

This course was developed as a part of the VLIR-UOS Cross-Cutting project s:

•Statistics: 2011-2016, 2017.

Statistics: 2017.

•Statistics for development : 2018-2020.

•The >eR-BioStat ITP: 2024-2026.



The >eR-Biostat initiative

Making R based education materials in statistics accessible for all

Applied Generalized Linear Models (GLM) using R (PART 2)

Developed by

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ER-BioStat

GitHub https://github.com/eR-Biostat





@erbiostat

Reference list

- Main reference
 - Dobson (2002): An introduction to generalized linear models.
- Other references:
 - McCillagh and Nelder (1983): Generalized linear models (first edition).
 - Collet D(1994): Modeling Binary data.
 - Lindsey (1997): Applying generalized linear models.



Software

- Two main R functions:
 - Linear models in R: the lm() function.
 - Generalized linear models in R: the glm() function in R.
- All R programs for the examples presented in the slides are available online.



Datasets

- Data are given as a part of R programs for the course.
- External datasets (which are not given as a part of the R code) and used for illustration are available online.

Topics (part 2)

- 11. Poisson Regression
- 12. Beyond Poisson and binomial distributions: models with different link functions and/or distributions
- 13. Poisson regression and log linear models
- 14. Over dispersion

Chapter 11: Poisson Regression

Donson: chapter 7.

Lindsey: Appendix B.

McCullagh & Nelder: chapter 2.

Count data

Count data:

- counts per unit of time/area/distance, etc
- contingency tables: counts cross-classified by categorical variables
- Covariates: categorical or continuous

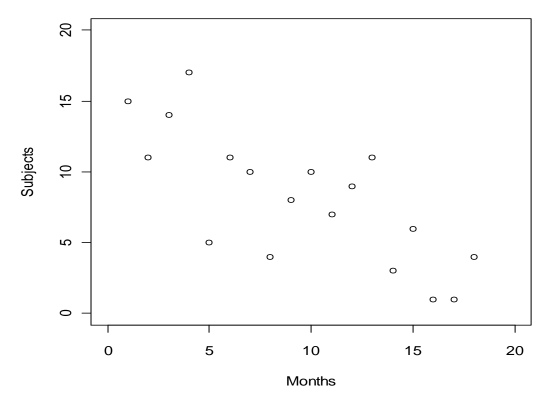
Example 1: Stress data

 One randomly chosen member from each randomly chosen household in a sample from Oakaland, California, USA was interviewed. In a list of 41 events, respondents were asked to note which had occurred within the last 18 months. The result is given as:

Month	1	2	3	4	5	6	7	8	9
Respondents	15	11	14	17	5	11	10	4	8
Month	10	11	12	13	14	15	16	17	18
Respondents	10	7	9	11	3	6	1	1	14

Data in R

```
> stress <- read.table("C:..../stress.txt", sep=",",header=TRUE)
> attach(stress)
plot(respondents ~ month, xlab = "Months",ylab = "Subjects",
    xlim=c(0,20), ylim=c(0,20))
```



Model formulation

The distribution of the response variable

$$Y_t \sim Poisson(\mu_t)$$

$$f(Y_t, \mu) = \frac{\mu^{y_t} e^{-\mu}}{Y_t!}$$

$$E(Y_t) = \mu_t$$

The dependency on the predictor

$$\mu_{t} = f(month)$$

A proposal?

$$\mu_t = \beta_0 + \beta_1 t$$

Model formulation

The distribution of the response variable

$$Y_t \sim Poisson(\mu_t)$$

$$E(Y_t) = \mu_t$$

The linear predictor

$$\mu_t = f(month)$$

$$\eta = \beta_0 + \beta_1 M_t$$

$$\mu_{\scriptscriptstyle t} = e^{\eta} = e^{\beta_0 + \beta_1 M_{\scriptscriptstyle t}}$$

$$g(E(Y_t)) = \log(\mu_t) = \eta$$

GLM for Poisson regression using glm()

$$\eta = \beta_0 + \beta_1 t$$

> respGLM <- glm(respondents ~ month, family=poisson, data=stress)

The relative risk

$$RR = \frac{E(Y_t \mid t+1)}{E(Y_t \mid t)} = \frac{e^{\beta_0 + \beta_1(t+1)}}{e^{\beta_0 + \beta_1 t}} = e^{\beta_1}$$

GLM for Poisson regression using glm()

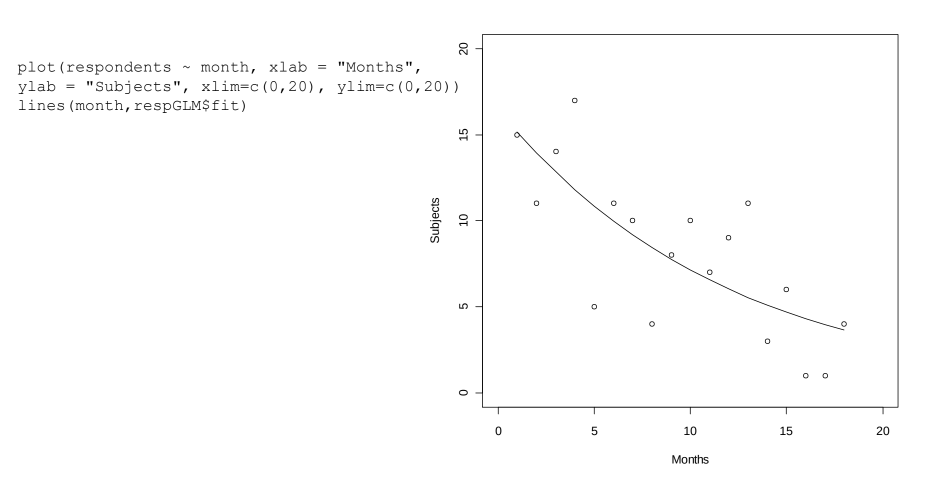
```
> summary(respGLM)
Call:
glm(formula = respondents ~ month, family = poisson, data = stress)
Deviance Residuals:
  Min
         10 Median
                       30
                            Max
-1.9886 -0.9631 0.1737 0.5131 2.0362
Coefficients:
      Estimate Std. Error z value Pr(>|z|)
(Intercept) 2.80316  0.14816  18.920 < 2e-16 ***
         month
Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
(Dispersion parameter for poisson family taken to be 1)
  Null deviance: 50.843 on 17 degrees of freedom
Residual deviance: 24.570 on 16 degrees of freedom
```

AIC: 95.825

Number of Fisher Scoring iterations: 5

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Data and fitted model



Example 2: Ministerial resignation

• On October 18, 1995, `The Independent' reported on the numbers of ministerial resignations because of different reason. The years start in 1945-1951, with a Labour government, and 7 Resignations.

Term	45-51	51-57	55-57	57-63	63-64	64-70	70-74	74-76	76-79	79-90	90-95	97-05
Gov	Lab	con	con	con	con	lab	con	lab	lab	con	con	lab
Res	7	1	2	7	1	5	6	5	4	1	1	1
Year	6	4	2	6	1	6	4	2	3	1	5	8

 Main question: Is there any difference between Government (Labor and Conservative) in the rate of resignations?

Model formulation

The distribution of the response variable

$$Y_t \sim Poisson(\mu_t)$$

$$E(Y_t) = \mu_t$$

The linear predictor

$$\mu_t = f(gov: L/C)$$

$$G_{t} = \begin{cases} 1 & L \\ 0 & C \end{cases}$$

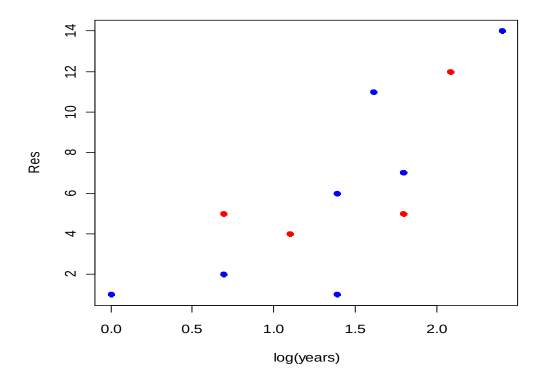
$$\eta = \beta_0 + \beta_1 G_t$$

$$\mu_t = e^{\eta} = e^{\beta_0 + \beta_1 G_t}$$

$$g(E(Y_t)) = \log(\mu_t) = \eta$$

The data in R

```
>Resignations <- read.table("C:..../resign.txt",header=T)
>attach(Resignations)
>plot(Res ~ log(years), pch=19, col=c(4,2)[Gov])
```



Model formulation (1)

> first.glm <- glm(Res ~ Gov , poisson) $\eta = \log(\mu_t) = \beta_0 + \beta_1 G_t$

The relative risk

$$RR = \frac{E(Y_{t} | L)}{E(Y_{t} | C)} = \frac{e^{\beta_{0} + \beta_{1}}}{e^{\beta_{0}}} = e^{\beta_{1}}$$

