Outline for today

- What is a generalized linear model
- Linear predictors and link functions
- Example: fit a constant (the proportion)
- Analysis of deviance table
- Example: fit dose-response data using logistic regression
- Example: fit count data using a log-linear model
- Advantages and assumptions of glm
- Quasi-likelihood models when there is excessive variance

Review: what is a linear model

A model of the following form:

$$Y = \beta_0 + \beta_1 X_1 + \beta_2 X_2 + \dots$$

- *Y* is the response variable
- The *X* 's are the explanatory variables
- The β 's are the parameters of the linear equation
- The errors are normally distributed with <u>equal variance</u> at all values of the *X* variables.
- Uses least squares to fit model to data and to estimate parameters
- lm in R

Review: fitting a linear model in R

Use the 1m package in R

Simplest linear model: fit a constant (the mean)

$$z < -lm(y \sim 1)$$

Linear regression

$$z < -lm(y \sim x)$$

Single factor ANOVA

$$z < - lm(y \sim A)$$

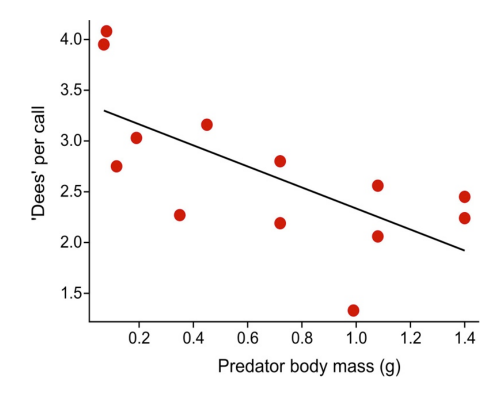
Review: what is a linear model

Eg: linear regression: $Y = \beta_0 + \beta_1 X + \text{error}$

The predicted Y-values, symbolized here by μ , is modeled as

$$\mu = \beta_0 + \beta_1 X$$

The part to the right of "=" is the linear predictor



What is a generalized linear model

A model whose predicted values are of the form

$$g(\mu) = \beta_0 + \beta_1 X_1 + \beta_2 X_2 + \dots$$

- The model still include a *linear predictor* (to right of "=")
- But the predicted Y-values are transformed
- $g(\mu)$ is called the "link function," of which there are several types
- Non-normal distributions of errors OK (specified by link function)
- Unequal error variances OK (specified by link function)
- Uses maximum likelihood to estimate parameters
- Uses log-likelihood ratio tests to test parameters
- Fit models using glm() in R

The two most common link functions

1) Natural log (i.e., base e)

$$\log(\mu) = \eta = \beta_0 + \beta_1 X_1 + \beta_2 X_2 + \dots$$

Usually used to model count data (e.g., number of mates, etc)

 $\log(\mu)$ is the link function.

The inverse function is $\mu=e^\eta$

The two most common link functions

2) Logistic or logit

$$\log \frac{\mu}{1 - \mu} = \eta = \beta_0 + \beta_1 X_1 + \beta_2 X_2 + \cdots$$

Used to model binary data (e.g., survived vs died)

The link function $\log \frac{\mu}{1-\mu}$ is also known as the log-odds

The inverse function is $\mu = \frac{e^{\eta}}{1+e^{\eta}}$

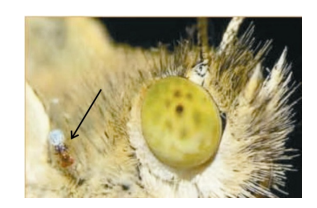
Example 1: Fit a constant to 0-1 data (estimate a proportion)

This example was used previously in Likelihood lecture. My goal with this example is to connect what glm() does with what we did by brute force last week.

The wasp, *Trichogramma brassicae*, rides on female cabbage white butterflies, *Pieris brassicae*. When a butterfly lays her eggs on a cabbage, the wasp climbs down and parasitizes the freshly laid eggs.

Fatouros et al. (2005) carried out trials to determine whether the wasps can distinguish mated female butterflies from unmated females. In each trial a single wasp was presented with two female cabbage white butterflies, one a virgin female, the other recently mated.

