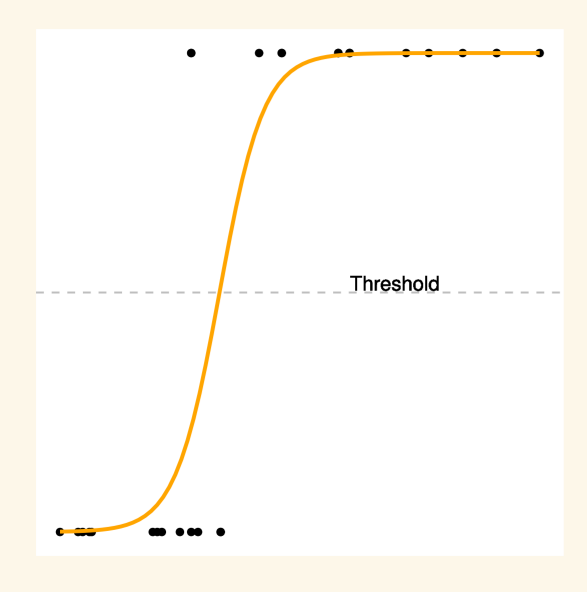
Binomial Logistic regression: deviance

Stat 230

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Overview



Today:

Deviance

Assumptions

Residuals and case influence

Review: A Logistic Regression Model for Binomial Count Data

For all $i=1,\ldots,n$,

$$y_i \sim ext{binomial}(m_i, \pi_i),$$

where m_i is a known number of trials for observation i,

$$\pi_i = rac{\expig(oldsymbol{x}_i'oldsymbol{eta}ig)}{1+\expig(oldsymbol{x}_i'oldsymbol{eta}ig)},$$

and y_1, \ldots, y_n are independent.

Overview: Binomial Distribution

Recall that for $y_i \sim \operatorname{binomial}(m_i, \pi_i)$, the probability mass function of y_i is

$$P\left(y_i=y
ight) = egin{cases} inom{m_i}{y} \pi_i^y (1-\pi_i)^{m_i-y} & ext{for } y \in \{0,\ldots,m_i\} \ 0 & ext{otherwise} \end{cases} \ E\left(y_i
ight) = m_i \pi_i, & ext{and } ext{Var}(y_i) = m_i \pi_i \left(1-\pi_i
ight). \end{cases}$$

The binomial log likelihood function is

$$\ell(oldsymbol{eta} \mid oldsymbol{y}) = \sum_{i=1}^n \left[y_i \log\!\left(rac{\pi_i}{1-\pi_i}
ight) + m_i \log(1-\pi_i)
ight] \ + ext{constant}$$

Deviance for Binomial responses

With Binomial responses, the likelihood function is

$$L(eta) = \prod_{i=1}^n \left(rac{m_i}{y_i}
ight) \pi(X_i)^{y_i} (1-\pi\left(X_i
ight))^{m_i-y_i}$$

and the deviance is

$$egin{aligned} G^2 &= 2[\ln L(ar{\pi}) - \ln L(\hat{\pi}(X))] \ &= 2\sum_{i=1}^n \left[y_i \ln\!\left(rac{y_i}{m_i \hat{\pi}\left(X_i
ight)}
ight) + \left(m_i - y_i
ight) \ln\!\left(rac{m_i - y_i}{m_i - m_i \hat{\pi}\left(X_i
ight)}
ight)
ight] \end{aligned}$$

- $L(\hat{\pi}(X))$: likelihood of the data that plugs in estimates $\hat{\pi}(X_i)$ from the logistic model.
- ullet $L(ar{\pi}):$ likelihood of the data that plugs in estimates $ar{\pi}_i=y_i/m_i$

$$L(ar{\pi}) \geq L(\hat{\pi}(X))$$

Logistic Regression Model vs Saturated model

Logistic Regression Model

- $ullet y_i \sim \mathrm{binomial}(m_i, \pi_i)$
- y_1,\ldots,y_n independent

$$\pi_i = rac{\expig(oldsymbol{x}_i'oldsymbol{eta}ig)}{1+\expig(oldsymbol{x}_i'oldsymbol{eta}ig)}$$

