# Wednesday, Mar 22

# Over-dispersion

Over-dispersion can occur for generalized linear models that assume a Poisson or binomial distribution for the response variable.

When we specify a distribution in a generalized linear model, what we are actually specifying is the *variance* structure

$$Var(Y_i) = \phi V[E(Y_i)],$$

where  $\phi$  is the dispersion parameter and V is the variance function.

Over-dispersion is when

$$Var(Y_i) > \phi V[E(Y_i)],$$

and underdispersion is when

$$Var(Y_i) < \phi V[E(Y_i)].$$

Over-dispersion is fairly common in practice, but under-dispersion is relatively rare.

### over-dispersion in Poisson Regression

If  $Y_i$  has a *Poisson* distribution, then

$$Var(Y_i) = E(Y_i),$$

so that it is implicitly assumed that  $\phi = 1$  and V(z) = z. Over-dispersion occurs if

$$Var(Y_i) > E(Y_i).$$

#### over-dispersion in Binomial Regression

If  $C_i$  has a binomial distribution, and  $Y_i = C_i/m_i$ , then

$$Var(Y_i) = E(Y_i)[1 - E(Y_i)]/m_i,$$

so that it is implicitly assumed that  $\phi = 1$  and  $V(z) = z(1-z)/m_i$ . over-dispersion occurs if

$$Var(Y_i) > E(Y_i)[1 - E(Y_i)]/m_i$$
.

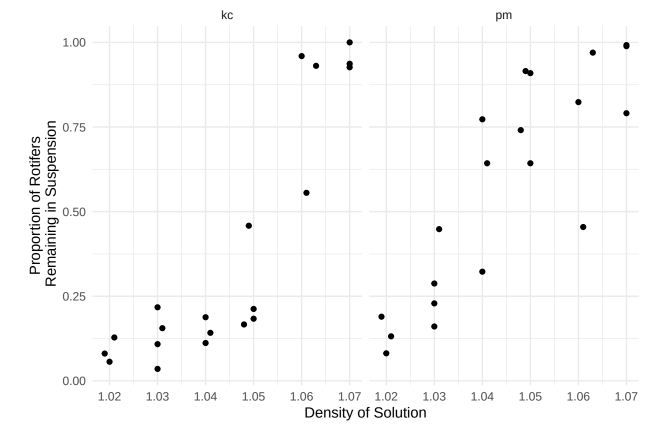
In general, failing to account for over-dispersion (or a misspefication of the variance structure in general) may yield incorrect standard errors (usually too small in the case of over-dispersion), leading to incorrect test statistics and confidence intervals.

## Causes of over-dispersion

- 1. Wrong assumed distribution for the response variable.
- 2. Unobserved explanatory variables that vary over observations.

Note: A misspecified *mean structure* (e.g., failing to transform an explanatory variable or omitting a strong interaction) may appear as overdisperson.

**Example**: Consider the following data from an experiment that investigated the proportion of rotifers of two species remaining in suspension in different solution densities after being put into a centrifuge.

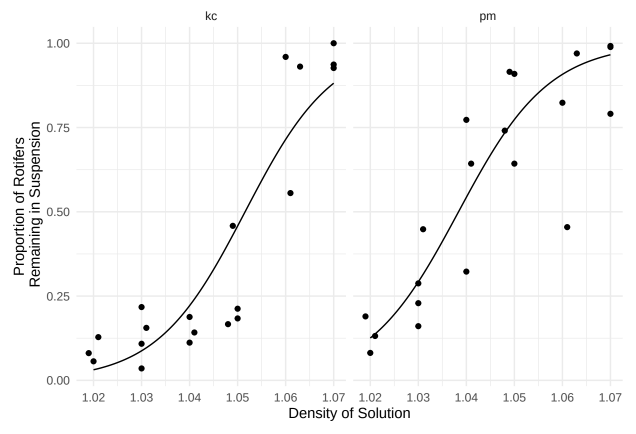


Logistic regression might be a reasonable model here.

```
m <- glm(cbind(y, total - y) ~ species * density,
    family = binomial, data = myrotifer)

d <- expand.grid(species = c("kc","pm"), density = seq(1.02, 1.07, length = 100))
d$yhat <- predict(m, newdata = d, type = "response")

p <- p + geom_line(aes(y = yhat), data = d)
plot(p)</pre>
```



Do these data exhibit over-dispersion for this model?

# Detection of over-dispersion

Standardized residuals can be used to detect over-dispersion. There are several types for GLMs.