Y = 23 of 32 wasps tested chose the mated female. What is the proportion p of wasps in the population choosing the mated female?



Number of wasps choosing the mated female fits a binomial distribution

Under random sampling, the number of "successes" in n trials has a binomial distribution, with p being the probability of "success" in any one trial.

To model these data, let "success" be "wasp chose mated butterfly"

Y = 23 successes

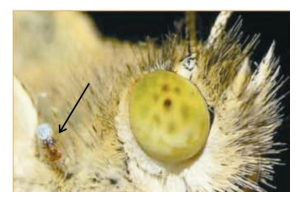
n = 32 trials

Goal: estimate p

Data are:

111011101010111110101111101110011



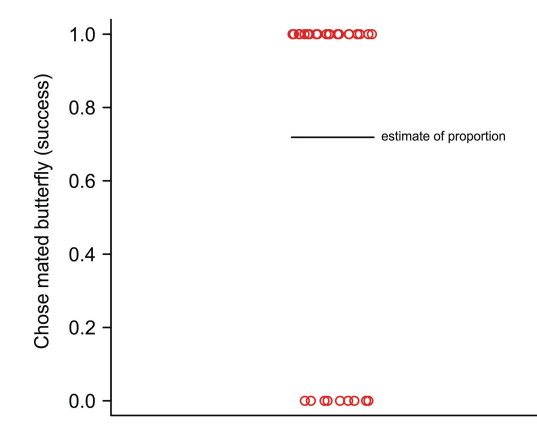


Use glm() to fit a constant, and so obtain the ML estimate of p

The data are binary. Each wasp has a measurement of 1 or 0 ("success" and "failure"): 111011101010111101011110011

z <- glm(choice ~ 1, family = binomial(link="logit"))</pre>

family specifies the error distribution (binomial) and the link function (logit).



Use glm() to fit a constant, and so obtain the ML estimate of p

Fits a model having only a constant. Use the link function appropriate for binary data:

$$\log \frac{\mu}{1-\mu} = \beta$$

 μ here refers to the population proportion (p) but let's stick with μ symbol here to use consistent notation for generalized linear models.

Fitting will yield the estimate, $\hat{\beta}$.

The estimate of proportion $\hat{\mu}$ is then obtained using the inverse function:

$$\hat{\mu} = \frac{e^{\hat{\beta}}}{1 + e^{\hat{\beta}}}$$

Use summary () for estimation

summary(z)

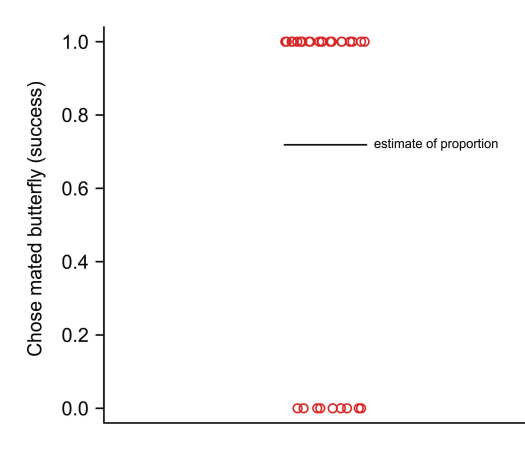
Coefficients:

Estimate Std. Error z value
$$Pr(>|z|)$$
 (Intercept) 0.9383 0.3932 $\frac{2.386}{0.017}$ *

0.9383 is the estimate of β (the constant on the **logit** scale). Convert back to ordinary scale (plug into inverse equation) to get estimate of population proportion:

$$\hat{\mu} = \frac{e^{\hat{\beta}}}{1 + e^{\hat{\beta}}} = \frac{e^{0.9383}}{1 + e^{0.9383}} = 0.719$$

This is the ML estimate of the population proportion. This is identical to the estimate obtained last week using likelihood function.