- p+1 β parameters
- ullet MLE: $\hat{\pi}_i = rac{\exp\left(oldsymbol{x}_i'\hat{oldsymbol{eta}}
 ight)}{1+\exp\left(oldsymbol{x}_i'\hat{oldsymbol{eta}}
 ight)}$

Saturated Model

- $ullet y_i \sim ext{binomial}(m_i, \pi_i) \ ullet y_1, \dots, y_n ext{ independent}$
- ullet $\pi_i \in [0,1]$ for $i=1,\ldots,n$ with no other restrictions
- *n* parameters
- $\mathsf{MLE} : y_i/m_i$

Deviance for Binomial responses

Deviance for binomial models can be used for two types of hypothesis tests:

- 1. Drop-in-deviance: Used to compare two models, just like in binary logistic models.
- 2. **Goodness-of-fit:** Used to test binomial response model adequacy.

Goodness-of-fit test

Our hypotheses for the GOF test are:

 H_0 : logistic model

 H_A : saturated model

 $H_0: Y_i \mid X_i \sim \mathrm{Binom}(m_i, \pi\left(X_i
ight))$

• $\pi(X_i)$ equals the logistic function of the p predictor terms.

 H_A : the saturated "model"

• uses the n empirical proportion of successes $\bar{\pi}=y_i/m_i$ for each case as the probability of success for all m_i trials.

Goodness-of-fit test

• The test statistic is the residual deviance of the logistic model

$$G^2=2[\ln L(ar{\pi})-\ln L(\hat{\pi}(X))]$$

- If the "fit" (likelihood) of the logistic model is "close" then G^2 is "close" to 0 and we can claim that the logistic model is adequate.
- If H_0 is true and m_i 's are large, G^2 will have an approximate chi-square distribution with n-(p+1) (model) degrees of freedom. The **p-value** is the probability of getting residual deviance values larger than the observed value:

$$p- ext{ value }=1-P\left(\chi^2>G^2
ight)=1- ext{pchisq}ig(G^2,df=n-(p+1)ig)$$

The suggested rule of thumb for "large m " is that we want most m_i 's to be at least 5 .

Goodness-of-fit test conclusions

Do not reject the null: (large p-value)

- Your logistic model is adequate.
- You don't have a large enough sample size *n* to have the power to detect inadequacies in your model.

Reject the null: (small p-value)

- You have outlier(s) that are inflating the residual deviance.
- Your logistic model is inadequate.

Goodness-of-fit test conclusions

Why might a model be inadequate?

- Your log-odds model is inadequate, it is ill-fitting and transformations are needed
- Extra-binomial variation: your response counts aren't well modeled by a Binomial model*.

*This could mean:

- trials are not independent for each case
- probability of success is not constant across trials for each case
- your choice of predictors **isn't** sufficient (i.e. you are missing key explanatory variables)

• **GOF** test hypotheses are

$$H_0: \log(ext{odds}) = eta_0 + eta_1 \log(ext{area}) \ H_A: \log(ext{odds}) = lpha_i(ext{saturated model})$$

We can conduct a GOF test because all our m_i 's (AtRisk), are above 5

island <- case2101
island\$AtRisk</pre>

[1] 75 67 66 51 28 20 43 31 28 32 30 20 31 16 15 33 40 6

Test stat: residual deviance of $G^2 = 12.062$

```
krunnit glm <- glm(Extinct/AtRisk ~ log(Area), family="binomial", weights=AtRisk, data = island)</pre>
summarv(krunnit glm)
Call:
glm(formula = Extinct/AtRisk ~ log(Area), family = "binomial",
   data = island, weights = AtRisk)
Deviance Residuals:
        1Q Median
    Min
                                          Max
                                  30
-1.71726 -0.67722 0.09726 0.48365 1.49545
Coefficients:
           Estimate Std. Error z value Pr(>|z|)
(Intercept) -1.19620 0.11845 -10.099 < 2e-16 ***
log(Area) -0.29710 0.05485 -5.416 6.08e-08 ***
Signif. codes: 0 '*** 0.001 '** 0.01 '* 0.05 '.' 0.1 ' ' 1
(Dispersion parameter for binomial family taken to be 1)
   Null deviance: 45.338 on 17 degrees of freedom
Residual deviance: 12.062 on 16 degrees of freedom
AIC: 75.394
Number of Fisher Scoring iterations: 4
```

