# MAST90104: A First Course in Statistical Learning

# Week 11 Lab and Workshop

# Practical questions

- 1. The cornnit dataset in the faraway package contains data on the effect of nitrogen on the yield of corn. Fit a gamma regression to this data, using the glm command and store the model fit as gmod, using the canonical link function. Hint: consider transforming the predictor variable first.
  - (a) Extract the Pearson residuals from the fitted model using the **residuals** function, then use them to estimate the dispersion parameter. Check that your answer agrees with the summary output from your model.

### Solution:

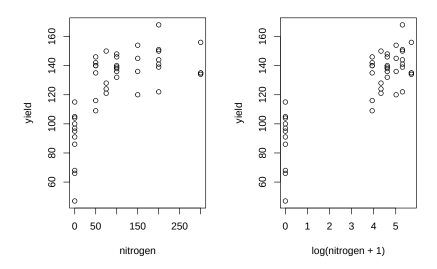


Figure 1: Scatter plot of yield against nitrogen on left panel and yield again log(nitrogen + 1) on right panel.

```
> library(faraway)
> data("cornnit")
> fivenum(cornnit$nitrogen)
[1]     0.0     25.0     87.5     175.0     300.0
> par(mfrow=c(1,2))
> plot(yield~nitrogen,data=cornnit)
> plot(yield~log(nitrogen+1),data=cornnit)
#Judging from scatter plots, log(nitrogen+1) would be a suitable transformation to #ensure linear relationship between response and predictor
> gmod <- glm(yield~log(nitrogen+1), family=Gamma, data = cornnit)
> summary(gmod)

Call:
glm(formula = yield ~ log(nitrogen + 1), family = Gamma, data = cornnit)
```

```
Coefficients:
```

```
Estimate Std. Error t value Pr(>|t|)
(Intercept) 1.106e-02 4.349e-04 25.439 < 2e-16 ***
log(nitrogen + 1) -7.902e-04 9.538e-05 -8.285 2.25e-10 ***
---
Signif. codes: 0 '***, 0.001 '**, 0.01 '*, 0.05 '., 0.1 ', 1
```

(Dispersion parameter for Gamma family taken to be 0.01874595)

Null deviance: 2.40614 on 43 degrees of freedom Residual deviance: 0.91727 on 42 degrees of freedom

AIC: 383.74

Number of Fisher Scoring iterations: 4
> sum(residuals(gmod, type = "pearson")^2)/(42)
[1] 0.01874588

The estimate for  $\phi$  is indeed 0.01875.

(b) The command anova(gmod, test="F") will compare your model against the intercept-only model, using an F test. Using the deviances and dispersion estimates reported by summary(gmod), check that the F statistic reported by the anova function is correct.

#### Solution:

```
> (2.40614-0.91727)/0.01874595/1
[1] 79.42356
> anova(gmod,test="F")
Analysis of Deviance Table
```

Model: Gamma, link: inverse

Response: yield

Terms added sequentially (first to last)

```
Df Deviance Resid. Df Resid. Dev F Pr(>F)

NULL 43 2.40614

log(nitrogen + 1) 1 1.4889 42 0.91727 79.423 3.11e-11 ***

---

Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
```

The F-statistic is indeed 79.423.

(c) Now do some diagnostic plots. Can you identify a potential outlier?

**Solution:** Based on the jack-knife residual plot and the Cook's distance plot, it is clear that both observation 21 is an influential outliers.

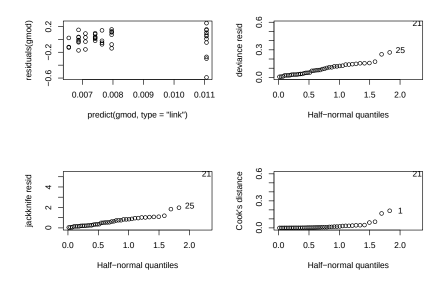


Figure 2: Residual plots for Gamma regression model

(d) Fit a linear model to the cornnit data. Which do you prefer, the linear model or the gamma model, and why?

### Solution:

```
> gmod.norm <- lm(yield ~ log(nitrogen+1), data=cornnit)</pre>
> summary(gmod.norm)
lm(formula = yield ~ log(nitrogen + 1), data = cornnit)
Residuals:
Min
         1Q Median
                         3Q
                                Max
-42.335 -10.261
                  2.126
                         10.558
                                 25.665
Coefficients:
                   Estimate Std. Error t value Pr(>|t|)
(Intercept)
                    89.335
                                 4.227
                                         21.13 < 2e-16 ***
                    10.201
                                 1.017
                                         10.03 1.03e-12 ***
log(nitrogen + 1)
                0 '***, 0.001 '**, 0.01 '*, 0.05 '., 0.1 ', 1
Signif. codes:
Residual standard error: 14.34 on 42 degrees of freedom
Multiple R-squared: 0.7055, Adjusted R-squared: 0.6985
F-statistic: 100.6 on 1 and 42 DF, p-value: 1.025e-12
> par(mfrow=c(2,2))
> plot(predict(gmod.norm, type="response"), residuals(gmod.norm))
> halfnorm(residuals(gmod.norm), ylab="deviance resid")
> halfnorm(rstudent(gmod.norm), ylab="jackknife resid")
> halfnorm(cooks.distance(gmod.norm), ylab="Cook's distance")
> plot(predict(gmod.norm, type="response")~predict(gmod, type = "response"))
> abline(0,1)
```