Confidence intervals

```
summary(z) Coefficients: Estimate Std. Error z value Pr(>|z|) (Intercept) 0.9383 0.3932 \frac{2.386}{2.386} 0.017 *
```

95% confidence limits:

```
myCI <- confint(z)  # on logit scale
exp(myCI)/(1 + exp(myCI)) # inverse logit scale

2.5 %  97.5 %
0.5501812 0.8535933</pre>
```

 $0.550 \le p \le 0.853$ is the same result we obtained last week for likelihood based confidence intervals using likelihood function (more decimal places this week).

Avoid using summary () for hypothesis testing

```
summary(z)
Coefficients:
Estimate Std. Error z value Pr(>|z|)
(Intercept) 0.9383 0.3932 \frac{2.386}{2.386} 0.017 *
```

The z-value (Wald statistic) and P-value test the null hypothesis that $\beta=0$. This is the same as a test of the null hypothesis that the true (population) proportion $\mu=0.5$, because

$$\frac{e^0}{1 + e^0} = 0.5$$

Agresti (2002, *Categorical data analysis*, 2nd ed., Wiley) says that for small to moderate sample size, the Wald test is <u>less reliable</u> than the log-likelihood ratio test.

Use anova() to test hypotheses

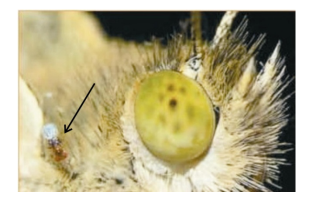
Last week we calculated the log-likelihood ratio test for these data "by hand". Here we'll use glm() to accomplish the same task.

```
"Full" model (\beta estimated from data):

z1 <- glm(y ~ 1, family = binomial(link="logit"))

"Reduced" model (\beta set to 0 by removing intercept from model):

z0 <- glm(y ~ 0, family = binomial(link="logit"))
```

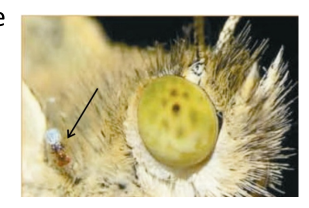


Use anova() to test hypotheses

```
anova(z0, z1, test = "Chi") # Analysis of deviance

Model 1: y ~ 0  # Reduced model

Model 2: y ~ 1  # Full model
```



Analysis of deviance table:

The <u>deviance</u> is the log-likelihood ratio statistic (*G*-statistic). It has an approximate χ^2 distribution under the null hypothesis.

Residual deviance is analogous to a residual sum of squares, and measures goodness of fit of the model to the data.

G = 6.227 is the identical result we obtained "by hand" last week.

Example 2: Logistic regression

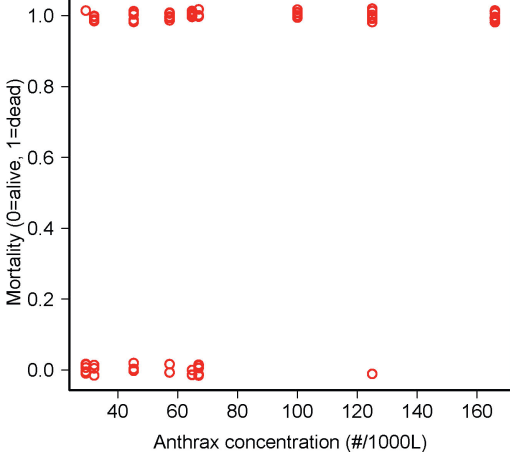
One of the most common uses of generalized linear models.

Goal is to model the relationship between a proportion and an explanatory variable

Data: 72 rhesus monkeys (Macacus rhesus) exposed for 1 minute to aerosolized

preparations of anthrax (Bacillus anthracis).

Goal is to estimate the relationship between dose and probability of death.