• **p-value:** Using 18 - 2 = 16 degrees of freedom

$$p ext{-value} = 1 - P\left(\chi^2 > 12.062\right) = 1 - pchisq(12.062, df = 16) = 0.7397$$

- The large p-value means that we do not reject the null hypothesis.
- Our model for the probability of extinction given log-area looks to be adequate.

```
krunnit_glm_nolog <- glm(Extinct/AtRisk ~ Area, family="binomial", weights=AtRisk, data = is</pre>
summary(krunnit_glm_nolog)
Call:
glm(formula = Extinct/AtRisk ~ Area, family = "binomial", data = island,
   weights = AtRisk)
Deviance Residuals:
   Min 1Q Median 3Q Max
-1.6526 -1.0661 -0.1877 1.0038 2.1860
Coefficients:
            Estimate Std. Error z value Pr(>|z|)
(Intercept) -1.305957 0.117339 -11.130 < 2e-16 ***
        -0.010121 0.002684 -3.771 0.000163 ***
Area
Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
(Dispersion parameter for binomial family taken to be 1)
   Null deviance: 45.338 on 17 degrees of freedom
Residual deviance: 24.661 on 16 degrees of freedom
AIC: 87,993
Number of Fisher Scoring iterations: 4
```

- Regression of extinction on area (not logged) has residual deviance of 24.661.
- The GOF p-value for this model is 0.076, which suggests that this model may not be adequate
 - agrees with our EDA for the log-odds suggests

```
1-pchisq(24.661, df=16)
```

[1] 0.07602884

Checking Assumptions

Binomial logistic model as the **same assumptions** as binary models:

- Independence of cases takes an understanding of how the data was collected.
- Log-odds **linearity** can be checked with an empirical log-odds plot against quantitative predictors and residual plots.

A third assumption is that the counts of successes Y_i has a **binomial distribution**:

- the m_i trials are independent events, and a success or failure for one trial doesn't affect the outcome of another trial, and
- the probability of success $\pi(X_i)$ is the same for all m_i trials.

Checking Assumptions

- If one, or both, of these assumptions is violated, then it often induces extra-binomial variation (a.k.a. **over dispersion**):
- the actual variation $SD\left(Y\mid X_{i}
 ight)$ is larger than the binomial SD of $\sqrt{m_{i}\pi\left(X_{i}
 ight)\left(1-\pi\left(X_{i}
 ight)
 ight)}$
- making our reported standard errors larger and p-values too small
- Check: use the goodness-of-fit test, when m_i are large enough, to check our binomial distribution assumption.

Checking Assumptions

If we do find evidence of lack-of-fit in our binomial model, then you should

- Check deviance residuals as case influence stats to see if an outlier(s) is affecting GOF results.
- Check the log odds form, change model structure, see if transformations of quantitative predictors are needed

If outliers and transformations aren't a concern, then consider an alternative model:

- binary logistic model if trial-level predictors are available
- quasi-binomial logistic model
- a model that allows for correlated trials (like a mixed-effects logistic model)

How might the Krunnit Island extinction counts violate the binomial counts model assumptions?

Independence:

- This assumption implies that the extinction, or not, of all at risk species on an island are independent events.
- This could be violated if the extinction of one species makes the extinction of a second more likely.

Probability:

- This assumption implies that the probability π (area\$\left._{i}\right)\$ of extinction on island i is the same for all at risk species on island i.
- This could be violated if, for example, species living primarily on the interior of the island had a lower chance of extinction than species living on the coastal region.

Residuals

Pearson residuals are basically response residuals standardized based on the binomial SD:

$$pr_{i}=rac{y_{i}-m_{i}\hat{\pi}\left(X_{i}
ight)}{\sqrt{m_{i}\hat{\pi}\left(X_{i}
ight)\left(1-\hat{\pi}\left(X_{i}
ight)
ight)}}$$

- resid(my_glm, type = "pearson")
- augment(my_glm, type.residuals = "pearson")

Residuals

Deviance residuals are each case's contribution to the residual deviance, with a \pm based on whether we over- or under-estimate a case's response (the \pm is denoted by $\operatorname{sign}(y_i - m_i \hat{\pi}(X_i))$):

$$ext{Dres}_i = ext{sign}(y_i - m_i \hat{\pi}\left(X_i
ight)) \sqrt{2\left[y_i \ln\!\left(rac{y_i}{m_i \hat{\pi}\left(X_i
ight)}
ight) + \left(m_i - y_i
ight) \ln\!\left(rac{m_i - y_i}{m_i - m_i \hat{\pi}\left(X_i
ight)}
ight)
ight]}$$

- resid(my_glm, type = "deviance")
- augment(my_glm, type.residuals = "deviance")

Residuals

- Pearson residuals are "easy" to interpret
- Deviance residuals are good to check if you find significant results in a GOF test.
- When m_i 's are large (at least 5), both types of residuals should be similar in value and have a N(0,1) distribution (approximately).
- Regardless of size of m_i , we should plot residuals vs. quantitative predictors to assess linearity of the log odds.

use the augment command to get both sets of residuals

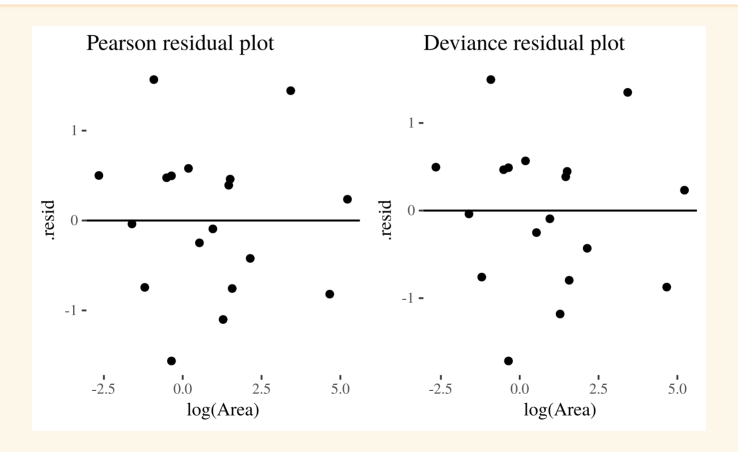
```
island_aug <- augment(krunnit_glm, type.residual="pearson")

plotA <- ggplot(island_aug, aes(x=`log(Area)`, y=.resid)) +
   geom_point() +
   geom_hline(yintercept = 0) +
   labs(title="Pearson residual plot")

island_aug <- augment(krunnit_glm, type.residual="deviance")

plotB <- ggplot(island_aug, aes(x=`log(Area)`, y=.resid)) +
   geom_point() +
   geom_hline(yintercept = 0) +
   labs(title="Deviance residual plot")</pre>
```

```
library(gridExtra)
grid.arrange(plotA, plotB, ncol=2)
```

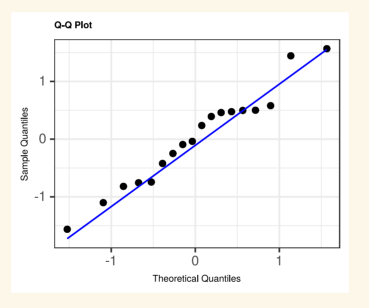


Residuals using ggResidpanel

- resid_xpanel(my_glm, type =): where type could be pearson or deviance (or response)
- resid_panel(my_glm, plots = "qq", type =): qq plot of residuals given by type

