = CCSO small, x = "TRUE")
##
## Deviance Residuals:
##
       Min
                 1Q
                      Median
                                    3Q
                                            Max
```

```
## -0.9393 -0.5485 -0.4817 -0.3391
                                        2.6527
##
## Coefficients:
##
                               Estimate Std. Error z value Pr(>|z|)
## RaceAsian/Pacific Islander -4.389865
                                          0.523612
                                                    -8.384
                                                             < 2e-16 ***
                              -1.876550
## RaceBlack
                                          0.144601 - 12.977
                                                             < 2e-16 ***
## RaceHispanic
                              -2.804549
                                          0.173349 - 16.179
                                                             < 2e-16 ***
                                          0.147160 -20.680
## RaceWhite
                              -3.043226
                                                             < 2e-16 ***
## SexMale
                               0.739834
                                          0.105380
                                                      7.021 2.21e-12 ***
## Age
                               0.007705
                                          0.003186
                                                      2.418
                                                              0.0156 *
##
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
##
##
  (Dispersion parameter for binomial family taken to be 1)
##
##
       Null deviance: 8201.3 on 5916
                                       degrees of freedom
## Residual deviance: 4668.7
                              on 5910
                                       degrees of freedom
  AIC: 4680.7
##
## Number of Fisher Scoring iterations: 6
```

The estimate column gives the estimate for  $\beta$  for the logistic model

$$logit(E(Y|X)) = X\beta$$

A unit increase in the predictor variable  $X_j$  corresponds to an increase of  $\beta_j$  (estimated by  $\hat{\beta}_j$ ) in the log-odds ratio with everything else being held fixed. A simpler interpretation is that  $\hat{\beta}_j > 0$  can be interpreted as: An increase in  $X_j$  implies that P(Y = 1 | X = x) increases.

Thus

- Race is significant when testing at reasonable significance levels. We observe that Black individuals
  are estimated to have comparatively larger propensity of incarcerations lasting one day or longer for
  "other traffic offenses". We would need to look into other factors such as socio-economic status, repeat
  offenders, and multiple offenses before we could conclude that race is the driver of longer incarcerations.
- Sex being Male is estimated to increase the propensity of incarcerations lasting longer than one day for "other traffic offenses".
- Age increasing also is estimated to increase the propensity of incarcerations lasting longer than one day
  for "other traffic offenses".

The standard error column gives the standard error of the estimate of the  $\beta$  coefficients. The Z-value and P-value help in detecting the significance of the covariates. At a level of  $\alpha = 0.05$  we can see that all the covariates are significant.

The null deviance and residual deviance give information about the goodness of fit of the null model (with no covariates) and the submodel we consider respectively. To check if the sub-model is better than the saturated model we can do a  $\chi^2$  test because under  $H_0$  (The submodel is a better fit)

$$D(y; \hat{\mu}) \sim \chi_{n-p}^2,$$

where D is the deviance.

```
pchisq(m1$deviance, df = m1$df.residual, lower = FALSE)
```

## [1] 1

Since the p-value is 1 this shows that the submodel is indeed a good fit to the data. We can also check if the null model  $(M_0)$  is better than the submodel  $(M_1)$  we choose.

 $H_0: M_0 \text{ true} \quad H_a: M_1 \text{ true}$ , but not  $M_0$ 

Then

$$D(y; \hat{\mu}_0) - D(y; \hat{\mu}_1) \sim \chi^2_{p_0 - p_1}$$

where  $D(y; \hat{\mu}_0)$  is the null deviance and  $D(y; \hat{\mu}_1)$  is the residual deviance.

```
pchisq(m1$null.deviance - m1$deviance, df = m1$df.null - m1$df.residual,
lower = FALSE)
```

#### ## [1] 0

Since the pvalue is 0 this means that the submodel we choose is a better fit than the null model.

(b) Summarize the summary tables produced by a call to summary(m1) in the Galapagos islands example in the count regression notes.

```
m2 <- glm(Species ~ Elevation + Nearest + Scruz + Adjacent + Size,
family = "poisson", data = gala, x = TRUE)
summary(m2)</pre>
```

```
##
## Call:
  glm(formula = Species ~ Elevation + Nearest + Scruz + Adjacent +
##
       Size, family = "poisson", data = gala, x = TRUE)
##
  Deviance Residuals:
##
##
        Min
                   1Q
                         Median
                                       3Q
                                                Max
##
  -10.3723
              -3.5214
                        -0.9947
                                   1.7193
                                            10.6627
##
## Coefficients:
##
                 Estimate Std. Error z value Pr(>|z|)
## (Intercept) 2.790e+00 8.108e-02
                                     34.410
                                              < 2e-16 ***
## Elevation
                9.361e-04
                           5.402e-05
                                      17.329
                                              < 2e-16 ***
                           1.748e-03
## Nearest
                6.469e-03
                                       3.702 0.000214 ***
               -6.266e-03
                           6.268e-04
                                      -9.997
## Scruz
                                              < 2e-16 ***
               -2.858e-04
                           2.961e-05
                                      -9.652
                                              < 2e-16 ***
## Adjacent
## Size2
                1.128e+00
                           9.535e-02
                                      11.826
                                              < 2e-16 ***
## Size3
                2.059e+00 9.419e-02
                                      21.856
                                              < 2e-16 ***
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## (Dispersion parameter for poisson family taken to be 1)
##
##
       Null deviance: 3510.73 on 29
                                      degrees of freedom
## Residual deviance: 594.18 on 23 degrees of freedom
## AIC: 769.01
##
## Number of Fisher Scoring iterations: 5
```

The estimate column gives the estimate for  $\beta$  for the logistic model

$$log(E(Y|X)) = X\beta$$

. A unit increase in the predictor variable  $X_j$  corresponds to an increase of  $\beta_j$  (estimated by  $\hat{\beta}_j$ ) in the log of the mean response with everything else being held fixed. A simpler interpretation is that  $\hat{\beta}_j > 0$  can be interpreted as: An increase in  $X_j$  implies that the mean response increases.

#### Thus

- Elevation is estimated to increase the expected number of plant species found on each island
- Distance to the nearest island is estimated to increases the expected number of plant species found on each island
- Distance to Scruz is estimated to decreases the number of expected plant species found on each island
- Larger adjacent islands are also estimated to decrease the number of expected plant species found on each island
- Medium and large islands are estimated to have more expected plant species found on each island

The standard error column gives the standard error of the estimate of the  $\beta$  coefficients. The Z-value and P-value help in detecting the significance of the covariates. At a level of  $\alpha = 0.05$  we can see that all the covariates are significant.

The null deviance and residual deviance give information about the goodness of fit of the null model (w ith no covariates) and the submodel we consider respectively. To check if the sub-model is better than the saturated model we can do a  $\chi^2$  test because under  $H_0$  (The submodel is a better fit)

$$D(y; \hat{\mu}) \sim \chi_{n-p}^2,$$

where D is the deviance.