1. Pearson residuals. Pearson residuals are defined as

$$\frac{y_i - \hat{y}_i}{\sqrt{\widehat{\operatorname{Var}}(Y_i)}}.$$

Dividing a Pearson residual by another term to account for the variance  $\hat{y}_i$  creates a *standardized* Pearson residual. These are obtained using rstandard(m, type = "pearson").

2. Deviance residuals. The residual deviance can be decomposed into a per-observation contribution so that  $D = \sum_{i=1}^{n} d_i$ . Then the residual deviance is defined as

$$sign(y_i - \hat{y}_i)\sqrt{d_i},$$

where

$$sign(z) = \begin{cases} 1, & \text{if } z > 0, \\ 0, & \text{if } z = 0, \\ -1, & \text{if } z < 0. \end{cases}$$

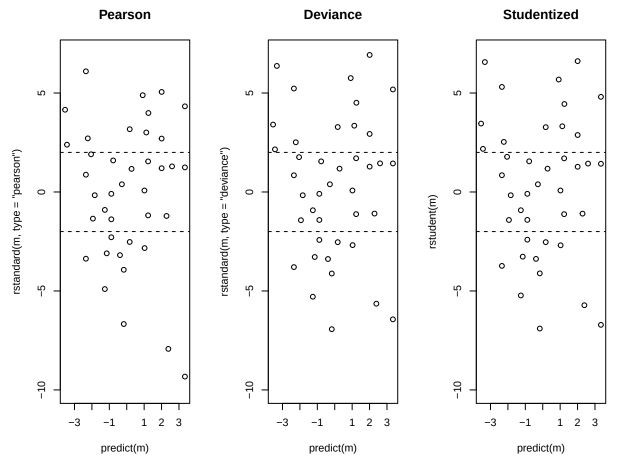
Dividing a deviance residual by another term to account for the variance  $\hat{y}_i$  creates a *standardized* deviance residual. These are obtained using rstandard(m, type = "deviance"). A numerical approximation to these residuals obtained when omitting the observation can be obtained using rstudent(m).

3. Studentized residuals. The function rstudent will produce approximate studentized residuals for GLMs.

Comment: If the model is correct the residuals might be approximately normally distributed with a mean of zero and standard deviation of one (i.e., "standard normal"), so an excess of values greater than two (in absolute value) may indicate over-dispersion or some other problem with the model. But with very coarse data (e.g., very small counts in a Poisson regression model or proportions with small  $m_i$  in a logistic regression model), the distribution of these residuals is not approximately normal.

**Example**: Let's look at the residuals for the rotifer model.

```
par(mfcol = c(1,3))
plot(predict(m), rstandard(m, type = "pearson"), ylim = c(-10, 7), main = "Pearson")
abline(h = c(-2,2), lty = 2)
plot(predict(m), rstandard(m, type = "deviance"), ylim = c(-10, 7), main = "Deviance")
abline(h = c(-2,2), lty = 2)
plot(predict(m), rstudent(m), ylim = c(-10, 7), main = "Studentized")
abline(h = c(-2,2), lty = 2)
```



Is there an explanation of the over-dispersion?

Another metric is to compare the residual deviance to the residual degrees of freedom in a GLM with a response variable with either a Poisson or binomial distribution. If the model is (approximately) correct then the ratio of the residual deviance to the residual degrees of freedom is approximately one.

**Example**: Consider the residual deviance and residual degrees of freedom for the rotifer model.

```
summary(m)
```

Call:

#### Deviance Residuals:

```
Min 1Q Median 3Q Max
-6.471 -2.338 0.599 2.438 6.235
```

#### Coefficients:

Estimate S	td. Error	z value	Pr(> z )	
-114.35	4.03	-28.35	<2e-16	***
4.63	6.60	0.70	0.48	
108.75	3.86	28.19	<2e-16	***
-3.08	6.33	-0.49	0.63	
	-114.35 4.63 108.75	-114.35 4.03 4.63 6.60 108.75 3.86	-114.35 4.03 -28.35 4.63 6.60 0.70 108.75 3.86 28.19	4.63 6.60 0.70 0.48 108.75 3.86 28.19 <2e-16

---

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

(Dispersion parameter for binomial family taken to be 1)