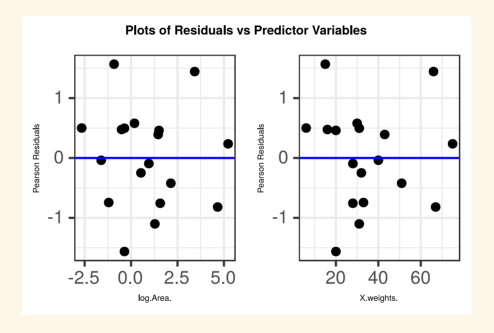
At risk counts m_i are rather large (all cases are 6 or larger) so residuals should be approximately N(0,1).

resid_panel(krunnit_glm, plots = "qq", type = "pearson", axis.text.size = 6, title.text.size = 6)



at risk counts m_i are the X.weights variable (we don't want residual values to depend on m_i)

resid_xpanel(krunnit_glm, type = "pearson", axis.text.size = 4, title.text.size = 6)



Case influence stats

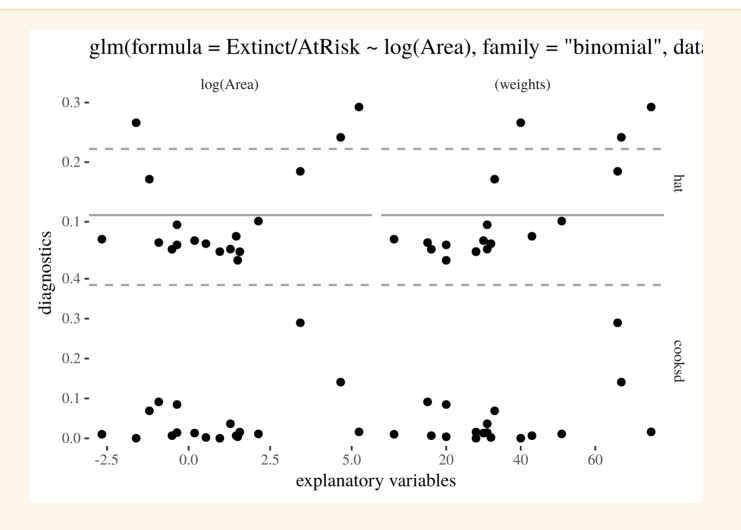
In a GLM, leverage measures

- both a cases's "extremeness" in terms of it's predictor values and
- the size of a case's weight m_i

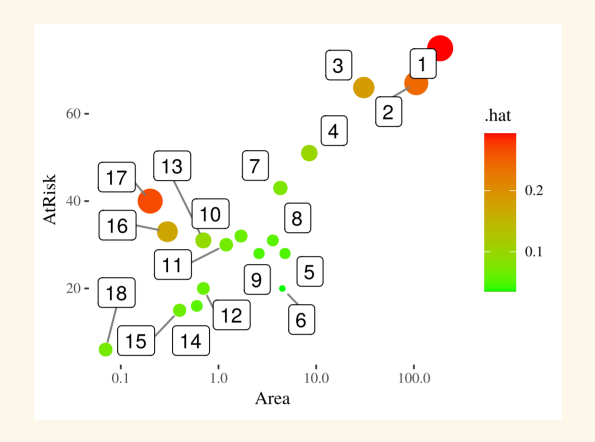
- cases with high values of m_i are given more weight, and hence higher leverage, in the fitted model
- Cook's distance also takes into account a cases leverage (measured both by predictor values and by m_i size) and a case's residual value.