```
pchisq(m2$deviance, df = m2$df.residual, lower = FALSE)
```

### ## [1] 7.617409e-111

Since the p-value is very small this shows that the submodel not really a good fit to the data. We prefer the saturated model over it. We can also check if the null model  $(M_0)$  is better than the submodel  $(M_1)$  we choose.

 $H_0: M_0 \text{ true } H_a: M_1 \text{ true }, \text{ but not } M_0$ 

Then

$$D(y; \hat{\mu}_0) - D(y; \hat{\mu}_1) \sim \chi^2_{p_0 - p_1}$$

where  $D(y; \hat{\mu}_0)$  is the null deviance and  $D(y; \hat{\mu}_1)$  is the residual deviance.

```
pchisq(m2$null.deviance - m2$deviance, df = m2$df.null - m2$df.residual,
lower = FALSE)
```

## ## [1] 0

Since the pvalue is 0 this means that the submodel we choose is a better fit than the null model.

**Problem 5** [10 points]: Derive expressions and compute standard errors  $se(\hat{\mu})$  in the logistic regression CCSO example without using predict.glm. Then construct Wald based confidence intervals for the estimated mean

value parameters. Also construct confidence intervals  $(g(\hat{\beta} - z_{\alpha/2}se(\hat{\beta})), g(\hat{\beta} + z_{\alpha/2}se(\hat{\beta})))$ . Comment on the differences between these two confidence intervals for  $\hat{\mu}$ .

###solution 5

We know that

$$\sqrt{n}(\hat{\beta} - \beta) \xrightarrow{d} N(0, \Sigma^{-1})$$

where  $\Sigma = X^T W X$ ,  $W = diag(\mu_i(1 - \mu_i))$  and  $\mu = e^{X\beta}/(1 + e^{X\beta})$ 

Now 
$$\mu_i = g(\beta) = e^{M_i^T \beta} / (1 + e^{M_i^T \beta}) \implies g'(\beta) = M_i * \mu_i (1 - \mu_i)$$

Thus by the delta method

$$\sqrt{n}(\hat{\mu}_i - \mu_i) \xrightarrow{d} N(0, \hat{\mu}_i^2 (1 - \hat{\mu}_i)^2 M_i^T \Sigma^{-1} M_i)$$

Thus our estimation of the SE is:

```
#Creating the model matrix
X = model.matrix(atleastone ~ -1 + Race + Sex + Age, data = CCSO_small)
#Calculating the se
se_pihat = sqrt(apply(X,1,function(j) t(j)%*%var_matrix_CCSO%*%j))*pi_CCSO*(1-pi_CCSO)
se_pihat[1:20]
```

```
## [1] 0.006773575 0.006392931 0.005695285 0.006773575 0.013586275 0.013586275

## [7] 0.011956810 0.013052798 0.005695285 0.006269441 0.010557231 0.010557231

## [13] 0.009911238 0.009911238 0.012797440 0.014138891 0.006552717 0.013037138

## [19] 0.006472744 0.006370288
```

The Wald based confidence intervals for the estimated mean value parameters are

$$\hat{\mu}_i(x) \pm z_{1-\alpha/2}\sigma_i$$
 where  $\sigma_i = \hat{\mu}_i(1-\hat{\mu}_i)\sqrt{M_i^T \Sigma^{-1} M_i}$ 

```
#Creating the conf intervals
conf_lower = pi_CCSO - qnorm(0.975)*se_pihat
conf_upper = pi_CCSO + qnorm(0.975)*se_pihat
waldci = cbind(conf_upper, conf_lower)
waldci = waldci %>% as.data.frame() %>%
    mutate(length = conf_upper - conf_lower)
head(waldci)
```

```
## V1 V2 length
## 1 0.11912305 0.09257112 0.02655193
## 2 0.12132956 0.09626973 0.02505983
## 3 0.06866721 0.04634211 0.02232511
## 4 0.11912305 0.09257112 0.02655193
## 5 0.16908475 0.11582753 0.05325722
## 6 0.16908475 0.11582753 0.05325722
```

And, for the other confidence interval type  $g(\hat{\beta} + z_{\alpha/2} se(\hat{\beta}))$ :

```
z = qnorm(p = 1-alpha)
n = nrow(M)
pici_upper = 1/(1 + exp( - M %*% (betahat + z*diag(vcov(m1))) ))
pici_lower = 1/(1 + exp( - M %*% (betahat - z*diag(vcov(m1))) ))
pici = cbind(pici_upper, pici_lower)
pici = pici %>% as.data.frame() %>%
  mutate(length = pici_upper - pici_lower)
head(pici)
##
             ۷1
                        ٧2
                                 length
## 1 0.11212335 0.09988262 0.012240731
## 2 0.11523654 0.10268058 0.012555961
## 3 0.05988461 0.05521374 0.004670872
## 4 0.11212335 0.09988262 0.012240731
## 5 0.15268704 0.13280332 0.019883714
## 6 0.15268704 0.13280332 0.019883714
Then, the average length of the Wald and plug-in approaches are given as:
avg_length_wald = round(mean(waldci$length), digits=4)
avg_length_wald
## [1] 0.0392
avg_length_pi = round(mean(pici$length), digits=4)
avg_length_pi
```

## [1] 0.0157

Then, the average Wald CI is somewhat larger than the average plug-in CI.

**Problem 6** [10 points]: Construct a nonparametric bootstrap procedure that estimates the uncertainty associated with both estimates of the average treatment effect (ATE) of online learning in the logistic regression notes. Do the conclusions change when we factor in the uncertainty obtained from the nonparametric bootstrap procedure? Explain.

### Solution 6:

Here we are estimating the ATE for the scores earned by students in online learning as opposed to in-person learning.

```
#Reading in the data

dat = read.csv("../notes/3_binary_response/online.csv")

dat_small <- dat %>% dplyr::select(Online, ACTMath, ACTMajor, ACT, Gender,
International, F17, S18, S19, Fa19, FR, S0, JR)

ATE_alt = NULL

#Taking 1000 bootstrap samples
for(i in 1:1000)
{
    rand = sample(nrow(dat_small),replace = T)
    m <- glm(Online ~., data = dat_small[rand,], family = "binomial")
    trt <- dat_small[rand,]$Online</pre>
```