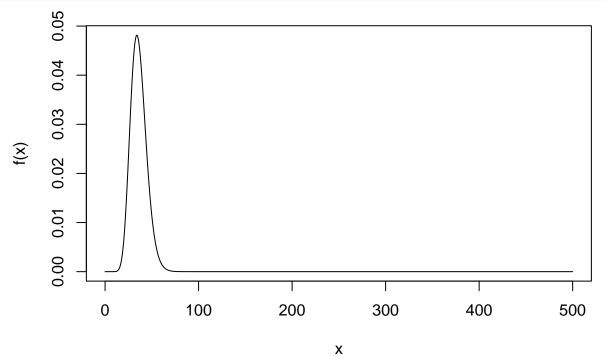
Null deviance: 3180.99 on 39 degrees of freedom Residual deviance: 434.02 on 36 degrees of freedom

AIC: 596.6

### Number of Fisher Scoring iterations: 5

If the model is correct and there is no over-dispersion, the residual deviance has approximate a  $\chi^2$  distribution with degrees of freedom equal to the residual degrees of freedom. We can use this as an informal test for over-dispersion.

```
f <- function(x) dchisq(x, 36)
curve(f, from = 0, to = 500, n = 1000)</pre>
```



```
1 - pchisq(434.02, df = 36)
```

#### [1] 0

Residuals are more informative, but the residual deviance is a quick way to check to see if over-dispersion may be an issue.

Note: For logistic regression, over-dispersion *cannot* be diagnosed in this way for *binary* data (and the residual deviance may not be reliable if the  $m_i$  are very small).

Example: Let's look again a the Poisson regression model for the trawling data.

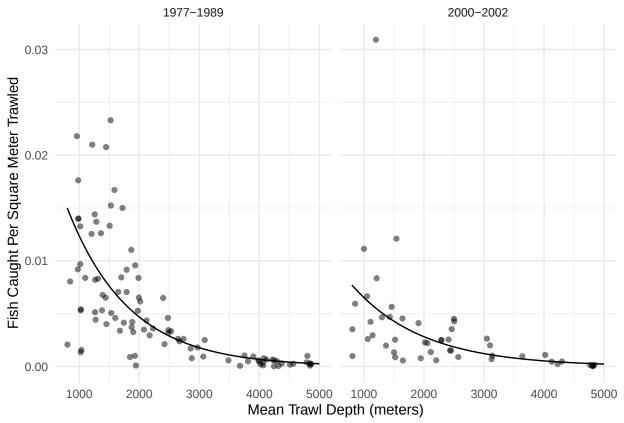
```
library(COUNT)
data(fishing)

m <- glm(totabund ~ period * meandepth + offset(log(sweptarea)),
    family = poisson, data = fishing)

d <- expand.grid(sweptarea = 1, period = levels(fishing$period),
    meandepth = seq(800, 5000, length = 100))

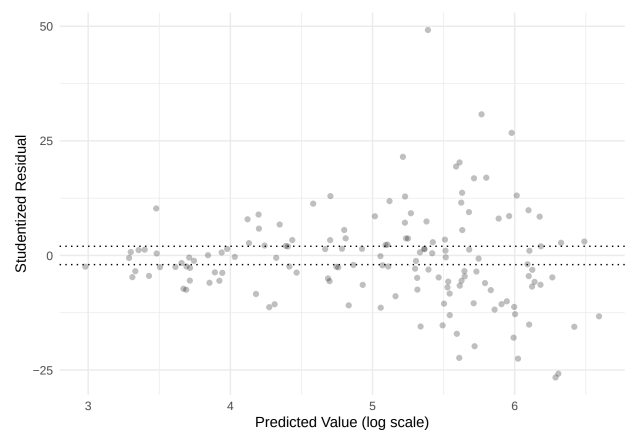
d$yhat <- predict(m, newdata = d, type = "response")

p <- ggplot(fishing, aes(x = meandepth, y = totabund/sweptarea)) +
    geom_point(alpha = 0.5) + facet_wrap(~ period) + theme_minimal() +
    labs(x = "Mean Trawl Depth (meters)",
        y = "Fish Caught Per Square Meter Trawled") +
    geom_line(aes(y = yhat), data = d)
plot(p)</pre>
```



```
summary(m)
```

```
Call:
glm(formula = totabund ~ period * meandepth + offset(log(sweptarea)),
   family = poisson, data = fishing)
Deviance Residuals:
  Min 1Q Median
                         3Q
                                  Max
-26.31 -5.95 -1.64 3.15
                                47.14
Coefficients:
                         Estimate Std. Error z value Pr(>|z|)
(Intercept)
                         -3.42e+00 1.49e-02 -229.67 <2e-16 ***
period2000-2002
                         -7.71e-01 2.97e-02 -25.94 <2e-16 ***
meandepth
                         -9.71e-04 7.96e-06 -121.94 <2e-16 ***
period2000-2002:meandepth 1.32e-04 1.52e-05 8.65 <2e-16 ***
Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
(Dispersion parameter for poisson family taken to be 1)
   Null deviance: 46176 on 146 degrees of freedom
Residual deviance: 14982 on 143 degrees of freedom
AIC: 15962
Number of Fisher Scoring iterations: 5
fishing$eta <- predict(m)</pre>
fishing$res <- rstudent(m)</pre>
p \leftarrow ggplot(fishing, aes(x = eta, y = res)) + theme_minimal() +
 geom_point(alpha = 0.25) +
 labs(x = "Predicted Value (log scale)",
   y = "Studentized Residual") +
 geom_hline(yintercept = c(-2, 2), linetype = 3)
plot(p)
```