Model 1 output in R

```
> first.glm <- glm(Res ~ Gov, poisson);</pre>
> summary(first.glm)
Call:
glm(formula = Res ~ Gov, family = poisson)
Deviance Residuals:
                               3Q
             10 Median
   Min
                                       Max
-2.5331 -1.2942 -0.3255 0.7548
                                   2.7793
Coefficients:
           Estimate Std. Error z value Pr(>|z|)
(Intercept) 1.79176 0.15430 11.61
                                       <2e-16 ***
Govlab
            0.09531 0.23262 0.41 0.682
Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
(Dispersion parameter for poisson family taken to be 1)
   Null deviance: 33.436 on 11 degrees of freedom
Residual deviance: 33.269 on 10 degrees of freedom
AIC: 78.459
Number of Fisher Scoring iterations: 5
```

Model formulation (2)

How can we take the number of government years into account?

$$\mu_t = e^{\eta} = e^{\beta_0 + \beta_1 G_t + \beta_2 \log(years_t)}$$

$$\eta = \log(\mu_t) = \beta_0 + \beta_1 G_t + \beta_2 \log(years_t)$$

GLM with Poisson family

> first.glm <- glm(Res ~ Gov + log(years), poisson)</pre>

The same slope for log(year)

$$\eta = \log(\mu_t) = \beta_0 + \beta_1 G_t + \beta_2 \log(years_t)$$

Model 2: Routput

```
glm(formula = Res ~ Gov + log(years), family = poisson)
Deviance Residuals:
             10 Median
   Min
                              30
                                     Max
-2.2417 -0.3469 -0.1250 0.3917 1.6513
Coefficients:
           Estimate Std. Error z value Pr(>|z|)
(Intercept) 0.29973 0.41598 0.721 0.471
Govlab
        0.03541 0.23271 0.152 0.879
log(years) 0.96636 0.22258 4.342 1.41e-05 ***
Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 '' 1
(Dispersion parameter for poisson family taken to be 1)
   Null deviance: 33.436 on 11 degrees of freedom
Residual deviance: 11.276 on 9 degrees of freedom
AIC: 58.466
Number of Fisher Scoring iterations: 4
```

Model 3: model formulation

Different intercepts and slopes

$$\eta = \beta_0 + \beta_1 G_t + \beta_2 \log(years_t) + \beta_3 G_t \log(year_t)$$

Model 3 output in R

```
Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
(Dispersion parameter for poisson family taken to be 1)
    Null deviance: 33.436 on 11 degrees of freedom
Residual deviance: 10.336 on 8 degrees of freedom
AIC: 59.526
Number of Fisher Scoring iterations: 4
```

Model 4: GLM with an offset variable: model formulation

Number of resignation per government year:

$$\frac{Y_t}{year_t} \sim Poisson(\mu_t)$$

$$Y_t \sim Poisson(years_t \times \mu_t)$$

Model 4: GLM with an offset variable - model formulation

$$Y_t \sim Poisson(years_t \times \mu_t)$$

$$E(Y_t) = years_t \times \mu_t = years_t \times e^{\beta_0 + \beta_1 G_t}$$

$$g(E(Y_t)) = g(years_t \times \mu_t) = \log(years_t \times e^{\beta_0 + \beta_1 G_t})$$

$$g(E(Y_t)) = \log(years_t) + \beta_0 + \beta_1 G_t = \eta$$

A model with offset in R

> next.glm<- glm(Res ~ Gov + offset(log(years)), poisson) $\beta_0 + \beta_1 G_t + \log(years_t)$

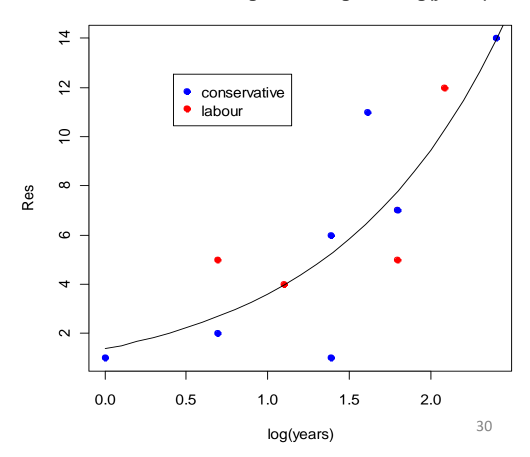
Model 4 GLM with offset output in R

```
summary(first.qlm4)
       Call:
       glm(formula = Res ~ Gov + offset(log(years)), family = poisson)
\hat{eta}_0 Estimate Std. Error z value Pr(>|z|) (Intercept) 0.24116 0.15430 1.563 0.118 \hat{eta}_1 Govlab 0.03647 0.23262 0.157 0.875
       (Dispersion parameter for poisson family taken to be 1)
            Null deviance: 11.323 on 11 degrees of freedom
       Residual deviance: 11.299 on 10 degrees of freedom
       AIC: 56.488
       Number of Fisher Scoring iterations: 4
```

Data and predicted models

```
plot(Res ~ log(years), pch=19, col=c(4,2)[Gov]) # Use palette() to find out which colour corresponds 
> legend(locator(1), legend= c("conservative", "labour"), col=c(4,2), pch=19) 
> 1 <- (0:25)/10 
> fv <- exp(0.3168 + 0.9654*1)# to plot fitted curve under last.glm 
> lines(1,fv)
```

Ministerial Resignations against log(years)



AIC for the different models

MODEL	No parameters	Deviance	AIC
1	2	33.269	78.45861
2	3	11.276	58.46574
3	4	10.336	59.52603
4	2	11.299	56.48846

$$g(E(Y_t)) = \log(years_t) + \beta_0 + \beta_1 G_t$$

Confidence interval for β_2 for model 2

ANOVA for model 3 and model 2

Model selection (II): the step() function in R

```
> step(first_qlm, direction = "backward")
                                                    AIC=56.49
                                             Step:
Start: AIC=59.53
                                             Res ~ log(years)
Res ~ log(years) + Gov + Gov:log(years)
                                                          Df Deviance
                                                                         ATC
                Df Deviance
                                                               11.299 56.489
                                ATC
                                             <none>
                                             -log(years) 1 33.436 76.626
- log(years):Gov 1
                     11.276 58.466
                                             -Call: glm(formula = Res ~
                     10.336 59.526
<none>
                                             log(years), family = poisson)
Step:
      AIC=58.47
                                             Coefficients:
Res ~ log(years) + Gov
                                             (Intercept) log(years)
                                                  0.3168
                                                               0.9654
             Df Deviance
                           AIC
                                             Degrees of Freedom: 11 Total (i.e.
                 11.299 56.489
- Gov
              1
                                             Null); 10 Residual
                 11,276 58,466
<none>
                                             Null Deviance:
                                                                 33.44
   log(years) 1 33.269 78.459
                                             Residual Deviance: 11.3
                                             ATC: 56.49
```

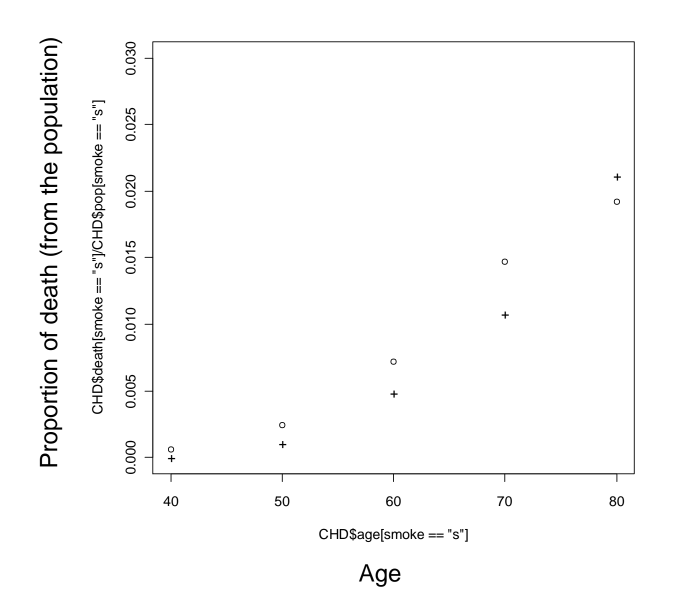
Model 4 with the offset variable is not included here.

Example 3: smoking and coronary death

CHD			
age	smoke	death	pop
40	S	32	52407
50	S	104	43248
60	S	206	28612
70	S	186	12663
80	S	102	5317
40	ns	2	18790
50	ns	12	10673
60	ns	28	5710
70	ns	28	2585
0 8 0	ns	31	1462
	age 40 50 60 70 80 40 50 60 70	age smoke 40 s 50 s 60 s 70 s 80 s 40 ns 50 ns 60 ns 70 ns	age smoke death 40 s 32 50 s 104 60 s 206 70 s 186 80 s 102 40 ns 2 50 ns 12 60 ns 28 70 ns 28

A study about tobacco consumption and coronary heart disease among British doctors.

Smoking and coronary death: the data



GLM with an offset variable: model formulation

Number of deaths per population size:

$$\frac{Y_i}{n_i} \sim Poisson(\mu_i)$$

$$Y_i \sim Poisson(n_i \times \mu_i)$$

$$g(\mu_i) = X\beta$$

Smoking and coronary death

- Is the death rate higher for smokers than non smokers?
- If so, by how much?
- Is there differential effects of age?

