Modeling binary data

Analysis of Ecological and Environmental Data

QERM 514

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Reminder

Your project plan is due today

Goals for today

- Understand the characteristics of binary data and the Bernoulli distribution
- Understand how to model binary data with logistic regression
- Understand approaches to inference in logistic regression
- Understand diagnostic measures for logistic regression

The Bernoulli distribution describes the probability of a single "event" y_i occurring

- present (1/1) or absent (0/1)
- alive (1/1) or dead (0/1)
- mature (1/1) or immature (0/1)

Binomial distribution

The binomial distribution is closely related to the Bernoulli

It describes the number of k "successes" in a sequence of n independent Bernoulli "trials"

For example, the number of heads in 4 coin tosses

Binomial distribution

For a population, these could be

- k survivors out of n tagged individuals
- \cdot k infected individuals out of n susceptible individuals
- *k* counts of allele A in *n* total chromosomes

Binomial distribution

The probability mass function

$$Pr(k; n, p) = \binom{n}{k} p^k (1 - p)^{n-k}$$
$$\binom{n}{k} = \frac{n!}{k!(n-k)!}$$

Special case of binomial with n = 1

$$\Pr(k; p) = p^{k} (1 - p)^{(1 - k)}$$

$$\downarrow \downarrow$$

$$k = \begin{cases} 1 \text{ if success (T, Y) with probability } p \\ 0 \text{ if failure (F, N) with probability } (1 - p) \end{cases}$$

where

$$Mean(k) = p$$
 $Var(k) = p(1 - p)$

Likelihood

$$\mathcal{L}(k;p) = \prod_{i=1}^{n} p^{k_i} (1-p)^{(1-k_i)}$$

$$\downarrow \downarrow$$

$$\log \mathcal{L}(k;p) = \log p \sum_{i=1}^{n} k_i + \log(1-p) \sum_{i=1}^{n} (1-k_i)$$

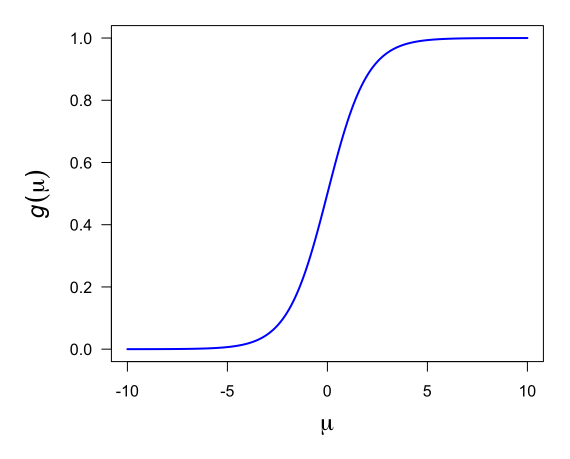
Canonical link is the logit

$$\log\left(\frac{\mu}{1-\mu}\right) = \mathbf{X}\boldsymbol{\beta}$$

$$\psi$$

$$\mu = \frac{\exp(\mathbf{X}\boldsymbol{\beta})}{1+\exp(\mathbf{X}\boldsymbol{\beta})}$$

Logit link



Logistic regression

Similar to other regression in that we assume

- the predictors are linear
- · the observations are independent of one another
- no(ish) multicollinearity among predictors

Logistic regression

Different from other regression in that

- the response is binary
- the relationship between response and predictors is often non-linear
- · the errors can be non-normal
- the errors can be heteroscedastic

Logistic regression is a GLM

We need 3 things to specify our GLM

- 1. Distribution of the data: $y \sim \text{Bernoulli}(p)$
- 2. Link function: $logit(p) = log(\frac{p}{1-p}) = \eta$
- 3. Linear predictor: $\eta = \mathbf{X}\boldsymbol{\beta}$

Logistic regression

The probability of a success is given by

$$p = \frac{\exp(\mathbf{X}\boldsymbol{\beta})}{1 + \exp(\mathbf{X}\boldsymbol{\beta})}$$

$$= \frac{1}{1 + \exp(-\mathbf{X}\boldsymbol{\beta})}$$

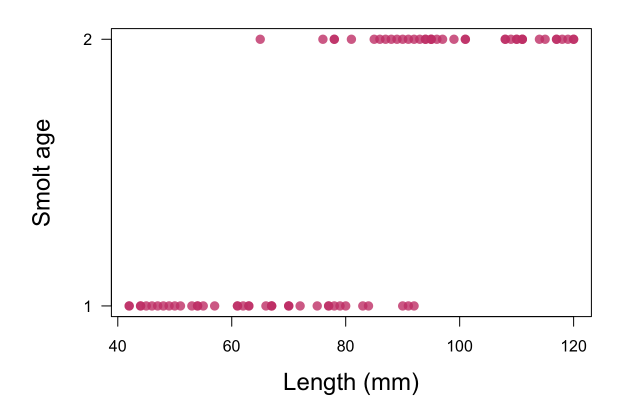
Logistic regression

Example

Sockeye salmon are born in freshwater and rear there for some time before migrating to the ocean as *smolts*