over-dispersion is not the only issue here. The variance of the residuals is not constant.

#### Solutions to over-dispersion

There are several potential solutions to over-dispersion.

- 1. Quasi-likelihood. Specify a variance structure other than the one implied by a specified distribution.
- 2. Specify a different distribution (possibly outside the exponential family).
- 3. Use a robust estimator of the standard errors.

#### Quasi-Likelihood Solutions to over-dispersion

The Poisson and binomial distributions assume the variance structures

$$Var(Y_i) = \phi E(Y_i)$$
 and  $Var(Y_i) = \phi E(Y_i)[1 - E(Y_i)]/m_i$ ,

respectively, where the dispersion parameter is fixed at  $\phi = 1$ . One solution is to allow  $\phi$  to be an unknown parameter to "relax" the variance structure and allow the variance to be larger than it would be for a Poisson or binomial distribution. The dispersion parameter can be estimated. R uses

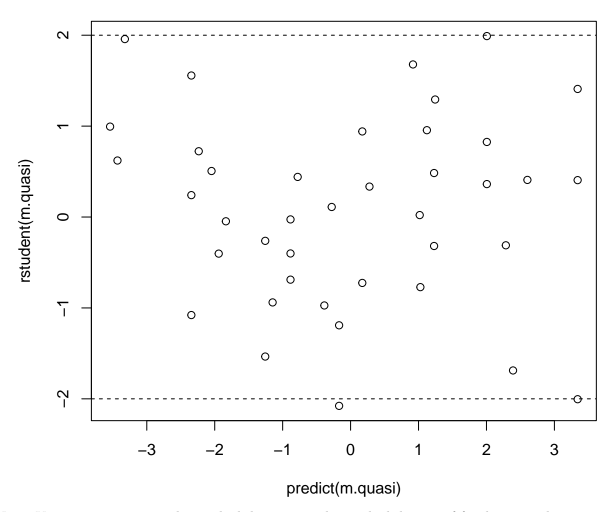
$$\hat{\phi} = \frac{1}{n-p} \sum_{i=1}^{n} \frac{(y_i - \hat{y}_i)^2}{V(\hat{y}_i)},$$

which is analogous to the estimate of  $\sigma^2$  in a normal linear model. This is a quasi-likelihood approach because the variance structures with  $\phi \neq 1$  do not correspond to a binomial or Poisson distribution. This kind of quasi-likelihood can be done with glm by using quasipoisson or quasibinomial instead of poisson or binomial, respectively, when specifying the family argument.

**Example**: Consider again the rotifer model.

```
m.quasi <- glm(cbind(y, total - y) ~ species + density + species:density,
  family = quasibinomial, data = myrotifer)
plot(predict(m.quasi), rstudent(m.quasi), main = "Residual Plot")
abline(h = c(-2,2), lty = 2)</pre>
```

# **Residual Plot**



Note: You **cannot** compare the residual deviance to the residual degrees of freedom as a diagnostic to determine if using quasi-likelihood was successful, but standardized residuals are still appropriate.

How does this impact our inferences?

```
m.binom <- glm(cbind(y, total - y) ~ species + density + species:density,
  family = binomial, data = myrotifer)
cbind(summary(m.binom)$coefficients, confint(m.binom))</pre>
```

```
Estimate Std. Error
                                       z value
                                                  Pr(>|z|)
                                                               2.5 %
                                                                       97.5 %
(Intercept)
                  -114.352
                                 4.034 -28.3454 9.534e-177 -122.420 -106.598
speciespm
                     4.629
                                         0.7016 4.830e-01
                                                              -8.464
                                 6.598
                                                                       17.431
                                        28.1910 7.535e-175
density
                   108.746
                                 3.857
                                                             101.332
                                                                      116.460
speciespm:density
                    -3.077
                                 6.329
                                       -0.4862 6.268e-01
                                                            -15.354
                                                                        9.487
```

cbind(summary(m.quasi)\$coefficients, confint(m.quasi))