```
preds <- predict(m, type = "response")</pre>
  weights_alt_trt <- 1 / sum(trt / preds) * trt /preds</pre>
  weights_alt_notrt \leftarrow 1 / sum((1 - trt)/(1 - preds)) * (1-trt)/(1-preds)
  dat_new <- data.frame(dat[rand,], weights = weights_alt_trt - weights_alt_notrt)</pre>
  ATE_alt <- c(ATE_alt,sum(weights_alt_trt * dat_new$ObjExam) -
sum(weights_alt_notrt * dat_new$ObjExam))
}
mean(ATE_alt)
## [1] 0.5886974
var(ATE_alt)
## [1] 0.45145
quantile(ATE_alt,prob = c(0.025,0.975))
##
         2.5%
                    97.5%
## -0.6685877 1.8941839
Since the mean and variance are small and the confidence intreval contains 0 we can still conclude that there
is no difference between the two types of learning.
ATE DR = NULL
for(i in 1:1000)
  rand = sample(nrow(dat),replace = T)
  trt = dat_small[rand,]$Online
  m <- glm(Online ~., data = dat_small[rand,], family = "binomial")</pre>
  preds <- predict(m, type = "response")</pre>
  dat boot = dat[rand,]
  m_trt <- lm(ObjExam ~ ACTMath + ACTMajor + ACT + International + Gender +
FR + SO + JR + F17 + S18 + S19,
data = dat_boot[trt == 1, ])
  Y_trt <- predict(m_trt, newdata = dat_boot)</pre>
  m_notrt <- lm(ObjExam ~ ACTMath + ACTMajor + ACT + International + Gender +
FR + SO + JR + F17 + S18 + S19,
data = dat_boot[trt == 0, ])
  Y_notrt <- predict(m_notrt, newdata = dat_boot)</pre>
  ATE_DR <- c(ATE_DR,mean( (dat_boot$ObjExam * trt - (trt - preds) * Y_trt) / preds -
(dat_boot$0bjExam * (1 - trt) + (trt - preds)*Y_notrt) / (1 - preds)))
}
mean(ATE_DR)
## [1] 0.4155045
var(ATE_DR)
## [1] 0.3864671
quantile(ATE_DR,prob = c(0.025,0.975))
         2.5%
                    97.5%
```

Since the mean and variance are small even for the robust estimate and the confidence intreval contains 0 we

## -0.8000527

1.6897111

can still conclude that there is no difference between the two types of learning.

#### Solution 7

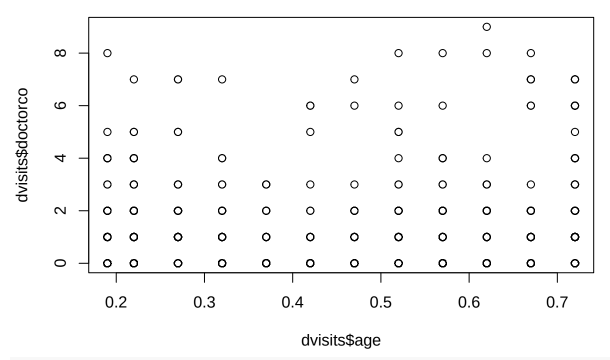
Problem 7 [15 points]: Use the dvisits data in the faraway package to answer the follow parts:

- (a) Make plots which show the relationship between the response variable, doctorco, and the potential predictors, age and illness.
- (b) Combine the predictors choond1 and choond2 into a single three-level factor. Make an appropriate plot showing the relationship between this factor and the response. Comment.
- (c) Build a Poisson regression model with doctorco as the response and sex, age, agesq, income, levyplus, freepoor, freerepa, illness, actdays, hscore and the three-level condition factor as possible predictor variables. Considering the deviance of this model, does this model fit the data?
- (d) Plot the residuals and the fitted values why are there lines of observations on the plot? Make a QQ plot of the residuals and comment.
- (e) Use a stepwise AIC-based model selection method. What sort of person would be predicted to visit the doctor the most under your selected model?
- (f) For the last person in the dataset, compute the predicted probability distribution for their visits to the doctor, i.e., give the probability they visit 0, 1, 2, etc. times.
- (g) Tabulate the frequencies of the number of doctor visits. Compute the expected frequencies of doctor visits under your most recent model. Compare the observed with the expected frequencies and comment on whether it is worth fitting a zero-inflated count model.
- (h) Fit a comparable (Gaussian) linear model and graphically compare the fits. Describe how they differ.

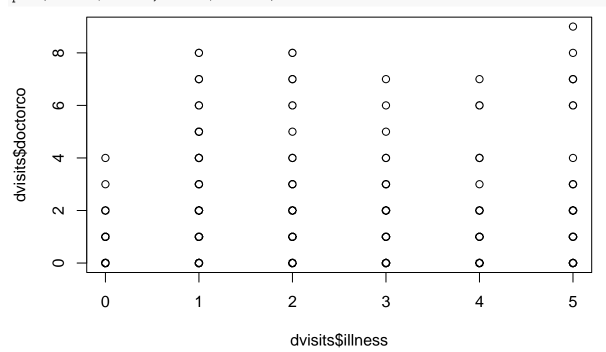
### Solution 7:

(a) Make plots which show the relationship between the response variable, doctorco, and the potential predictors, age and illness.

data(dvisits)
plot(dvisits\$age,dvisits\$doctorco)



plot(dvisits\$illness,dvisits\$doctorco)

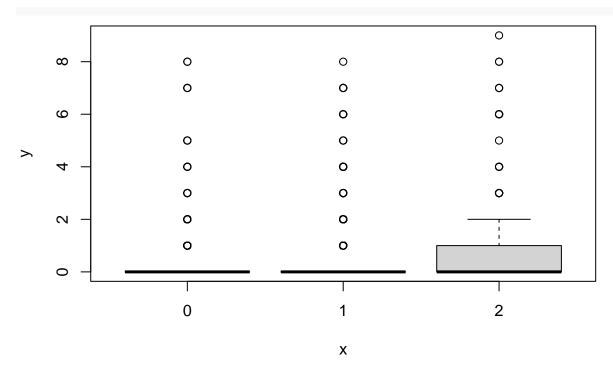


(b) Combine the predictors choond1 and choond2 into a single three-level factor. Make an appropriate plot showing the relationship between this factor and the response. Comment.

We create a new variable choond which takes 3 factor values

- 1 if the patient has a chronic condition(s) but is not limited in activity
- 2 if the patient has a chronic condition(s) but is limited in activity
- 0 Otherwise

chcond = as.factor(dvisits\$chcond1+2\*dvisits\$chcond2)
plot(chcond,dvisits\$doctorco)



From the plot we can see that patients which chronic conditions which limit activity have more visits to the doctor. This possible is a factor which influences the response variable.

(c) Build a Poisson regression model with doctorco as the response and sex, age, agesq, income, levyplus, freepoor, freerepa, illness, actdays, hscore and the three-level condition factor as possible predictor variables. Considering the deviance of this model, does this model fit the data?

