# R Code And Tasks Chapter 6 (MAS 6003)

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# 6.1 Types of 2 way tables - response / control variables

#### 6.1.1 Case(a): Skin cancer (melanoma) data - 2 response variables

**Cross sectional** study of malignant melanoma. Both tumor type and site are **response variables** because none of the row or column totals were fixed in advance of the data collection.

```
library(reshape2)
rm(list=ls())
load("data/MAS367-GLMs.RData", envir = e <- new.env())</pre>
Mela <- e$Mela
head(Mela)
##
     number tumour.type
## 1
         22
                       A Head.and.Neck
## 2
                       B Head.and.Neck
## 3
         19
                       C Head.and.Neck
## 4
         11
                       D Head.and.Neck
## 5
          2
                                  Trunk
                       Α
## 6
         54
                       В
                                  Trunk
dcast(Mela,tumour.type~ site, value.var="number")
     tumour.type Head.and.Neck Trunk Extremities
##
## 1
                Α
                              22
                                      2
                                                  10
## 2
                В
                              16
                                    54
                                                 115
## 3
                С
                              19
                                    33
                                                  73
## 4
                D
                              11
                                    17
                                                  28
```

#### 6.1.2 Case(b): Flu vaccine data - 1 response and 1 control variable

Patients were randomly assigned to the two groups (Placebo, Vaccine), and the response (levels of an antibody found in the blood six weeks after vaccination) was determined. Antibody level is the **response** and vaccine group is a **controlled variable** (with totals fixed by experimental design).

```
vaccine <- e$vaccine
head(vaccine)</pre>
```

```
##
     count response treatment
## 1
        25
              small
                       placebo
## 2
               small
                       vaccine
## 3
         8 moderate
                       placebo
## 4
        18 moderate
                       vaccine
## 5
         5
              large
                       placebo
## 6
        11
               large
                       vaccine
```

#### dcast(vaccine, treatment ~ response, value.var="count")

```
## treatment small moderate large
## 1 placebo 25 8 5
## 2 vaccine 6 18 11
```

#### 6.2.1 Association, Independence and Homogeneity

Independence:

#### Case(a) - skin cancer

Probabilities of interest  $\pi_{ij} = P(A = i, B = j)$ 

$$P(A = i, B = j) = P(A = i) \times P(B = j)$$

for all i and j, where  $\pi_i = P(A=i)$  and  $\pi_j = P(B=j)$  are the marginal probabilities of row i and column j

#### Case(b) - flue vaccine data

Probabilities of interest are conditional probabilities  $\pi i j = P(B = j | A = i)$ .

The interest is in whether the probability distribution of the response (antibody level) is the same in each level of the controlled variable (drug group). If it doesn't depend on i then we can write

$$(\pi_{ij} = \pi_{.j})$$

.

This is known as **homogeneity**.

# 6.3 Distribution for two-way tables

6.3.1 Case(a): two response variables

6.3.2 Case(b): one response variable

6.3.3 Case(c): independent poisson (no fixed margins).

6.3.4 Expected values

### 6.4 GLMs and two-way contingency tables

6.4.1 Natural hypothesis are log-linear models

6.4.2 Poisson log-linear modelling for two-way tables

# 6.4.3 Maximum likelihood estimation for $\pi_{ij}$ in case (a)

#### Task 19

Verify the maximum likelihood estimate for  $\pi_{ij}$  for the A + B model for case(a).

Verify that  $\pi_{ij} = \frac{y_{i} \cdot y_{\cdot j}}{n^2}$ 

and somehow use information that  $\mu_{ij} = n\pi_i\pi_j$ 

#### Task 20

Verify the maximum likelihood estimate for  $\pi_{ij}$  for the A + B model for case(b).

Verify that  $\pi_{ij} = \frac{y \cdot j}{n}$ 

and somehow use information that  $\mu_{ij} = n_i \pi_j$ 

$$\partial l(\mu_i)/\partial \pi_j = \left(\sum_i \sum_j -n_i \pi_j + y_{ij} \log(n_i \pi_j) - \log(y_{ij}!)\right)'$$

$$= \sum_i -n_i + \frac{y_{ij}}{\pi_j}$$

$$= n + \frac{y_{ij}}{\pi_j} = 0$$

$$\pi_{ij} = \frac{y_{\cdot j}}{n}$$

# 6.5 Interaction plots and examination of residues.

interaction.plot honorable mentioned.