GLM with an offset variable: model formulation

Number of deaths per population size:

$$\frac{Y_i}{n_i} \sim Poisson(\mu_i)$$

 $Y_i \sim Poisson(n_i \times \mu_i)$

Linear predictor:

$$\eta = f(age, smoking, pop.)$$

3 models

$$M_{1}: \eta = \log(pop.) + \beta_{1}age + \beta_{2}smoke$$

$$M_{2}: \eta = \log(pop.) + \beta_{1}age + \beta_{2}smoke + \beta_{3}age \times smoke$$

$$M_{3}: \eta = \log(pop.) + \beta_{11}age + \beta_{12}age^{2} + \beta_{2}smoke + \beta_{3}age \times smoke$$

```
>fit.chd1<-glm(death ~ age + smoke+offset(log(pop)), poisson)
>fit.chd2<-glm(death ~ age + smoke+age:smoke+offset(log(pop)), poisson)
>age2<-age^2
>fit.chd3<-glm(death ~ age+age2+smoke+age:smoke+offset(log(pop)), poisson)</pre>
```

Model selection

```
> extractAIC(fit.chd1, k=2)
[1]   3.0000 130.2500
> extractAIC(fit.chd2, k=2)
[1]   4.0000 122.9614
> extractAIC(fit.chd3, k=2)
[1]   5.00000 66.70331
```

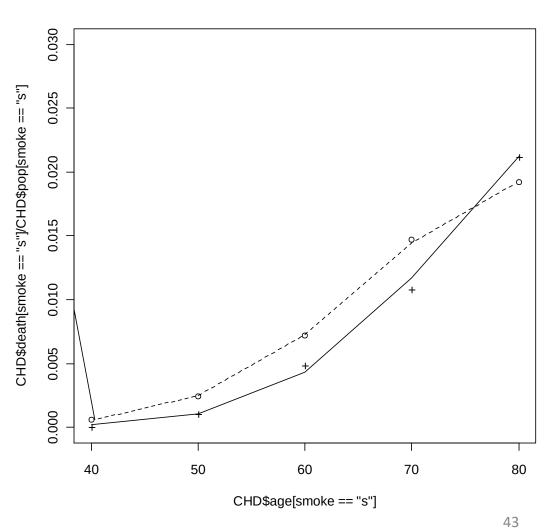
The model with quadratic age effect has the best goodness-of-fit.

R output (model 3)

```
> summary(fit.chd3)
Call:
qlm(formula = death ~ age + age2 + smoke + age:smoke + offset(log(pop)),
   family = poisson)
Coefficients:
             Estimate Std. Error z value Pr(>|z|)
(Intercept) -1.970e+01 1.253e+00 -15.717 < 2e-16 ***
        3.563e-01 3.632e-02 9.810 < 2e-16 ***
age
age2 -1.977e-03 2.737e-04 -7.223 5.08e-13 ***
smokes 2.364e+00 6.562e-01 3.602 0.000316 ***
age:smokes -3.075e-02 9.704e-03 -3.169 0.001528 **
               0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
Signif. codes:
(Dispersion parameter for poisson family taken to be 1)
   Null deviance: 935.0673 on 9 degrees of freedom
Residual deviance: 1.6354 on 5 degrees of freedom
AIC: 66.703
Number of Fisher Scoring iterations: 4
```

Data and predicted model

$$\hat{E}(Y_i) = n_i \times \exp(\mu_i) = n_i \times \exp(X\hat{\beta})$$



Chapter 12:

Beyond Poisson and binomial distributions: Models with different link functions and/or distributions

Lindsey: Chapter 4.

Example 1: Employment duration

• The employment duration of staff, age 25 to 44, recruited to the British post office in the first quarter of 1973 and classified in to two grades.

Month	Grade 1	Grade2	Month	Grade1	Grade2
1	22	30	13	0	1
2	18	28	14	0	0
3	19	31	15	0	0
4	13	14	16	1	1
5	5	10	17	1	1
6	6	6	18	1	0
7	3	5	19	3	2
8	2	2	20	1	0
9	2	3	21	1	3
10	1	0	22	0	1
11	0	0	23	0	1
12	1	1	24	0	0

The data in R

```
> employ <- read.table("C:.... /employ.txt",header=TRUE)</pre>
> attach(employ)
> employ
                                                                    log
                                          Original sacle.
    month grade duration
                                                                    transformation.
                  1
                              22
                              30
3
                              18
                                       33
                              28
                                       8
                                                                  2.0
5
                              19
                                      5
                                                                  1.5
6
          3
                              31
                                       9
                                                                  0:
                              13
                                                                  0:0
                                                     000 0000
8
                              14
                                                       20
                                                                           10
                                                                               15
                                                                                  20
9
                                                month
                                                                           month
10
                              10
```

Model formulation

Models with normal error.

$$Y_t \sim N(\mu_t, \sigma^2)$$

$$\eta = \beta_0 + \beta_1 t + \beta_2 G$$

$$\mu = \beta_0 + \beta_1 t + \beta_2 G$$

$$\log(\mu) = \beta_0 + \beta_1 t + \beta_2 G$$

$$\frac{1}{\mu} = \beta_0 + \beta_1 t + \beta_2 G$$

$$\mu = e^{\beta_0 + \beta_1 t + \beta_2 G}$$

$$\mu = \frac{1}{\beta_0 + \beta_1 t + \beta_2 G}$$

identity link

Log link

invesre link

Models with normal error in R

```
m.normal.idt <- glm(duration ~ month + grade + month:grade,data = employ,
family = gaussian(link = identity))

m.normal.inv <- glm(duration ~ month + grade + month:grade,data = employ,
family = gaussian(link = inverse))

m.normal.log <- glm(duration ~ month + grade + month:grade,data = employ,
family = gaussian(link = log))

m.normal.log1 <- glm(duration ~ month + grade ,data = employ,
family = gaussian(link = log))</pre>
```

Model selection

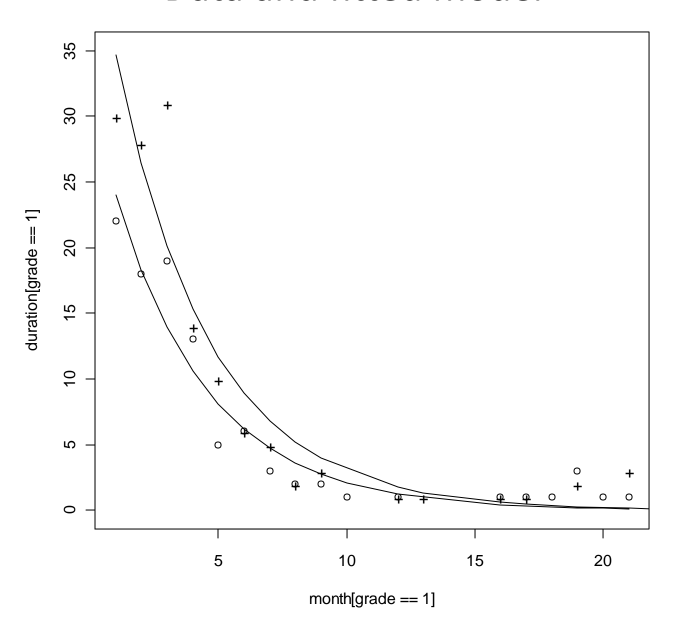
```
> extractAIC(m.normal.idt, k=2)
[1]    4.0000 228.5094
> extractAIC(m.normal.inv, k=2)
[1]    4.0000 198.3163
> extractAIC(m.normal.log, k=2)
[1]    4.0000 173.8542
> extractAIC(m.normal.log1, k=2)
[1]    3.0000 171.8545
```

The model with log link has the smallest AIC value.