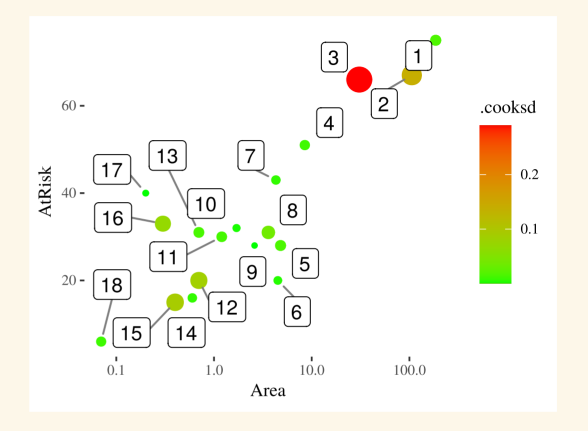
 rows 1,2 and 17 have largest leverage while case 3 looks to have the highest Cook's distance value. plot(krunnit_glm, which=5, id.n=18)

```
library(GGally)
ggnostic(krunnit_glm, columnsY = c(".hat",".cooksd"))
```



```
island_aug <- augment(krunnit_glm, data=island, type.predict = "response")</pre>
island_aug <- island_aug %>% mutate(ID = row_number())
island_aug %>% slice(1,2,3,17) %>% select(-.sigma)
# A tibble: 4 × 10
 Island Area AtRisk Extinct .fitted .resid .std.resid .hat .cooksd
                                                                    TD
 <fct> <dbl> <int>
                       <int> <dbl> <dbl> <dbl> <dbl> <dbl> <int>
1 Ulkokrunni 186.
                   75
                           5 0.0602 0.233 0.277 0.293 1.63e-2
2 Maakrunni 106. 67 3 0.0704 -0.874 -1.00 0.242 1.41e-1
                                                                     2
3 Ristikari 30.7 66 10 0.0985 1.35 1.49 0.185 2.89e-1
                       13 0.328 -0.0381 -0.0445 0.266 3.59e-4
4 Tiirakari 0.2 40
                                                                    17
```

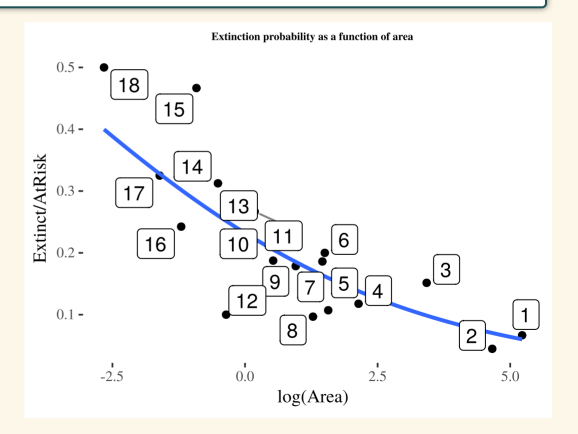




Case 1 (Ulkokrunni):

- largest area and the largest number at risk m_i .
- has largest leverage but it doesn't have an large residual value so it doesn't have high Cook's distance.

```
library(ggrepel)
island <- island %>% mutate(ID = row_number())
plot <- ggplot(island, aes(x=log(Area),</pre>
                         v = Extinct/AtRisk.
                         weight = AtRisk)) +
geom_point() +
 geom_smooth(method="glm", se=FALSE,
method.args = list(family="binomial")) +
 labs(title="Extinction probability as a functi
theme(plot.title = element text(hjust=0.5, siz
                                  face='bold'))
      geom_label_repel(aes(label = ID),
plot +
                  box.padding = 0.15,
                  point.padding = 0.3,
                  segment.color = 'grey50')
```



Case 17 (Tiirakari)

- second smallest area but a large number of at risk species given its small size.
- has larger leverage than case 18 which has the smallest area but smaller number at risk.
- has a small residual and low Cook's distance.

Case 3 (Ristikari)

- third largest area (30.7) but it's number at risk (66) is only one smaller than case 2 (Maakrunni) which is the second largest area (106).
- has a much larger residual than case 2, which results in it having the highest Cook's distance value in the data set.

• None of these cases is overly influential in the model it and removal of case 3, the highest Cook's distance, changes the estimate of β_1 from 0.30 to 0.33 and it's significance doesn't change.