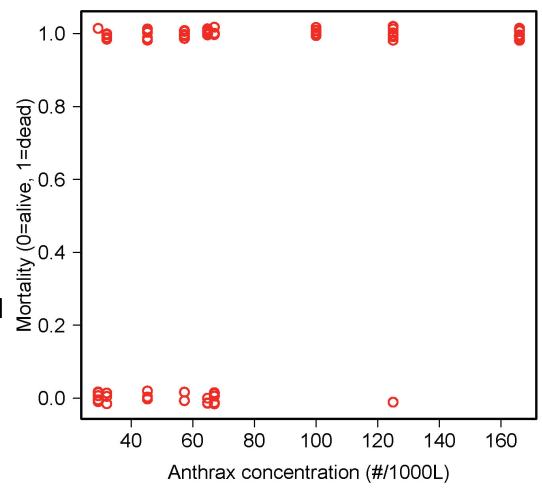


Logistic regression

Measurements of individuals are 1 (dead) or 0 (alive)

Ordinary linear regression model not appropriate because

- For each X the Y observations are binary, not normal
- For every X the variance of Y is not constant
- A linear relationship is not bounded between 0 and 1
- 0, 1 data can't simply be transformed



$$g(\mu) = \beta_0 + \beta_1 X_1$$

 μ is the <u>probability</u> of death, which depends on concentration X.

 $g(\mu)$ is the link function.

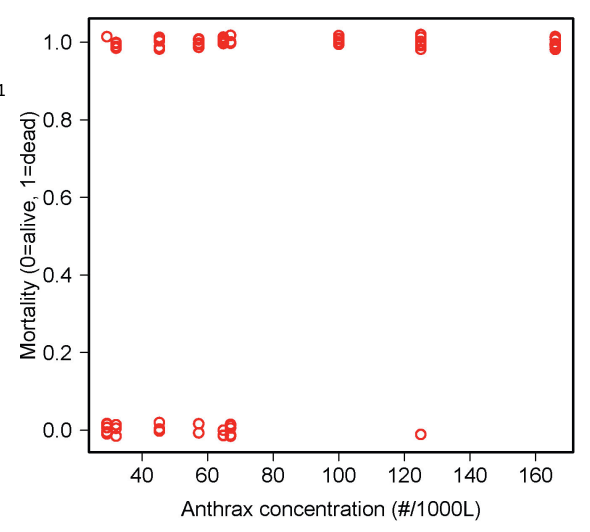
Linear predictor (right side of equation) is like an ordinary linear regression, with intercept β_0 and slope β_1

Logistic regression uses the logit link function

$$g(\mu) = \beta_0 + \beta_1 X$$

glm() uses maximum likelihood: the method finds those values of β_0 and β_1 for which the data have maximum probability of occurring. These are the maximum likelihood estimates.

No formula for the solution. glm() uses an iterative procedure to find the maximum likelihood estimates on the likelihood surface.



Use summary() for estimation

```
z <- glm(mortality ~ concentration, family = binomial(link = "logit")) summary(z) Estimate Std. Error z value Pr(>|z|) (Intercept) -1.74452 0.69206 -2.521 0.01171 * concentration 0.03643 0.01119 3.255 0.00113 **
```

Number of Fisher Scoring iterations: 5

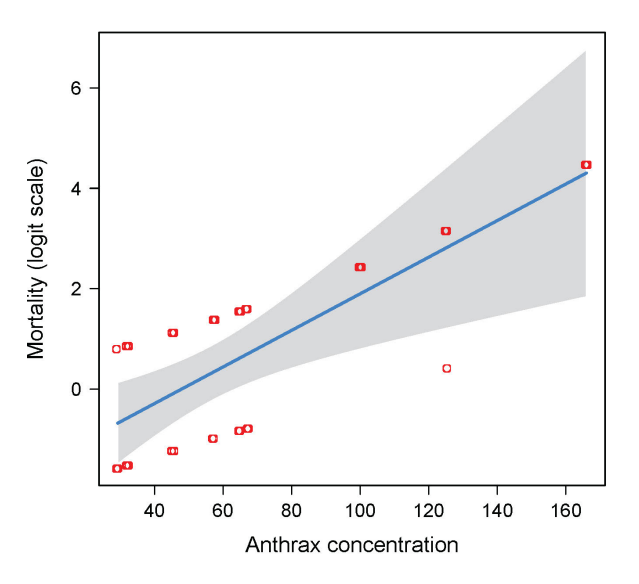
Numbers in red are the estimates of β_0 and β_1 (intercept and slope) which predict $\log(\mu/1-\mu)$.

Number of Fisher Scoring iterations refers to the number of iterations used before the algorithm used by glm() converged on the maximum likelihood solution.