GLM with normal error and log link in R

```
> m.normal.log <- glm(duration ~ month + grade + month:grade,data = employ1, family =
qaussian(link = loq))
> summary(m.normal.log)
Call:
glm(formula = duration ~ month + grade + month:grade, family = gaussian(link = log),
   data = employ1)
Deviance Residuals:
   Min
             10 Median
                              30
                                      Max
-4.6676 -1.5332 -0.2005 0.8915 10.8676
Coefficients:
              Estimate Std. Error t value Pr(>|t|)
(Intercept) 3.4476971 0.1304802 26.423 < 2e-16 ***
           -0.2709521 0.0444746 -6.092 1.08e-06 ***
month
grade2 0.3698490 0.1591805 2.323 0.0271 *
month:grade2 -0.0007872 0.0545143 -0.014 0.9886
Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 '' 1
(Dispersion parameter for gaussian family taken to be 8.219982)
   Null deviance: 2771.88 on 33 degrees of freedom
Residual deviance: 246.59 on 30 degrees of freedom
ATC: 173.85
Number of Fisher Scoring iterations: 7
```

Data and fitted model



Other models

- Different distributions:
 - Normal.
 - Gamma.
 - ...

$$Y_t \sim H(\mu_t)$$

$$g(\mu_t) = \eta = \beta_0 + \beta_1 t + \beta_2 G$$

$$\mu = \beta_0 + \beta_1 t + \beta_2 G$$

$$\log(\mu) = \beta_0 + \beta_1 t + \beta_2 G$$

$$\frac{1}{\mu} = \beta_0 + \beta_1 t + \beta_2 G$$

$$\mu = e^{\beta_0 + \beta_1 t + \beta_2 G}$$

$$\mu = \frac{1}{\beta_0 + \beta_1 t + \beta_2 G}$$

identity link

Log link

invesre link

Model formulation (model 1)

m.normal.log <- glm(duration ~ month + grade + month:grade,
data = employ1, family = gaussian(link = log))</pre>

$$\eta = \log(\mu_i) = \beta_0 + \beta_2 t_i + \beta_1 G_i + \beta_3 t_i \times G_i$$

A model with normal error and log link:

$$Y_i \sim N(\mu_i, \sigma^2)$$
$$g(\mu) = \eta, \log(\mu) = \eta$$

R output: model 1

```
> m.normal.log <- glm(duration ~ month + grade + month:grade, data = employ, family =
   gaussian(link = log))
> summary(m.normal.log)
Call:
glm(formula = duration ~ month + grade + month:grade, family = gaussian(link = log),
   data = employ)
Deviance Residuals:
             10 Median 30
   Min
                                     Max
-4.6676 -1.5332 -0.2005 0.8915 10.8676
Coefficients:
             Estimate Std. Error t value Pr(>|t|)
(Intercept) 3.0778481 0.2764302 11.134 3.54e-12 ***
           -0.2701650 0.0943705 -2.863 0.00759 **
month
grade 0.3698490 0.1591805 2.323 0.02712 *
month:grade -0.0007872 0.0545143 -0.014 0.98857
Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 '' 1
(Dispersion parameter for gaussian family taken to be 8.219982)
   Null deviance: 2771.88 on 33 degrees of freedom
Residual deviance: 246.59 on 30 degrees of freedom
ATC: 173.85
```

Model with gamma error and inverse link function: model formulation (model 2)

$$\eta = (1/\mu_i) = (\beta_0 + \beta_2 t_i + \beta_1 G_i + \beta_3 t \times G_i)$$

A model with gamma error and inverse:

$$Y_i \sim Gamma(\mu_i)$$

$$\frac{1}{\mu_i} = \eta_i$$

R output: model 2

```
> m.gamma.inv <- glm(duration ~ month + grade + month:grade, data = employ, family =
   Gamma(link = inverse))
> summary(m.gamma.inv )
Call:
glm(formula = duration ~ month + grade + month:grade, family = Gamma(link = inverse),
    data = employ)
Coefficients:
             Estimate Std. Error t value Pr(>|t|)
(Intercept) -0.0122937 0.0396403 -0.310 0.75861
        0.0492390 0.0154143 3.194 0.00329 **
month
           -0.0008403 0.0226236 -0.037 0.97062
grade
month:grade -0.0089011 0.0090337 -0.985 0.33235
Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 '' 1
(Dispersion parameter for Gamma family taken to be 0.3661260)
   Null deviance: 51.752 on 33 degrees of freedom
Residual deviance: 9.887 on 30 degrees of freedom
AIC: 147.27
```

Model with gamma error and inverse link function: model formulation (model 3)

Grade is not included in the model:

$$\eta = 1/\mu_i = \beta_0 + \beta_2 t_i$$

A model with gamma error and inverse:

$$Y_i \sim Gamma(\mu_i)$$

$$\frac{1}{\mu_i} = \eta_i$$

R output: model 3

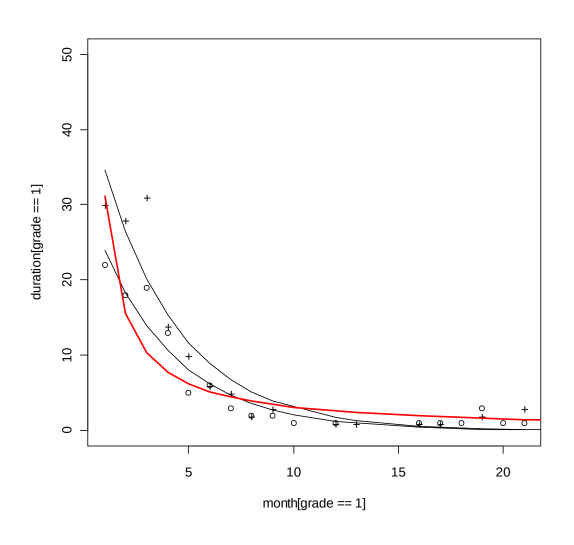
```
> m.gamma.inv1 <- glm(duration ~ month , data = employ, family = Gamma(link = inverse))</pre>
> summary(m.gamma.inv1)
Call:
qlm(formula = duration ~ month, family = Gamma(link = inverse),
   data = employ)
Deviance Residuals:
            10 Median 30
   Min
                                      Max
-0.9163 \quad -0.5284 \quad -0.2795 \quad 0.2599 \quad 1.2685
Coefficients:
            Estimate Std. Error t value Pr(>|t|)
(Intercept) -0.014033 0.010821 -1.297 0.204
        month
Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 '' 1
(Dispersion parameter for Gamma family taken to be 0.3762360)
   Null deviance: 51.752 on 33 degrees of freedom
Residual deviance: 10.562 on 32 degrees of freedom
AIC: 145.63
Number of Fisher Scoring iterations: 6
```

Model selection

```
> AIC(m.normal.log)
[1] 173.8542
> AIC(m.gamma.inv)
[1] 147.2748
> AIC(m.gamma.inv1)
[1] 145.6314
> AIC(m.gamma.inv2)
[1] 145.3437
```

The model which give small AIC is the gamma model with inverse line and only duration in the model

Data and fitted models



Chapter 13: Poisson regression and log linear models

Based on Dobson: Chapter 9

Log linear models

The general frame work of Poisson regression is given by:

$$Y_i \sim Poisson(\mu_i)$$

$$g(\mu_i) = \log(\mu_i) = X\beta$$

Example 1: melanoma

	Site			
Tumor Type	Head & neck	Trunk	Extere mities	total
Hutchinson	22	2	10	34
Superficial melanoma	16	54	115	185
Nodular	19	33	73	125
Indeterminate	11	17	28	56
Total	68	106	226	400

Cross sectional study of patients with form of skin cancer.

Different sites & different tumor types.

Example 1

	Site			
Tumor Type	Head & neck	Trunk	Extere mities	total
Hutchinson	22	2	10	34
Superficial melanoma	16	54	115	185
Nodular	19	33	73	125
Indeterminate	11	17	28	56
Total	68	106	226	400

$$Y_{ij}$$
 Frequency of subjects with tumor type i and site j.

$$Y_{ij} \sim Poisson(\mu_{ij})$$

 \mathcal{H} Sample size.

Example 1: independence?

	Site			
Tumor Type	Head & neck	Trunk	Extere mities	total
Hutchinson	22	2	10	34
Superficial melanoma	16	54	115	185
Nodular	19	33	73	125
Indeterminate	11	17	28	56
Total	68	106	226	400

$$Y_{.j}$$

Chi-squared statistic for independence:

$$X^{2} = \frac{\sum_{ij} (Y_{ij} - e_{ij})^{2}}{e_{ij}}$$

$$e_{ij} = \frac{Y_{i.} \times Y_{.j}}{n}$$

$$e_{11} = \frac{34 \times 68}{400}$$

Example 1: independence?

$$X^{2} = \frac{\sum_{ij} (Y_{ij} - e_{ij})^{2}}{e_{ij}} = \frac{(22 - 5.78)^{2}}{5.78} + \dots + \frac{(28 - 31.64)^{2}}{31.64} = 65.8$$

Model formulation

$$Y_{ij} \sim Poisson(\mu_{ij})$$

$$g(\mu_{ij}) = \mu$$

$$g(\mu_{ij}) = \mu + \alpha_i + \beta_j$$

$$g(\mu_{ij}) = \mu \qquad g(\mu_{ij}) = \mu + \alpha_i + \beta_j \qquad g(\mu_{ij}) = \mu + \alpha_i + \beta_j + \alpha \beta_{ij}$$

The minimal model

Independence model

M1

M2

Second ordered interaction model

Data in R

```
> melanoma
   Tumor
           Type
                22
      Hu
             HN
      Hu
          Trunk
3
     Hu Extrem 10
4
                 16
             HN
      Su
5
      Su Trunk
                45
6
      Su Extrem 115
    Nod
         HN
                19
8
                33
     Nod Trunk
9
                73
     Nod Extrem
10
                 11
     Ind
             HN
11
     Ind Trunk
                17
12
                28
     Ind Extrem
```

Models in R

```
> M1<-glm(y~1, family=poisson, data=melanoma)
             g(\mu_{ii}) = \mu
> M2<-glm(y~Tumor+Type, family=poisson, data=melanoma)
             g(\mu_{ii}) = \mu + \alpha_i + \beta_i
> M3<-glm(y~Tumor+Type+Tumor:Type, family=poisson,
             data=melanoma)
             g(\mu_{ii}) = \mu + \alpha_i + \beta_i + \alpha \beta_{ii}
> AIC (M1)
[1] 348.8361
> AIC (M2)
[1] 121.5482
> AIC(M3)
[1] 82.9297
```

Likelihood ratio test

Example 2: Death penalty

Death penalty verdicts for cases involving multiple murders in Florida between 1976 and 1987. This data is from an article that studied effects of racial characteristics on whether persons convicted of homicide received the death penalty. The 674 subjects classified is in to a 2x2x2 contingency table-two rows, two columns, and two layers.