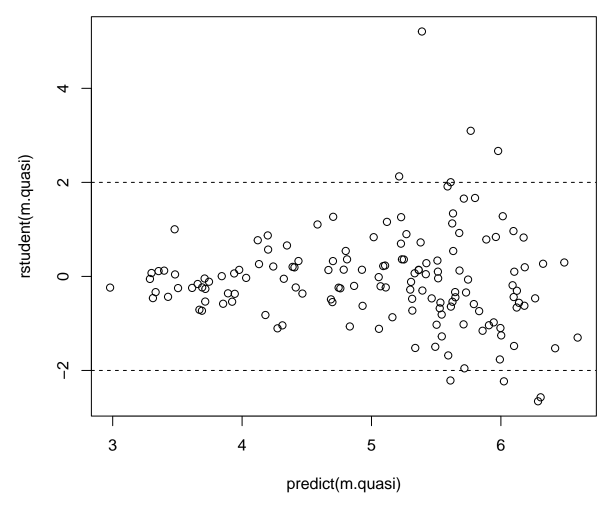
```
Estimate Std. Error t value Pr(>|t|)
                                                         2.5 % 97.5 %
(Intercept)
                  -114.352
                                14.95 -7.6472 4.736e-09 -146.02 -87.01
                                24.46 0.1893 8.509e-01 -46.15 51.31
speciespm
                     4.629
density
                   108.746
                                14.30 7.6056 5.358e-09
                                                        82.60 139.02
speciespm:density
                   -3.077
                                23.46 -0.1312 8.964e-01 -47.81 45.70
# odds ratios for effect of a 0.01 unit increase in density
trtools::contrast(m.binom,
  a = list(species = c("kc","pm"), density = 0.02),
  b = list(species = c("kc", "pm"), density = 0.01),
cnames = c("kc","pm"), tf = exp)
   estimate lower upper
      2.967 2.751 3.200
      2.877 2.607 3.174
# odds ratios for effect of a 0.01 unit increase in density
trtools::contrast(m.quasi,
 a = list(species = c("kc", "pm"), density = 0.02),
  b = list(species = c("kc", "pm"), density = 0.01),
cnames = c("kc","pm"), tf = exp)
   estimate lower upper
      2.967 2.220 3.965
kc
      2.877 1.973 4.195
pm
```

Note that point estimates are unchanged, but standard errors, tests, and confidence intervals are affected.

**Example:** Now let's try the same approach with trawling data.

```
m.quasi <- glm(totabund ~ period * meandepth + offset(log(sweptarea)),
    family = quasipoisson, data = fishing)
plot(predict(m.quasi), rstudent(m.quasi), main = "Residual Plot")
abline(h = c(-2,2), lty = 2)</pre>
```

# **Residual Plot**



That was maybe somewhat less successful. Note the "megaphone" pattern. The assumed variance structure is

$$Var(Y_i) = \phi E(Y_i).$$

We could relax this by assuming instead

$$Var(Y_i) = \phi E(Y_i)^p$$
.

for some p > 1. If p = 1, 2, or 3 then we can use quasi. Here we are using it for p = 2.

```
m.quasi <- glm(totabund ~ period * meandepth + offset(log(sweptarea)),
    family = quasi(link = "log", variance = "mu^2"), data = fishing)
summary(m.quasi)$coefficients</pre>
```

```
Estimate Std. Error t value Pr(>|t|)

(Intercept) -3.250e+00 1.592e-01 -20.4180 3.187e-44

period2000-2002 -6.041e-01 2.720e-01 -2.2212 2.791e-02

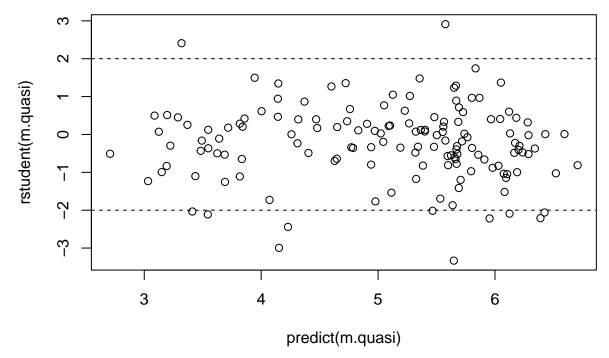
meandepth -1.041e-03 5.866e-05 -17.7403 5.988e-38

period2000-2002:meandepth 7.272e-05 9.992e-05 0.7278 4.679e-01

plot(predict(m.quasi), rstudent(m.quasi), main = "Residual Plot")

abline(h = c(-2,2), lty = 2)
```