Use predict(z) to obtain predicted values on the logit scale

$$g(\widehat{\mu}) = \widehat{\eta} = -1.74 + 0.036X$$

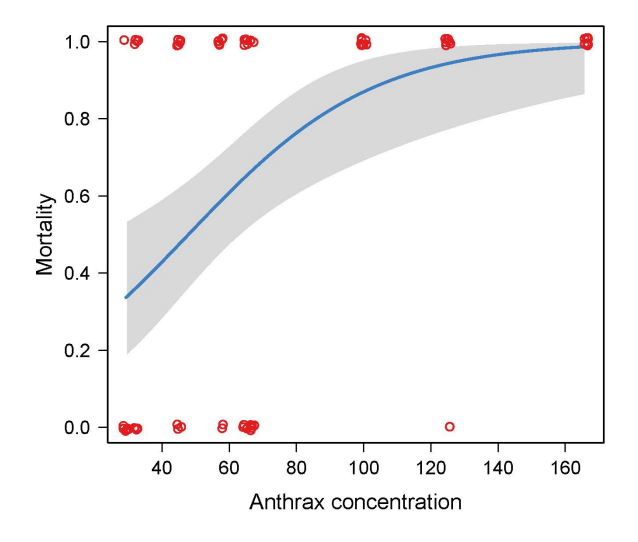
visreg(z) uses predict to plot predicted values, with confidence limits, on logit scale. See that the function is a line. The points on this scale are not the logit-transformed data. Instead, glm() creates "working" values to fit model to data on transformed scale.



Use fitted(z) to obtain predicted values on the original scale

$$\hat{\mu} = rac{e^{\hat{\eta}}}{1 + e^{\hat{\eta}}}$$

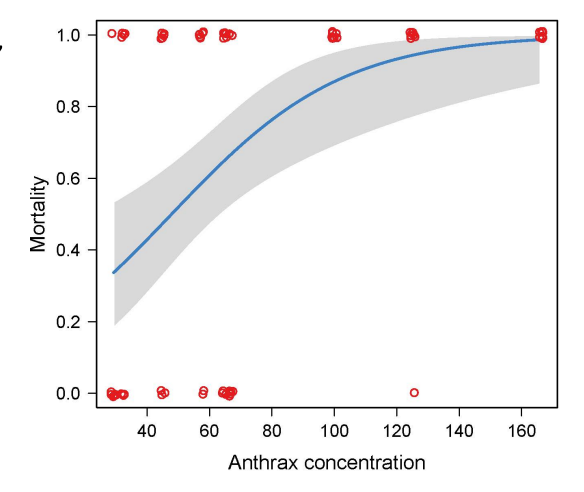
Use
visreg(z,
 scale = 'response')
to get fitted curve with
confidence bands on the
original scale.



$$LD_{50} = -\frac{intercept}{slope} = -\frac{0.03643}{-1.7445} = 47.88$$

The parameter estimates from the model fit can be used to estimate LD₅₀, the estimated concentration at which 50% of individuals are expected to die.

library(MASS)
dose.p(z)



Use anova() to test hypotheses

Analysis of deviance table gives log-likelihood ratio test of the null hypothesis that there is no differences among years in mean number of offspring.

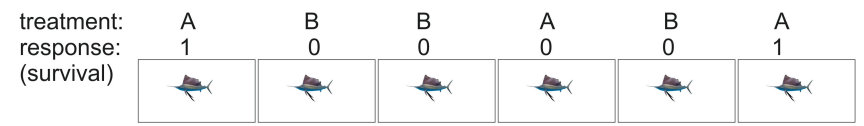
anova(z, test="Chisq")

As with lm(), terms are tested using model comparison (always a "full" vs "reduced" model). Default program of action is to fit terms sequentially ("Type 1 sums of squares"), as with lm().

