MATH3823 Generalized Linear Models

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Weekly schedule

i Week 11 (8 - 12 May)

Revision and examination preparation.

i Week 10 (1 - 5 May)

Any remaining module material. To be confirmed after Week 9.

- i Week 9 (27 31 March)
 - Before next Lecture: Re-read Chapter 6: Log-linear Models.
 - Lecture on Tuesday: Start Chapter 7: Extensions to Loglinear models.
 - Lecture on Thursday: Continue Chapter 7: Extensions to Loglinear models.
 - Weekly feedback: Relevant exercise questions from last year (solutions online).
- Coursework Practical Sessions (20 24 March)
 - Coursework for this module involves a single written report worth 20% of the module grade. This will mainly involve investigating different models using R and interpreting the results. Tasks are expected to be handed out on 14 March with **extended** hand-in deadline expect to be **5 April**. See *Learning Resources* / *Practical Assessment* for details of the tasks and for submission links.
- i Week 8 (20 24 March)
 - Before next Lecture: Re-read Chapter 5: Sections 5.1-5.3.
 - Lecture on Tuesday: Cancelled due to UCU strike.
 - Computer Practical on Tuesday: Cancelled due to UCU strike.
 - Computer Practical on Wednesday: Cancelled due to UCU strike.
 - Lecture on Thursday: Continue Chapter 6: Loglinear Modelling.
 - Weekly feedback: Relevant exercise questions from last year (solutions online).

i Week 7 (13 - 17 March)

- Before next Lecture: Re-read Chapter 5: Sections 5.1-5.3.
- Lecture on Tuesday: Complete Chapter 5: by looking at Section 5.4 Application to dose-response experiments.
- Lecture on Thursday: Cancelled due to UCU strike. Please self-study Chapter 6: Loglinear Modelling with Section 6.1 Overview and Section 6.2 Motivating Examples.
- Weekly feedback: Relevant exercise questions from last year (solutions online).

i Week 6 (6 - 10 March)

- Before next Lecture: Re-read Chapter 4: Sections 4.1-4.4.
- Lecture on Tuesday: Complete *Chapter 4:* with Section 4.5 Fitting GLMs in R
- Lecture on Thursday: Start Chapter 5: Logistic Regression with Sections 5.1 & 5.2.
- Weekly feedback: Relevant exercise questions from last year (solutions online).

Week 5 (27 February - 3 March)

- Before next Lecture: Re-read Chapter 4: Section 4.1.
- Lecture on Tuesday: Cancelled due to illness. Please read *Chapter 4: Section* 4.2.
- Lecture on Thursday: Chapter 4: Sections 4.3, 4.4 & 4.5.
- Weekly feedback: Start Exercises in Chapter 4.

i Week 4 (20 - 24 February)

- Before next Lecture: Be confident with all material up to, and including, Section 3.4 Moments of exponential-family distributions.
- Lecture on Tuesday: Chapter 3: Sections 3.5 & 3.6
- Lecture on Thursday: Start Chapter 4 by covering Section 4.1.
- Weekly feedback: Complete Exercises in *Chapter 3*.

Week 3 (13 - 17 February)

- Before next Lecture: Be confident with material in Chapter 2: Essentials of Normal Linear Models.
- Lecture on Tuesday: Cancelled due to UCU strike. Instead, self-study *Chapter 3: Sections 3.1 & 3.2.*
- Lecture on Thursday: Cancelled due to UCU strike. Instead, self-study *Chapter 3: Sections 3.3 & 3.4.*

• Before next Lecture: Complete questions and check solutions, including video(s), for all Exercises in *Chapters 1 and 2*. Start Exercises in *Chapter 3*.

i Week 2 (6 - 10 February)

- Before next Lecture: Please re-read Section 2.1: Overview and read Section 2.2: Types of normal linear model.
- Lecture on Tuesday: We will briefly cover all remaining material in *Chapter* 2: Essentials of Normal Linear Models.
- Before next Lecture: Please re-read Chapter 2 carefully.
- Lecture on Thursday: Cancelled due to UCU strike.
- Weekly feedback: Self-study the Exercises in Section 2.6 solutions to be posted during Week 3.

i Week 1 (30 January - 3 February)

- Before next Lecture: Please read the Overview.
- Lecture on Tuesday: We will briefly cover all material in *Chapter 1: Introduction*.
- Before next Lecture: Please re-read *Chapter 1* carefully.
- Lecture on Thursday: Start Chapter 2: Essentials of Normal Linear Models with Section 2.1: Overview.
- Weekly feedback: Self-study the Exercises in Section 1.5 solutions to be posted during Week 1.

Overview

Preface

These lecture notes are produced for the University of Leeds module MATH3823 - Generalized Linear Models for the academic year 2022-23. Please note that this material also forms part of the module MATH5824 - Generalized Linear and Additive Models. They are based on those used previously for this module and I am grateful to previous module lecturers for their considerable effort: Lanpeng Ji, Amanda Minter, John Kent, Wally Gilks, and Stuart Barber. This is the first year, however, that they have been produced in accessible format and hence some errors might occur during this conversion process. For information, I am using Quarto (a successor to RMarkdown) from RStudio to produce both the html and PDF, and then GitHub to create the website which can be accessed at rgaykroyd.github.io/MATH3823/. Please note that the PDF versions will only be made available on the University of Leeds Minerva system. Although I am a long-term user of RStudio, I have not previously used Quarto/RMarkdown nor Github and hence please be patient if there are hitches along the way.

RG Aykroyd, Leeds, November 22, 2022



Warning

Statistical ethics and sensitive data

Please note that from time to time we will be using data sets from situations which some might perceive as sensitive. All such data sets will, however, be derived from real-world studies which appear in textbooks or in scientific journals. The daily work of many statisticians involves applying their professional skills in a wide variety of situations and as such it is important to include a range of commonly encountered examples in this module. Whenever possible, sensitive topics will be signposted in advance. If you feel that any examples may be personally upsetting then, if possible, please contact the module lecturer in advance. If you are significantly effected by any of these situations, then you can seek support from the Student Counselling and Wellbeing service.

Official Module Description

Module summary

Linear regression is a tremendously useful statistical technique but is very limited. Generalised linear models extend linear regression in many ways - allowing us to analyse more complex data sets. In this module we will see how to combine continuous and categorical predictors, analyse binomial response data and model count data.

Objectives

On completion of this module, students should be able to:

- a) carry out regression analysis with generalised linear models including the use of link functions;
- b) understand the use of deviance in model selection;
- c) appreciate the problems caused by overdispersion;
- d) fit and interpret the special cases of log linear models and logistic regression;
- e) use a statistical package with real data to fit these models to data and to write a report giving and interpreting the results.

Syllabus

Generalised linear model; probit model; logistic regression; log linear models.

University Module Catalogue

For any further details, please see MATH3823 Module Catalogue page

1 Introduction

1.1 Overview

In previous modules you have studied linear models with a normally distributed error term, such as simple linear regression, multiple linear regression and ANOVA for normally distributed observations. In this module we will study **generalized** linear models.

Outline of the module:

- 1. Revision of linear models with normal errors.
- 2. Introduction to generalized linear models, GLMs.
- 3. Logistic regression models.
- 4. Loglinear models, including contingency tables.

Important

This module will make extensive use of \mathbf{R} and hence it is very important that you are comfortable with its use. If you need some revision, then material is available on Minerva under $RStudio\ Support$.

The purpose of a generalized linear model is to describe the dependence of a response variable y on a set of p explanatory variables $x = (x_1, x_2, \dots, x_p)$ where, conditionally on x, observation y has a distribution which is **not necessarily** normal.

Note that in these notes we may use lowercase letters, for example y or y_i , to denote both observed values or random variables, which is being considered should be clear from the context.

Important

This module will make extensive use of many basic ideas from statistics. If you need some revision, then see *Appendix A: Basic material* on Minerva under *Basic Prerequisite Material*.

1.2 Motivating example

Table 1.1 shows data¹ on the number of beetles killed by five hours of exposure to 8 different concentrations of gaseous carbon disulphide.

Table 1.1: Numbers of beetles killed by five hours of exposure to 8 different concentrations of gaseous carbon disulphide

Dose	No. of beetle	No. killed
x_i	m_i	y_{i}
1.6907	59	6
1.7242	60	13
1.7552	62	18
1.7842	56	28
1.8113	63	52
1.8369	59	53
1.8610	62	61
1.8839	60	60

Figure 1.1a shows the same data with a linear regression line superimposed. Although this line goes close to the plotted points, we can see some fluctuations around it. More seriously, this is a stupid model: it would predict a mortality rate of greater than 100% at a dose of 1.9 units, and a negative mortality rate at 1.65 units!

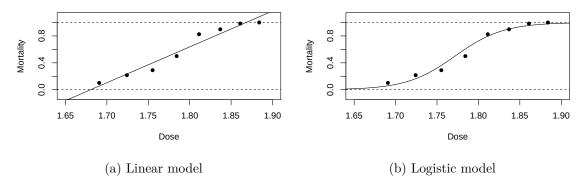


Figure 1.1: Beetle mortality rates with fitted dose- response curves.

A more sensible dose—response relationship for the beetle mortality data might be based on the *logistic* function (to be defined later), as plotted in Figure 1.1b. The resulting curve is a closer, more-sensible, fit. Later in this module we will see how this curve was fitted using maximum likelihood estimation for an appropriate generalized linear model.

¹Dobson and Barnett, 3rd edn, p.127

This is an example of a dose-response experiment which are widely used in medical and pharmaceutical situations.

Warning

Warning of potentially sensitive material. For further information on doseresponse experiments see, for example, www.britannica.com/science/dose-responserelationship.

1.3 Revision of least-squares estimation

Suppose that we have n paired data values $(x_1, y_1), \dots, (x_n, y_n)$ and that we believe these are related by a linear model

$$y_i = \alpha + \beta x_i + \epsilon_i$$

for all $i \in \{1, 2, \dots, n\}$, where $\epsilon_1, \dots, \epsilon_n$ are independent and identically distributed (iid) with $\mathbf{E}(\epsilon_i) = 0$ and $\mathrm{Var}(\epsilon_i) = \sigma^2$. The aim will be to find values of the model parameters, α, β and σ^2 using the data. Specifically, we will estimate α and β using the values which minimize the residual sum of squares (RSS)

$$RSS(\alpha, \beta) = \sum_{i=1}^{n} (y_i - (\alpha + \beta x_i))^2.$$
 (1.1)

This measures how close the data points are around the regression line and hence the resulting estimates, $\hat{\alpha}$ and $\hat{\beta}$, will give us a fitted regression line which is *closest* to the data.

It can be shown that Equation 1.1 takes its minimum when the parameters are given by

$$\hat{\alpha} = \bar{y} - \hat{\beta}\bar{x}, \quad \text{and} \quad \hat{\beta} = \frac{s_{xy}}{s_x^2}$$
 (1.2)

where \bar{x} and \bar{y} are the sample means,

$$s_{xy} = \frac{1}{n-1} \sum_{i=1}^n (x_i - \bar{x})(y_i - \bar{y})$$

is the sample covariance and

$$s_x^2 = \frac{1}{n-1} \sum_{i=1}^n (x_i - \bar{x})^2$$

is the sample variance of the x values. It can be shown that these estimators are unbiased, that is $E[\hat{\alpha}] = \alpha$ and $E[\hat{\beta}] = \beta$ – see Section 1.5.

The fitted regression lines is then given by $\hat{y} = \hat{\alpha} + \hat{\beta}x$, the fitted values by $\hat{y}_i = \hat{\alpha} + \hat{\beta}x_i$, and the model residuals by $r_i = \hat{\epsilon}_i = y_i - \hat{y}_i$ for all $i \in \{1, \dots, n\}$.

To complete the model fitting, we also estimate the error variance, σ^2 , using

$$\hat{\sigma}^2 = \frac{1}{n-2} \sum_{i=1}^n r_i^2. \tag{1.3}$$

Note that, by construction, $\bar{r} = 0$ and, further, it can be shown that $\hat{\sigma}^2$ is an unbiased estimator of σ^2 , that is $E[\hat{\sigma}^2] = \sigma^2$.

Returning to the above beetle data example, we have $\hat{\alpha} = -8.947843$, $\hat{\beta} = 5.324937$, and $\hat{\sigma}^2 = 0.0075151$.

We will interpret the output later, but in R, the fitting can be done with a single command with corresponding fitting output from a second command:

Call:

lm(formula = mortality ~ dose)

Residuals:

Min 1Q Median 3Q Max -0.10816 -0.06063 0.00263 0.05119 0.12818

Coefficients:

Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1

Residual standard error: 0.08669 on 6 degrees of freedom Multiple R-squared: 0.9524, Adjusted R-squared: 0.9445 F-statistic: 120.2 on 1 and 6 DF, p-value: 3.422e-05

Important

You should have met R output like this in previous statistics modules, but if you need some revision then see Appendix-C: Background to Analysis of Variance on Minerva under Basic Pre-requisite Material.

1.4 Types of variables

The way a variable enters a model will depends on its type. The most common five types of variable are:

1. Quantitative

- a. Continuous: for example, height; weight; duration. Real valued. Note that although recorded data is rounded it is still usually best regarded as continuous.
- b. Count (discrete): for example, number of children in a family; accidents at a road junction; number of items sold. Non-negative and integer-valued.

2. Qualitative

- a. Ordered categorical (ordinal): for example, severity of illness (Mild/ Moderate/Severe); degree classification (first/ upper-second/ lower-second/ third).
- b. Unordered categorical (nominal):
 - Dichotomous (binary): two categories: for example sex (M/F); agreement (Yes/No); coin toss (Head/Tail).
 - Polytomous (also known as polychotomous): more than two categories: for example blood group (A/ B/ O); eye colour (Brown/ Blue/ Green).

Note that although dichotomous is clearly a special case of polytomous, making the distinction is usually worthwhile as it often leads to a simplified modelling and testing approach.

1.5 Exercises

Important

Unless otherwise stated, data files will be available online at: rgaykroyd.github.io/MATH3823/Datasets/filename.ext, where filename.ext is the stated filename with extension.

1.1 Consider again the beetle data in Table 1.1. Perform the calculations by hand and then check the answers using R – a copy of the data is available in the file beetle.txt. Finally plot the fitted regression line on a scatter plot of the data. [Hint: See the code chunk used to produce Figure 1.1.]

1.2 Consider the following synthetic data:

	i = 1	i = 2	i = 3	i = 4	i = 5	i = 6	i = 7	i = 8
x_i	-1	0	1	2	2.5	3	4	6
\boldsymbol{y}_i	-2.8	-1.1	7.2	8.0	8.9	9.2	14.8	24.7

Plot the data to check that a linear model is suitable and then fit a linear regression model. Do you think that the fitted model can be reliably used to predict the values of y when x = 5 and x = 10? Justify your answers.

1.3 Starting from Equation 1.1, derive the estimation equations given in Equation 1.2. Further, show that $\hat{\alpha}$ and $\hat{\beta}$ are unbiased estimators of α and β . [Hint: Check your MATH1712 lecture notes.]

What can be said about $\hat{\sigma}^2$ as an estimator of σ^2 ? [Hint: There is a careful theoretical proof, but here only an intuitive explanation is expected.]

1.4 The Brownlee's Stack Loss Plant Data² is already available in **R**, with background details on the help page, ?stackloss. [Hint: You already met this example in MATH1712.]

After plotting all pairs of variables, which of Air.Flow, Water.Temp and Acid.Conc do you think could be used to model stack.loss using a linear regression? Justify your answer.

Perform a simple linear regression with using stack.loss as the response variable and your chosen variable as the explanatory variable. Add the fitted regression line to a scatter plot of the data and comment.

1.5 In an experiment conducted by de Silva et al. in 2020³ data was obtained to investigate falling objects and gravity, as first consider by Galileo and Newton. A copy of the data is available in the file physics_from_data.csv.

Read the data file into R and perform a simple linear regression of the maximum Reynolds number as the response variable and, in turn, each of the other variables as the explanatory variable.

Plot the data and add the corresponding fitted linear models. Which variable do you think helps explain Reynolds number the best? Why do you think this?

Here are an infinite number of further numerical examples from **maths e.g.** (thanks to https://www.mathcentre.ac.uk/):

Finding the intersercept

Finding the slope - Part 1

Finding the slope - Part 2

²Brownlee, K. A. (1960, 2nd ed. 1965) Statistical Theory and Methodology in Science and Engineering. New York: Wiley. pp. 491–500.

³de Silva BM, Higdon DM, Brunton SL, Kutz JN. Discovery of Physics From Data: Universal Laws and Discrepancies. Front Artif Intell. 2020 Apr 28;3:25. doi: 10.3389/frai.2020.00025. PMID: 33733144; PMCID: PMC7861345.

2 Essentials of Normal Linear Models

2.1 Overview

In many fields of application, we might assume the response variable is normally distributed. For example: heights, weights, log prices, etc.