# 6.6 Analysis of the skin cancer data (case(a)) using log-linear models

Task 21

#### 6.6.1 Fitted values for the skin cancer data

•  $A \times B$  saturated model.

```
glm.sat <- glm(number ~ factor(tumour.type) * factor(site), family = poisson(log), data=Mela)</pre>
matrix(glm.sat$fitted.values,ncol=3)
        [,1] [,2] [,3]
##
## [1,]
          22
## [2,]
               54 115
          16
                     73
## [3,]
          19
               33
## [4,]
                     28
          11
  • A + B :independence
glm.indep <- glm(number ~ factor(tumour.type) + factor(site), family = poisson(log), data=Mela)</pre>
matrix(glm.indep$fitted.values,ncol=3)
##
         [,1]
                 [,2]
                         [,3]
## [1,] 5.78 9.010 19.210
## [2,] 31.45 49.025 104.525
## [3,] 21.25 33.125 70.625
## [4,] 9.52 14.840 31.640
  • A: independence and the same probability for each column category
glm.indepCol <- glm(number ~ factor(tumour.type), family = poisson(log), data=Mela)</pre>
matrix(glm.indepCol$fitted.values,ncol=3)
##
             [,1]
                      [,2]
                                [,3]
## [1,] 11.33333 11.33333 11.33333
## [2,] 61.66667 61.66667 61.66667
## [3,] 41.66667 41.66667 41.66667
## [4,] 18.66667 18.66667 18.66667
  • B: independence and the same probability for each row category
Calculate the table of fitted values for the linear predictor containing B for case(a)
glm.indepRow <- glm(number ~ factor(site), family = poisson(log), data=Mela)</pre>
res<-matrix(glm.indepRow$fitted.values,ncol=3)
rowSums(res)
## [1] 100 100 100 100
colSums(res)
## [1] 68 106 226
```

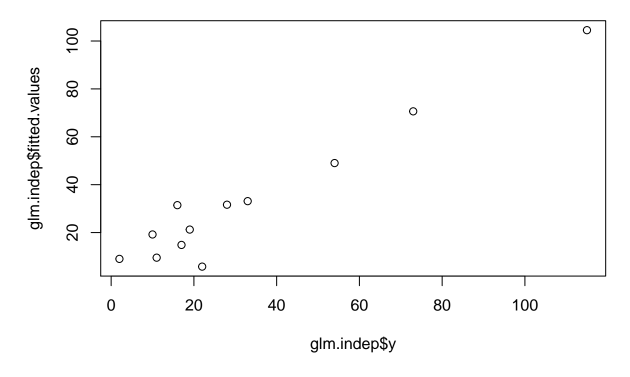
#### 6.6.2

•  $A \times B$ : saturated model

```
void <- glm(number ~ factor(tumour.type) * factor(site), family = poisson(log), data=Mela)</pre>
```

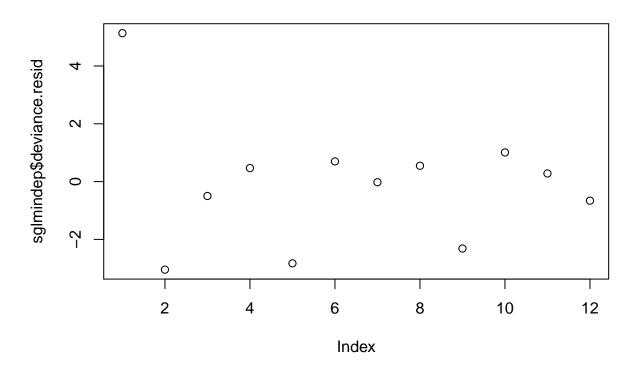
# 6.6.3 Skin cancer data case(a) revisited

```
• 1 The test of independence based on the log-linear model A+B
1 - pchisq(glm.indep$deviance,6)
## [1] 2.050453e-09
qchisq(0.975,6)
## [1] 14.44938
  • 2 The usual Pearson \chi^2 test for independence
chisq.test(matrix(Mela$number,nrow=4))
##
##
   Pearson's Chi-squared test
##
## data: matrix(Mela$number, nrow = 4)
## X-squared = 65.813, df = 6, p-value = 2.943e-12
1-pchisq(65.813,6)
## [1] 2.94309e-12
  • 5
Deviance residuals see 3.6.4 in the notes. Pearson residuals see Task 9 and page 91.
plot(glm.indep$y,glm.indep$fitted.values)
```