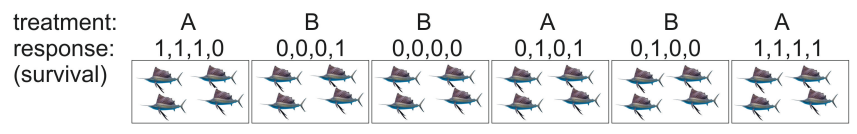
Advantages of generalized linear models

- More flexible than simply transforming variables. (A given transformation of the raw data may not accomplish both linearity and homogeneity of variance.)
- Yields more familiar measures of the response variable than data transformations.
- Avoids the problems associated with transforming 0's and 1's. For example, the logit transformation of 0 or 1 can't be computed.
- Retains the same analysis framework as linear models.

When glm() is appropriate and when it is not



Glm is appropriate (individual is the replicate)



Glm is not appropriate (tank is the replicate)

In second case, analyze summary statistic (fraction surviving) with lm(). Fitting generalized linear mixed models is possible in lme4 package using glmm() method.

Assumptions of generalized linear models

- Statistical independence of data points.
- Correct specification of the link function for the data.
- The variances of the residuals correspond to that assumed by the link function. (Later, I will show a method for dealing with violations of this assumption).

Example 3: Analyzing count data with log-linear regression

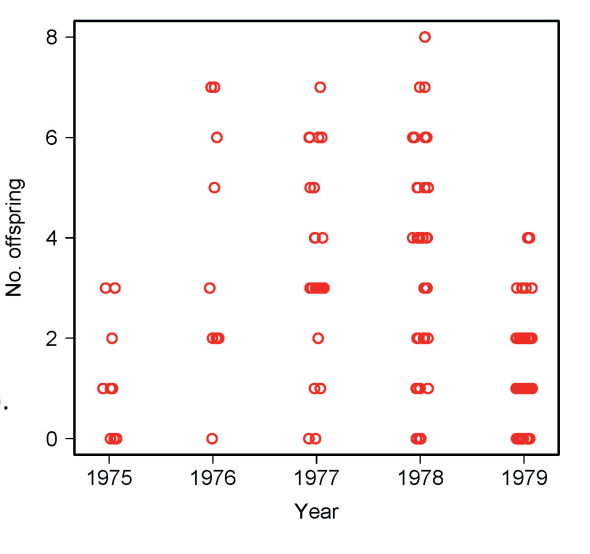
Estimate mean number of offspring fledged by female song sparrows on Mandarte Island, BC.



http://commons.wikimedia.org/wiki/ File:Song Sparrow-27527-2.jpg

Linear model assumptions not met:

Data are discrete counts (non-normal). Variance increases with mean.



Example 3: Analyzing count data with log-linear regression

No. offspring

Estimate mean number of offspring fledged by female song sparrows on Mandarte Island, BC.

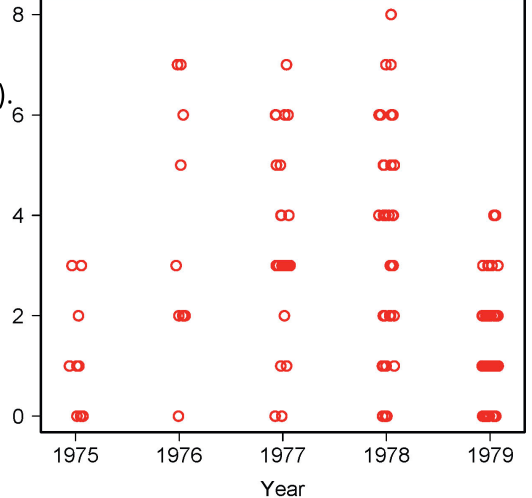
Linear model assumptions not met:

Data are discrete counts (non-normal).
Variance increases with mean.

Two solutions:

1. Transform data: $X' = \log(X + 1)$

Generalized linear model.
 Poisson distribution might be appropriate for error distribution.
 So try log link function.



Example 3: Analyzing count data with log-linear regression

Log-linear regression (a.k.a. Poisson regression) uses the log link function

$$\log(\mu) = \eta = \beta_0 + \beta_1 X_1 + \beta_2 X_2 + \cdots$$

 η is the response variable on the log scale (here, mean of each group on log scale).

Year is a categorical variable. So is analogous to single factor ANOVA.

Categorical variables are modeled in R using "dummy" indicator variables, as with lm().