		Victim's Race					
			White	Black			
		Defendant's Race		Defendant's Race			
		White	Black	White	Black		
Death	Yes	53	11	0	4		
Penalty	No	414	37	16	139		
Percent Yes		11.3	22.9	0.0	2.8		

Example 3: Death penalty

		Victim's Race				
			White	Black		
		Defer	ndant's Race	Defendant's Race		
		White	Black	White	Black	
Death	Yes	53	11	0	4	
Penalty	No	414	37	16	139	
Percent Yes		11.3	22.9	0.0	2.8	

The response variable: death penalty verdicts

$$Y_{ijk} \sim Poisson(\mu_{ijk})$$

Example 3: Death penalty

		Victim's Race				
			White	Black		
		Defe	ndant's Race	Defendant's Race		
		White	Black	White	Black	
Death Penalty	Yes	53	11	0	4	
	No	414	37	16	139	

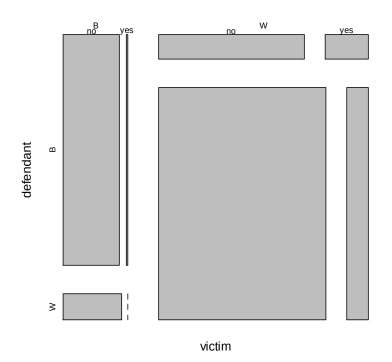
The mean structure

$$log(\mu_{ijk}) = \mu + v_i + d_j + p_k + interactio n$$

$$\log(\mu_{ijk}) = \mu + v_i + d_j + p_k + vd_{ij} + vp_{ik} + dp_{jk} + vdp_{ijk}$$

Exploring the Data in R

xtabs(number ~ victim + defendant + death, deathpenalty)



The model in R

$$\log(\mu_{ijk}) = \mu + v_i + d_j + p_k + vd_{ij} + vp_{ik} + dp_{jk} + vdp_{ijk}$$

Saturated model with Poisson family

```
> M1<-glm(number~victim*defendant*death, family=poisson, data=deathpenalty)
> summary(M1)
> Call:
Estimate Std. Error z value Pr(>|z|)
(Intercept)
                           4.934e+00 8.482e-02 58.177 < 2e-16 ***
victimW
                        -1.324e+00 1.850e-01 -7.155 8.38e-13 ***
                          -2.162e+00 2.640e-01 -8.189 2.63e-16 ***
defendantW
                         -3.548e+00 5.071e-01 -6.996 2.63e-12 ***
deathyes
victimW:defendantW
                         4.577e+00 3.149e-01 14.536 < 2e-16 ***
                          2.335e+00 6.125e-01 3.813 0.000137 ***
victimW:deathyes
defendantW:deathyes
                    -2.153e+01 4.225e+04 -0.001 0.999593
victimW:defendantW:deathyes 2.068e+01 4.225e+04 0.00049 0.999609
Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 '' 1
(Dispersion parameter for poisson family taken to be 1)
   Null deviance: 1.2251e+03 on 7 degrees of freedom
Residual deviance: 4.1224e-10 on 0
                                   degrees of freedom
AIC: 54.04
Number of Fisher Scoring iterations: 20
```

Using the step function in R to identify the best model

step (the three way interaction model)

- A stepwise selection based on AIC.
- Starting point: the three way interaction model.

Step () function for model selection

```
> step(M1)
                                                      Starting
Start: AIC=54.04
                                                      point (see
number ~ victim * defendant * death
                                                      also slide 79).
                          Df Deviance
                                        ATC
- victim:defendant:death 1 0.37984 52.42
                              0.00000 54.04
                                                      Three was
<none>
                                                      interaction is
Step: AIC=52.42
                                                      excluded.
number ~ victim + defendant + death +
victim:defendant + victim:death +
    defendant:death
                    Df Deviance
                                   ATC
<none>
                           0.38
                                52.42
                                                      Two way
- defendant:death 1 5.39 55.43
                                                      interaction
- victim:death 1 20.73
                                 70.77
                                                      are excluded
- victim:defendant
                         384.43
                                434.47
                                                      (one at the
                                                      time).
```

The best model includes all two way interactions:

$$\log(\mu_{ijk}) = \mu + v_i + d_j + p_k + vd_{ij} + vp_{ik} + dp_{jk}$$

Output of the Step() function

```
Call: qlm(formula = number ~ victim + defendant + death + victim:defendant +
                         victim:death + defendant:death,
                         family = poisson, data = deathpenalty)
Coefficients:
                                     victimW
                                                          defendant.W
         (Intercept)
                                     -1.3298
              4.9358
                                                             -2.1746
            deathyes
                       victimW:defendantW
                                                   victimW:deathyes
             -3.5961
                                      4.5950
                                                              2.4044
defendantW:deathyes
             -0.8678
Degrees of Freedom: 7 Total (i.e. Null); 1 Residual
Null Deviance:
                      1225
                                    AIC: 52.42
Residual Deviance: 0.3798
```

The two way interaction model in R

$$\log(\mu_{ijk}) = \mu + v_i + d_j + p_k + vd_{ij} + vp_{ik} + dp_{jk}$$

$$\text{M2} < -\text{glm (number} \sim \text{victim+defendant+death} \\ + \text{victim:defendant} \\ + \text{victim:death} \\ + \text{defendant:death,}$$

family=poisson, data=deathpenalty)

Output in R

```
> summary(M2)
Call:
qlm(formula = number ~ victim + defendant + death + victim:defendant +
   victim:death + defendant:death, family = poisson, data = deathpenalty)
Coefficients:
                   Estimate Std. Error z value Pr(>|z|)
                               0.08471 58.265 < 2e-16 ***
(Intercept)
                    4.93578
victimW
                   -1.32980
                               0.18479 -7.196 6.19e-13 ***
defendant.W
                   -2.17465
                               0.26377 -8.245 < 2e-16 ***
                   -3.59610
deathves
                               0.50691 -7.094 1.30e-12 ***
                    4.59497
                               0.31353 14.656 < 2e-16 ***
victimW:defendantW
victimW:deathyes
                    2.40444
                               0.60061 4.003 6.25e-05 ***
defendantW:deathyes -0.86780
                               0.36707 -2.364 0.0181 *
Signif. codes:
               0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
(Dispersion parameter for poisson family taken to be 1)
   Null deviance: 1225.07955 on 7 degrees of freedom
Residual deviance:
                     0.37984 on 1 degrees of freedom
AIC: 52.42
```

Example 3: Antibiotic prescription

Diagnosis of Respiratory Tract Infections Hueston and Stott (2000) report a study of clinicians' diagnoses of respiratory tract infections over a 14-month period. The aim was to determine whether a reduction in prescription of antibiotics to acute bronchitis patients was due to clinicians assigning an alternative diagnosis.

	Time period					
Diagnosis	1-3/96				1-2/97	
Acute bronchitis	113	58	40	108	100	
Acute sinusitis	99	37	23	50	32	
URI	410	228	125	366	304	
Pneumonia	60	43	30	56	45	
Total	682	366	218	580	481	

The data in R

Example 3: Antibiotic prescription

	Time period					
Diagnosis	1-3/96	4-6/96	7-9/96	10-12/96	1-2/97	
Acute bronchitis	113	58	40	108	100	
Acute sinusitis	99	37	23	50	32	
URI	410	228	125	366	304	
Pneumonia	60	43	30	56	45	

Research question: Diagnostic and time period are independent?

Data in R

```
> rt
   diag time count
1 bron win96
                113
2 sinus win96
                 99
3 URI win96
                410
4 pneu win96
                60
5 bron spr96
                58
6 sinus spr96
                37
   URI spr96
                228
  pneu spr96
                 43
   bron sum96
                40
10 sinus sum96
                2.3
                125
11 URI sum96
12 pneu sum96
                 30
  bron aut96
                108
14 sinus aut96
                 50
   URI aut96
                366
  pneu aut96
                56
   bron spr97
                100
18 sinus spr97
                 32
19
    URI spr97
                304
   pneu spr97
20
                4.5
```

Models formulation

$$Y_{ij} \sim Poisson(\mu_{ij})$$

$$g(\mu_{ij}) = \mu$$

$$g(\mu_{ij}) = \mu + D_i + T_j$$

 $g(\mu_{ij}) = \mu$ $g(\mu_{ij}) = \mu + D_i + T_j$ $g(\mu_{ij}) = \mu + D_i + T_j + DT_{ij}$

Minimal model

Independence model

Second ordered interaction

Models in R

```
> M1 <- glm(count ~ 1, family=poisson, data=rt)
> M2 <- glm(count ~ diag+time, family=poisson, data=rt)
> M3 <- glm(count ~ diag+time+diag:time,family=poisson, data=rt)
>
> AIC(M1)
[1] 1915.304
> AIC(M2)
[1] 169.8704
> AIC(M3)
[1] 164.2791
```

Likelihood ratio test

Independence model is rejected.