```
dat = dvisits %>% dplyr::select(doctorco, sex, age, agesq, income,
levyplus, freepoor, freerepa, illness, actdays, hscore)
dat = cbind(dat, "chcond" = chcond)
head(dat)
##
     doctorco sex age agesq income levyplus freepoor freerepa illness actdays
## 1
             1
                 1 0.19 0.0361
                                  0.55
                                               1
                                                         0
                                                                  0
## 2
                 1 0.19 0.0361
                                  0.45
                                               1
                                                                           1
                                                                                    2
             1
                                                         0
## 3
             1
                 0 0.19 0.0361
                                  0.90
                                                                  0
                                                                           3
                                                                                    0
## 4
                 0 0.19 0.0361
                                  0.15
                                               0
                                                         0
                                                                  0
                                                                           1
                                                                                    0
             1
## 5
             1
                 0 0.19 0.0361
                                  0.45
                                               0
                                                         0
                                                                   0
                                                                           2
                                                                                    5
## 6
             1
                 1 0.19 0.0361
                                  0.35
                                               0
                                                         0
                                                                           5
                                                                                    1
     hscore chcond
##
## 1
          1
## 2
          1
                  0
## 3
                  0
          0
                  0
## 4
          0
          1
                  1
## 5
## 6
          9
#Creating the model
mod = glm(doctorco~.,data = dat,family = "poisson",x =TRUE)
summary(mod)
##
```

## Call:

```
## glm(formula = doctorco ~ ., family = "poisson", data = dat, x = TRUE)
##
## Deviance Residuals:
##
       Min
                 1Q
                      Median
                                    3Q
                                            Max
##
   -2.9170 -0.6862 -0.5743
                              -0.4839
                                         5.7005
##
## Coefficients:
##
                Estimate Std. Error z value Pr(>|z|)
## (Intercept) -2.223848
                           0.189816 -11.716
                                               <2e-16 ***
## sex
                0.156882
                           0.056137
                                       2.795
                                               0.0052 **
## age
                1.056299
                           1.000780
                                       1.055
                                               0.2912
                                      -0.787
## agesq
               -0.848704
                           1.077784
                                               0.4310
## income
               -0.205321
                           0.088379
                                     -2.323
                                               0.0202 *
                                               0.0855 .
## levyplus
                0.123185
                           0.071640
                                      1.720
                                      -2.447
                                               0.0144 *
## freepoor
               -0.440061
                           0.179811
## freerepa
                0.079798
                           0.092060
                                      0.867
                                               0.3860
                0.186948
                                     10.227
                                               <2e-16 ***
## illness
                           0.018281
## actdays
                0.126846
                           0.005034
                                      25.198
                                               <2e-16 ***
                                       2.979
                                               0.0029 **
## hscore
                0.030081
                           0.010099
## chcond1
                0.114085
                           0.066640
                                       1.712
                                               0.0869 .
## chcond2
                0.141158
                           0.083145
                                       1.698
                                               0.0896 .
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
##
##
  (Dispersion parameter for poisson family taken to be 1)
##
##
       Null deviance: 5634.8 on 5189
                                        degrees of freedom
## Residual deviance: 4379.5 on 5177
                                       degrees of freedom
## AIC: 6737.1
##
## Number of Fisher Scoring iterations: 6
#Testing for the goodness of fit of the model against the saturated moddel
pchisq(mod$deviance, df = mod$df.residual, lower = FALSE)
```

### ## [1] 1

Since the pvalue is 1 we can conclude that the model is indeed a better fit than the saturated model.

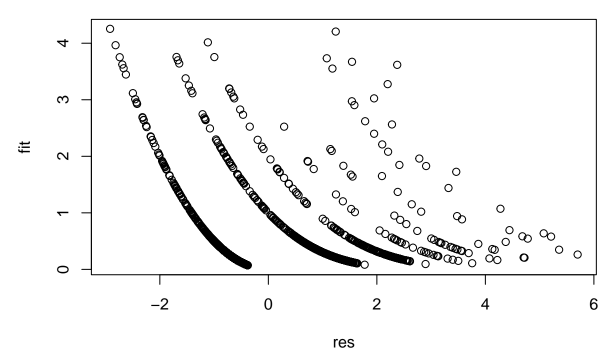
```
pchisq(mod$null.deviance - mod$deviance, df = mod$df.null - mod$df.residual,
lower = FALSE) %>% round(4)
```

### ## [1] 0

Since the pvalue is nearly 0 we can conclude that the model is also better than the null model. Thus it is a appropriate fit to the data.

(d) Plot the residuals and the fitted values — why are there lines of observations on the plot? Make a QQ plot of the residuals and comment.

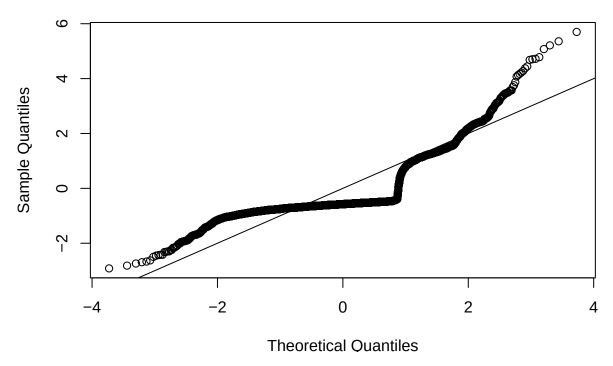
```
res = residuals(mod)
fit = fitted(mod)
plot(res,fit)
```



We observe lines of observations because most of the variables are factor variables with a small number of levels.

```
qqnorm(res)
abline(0, 1)
```

# Normal Q-Q Plot



The QQ-plot shows that the residuals do not follow a normal distribution very well indicating the normality of residuals assumption is not reasonable.

(e) Use a stepwise AIC-based model selection method. What sort of person would be predicted to visit the

doctor the most under your selected model?

```
#Selecting the model
library(MASS)
mod_Select = stepAIC(mod)
## Start: AIC=6737.08
## doctorco ~ sex + age + agesq + income + levyplus + freepoor +
##
      freerepa + illness + actdays + hscore + chcond
##
##
             Df Deviance
                           AIC
## - agesq
              1 4380.1 6735.7
## - freerepa 1 4380.3 6735.8
## - age
              1 4380.6 6736.2
## - chcond
              2 4383.2 6736.7
## <none>
                 4379.5 6737.1
## - levyplus 1 4382.5 6738.1
              1 4385.0 6740.5
## - income
## - freepoor 1 4386.2 6741.8
## - sex
             1 4387.4 6743.0
## - hscore 1 4388.1 6743.7
## - illness 1 4481.8 6837.4
## - actdays 1 4917.1 7272.7
##
## Step: AIC=6735.7
## doctorco ~ sex + age + income + levyplus + freepoor + freerepa +
##
      illness + actdays + hscore + chcond
##
             Df Deviance
                           AIC
## - freerepa 1 4381.0 6734.5
## <none>
                 4380.1 6735.7
## - chcond
              2 4384.2 6735.8
              1 4383.0 6736.5
## - age
## - levyplus 1 4383.3 6736.9
## - income
              1 4385.0 6738.6
## - freepoor 1 4386.8 6740.4
             1 4388.0 6741.5
## - sex
## - hscore
              1 4389.1 6742.7
## - illness
              1 4481.9 6835.4
## - actdays 1 4917.1 7270.7
##
## Step: AIC=6734.53
## doctorco ~ sex + age + income + levyplus + freepoor + illness +
##
      actdays + hscore + chcond
##
##
             Df Deviance
                           AIC
## <none>
                 4381.0 6734.5
## - levyplus 1
                4383.4 6735.0
## - chcond
                4385.5 6735.0
              2
## - income
                4386.7 6738.2
              1
## - age
              1 4387.1 6738.7
## - freepoor 1 4389.1 6740.6
## - sex
              1 4389.5 6741.0
              1 4390.2 6741.8
## - hscore
## - illness 1 4482.7 6834.2
```