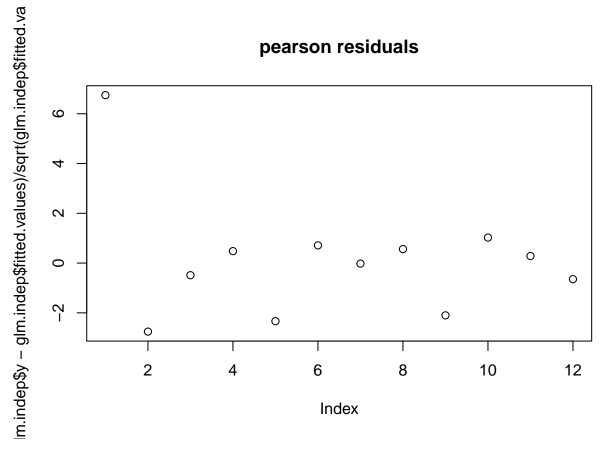


```
sglmindep<-summary(glm.indep)
plot(sglmindep$deviance.resid , main="deviance residuals")</pre>
```

# deviance residuals

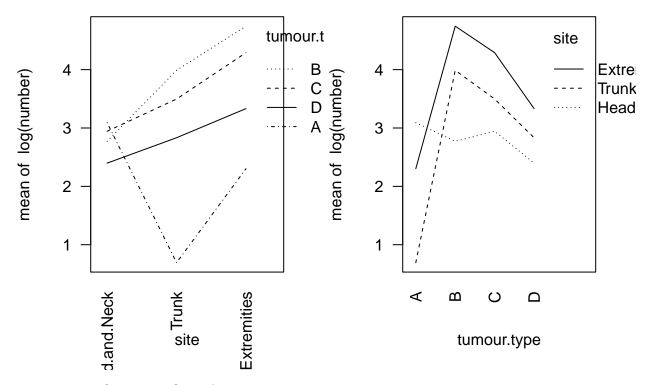


plot((glm.indep\$y-glm.indep\$fitted.values)/sqrt(glm.indep\$fitted.values), main = "pearson residuals")



 $\bullet$  interaction plot

```
par(mfrow=c(1,2))
with(Mela, {
  interaction.plot(site, tumour.type, log(number),las=2)
  interaction.plot(tumour.type, site, log(number),las=2)
})
```



• removing first row or first column

```
remove cancer type A
MelaNoA <- Mela[Mela$tumour.type!="A",]</pre>
glm.indepNoA <- glm(number ~ factor(tumour.type) + factor(site), family = poisson(log), data=MelaNoA)</pre>
glm.indepNoA
##
   Call: glm(formula = number ~ factor(tumour.type) + factor(site), family = poisson(log),
##
##
       data = MelaNoA)
##
   Coefficients:
##
                (Intercept)
                                 factor(tumour.type)C
                                                           factor(tumour.type)D
##
##
                     3.1464
                                              -0.3920
                                                                         -1.1950
         factor(site)Trunk factor(site)Extremities
##
##
                     0.8157
                                               1.5466
##
## Degrees of Freedom: 8 Total (i.e. Null);
                                               4 Residual
## Null Deviance:
                         203.3
## Residual Deviance: 6.509
                                 AIC: 63.91
qchisq(0.975, 4)
```

## [1] 11.14329

remove site Head

```
MelaNoHead <- Mela[Mela$site!="Head.and.Neck",]</pre>
glm.indepNoHead <- glm(number ~ factor(tumour.type) + factor(site), family = poisson(log), data=MelaNoH
glm.indepNoHead
##
## Call: glm(formula = number ~ factor(tumour.type) + factor(site), family = poisson(log),
       data = MelaNoHead)
##
##
## Coefficients:
##
               (Intercept)
                                factor(tumour.type)B
                                                          factor(tumour.type)C
##
                     1.3432
                                              2.6450
                                                                        2.1785
##
      factor(tumour.type)D factor(site)Extremities
##
                    1.3218
                                              0.7571
##
## Degrees of Freedom: 7 Total (i.e. Null); 3 Residual
## Null Deviance:
                        237.2
## Residual Deviance: 2.165
                                 AIC: 52.68
qchisq(0.975, 3)
```