Output model 3

> summary(M3) Coefficients: Estimate Std. Error z value Pr(>|z|)0.09407 50.253 < 2e-16 *** 4.72739 (Intercept) diagsinus -0.13227 0.13766 -0.961 0.33664 diagURI 1.28877 0.10625 12.130 < 2e-16 *** diagpneu -0.63304 0.15974 -3.963 7.40e-05 *** timespr96 -0.66694 0.16153 -4.129 3.64e-05 *** timesum96 -1.03851 0.18398 -5.645 1.66e-08 *** timeaut96 -0.04526 0.13457 -0.336 0.73664 timespr97 -0.12222 0.13729 -0.890 0.37336 diagsinus:timespr96 -0.31726 0.25143 -1.262 0.20702 0.442 0.65872 diagURI:timespr96 0.08013 0.18143 diagpneu:timespr96 1.299 0.19388 0.33380 0.25693 diagsinus:timesum96 -0.42112 0.29568 - 1.424 0.15438diagURI:timesum96 -0.14934 0.21045 -0.710 0.47795 diagpneu:timesum96 0.34536 0.28957 1.193 0.23300 diagsinus:timeaut96 -0.63784 0.21957 - 2.905 0.00367 **diagURI:timeaut96 -0.06827 0.15258 -0.447 0.65457 diagpneu:timeaut96 -0.02374 0.22942 -0.103 0.91760 diagsinus:timespr97 -1.00717 0.24536 -4.105 4.05e-05 *** diagURI:timespr97 -0.17691 0.15677 -1.128 0.25913 diagpneu:timespr97 -0.16546 0.24029 -0.689 0.49107 Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1 (Dispersion parameter for poisson family taken to be 1) Null deviance: 1.7890e+03 on 19 degrees of freedom Residual deviance: 1.5765e-14 on 0 degrees of freedom AIC: 164.28 Number of Fisher Scoring iterations: 3

Chapter 14 Over dispersion

Lindsey: Chapter 3

Over dispersion parameter

The general form of exponential family is defined as:

$$f(y) = \exp\left\{\frac{y_i \theta_i - b(\theta_i)}{a_i(\phi)} + c(y_i, \phi)\right\}$$

- Where Ø is the dispersion parameter.
- $a(\emptyset)$: scale parameter.

Example: normal distribution

$$f(y_i; \mu_i, \sigma^2) = \frac{1}{\sqrt{2\pi\sigma^2}} e^{\frac{-(y_i - \mu_i)^2}{2\sigma^2}}$$
$$= \exp\left\{ \left[y_i \mu_i - \frac{\mu_i^2}{2} \right] \frac{1}{\sigma^2} - \frac{y_i^2}{2\sigma^2} - \frac{1}{2} \log(2\pi\sigma^2) \right\}$$



The mean and the variance are separated parameters.

$$\theta_{i} = \mu_{i},$$

$$b(\theta_{i}) = \theta_{i}^{2} / 2$$

$$a_{i}(\phi) = \sigma^{2}$$

$$c(y_{i}, \phi) = -\left[y_{i}^{2} / \phi + \log(2\pi\phi)\right] / 2.$$

Example: Binomial distribution

$$Z_{i} = \begin{cases} 1 \\ 0 \end{cases} \qquad Y_{i} = \sum_{i=1}^{n} Z_{i} \qquad Y_{i} \sim B(n, \pi_{i})$$

$$p(y_{i} \mid \theta) = \binom{n_{i}}{y_{i}} \theta^{y_{i}} (1 - \theta)^{n - y} =$$

$$\exp \left\{ y_{i} \log \left[\frac{\theta_{i}}{1 - \theta_{i}} \right] + n_{i} \log(1 - \theta_{i}) + \log \binom{n_{i}}{y_{i}} \right\}$$

The variance is a function of the mean.

$$a_{i}(\phi) = 1, \quad b(\theta_{i}) = \log(1 + \exp(\theta_{i}))$$

$$c(y) = \log \binom{n_{i}}{y_{i}}$$

$$E(y) = \mu = b'(\theta_{i}) = e^{\theta} (1 + \exp(\theta_{i}))^{-1}$$

$$\text{var}(y) = n\mu(1 - \mu)$$
95

Poisson distribution

$$Y_i \sim Poisson(\mu)$$

$$f(y_i, \theta_i) \frac{\theta_i^{y_i} e^{-\theta}}{y_i!}$$

$$E(y)=V(y)$$
.

$$a_{i}(\phi) = 1$$

$$b(\theta) = \exp(\theta)$$

$$c(y) = -\log(y!)$$

$$E(y) = \mu = b'(\theta) = \exp(\theta)$$

$$\operatorname{var}(y) = \mu$$

Overdispersion

The binomial and Poisson distribution are a members of one parameter exponential family.

$$Y_t \sim Poisson(\mu_t)$$

$$E(Y_t) = V(Y_t) = \mu_t$$

$$Y_i \sim B(n,\pi)$$

$$E(Y_i) = n\pi$$

$$V(Y_i) = n\pi(1-\pi)$$

$$a_i(\phi) = 1$$

Overdispersion

Often, we observed extra variability than expected:

$$V(Y_i) > E(Y_i)$$

$$V(Y_i) > n\pi(1-\pi)$$

$$V(Y_i) = \phi E(Y_i)$$

$$V(Y_i) = \phi n \pi (1 - \pi)$$

$$\phi = 1$$
 No problem with overdispersion

Estimating over dispersion

Formula

$$\phi = \frac{\chi_p^2}{n - p}$$

where

p = number of parametr in the modeln = number of observations

Example 1: Germination of seeds from Orobanche

o. aegyptiaco 75				o. aegyptiaco 73			
bean		cucumber		bean		cucumber	
germ.	total	germ.	total	germ.	total	germ.	total
10	39	5	6	8	16	3	12
23	62	53	74	10	30	22	41
23	<u>81</u>	55	72	8	28	15	30
26	51	32	51	23	45	32	51
17	39	46	79	0	<u>4</u>	3	7
		10	13				

Data in R

```
> fac<-read.table("C:..../seed.txt", header = TRUE)</pre>
> fact <- transform(fac, prop = germ/total, extr.seed = interaction(extract, seed))</pre>
            fact
                    extract germ total
                                                       extr.seed
               seed
                                             prop
                                     39 0.2564103
             aeg75
                       bean
                               10
                                                     bean.aeg75
                                  62 0.3709677
           2
             aeq75
                      bean
                               23
                                                    bean.aeq75
                                                    bean.aeg75
           3
             aeg75
                       bean
                                    81 0.2839506
             aeg75
                      bean
                                    51 0.5098039
                                                     bean.aeg75
                               17
                                    39 0.4358974
             aeg75
                                                     bean.aeg75
                       bean
             aeq75 cucumber
                                    6 0.8333333 cucumber.aeq75
             aeg75 cucumber
                                    74 0.7162162 cucumber.aeg75
             aeq75 cucumber
                               55
                                    72 0.7638889 cucumber.aeq75
             aeq75 cucumber
                               32
                                     51 0.6274510 cucumber.aeq75
          10 aeg75 cucumber
                               46
                                     79 0.5822785 cucumber.aeq75
          11 aeg75 cucumber
                               10
                                    13 0.7692308 cucumber.aeq75
                                    16 0.5000000
          12 aeq73
                       bean
                                                     bean.aeq73
          13 aeg73
                               10
                                    30 0.3333333
                                                   bean.aeq73
                     bean
                                    28 0.2857143
                                                    bean.aeg73
          14 aeq73
                      bean
          15 aeg73
                       bean
                               23
                                     45 0.5111111
                                                    bean.aeg73
          16 aeg73
                                     4 0.0000000
                                                     bean.aeq73
                       bean
          17 aeg73 cucumber
                                    12 0.2500000 cucumber.aeg73
          18 aeg73 cucumber
                                    41 0.5365854 cucumber.aeg73
          19 aeg73 cucumber
                               15
                                     30 0.5000000 cucumber.aeg73
          20 aeg73 cucumber
                               32
                                     51 0.6274510 cucumber.aeg73
```

7 0.4285714 cucumber.aeg73

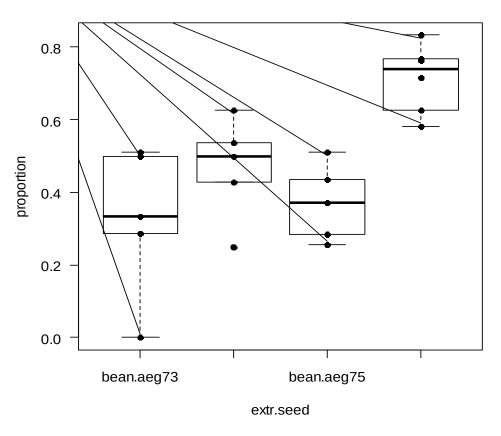
21 aeg73 cucumber

3

Exploring the data: Box-Plot

```
> plot(prop ~ extr.seed, data = fact, las = 1, ylab = "proportion")
> points(prop ~ extr.seed, data = fact, pch = 16)
> title("proportion of germinating seeds")
```

proportion of germinating seeds



Model 1 formulation

Binomial model

$$yi \sim Bin(n_i, \pi_i)$$

 $g(\pi_i) = \eta, \ \log it(\pi_i) = \eta$
 $\log it(\pi_i) = \beta_0 + \beta_1 ext + \beta_2 seed + \beta_3 ext \times seed$