# **Residual Plot**



Note that quasi(link = "log", variance = "mu") is the same as quasipoisson. For more options consider family = tweedie. The tweedie family defines power functions for link and variance functions of the form

$$E(Y_i)^q = \eta_i$$
 and  $Var(Y_i) = \phi E(Y_i)^p$ ,

where  $E(Y_i)^0 \equiv \log E(Y_i)$  when using tweedie (not mathematically of course — this is just for interface purposes). For example, to replicate the quasi-likelihood model above we can use the following.

```
library(statmod) # for tweedie "family"
m.tweedie <- glm(totabund ~ period * meandepth + offset(log(sweptarea)),
   family = tweedie(link.power = 0, var.power = 2), data = fishing)
summary(m.tweedie)$coefficients</pre>
```

```
Estimate Std. Error t value Pr(>|t|)
(Intercept) -3.250e+00 1.592e-01 -20.4180 3.187e-44
period2000-2002 -6.041e-01 2.720e-01 -2.2212 2.791e-02
meandepth -1.041e-03 5.866e-05 -17.7403 5.988e-38
period2000-2002:meandepth 7.272e-05 9.992e-05 0.7278 4.679e-01
```

The powers p and q are not required to be integers when using tweedie.

Whether or not we use quasi-likelihood will affect the standard errors, as well as tests and confidence intervals. Failing to account for substantial over-dispersion can result in biased standard errors, and thus incorrect tests and confidence intervals. Estimates of parameters (of functions thereof such as what we get from contrast) may or may not change, depending on the variance structure.

```
m.poisson <- glm(totabund ~ period * meandepth + offset(log(sweptarea)),
    family = poisson, data = fishing)
# rate ratios for year
trtools::contrast(m.poisson,
    a = list(sweptarea = 1, meandepth = c(1000,2000,3000,4000,5000), period = "2000-2002"),
    b = list(sweptarea = 1, meandepth = c(1000,2000,3000,4000,5000), period = "1977-1989"),</pre>
```

```
cnames = c("1000m","2000m","3000m","4000m","5000m"), tf = exp)
      estimate lower upper
1000m
       0.5277 0.5100 0.5460
2000m
       0.6020 0.5861 0.6183
3000m
       0.6869 0.6565 0.7187
4000m
       0.7837 0.7293 0.8421
       0.8941 0.8087 0.9885
5000m
trtools::contrast(m.tweedie,
  a = list(sweptarea = 1, meandepth = c(1000,2000,3000,4000,5000), period = "2000-2002"),
  b = list(sweptarea = 1, meandepth = c(1000, 2000, 3000, 4000, 5000), period = "1977-1989"),
 cnames = c("1000m","2000m","3000m","4000m","5000m"), tf = exp)
      estimate lower upper
       0.5878 0.4046 0.8540
1000m
2000m
       0.6321 0.4869 0.8206
3000m
       0.6798 0.5173 0.8935
4000m
       0.7311 0.4905 1.0897
       0.7863 0.4458 1.3867
5000m
```

### Inferences With Quasi-Likelihood

Using quasi-likelihood instead of maximum likelihood changes how inferences are made in several ways.