The age at which sockeye smolt can vary from 1 to 2 years, which is thought to depend on their body size

Let's examine the relationship between fish length and its probability of smolting at age-2 instead of age-1



In R we use glm() to fit logistic regression models (and other GLMs)

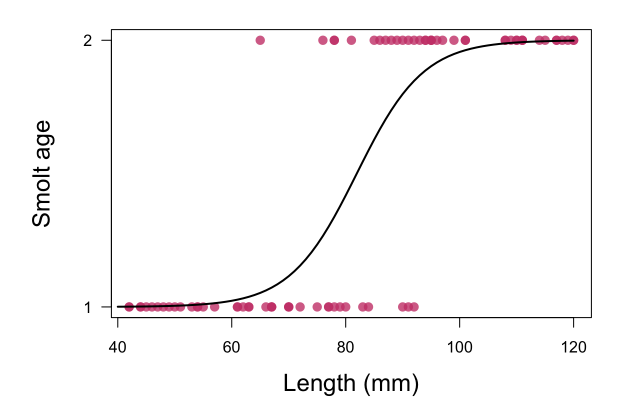
The probability of smolting at age-2 is given by

$$p_i = \frac{1}{1 + \exp(-\mathbf{X}_i \boldsymbol{\beta})}$$

$$\approx \frac{1}{1 + \exp(14 - 0.17L_i)}$$

We can get the fitted values with predict()

```
## get fitted values
newdata <- data.frame(length = seq(40, 120))
p_hat <- 1 / (1 + exp(-predict(fit_mod, newdata)))</pre>
```



The length at which the probability of smolting at age-2 is 0.5

$$0.5 = \frac{1}{1 + \exp(14 - 0.17L_{0.5})}$$

$$2 = 1 + \exp(14 - 0.17L_{0.5})$$

$$1 = \exp(14 - 0.17L_{0.5})$$

$$0 = 14 - 0.17L_{0.5}$$

$$0.17L_{0.5} = 14$$

$$L_{0.5} \approx 82 \text{ mm}$$

Logistic regression and odds

We have talked a bit about odds with respect to evidence ratios

Odds o are an unbounded alternative to probability p

If we represent the k-to-1 odds against something as 1/k, then the following holds

$$o = \frac{1}{1 - p} \implies p = \frac{o}{1 + o}$$

Logistic regression and odds

$$\log\left(\frac{p}{1-p}\right) = \mathbf{X}\boldsymbol{\beta}$$

$$\downarrow \mathbf{b}$$

$$\log(\text{odds}) = \mathbf{X}\boldsymbol{\beta}$$

$$\downarrow \mathbf{b}$$

$$0 \text{odds} = \exp(\mathbf{X}\boldsymbol{\beta})$$

$$\log\left(\frac{p}{1-p}\right) = -14 + 0.17L$$

$$\downarrow \qquad \qquad \downarrow$$

$$\log(\text{odds}) = -14 + 0.17L$$

A unit increase in L increases the log-odds by 0.17

$$\log\left(\frac{p}{1-p}\right) = -14 + 0.17L$$

$$\downarrow 0 \log(\text{odds}) = -14 + 0.17L$$

$$\downarrow 0 \log(\text{odds}) = \exp(-14 + 0.17L)$$

A unit increase in L increases odds by $\exp(0.17) \approx 1.19 = 19\%$

QUESTIONS?