Model 1: binomial family

```
> g <- glm(cbind(germ, total - germ) ~ extract + seed +
    extract:seed, family = binomial, data = fact)
> r.pears<-residuals(g, type="pearson")
> summary(g)
```

The assumption here is over dispersion parameter is $\Phi=1$

Model 1 output

```
Call:
glm(formula = cbind(germ, total - germ) ~ extract + seed + extract:seed,
   family = binomial, data = fact)
Deviance Residuals:
    Min 10 Median 30
                                         Max
-2.01617 \quad -1.24398 \quad 0.05995 \quad 0.84695 \quad 2.12123
Coefficients:
                       Estimate Std. Error z value Pr(>|z|)
(Intercept)
                        -0.4122 0.1842 -2.238 0.0252 *
                        extractcucumber
                         -0.1459 0.2232 -0.654 0.5132
seedaeq75
extractcucumber:seedaeg75  0.7781  0.3064  2.539  0.0111 *
Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 '' 1
(Dispersion parameter for binomial family taken to be 1)
   Null deviance: 98.719 on 20 degrees of freedom
Residual deviance: 33.278 on 17 degrees of freedom
AIC: 117.87
Number of Fisher Scoring iterations: 4
```

Estimating over dispersion in R

```
> X2 <- sum(residuals(g, type = "pearson")^2)
> X2
[1] 31.65114
> phi <- X2/g$df.residual
> phi
[1] 1.861832
> phi <- g$deviance/g$df.residual
> phi
[1] 1.957517
```

As we can see from the R output, the over dispersion parameter is greater than 1 and hence this is an indication of the presence of over dispersion

Confidence interval for model 1

Taking into account overdispersion in R

```
glm(model, family = quasibinomial,...)
```

Output quasi-binomial model 2

```
> g.over <- glm(cbind(germ, total - germ) ~ extract + seed + extract:seed, family =
    quasibinomial, data = fact)
> summary(q.over)
Call:
glm(formula = cbind(germ, total - germ) ~ extract + seed + extract:seed,
   family = quasibinomial, data = fact)
Deviance Residuals:
    Min
               10 Median
                                  30
                                           Max
-2.01617 -1.24398 0.05995 0.84695 2.12123
Coefficients:
                         Estimate Std. Error t value Pr(>|t|)
                                    0.2513 - 1.640
(Intercept)
                          -0.4122
                                                     0.1193
extractcucumber
                          0.5401
                                    0.3409 1.584 0.1315
                          -0.1459 0.3045 -0.479 0.6379
seedaeq75
extractcucumber:seedaeg75 0.7781 0.4181 1.861
                                                      0.0801 .
Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 '' 1
(Dispersion parameter for quasibinomial family taken to be 1.861832)
   Null deviance: 98.719 on 20 degrees of freedom
Residual deviance: 33.278 on 17 degrees of freedom
AIC: NA
Number of Fisher Scoring iterations: 4
> summary(g.over)$dispersion
[1] 1.861832
```

Confidence Interval model2

```
> library(MASS)
> fact$prop <- with(fact, germ/total)</pre>
> g.over.alt <- glm(prop ~ extract + seed +
  extract: seed,
+ weights = total, family = quasibinomial, data =
  fact)
> confint(g.over.alt)
Waiting for profiling to be done ...
                                 2.5 % 97.5 %
                           -0.91562380 0.07457178
(Intercept)
extractcucumber
                           -0.12415989 1.21529201
                           -0.74043242 0.45663323
seedaeg75
extractcucumber:seedaeg75 -0.04413964 1.59702376
```

Estimates and CI for proportion with and without over dispersion

_	$\hat{\pi}$	2.5%	97.5%
$\phi = 1$	0.36	0.31	0.42
$\hat{\phi} = 1.86$	0.36	0.28	0.45

To calculate this you need to install the R packages:

library(doBy)
library(R2HTML)

Example 2: Habitat preferences of lizards

- A study consists of two lizards type: Grohami and Opalinus.
- Response: number of sites (from the total) occupied by Grahami lizards.
- Covariates:
- 1. Height of the site (H).
- 2. Diameter (D).
- 3. Sun condition of the site (S, sun/ shade).
- 4. Time of the day (T).

Habitat preferences of lizards

```
> habitat
    G Total
             S
                D
   20
         22 S1 D1 H1 Early
          9 S1 D1 H1
                       Mid
          8 S1 D1 H1
                      Late
   13
         13 S1 D1 H2 Early
  8
         8 S1 D1 H2
                       Mid
  12
         12 S1 D1 H2
                      Late
         11 S1 D2 H1 Early
          5 S1 D2 H1
                        Mid
    5
          8 S1 D2 H1
                      Late
10
          6 S1 D2 H2 Early
11
          0 S1 D2 H2
                       Mid
12
       2 S1 D2 H2
                      Late
13 34
      45 S2 D1 H1 Early
14 69
      89 S2 D1 H1
                        Mid
15 18
         28 S2 D1 H1
                      Late
         36 S2 D1 H2 Early
16 31
17 55
         59 S2 D1 H2
                       Mid
         16 S2 D1 H2
18 13
                      Late
19 17
         32 S2 D2 H1 Early
20 60
         92 S2 D2 H1
                       Mid
21 8
         16 S2 D2 H1
                       Late
         13 S2 D2 H2 Early
22 12
23 21
         26 S2 D2 H2
                        Mid
24 4
          8 S2 D2 H2
                      Late
```

S: sun conditions sun / shade).

D: diameter (<2/>2).

H: hight (< 5 / > 5).

T: time of day (early/ mid day/late).

Habitat preferences of lizards: model formulation

$$y_{ijkl} \sim B(n_{ijkl}, \pi_{ijkl})$$
Total ample size.

Number of sites occupied by Grahami lizards.

 $\pi_{ijkl}=$ The probability that a site is occupied by Grahami lizards.

$$g(\pi_{ijkl}) = \mu + \alpha_i + \beta_j + \gamma_k + \delta_l + \dots$$

Habitat preferences of lizards: model formulation in R

Main effects model in R

$$g(\pi_{ijkl}) = \mu + \alpha_i + \beta_j + \gamma_k + \delta_l$$

> f1<-glm((G/Total)~H+D+S+T, family="binomial", data=habitat)

R output

```
summary(f1)
Call:
glm(formula = (G/Total) \sim H + D + S + T, family = "binomial",
   data = habitat)
Deviance Residuals:
                 Median
              10
    Min
                               30
                                       Max
-0.50878 -0.11019 0.02009 0.26466 0.52322
Coefficients:
          Estimate Std. Error z value Pr(>|z|)
(Intercept) 2.0618
                  1.4060 1.466 0.143
          1.0631 1.1222 0.947 0.343
HH2
DD2
           -0.8798 1.0841 -0.812 0.417
SS2
          -0.6415 1.0884 -0.589 0.556
TLate
          -1.2054 1.2761 -0.945 0.345
TMid 0.0587 1.4590 0.040 0.968
(Dispersion parameter for binomial family taken to be 1)
   Null deviance: 4.6730 on 22 degrees of freedom
Residual deviance: 1.5417 on 17 degrees of freedom
  (1 observation deleted due to missingness)
AIC: 28.658
```

Interpretation

```
Coefficients:
            Estimate Std.
                           Error
                                 z value Pr(>|z|)
                          1.4060
                                   1.466
(Intercept)
              2.0618
                                             0.143
              1.0631
                                   0.947
                                             0.343
HH2
                          1.1222
DD2
             -0.8798
                          1.0841
                                  -0.812
                                             0.417
                          1.0884
SS2
             -0.6415
                                  -0.589
                                             0.556
TLate
             -1.2054
                          1.2761
                                  -0.945
                                             0.345
TMid
              0.0587
                                             0.968
                          1.4590
                                   0.040
```

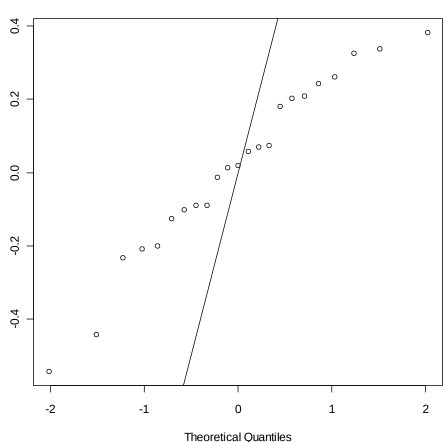
All the parameters estimates are not significant.