- 1. The standard errors are multiplied by  $\sqrt{\hat{\phi}}$ . If  $\hat{\phi} > 1$  (which it probably is if over-dispersion is present) then the standard errors will be larger (and thus failing to account for over-dispersion leads us to usually underestimate them). Note that this adjustment is made automatically when using quasi-likelihood.
- 2. Wald confidence intervals and tests for a single parameter or function of parameters are based on the t distribution rather than the standard normal distribution. The t distribution is believed to provide more accurate results, although it is still an approximation.
- 3. Using confint or anova use the F distribution rather than the  $\chi^2$  distribution. The underlying test statistic is similar to the F test statistic used in normal linear models. When using anova you should use test = "F" rather than test = "LRT" if you are using quasi-likelihood.
- 4. Function in **emmeans** do not adjust the degrees of freedom for estimating the dispersion parameter when using quasi-likelihood. This does not make much difference unless n is small. But you can specify the it manually via the df argument (use the degrees of freedom for the residual deviance from summary or extract it with modelname\$df.residual). But contrast and lincon do not require manual specification, although you can via the df argument for those functions.

```
library(emmeans)
m.quasi <- glm(cbind(y, total - y) ~ species + density + species:density,
  family = quasibinomial, data = myrotifer)
trtools::contrast(m.quasi,
  a = list(species = c("kc","pm"), density = 0.02),
  b = list(species = c("kc","pm"), density = 0.01),
  cnames = c("kc","pm"), tf = exp)

  estimate lower upper
kc   2.967  2.220  3.965
pm   2.877  1.973  4.195

pairs(emmeans(m.quasi, ~density|species, at = list(density = c(0.02, 0.01)),
  type = "response"), infer = TRUE) # wrong df</pre>
```

```
species = kc:
                                         SE df asymp.LCL asymp.UCL null z.ratio p.value
 contrast
                           odds.ratio
 density0.02 / density0.01
                                 2.97 0.424 Inf
                                                     2.24
                                                               3.93
                                                                       1
                                                                           7.606 < .0001
species = pm:
 contrast
                                         SE df asymp.LCL asymp.UCL null z.ratio p.value
                           odds.ratio
 density0.02 / density0.01
                                 2.88 0.535 Inf
                                                     2.00
                                                                4.14
                                                                           5.681 <.0001
Confidence level used: 0.95
Intervals are back-transformed from the log odds ratio scale
Tests are performed on the log odds ratio scale
pairs(emmeans(m.quasi, ~density|species, at = list(density = c(0.02, 0.01)),
type = "response"), infer = TRUE, df = m.quasi$df.residual) # correct df
species = kc:
contrast
                           odds.ratio
                                         SE df lower.CL upper.CL null t.ratio p.value
 density0.02 / density0.01
                                 2.97 0.424 36
                                                   2.22
                                                            3.96
                                                                    1
                                                                       7.606 < .0001
species = pm:
 contrast
                           odds.ratio
                                         SE df lower.CL upper.CL null t.ratio p.value
                                 2.88 0.535 36
 density0.02 / density0.01
                                                   1.97
                                                            4.20
                                                                     1
                                                                         5.681 <.0001
Degrees-of-freedom method: user-specified
Confidence level used: 0.95
Intervals are back-transformed from the log odds ratio scale
Tests are performed on the log odds ratio scale
Admittedly it does not make much difference here.
```

# Misspecified Mean Structures and over-dispersion

A poorly specified mean structure may be mistaken for over-dispersion.

```
library(trtools)
ceriodaphniastrain$strain$strain <- factor(ceriodaphniastrain$strain, labels = c("a", "b"))</pre>
m <- glm(count ~ strain + sqrt(concentration), family = poisson, data = ceriodaphniastrain)</pre>
summary(m)
Call:
glm(formula = count ~ strain + sqrt(concentration), family = poisson,
    data = ceriodaphniastrain)
Deviance Residuals:
          10 Median
                            30
                                   Max
-3.742 -1.098 -0.225 0.938
                                 3.138
Coefficients:
                    Estimate Std. Error z value Pr(>|z|)
(Intercept)
                      4.5284
                                 0.0400 113.08 < 2e-16 ***
strainb
                     -0.2750
                                 0.0484
                                          -5.68 1.3e-08 ***
sqrt(concentration) -1.6576
                                 0.0474 -34.99 < 2e-16 ***
Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
```

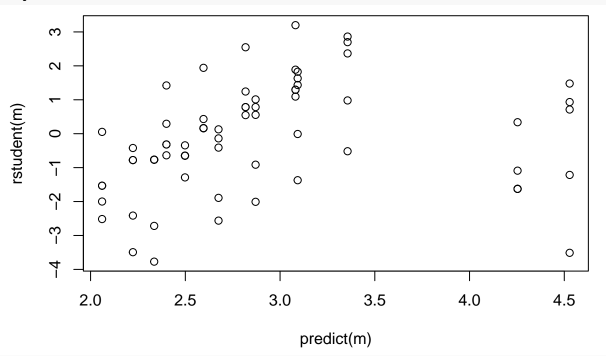
(Dispersion parameter for poisson family taken to be 1)

Null deviance: 1359.38 on 69 degrees of freedom Residual deviance: 164.28 on 67 degrees of freedom

AIC: 493.9

Number of Fisher Scoring iterations: 4

plot(predict(m), rstudent(m))



m <- glm(count ~ strain + concentration, family = poisson, data = ceriodaphniastrain)
summary(m)</pre>

#### Call:

# Deviance Residuals:

Min 1Q Median 3Q Max -2.680 -0.677 0.153 0.679 2.077

#### Coefficients:

Estimate Std. Error z value Pr(>|z|)