Inference

Consider 2 models, A & B, such that B is a subset of A

$$A = f(x_1, x_2)$$

$$\mathsf{B} = g(x_1)$$

We have seen that we can compare A & B via a likelihood ratio test

$$\lambda = -2 \log \frac{\mathcal{L}_A}{\mathcal{L}_B} \sim \chi_{df=k_A-k_B}^2$$

Log-likelihood

The log-likelihood using a logit link is

$$\log \mathcal{L}(k; p) = \log p \sum_{i=1}^{n} k_i + \log(1-p) \sum_{i=1}^{n} (1-k_i)$$

Deviance

Deviance D is a goodness-of-fit statistic

It's a generalization of using the sum-of-squares of residuals in ordinary least squares to cases where model-fitting is achieved by maximum likelihood

$$D = -2\log \mathcal{L}$$

Deviance for logistic regression

$$D = -2 \left[\log p \sum_{i=1}^{n} k_i + \log(1-p) \sum_{i=1}^{n} (1-k_i) \right]$$
$$= -2 \sum_{i=1}^{n} \left[p_i \operatorname{logit}(p_i) + \log(1-p_i) \right]$$

Likelihood ratio test

$$\lambda = -2\log \frac{\mathcal{L}_A}{\mathcal{L}_B} \sim \chi_{df=k_A-k_B}^2$$

$$\downarrow \downarrow$$

$$\lambda = -2(\log \mathcal{L}_A - \log \mathcal{L}_B) \sim \chi_{df=k_A-k_B}^2$$

$$\downarrow \downarrow$$

$$\lambda = D(B) - D(A) \sim \chi_{df=k_A-k_B}^2$$

The output from glm() includes the deviances for the full model and a null model with no predictors

Likelihood ratio test for $H_0: \beta_1 = 0$

```
## deviance of full model
D_full <- summary(fit_mod)$deviance
## deviance of null model
D_null <- summary(fit_mod)$null.deviance
## test statistic
lambda <- D_null - D_full
## LRT with df = 1
(p_value <- pchisq(lambda, 1, lower.tail = FALSE))</pre>
```

```
## [1] 1.062116e-16
```

Model selection via AIC

$$AIC = 2k - 2\log \mathcal{L}$$
$$= 2k + D$$

```
## AIC
AIC(fit_mod)
## AIC via likelihood
(2 * 2) - 2 * logLik(fit_mod)
## AIC via deviance
(2 * 2) + summary(fit_mod)$deviance
```

```
## [1] 46.05294
## 'log Lik.' 46.05294 (df=2)
## [1] 46.05294
```

Smolt age versus length

Compare to a null model with no predictors

Model selection via AIC

```
## difference in AIC
AIC(fit_null) - AIC(fit_mod)
```

[1] 66.85061

Significance test for β_i

An alternative to the χ^2 test is a z test

$$z = \frac{\hat{\beta_i}}{\text{SE}(\hat{\beta_i})} \sim z_{\alpha/2}$$

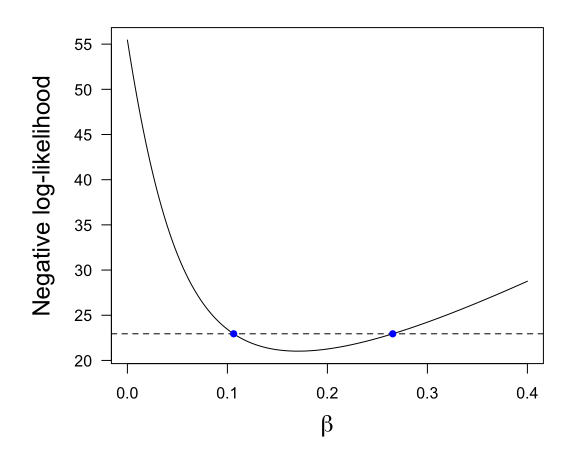
Significance test for β_i

We can also compute a 100(1 - α)% confidence interval

$$\hat{\beta}_i \pm z_{\alpha/2} SE(\hat{\beta}_i)$$

```
## beta
beta_1 <- coef(fit_mod)[2]
## SE of beta
se_beta_1 <- sqrt(diag(vcov(fit_mod)))[2]
## 95% CI
beta_1 + c(-1,1) * 1.96 * se_beta_1
## [1] 0.09266613 0.24862556</pre>
```

Due to possible bias in $SE(\beta)$, we can compute Cl's based on the *profile* likelihood



We can compute CI's based on the profile likelihood with confint()

```
## 95% CI via profile likelihood
confint(fit_mod)

## Waiting for profiling to be done...

## 2.5 % 97.5 %

## (Intercept) -21.8553251 -8.6351047

## length 0.1062832 0.2653229
```

Model diagnostics

As with other models, it's important to examine diagnostic checks for our fitted models

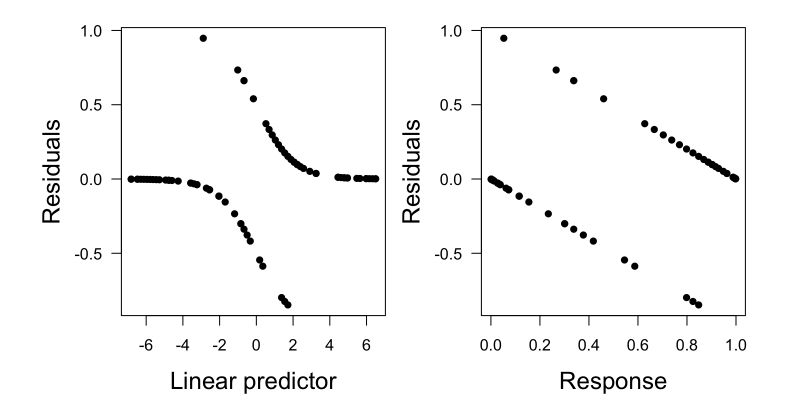
Residuals

We usually think about residuals \emph{e} as

$$e = y - \hat{y}$$

With logistic regression, the response can take 1 of 2 possible values

Residuals



Deviance residuals

We can instead use the *deviance residuals*

$$e_i = (2y_i - 1)D_i$$

$$2y - 1$$
 is 1 (-1) if y is 1 (0)

This is the default for residuals()

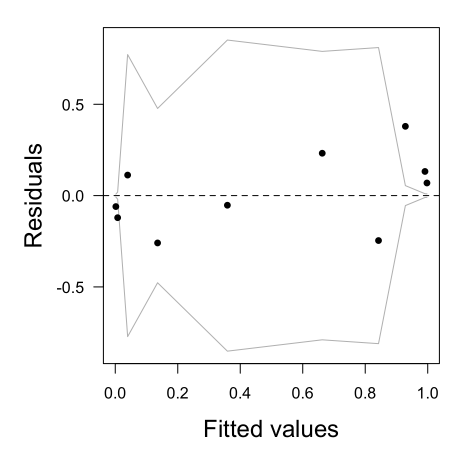
Deviance residuals

We then place the deviance residuals into bins for easier inspection

- Sensitive to the number of bins (~30/bin is good)
- Mean of *e* not constrained to 0
- Check to see that ~95% of points fall within the CI

Can use binnedplot() from the arm package to do this

Deviance residuals

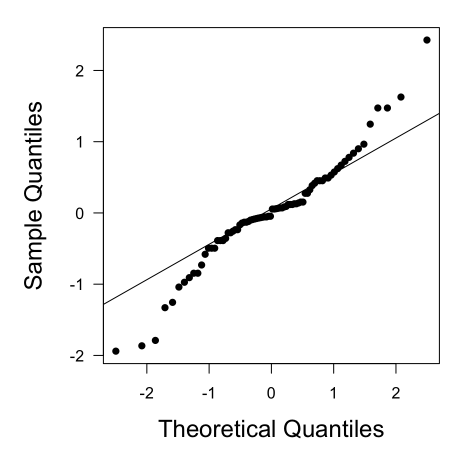


Q-Q plots

We can examine a Q-Q plot, but there is no assumption that the e are normal

It can help to identify unusual points

Q-Q plots



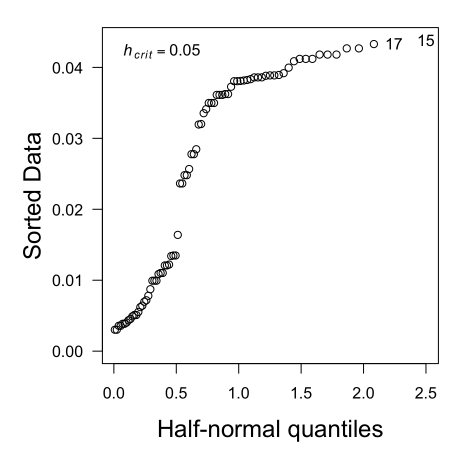
Leverage

We can also calculate the leverages h to look for unusual observation in $predictor\ space$

Recall we are potentially concerned about $h > 2\frac{k}{n}$

We can use faraway::halfnorm()

Leverage



Cook's Distance

Recall that we can use Cook's D to identify potentially influential points

$$D_i = e_i^2 \frac{1}{k} \left(\frac{h_i}{1 - h_i} \right)$$

Cook's Distance