The data¹ in Table 2.1 record the birth weights of 12 girls and 12 boys and their gestational ages (time from conception to birth).

A key question is, can we predict the birth weight of a baby born at a given gestational age using these data. For this we will need to make assumptions about the relationship between birth weight and gestational age, and any associated natural variation – that is we require a model.

First we should explore the data. Figure 2.1a shows a histogram of the birth weights indicating a spread around modal group 2800-3000 grams; Figure 2.1b indicates slightly higher birth weights for the boys than the girls; and Figure 2.1c shows an increasing relationship between weight and age. Together, these suggest that gestational age and sex are likely to be important for predicting weight.

Before considering possible models, Figure 2.2 again shows the relationship between weight and age but this time with the points coloured according to the baby's sex. This, perhaps, shows the boys to have generally higher weights across the age range than girls.

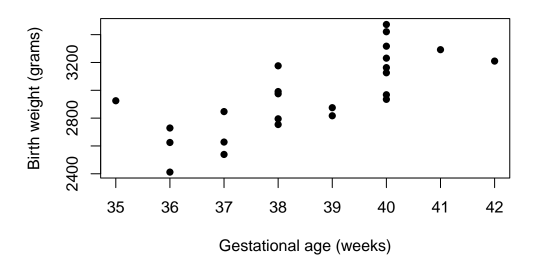
Of course, there are very many possible models, but here we will consider the following:

```
\begin{array}{lll} \hline \text{Model 0:} & \text{Weight} = \alpha \\ \hline \text{Model 1:} & \text{Weight} = \alpha + \beta. \\ \hline \text{Model 2:} & \text{Weight} = \alpha + \beta. \\ \hline \text{Model 3:} & \text{Weight} = \alpha + \beta. \\ \hline \text{Age} + \gamma. \\ \hline \text{Sex} + \delta. \\ \hline \text{Age.Sex} \\ \hline \end{array}
```

In these models, Weight is called the *response* variable (sometimes called the *dependent* variable) and Age and Sex are called the *covariates* or *explanatory* variables (sometimes called the *predictor* or *independent* variables). Here, Age is a continuous variable whereas Sex is coded as a *dummy* variable taking the value 0 for girls and 1 for boys; it is an example of a *factor*, in this case with just two *levels*: Girl and Boy.

¹Dobson and Barnett, 3rd edition, Table 2.3.





(c) Relationship beween variables

Figure 2.1: Birthweight and gestational age for 24 babies.

Table 2.1: Gestational ages (in weeks) and birth weights (in grams) for 24 babies (12 girls and 12 boys).

(a) Gi	rls		(b) Bo	oys
Gestational Age	Gestational Age Birth weight		Gestational Age	Birth weight
40	3317		40	2968
36	2729		38	2795
40	2935		40	3163
37	2754		35	2925
42	3210		36	2625
39	2817		37	2847
40	3126		41	3292
37	2539		40	3473
36	2412		37	2628
38	2991		38	3176
39	2875		40	3421
40	3231		38	3975



Figure 2.2: Birthweight and gestational age for 12 girls (red squares) and 12 boys (black dots).

Note that Model 0 is a special case of Model 1 (consider the situation when $\beta=0$) and that Model 1 is a special case of Model 2 (consider the situation when $\gamma=0$) and finally that Model 2 is a special case of Model 3 (consider the situation when $\delta=0$) – such models are called *nested*.

In these models, α , β , γ and δ are model parameters. Parameter α is called the *intercept* term; β is called the main effect of Age; and is interpreted as the effect on birth weight per week of gestational age. Similarly, γ is the main effect of Sex, interpreted as the effect on birth weight of being a boy (because girl is the baseline category).

Parameter δ is called the *interaction effect* between Age and Sex. Take care when interpreting an interaction effect. Here, it does not mean that age somehow affects sex, or vice-versa. It means that the effect of gestational age on birth weight depends on whether the baby is a boy or a girl.

These models can be fitted to the data using (Ordinary) *Least Squares* to produce the results presented in Figure 2.3.

Which model should we use?

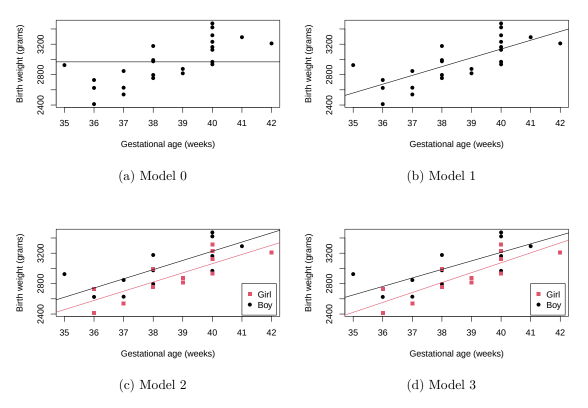


Figure 2.3: Birthweight and gestational age data with superimposed fitted regression lines from various competing models.

We know from previous modules that statistical tests can be used to check the importance of regression coefficients and model parameters, but it is also important to use the graphical results, as in Figure 2.3, to guide us.

Model 0 says that there is no change in birth weight with gestational age which means that we would use the average birth weight as the prediction whatever the gestational age – this makes no sense. As we can easily see from the scatter plot of the data, the fitted line in this case is clearly inappropriate.

Model 1 does not take into account whether the baby is a girl or a boy, but does model the relationship between birth weight and gestational age. This does seem to provide a good fit and might be adequate for many purposes. Recall from Figure 2.1b and Figure 2.2, however, that for a given gestational age the boys seem to have a higher birth weight than the girls.

Model 2 does take the sex of the baby into account by allowing separate intercepts in the fitted lines – this means that the lines are parallel. By eye, there is a clear difference between these two lines but it might not be important.

Model 3 allows for separate slopes as well as intercepts. There is a slight difference in the slopes, with the birth weight of the girls gradually catching-up as the gestational age increases. It is difficult to see, however, if this will be a general pattern or if it is only true for this data set – especially given the relatively small sample size.

Here, it is not clear by eye which of the fitted models will be the best and hence we should use a statistical test to help. In particular, we can choose between the models using F-tests.

Let y_i denote the value of the dependent variable Weight for individual $i=1,\ldots,n,$ and let the four models be indexed by k=0,1,2,3.

Let R_k denote the residual sum of squares (RSS) for Model k:

$$R_k = \sum_{i=1}^n (y_i - \hat{\mu}_{ki})^2, \tag{2.1}$$

where $\hat{\mu}_{ki}$ is the fitted value for individual *i* under Model *k*. Let r_k denote the corresponding residual degrees of freedom for Model *k* (the number of observations minus the number of model parameters).

Consider the following hypotheses:

$$H_0: \text{Model } 0 \text{ is true}; \quad H_1: \text{Model } 1 \text{ is true}.$$

Under the null hypothesis H_0 , the difference between R_0 and R_1 will be purely random, so the between-models mean-square $(R_0-R_1)/(r_0-r_1)$ should be comparable to the residual mean-square R_1/r_1 . Thus our test statistic for comparing Model 1 to the simpler Model 0 is:

$$F_{01} = \frac{(R_0 - R_1)/(r_0 - r_1)}{R_1/r_1}. (2.2)$$

It can be shown that, under the null hypothesis H_0 , the statistic F_{01} will have an F-distribution on $r_0 - r_1$ and r_1 degrees of freedom, which we write: $F_{r_0 - r_1, r_1}$. Under the alternative hypothesis H_1 , the difference $R_0 - R_1$ will tend to be larger than expected under H_0 , and so the observed value F_{01} will probably lie in the upper tail of the $F_{r_0 - r_1, r_1}$ distribution.

Returning to the birth weight data, we obtain the following output from R when we fit Model 1:

```
(Intercept) age
-1484.9846 115.5283
```

Analysis of Variance Table

```
Response: weight

Df Sum Sq Mea
```

Residuals 22 816074 37094

Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1

Thus we have parameter estimates: $\hat{\alpha} = -1484.98$ and $\hat{\beta} = 115.5$. The Analysis of Variance (ANOVA) table gives: $R_0 - R_1 = 1013799$ with $r_0 - r_1 = 1$ and $R_1 = 816074$ with $r_1 = 22$.

If we wanted R_0 and r_0 then we can either fit Model 0 or get them by subtraction.

The F_{01} statistic, Equation 2.2, is then

$$F_{01} = \frac{113799/1}{816074/22} = 27.33,$$

which can be read directly from the ANOVA table in the column headed 'F value'.

Is $F_{01}=27.33$ in the upper tail of the $F_{1,22}$ distribution? (See Figure 2.4 and note that 27.33 is very far to the right.) The final column of the ANOVA table tells us that the probability of observing $F_{01}>27.33$ is only 3.04×10^5 – this is called a p-value. The *** beside this p-value highlights that its value lies between 0 and 0.001. This indicates that we should reject H_0 in favour of H_1 – there is very strong evidence for the more complicated model. Thus we would conclude that the effect of gestational age is statistically significant in these data.

Next, consider the following hypotheses:

$$H_0: Model 1 \text{ is true}; \quad H_1: Model 2 \text{ is true}.$$

Under the null hypothesis H_0 , the difference between R_1 and R_2 will be purely random, so the between-models mean-square $(R_1 - R_2)/(r_1 - r_2)$ should be comparable to the residual



Figure 2.4: Probability density function of ${\cal F}_{01}$ distribution.

mean-square R_2/r_2 . Thus our test statistic for comparing Model 2 to the simpler Model 1 is:

$$F_{12} = \frac{(R_1 - R_2)/(r_1 - r_2)}{R_2/r_2}. (2.3)$$

Under the null hypothesis H_0 , the statistic F_{12} will have an F-distribution on r_1-r_2 and r_2 degrees of freedom, which we write: $F_{r_1-r_2,r_2}$. Under the alternative hypothesis H_1 , the difference R_1-R_2 will tend to be larger than expected under H_0 , and so the observed value F_{12} will probably lie in the upper tail of the $F_{r_1-r_2,r_2}$ distribution.

Returning to the birth weight data, we obtain the following output from R (where sexM denotes Boy):

```
(Intercept) age sexM
-1773.3218 120.8943 163.0393
```

Analysis of Variance Table

Response: weight

Df Sum Sq Mean Sq F value Pr(>F)
age 1 1013799 1013799 32.3174 1.213e-05 ***
sex 1 157304 157304 5.0145 0.03609 *
Residuals 21 658771 31370

Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1

Thus we have parameter estimates: $\hat{\alpha} = -1773.3$, $\hat{\beta} = 120.9$ and $\hat{\gamma} = 163.0$, the latter being the effect of being a boy compared to the baseline category of being a girl.

The Analysis of Variance (ANOVA) table gives: $R_1 - R_2 = 157304$ with $r_1 - r_2 = 1$, and $R_2 = 658771$ with $r_2 = 21$. The F_{12} statistic, Equation 2.3, is then

$$F_{12} = \frac{157304/1}{658771/21} = 5.0145,$$

which can be read directly from the ANOVA table in the column headed 'F value'. Is $F_{12} = 5.01$ in the upper tail of the $F_{1,21}$ distribution?

The final column of the ANOVA table tells us that the probability of observing $F_{12} > 5.01$ is only 0.03609 – this is called a p-value. The * beside this p-value highlights that its value lies between 0.01 and 0.05. This indicates that we should reject H_0 in favour of H_1 – there is evidence for the more complicated model. Thus we would conclude that the effect of the sex of the baby, after controlling for gestational age, is statistically significant in these data.

To complete the analysis, we should now compare Model 2 with Model 3 - see Exercises.

2.2 Types of normal linear model

Here we consider how normal linear models can be set up for different types of explanatory variable. The dependent variable y is modelled as a linear combination of p explanatory variables $x=(x_1,x_2,\ldots,x_p)$ plus a random error $\epsilon \sim N(0,\sigma^2)$, where '~' means 'is distributed as'. Several models are of this kind, depending on the number and type of explanatory variables. Table 2.3 lists some types of normal linear models with their explanatory variable types.

Table 2.3: Types of normal linear model and their explanatory variable types where indicator function I(x = j) = 1 if x = j and 0 otherwise.

p	Explanatory variables	Model
1	Quantitative	Simple linear regression $y = \alpha + \beta x + \epsilon$
>1	Quantitative	Multiple linear regression $y = \alpha + \sum_{i=1}^{p} \beta_i x_i + \epsilon$
1	Dichotomous $(x = 1 \text{ or } 2)$	Two-sample t-test $y = \alpha + \delta I(x = 2) + \epsilon$
1	Polytomous, k levels $(x = 1, \dots, k)$	One-way $\text{ANOVA} y = \alpha + \sum_{j=1}^k \delta_j \ I(x=j) + \epsilon$

Table 2.3: Types of normal linear model and their explanatory variable types where indicator function I(x = j) = 1 if x = j and 0 otherwise.

. 1	0 1: .:	n-way ANOVA	
>1	Qualitative	p-way ANOVA	

For the two-sample t-test model², observations in the two groups have means $\alpha + \beta_1$ and $\alpha + \beta_2$. Notice, however, that we have three parameters with only two group sample means and hence parameter estimation is not possible. To avoid this identification problem, we either impose a 'corner' constraint: $\beta_1 = 0$ and then β_2 represents the difference in the Group 2 mean relative to a baseline of Group 1. Alternatively, we may impose a 'sum-to-zero' constraint: $\beta_1 + \beta_2 = 0$, the values $\beta_1 = -\beta_2$ then give differences in the groups means relative to the overall mean. Table 2.4 shows the effect of the parameter constraint on the group means.

Table 2.4: Parameters in the two-sample t-test model after imposing parameter constraint to avoid the identification problem.

Constraint	Group 1 mean	Group 2 mean
$\beta_1 = 0$	α	$\alpha + \beta_2$
$\beta_1 + \beta_2 = 0$	$\alpha-\beta_2$	$\alpha + \beta_2$

For the general one-way ANOVA model with k groups, observations in Group j have mean $\alpha+\delta_j$, for $j=1,\ldots,k$ – that leads to k+1 parameters describing k group means. Again we can impose the 'corner' constraint: $\delta_1=0$ and then δ_j represents the difference in means between Group j and the baseline Group 1. Alternatively, we may impose a 'sum-to-zero' constraint: $\sum_{j=1}^k \delta_j = 0$ and again $(\delta_1,\delta_2,\ldots,\delta_k)$ then represents an individual group effect relative to the overall data mean.

2.3 Matrix representation of linear models

All of the models in Table 2.3 can be fitted by least squares (OLS). To describe this, a matrix formulation will be most convenient:

$$\mathbf{Y} = X\beta + \epsilon \tag{2.4}$$

where

- Y is an $n \times 1$ vector of observed response values with n being the number of observations.
- X is an $n \times p$ design matrix, to be discussed below.
- β is a $p \times 1$ vector of parameters or coefficients to be estimated.

²Notice that this is a special case of the one-way ANOVA when there are only two-groups.

• ϵ is an $n \times 1$ vector of independent and identically distributed (IID) random variables, which here $\epsilon \sim N(0, \sigma^2)$ and is called the "error" term.

2.4 Construction of the design matrix

Creating the design matrix is a key part of the modelling as it describes the important structure of investigation or experiment. The design matrix can be constructed by the following process.

- 1. Begin with an X containing only one column: a vector of ones for the overall mean or intercept term (the α in Table 2.3).
- 2. For each explanatory variable x_i , do the following:
 - a. If a variable x_i is quantitative, add a column to X containing the values of x_i .
 - b. If x_j is qualitative with k levels, add k "dummy" columns to X, taking values 0 and 1, where a 1 in the ℓ th dummy column identifies that the corresponding observation is at level ℓ of factor x_j . For example, suppose we have a factor $\mathbf{x}_j = (M, M, F, M, F)$ representing the sex of n = 5 individuals. This information can be coded into two dummy columns of X:

$$\begin{array}{ccc}
F & M \\
0 & 1 \\
0 & 1 \\
1 & 0 \\
0 & 1 \\
1 & 0
\end{array}$$

3. When qualitative variables are present, X will be singular – that is, there will be linear dependencies between the columns of X. For example, the sum of the two columns above is a vector of ones, the same as the intercept column. We resolve this identification problem by deleting some columns of X. This is equivalent to applying the corner constraint $\delta_1 = 0$ in the one-way ANOVA.

In the above example, after removing a column, we get:

$$\mathbf{X} = \begin{bmatrix} 1 & 1 \\ 1 & 1 \\ 1 & 0 \\ 1 & 1 \\ 1 & 0 \end{bmatrix}.$$

4. Each column of X represents either a quantitative variable, or a level of a qualitative variable. We will use $i=1,\ldots,n$ to label the observations (rows of X) and $j=1,\ldots,p$ to label the columns of X.