```
## - actdays
               1
                   4917.6 7269.2
#Outputting the best models
mod Select$anova
## Stepwise Model Path
## Analysis of Deviance Table
##
## Initial Model:
## doctorco ~ sex + age + agesq + income + levyplus + freepoor +
##
       freerepa + illness + actdays + hscore + chcond
##
## Final Model:
## doctorco ~ sex + age + income + levyplus + freepoor + illness +
       actdays + hscore + chcond
##
##
##
##
           Step Df Deviance Resid. Df Resid. Dev
                                                        AIC
## 1
                                   5177
                                          4379.515 6737.083
        - agesq 1 0.6180113
                                   5178
                                          4380.133 6735.701
## 3 - freerepa 1 0.8279216
                                   5179
                                          4380.961 6734.529
# The beta coefficients of our best moddel
beta = mod_Select$coefficients
beta
## (Intercept)
                                                                    freepoor
                                             income
                                                       levyplus
                       sex
                                    age
## -2.08906349
                            0.35513074 -0.19980641
                                                     0.08368852 -0.46959634
                0.16199995
       illness
##
                                                        chcond2
                   actdays
                                hscore
                                            chcond1
   0.18610078
               0.12661065
                           0.03111559
                                        0.12110045
                                                     0.15889355
```

We can see that the number of doctor consultations increases when the patient is female, with increasing age, with low income, if covered by private health insurance, not covered by the government insurance for low income, high number of illness, high number of days of reduced activity, bad health score and with presence of chronic conditions. This indicates a poorer older woman with private insurance and higher number of illness is predicted to visit doctor more often.

(f) For the last person in the dataset, compute the predicted probability distribution for their visits to the doctor, i.e., give the probability they visit 0, 1, 2, etc. times.

```
options (scipen = 99)
X = mod_Select$x
Y = mod_Select$y
hat_lambda_last = exp(t(X[nrow(dat),])%*%beta)
data.frame("Value" = 0:9, "Prob" = dpois(0:9,hat_lambda_last) %>% round(6))
```

```
##
      Value
## 1
          0 0.858916
## 2
          1 0.130628
## 3
          2 0.009933
          3 0.000504
## 4
## 5
          4 0.000019
          5 0.000001
## 6
## 7
          6 0.000000
          7 0.000000
## 8
## 9
          8 0.000000
          9 0.000000
## 10
```

(g) Tabulate the frequencies of the number of doctor visits. Compute the expected frequencies of doctor visits under your most recent model. Compare the observed with the expected frequencies and comment on whether it is worth fitting a zero-inflated count model.

```
observed_freq <- with(dvisits, table(doctorco))
est <- matrix(nrow=dim(dvisits)[1], ncol=10)
for(i in 1:dim(dvisits)[1]){
est[i,] <- dpois(0:9, fitted.values(mod_Select)[i])
}
expected_freq <- colMeans(est)*dim(dvisits)[1]
cbind.data.frame(observed_freq, expected_freq)</pre>
```

```
##
      doctorco Freq expected_freq
## 1
              0 4141
                       4013.6020569
## 2
                 782
                        928.3492327
              1
## 3
              2
                 174
                        168.0095991
              3
                  30
                         45.4859546
## 5
              4
                  24
                         18.9118479
              5
## 6
                   9
                          8.8170066
## 7
              6
                  12
                          4.0118473
              7
## 8
                  12
                          1.7230627
## 9
              8
                   5
                          0.6931622
                    1
                          0.2608052
```

From the two tables, the observed and expected frequencies are close enough and thus it does not seem worth fitting a zero inflated model.

(h) Fit a comparable (Gaussian) linear model and graphically compare the fits. Describe how they differ.

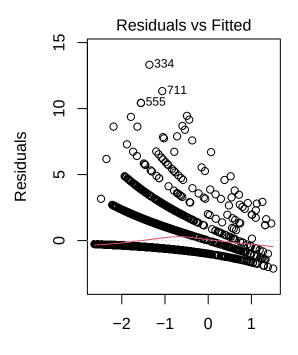
```
dvisitsmod_lm <- lm(Y ~ X)
summary(dvisitsmod_lm)</pre>
```

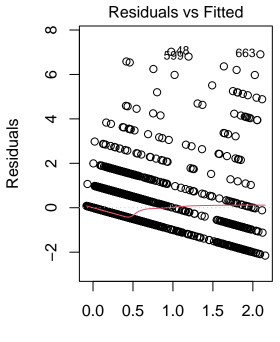
```
##
## Call:
## lm(formula = Y ~ X)
##
## Residuals:
##
       Min
                 10 Median
                                 3Q
                                         Max
  -2.1543 -0.2584 -0.1440 -0.0434
                                     7.0211
##
## Coefficients: (1 not defined because of singularities)
##
                  Estimate Std. Error t value
                                                            Pr(>|t|)
## (Intercept)
                  0.036781
                             0.035794
                                         1.028
                                                            0.304201
## X(Intercept)
                                   NA
                                            NA
                        NΑ
                                                                  NΑ
## Xsex
                  0.035574
                             0.021505
                                         1.654
                                                            0.098137
## Xage
                  0.180024
                             0.055912
                                                            0.001291 **
                                         3.220
## Xincome
                 -0.061208
                             0.030522
                                        -2.005
                                                            0.044975
## Xlevyplus
                 0.024041
                             0.021235
                                         1.132
                                                            0.257626
## Xfreepoor
                 -0.111650
                             0.051532
                                        -2.167
                                                            0.030308 *
## Xillness
                  0.060148
                             0.008332
                                         7.219
                                                  0.00000000000602 ***
## Xactdays
                  0.103140
                             0.003656
                                        28.213 < 0.0000000000000000 ***
## Xhscore
                  0.017064
                             0.005180
                                         3.294
                                                            0.000994 ***
## Xchcond1
                  0.005006
                             0.023709
                                         0.211
                                                            0.832776
## Xchcond2
                  0.045319
                             0.035352
                                         1.282
                                                            0.199921
## Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
```

```
##
## Residual standard error: 0.7138 on 5179 degrees of freedom
## Multiple R-squared: 0.2017, Adjusted R-squared: 0.2002
## F-statistic: 130.9 on 10 and 5179 DF, p-value: < 0.0000000000000000022
par(mfrow=c(1,2))
plot(mod_Select, which=1)
title(main = "Poisson regression\n")
plot(dvisitsmod_lm, which=1)
title(main = "linear regression\n")</pre>
```

# **Poisson regression**

# linear regression

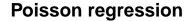




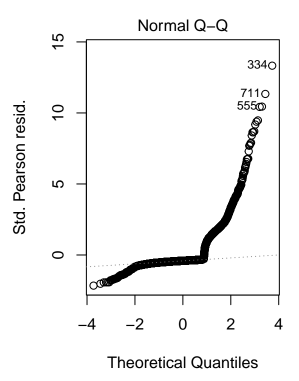
# Predicted values

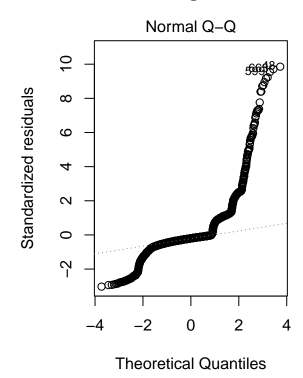
Fitted values

```
par(mfrow=c(1,2))
plot(mod_Select, which=2)
title(main = "Poisson regression\n")
plot(dvisitsmod_lm, which=2)
title(main = "linear regression\n")
```



# linear regression





This seems to indicate how the Poisson regression is a little better because the line for the residuals plot is much more linear around zero, along with the Q-Q plot. In conclusion, we can see that even though we need to account for overdispersion in the Poisson model it is still a better fit than the linear model.

**Problem 8** [20 points]: Analyze the CCSO data set with Days in Jail as the response variable. You are allowed to dichotomize the response into a binary variable. Restrict attention to other traffic offenses as done in class. Your analysis needs to consider the variables considered in class as well as repeat offenders, multiple offenses, released reason, and agency. The determination of repeat offenders and multiple offenses can be done via the jacket number variable. Report interesting significant and null findings and determine that your final model is appropriate. You are allowed to use other inferential techniques than GLM. If you do so, then you need to justify your choices.

#### Solution 8

**Data Wrangling:** We first select the necessary data, here we focus on OTHER TRAFFIC OFFENSES type crimes. Our response variable is the binary variable denoting whether the offender stayed in jail for at least one day or not. We also restrict ourselves with some specific Race, Sex, and Arrest Agencies.

```
CCSO <- fread("https://uofi.box.com/shared/static/9elozjsg99bgcb7gb546wlfr3r2gc9b7.csv")
```

The full data has 67764 observations and 35 features.

```
mutate(atleastone = ifelse(Days > 0,1,0)) %>%
  filter(Crime == "OTHER TRAFFIC OFFENSES") %>%
  filter(Race %in% c("Asian/Pacific Islander", "Black", "White", "Hispanic")) %>%
  filter(Sex %in% c("Female", "Male")) %>%
  ## Agency (and possible interactions with Agency) is interesting if you want to go deepe filter(Agenc
  filter(Agency %in% c( "Illinois State Police", "Rantoul Police Department",
         "University of Illinois Police Department", "Urbana Police Department",
         "Champaign County Sherriff's Office", "Champaign Police Department") ) %>%
  dplyr::select(atleastone, Age, Sex, Date, Race, Agency, JNum, `RELEASED REASON`, `EMPLOYMENT STATUS`,
  mutate(Race = fct_drop(Race), Sex = fct_drop(Sex), Agency = fct_drop(Agency))
CCSO_imp <- CCSO_imp[complete.cases(CCSO_imp), ]</pre>
Now, we will combine categories for the features RELEASED REASON, EMPLOYMENT STATUS, INCARCERATION
REASON in order to reduce number of factors in these features.
CCSO_imp$`RELEASED REASON` [CCSO_imp$`RELEASED REASON` %in% c("Cash Bond Posted", "Paid Fine, Court Costs
CCSO imp$'RELEASED REASON' [CCSO imp$'RELEASED REASON' %in% c("Credit Card Bond Posted")] = "CardBonds"
CCSO_imp$`RELEASED REASON`[CCSO_imp$`RELEASED REASON` %in% c("Placed on Probation
CCSO_imp$`RELEASED REASON`[CCSO_imp$`RELEASED REASON` %in% c("Release on Personal Recognizance
CCSO_imp$`RELEASED REASON`[CCSO_imp$`RELEASED REASON` %in% c("Served Sentence of Incarceration
CCSO_imp$`RELEASED REASON` [CCSO_imp$`RELEASED REASON` %in% c("Transfer to other county/state authoritie
CCSO_imp$`RELEASED REASON` [!(CCSO_imp$`RELEASED REASON` %in% c("Bonds paid", "Conditional Release", "Rele
CCSO_imp$`EMPLOYMENT STATUS` [CCSO_imp$`EMPLOYMENT STATUS` %in% c("Employed - Full Time", "Employed - Par
CCSO_imp$`EMPLOYMENT STATUS`[CCSO_imp$`EMPLOYMENT STATUS` %in% c("Student", "Retired")] = "Student/Reti
CCSO_imp$`EMPLOYMENT STATUS` [CCSO_imp$`EMPLOYMENT STATUS` %in% c("Unemployed", "Laid Off")] ="Unemployed"
CCSO_imp$`INCARCERATION REASON`[CCSO_imp$`INCARCERATION REASON` %in% c("Arrest - Without Warrant", "Arre
CCSO_imp$\incarceration reason\[CCSO_imp$\incarceration reason\] %in% c("FTA - CITY WARRANT (OV)","FTA -
CCSO_imp$\incarceration reason\[CCSO_imp$\incarceration reason\] %in% c("Sentenced - EHD", "Sentenced")] =
CCSO imp$\incarceration reason\[!(CCSO imp$\incarceration reason\] %in% c("Arrest", "FTA", "Sentenced"))] =
## Category Recoding
## 1. only keep arrest data; other categories are sparse, unknown, or have
      a disruptive effect on modeling (sentencing for example)
## 2. remove Conditional Release, Served/Acquitted, and Transfered from
      release reason. These levels seem to disrupt the model (have abnormally
      high propensity for incarcerations lasting one or more days)
CCSO_small <- CCSO_imp %>% rename(Inc_Reason = "INCARCERATION REASON",
                            Emp_Status = "EMPLOYMENT STATUS",
                            Rls_Reason = "RELEASED REASON") %>%
  filter(Inc Reason %in% "Arrest") %>%
  filter(!Rls_Reason %in% c("Conditional Release", "Served/Acquitted",
                            "Transfered")) %>%
  dplyr::select(-Inc_Reason)
```

Next, we add two derived features as directed in the problem: One denoting whethere the offender did multiple offenses in the same day (called multiple), and the other denoting whether they are repeat offenders (called repeatOffender)

```
date_and_jnum <- paste(CCSO_small$Date, CCSO_small$JNum)
dup_date_and_jnum <- date_and_jnum[duplicated(date_and_jnum)]
dup_date_and_jnum <- unique(dup_date_and_jnum)
CCSO_small <- cbind(CCSO_small, as.numeric(is.element(date_and_jnum, dup_date_and_jnum)))
CCSO_small <- CCSO_small %>% rename(multiple = 'V2')

CCSO_small <- cbind(CCSO_small, as.numeric(duplicated(CCSO_small$JNum)))
CCSO_small <- CCSO_small %>% rename(repeatOffender = 'V2')

CCSO_small <- CCSO_small %>% dplyr::select(-c('JNum', 'Date'))
```

We show three rows from our final dataset. It has 5193 observations and 9 features

```
head(CCSO_small, 3)
```

```
##
      atleastone Age
                       Sex Race
                                                             Agency Rls_Reason
## 1:
              0 22
                                        Champaign Police Department
                     Male White
                                                                         Other
## 2:
              0 26 Male White Champaign County Sherriff's Office
                                                                         Other
              0 32 Female White
                                        Champaign Police Department
## 3:
                                                                      Released
##
          Emp_Status multiple repeatOffender
## 1: Student/Retired
                            0
                            0
                                           0
## 2:
            Employed
                            0
                                           0
## 3:
            Employed
```

## **Model Fitting**

Let us fit our full model here

Let us first test the goodness of the model against the null model and a model with no covariates (iid Bernoulli trials)

```
## [1] 0
```

**Model Selection** We can see not all the features considered show a significant effect on the response. Thus, we can do feature selection via stepwise method

```
mod_Select = stepAIC(mymod)

## Start: AIC=2839.14

## atleastone ~ -1 + (Age + Sex + Race + Agency + Rls_Reason + Emp_Status +
```

```
##
      multiple + repeatOffender)
##
##
                   Df Deviance
                                 AIC
                        2805.6 2837.6
## - Age
                    1
## - Rls_Reason
                        2806.5 2838.5
## <none>
                        2805.1 2839.1
## - repeatOffender 1 2807.4 2839.4
## - multiple
                    1 2813.5 2845.5
## - Agency
                    5
                       2846.3 2870.3
## - Sex
                    2 2849.0 2879.0
## - Race
                    3 2892.9 2920.9
                    3 2904.5 2932.5
## - Emp_Status
##
## Step: AIC=2837.64
## atleastone ~ Sex + Race + Agency + Rls_Reason + Emp_Status +
##
      multiple + repeatOffender - 1
##
##
                   Df Deviance
                                 AIC
## - Rls_Reason
                    1 2806.9 2836.9
## <none>
                        2805.6 2837.6
## - repeatOffender 1 2807.9 2837.9
## - multiple
                   1 2813.9 2843.9
                    5 2847.7 2869.7
## - Agency
## - Sex
                    2 2849.2 2877.2
## - Race
                    3 2894.1 2920.1
## - Emp_Status
                    3 2905.3 2931.3
##
## Step: AIC=2836.93
## atleastone ~ Sex + Race + Agency + Emp_Status + multiple + repeatOffender -
##
##
##
                   Df Deviance
                                 AIC
## <none>
                        2806.9 2836.9
## - repeatOffender 1
                        2809.2 2837.2
## - multiple
                    1
                      2815.5 2843.5
                    5 2849.7 2869.7
## - Agency
## - Sex
                    2 2849.6 2875.6
## - Race
                    3 2896.7 2920.7
## - Emp_Status
                    3 2907.7 2931.7
summary(mod_Select)
##
## glm(formula = atleastone ~ Sex + Race + Agency + Emp_Status +
##
      multiple + repeatOffender - 1, family = "binomial", data = CCSO_small,
##
      x = "TRUE")
##
## Deviance Residuals:
           1Q Median
                                 3Q
      Min
                                         Max
## -1.2735 -0.4534 -0.3169 -0.2628
                                      2.7644
##
## Coefficients:
##
                                                Estimate Std. Error z value
## SexFemale
                                                -3.34684 0.66755 -5.014
```

```
## SexMale
                                                   -2.88231
                                                               0.65704 -4.387
## RaceBlack
                                                    1.59003
                                                               0.52130
                                                                         3.050
## RaceHispanic
                                                    0.41957
                                                               0.54283
                                                                         0.773
## RaceWhite
                                                    0.69330
                                                               0.52209
                                                                         1.328
## AgencyChampaign Police Department
                                                    0.14061
                                                               0.15919
                                                                         0.883
## AgencyIllinois State Police
                                                    0.26231
                                                               0.15085
                                                                         1.739
## AgencyRantoul Police Department
                                                    1.21401
                                                               0.19390
                                                                         6.261
## AgencyUniversity of Illinois Police Department 0.00388
                                                               0.20902
                                                                         0.019
## AgencyUrbana Police Department
                                                    0.47956
                                                               0.17086
                                                                         2.807
## Emp_StatusEmployed
                                                   -0.77713
                                                               0.39418 -1.972
## Emp_StatusStudent/Retired
                                                   -0.75458
                                                               0.43082 - 1.751
## Emp_StatusUnemployed
                                                    0.30138
                                                               0.39581
                                                                         0.761
## multiple
                                                   -0.36809
                                                               0.12637
                                                                        -2.913
## repeatOffender
                                                    0.18779
                                                               0.12567
                                                                         1.494
##
                                                         Pr(>|z|)
## SexFemale
                                                   0.000000534185 ***
## SexMale
                                                   0.000011503639 ***
## RaceBlack
                                                          0.00229 **
## RaceHispanic
                                                          0.43957
## RaceWhite
                                                          0.18420
## AgencyChampaign Police Department
                                                          0.37707
## AgencyIllinois State Police
                                                          0.08207 .
                                                   0.00000000382 ***
## AgencyRantoul Police Department
## AgencyUniversity of Illinois Police Department
                                                          0.98519
## AgencyUrbana Police Department
                                                          0.00500 **
## Emp_StatusEmployed
                                                          0.04866 *
## Emp_StatusStudent/Retired
                                                          0.07986
## Emp_StatusUnemployed
                                                          0.44640
                                                          0.00358 **
## multiple
## repeatOffender
                                                          0.13511
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## (Dispersion parameter for binomial family taken to be 1)
##
##
       Null deviance: 7199.0 on 5193 degrees of freedom
## Residual deviance: 2806.9 on 5178 degrees of freedom
## AIC: 2836.9
##
## Number of Fisher Scoring iterations: 6
```

The Age and 'release reason" features are removed. Let us examine if this is a good fit. We can notice this model is significantly better than the null model and the full model is not significantly better than this model. We are satisfied with this selected model

```
#Testing against the saturated model
pchisq(mod_Select$deviance - mymod$deviance, df = mod_Select$df.residual - mymod$df.residual, lower = F.
## [1] 0.4084187
#Testing against the null model
pchisq(mymod$null.deviance - mod_Select$deviance, df = mymod$df.null - mod_Select$df.residual,
lower = FALSE)
```

However, our graphical diagnostics tell a different story. It appears that there is some lack of model fit over

## [1] 0

the bulk of the data and some individuals with predictive probability close to 1.

```
library(heatmapFit)
y <- CCSO_small$atleastone
p1 <- predict(mod_Select, type = "response", se.fit = FALSE)
heatmap.fit(y, p1)
##
## Calculating optimal loess bandwith...
## aicc Chosen Span = 0.9504823
##
## Generating Bootstrap Predictions...
##
                              Predicted Probability Deviation
                        Model Predictions vs. Empirical Frequency
                                                                             p-Value
                            Heat Map Plot
                                                                             Legend
                                                                      0.5
Smoothed Empirical Pr(y=1)
                    heat map line
                                                                      0.4
                    perfect fit
     9.0
                                                                      0.3
     0.2
                                                                      0.2
     0.0
                                                                      0.1
          0.0
                   0.1
                            0.2
                                     0.3
                                              0.4
                                                       0.5
                                                                    0.01
                        Model Prediction, Pr(y=1)
##
##
## 74.48488% of Observations have one-tailed p-value <= 0.1
## Expected Maximum = 20%
```

The variable multiple offenses is inherently vague in nature. Some offenses can be more severe than others so this might disrupt our model. Let's remove everyone with multiple offenses and restrict attention to traffic offenses only. When we do this, the model is better fitting according to our visual diagnostics and it appears that some previous conclusions change (note that violations in fit appear where there is not much data and is extrapolated to where there is no data). Namely, race exhibits a much weaker association with propensity for incarcerations lasting one or more days, and repeat offenders seem to be punished more.

```
m1 <- glm(atleastone ~ Sex + Race + Agency + repeatOffender,</pre>
    family = "binomial", data = CCSO_small %>% filter(multiple == 0))
m2 <- glm(atleastone ~ Sex + Agency + repeatOffender,</pre>
    family = "binomial", data = CCSO_small %>% filter(multiple == 0))
anova(m2, m1, test = "LRT")
## Analysis of Deviance Table
##
## Model 1: atleastone ~ Sex + Agency + repeatOffender
## Model 2: atleastone ~ Sex + Race + Agency + repeatOffender
     Resid. Df Resid. Dev Df Deviance
                                            Pr(>Chi)
## 1
          1775
                   1292.2
## 2
          1772
                    1256.1 3
                                36.117 0.00000007074 ***
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
heatmap.fit(CCSO_small %% filter(multiple == 0) %>% pull(atleastone),
            predict(m1, type = "response", se.fit = FALSE))
##
## Calculating optimal loess bandwith...
## aicc Chosen Span = 0.9584928
## Generating Bootstrap Predictions...
                                                                                       1
##
                             Predicted Probability Deviation
                       Model Predictions vs. Empirical Frequency
                                                                           p-Value
                           Heat Map Plot
                                                                           Legend
                                                                    0.5
Smoothed Empirical Pr(y=1)
                   heat map line
     0.4
                                                                    0.4
                   perfect fit
                                                                    0.3
     0.2
                                                                    0.2
```

Model Prediction, Pr(y=1)

0.3

0.2

##

0.1

0.0

##

0.4

0.5

0.1

0.01

```
## 15.5917% of Observations have one-tailed p-value <= 0.1
## Expected Maximum = 20%
## ************
## compare with saturated
pchisq(m1$deviance, m1$df.residual, lower = FALSE)
## [1] 1
Summary table for final model
summary(m1)
##
## Call:
  glm(formula = atleastone ~ Sex + Race + Agency + repeatOffender,
       family = "binomial", data = CCSO_small %>% filter(multiple ==
##
          0))
##
## Deviance Residuals:
      Min
                10
                     Median
                                   3Q
                                           Max
  -1.1960 -0.5328 -0.4264 -0.3339
##
                                        2.4723
##
## Coefficients:
##
                                                  Estimate Std. Error z value
## (Intercept)
                                                  -3.47204
                                                             0.76869 - 4.517
## SexMale
                                                   0.54505
                                                             0.19623
                                                                        2.778
## RaceBlack
                                                   1.23425
                                                              0.74135
                                                                       1.665
## RaceHispanic
                                                  -0.08086
                                                              0.77987 -0.104
## RaceWhite
                                                   0.55369
                                                              0.74313
                                                                       0.745
## AgencyChampaign Police Department
                                                              0.22179
                                                                       0.670
                                                   0.14864
## AgencyIllinois State Police
                                                   0.24726
                                                              0.25868
                                                                       0.956
## AgencyRantoul Police Department
                                                   1.03533
                                                              0.28236
                                                                        3.667
## AgencyUniversity of Illinois Police Department 0.02118
                                                              0.30176
                                                                        0.070
## AgencyUrbana Police Department
                                                   0.33929
                                                              0.23078
                                                                        1.470
## repeatOffender
                                                   0.70111
                                                              0.26443
                                                                        2.651
##
                                                    Pr(>|z|)
## (Intercept)
                                                  0.00000628 ***
## SexMale
                                                    0.005475 **
## RaceBlack
                                                    0.095936 .
## RaceHispanic
                                                    0.917424
## RaceWhite
                                                    0.456224
## AgencyChampaign Police Department
                                                    0.502732
## AgencyIllinois State Police
                                                    0.339133
## AgencyRantoul Police Department
                                                    0.000246 ***
## AgencyUniversity of Illinois Police Department
                                                    0.944039
## AgencyUrbana Police Department
                                                    0.141502
## repeatOffender
                                                    0.008017 **
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## (Dispersion parameter for binomial family taken to be 1)
##
##
       Null deviance: 1324.5 on 1782 degrees of freedom
## Residual deviance: 1256.1 on 1772 degrees of freedom
## AIC: 1278.1
```

#### ##

## Number of Fisher Scoring iterations: 5

**Discussion** Now, finally - if we look at the summary table - the features that are most strongly associated with propensity of incarcerations lasting one or more days:

- Race seems to be a factor, and being black seems to be associated with longer incarcerations.
   However, the association between race and propensity of incarcerations lasting one or more days is not very strong when we consider important confounding factors.
- One agency 'Rantoul Police Department' stands out among all agencies. It appears people arrested by this agency tends to have longer incarcerations.
- — Repeat offenses is strongly associated with longer incarcerations.
- — Males are strongly associated with longer incarcerations.

We should remember this is observational data and we can not make causal connections with confidence.