Observation 21 appears to be an influential observation under both the gamma and Gaussian regression models, so in this respect linear regression doesn't improve the model fit. However, the deviance residuals are much smaller for the Gamma regression than in the linear regression. Hence, we prefer the gamme regression model.

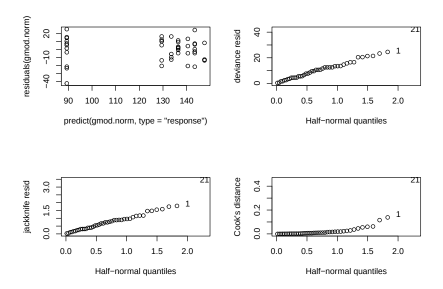


Figure 3: Residual plots for linear (Gaussian) regression model

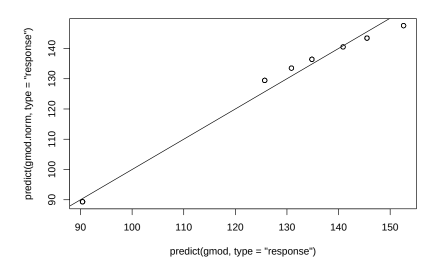


Figure 4: Correspondence of predicted values between gamma regression and linear regression models

- 2. The Articles dataset in the Rchoice package contains data on the publication counts (art) of research scientists and their respective gender (fem), marital status (mar), number of children (kid5), prestige of graduate program (phd), and the number of articles published by their mentors (ment).
  - (a) Fit a Poisson regression model with art as the response variable using the canonical link function.

```
#install.packages("Rchoice")
> library(Rchoice)
> data("Articles")
> mod.pois <- glm(art~., family=poisson, data = Articles)</pre>
```

(b) Perform stepwise selection using AIC criterion starting from the full Poisson regression model with all predictors. Write down the equation of your your final regression model.

```
> step(mod.pois)
Start: AIC=3314.11
art ~ fem + mar + kid5 + phd + ment
Df Deviance
             AIC
- phd 1 1634.6 3312.3
<none>
          1634.4 3314.1
- mar 1 1640.8 3318.5
- fem 1 1651.5 3329.2
- kid5 1 1656.5 3334.2
- ment 1 1766.2 3444.0
Step: AIC=3312.35
art ~ fem + mar + kid5 + ment
Df Deviance
             AIC
<none>
          1634.6 3312.3
- mar 1 1640.8 3316.6
      1 1651.8 3327.5
- fem
         1656.7 3332.4
- kid5 1
- ment 1 1776.7 3452.5
Call: glm(formula = art ~ fem + mar + kid5 + ment, family = poisson,
data = Articles)
Coefficients:
(Intercept)
                                          kid5
                  fem
                         mar
0.34517 -0.22530 0.15218 -0.18499
ment
0.02576
Degrees of Freedom: 914 Total (i.e. Null); 910 Residual
Null Deviance:
                1817
Residual Deviance: 1635 AIC: 3312
> mod.pois.final <- glm(art ~ fem + mar + kid5 + ment, family = poisson,data = Articles)
> summary(mod.pois.final)
Call:
glm(formula = art ~ fem + mar + kid5 + ment, family = poisson,
data = Articles)
Coefficients:
Estimate Std. Error z value Pr(>|z|)
(Intercept) 0.34517 0.06012 5.741 9.41e-09 ***
          -0.22530
                     0.05461 -4.125 3.70e-05 ***
fem
mar
           -0.18499
                     0.04014 -4.609 4.05e-06 ***
kid5
           0.02576
                     0.00195 13.212 < 2e-16 ***
ment
Signif. codes:
0 '***, 0.001 '**, 0.01 '*, 0.05 '., 0.1 ', 1
(Dispersion parameter for poisson family taken to be 1)
Null deviance: 1817.4 on 914 degrees of freedom
Residual deviance: 1634.6 on 910 degrees of freedom
```

AIC: 3312.3

Number of Fisher Scoring iterations: 5

The fitted final regression model is  $y_i \sim \text{Poisson}(\mu_i)$ , i = 1, ..., n, where

$$\log(\mu_i) = \beta_0 + \beta_1 \text{fem}_i + \beta_2 \text{mar}_i + \beta_3 \text{kid5}_i + \beta_4 \text{ment}_i.$$

(c) The glm.nb command in the MASS library fits the following model:  $y_i \sim \text{NegBin}(\mu_i, k)$ , where  $\mu_i > 0, k > 0$ , and

$$p(y_i = y) = \frac{\Gamma(y+k)}{y!\Gamma(k)} \left(\frac{\mu_i}{\mu_i + k}\right)^y \left(\frac{k}{\mu_i + k}\right)^k, \ y = 0, 1, 2, \dots$$

Here,  $E(y_i) = \mu_i$  and  $Var(y_i) = \mu_i + \mu_i^2/k$ . The default log link function is  $g(\mu) = \log(\mu)$ . Using glm.nb, fit a Negative Binomial regression model with art as the response variable using the log link function.

- > library(MASS)
- > mod.nb <- glm.nb(art~., data = Articles)
- (d) Perform stepwise selection using AIC criterion starting from the full Negative Binomial regression model with all predictors. Write down the equation of your your final regression model.

```
> step(mod.nb,scope=~.)
Start: AIC=3133.92
```

art ~ fem + mar + kid5 + phd + ment

```
Df Deviance
               AIC
           1004.5 3132.1
- phd
<none>
            1004.3 3133.9
           1007.6 3135.3
- mar
        1
- fem
        1
            1013.2 3140.8
            1015.5 3143.1
- kid5 1
- ment 1
            1079.1 3206.7
```

Step: AIC=3132.1

art ~ fem + mar + kid5 + ment

```
AIC
Df Deviance
<none>
           1004.4 3132.1
- mar
          1007.7 3133.3
       1
+ phd
          1004.2 3133.9
      1
           1013.3 3139.0
- fem
       1
- kid5 1
           1015.6 3141.3
- ment 1
           1085.3 3211.0
```

Call: glm.nb(formula = art ~ fem + mar + kid5 + ment, data = Articles,
init.theta = 2.264120074, link = log)

## Coefficients:

```
(Intercept) fem mar kid5
0.30333 -0.21667 0.14694 -0.17680
ment
0.02943
```

Degrees of Freedom: 914 Total (i.e. Null); 910 Residual

Null Deviance: 1109

Residual Deviance: 1004 AIC: 3134

> mod.nb.final <- glm.nb(formula = art ~ fem + mar + kid5 + ment, data = Articles)

> summary(mod.nb.final)

```
Call:
```

```
glm.nb(formula = art ~ fem + mar + kid5 + ment, data = Articles,
init.theta = 2.26411693, link = log)
```

#### Coefficients:

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

```
(Intercept) 0.303328
                       0.081389
                                  3.727 0.000194 ***
                       0.072624 -2.983 0.002850 **
fem
           -0.216673
            0.146944
                       0.081765
                                 1.797 0.072312 .
mar
kid5
           -0.176797
                       0.052826 -3.347 0.000818 ***
            0.029430
                       0.003108
                                  9.470 < 2e-16 ***
ment
```

---

Signif. codes:

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

(Dispersion parameter for Negative Binomial(2.2641) family taken to be 1)

Null deviance: 1108.9 on 914 degrees of freedom Residual deviance: 1004.4 on 910 degrees of freedom

AIC: 3134.1

Number of Fisher Scoring iterations: 1

Theta: 2.264 Std. Err.: 0.271

2 x log-likelihood: -3122.096

The final fitted regression model is:  $y_i \sim \text{NegBin}(\mu_i, k)$ , where

$$\log(\mu_i) = \beta_0 + \beta_1 \mathtt{fem}_i + \beta_2 \mathtt{mar}_i + \beta_3 \mathtt{kid5}_i + \beta_4 \mathtt{ment}_i.$$

(e) Which model would you prefer – the Poisson or Negative Binomial? Justify your answer with a suitable residual plot.

By assessing the deviance residual-half normal quantile plots, the Poisson regression yields more observations with extreme residuals. Furthermore, the AIC of the final NegBin regression model is lower than that for the final Poisson regression model. Hence, we prefer the NegBin model.

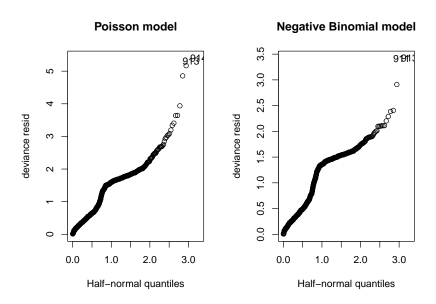


Figure 5: Deviance residual plots for final Poisson (left) and negative binomial regression (right) models