(Intercept) 4.4546 0.0391 113.82 < 2e-16 \*\*\*

strainb -0.2750 0.0484 -5.68 1.3e-08 \*\*\*

concentration -1.5431 0.0466 -33.11 < 2e-16 \*\*\*

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

(Dispersion parameter for poisson family taken to be 1)

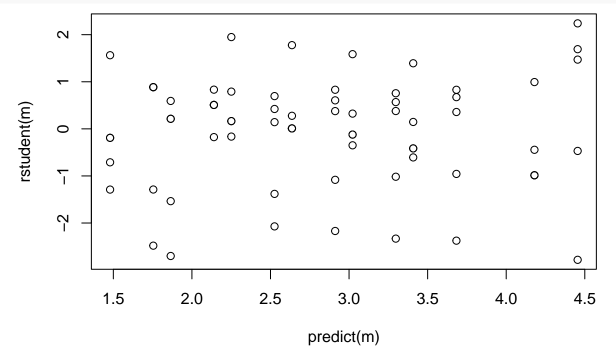
Null deviance: 1359.381 on 69 degrees of freedom

Residual deviance: 86.376 on 67 degrees of freedom

AIC: 416

Number of Fisher Scoring iterations: 4

plot(predict(m), rstudent(m))



# Quasi-Likelihood and Nonlinear Regression

Quasi-likelihood for a GLM is essentially the same as using (nonlinear) regression with iteratively weighted least squares to account for heteroscedasticity. The weights are

$$w_i = \frac{1}{V(\hat{y}_i)},$$

where V is the variance function.

**Example**: Consider the model for the trawling data where the variance is proportional to  $E(Y_i)^2$ . To estimate this model using iteratively weighted least squares we use weights of  $w_i = 1/E(Y_i)^2$ .

```
m.quasi <- glm(totabund ~ period * meandepth + offset(log(sweptarea)),
  family = quasi(link = "log", variance = "mu^2"), data = fishing)
summary(m.quasi)$coefficients</pre>
```

```
fishing$w <- 1 / predict(m.iwls)^2
}
summary(m.iwls)$coefficients</pre>
```

```
Estimate Std. Error t value Pr(>|t|)

b0 -3.250e+00 1.592e-01 -20.4179 3.189e-44

b1 -6.041e-01 2.720e-01 -2.2213 2.790e-02

b2 -1.041e-03 5.866e-05 -17.7402 5.991e-38

b3 7.273e-05 9.992e-05 0.7279 4.679e-01
```

**Example:** Consider the model for the rotifer data. Here the variance is proportional to  $E(Y_i)[1 - E(Y_i)_i]$ . To estimate this model using iteratively weighted least squares we use weights of

$$w_i = \frac{m_i}{E(Y_i)[1 - E(Y_i)]}.$$

```
m.binomial <- glm(cbind(y, total - y) ~ species * density,
    family = quasibinomial, data = myrotifer)
summary(m.binomial)$coefficients</pre>
```

Estimate Std. Error t value Pr(>|t|)

```
14.95 -7.6472 4.736e-09
(Intercept)
                  -114.352
speciespm
                     4.629
                                 24.46 0.1893 8.509e-01
density
                    108.746
                                 14.30 7.6056 5.358e-09
                                 23.46 -0.1312 8.964e-01
speciespm:density
                    -3.077
myrotifer$w <- 1
for (i in 1:20) {
    m <- nls(y/total \sim plogis(b0 + b1*(species == "pm") + b2*density +
      b3*(species == "pm")*density), data = myrotifer, weights = w,
      start = list(b0 = -114, b1 = 4.6, b2 = 109, b3 = -3))
    myrotifer$yhat <- predict(m)</pre>
    myrotifer$w <- myrotifer$total / (myrotifer$yhat * (1 - myrotifer$yhat))</pre>
}
```

summary(m)\$coefficients

Note that plogis is the function  $e^x/(1+e^x)$ . The model can be written as

$$E(Y_i) = \frac{e^{\eta_i}}{1 + e^{\eta_i}}$$

where  $Y_i$  is the observed proportion, and

$$\eta_i = \beta_0 + \beta_1 s_i + \beta_2 d_i + \beta_3 s_i d_i,$$

where  $s_i$  is an indicator variable for the pm species, and  $d_i$  is the density.

Using iteratively weighted least squares is not necessary if we can use quasi or tweedie, but it is a useful option for cases where the variance structure is outside what can be done with quasi or tweedie (although one can *program* new variance structures).