diagnostic

>r.pearson<-resid(f1, type="pearson")</pre>

- > par(mfrow=c(1,1))
- > qqnorm(r.pearson)
- > abline(0,1)

$$r_i^P = \frac{y_i - \hat{\mu}_i}{\sqrt{V(\hat{\mu}_i)}} \sim N(0,1)$$

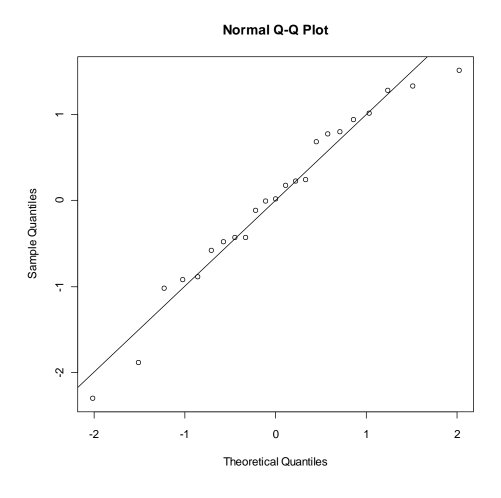


Normal Q-Q Plot

diagnostic

```
> mean(r.pearson)
[1] 0.01454735
  var(r.pearson)
[1] 0.05871372
|>r.s<-(r.pearson-mean(r.pearson))/</pre>
        sqrt((var(r.pearson)))
> qqnorm(r.s)
\geq abline (0,1)
```

The variance of pearson residual is much smaller than 1



Taking into account overdispersion

glm((G/Total)~H+D+S+T, family="binomial", data=habitat)



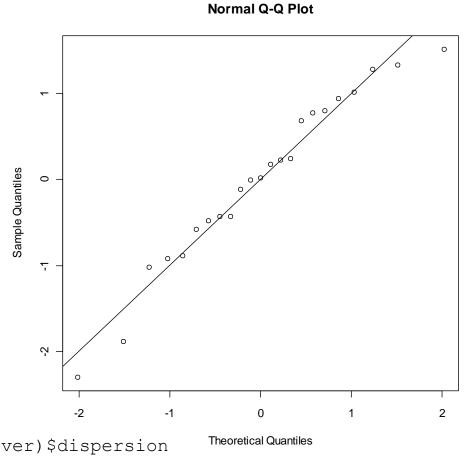
glm((G/Total)~H+D+S+T, family=quasibinomial, data= habitat)

Taking into account overdispersion

```
> f1.over <- glm((G/Total)~H+D+S+T, family=quasibinomial, data= habitat)
> summary(f1.over)
Call:
qlm(formula = (G/Total) \sim H + D + S + T, family = quasibinomial,
    data = habitat)
Deviance Residuals:
    Min
               10
                  Median
                                   30
                                           Max
-0.50878 -0.11019 0.02009 0.26466 0.52322
Coefficients:
           Estimate Std. Error t value Pr(>|t|)
                       0.3883 5.310 5.76e-05 ***
           2.0618
(Intercept)
                       0.3099 3.430 0.00319 **
HH2
            1.0631
                       0.2994 -2.939 0.00918 **
DD2
            -0.8798
                       0.3006 -2.134 0.04768 *
SS2
            -0.6415
                       0.3524 -3.420 0.00326 **
Tlate
           -1.2054
                       0.4029 0.146 0.88588
TMid
            0.0587
Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 '' 1
(Dispersion parameter for quasibinomial family taken to be 0.07626879)
   Null deviance: 4.6730 on 22 degrees of freedom
Residual deviance: 1.5417 on 17 degrees of freedom
  (1 observation deleted due to missingness)
AIC: NA
```

Diagnostic

The variance of pearson residual is much smaller than 1



> summary(f1.over)\$dispersion
[1] 0.07626879

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Example 3: ship damage

- The ships data from the MASS package concern a type of damage caused by waves to the forward section of cargocarrying vessels.
- The variables are
 - incidents number of damage incidents
 - service aggregate months of service
 - period period of operation: 1960-74, 75-79
 - year year of construction: 1960-64, 65-69, 70-74, 75-79
 - type type: "A" to "E"
- Here it makes sense to model the expected number of incidents per aggregate months of service.

Data in R

The data is available in the R the MASS library as:

```
> library(MASS)
> data(ships)
> ships2 <- subset(ships, service > 0)
> ships2$year <- as.factor(ships2$year)</pre>
   ships2$period <- as.factor(ships2$period)</pre>
> ships
   type year period service incidents
           60
1
      Α
                   60
                          127
                                        0
           60
                  75
                           63
      Α
3
           65
                  60 1095
      Α
           65
                  75 1095
4
      Α
5
           70
                   60
                         1512
      Α
                                        6
                  •
37
           70
                  60
                         1157
      \mathbf{E}
38
           70
                  75
                         2161
                                      12
      Ε
39
          75
                   60
                            0
      Ε
                                       0
      Ε
           75
                  75
                          542
40
```

Mean structure and model formulation in R

Model formulation:

$$Y_{ijk} \sim Poisson(\mu_{ijk})$$

$$g(\mu_{ijk}) = \mu + Tyep_i + Year_j + Period_k + \log(servise)$$
 offset

Model formulation in R:

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R output

```
> glm1 <- glm(formula = incidents ~ type + year + period,</pre>
     family = poisson(link = "log"), data = ships2,
     offset = log(service))
> summary(glm1)
Call:
glm(formula = incidents ~ type + year + period, family = poisson(link = "log"),
   data = ships2, offset = log(service))
Deviance Residuals:
            10 Median
   Min
                              3Q
                                      Max
-1.6768 -0.8293 -0.4370 0.5058 2.7912
Coefficients:
           Estimate Std. Error : value Pr(>|z|)
                      0.21744 -29.460 < 2e-16 ***
(Intercept) -6.4059(
           -0.54334
                      0.17759 -3.060 0.00222 **
typeB
           -0.6874
                      0.32904 -2.089 0.03670 *
typeC
                      0.29058 -0.261 0.79377
typeD
           -0.07596
                      0.23588 1.380 0.16750
typeE
           0.32558
                      0.14964 4.659 3.18e-06 ***
           0.69714
year65
                      0.16977 4.821 1.43e-06 ***
year70
           0.81843
year75
           0.45343
                      0.23317 1.945 0.05182 .
                       0.11827 3.251 0.00115 **
period75
          0.38447
Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
(Dispersion parameter for poisson family taken to be 1)
   Null deviance: 146.328 on 33 degrees of freedom
Residual deviance: 38.695 on 25 degrees of freedom
AIC: 154.56
Number of Fisher Scoring iterations: 5
```

Model 1: quasi-poisson log linear

```
> glm2 <- update(glm1, family = quasipoisson(link = "log"))</pre>
                                             \phi \neq 1
> summary(glm2)
> Call:
qlm(formula = incidents ~ type + year + period, family = quasipoisson(link = "loq"),
   data = ships2, offset = log(service))
Coefficients:
           Estimate Std. Error t value Pr(>|t|)
                                                      Standard errors are
                      0.28276 -22.655 < 2e-16 ***
(Intercept) -6.40590
typeB
          -0.54334
                     0.23094
                              -2.353 0.02681 *
                                                      changed since \Phi>1.
          -0.68740
                     0.42789 -1.607 0.12072
typeC
typeD -0.07596
                     0.37787 +0.201 0.84230
      0.32558
                     0.30674 1.061 0.29864
typeE
year65
           0.69714
                     0.19459
                               3.583 0.00143 **
                              3.707 0.00105 **
year70
           0.81843
                      0.22077
            0.45343
year75
                      0.30321
                               1.495 0.14733
            0.38447
                               2.500 0.01935 *
period75
                      0.15380
              0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
Signif. codes:
(Dispersion parameter for quasipoisson family taken to be 1.691028)
   Null deviance: 146.328 on 33 degrees of freedom
Residual deviance: 38.695 on 25 degrees of freedom
AIC: NA
Number of Fisher Scoring iterations: 5
```

Estimation of over dispersion parameter

```
> X2 <- sum(residuals(glm2, type = "pearson")^2)
> X2
[1] 42.27525
> phi <- X2/glm2$df.residual
> phi
[1] 1.69101
> phi <- g$deviance/glm2$df.residual
> phi
[1] 1.331111
```

CI for the parameters

> confint(glm2)

```
97.5 %
               2.5 %
            -6.9789252 -5.86832189
(Intercept)
            -0.9793495 -0.07040017
typeB
            -1.6043341
                        0.09972077
typeC
            -0.8628299
                        0.63543426
typeD
            -0.2880598
                        0.92322467
typeE
year65
             0.3217115
                        1.08674462
             0.3882167
                        1.25564720
year70
            -0.1562814
                        1.03712256
year75
                        0.68792187
period75
             0.0841713
```

Summary

ANOVA table without and with assuming over dispersion

Effect	DF	$\phi = 1$	$\hat{\phi} = 1.67$
		$P(> \chi^2)$	P(>F)
Туре	4	2.63E-11	2.29E-04
Period	1	1.1E-03	1.888E-02
Year	3	5.038e-09	5.777E-04