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2.4.1 Example: Simple linear regression

Consider the simple linear regression model $y = \alpha + \beta x + \epsilon$ with $\epsilon \sim N(0, \sigma^2)$. Given data on n pairs $(x_i, y_i), i = 1, ..., n$, we write this as

$$y_i = \alpha + \beta x_i + \epsilon_i, \quad \text{for } i = 1, 2, \dots, n, \tag{2.5}$$

where the ϵ_i are IID $N(0, \sigma^2)$. In matrix form, this becomes

$$\mathbf{Y} = X\beta + \epsilon \tag{2.6}$$

with

$$\mathbf{Y} = \begin{bmatrix} y_1 \\ \vdots \\ y_n \end{bmatrix}, \quad X = \begin{bmatrix} 1 & x_1 \\ \vdots & \vdots \\ 1 & x_n \end{bmatrix}, \quad \beta = \begin{bmatrix} \beta_1 \\ \beta_2 \end{bmatrix} = \begin{bmatrix} \alpha \\ \beta \end{bmatrix}, \quad \epsilon = \begin{bmatrix} \epsilon_1 \\ \vdots \\ \epsilon_n \end{bmatrix}.$$

The *i*th row of Equation 2.6 has the same meaning as Equation 2.5:

$$y_i = 1 \times \beta_1 + x_i \times \beta_2 + \epsilon_i = \alpha + \beta x_i + \epsilon_i, \ \text{ for } i = 1, 2, \dots, n.$$

2.4.2 Example: One-way ANOVA

For one-way ANOVA with k levels, the model is

$$y_i = \alpha + \sum_{i=1}^k \delta_j \ I(x_i = j) + \epsilon_i, \quad \text{for } i = 1, 2, \dots, n,$$

where x_i denotes the group level of individual i. So if y_i is from the jth group then $y_i \sim N(\alpha + \delta_j, \sigma^2)$. Here α is the intercept and the $(\delta_1, \delta_2, \dots, \delta_k)$ represent the "main effects".

We can store the information about the levels of g in a dummy matrix $X^* = (x_{ij}^*)$ where

$$x_{ij}^* = \begin{cases} 1, & g_i = j, \\ 0, & \text{otherwise.} \end{cases}$$

Then set $X = [1, X^*]$, where 1 is an *n*-vector of 1's. For the male–female example at (1.12), we have n = 5 and a sex factor:

$$g = \begin{bmatrix} 1 \\ 1 \\ 2 \\ 1 \\ 2 \end{bmatrix}, \quad X = \begin{bmatrix} 1 & 1 & 0 \\ 1 & 1 & 0 \\ 1 & 0 & 1 \\ 1 & 1 & 0 \\ 1 & 0 & 1 \end{bmatrix}, \quad \beta = \begin{bmatrix} \beta_1 \\ \beta_2 \\ \beta_3 \end{bmatrix} = \begin{bmatrix} \alpha \\ \delta_1 \\ \delta_2 \end{bmatrix}.$$

Then the *i*th row of X becomes $\beta_1 + \beta_2 = \alpha + \delta_1$ if $g_i = 1$ and $\beta_1 + \beta_3 = \alpha + \delta_2$ if $g_i = 2$. That is, the *i*th row of X is

$$\alpha + \sum_{i=1}^{2} \delta_{j} I(g_{i} = j)$$

so this model can be written $Y = X\beta + \epsilon$. Here, X is singular: its last two columns added together equal its first column. Statistically, the problem is that we are trying to estimate two means (the mean response for Boys and the mean response for Girls) with three parameters $(\alpha, \delta_2 \text{ and } \delta_2)$.

In practice, we often resolve this aliasing or identification problem by setting one of the parameters to be zero, that is $\delta_1 = 0$, which corresponds to deleting the second column of X).

2.5 Model shorthand notation

In R, a qualitative (categorical) variable is called a *factor*, and its categories are called *levels*. For example, variable Sex in the birth weight data (above) has levels coded "M" for 'Boy' and "F" for 'Girl'. It may not be obvious to R whether a variable is quantitative or qualitative. For example, a qualitative variable called **Grade** might have categories 1, 2 and 3. If **grade** was included in a model, R would treat it as quantitative unless we declare it to be a factor, which we can do with the command:

grade = as.factor(grade)

A convenient model-specification notation has been developed from which the design matrix X can be constructed. Below, E, F, \dots denote generic quantitative (continuous) or qualitative (categorical) variables. Terms in this notation may take the following forms:

- a. 1: a column of 1's to accommodate an intercept term (the α 's of Table 2.3). This is included in the model by default.
- b. E: variable E is included in the model. The design matrix includes k_E columns for E. If E is quantitative, $k_E = 1$. If E is qualitative, k_E is the number of levels of E minus 1.
- c. E+F: both E and F are included the model. The design matrix includes k_E+k_F columns accordingly.
- d. E: F (sometimes $E \cdot F$): the model includes an interaction between E and F; each column that would be included for E is multiplied by each column for F in turn. The design matrix includes $k_E \times k_F$ columns accordingly.
- e. E * F: shorthand for 1 + E + F + E : F: useful for crossed models where E and F are different factors. For example, E labels age groups; F labels medical conditions.

- f. E/F: shorthand for 1+E+E:F: useful for nested models where F is a factor whose levels have meaning only within levels of factor E. For example, E labels different hospitals; F labels wards within hospitals.
- g. $\operatorname{poly}(E;\ell)$: shorthand for an orthogonal polynomial, wherein x contains a set of mutually orthogonal columns containing polynomials in E of increasing order, from order 1 through order ℓ .
- h. -E: shorthand for removing a term from the model; for example E * F E is short for 1 + F + E : F.
- i. I(): shorthand for an arithmetical expression (not to be confused with the indicator function defined above). For example, I(E+F) denotes a new quantitative variable constructed by adding together quantitative variables E and F. This would cause an error if either E or F has been declared as a factor. What would happen in this example if we omitted the $I(\cdot)$ notation?

The notation uses "~" as shorthand for "is modelled by" or "is regressed on". For example,

• Weight is regressed on age-group and sex with no interaction between them:

$${\tt Weight} \sim {\tt Age} + {\tt Sex}$$

as for the birthweight data in Figure 1.2c.

• Well being is regressed on age-group and income-group, where income is thought to affect wellbeing differentially by age:

Wellbeing
$$\sim$$
 Age $*$ Income

• Class of degree is regressed on school of the university and on degree subject within the school:

$${\tt DegreeClass} \sim {\tt School/Subject}$$

• Yield of wheat is regressed on seed-variety and annual rainfall:

Yield
$$\sim$$
 Variety + poly(Rainfall, 2)

• Profit is regressed on amount invested:

Profit
$$\sim$$
 Investment -1

(no intercept term, that is a regression through the origin).

2.6 Exercises

Important

Please note that these questions are currently unchecked. I aim to look at them very soon but a apologize is anything is incomplete or there are errors.

2.1. An extra model which could have been considered for the Birth weight data example would be one that says that Weight is different for girls and boys, but does not depend on gestational age.

Write down the equation corresponding to this model. Then, load the birth weight data into RStudio and fit the model. How are the fitted model parameters related to the overall birth weight mean and the mean birth weights of the girls and boys? Is this a good fit to the data? Is Sex statistically significant?

2.2. In an experiment to investigate Ohm's Law, V = IR where V is Voltage, I is current and R is resistance of the material, the following data³ were recorded:

Table 2.5: Experimental verification of Ohm's Law

Voltage (Volts)	4	8	10	12	14	18	20	24
Current (mAmps)	11	24	30	36	40	53	58.5	70

Does this data support Ohm's Law? What is the resistance of the material used?

2.3 In an investigation⁴ into the effect of eating on pulse rate, 6 men and 6 women were tested before and after a meal, with the following results:

Table 2.6: At rest pulse rate before and after a meal for men and women

Men	before	105	79	79	103	87	97
	after	109	87	86	109	100	101
Women	before	74	73	82	78	86	77
	after	82	80	90	90	93	81

Suggest a suitable model for this situation and write down the corresponding design matrix. Calculate the parameter estimates using the matrix regression estimation equation.

Perform an appropriate analysis in R to find out if there is evidence to suggest that the change in pulse rate due to a meal is the same for men and women.

2.4 A laboratory experiment⁵ was performed into the effect of seasonal floods on the growth of barley seedlings in a incubator, as measured by their height in mm. Three types of barley seed (Goldmarker, Midas, Igri) were used with two watering condition (Normal and Waterlogged). Further, each combination was repeated four times on different shelves in

³Aykroyd, P.J. (1956). Unpublished.

⁴Source unknown.

 $^{^5\}mathrm{Source}$ unknown.

the laboratory incubator (Top, Second, Third and Bottom shelf). The data are available in the file barley.csv

Suggest a suitable model for this situation. Identify the response and explanatory variables and list the levels for any qualitative variables. Write down the design matrix for each model you consider.

Perform appropriate analyses to test if each of the following are important: (a) watering condition, (b) type of barley seed, and (c) shelf position.

In the analysis, do not include any interactions involving shelf position. If you find a significant interaction between watering condition and type of barley seed, carefully interpret the parameter estimates.

3 GLM Theory

3.1 Motivating examples

We cannot always assume that the dependent variable Y is normally distributed. For example, for the beetle mortality data in Table 1.1, suppose each beetle subjected to a dose x_i has a probability p_i of being killed. Then, the number of beetles killed Y_i out of a total number m_i at dose-level x_i will have a $Bin(m_i, p_i)$ distribution with probability mass function

$$\Pr(y_i;\ p_i,m_i) = \left(\begin{array}{c} m_i \\ y_i \end{array}\right) p_i^{y_i} (1-p_i)^{m_i-y_i} \eqno(3.1)$$

where y_i takes values in $\{0, 1, ..., m_i\}$.

Table 3.1 contains seasonal data on tropical cyclones for 13 seasons. Suppose that, within season i, there is a constant probability $\lambda_i dt$ of a cyclone occurring in any short time-interval dt. Then the total number of cyclones Y_i during season i will have a Poisson distribution with mean λ_i , that is $Y_i \sim \text{Po}(\lambda_i)$, with probability mass function

$$\Pr(y_i; \ \lambda_i) = \frac{\lambda_i^{y_i} e^{-\lambda_i}}{y_i!} \tag{3.2}$$

where y_i takes values in $\{0, 1, 2, ...\}$.

Table 3.1: Numbers of tropical cyclones in n = 13 successive seasons¹

Season	1	2	3	4	5	6	7	8	9	10	11	12	13
No of	6	5	4	6	6	3	12	7	4	2	6	7	4
cyclones													

In these two examples, we have non-normal data and would like to know whether and how the dependent variable Y_i depends on the covariate x_i or i.

Generalized linear models provide a modelling framework for data analysis in the nonnormal setting. We will revisit the beetle mortality and cyclone data sets after describing the structure of a generalized linear model.

¹Dobson and Barnett, 3rd edn, Table 1.2

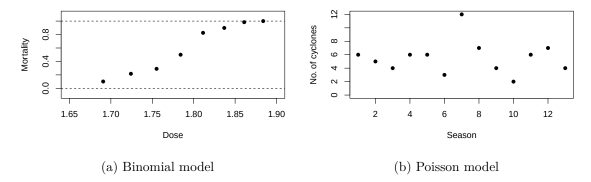


Figure 3.1: Examples of non-normally distributed data.

3.2 The GLM structure

A generalized linear model relates a continuous or discrete response variable Y to a set of explanatory variables $\mathbf{x} = (x_1, \dots, x_p)$. The model contains three parts:

Random part: The probability (mass or density) function of Y is assumed to belong to the two-parameter exponential family of distributions with parameters θ and ϕ :

$$f(y;\theta,\phi) = \exp\left\{\frac{y\theta - b(\theta)}{\phi} + c(y,\phi)\right\},\tag{3.3}$$

where $\phi > 0$. Here, θ is called the *canonical* or *natural* parameter of the distribution and ϕ is called the *scale* parameter. We show below that the mean E[Y] depends only on θ , and Var[Y] depends on ϕ and possibly also θ . Various choices for functions $b(\cdot)$ and $c(\cdot)$ produce a wide variety of familiar distributions (see below). Sometimes we may set $\phi = 1$; then Equation 3.3 is called the *one-parameter exponential family*.

Further, note that in some references to generalized linear models (such as Dobson and Barnett, 3rd edn.), ϕ does not appear at all in the exponential family formula Equation 3.3, instead it is absorbed into θ and $b(\theta)$.

In this module, we will generally assume that each observation Y_i , $i=1,\ldots,n$, is independently drawn from an exponential family where θ depends on the covariates. Thus we write

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

Note the subscripts on both y and θ , and hence the observations may not be identically distributed.

Systematic part: This is a linear predictor:

$$\eta = \sum_{j=1}^{p} \beta_j x_j. \tag{3.4}$$

Note that the symbol η is pronounced eta.

Link function: This is a one-to-one function providing the link between the linear predictor η and the mean $\mu = E[Y]$:

$$\eta = g(\mu), \text{ and } \mu = g^{-1}(\eta) = h(\eta).$$
(3.5)

Here, $g(\mu)$ is called the *link function*, and $h(\eta)$ is called the *inverse link function*.

We will now discuss each of these parts in more detail.

3.3 The random part of a GLM

We begin with some examples of exponential family members.

Example: Poisson distribution

If Y has a Poisson distribution with parameter λ , that is $Y \sim \text{Po}(\lambda)$, then Y takes values in $\{0, 1, 2, ...\}$ and has probability mass function:

$$f(y) = \frac{e^{-\lambda} \lambda^y}{y!} = \exp\left\{y \log \lambda - \lambda - \log y!\right\},\tag{3.6}$$

which has the form of Equation 3.3 with components as in Table 3.2.

Table 3.2: Exponential model components for the Poisson

$$\frac{\theta \qquad \phi \qquad b(\theta) \qquad c(y,\phi)}{\log \lambda \quad 1 \quad \lambda = e^{\theta} \quad -\log y!}$$

For example, to model the cyclone data in Table 3.1, we might simply assume that the number of cyclones in each season has a Poisson distribution, assuming a constant rate λ across all seasons i. That is $Y_i \sim \text{Po}(\lambda)$. The parameter would be simply estimated by the sample mean, $\hat{\lambda} = \bar{y} = 5.5384615$.

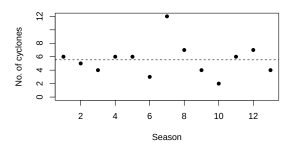
Example: Binomial distribution

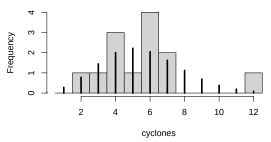
Let Y have a Binomial distribution, that is $Y \sim \text{Bin}(m, p)$, with m fixed. Then Y is discrete, taking values in $\{0, 1, ..., m\}$, and has probability mass function:

$$f(y)={m\choose y}p^y(1-p)^{m-y}={m\choose y}\left(\frac{p}{1-p}\right)^y(1-p)^m$$

which can be re-written as

$$f(y) = \exp\left\{y \text{ logit } p + m \log(1-p) + \log {m \choose y}\right\}, \tag{3.7}$$





(a) No. of cyclones against season with mean (b) Fitted Poisson model assuming constant rate Figure 3.2: Poisson model fitted to cyclone data.

which has the form of Equation 3.3 with,

$$\theta = \text{logit } p = \log\left(\frac{p}{1-p}\right),$$

and with components as in Table 3.3.

Table 3.3: Exponential model components for the Binomial

θ	ϕ	$b(\theta)$	$c(y,\phi)$
$\overline{\text{logit } p}$	1	$m\log(1+e^{\theta})$	$\log \binom{m}{y}$

Note that it can be shown that $-m\log(1-p)=m\log(1+e^{\theta})$ – see Exercises.

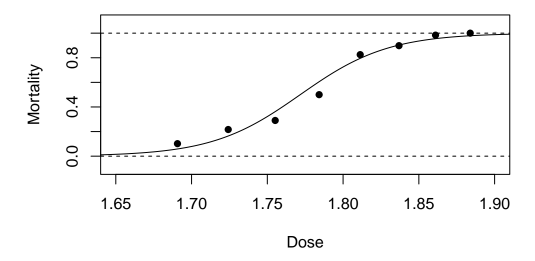


Figure 3.3: Binomial model fitted to beetle data.

Example: Normal distribution 33

Let Y have a Normal distribution with mean μ and variance σ^2 , that is $Y \sim N(0, \sigma^2)$. Then Y takes values on the whole real line and has probability density function

$$\begin{split} f(y;\mu,\sigma^2) &= \frac{1}{\sqrt{2\pi\sigma^2}} \exp\left\{\frac{-1}{2\sigma^2}(y-\mu)^2\right\}, \\ &= \frac{1}{\sqrt{2\pi\sigma^2}} \exp\left\{-\frac{y^2}{2\sigma^2} + \frac{y\mu}{\sigma^2} - \frac{\mu^2}{2\sigma^2}\right\} \\ &= \exp\left\{\frac{y\mu - \mu^2/2}{\sigma^2} + \left[\frac{-y^2}{2\sigma^2} - \frac{1}{2}\log(2\pi\sigma^2)\right]\right\}, \end{split}$$

which has the form of Equation 3.3 with components as in Table 3.4.