Use summary() for estimation

```
z <- glm(noffspring ~ year, family=poisson(link="log"))
summary(z)</pre>
```

```
Estimate Std. Error z value Pr(>|z|) (Intercept) 0.24116 0.26726 0.902 0.366872 year1976 0.96665 0.28796 3.301 0.000963 *** year1978 0.97700 0.28013 3.488 0.000487 *** year1979 0.03572 0.29277 -0.122 0.902898
```

(Dispersion parameter for poisson family taken to be 1)

Numbers in red are the parameter estimates on the <u>log scale</u>. Intercept refers to mean of the first group (1975) and the rest of the coefficients are differences between each given group (year) and the first group.

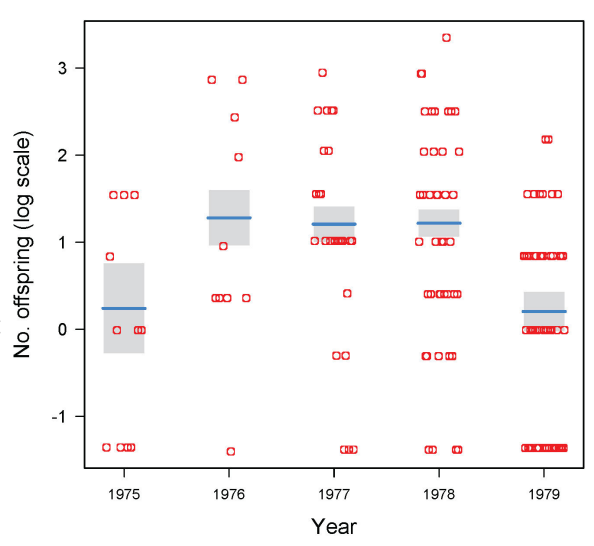
Dispersion parameter of 1 states assumption that in every year, variance = mean.

Predicted values on the transformed scale

Predicted values on the transformed (log) scale: predict(z)

visreg(z) uses predict to plot the predicted values, with confidence limits, on this transformed scale.

See that the "data points" on this scale are not just the transformed data (we can't take log of 0). Instead, glm() creates "working" values to fit the model to the data on the transformed scale.



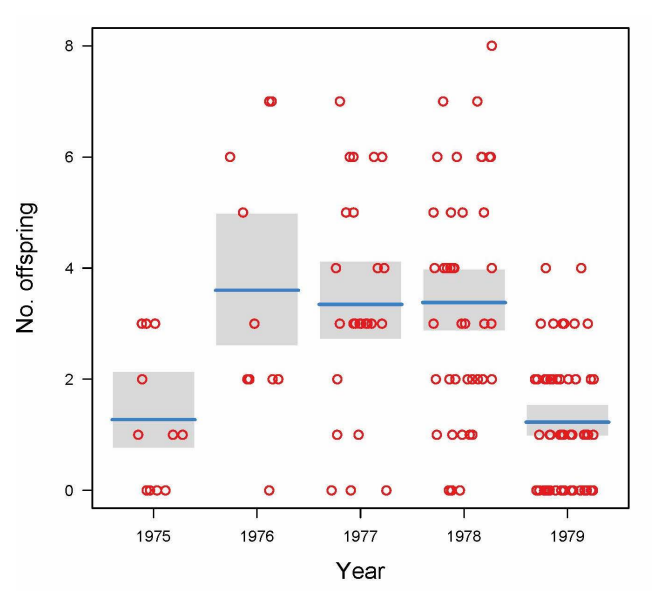
Predicted values on the original scale

Predicted values on original scale: fitted.values(z)

$$\hat{\mu} = e^{\hat{\eta}}$$

I have plotted them here using visreg() and superimposed the original data points.

Note that the fitted values aren't the means of the original data. Fitted values are the transformed means estimated on the log scale (geometric means).