#### ## [1] 9.348404

- 9 and 10 is about pooling the B, C, D or Trunk and Extremities
- 11

A log-linear model can be fitted which is additive in the factors, but includes a term for the (1,1) cell — an indicator variable for that cell (that is, treats it as an outlier).

```
head(Mela)
```

factor(labelAHead)1

##

```
##
     number tumour.type
## 1
         22
                       A Head.and.Neck
## 2
         16
                       B Head.and.Neck
                       C Head.and.Neck
## 3
         19
## 4
         11
                       D Head.and.Neck
                                 Trunk
## 5
          2
                       Α
## 6
         54
                       В
                                 Trunk
MelaFix <- Mela
MelaFix$labelAHead <- rep(0,nrow(Mela))</pre>
MelaFix$labelAHead[1] <- 1</pre>
MelaFix$labelAHead <- as.factor(MelaFix$labelAHead)</pre>
glm.indepNoHead <- glm(number ~ factor(tumour.type) + factor(site) + factor(labelAHead), family = poiss
glm.indepNoHead
##
## Call: glm(formula = number ~ factor(tumour.type) + factor(site) + factor(labelAHead),
##
       family = poisson(log), data = MelaFix)
##
## Coefficients:
##
                (Intercept)
                                factor(tumour.type)B
                                                           factor(tumour.type)C
##
                     0.5452
                                               2.6011
                                                                          2.2091
##
      factor(tumour.type)D
                                    factor(site)Trunk factor(site)Extremities
##
                                               0.7980
```

```
## 2.5458
##
## Degrees of Freedom: 11 Total (i.e. Null); 5 Residual
## Null Deviance: 295.2
## Residual Deviance: 8.002 AIC: 81.11
```

#### Task 22 Verify the analysis (See above)

#### Task 23

For the  $4 \times 3$  table in Example 6.1.1, and the independence model, show directly (with- out fitting a log-linear model) that  $\mu_{11} = 5.780$ ,  $e_{P,11} = 6.747$ , and  $e_{D,11} = 5.135$ .

#### Task 24

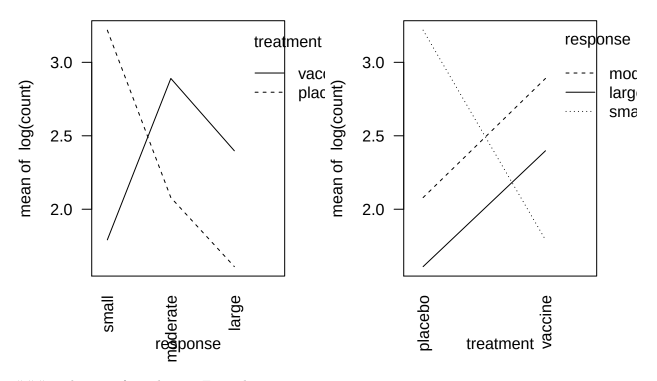
## [1] 4

# 6.7 Flu vaccine data (case(b)) revisited

• minimal model

```
head(vaccine)
     count response treatment
##
                      placebo
## 1
        25
              small
## 2
         6
              small
                      vaccine
        8 moderate
                      placebo
## 4
        18 moderate
                      vaccine
## 5
        5
              large
                      placebo
## 6
        11
              large
                      vaccine
dim(vaccine)
## [1] 6 3
glm.null <- glm(count ~ 1 , family = poisson(log), data=vaccine)</pre>
glm.null$df.null
## [1] 5
glm.min <- glm(count ~ treatment , family = poisson(log), data=vaccine)</pre>
glm.min
##
## Call: glm(formula = count ~ treatment, family = poisson(log), data = vaccine)
##
## Coefficients:
##
        (Intercept) treatmentvaccine
##
            2.53897
                              -0.08224
##
## Degrees of Freedom: 5 Total (i.e. Null); 4 Residual
## Null Deviance:
                         23.81
## Residual Deviance: 23.68
                                 AIC: 52.81
glm.min$df.residual
```

```
qchisq(0.95,4)
## [1] 9.487729
  • homogeneity model (A+B)
head(vaccine)
     count response treatment
## 1
        25
              small
                     placebo
## 2
        6
              small
                     vaccine
## 3
        8 moderate placebo
## 4
        18 moderate
                     vaccine
## 5
        5
              large
                      placebo
## 6
        11
              large
                      vaccine
glm.homo <- glm(count ~ response + treatment , family = poisson(log), data=vaccine)</pre>
glm.homo$df.residual
## [1] 2
glm.homo$deviance
## [1] 18.64253
qchisq(0.95,4)
## [1] 9.487729
  • not much of an improvement on 2 degrees of freedom
glm.min$deviance - glm.homo$deviance
## [1] 5.041382
qchisq(0.95,2)
## [1] 5.991465
  • groups differ in their response
Again interaction.plot can be used.
par(mfrow=c(1,2))
with(vaccine, {
interaction.plot(response, treatment, log(count),las=2)
interaction.plot(treatment, response, log(count),las=2)
})
```



### Task 25 verify analysis in Example 6.7

Task 26 What is the largest pearson residual for the A+B model?