Table 3.4: Exponential model components for the Gaussian

θ	ϕ	b(heta)	$c(y,\phi)$
μ	σ^2	$\theta^2/2$	$-\frac{y^2}{2\phi} - \frac{1}{2}\log(2\pi\phi)$

From the usual regression point of view, we write $y = \alpha + \beta x + \epsilon$, with $\epsilon \sim N(0, \sigma^2)$. From the point of view of a generalized linear model, we write $Y \sim N(\mu, \sigma^2)$ where $\mu(x) = \alpha + \beta x$.

3.4 Moments of exponential-family distributions

It is straightforward to find the mean and variance of Y in terms of $b(\theta)$ and ϕ . Since we want to explore the dependence of E[Y] on explanatory variables, this property makes the exponential family very convenient.

Proposition 3.1. For random variables in the exponential family:

$$E[Y] = b'(\theta), \quad and \quad Var[Y] = b''(\theta)\phi.$$
 (3.8)

Proof We give the proof for a continuous random variables. For the discrete case, replace all integrals by sums – see Exercises.

Starting with the simple property that all probability density functions integrate to 1, we have

$$1 = \int \exp\left\{\frac{y\theta - b(\theta)}{\phi} + c(y, \phi)\right\} dy$$

and then differentiating both sides with respect to θ gives

$$0 = \int \left[\frac{y - b'(\theta)}{\phi} \right] \exp \left\{ \frac{y\theta - b(\theta)}{\phi} + c(y, \phi) \right\} dy.$$
 (3.9)

Next, using the definition of the exponential family to simplify the equation gives

$$0 = \int \left[\frac{y - b'(\theta)}{\phi} \right] f(y; \theta) \ dy$$

and expanding the brackets leads to

$$0 = \frac{1}{\phi} \left(\int y f(y;\theta) dy - b'(\theta) \, \int f(y;\theta) \, \, dy \right).$$

The first integral is simply the expectation of Y and the second is the integral of the probability density function of Y, and hence

$$0 = \frac{1}{\phi} \left(\mathbf{E}[Y] - b'(\theta) \right)$$

which implies that

$$E[Y] = b'(\theta), \tag{3.10}$$

which proves the first part of the proposition.

Differentiating Equation 3.9 by parts and then using the definition of the exponential family to simplify again yields

$$0 = \int \left\{ -\frac{b''(\theta)}{\phi} + \left[\frac{y - b'(\theta)}{\phi} \right]^2 \right\} f(y; \theta) \ dy$$

and using Equation 3.10 gives,

$$0 = -\frac{b''(\theta)}{\phi} + \int \left[\frac{y - \mathbf{E}[Y]}{\phi} \right]^2 f(y; \theta) \ dy$$
$$0 = -\frac{b''(\theta)}{\phi} + \frac{\mathbf{Var}[Y]}{\phi^2}$$

which implies that

$$Var[Y] = \phi \ b''(\theta).$$

which proves the second part of the proposition.

Together, these two results allow us to write down the expectation and variance for any random variable once we have shown that it is a member of the exponential family.

Example: Poisson and normal distribution moments

Table 3.5: Summary of moment calculations via exponential family properties

				E[Y] =		Var[Y] =
	heta	$b(\theta)$	ϕ	$b'(\theta)$	$b''(\theta)$	$b''(\theta)\phi$
Poisson, $Po(\lambda)$	$\log \lambda$	e^{θ}	1	$e^{\theta} = \lambda$	e^{θ}	$e^{\theta} \times 1 = \lambda$
Normal, $N(\mu, \sigma^2)$	μ	$\theta^2/2$	σ^2	$\theta = \mu$	1	$1 \times \sigma^2 = \sigma^2$

3.5 The systematic part of the model

The second part of the generalized linear model, the linear predictor, is given in as $\eta = \sum_{j=1}^{p} \beta_{j} x_{j}$, where x_{j} is the jth explanatory variable (with $x_{1} = 1$ for the intercept). Now, for each observation y_{i} , $i = 1, \ldots, n$, the explanatory variables may differ. To make explicit this dependence on i, we write:

$$\eta_i = \sum_{i=1}^p \beta_j x_{ij}, \tag{3.11}$$

where x_{ij} is the value of the jth explanatory variable on individual i (with $x_{i1} = 1$). Rewriting this in matrix notation:

$$\eta = X\beta,
\tag{3.12}$$

where now $\eta = (\eta_1, \dots, \eta_n)$ is a vector of linear predictor variables, $\beta = (\beta_1, \dots, \beta_p)$ is a vector of regression parameters, and X is the $n \times p$ design matrix.

Recall from Section 1.4 and Section 2.2 that we are concerned with two kinds of explanatory variable:

- Quantitative for example, $x \in (-\infty, \infty)$ etc.
- Qualitative for example, $x \in \{A, B, C\}$ etc.

As discussed in Section 2.4, each quantitative variable is represented in X by an $n \times 1$ column vector. Each qualitative variable, with k+1 levels, say, is represented by a dummy $n \times k$ matrix (one column, usually the first, being dropped to avoid identification problems) of 0's and 1's.

3.6 The link function

In Section 3.2 we saw that the random part of an observation, y, might be described by a member of the exponential family. We also saw, that the systematic part of y might be described using a linear predictor, η , of the explanatory variables. Further, we introduced the notion of a link function $\eta = g(\mu)$ to bring these two parts together, where μ is the mean of y.

Occasionally, the choice of link function $g(\mu)$ is motivated by theory underlying the data at hand. For example, in a dose–response setting, the appropriate model might be motivated by the solution to a set of partial differential equations describing the flow through the body of a dose of a drug.

When there is no compelling underlying theory, however, we typically choose a link function that will transform a restricted range of the dependent variable onto the whole real line. For example, when observations are typically positive, so we have $\mu > 0$, we might choose the logarithmic link:

$$g(\mu) = \log(\mu). \tag{3.13}$$

When observations are binomial counts from B(m,p), $0 , with mean <math>\mu = mp$, we might choose the logit link from

$$\eta = g(\mu) = \text{logit}(\mu/m) = \text{logit}(p) = \log\{p/(1-p)\}$$
(3.14)

or the *probit* link which is the inverse of the cumulative distribution function of the N(0,1) distribution:

$$\eta = g(\mu) = \Phi^{-1}(\mu/m) = \Phi^{-1}(p), \tag{3.15}$$

or the complementary log-log (cloglog) link:

$$\eta = g(\mu) = \log(-\log(1 - \mu/m)) = \log(-\log(1 - p)), \tag{3.16}$$

or the *cauchit* link which is the inverse of the cumulative distribution function of the Cauchy (t_1) distribution:

$$\eta = g(\mu) = \tan(\pi(\mu/m - \frac{1}{2})) = \tan(\pi(p - \frac{1}{2})).$$
(3.17)

Figure 3.4 shows these link functions for proportions fitted to the beetle mortality data. This demonstrates that the logit and probit links are very similar, that the complementary log-log link fits these data slightly better in the extremes, but that the cauchit link fits these data quite poorly in the extremes.

A mathematically and computationally convenient choice of link function $g(\mu)$ can be constructed by setting:

$$\theta = \eta, \tag{3.18}$$

where θ is the canonical parameter of the exponential family as defined in Equation 3.3. Then, Equation 3.8 shows that the mean μ is a function of θ and therefore, Equation 3.18 indirectly provides a link between μ and η . That is, Equation 3.18 implicitly defines a link function $\eta = g(\mu)$. But what is the form of this $g(\cdot)$?

From Equation 3.8,

$$\mu = b'(\theta)$$
.

So, provided function $b'(\cdot)$ has an inverse $(b')^{-1}(\cdot)$, we may write

$$\theta = (b')^{-1}(\mu). \tag{3.19}$$

Now, from Equation 3.5, $g(\mu) = \eta$, so using Equation 3.18:

$$g(\mu) = \theta = (b')^{-1}(\mu),$$
 (3.20)



Figure 3.4: Dose–response curves fitted to the beetle mortality data from Table 1.1 with different choices of link function.

from Equation 3.19. This makes explicit the $g(\mu)$ that is implicitly asserted by Equation 3.18. The function produced form Equation 3.20 is called the *canonical* link function.

Proposition 3.2. For the canonical link function,

$$g'(\mu) = 1/b''(\theta).$$

Proof: From Proposition 3.1, $\mu = E[Y] = b'(\theta)$, so

$$\frac{\mathrm{d}\mu}{\mathrm{d}\theta} = b''(\theta).$$

From Equation 3.20, for the canonical link function, we have $\theta = g(\mu)$, so

$$\frac{\mathrm{d}\theta}{\mathrm{d}\mu} = g'(\mu).$$

Now $d\theta/d\mu = (d\mu/d\theta)^{-1}$ and hence

$$g'(\mu) = 1/b''(\theta).$$

Which proves the proposition.

Example: Poisson canonical link function

For the Poisson distribution $Po(\lambda)$, we have from Table 3.2 that $b(\theta) = e^{\theta}$. Therefore,

$$b'(\theta) = e^{\theta}$$
.

so the inverse of function $b'(\cdot)$ exists and is the inverse of the exponential function, which is the logarithmic function. Then, applying Equation 3.20

$$q(\mu) = \log(\mu)$$

Thus the canonical link for the Poisson distribution is log.

Example: Normal canonical link function

For the Normal distribution $N(\mu, \sigma^2)$, we have from Table 3.4 that $b(\theta) = \theta^2/2$. Therefore

$$b'(\theta) = \theta$$

so the inverse of function $b'(\cdot)$ exists and is the inverse of the identity function, which is the identity function. (The identity function is that which maps a value onto itself.) Then, applying Equation 3.20,

$$g(\mu) = \mu$$
.

Thus the canonical link for the Normal distribution is the identity function.

For many models, μ has a restricted range, but we would like η to have unlimited range. It turns out, for several members of the exponential family, that the canonical link function provides η with unlimited range. However, Table 3.6 shows that this is not always so.

Table 3.6: Canonical link functions and their ranges (see McCullagh and Nelder, 2nd Edn., p291 with †binomial distribution with index m and mean μ and ‡gamma distribution with mean μ (see Exercises for details).

				Canonical	
f(y)	Range of μ	b(heta)	$\mu = b'(\theta)$	link, $g(\mu)$	Range of η
Normal	$(-\infty, \infty)$	$\frac{1}{2}\theta^2$	θ	μ	$(-\infty,\infty)$
Poisson	$(0, \infty)$	e^{θ}	$e^{ heta}$	$\log \mu$	$(-\infty, \infty)$
Binomial†	(0,m)	$m\log(1-e^{\theta})$	$m/(1+e^{-\theta})$	$\operatorname{logit}(\mu/m)$	$(-\infty, \infty)$
Gamma‡	$(0, \infty)$	$-\log(-\theta)$	$-\theta^{-1}$	$-\mu^{-1}$	$(-\infty,0)$

Why is the canonical link function Equation 3.20 convenient? The assertion Equation 3.18 means that, in the exponential-family formula Equation 3.3, we can simply substitute the linear predictor

$$\eta = \sum_{j} \beta_{j} x_{j}$$

from Equation 3.4 in place of θ , to give:

$$f(y; \mathbf{x}, \beta, \phi) = \exp\left\{\frac{y\left[\sum_{j} \beta_{j} x_{j}\right] - b\left(\left[\sum_{j} \beta_{j} x_{j}\right]\right)}{\phi} + c(y, \phi)\right\},\tag{3.21}$$

where
$$\mathbf{x} = \{x_i, j = 1, ..., p\}$$
 and $\beta = \{\beta_i, j = 1, ..., p\}$.

Further, suppose we have n independent observations, $\{y_i, i=1,\ldots,n\}$. As discussed in Section 3.5, the explanatory variables (x_1,\ldots,x_p) will depend on i, and so η will also depend on i. Therefore, we attach subscript i to y and to each x_j , giving:

$$f(y_i; \mathbf{x}_i, \beta, \phi) = \exp\left\{\frac{y_i \left[\sum_j \beta_j x_{ij}\right] - b\left(\left[\sum_j \beta_j x_{ij}\right]\right)}{\phi} + c(y_i, \phi)\right\}. \tag{3.22}$$

where $\mathbf{x}_i = \{x_{ij}, j = 1, \dots, p\}$ and $\beta = \{\beta_j, j = 1, \dots, p\}.$

By independence, the joint distribution of all observations $\mathbf{y} = \{y_i, i = 1, ..., n\}$, with design matrix $X = \{x_{ij}, i = 1, ..., n; j = 1, ..., p\}$, is:

$$f(\mathbf{y}; X, \beta, \phi) = \prod_{i=1}^{n} f(y_i; \theta_i, \phi),$$

so

$$\log f(\mathbf{y}; X, \beta, \phi) = \sum_{i=1}^{n} \log f(y_i; \theta_i, \phi)$$

then substituting in using Equation 3.22 gives

$$\log f(\mathbf{y}; X, \beta, \phi) = \sum_{i=1}^{n} \left\{ \frac{y_i \left[\sum_{j} \beta_j x_{ij} \right] - b \left(\left[\sum_{j} \beta_j x_{ij} \right] \right)}{\phi} + c(y_i, \phi) \right\}$$

and finally simplifying to give

$$\log f(\mathbf{y}; X, \beta, \phi) = \frac{\sum_{j} \beta_{j} S_{j} - \sum_{i} b\left(\left[\sum_{j} \beta_{j} x_{ij}\right]\right)}{\phi} + \sum_{i} c(y_{i}, \phi)$$
(3.23)

where

$$S_j = \sum_{i=1}^n y_i x_{ij}.$$

Thus, in the log-likelihood Equation 3.23, it is only the first term that involves both the observations $\mathbf{y} = \{y_i, i = 1, ..., n\}$ and the parameters $\beta = \{\beta_j, j = 1, ..., p\}$, and this term depends on the observations only through the statistics $\mathbf{S} = \{S_j, j = 1, ..., p\}$ – these are called *sufficient statistics*, and their appearance in Equation 3.23 confers both theoretical and practical advantages.

3.7 Exercises

- 3.1 Consider the beetle data again, see Table 1.1, but suppose that we had only been given the y values, that is the number killed, and misinformed that each came from a sample of size 62. Further, suppose that we did not know that different doses had been used. That is, we where given data: $\mathbf{y} = \{6, 13, 18, 28, 52, 53, 61, 60\}$ and led to believe that the model $Y \sim \text{Bin}(62, p)$ was appropriate. Use the given data to estimate p. Then, calculate the fitted probabilities and superimpose them on a histogram of the data. [Hint: see the code chunk used to create Figure 3.2.]
- 3.2 Verify that, in general, if $q = 1/(1 + e^{-x})$ then $x = \log(q/(1-q))$ and then for the binomial distribution, $Y \sim \text{Bin}(m, p)$, show that

$$-m\log(1-p) = m\log(1+e^{\theta})$$

where $\theta = \text{logit } p = \log(p/(1-p))$.

3.3 Suppose that Y has a gamma distribution with parameters α and β , that is $Y \sim \text{Gamma}(\alpha, \beta)$, with probability density function

$$f(x;\alpha,\beta) = \frac{\beta^{\alpha} x^{\alpha-1} e^{-\beta x}}{\Gamma(\alpha)} \qquad x > 0; \alpha,\beta > 0.$$

Write this in the form of the exponential family and clearly identify θ , ϕ , $b(\theta)$ and $c(y, \phi)$ – as in Table 3.2 and Table 3.3.

3.4 Express the geometric distribution, $Y \sim \text{Geom}(p), 0 , with probability mass function$

$$f(y) = (1-p)^{y-1}p;$$
 $y = 1, 2, 3...$

as an exponential family distribution, possibly with a scale parameter.

- 3.5 Prove Proposition 3.1 assuming that Y follows a discrete distribution. Verify the results for the Poisson in Table 3.5 and then derive similar results for the binomial, $Y \sim \text{Bin}(n,p)$. From these results, what are the mean and variance of Y.
- 3.6 Use the properties of exponential families to find the expectation and variance of each of the geometric and gamma distributions defined above.

[Hint for the gamma, treat $1/\alpha$ as a scale parameter and let θ be a suitable function of λ and α .]

3.7 Using Proposition 3.2, verify the canonical link function, $g(\mu)$, for the binomial and gamma distributions shown in Table 3.6.

4 GLM Estimation

Warning

Please note that to make these lecture notes available on time, there has not been sufficient time to check the following sections. This, in particular, may mean that some of the cross-referencing is not accurate, as well as small typos. Further, the final few sections are not yet complete.

Throughout this module we use the principle of maximum likelihood estimation (MLE) to estimate model parameters and will consider two cases.

4.1 The identically distributed case

4.1.1 Maximum likelihood estimation

Suppose we have n independent and identically distributed (i.i.d.) observations y_i , i = $1, \ldots, n$, where each y_i is sampled from the same exponential family density

$$f(y_i;\theta,\phi) = \exp\left\{\frac{y_i\theta - b(\theta)}{\phi} + c(y_i,\phi)\right\}, \tag{4.1}$$

for $i=1,\ldots,n$. In this case, the canonical parameter θ does not depend on i.

By independence, the joint distribution of all the observations $\mathbf{y} = \{y_i, i = 1, \dots, n\}$ is:

$$f(\mathbf{y}; \theta, \phi) = \prod_{i=1}^{n} f(y_i; \theta, \phi).$$

So, taking logs and then substituting for the probability function using the exponential family form, Equation 3.3, gives

$$\log f(\mathbf{y}; \theta, \phi) = \sum_{i=1}^{n} \log f(y_i; \theta, \phi) = \sum_{i=1}^{n} \left[\frac{y_i \theta - b(\theta)}{\phi} + c(y_i, \phi) \right].$$

Regarding the observations y as constants (which they are, once we have them) and the scale parameter ϕ as a fixed nuisance parameter (whose value we may not know), the loglikelihood as a function of the parameter θ of interest is:

$$l(\theta; \mathbf{y}, \phi) = n \left(\frac{\bar{y} \theta - b(\theta)}{\phi} \right) + \text{constant},$$
 (4.2)

where $\bar{y} = \sum y_i/n$.

We estimate θ by maximizing the log likelihood – i.e. given the data \mathbf{y} , we estimate the value of θ to be that value for which the likelihood, and hence the log-likelihood, is greatest.

We maximize the log-likelihood by differentiating it and setting it to zero:

$$\frac{dl(\boldsymbol{\theta};\mathbf{y},\boldsymbol{\phi})}{d\boldsymbol{\theta}} = n\left(\frac{\bar{y} - b'(\boldsymbol{\theta})}{\boldsymbol{\phi}}\right)$$

and hence the MLE for θ , which we denote $\hat{\theta}$, satisfies

$$b'(\hat{\theta}) = \bar{y} \tag{4.3}$$

and hence

$$\hat{\theta} = (b')^{-1}(\bar{y}). \tag{4.4}$$

Further, we showed in Proposition 3.1 that $E[Y] = \mu = b'(\theta)$ and if we let $\hat{\mu}$ denote the MLE of μ , then $\hat{\mu} = b'(\hat{\theta})^1$, hence we have $\hat{\mu} = \bar{y}$. So we find that $\hat{\theta}$ is the value of θ for which the theoretical mean $\hat{\mu} = b'(\hat{\theta})$ matches the sample mean \bar{y} .

Example: MLE of the Poisson distribution

For the Poisson distribution, Po(λ), we have found that $b(\theta) = e^{\theta}$ and therefore $b'(\theta) = e^{\theta}$. Hence, the MLE of natural parameter θ is found as the solution of $b'(\hat{\theta}) = e^{\hat{\theta}} = \bar{y}$, that is $\hat{\theta} = \log(\bar{y})$.

4.1.2 Estimation accuracy

For our i.i.d. sample y_i , $i=1,\ldots,n$, we have $b'(\hat{\theta})=\hat{\mu}=\bar{y}$. Let θ_0 be the true value of θ with corresponding mean μ_0 , i.e.

$$b'(\theta_0) = \mu_0. \tag{4.5}$$

How accurate is $\hat{\theta}$? We know that

$$E[\bar{Y}] = E\left[\frac{1}{n}\sum_{i=1}^{n}Y_{i}\right] = \frac{1}{n}\sum_{i=1}^{n}E[Y_{i}] = \mu_{0} = b'(\theta_{0}), \tag{4.6}$$

using Equation 4.5, and

$$\operatorname{Var}[\bar{Y}] = \operatorname{Var}\left[\frac{1}{n}\sum_{i=1}^{n}Y_{i}\right] = \frac{1}{n^{2}}\sum_{i=1}^{n}\operatorname{Var}[Y_{i}]$$

because the observations are independent. Then, using the result Equation 3.8

$$\operatorname{Var}[\bar{Y}] = \frac{1}{n} b''(\theta_0)\phi. \tag{4.7}$$

¹Using the result that the MLE of any function of a parameter is given by the same function applied to the MLE of the parameter.

We can use Taylor's theorem to expand $b'(\hat{\theta})$ about θ_0 :

$$\bar{y} = b'(\hat{\theta}) \approx b'(\theta_0) + (\hat{\theta} - \theta_0)b''(\theta_0),$$

which implies that

$$(\hat{\theta} - \theta_0) \approx b''(\theta_0)^{-1} \{b'(\hat{\theta}) - b'(\theta_0)\} = b''(\theta_0)^{-1} (\bar{Y} - \mu_0), \tag{4.8}$$

using Equation 4.3 and Equation 4.5. We can use Equation 4.8 to get approximations to the mean and variance of $\hat{\theta}$:

$$\mathrm{E}[\hat{\theta} - \theta_0] \approx b''(\theta_0)^{-1} \mathrm{E}[\bar{Y} - \mu_0] = 0,$$

using Equation 4.6, so

$$E[\hat{\theta}] \approx \theta_0, \tag{4.9}$$

and

$$\mathrm{Var}(\hat{\theta}) \approx \mathrm{E}\left[(\hat{\theta} - \theta_0)^2\right]$$

using Equation 4.9,

$$\mathrm{Var}(\hat{\theta}) \approx \mathrm{E}\left[\left(b''(\theta_0)^{-1}(\bar{Y} - \mu_0)\right)^2\right]$$

using Equation 4.8,

$$\mathrm{Var}(\hat{\theta}) \approx \left(b''(\theta_0)\right)^{-2} \mathrm{Var}[\bar{Y}]$$

using Equation 4.6,

$$Var(\hat{\theta}) = \frac{\phi}{n \ b''(\theta_0)} \tag{4.10}$$

using Equation 4.7.

Thus we see that the first two derivatives of $b(\theta)$ play a key role in inference.

Example: Accuracy for the Poisson distribution

For the Poisson distribution, $Po(\lambda)$, we have found that $\hat{\theta} = \log(\bar{Y})$. Using Equation 4.9 we know that $\hat{\theta}$ is, at least, approximately unbiased. Then, using that $\phi = 1$ (Table 3.2), $b'(\theta) = e^{\theta}$ (Table 3.6) hence $b''(\theta) = e^{\theta}$, and using Equation 4.10, leads to the result

$$\operatorname{Var}(\hat{\theta}) = \frac{\phi}{n \ b''(\theta_0)} = \frac{1}{ne^{\theta_0}}.$$

4.2 The general case

4.2.1 MLE Estimation

Suppose that now the n independent observations $\{y_i, i = 1, ..., n\}$ are not identically distributed. They are, however, sampled from the same exponential family density but with differing parameters, that is

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

for $i=1,\ldots,n$. In this case, the canonical parameter does depend on i – but we assume that the scale parameter ϕ does not – and let $\theta = \{\theta_i, i=1,\ldots,n\}$.

In most applications, we are not interested in estimation of θ but instead we are interested in the linear predictor parameters $\beta = \{\beta_1, \dots, \beta_p\}$. Note, however, that each θ_i will depend on all β_1, \dots, β_p . This is most obvious for the canonical parameter case where a convenient choice of link function is obtained using $\theta = \eta = X\beta$, hence $\theta_i = \mathbf{x}_i^T \beta = \sum_{j=1}^p x_{ij}\beta_j$ and each θ_i clearly depends on all β_1, \dots, β_p .

The principle of maximum likelihood will be used to estimate the model parameters β . Using the independence of $y_i, i = 1, ..., n$, given the parameters β , the likelihood function is:

$$L(\beta; \mathbf{y}, \phi) = f(\mathbf{y}; X, \beta, \phi) = \prod_{i=1}^{n} f(y_i; \mathbf{x}_i, \beta, \phi)$$
(4.11)

and the log-likelihood by

$$l(\beta; \mathbf{y}, \phi) = \log L(\beta; \mathbf{y}, \phi) = \sum_{i=1}^{n} \log f(y_i; \mathbf{x}_i, \beta, \phi). \tag{4.12}$$

Then we wish to find the value of β which maximizes the log-likelihood function,

$$\hat{\beta} = \max_{\beta} l(\beta; \mathbf{y}, \phi).$$

For generalized linear models there is usually no closed-form expression for the MLE $\hat{\beta}$. Instead, an iterative approach based on the Newton–Raphson algorithm is usually adopted.

For the normal linear regression model, however, that is where the exponential family is Gaussian and the link function $g(\mu)$ is the identity function, we have the familiar closed-form expression

$$\hat{\boldsymbol{\beta}} = \left(X^T X\right)^{-1} X^T \mathbf{y}. \tag{4.13}$$

4.2.2 The score function and Fisher information

We define the *score function*:

$$U(\beta) = \frac{\partial l(\beta; \mathbf{y}, \phi)}{\partial \beta},\tag{4.14}$$

which is a $p \times 1$ vector. We define the observed Fisher information:

$$I(\beta) = -\frac{\partial U(\beta)}{\partial \beta^T} = -\frac{\partial^2 l(\beta; \mathbf{y}, \phi)}{\partial \beta \partial \beta^T}, \tag{4.15}$$

which is a $p \times p$ matrix whose (j,k)th element is: $-\frac{\partial^2 l(\beta;\mathbf{y},\phi)}{\partial \beta_j \partial \beta_k}$. We also define the expected Fisher information:

$$J(\beta) = \mathbf{E} \left[-\frac{\partial^2 l(\beta; \mathbf{y}, \phi)}{\partial \beta \partial \beta^T} \right], \tag{4.16}$$

which is also a $p \times p$ matrix.

Proposition 4.1. With definitions Equation 4.14, Equation 4.15, Equation 4.16 above,

- $E[U(\beta)] = 0$,
- $J(\beta) = E[U(\beta)U^T(\beta)].$

Proof We give the proof for continuous random variables. For the discrete case, replace integration by sums – see Exercises.

To start the proof, notice that we can re-write the joint density of the data given the parameters as

$$f(\mathbf{y}; X, \beta, \phi) = L(\beta; \mathbf{y}, \phi) = \exp\{l(\beta; \mathbf{y}, \phi)\}\$$

and then

$$1 = \int f(\mathbf{y}; X\beta, \phi) d\mathbf{y} = \int \exp\{l(\beta; \mathbf{y}, \phi)\} d\mathbf{y},$$

where $d\mathbf{y}=dy_1\cdots dy_n$. Differentiating this with respect to $\boldsymbol{\beta}=(\beta_1,\dots,\beta_p)^T$ gives:

$$0 = \int \frac{\partial l(\beta; \mathbf{y}, \phi)}{\partial \beta} \exp\{l(\beta; \mathbf{y}, \phi)\} d\mathbf{y}$$

$$= \int U(\beta) f(\mathbf{y}; X, \beta, \phi) d\mathbf{y}$$

$$= \operatorname{E} [U(\beta)]. \tag{*}$$

Proving the first part.

Next, differentiating (\star) by parts with respect to β^T ,

$$0 = \int \frac{\partial U(\beta)}{\partial \beta^{T}} f(\mathbf{y}; X, \beta, \phi) + \frac{\partial l(\beta; \mathbf{y}, \phi)}{\partial \beta} \frac{\partial l(\beta; \mathbf{y}, \phi)}{\partial \beta^{T}} f(\mathbf{y}; X, \beta, \phi) d\mathbf{y}$$
$$= \int -I(\beta) f(\mathbf{y}; X, \beta, \phi) d\mathbf{y} + \int U(\beta) U^{T}(\beta) f(\mathbf{y}; X, \beta, \phi) d\mathbf{y}$$
$$= -J(\beta) + \mathbb{E} \left[U(\beta) U^{T}(\beta) \right].$$

proving the second part.

Proposition 4.2. Under some regularity conditions, the MLE $\hat{\beta}$ of β has the following asymptotic properties:

- E(β) = β; i.e. β is unbiased for β.
 Var(β) = J⁻¹(β).
 β follows a p-dimensional Normal distribution.

Combining these three statements we have that asymptotically

$$\hat{\boldsymbol{\beta}} \sim N_p(\boldsymbol{\beta}, J^{-1}(\boldsymbol{\beta})). \tag{4.17}$$

Proof: Omitted. Similar to the proof using Taylor's theorem in Section 4.1.

From Equation 4.17, variances $Var(\hat{\beta}_k)$, standard errors $se(\hat{\beta}_k)$ and correlations between parameter estimates $Corr(\hat{\beta}_k, \hat{\beta}_h)$ can be estimated.

4.2.3 The saturated case

Again we assume the observations y_i , $i=1,\ldots,n$ are independent but now we assume that y_i is sampled from an exponential family probability function with canonical parameter θ_i ,

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

for $i=1,\ldots,n$. We can form the log-likelihood in the usual way to give

$$l(\theta; \mathbf{y}, \phi) = \sum_{i=1}^n \log f(y_i; \theta_i, \phi) = \sum_{i=1}^n \left[\frac{y_i \theta_i - b(\theta_i)}{\phi} + c(y_i, \phi) \right]$$

and so we find the the MLE of θ using

$$\hat{\theta}_i = (b')^{-1}(y_i), \quad i = 1, \dots, n.$$

Note that each parameter is only a function of the corresponding observation.

Further, from Proposition 3.1, $E[Y_i] = \mu_i = b'(\theta_i)$ and if we again let $\hat{\mu}_i$ denote the MLE of μ_i , then $\hat{\mu}_i = b'(\hat{\theta}_i)$, hence we have $\hat{\mu}_i = y_i$. Thus we see that, under the saturated model, the mean of the distribution of y_i is estimated to be equal to y_i itself. That is, the data are fitted exactly by the model. Of course this model is quite useless for explanation or prediction, since it misinterprets random variation as systematic variation. Nevertheless, the saturated model is useful as a benchmark for comparing models, as we will see later.

It is worth noting that the same situation can occur even when modelling in terms of the regression parameters β_j , $j=1,\ldots,p$. When $p\geq n$, and if the covariates are linearly independent, each θ_i can take on any value independently of the others and so estimating the β_j 's is equivalent to estimating the θ_i 's. That is we are also considering the saturated or full model. This highlights the danger of putting too many covariates into the model. There is a big literature on how to deal with more parameters then data using techniques of regularized regression.

4.3 Model deviance

The *deviance* is a quantity we use to assess the fit of a model to the data. Let M be a model of interest with fitted parameters $\hat{\theta}$ and corresponding fitted values $\hat{\mu}$. Also consider the saturated model with fitted parameters $\tilde{\theta}$ and fitted values $\tilde{\mu}$.

The deviance of model M is defined as twice the difference between the log-likelihood of the saturated model, $l(\tilde{\theta}; \mathbf{y}, \phi)$, and the log-likelihood of model M, $l(\hat{\theta}; \mathbf{y}, \phi)$, multiplied by ϕ ,

$$D = 2\phi \left\{ l(\tilde{\theta}; \mathbf{y}, \phi) - l(\hat{\theta}; \mathbf{y}, \phi) \right\}$$

$$= \begin{cases} \sum_{i=1}^{n} (y_i - \hat{\mu}_i)^2 = \text{Residual sum of squares} & \text{Normal} \\ 2\sum_{i=1}^{n} \left\{ y_i \log \left(\frac{y_i}{\hat{\mu}_i} \right) + (m_i - y_i) \log \left(\frac{m_i - y_i}{m_i - \hat{\mu}_i} \right) \right\} & \text{Binomial} \\ 2\sum_{i=1}^{n} \left\{ y_i \log \left(\frac{y_i}{\hat{\mu}_i} \right) - y_i + \hat{\mu}_i \right\} & \text{Poisson} \end{cases}$$

$$(4.18)$$

Note that Dobson, and others, call $D^* = D/\phi = 2\{l(\tilde{\theta}; y, \phi) - l(\hat{\theta}; y, \phi)\}$ the scaled deviance.

We now consider two situations:

Scale parameter ϕ **known.** For some data-types (e.g. Poisson, Binomial), we know $\phi = 1$. Consider two nested models M_1 and M_2 with r_1 and r_2 parameters respectively where the parameters in M_1 are a subset of those in M_2 and hence $r_1 < r_2$. Further, let D_1 and D_2 be the deviances of model M_1 and M_2 respectively.

Then, asymptotically,

- the log likelihood-ratio statistic $D_1 D_2 \sim \chi^2_{r_2 r_1}$ can be used to test the importance of the extra parameters in M_2 not included in M_1 ;
- a goodness-of-fit test for M_2 can be done based on $D_2 \sim \chi^2_{n-r_2}$.

The quality of the approximations involved depends on there being a large amount of *in-formation*, for example, large counts for Binomial and Poisson data, or a large sample size for Normal data.

Scale parameter ϕ **unknown.** For some data-types (e.g. Normal, Gamma), ϕ is not known (typically $\phi = \sigma^2$). We must find a model M_3 big enough to be believed, then estimate ϕ by the residual mean square:

$$\hat{\phi} = \frac{D_3}{n - r_3}. (4.19)$$

Then test M_1 against M_2 using

$$\mathbf{F} = \frac{(D_1 - D_2)/(r_2 - r_1)}{\hat{\phi}} = \frac{(D_1 - D_2)/(r_2 - r_1)}{D_3/(n - r_3)} \tag{4.20}$$

with

$${\bf F} \sim F_{r_2-r_1,n-r_3}. \eqno(4.21)$$

So if the observed value of the statistic Equation 4.20 was within the upper (say 5%) tail of the F-distribution Equation 4.21, we would infer that Model M_2 is better than Model M_1 .

4.4 Model residuals

Consider a generalized linear model with observed values $y_i, i = 1, ..., n$ and fitted values $\hat{\mu}_i$. Then the raw or response residuals are defined by

$$e_i^{\text{raw}} = y_i - \hat{\mu}_i$$
.

More useful are the *standardized* or *Pearson* residuals defined by

$$e_i^{\text{std}} = e_i^{\text{P}} = \frac{y_i - \hat{\mu}_i}{\sqrt{b''(\theta_i)}}.$$

Recall from Equation 3.8 that $Var(Y_i) = \phi b''(\theta_i)$.

Deviance residuals are defined so that the sum of squared deviance residuals equals the total deviance. Thus we set

$$e_i^{\text{dev}} = \text{sign}(y_i - \hat{\mu}_i) \sqrt{d_i},$$

where d_i is the contribution of observation i to the deviance, D. For example, when y_i has a Poisson distribution with estimated mean $\hat{\mu}_i$, we have

$$e_i^{\text{dev}} = \text{sign}(y_i - \hat{\mu}_i) \sqrt{2 \left[y_i \log \left(\frac{y_i}{\hat{\mu}_i} \right) - y_i + \hat{\mu}_i \right]}.$$

Residuals are useful for assessing the overall fit of a model to the data, and for identifying where the model might need to be improved.

4.5 Fitting generalized linear models in R

4.5.1 GLM-related R commands

The function used to fit a generalized linear model in \mathbf{R} is

Let x,y,z,a,b,c, ... be a set of vectors all of the same length n (perhaps read in from a data file using the read.table and attach commands). If a,b,c are qualitative variables, then they first need to be declared as factors by a = as.factor(a), etc.

The formula argument of glm specifies the required model in compact notation, e.g. $y \sim x*a$ or $y \sim x + z*a$ where \sim , +, * have the same meaning as in Section 2.5.

The family argument specifies which exponential family is to be used. We shall use gaussian, poisson and binomial; gaussian is the default. Other options are available; see help(family) for further information.

Along with the family, a link function can be specified. The possible choices are:

- gaussian "identity" (default)
- poisson "log" (default), "sqrt", "identity"
- binomial "logit" (default), "probit", "cloglog".

R assumes the default options unless we state otherwise. For example,

```
glm(y \sim a+b) \# Gaussian errors, identity link <math>glm(y \sim a+b, poisson) \# Poisson errors, log link <math>glm(y \sim a+b, poisson("sqrt")) \# Poisson errors, sqrt link
```

Note that for the binomial case, the response variable should be an $n \times 2$ matrix ym, say, not a vector, where the first column contains the numbers of successes and the second column the numbers of failures, for example:

```
glm(ym \sim a+b, binomial) \# binomial errors, logit link
```

To extract information about a fitted generalized linear model, it is best to store the result of glm as a variable and then to use the following functions:

- To fit a GLM and store the result in y.glm (for example):
 y.glm = glm(y ~ a*b, poisson("sqrt"))
- To print various pieces of information including deviance residuals, parameter estimates and standard errors, deviances, and (if specified) correlations of parameter estimates:

```
summary(y.glm, correlation=T)
```

- To print the anova table of the fitted model: anova(y.glm)
- To print the deviance of the fitted model: deviance(y.glm)
- To print the residual degrees of freedom of the fitted model: df.residual(y.glm)
- To print the vector of fitted values under the fitted model: fitted.values(y.glm)
- To print the residuals from the fitted model:

```
residuals(y.glm, type)
```

Note: type should be "deviance" (default), "pearson", or "response"

- To print the parameter estimates from the fitted model: coefficients(y.glm)
- To print the design matrix for a specified model formula: model.matrix(y ~ a*b)

The functions summary, anova, and possibly model.matrix are the most useful for printing out information about the fitted model. The results of the other functions can be saved as variables for further computation, if desired.

4.5.2 Example of fitting Poisson GLM in R

Here is a toy example of \mathbf{R} commands for modelling a response in terms of two qualitative explanatory variables (that is factors). The model assumes the data are Poisson-distributed and uses the logarithmic link function.

```
Call:
glm(formula = y ~ a + b, family = poisson)

Deviance Residuals:
    Min    1Q    Median    3Q    Max
```

```
-1.8740 -0.6834 0.1287 0.7151 1.5956
```

Coefficients:

Estimate Std. Error z value Pr(>|z|)1.1408 0.3351 3.404 0.000663 *** (Intercept) 0.3522 -0.867 0.385934 a2 -0.3054 0.1466 0.3132 0.468 0.639712 a3 0.4568 0.2932 1.558 0.119269 a4 0.3162 2.898 0.003761 ** b2 0.9163 b3 0.9445 0.3150 2.999 0.002712 **

Signif. codes: 0 '*** 0.001 '** 0.01 '* 0.05 '.' 0.1 ' ' 1

(Dispersion parameter for poisson family taken to be 1)

Null deviance: 31.725 on 11 degrees of freedom Residual deviance: 13.150 on 6 degrees of freedom

AIC: 68.227

Number of Fisher Scoring iterations: 5

Correlation of Coefficients:

(Intercept) a2 a3 a4 b2 a2 -0.45 a3 -0.50 0.48 a4 -0.54 0.51 0.57 b2 -0.67 0.00 0.00 0.00 b3 -0.68 0.00 0.00 0.00 0.72

Analysis of Deviance Table

Model: poisson, link: log

Response: y

Terms added sequentially (first to last)

Df Deviance Resid. Df Resid. Dev NULL 11 31.725 a 3 6.2793 8 25.445 b 2 12.2947 6 13.150

[1] 13.15047

[1] 6

```
1 2 3 4 5 6 7 8
3.129412 2.305882 3.623529 4.941176 7.823529 5.764706 9.058824 12.352941
9 10 11 12
8.047059 5.929412 9.317647 12.705882
```

4.6 Exercises

- 4.1 Use Equation 4.4 to obtain estimation equations for the natural parameter θ , based on a sample $\mathbf{y} = \{y_1, \dots, y_n\}$, for each of the following situations:
 - (a) the binomial, $Y \sim Bin(n, p)$,
 - (b) the geometric, $Y \sim Ge(p)$,
 - (c) the exponential, $Y \sim \text{Exp}(\lambda)$.

Are these estimators unbiased for θ ? What is the variance of the estimator in each case?

- 4.2 For a sample of size n from the normal distribution, $Y \sim N(\mu, \sigma^2)$, how do the results produced using Equation 4.9 and Equation 4.10 compare with the familiar results $\hat{\mu} = \bar{y}$, $\mathrm{E}[\hat{\mu}] = \mu$, and $\mathrm{Var}[\hat{\mu}] = \sigma^2/n$?
- 4.3 For the normal linear regression model, $\mathbf{Y} = X\beta + \epsilon$ where $\epsilon \sim N_n(0, \sigma^2 I_n)$ use the principle of maximum likelihood to show that the MLE has the closed form given in Equation 4.13.

Further questions to be added later

5 Modelling Proportions

5.1 Introduction

In this chapter we will focus on applications of generalized linear modelling where the response variable follows a binomial distribution. This can arise when the outcome is binary, that is it can take one of only two possible values, or is the sum of a set of such binary outcomes. These two outcomes record whether some event of interest has occurred or not. In the simplest case, the response variable, B say, is defined as

$$B = \begin{cases} 1 & \text{if the event has occurred} \\ 0 & \text{if the event has not occurred} \end{cases}$$
 (5.1)

and we set Pr(B=1)=p, and hence Pr(B=0)=1-p. This is, of course, the definition of a Bernoulli trial leading to a Bernoulli random variables, $B \sim \text{Bernoulli}(p)$.

Suppose now that there are m similar binary outcomes, B_i , $i=1,\ldots,m$ with $Pr(B_i=1)=p$, and that the total number of times that the event occurred is recorded as the response Y. Assuming that m is known before the trials start, that p is fixed and that the individual Bernoulli trials are independent then Y follows a binomial distribution, $Y \sim B(m,p)$ with probability mass function

$$f(y) = {m \choose y} p^y (1-p)^{m-y}. \tag{5.2}$$

Note that the binomial random variable can be thought of as the sum of i.i.d Bernoulli random variables, $Y = B_1 + \dots + B_m$, if that is helpful.

The Binomial distribution B(m, p) is often described in terms of *success* and *failure* and a binomial distribution in terms of the number of successes in m independent trials, where p is the probability of success in each trial. The term *success* need not correspond to a favourable outcome; it is merely the language traditionally used in connection with this model. For example, success might correspond to death.

Of course, the special case with m=1 reduces to the Bernoulli distribution. Further, if $Y_1 \sim \mathrm{B}(m_1,p)$ and independently $Y_2 \sim \mathrm{B}(m_2,p)$, then $Y_1 + Y_2$ also follows a binomial distribution, $\mathrm{B}(m_1+m_2,p)$ – note that is only valid when p is common.

Recall that the binomial can be re-written in the exponential family form of Equation 3.3 with

$$f(y) = \exp\left\{y \text{ logit } p + m \log(1-p) + \log{m \choose y}\right\},\,$$

with natural or canonical parameter $\theta = \text{logit } p = \log\left(p(1-p)\right)$,, scale parameter $\phi = 1$, $b(\theta) = m\log(1+e^{\theta})$ and $c(y,\phi) = \log\binom{m}{y}$ as in Table 3.3.

5.2 The linear logistic model

Throughout this module we have assumed that a response variable, Y, depends on a set of explanatory variables, $\mathbf{x} = \{x_1, \dots, x_p\}$. In particular, for binomial counts from $\mathbf{B}(m,p)$, $0 with mean <math>\mu = mp$ we set the link function, $g(\mu)$, equal to the linear predictor $\eta = \sum \beta_j x_j$ and hence have

$$g(\mu) = \sum_{i=1}^{p} \beta_j x_j = \mathbf{x}^T \beta$$

where $\beta = \{\beta_1, \dots, \beta_p\}$ are the linear predictor parameters. Recall that the canonical logit link Equation 3.14 for the binomial leads to the systematic part of the model

$$logit(p) = log\left(\frac{p}{1-p}\right) = \mathbf{x}^T \beta \tag{5.3}$$

but that other links function are possible, such as the probit, Equation 3.15, the complementary log-log, Equation 3.16, and the cauchit Equation 3.17.

This model can alternatively be written

$$Y \sim B(m, p), \text{ where } p = \frac{\exp\{\mathbf{x}^T \beta\}}{1 + \exp\{\mathbf{x}^T \beta\}}$$
 (5.4)

which makes the dependence of Y on \mathbf{x} , and β , more explicit.

In general, we might want to consider n independent binomial random variables representing subgroups of the sample with $Y_1 \sim \mathrm{B}(m_1, p_1), \ldots Y_n \sim \mathrm{B}(m_n, p_n)$, see Table 5.1 and hence

$$\mathbf{Y} \sim \mathrm{B}(\mathbf{m}, \mathbf{p}), \quad \text{and } \mathbf{p} = \frac{\exp\{X\beta\}}{1 + \exp\{X\beta\}},$$

with response $\mathbf{Y} = \{Y_1, \dots, Y_n\}$, $\mathbf{m} = \{m_1, \dots, m_n\}$, $\mathbf{p} = \{p_1, \dots, p_n\}$, X being the $n \times p$ design matrix and $\beta = \{\beta_1, \dots, \beta_p\}$ the linear predictor parameters.

Table 5.1: Linear logistic model

	Group 1	Group 2	 Group n
Successes	Y_1	Y_2	 Y_n
Failures	$m_1 - Y_1$	$m_2 - Y_2$	 $m_n - Y_n$
Total	m_1	m_2	 m_n

Then we can write down the log likelihood of β using Equation 4.12 and Equation 5.2 as

$$l(\beta; \mathbf{y}) = \sum_{i=1}^{n} \left\{ y_i \, \log(p_i) + (m_i - y_i) \log(1 - p_i) + \log {m_i \choose y_i} \right\} \tag{5.5}$$

where

$$p_i = \frac{\exp\{\mathbf{x}_i^T \boldsymbol{\beta}\}}{1 + \exp\{\mathbf{x}_i^T \boldsymbol{\beta}\}}.$$

We would then use the principle of maximum likelihood to estimate β

$$\hat{\boldsymbol{\beta}} = \arg\max_{\boldsymbol{\beta}} \ l(\boldsymbol{\beta}; \mathbf{y}).$$

Then, the fitted probability are given by

$$\hat{p}_i = \frac{\exp\{\mathbf{x}_i^T \hat{\boldsymbol{\beta}}\}}{1 + \exp\{\mathbf{x}_i^T \hat{\boldsymbol{\beta}}\}}$$

and corresponding fitted values

$$\hat{y}_i = m_i \, \hat{p}_i$$
.

The Pearson residuals for binomial data take the form

$$e_i^P = \frac{y_i - m_i \hat{p}_i}{\sqrt{m_i \hat{p}_i (1 - \hat{p}_i)}}.$$

From the general result $\text{Var}(Y_i) = \phi b''(\theta)$ in Proposition 3.1, it can be shown that $\text{Var}(Y_i) = m_i p_i (1 - p_i)$. Then, using the estimate of p_i leads to the denominator above.

For large m_i , the usual Normal approximation to the Binomial means that the Pearson residuals are approximately N(0,1) distributed.

It can be shown that the deviance for the fitted model is

$$D = 2\sum_{i=1}^n \left\{ y_i \log \left(\frac{y_i}{m_i \hat{p}_i} \right) + (m_i - y_i) \log \left(\frac{m_i - y_i}{m_i (1 - \hat{p}_i)} \right) \right\}, \tag{5.6}$$

which is approximately χ^2_{n-r} distributed if the model is correct, where r is the number of degrees of freedom in the model (i.e. the number of columns of the design matrix).

This formula can be shown to be equivalent to

$$D = 2\sum_{j=1}^{2} \sum_{i=1}^{n} O_{ji} \log \frac{O_{ji}}{E_{ji}}$$

where O_{ji} denotes the observed value and E_{ji} denotes the expected value in cell (j, i) of the $2 \times n$ table of successes and failures:

Table 5.2: Observed and expected frequencies in the linear logistic model

	Group 1	Group 2	 Group n
Successes, $j = 1$	O_{11}	O_{12}	 O_{1n}
	E_{11}	E_{12}	 E_{1n}
Failures, $j=2$	O_{21}	O_{22}	 O_{2n}
	E_{21}	E_{22}	 E_{2n}

Another goodness-of-fit statistic is the Pearson chi-squared statistic:

$$X^2 = \sum_{j=1}^2 \sum_{i=1}^n \frac{(O_{ji} - E_{ji})^2}{E_{ji}}.$$

This is asymptotically equivalent to the deviance Equation 5.6 (proof is by Taylor series expansion; omitted). Thus, asymptotically, X^2 is also approximately χ^2_{n-r} distributed. Both approximations can be poor if the expected frequencies are small, but X^2 copes slightly better with this problem. See Dobson, p.136 for more details.

5.3 Overdispersion

Examination of residuals and deviances may indicate that a model is not an adequate fit to the data. One possible reason is *overdispersion*. This can occur for any error distribution where the variance is linked to the mean — e.g. Binomial, Poisson. In the binomial case, overdispersion is called *extra-Binomial* variation.

Recall that if $Y_i \sim \text{Bin}(m_i, p_i)$, $\text{Var}(Y_i) = m_i p_i (1-p_i)$. Overdispersion occurs if observations which have been modelled by a $\text{Bin}(m_i, \hat{p}_i)$ distribution have substantially greater variation than $m_i \hat{p}_i (1-\hat{p}_i)$. This will lead to a value of D substantially greater than the expected value of n-r. This can occur if the model is missing appropriate explanatory variables or has the wrong link function, or if the Y_i are not independent.

One solution is to include an extra parameter τ in the model so that $\mathrm{Var}(Y_i) = \tau \times m_i p_i (1-p_i)$. For more details, see Section 7.7 of Dobson or Chapter 6 of Collett (1991) *Modelling Binary data*, Chapman & Hall.

The glm function in **R** allows for *extra-Binomial* variation by setting family=quasibinomial() with the usual link functions available.

5.4 Application to dose-response experiments

A variable dose of some reagent is administered to each study subject, and the occurrence of a specific response is recorded. This is a *dose-response* experiment, one of the first uses of regression models for Bernoulli (or Binomial) responses.

For example, the Table 5.3 (including the information from Table 1.1) gives the number of beetles killed y_i and the number not killed $(m_i - y_i)$ out of a total number m_i that were exposed to a dose x_i of gaseous carbon disulphide, for n = 8 dose levels i = 1, ..., 8 (Dobson: pp.109 in 1st edn; pp.119 in 2nd edn; pp.127 in 3rd edn). The proportion killed $p_i = y_i/m_i$ at each dose level i is also shown in Table 5.3 and plotted in Figure 5.1.

Table 5.3: Numbers of beetles killed by five hours of exposure to 8 different concentrations of gaseous carbon disulphide

Dose	No. of beetle	No. killed	No. not killed	Proportion
x_i	m_i	y_i	$m_i - y_i$	$p_i = y_i/m_i$
1.6907	59	6	53	0.10
1.7242	60	13	47	0.22
1.7552	62	18	44	0.29
1.7842	56	28	28	0.50
1.8113	63	52	11	0.83
1.8369	59	53	6	0.90
1.8610	62	61	1	0.98
1.8839	60	60	0	1.00

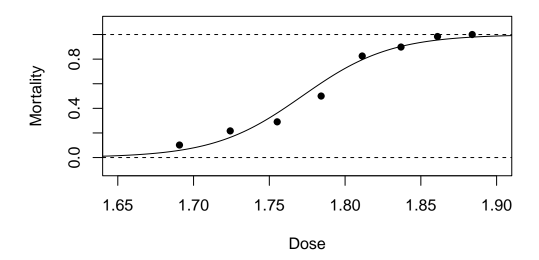


Figure 5.1: Beetle mortality rates with fitted dose-response curves.

Now Equation 5.4 motivates modelling the beetle data as

$$Y_i \sim B(m_i, p_i), \text{ for } i = 1, ..., n = 8$$

where

$$\eta_i = \alpha + \beta x_i$$

and so

$$p_i = \frac{\exp\{\alpha + \beta x_i\}}{(1 + \exp\{\alpha + \beta x_i\})}.$$

To fit this model in \mathbf{R} , a matrix with columns containing the numbers killed y_i and the numbers not killed m_i-y_i is first calculated

	[,1]	[,2]
[1,]	6	53
[2,]	13	47
[3,]	18	44
[4,]	28	28
[5,]	52	11
[6,]	53	6

which is then used in the glm command

The model parameter estimates are given by

and hence

$$\hat{p}_i = \frac{\exp\{-60.7 + 34.3x_i\}}{(1 + \exp\{-60.7 + 34.3x_i\})}.$$

Further, the deviance if given by

[1] 11.23223

or more helpfully a full summary, which contains parameter estimates and deviance values using

```
Call:
glm(formula = y ~ dose, family = binomial)
```

Deviance Residuals:

```
Min 1Q Median 3Q Max -1.5941 -0.3944 0.8329 1.2592 1.5940
```

Coefficients:

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

(Dispersion parameter for binomial family taken to be 1)

Null deviance: 284.202 on 7 degrees of freedom Residual deviance: 11.232 on 6 degrees of freedom

AIC: 41.43

Number of Fisher Scoring iterations: 4

or the anova table

Analysis of Deviance Table

Model: binomial, link: logit

Response: y

Terms added sequentially (first to last)

Clearly, many of the results are repeated in the various \mathbf{R} output.

When considering testing model goodness of fit, it is important to distinguish between deviance values which compare a given model with the saturated model and deviance values which compare two competing models. For example, in the Analysis of Deviance Table, the right hand values compare the Null model with the saturated and the model including dose with the saturated. Whereas the left hand value is relevant to the comparison of Null model with the mode including dose.

Summary of results:

From the output, we can first test if the Null model, which contains only a constant term, is a good fit to the data using the hypotheses:

 H_0 : Null model is true against H_1 : Null model is false.

From the output, note that the Null deviance is $D_0=284.202$ and that this model has $r_0=1$ parameters. Therefore, D_0 follows a χ^2 distribution with $n-r_0=8-1=7$ degrees of freedom and hence the p-value is:

[1] 1.425247e-57

This means that we reject H_0 as the observed value of D is in the upper tail of the χ^2 distribution.

If we had obtained a non-significant result, then we would stop the analysis and conclude that the Null model is an adequate fit.

Next, consider testing the model which includes the explanatory variable where the the hypotheses are

 H_0 : Null model is true against H_1 : Model with dose is true.

From the output, the deviance of the proposed model is $D_1=11.232$ with $r_1=2$ parameters. The test statistics is then $D_0-D_1=284.202-11.232=272.97$ which follows a χ^2 distribution with $r_1-r_0=2-1=1$ degrees of freedom. The p-value is then

[1] 2.556369e-61

and we reject H_0 in favour of H_1 and conclude that dose is important.

We can finish the testing by checking if this model is a good fit to the data. That is,

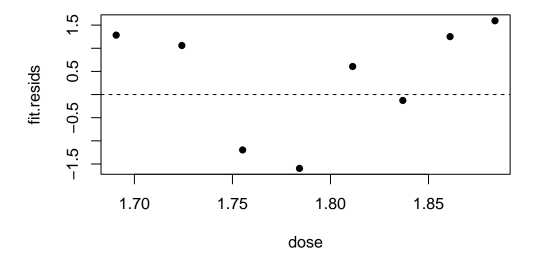
 ${\cal H}_0$: The model with dose is true against ${\cal H}_1$: The model with dose is false

The deviance of this is $D_1=11.232$ with $r_1=2$ parameters which follows a χ^2 distribution with $n-r_1=8-2=6$ degrees of freedom and has p=value

[1] 0.08146544

which means that we accept H_0 at the 5% level and conclude that this model is an adequate fit to the data.

Finally, we can look at the residuals:



These show a very slight u-shaped pattern and hence it would be work exploring fitting using the other link functions. For this data set, the complementary log-log (cloglog) link has a slightly better fit, see Figure 3.4, but a slight pattern remains.

Before finishing, suppose that we wish to predict, by hand, the probability for dose level $x_0=1.85$, say, and the expected number of beetles killed when $m_0=60$ beetles, say, are exposed.

The probability is predicted as

$$\hat{p}_i = \frac{\exp\{-60.7 + 34.3x_0\}}{(1 + \exp\{-60.7 + 34.3x_0\})} = 0.936$$

and the expected number of beetles

$$m_0 \, \hat{p}_i = 56.2$$

Finally, suppose that we wish to find the minimum dose which kills 95% of the beetles, that is $p_0=0.95$ which requires us to invert the logistic equation.

In general, suppose we wish to find x_p which corresponds to proportion p then

$$x_p = \frac{1}{\beta} \left\{ \log \left(\frac{p}{1-p} \right) - \alpha \right\}$$

and hence here $\hat{x}_p = 1.856$.

5.5 Exercises

To be added later.

6 Loglinear Models

6.1 Overview

In this chapter, we deal with data sets in which the response variable y is a count and the explanatory variables are all factors – i.e. qualitative variables. Initially we assume y has a Poisson distribution.

See Chapter 9 of Dobson and Barnett (2008). See also Agresti (1996) An introduction to categorical data analysis.

We assume a generalized linear model with

- responses (counts) having independent Poisson distributions;
- a logarithmic link function (hence the name log-linear model).

Consider, for example, the two-way contingency table in Table 6.1}:

Table 6.1: A two-way contingency table with k_1 rows and k_2 columns, where each entry y_{ij} is a count.

	1	2	•••	j	•••	k_2	Total
1	y_{11}	y_{12}		y_{1j}		y_{1k_2}	y_{1+}
		•••		•••	•••		
i	y_{i1}	y_{12}	•••	y_{1j}	•••	y_{ik_2}	y_{i+}
	•••	•••					
k_1	$y_{k_{1}1}$	$y_{k_1 2}$		y_{1j}		$y_{k_2k_2}$	y_{1k_1}
Total	y_{+1}	y_{+2}		y_{1j}		y_{+k_2}	y_{++}

In row i and column j of Table 6.1 we assume

$$Y_{ij} \sim \text{Po}(\lambda_{ij})$$

where

$$\log \lambda_{ij} = \mu + \alpha_i + \beta_j + (\alpha \beta)_{ij}. \tag{6.1}$$

Here μ is called the main effect; α_i is a row effect; β_j is a column effect; and $(\alpha\beta)_{ij}$ denotes an interaction effect parameter. Some or all of these effects must be present in the model, with constraints to ensure that the model is identifiable – this is sometimes referred ot as the identifiability or aliasing problem. Generally, $\mathbf R$ function glm automatically sets the first level of each effect to zero to achieve model identification. Thus, for the two-way model,

$$\alpha_1=0,\quad \beta_1=0,\quad (\alpha\beta)_{11}=(\alpha\beta)_{1j}=(\alpha\beta)_{k_1}=0.$$

6.2 Motivating examples

6.2.1 Malignant melanoma

The data in Table 6.2 are from a study of 400 patients with malignant melanoma, a particular form of skin cancer (see Dobson and Barnett, p.172). For each tumour, its type and site were recorded. The data in Table 6.2 comprise the numbers of tumours y in each combination of site and tumour-type. This is a two-way contingency table. We want to know how melanoma frequency depends on site and type.

Table 6.2: Melanoma counts by type and site.

Type	Head	Trunk	Extremities	Total
Hutchinson's melanotic freckle	22	2	19	34
Superficial spreading melanoma	16	54	115	185
Nodular	19	33	73	125
Indeterminate	11	17	28	56
Total	68	106	226	400

Table 6.3: Percentages across columns within rows for melanoma data.

Type	Head	Trunk	Extremities	Total
Hutchinson's melanotic freckle	64.7	5.9	29.4	100
Superficial spreading melanoma	8.6	29.2	62.2	100
Nodular	15.2	26.4	58.4	100
Indeterminate	19.6	30.4	50.0	100
Total	17.0	26.5	56.5	100

Table 6.4: Percentages across rows within columns for melanoma data.

Type	Head	Trunk	Extremities	Total
Hutchinson's melanotic freckle	32.4	1.9	4.4	8.5
Superficial spreading melanoma	23.5	50.9	50.9	46.3
Nodular	27.9	31.1	32.3	31.3
Indeterminate	16.2	16.0	12.4	14.0
Total	100.0	99.9	100.0	100.0

Although we have two factors, Type and Site, that we may use as predictors, standard ANOVA regression methods are inappropriate here as the dependent variable is not continuous but is instead a count. We will use *log-linear regression*, a type of generalized linear model, to analyse these data.

Table 6.3 shows row and Table 6.4 column percentages for these data. For example, 15.2% of nodular melanomas occurred in the head and neck, 26.4% in the trunk, and 58.4% in the extremities. Compare this to the equivalent figures for Hutchinson's melanotic freckles: 64.7%, 5.9%, and 29.4% - strikingly different. So different types of melanomas are more likely to occur in different locations.

6.2.2 Flu vaccine

The data in Table 6.5 are from a randomized controlled trial in which 73 patients were randomized into two groups (see Dobson and Barnett, 2008, p.173). The treatment group was given a flu vaccine, while the control group was given a placebo. Levels of an antibody (HIA) were measured after 6 weeks and classified into three groups: Low, Moderate, and High.

Table 6.5: Antibody responses to flu vaccine from a randomized controlled trial.

Group	Low	Moderate	High
Placebo	25	8	5
Vaccine	6	18	11
Total	31	26	16

Is the pattern of response the same for each treatment group? The percentages in Table 6.6 suggest not - row percentages indicate lower responses in the placebo group.

Table 6.6: Percentages across rows within columns for antibody responses to flu vaccine.

Group	Low	Moderate	High	Total
Placebo	65.8	21.1	13.2	100
Vaccine	17.1	51.4	31.4	100
Total	42.4	35.6	22.0	100

Table 6.7: Percentages across rows within columns for antibody responses to flu vaccine.

Group	Low	Moderate	High
Placebo	80.6	30.8	31.2
Vaccine	19.4	69.2	68.8
Total	100	100	100

6.3 Maximum likelihood estimation

Recall that for each cell $Y_{ij} \sim \text{Po}(\lambda_{ij})$ so $\text{E}[Y_{ij}] = \lambda_{ij}$. However, we estimate $\hat{\lambda}_{ij} = y_{ij}$ only for the saturated model given by Equation 6.1. In general, for non-saturated models, the estimate $\hat{\lambda}_{ij} \neq y_{ij}$.

Consider the independence model for a 2-way table, that is where $(\alpha\beta)_{ij} = 0$ for all i, j. Here,

$$\log \lambda_{ij} = \mu + \alpha_i + \beta_j \tag{6.2}$$

and y_{ij} is the observed count for cell (i, j), so the likelihood is given by

$$L(\lambda; y) = \prod_{i,j} \frac{e^{-\lambda_{ij}} \lambda_{ij}^{y_{ij}}}{y_{ij}!}$$

and the log-likelihood is

$$l(\lambda;y) = \sum_{i,j} \left\{ y_{ij} \log \lambda_{ij} - \lambda_{ij} - \log y_{ij}! \right\}.$$

Next, using Equation 6.2, gives

$$l(\lambda;y) = \sum_{i,j} \left\{ y_{ij}(\mu + \alpha_i + \beta_j) - \exp(\mu + \alpha_i + \beta_j) - \log y_{ij}! \right\}$$

which can be re-written as

$$l(\lambda;y) = \mu y_{++} + \sum_{i} \alpha_{i} y_{i+} + \sum_{j} \beta_{j} y_{+j} - e^{\mu} \left(\sum_{i} e^{\alpha_{i}} \right) \left(\sum_{j} e^{\beta_{j}} \right) - \sum_{ij} \log y_{ij}!. \tag{6.3}$$

The maximum likelihood estimates of the model parameters are obtained in the usual way. Differentiating Equation 6.3 with respect to μ and setting the result to zero gives, at the MLE,

$$y_{++} = e^{\hat{\mu}} \left(\sum_{i} e^{\hat{\alpha}_{i}} \right) \left(\sum_{j} e^{\hat{\beta}_{j}} \right).$$

Differentiating Equation 6.3 with respect to α_i (where $i \neq 1$ because $\alpha_1 = 0$ to avoid an identifiability problem) and setting the result to zero gives, at the MLE,

$$y_{i+} = e^{\widehat{\mu}} e^{\widehat{\alpha}_i} \left(\sum_j e^{\widehat{\beta}_j} \right).$$

Differentiating Equation 6.3 with respect to β_j (where $j \neq 1$ because $\beta_1 = 0$) and setting the result to zero gives, at the MLE,

$$y_{+j} = e^{\widehat{\mu}} e^{\widehat{\beta}_j} \left(\sum_i e^{\widehat{\alpha}_i} \right).$$

Then

$$\frac{y_{i+} y_{+j}}{y_{++}} = e^{\widehat{\mu} + \widehat{\alpha}_i + \widehat{\beta}_j} = \widehat{\lambda}_{ij}. \tag{6.4}$$

It follows that

$$\hat{\lambda}_{i+} = y_{i+}$$
 $\hat{\lambda}_{+i} = y_{+i}$ $\hat{\lambda}_{++} = y_{++}$.

Thus, the total fitted count in row i is identical to the total observed count in row i. Further, the total fitted count in column j is equal to the total observed count in column j and the total fitted count equals the total observed count.

6.4 Model fitting in R

Consider an analysis of Melanoma data introduced in Section 6.2.1.

To test the independence of Type and Size, we fit the model

Count
$$\sim$$
 Type + Site

assuming Poisson counts and a logarithmic link function, using the commands:

Call:

glm(formula = count ~ type.F + site.F, family = "poisson")

Deviance Residuals:

Min 1Q		Median 3Q		Max	
-3 0453	-1 0741	0 1297	0 5857	5 1354	

Coefficients:

	Estimate	Std. Error z	value	Pr(> z)	
(Intercept)	1.7544	0.2040	8.600	< 2e-16	***
type.F2	1.6940	0.1866	9.079	< 2e-16	***
type.F3	1.3020	0.1934	6.731	1.68e-11	***
type.F4	0.4990	0.2174	2.295	0.02173	*
site.F2	0.4439	0.1554	2.857	0.00427	**
site.F3	1.2010	0.1383	8.683	< 2e-16	***

Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1

(Dispersion parameter for poisson family taken to be 1)

Null deviance: 295.203 on 11 degrees of freedom Residual deviance: 51.795 on 6 degrees of freedom

AIC: 122.91

Number of Fisher Scoring iterations: 5

We see that the residual deviance for this model is 51.795 on 6 degrees of freedom. If the model is true, the residual deviance will have an approximate χ^2 distribution on 6 degrees of freedom. If the model is not true, the residual deviance will probably be too large to correspond to this distribution. Thus we calculate the p-value in the upper tail of the χ_6^2 distribution, which can be computed using the command:

[1] 2.050465e-09

This strongly indicates that the independence model is inadequate. Therefore, unless we can spot alternative simplifications, we will have to use the saturated model.

We can look at residuals from the model see where the departures from independence occur. The largest residual is for Hutchinson's freckle on the head and neck, which occurs more often than would be expected under independence.

For the saturated model, $\hat{\lambda}_{ij} = y_{ij}$. In this example, $\sum_{ij} \hat{\lambda}_{ij} = \sum_{ij} y_{ij} = 400$, so the probability of a tumour being in category (i,j) is $y_{ij}/400$ — just the observed proportions.

Note that in Table 6.2 we have a total of $y_{++} = 400$ observations. If the data were truly Poisson, this would be a suspiciously round number. In reality, this total was fixed by design, so we should take into account the fact that $y_{++} = 400$ and fit a more suitable model, such as the *multinomial* model which we will meet in the next chapter.

6.5 Multi-way contingency tables

We can generalize the model notation introduced in Equation 6.1 to accommodate multiway contingency tables; that is tables of counts indexed by multiple factors. For example, for a 3-way table of cell counts y_{ijk} , the saturated model would be written:

$$Y_{ijk} \sim \text{Po}(\lambda_{ijk})$$

where

$$\log \lambda_{ijk} = \mu + \alpha_i + \beta_j + \gamma_k + (\alpha\beta)_{ij} + (\alpha\gamma)_{ik} + (\beta\gamma)_{jk} + (\alpha\beta\gamma)_{ijk}.$$

Here, α_i , β_j and γ_j are main effects; $(\alpha\beta)_{ij}$, $(\alpha\gamma)_{ik}$, and $(\beta\gamma)_{jk}$ are two-way interaction effects; and $(\alpha\beta\gamma)_{ijk}$ is a three-way interaction.

This approach is easily extended to tables of any number of factors. Note, however, that not all terms need be present in a model, thought 3-way or higher-order interactions might sometimes be required.

A hierarchical model is one in which each term in the model is accompanied by all lower-order terms. For example, including the term $(\alpha\beta\gamma)_{ijk}$ would require inclusion of each of the terms $\alpha_i, \beta_j, \gamma_k, (\alpha\beta)_{ij}, (\alpha\gamma)_{ik}, (\beta\gamma)_{jk}$. We will be concerned only with hierarchical contingency table models. Non-hierarchical models are sometimes appropriate, depending on the nature of the factors involved, but hierarchical models are more generally applicable and easier to interpret.

6.6 Exercises

Exercises to be added later.

7 Extensions to Loglinear models

7.1 Overview

In the melanoma example, see Table 6.2, the overall total number of observations was exactly 400. If the data were truly Poisson, this would be a suspiciously round number. In reality, this total was fixed by design, so we should take into account the fact that $y_{++} = 400$ and fit a more appropriate model. Also, recall the flu vaccine example of Table 6.5. Suppose that some of the marginal totals are fixed by design. For example, that the number of patients in each arm of the trial (38 in the Placebo group, 35 in the Vaccine group) was fixed before data collection started. In this case, we should also take into account the fact that $y_{1+} = 38$ and $y_{2+} = 35$ and fit a more suitable model.

Before we can re-consider these data sets, we first need to establish some theoretical results to deal with such *conditional distribution* cases.

7.2 Contingency tables with fixed marginals

Consider, again, a general two-way contingency table such as that in Table 7.1. The fixed marginals are the column sums: y_{+j} , the row sums: y_{i+} , and the overall sum: y_{++} .

Table 7.1: A two-way contingency table with k_1 rows and k_2 columns, where each entry y_{ij} is a count.

	1	2	•••	j	•••	k_2	Total
1	y_{11}	y_{12}	•••	y_{1j}	•••	y_{1k_2}	y_{1+}
i	y_{i1}	y_{12}		y_{1j}	•••	y_{ik_2}	y_{i+}
k_1	$y_{k_{1}1}$	$y_{k_{1}2}$		y_{k_1j}		$y_{k_1k_2}$	y_{k_1+}
Total	y_{+1}	y_{+2}		y_{+j}		y_{+k_2}	y_{++}

Suppose that we have fixed row totals y_{i+} , or a fixed overall total y_{++} , how would such constraints affect the distribution of the random variables Y_{ij} ?

To answer this question, we will need the following results.

Proposition 7.1. Let Y_{ij} , for $i=1,\ldots,k_1$ and $j=1,\ldots,k_2$, be the responses in a two-way contingency table, with k_1 rows and k_2 columns. Further, assume that the counts follow independent Poisson distributions with corresponding means λ_{ij} , that is $Y_{ij} \sim Po(\lambda_{ij})$. Then, the distributions of the row sums are,

$$Y_{i+} \sim Po(\lambda_{i+}),$$

where
$$Y_{i+} = \sum_{j=1}^{k_2} Y_{ij} \text{ and } \lambda_{i+} = \sum_{j=1}^{k_2} \lambda_{ij}.$$

Notes:

- Here we are not yet assuming that the marginal totals are fixed.
- Similar results hold when considering column sums and the overall sum.

Proof: See MATH2715, Chapter 7 on MGF Properties of Linear Functions.

Proposition 7.2. Let $Y_{ij} \sim Po(\lambda_{ij})$ for $i = 1, ..., k_1$ and $j = 1, ..., k_2$ then the conditional distribution of each count in a row given that the row sum is fixed is given by

$$\left(Y_{ij} \,|\, Y_{i+} = m\right) \sim Bin\left(m, \frac{\lambda_{ij}}{\lambda_{i+}}\right).$$

That is, conditioning on the sum of independent Poisson random variables induces a Binomial distribution.

Proof: First define $Y_{i-j} = Y_{i+} - Y_{ij}$ with correspondingly $y_{i-j} = m - y_{ij}$ and also $\lambda_{i-j} = \lambda_{i+} - \lambda_{ij}$.

Consider the conditional probability mass function

$$\begin{split} p(Y_{ij} = y_{ij} | Y_{i+} = m) &= \frac{p(Y_{ij} = y_{ij}, Y_{i+} = m)}{p(Y_{i+} = m)} \quad \text{by conditional probability definition} \\ &= \frac{p(Y_{ij} = y_{ij}, Y_{i-j} = m - y_{ij})}{p(Y_{i+} = m)} \quad \text{rearranging} \\ &= \frac{p(Y_{ij} = y_{ij}) p(Y_{i-j} = m - y_{ij})}{p(Y_{i+} = m)} \quad \text{by independence.} \end{split}$$

Then, each of these follows a Poisson distribution: $Y_{ij} \sim \text{Po}(\lambda_{ij}), Y_{i-j} \sim \text{Po}(\lambda_{i-j}),$ and $Y_{i+} \sim \text{Po}(\lambda_{i+}).$ Therefore,

$$\begin{split} p(Y_{ij} = y_{ij} | Y_{i+} = m) &= \frac{\lambda_{ij}^{y_{ij}} e^{-\lambda_{ij}}}{y_{ij}!} \frac{\lambda_{i-j}^{m-y_{ij}} e^{-\lambda_{i-j}}}{m - y_{ij}!} \left/ \frac{\lambda_{i+}^{m} e^{-\lambda_{i+}}}{m!} \right. \\ &= {m \choose y_{ij}} \left(\frac{\lambda_{ij}}{\lambda_{i+}} \right)^{y_{ij}} \left(\frac{\lambda_{i-j}}{\lambda_{i+}} \right)^{m-y_{ij}} \\ &= {m \choose y_{ij}} \pi^{y_{ij}} \left(1 - \pi \right)^{m-y_{ij}} \end{split}$$

which is the probability mass function of a $Bin(m, \pi)$ random variable, where $\pi = \lambda_{ij}/\lambda_{i+}$.

Proposition 7.3. Let $Y_{ij} \sim Po(\lambda_{ij})$ for $i = 1, ..., k_1$ and $j = 1, ..., k_2$ then the **joint** conditional distribution of the counts in a row given that the row sum is fixed is given by

$$p(Y_{i1},\dots,Y_{ik_2}\,|\,Y_{i+}=m)=\frac{m!}{y_{i1}!\cdots y_{ik_2}!}\pi_{i1}^{y_{i1}}\cdots\pi_{ik_2}^{y_{ik_2}}$$

with

$$\pi_{ij} = \lambda_{ij}/\lambda_{i+}$$
 and $\sum_{j=1}^{k_2} \pi_{ij} = 1$.

This distribution is called the Multinomial distribution, with index m and parameters π_{ij} , $j = 1, ..., k_2$, which can be denoted

$$(Y_{i1},\ldots,Y_{ik_2}\,|\,Y_{i+}=m)\sim \mathit{Mult}(m,\pi_{i1},\ldots\pi_{ik_2}).$$

Proof: Omitted.

Proposition 7.4. Let $Y_{ij} \sim Po(\lambda_{ij})$ for $i = 1, ..., k_1$ and $j = 1, ..., k_2$ then the conditional distribution of the counts given that the overall sum is fixed is

$$(Y_{11},\ldots,Y_{k_1k_2}\,|\,Y_{++}=m)\sim \mathit{Mult}(m,\pi_{11},\ldots,\pi_{k_1k_2}).$$

with probability mass function given by

$$p(Y_{11},\ldots,Y_{k_1k_2}\,|\,Y_{++}=m)=\frac{m!}{y_{11}!\cdots y_{k_1k_2}!}\pi_{11}^{y_{11}}\cdots\pi_{k_1k_2}^{y_{k_1k_2}}$$

where $\pi_{ij} = \lambda_{ij}/\lambda_{++}$.

Proof: Omitted.

7.3 Product-multinomial models

The Poisson contingency table model assumes that each cell count (and therefore row and column total) is random – specifically, Poisson.

However, in many examples, some of the marginal totals are fixed by design.

For example, as discussed in the Flu-vaccine example of Section 6.2.2, the numbers in the placebo (38) and vaccine (35) groups are not random, but fixed by the experimenter. Further, we are not interested in modelling these row totals but instead the response to

treatment (placebo or vaccine). Thus, in the Flu vaccine example, we should condition the Poisson model on the row totals.

From Equation 6.1 the saturated model for a two-factor table is

$$\lambda_{ij} = \exp\left\{\mu + \alpha_i + \beta_j + (\alpha\beta)_{ij}\right\}.$$

Then, using the result in Proposition 7.3, we see that

$$\pi_{ij} = \frac{\lambda_{ij}}{\lambda_{i+}} = \frac{\exp\left\{\mu + \alpha_i + \beta_j + (\alpha\beta)_{ij}\right\}}{\sum_{j} \exp\{\mu + \alpha_i + \beta_j + (\alpha\beta)_{ij}\}}$$

which simplifies to give

$$\pi_{ij} = \frac{\exp\left\{\beta_j + (\alpha\beta)_{ij}\right\}}{\sum_{i} \exp\left\{\beta_j + (\alpha\beta)_{ij}\right\}}.$$
(7.1)

Thus, for each row, that is each i, this multinomial distribution depends on β_j and $(\alpha\beta)_{ij}$ for $j=1,\ldots,k_2$, but not on μ and α_i . This means that the data in each row of the table are independent and hence, overall, the model likelihood is given by

$$L(\beta; \mathbf{y}) = \prod_{i=1}^{k_1} p(Y_{i1}, \dots, Y_{ik_2} \, | \, Y_{i+} = y_{i+}),$$

with $\beta=\{\mu,\alpha_1,\dots,\alpha_{k_1},\beta_1,\dots,\beta_{k_2},(\alpha\beta)_{11},\dots,(\alpha\beta)_{k_1k_2}\}$ and where

$$p(Y_{i1},\ldots,Y_{ik_2}\,|\,Y_{i+}=y_{i+})=\frac{y_{i+}!}{y_{i1}!\cdots y_{ik_2}!}\pi_{i1}^{y_{i1}}\cdots\pi_{ik_2}^{y_{ik_2}}.$$

Further, using Equation 7.1 the log-likelihood is given by

$$l(\beta; \mathbf{y}) = \mathrm{const} + \sum_{i=1}^{k_1} \sum_{j=1}^{k_2} y_{ij} \left\{ \beta_j + (\alpha \beta)_{ij} - \log \sum_j \exp \left\{ \beta_j + (\alpha \beta)_{ij} \right\} \right\}.$$

Recall that the saturated model will have the fitted values equal to the data values in all cases. Therefore, we concentrate on the product multinomial models with interactions set to zero, that is $(\alpha\beta)_{ij} = 0$, then

$$l(\boldsymbol{\beta}; \mathbf{y}) = \operatorname{const} + \sum_{i=1}^{k_1} \sum_{j=1}^{k_2} y_{ij} \left\{ \beta_j - \log \sum_j \exp\left\{\beta_j\right\} \right\}$$

which simplifies to

$$l(\beta; \mathbf{y}) = \mathrm{const} + \sum_{j=1}^{k_2} y_{+j} \left\{ \beta_j - y_{++} \log \sum_j \exp\left\{\beta_j\right\} \right\}.$$

Differentiating this with respect to single parameter β_k and setting to zero gives

$$\frac{y_{+k}}{y_{++}} = e^{\hat{\beta}_k} / \sum_{i} e^{\hat{\beta}_j}$$

which is identical to the parameter estimate for the previous independence Poisson model. Thus, in this case, if we want to fit the product multinomial model (which is hard) we can instead fit the Poisson model (which is easy). Note, however, that these estimates only coincide when $\alpha_1, \ldots, \alpha_{k_1}$ are included in the Poisson independence model, even though we see that these are eliminated from the multinomial model. In fact, these parameter estimates reflect the fixed marginal totals in the multinomial situation.

Although this derivation has only considered the cases of the independence Poisson model: $\log \lambda_{ij} = \mu + \alpha_i + \beta_j$ and the product multinomial model with $(\alpha\beta)_{ij} = 0$, the following result gives general guidance.

Theorem 7.1. Tables with fixed margin sums can be analyzed using a multinomial or product-multinomial model as though they were independent Poisson models, with log-linear predictor, provided terms corresponding to the fixed margins are included in the model. Then the MLEs for the two models yield the same values for the parameters of interest and have the same deviances.

For a 2-way table:

Multinomial conditioning on	Poisson model must include
y_{++}	μ
y_{i+}	$\mu, lpha_i$

For a 3-way table:

Multinomial conditioning on	Poisson model must include
y_{+++}	μ
y_{i++}	μ, α_i
y_{ij+}	$\mu, \alpha_i, \beta_j, (\alpha \beta)_{ij}$
$y_{ij+},\ y_{+jk}$	$\mu, \alpha_i, \beta_j, \gamma_k, (\alpha\beta)_{ij}, (\beta\gamma)_{jk}$

Proof: Omitted. For details see Birch, 1963, JRSS(B), 220–233. The proof is not especially difficult but is a bit messy. It depends on the property of the Poisson model with log link function that observed totals equals fitted totals for the effects present in the Poisson model.

7.4 Model fitting in R

Consider again he flu vaccine data described in Section 6.2.2 with data repeated below. Recall that 38 people were recruited to the Placebo group and 35 to the Vaccine group and that we assume these were fixed before data collection started.

Table 7.4: Antibody responses to flu vaccine from a randomized controlled trial.

Group	Low	Moderate	High	Total
Placebo	25	8	5	38
Vaccine	6	18	11	35
Total	31	26	16	73

The fixed row sums mean that a product-multinomial model is appropriate but this can be fitted using the usual same command as the Independence Poisson model as long as we include the term for the row effect, that is for Group.

Call:

glm(formula = count ~ antibodyF + groupF, family = "poisson")

Deviance Residuals:

Coefficients:

Estimate Std. Error z value Pr(>|z|)
(Intercept) 3.0147 0.1985 15.189 < 2e-16 ***

antibodyF1 -0.1759 0.2659 -0.661 0.50835 antibodyF2 -0.6614 0.3078 -2.149 0.03167 * groupF1 -0.6523 0.2466 -2.645 0.00817 **

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

(Dispersion parameter for poisson family taken to be 1)

Null deviance: 23.807 on 5 degrees of freedom Residual deviance: 11.394 on 2 degrees of freedom

AIC: 44.522

Number of Fisher Scoring iterations: 5

Hence, our proposed model has a residual deviance of 18.643 on 2 degrees of freedom. The chi-square test gives a p-value of

$$p = \Pr(\chi_2^2 > 18.643) = 8.95 \times 10^{-5},$$

so we reject the independence model in favour of the saturated model.

For the independence model

$$\hat{\pi}_{ij} = e^{\hat{\beta}_j} / \sum_j e^{\hat{\beta}_j}$$

whose right-hand side does not depend on i and hence is the same for all rows. The corresponding fitted values are then given by $\hat{y}_{ij} = y_{i+} \hat{\pi}_{ij}$.

7.5 Exercises

To be added later.