Use emmeans () to obtain model predicted values on original scale

```
emmeans(z, "year", type = "response")
```

Yields model-fitted predicted values and approximate 95% confidence intervals.

year	rate	SE	df	asymp.LCL	asymp.UCL
1975	1.272727	0.3401496	Inf	0.7537770	2.148957
1976	3.600000	0.6000000	Inf	2.5967825	4.990791
1977	3.346154	0.3587453	Inf	2.7119865	4.128614
1978	3.380952	0.2837232	Inf	2.8681893	3.985385
1979	1.228070	0.1467822	Inf	0.9715952	1.552248

Uses a large-sample approximation (degrees of freedom (df) are shown as infinite), and confidence limits might not be accurate for small sample sizes.

Use anova() to test hypotheses

Analysis of deviance table gives log-likelihood ratio test of the null hypothesis that there is no differences among years in mean number of offspring.

```
anova(z, test="Chisq")

Terms added sequentially (first to last)

Df Deviance Resid. Df Resid. Dev P(>|Chi|)

NULL 145 288.656

year 4 75.575 141 213.081 1.506e-15 ***
```

As with lm(), terms are tested using model comparison (always a "full" vs "reduced" model). Default program of action is to fit terms sequentially ("Type 1 sums of squares"), as with lm().

Evaluating assumptions of the glm() fit

Do the variances of the residuals correspond to those assumed by the chosen link function?

The log link function assumes that the Y values are Poisson distributed at each X.

A key property of the Poisson distribution is that within each treatment group the variance and mean are equal (i.e., the glm() dispersion parameter = 1).

Similarly, when analyzing binary data, the logit link function also assumes a strict mean-variance relationship, specified by binomial distribution, when dispersion parameter = 1. This is rarely violated with 0/1 data, so I focus on log-linear model here.

Evaluating assumptions of the glm() fit

A central property of the Poisson distribution is that the variance and mean are equal (i.e., the glm() dispersion parameter = 1).

Let's check the sparrow data:

```
tapply(noffspring, year, mean)
tapply(noffspring, year, var)
```

```
1975 1976 1977 1978 1979
1.272727 3.600000 3.346154 3.380952 1.228070 # mean
1.618182 6.044444 3.835385 4.680604 1.322055 # variance
```

Variances slightly, but not alarmingly, larger than means.

Finding excessive variance ("overdispersion") is common when analyzing count data. Excessive variance occurs because variables not included in the model also affect the response variable.

In the workshop we will analyze an example where the problem is more severe than in the case of the song sparrow data here.

Excessive variance can be accommodated with glm() by using a different link function, one that incorporates a dispersion parameter (which must also be estimated). If the estimated dispersion parameter is >> 1 then there is likely excessive variance.

The glm() procedure to accomplish over (or under) dispersion uses the observed relationship between mean and variance rather than an explicit probability distribution for the data. In the case of count data,

variance = dispersion parameter × mean

Method generates "quasi-likelihood" estimates that behave like maximum likelihood estimates.

Dispersion parameter for quasipoisson family taken to be 1.230689

The dispersion parameter is reasonably close to 1 for these data. But typically it is much larger than 1 for count data, so use family = quasipoisson.

Lets try it with the song sparrow data
z <- glm(noffspring ~ year, family = quasipoisson)</pre>

summary(z)

```
Estimate Std. Error t value Pr(>|t|) Intercept) 0.24116 0.29649 0.813 0.41736 year1976 1.03977 0.34942 2.976 0.00344 ** year1977 0.96665 0.31946 3.026 0.00295 ** year1978 0.97700 0.31076 3.144 0.00203 ** year1979 -0.03572 0.32479 -0.110 0.91259
```

Dispersion parameter for quasipoisson family taken to be 1.230689

The point estimates are identical with those obtained using family=poisson instead, but the standard errors (and resulting confidence intervals) are wider.

Other uses of generalized linear models

We have used glm() to model binary frequency data, and count data.

The method is commonly used to model r x c (and higher order) contingency tables, in which cell counts depend on two (or more) categorical variables each of which may have more than two categories or groups (see R tips "Fit model" page).

glm() can handle data having other probability distributions than the ones used in my examples, including exponential and gamma distributions.

Discussion paper for next week:

Whittingham et al (2006) Why do we still use stepwise modelling?

Download from "assignments" tab on course web site.

Presenters:

Moderators: