# Friday, Feb 16

## Solutions for Heteroscedasticity

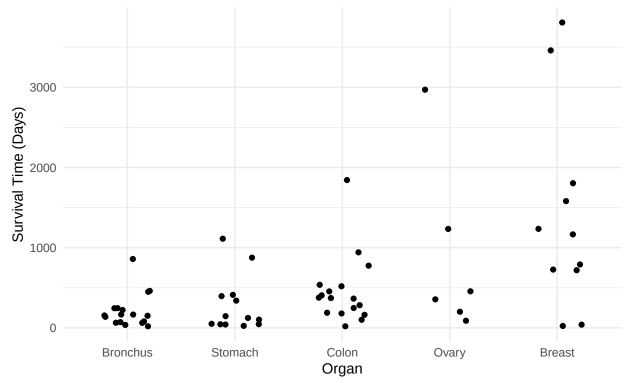
We will discuss four solutions to heteroscedasticity in linear and nonlinear regression: variance-stabilizing transformations, weighted least squares, robust standard errors, and models that do not assume homoscedasticity.

#### Variance-Stabilizing Transformations

The idea is to use  $Y_i^* = g(Y_i)$  instead of  $Y_i$  as the response variable, where g is a variance-stabilizing transformation.

Example: Consider again the cancer survival time data.

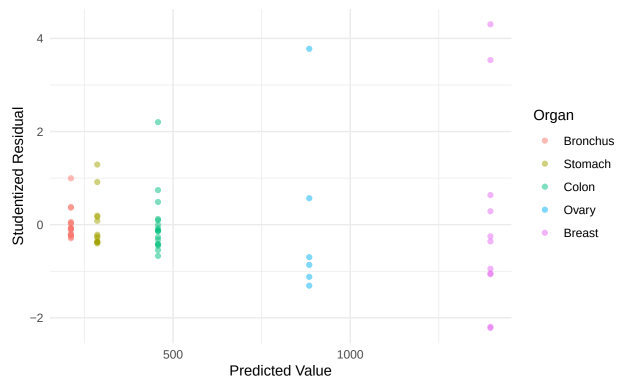
```
library(Stat2Data)
data(CancerSurvival)
CancerSurvival$Organ <- with(CancerSurvival, reorder(Organ, Survival, mean))
p <- ggplot(CancerSurvival, aes(x = Organ, y = Survival)) +
    geom_jitter(height = 0, width = 0.25) +
    labs(y = "Survival Time (Days)") + theme_minimal()
plot(p)</pre>
```



```
m <- lm(Survival ~ Organ, data = CancerSurvival)
CancerSurvival$yhat <- predict(m)</pre>
```

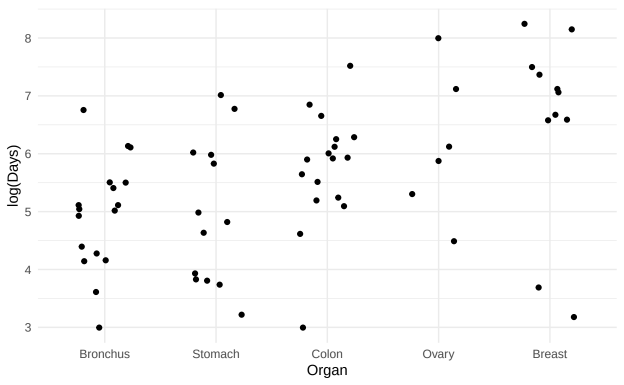
```
CancerSurvival$rest <- rstudent(m)

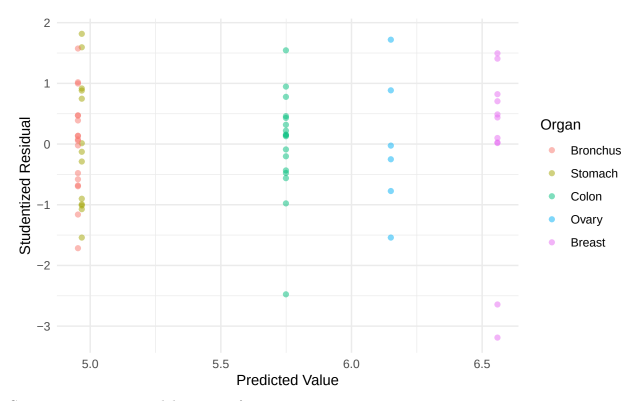
p <- ggplot(CancerSurvival, aes(x = yhat, y = rest, color = Organ)) +
    geom_point(alpha = 0.5) + theme_minimal() +
    labs(x = "Predicted Value", y = "Studentized Residual")
plot(p)</pre>
```



A model for  $\log$  time might exhibit something closer to homoscedasticity.

```
p <- ggplot(CancerSurvival, aes(x = Organ, y = log(Survival))) +
    geom_jitter(height = 0, width = 0.25) +
    labs(y = "log(Days)") + theme_minimal()
plot(p)</pre>
```





Comments on variance-stabilizing transformations.

- 1. Depending on the situation, other transformations may exhibit variance-stabilizing properties. Some common transformations are  $\sqrt{Y_i}$ ,  $\log(Y_i)$ ,  $1/\sqrt{Y_i}$  and  $1/Y_i$  for right-skewed response variables, and  $n_i \sin^{-1} \sqrt{Y_i}$  when  $Y_i$  is a proportion with a denominator of  $n_i$ .
- 2. A limitation of variance stabilizing transformations is that it is often difficult (and undesirable) to to *interpret* the model in terms of the transformed response variable (although there are exceptions as we will later see with the log transformation in the context of accelerated failure time models for survival data).
- 3. It is important to note that for any *nonlinear* transformation that  $E[g(Y)] \neq g[E(Y)]$  (i.e., the expected transformed response does not necessarily equal the transformed expected response). For example, the expected log of survival time does not equal the log of the expected survival time. So we cannot obtain inferences for the expected response by applying the inverse function. For example, while we have that  $\exp[\log(Y)] = Y$ , this **does not** imply that  $\exp\{E[\log(Y)]\} = E(Y)$ .

### Weighted Least Squares

A weighted least squares (WLS) estimator of the regression model parameters minimizes

$$\sum_{i=1}^{n} w_i (y_i - \hat{y}_i)^2,$$

were  $w_i > 0$  is the weight for the i-th observation. So-called ordinary least squares (OLS) or unweighted least squares is a special case where all  $w_i = 1$ .

To account for heteroscedasticity, the weights should be inversely proportional to the variance of the response so that

$$w_i \propto \frac{1}{\operatorname{Var}(Y_i)}.$$

Estimation is *efficient* meaning that the *true* standard errors (which are not necessarily the *reported* standard errors shown by software since these are estimates and may be biased without using weights as defined above)

are as small as they can be when using weighted least squares.

**Example**: Consider the following data.

```
turkeys <- data.frame(
  weight = c(674, 764, 795, 796, 826, 782, 834, 836, 830),
  pens = c(10, 5, 2, 2, 5, 5, 2, 2, 5),
  dosea = c(c(0, 0.12, 0.22, 0.32, 0.44), rep(0, 4)),
  doseb = c(rep(0, 5), c(0.12, 0.22, 0.32, 0.44))
)
turkeys</pre>
```

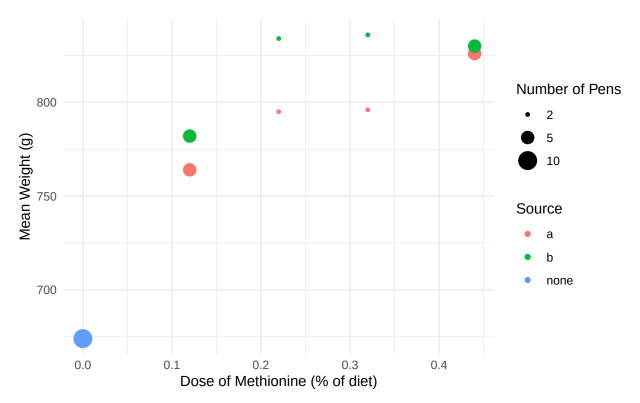
```
weight pens dosea doseb
        10 0.00 0.00
    674
1
2
          5 0.12 0.00
    764
3
    795
          2 0.22 0.00
4
    796
        2 0.32 0.00
5
    826
          5 0.44 0.00
6
    782
          5 0.00 0.12
7
    834
          2 0.00 0.22
8
    836
           2 0.00 0.32
9
          5 0.00 0.44
    830
```

For plotting and modeling convenience we will rearrange the data a bit.

```
library(dplyr)
turkeys <- turkeys %>%
  mutate(dose = dosea + doseb) %>%
  mutate(source = case_when(
    dose == 0 ~ "none",
    dosea > 0 ~ "a",
    doseb > 0 ~ "b")
) %>%
  select(-dosea, -doseb)
turkeys
```

```
weight pens dose source
           10 0.00
     674
                     none
1
2
     764
           5 0.12
3
    795
            2 0.22
                        а
     796
            2 0.32
4
                        a
5
     826
            5 0.44
6
    782
            5 0.12
7
            2 0.22
     834
                        b
8
     836
            2 0.32
                        b
    830
            5 0.44
```

```
library(ggplot2)
p <- ggplot(turkeys, aes(x = dose, y = weight, color = source)) +
    theme_minimal() + geom_point(aes(size = pens)) +
    scale_size(breaks = c(2, 5, 10)) +
    labs(x = "Dose of Methionine (% of diet)", y = "Mean Weight (g)",
        color = "Source", size = "Number of Pens")
plot(p)</pre>
```

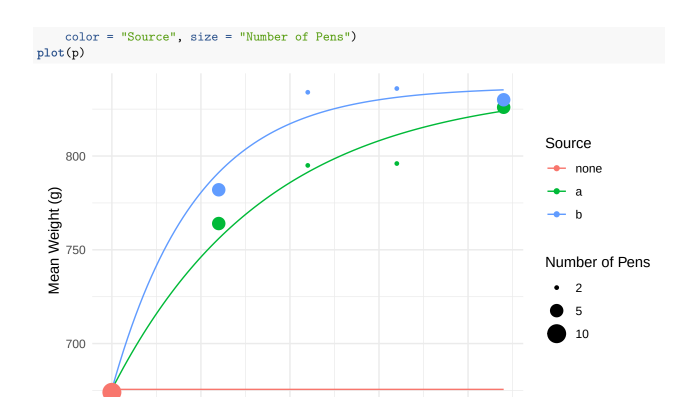


Suppose we want to estimate the following model.

$$E(W_i) = \begin{cases} \gamma, & \text{if no methionine was given,} \\ \alpha + (\gamma - \alpha) 2^{-d_i/\lambda_a}, & \text{if methionine was given from source } a, \\ \alpha + (\gamma - \alpha) 2^{-d_i/\lambda_b}, & \text{if methionine was given from source } b. \end{cases}$$

Note that this is like the Von Bertalanffy growth model. Here  $\alpha$  is an asymptote for expected weight we are approach as dose increases,  $\gamma$  is the expected response for a zero dose, and  $\lambda_a$  and  $\lambda_b$  are "half-life" parameters. We can estimate this model as follows using *ordinary* (i.e., unweighted) least squares as follows.

```
m.ols <- nls(weight ~ case_when(</pre>
  source == "none" ~ gamma,
  source == "a" ~ alpha + (gamma - alpha) * 2^(-dose/lambdaa),
  source == "b" ~ alpha + (gamma - alpha) * 2^(-dose/lambdab)),
  start = list(alpha = 825, gamma = 675, lambdaa = 0.1, lambdab = 0.1),
  data = turkeys)
summary(m.ols)$coefficients
         Estimate Std. Error t value Pr(>|t|)
                              94.278 2.545e-09
alpha
        836.87362
                      8.87663
gamma
        675.56291
                     10.92399
                               61.842 2.092e-08
          0.12049
                      0.02250
                                5.355 3.051e-03
lambdaa
lambdab
          0.06589
                      0.01601
                                4.115 9.222e-03
d \leftarrow expand.grid(source = c("none", "a", "b"), dose = seq(0, 0.44, length = 100))
d$yhat <- predict(m.ols, newdata = d)</pre>
p <- ggplot(turkeys, aes(x = dose, y = weight, color = source)) +</pre>
  geom_line(aes(y = yhat), data = d) +
  theme_minimal() + geom_point(aes(size = pens)) +
  scale_size(breaks = c(2, 5, 10)) +
  labs(x = "Dose of Methionine (% of diet)", y = "Mean Weight (g)",
```



The response variable is an mean of several observations so that

0.1

0.0

$$Y_i = \frac{Z_{i1} + Z_{i2} + \dots + Z_{in_i}}{n_i}$$

Dose of Methionine (% of diet)

0.4

where  $Z_{ij}$  is the length of the j-th pen that goes into the i-th average, and a total of  $n_i$  pens go into the i-th average. If  $Var(Z_{ij}) = \sigma^2$  then  $Var(Y_i) = \sigma^2/n_i$ . Thus the weights should be

$$w_i \propto \frac{1}{\sigma^2/n_i} = \frac{n_i}{\sigma^2}.$$

Since  $1/\sigma^2$  is a constant for all observations, we can define the weights as  $w_i = n_i$ . The weights can be specified in lm and nls (and other functions for regression) using the weights argument.

```
m.wls <- nls(weight ~ case_when(
   source == "none" ~ gamma,
   source == "a" ~ alpha + (gamma - alpha) * 2^(-dose/lambdaa),
   source == "b" ~ alpha + (gamma - alpha) * 2^(-dose/lambdab)),
   start = list(alpha = 825, gamma = 675, lambdaa = 0.1, lambdab = 0.1),
   data = turkeys, weights = pens)
summary(m.wls)$coefficients</pre>
```

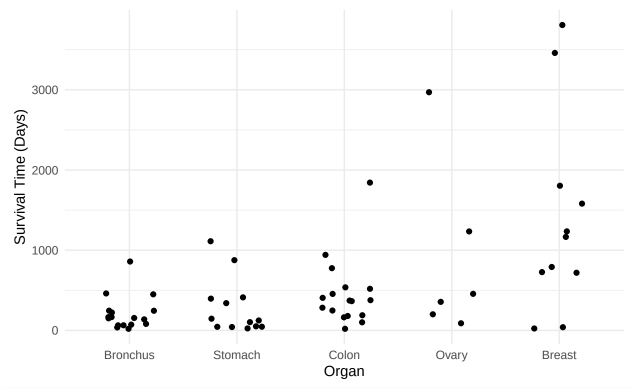
```
Estimate Std. Error t value Pr(>|t|)
alpha 834.78331 6.7761 123.195 6.684e-10
gamma 674.30393 5.5420 121.671 7.113e-10
lambdaa 0.10997 0.0163 6.746 1.086e-03
lambdab 0.06857 0.0117 5.860 2.051e-03
```

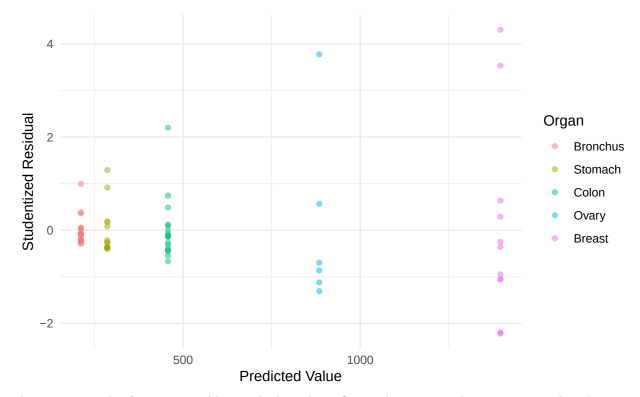
### summary(m.ols)\$coefficients

```
Estimate Std. Error t value Pr(>|t|)
alpha 836.87362 8.87663 94.278 2.545e-09
gamma 675.56291 10.92399 61.842 2.092e-08
lambdaa 0.12049 0.02250 5.355 3.051e-03
lambdab 0.06589 0.01601 4.115 9.222e-03
```

**Example:** Consider again the cancer survival time data.

```
p <- ggplot(CancerSurvival, aes(x = Organ, y = Survival)) +
    geom_jitter(height = 0, width = 0.25) +
    labs(y = "Survival Time (Days)") + theme_minimal()
plot(p)</pre>
```





There are a couple of ways we could go with these data. One is that since we have a categorical explanatory variable with multiple observations per category, we could *estimate* the variance of  $Y_i$  of each organ, and then set the weights to the reciprocals of these estimated variances.

```
library(dplyr)
CancerSurvival %>% group_by(Organ) %>%
  summarize(variance = var(Survival), weight = 1/var(Survival))
# A tibble: 5 x 3
  Organ
           variance
                         weight
  <fct>
              <dbl>
                           <dbl>
             44041. 0.0000227
1 Bronchus
            119930. 0.00000834
2 Stomach
3 Colon
            182473. 0.00000548
           1206875. 0.000000829
4 Ovary
5 Breast
           1535038. 0.000000651
```

We can use the following to compute weights and add them to the data frame.

```
CancerSurvival <- CancerSurvival %>%
  group_by(Organ) %>% mutate(w = 1/var(Survival))
head(CancerSurvival)
```

```
# A tibble: 6 x 3
# Groups:
            Organ [1]
  Survival Organ
     <int> <fct>
                         <dbl>
       124 Stomach 0.00000834
1
2
        42 Stomach 0.00000834
3
        25 Stomach 0.00000834
4
        45 Stomach 0.00000834
5
       412 Stomach 0.00000834
```

#### 6 51 Stomach 0.00000834

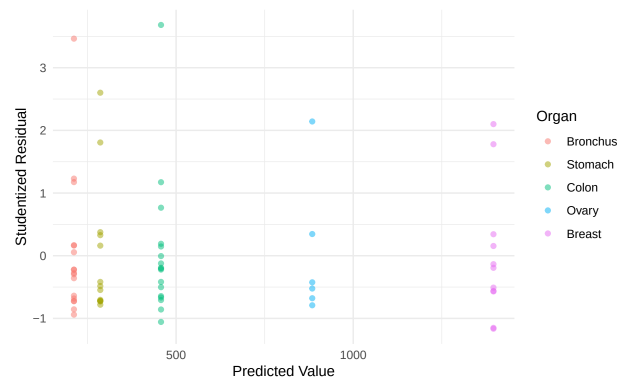
Now let's estimate the model using weighted least squares with these weights and inspect the residuals.

```
m.wls <- lm(Survival ~ Organ, weights = w, data = CancerSurvival)

CancerSurvival$yhat <- predict(m.wls)

CancerSurvival$rest <- rstudent(m.wls)

p <- ggplot(CancerSurvival, aes(x = yhat, y = rest, color = Organ)) +
    geom_point(alpha = 0.5) + theme_minimal() +
    labs(x = "Predicted Value", y = "Studentized Residual")
plot(p)</pre>
```



Note how this affects our inferences.

```
cbind(summary(m.ols)$coefficients, confint(m.ols))

Estimate Std. Error t value Pr(>|t|) 2.5 % 97.5 %
```

```
(Intercept)
              211.59
                          162.4 1.3030 0.1976373 -113.34 536.5
OrganStomach
               74.41
                          246.7 0.3017 0.7639784 -419.20 568.0
OrganColon
              245.82
                          229.6 1.0704 0.2887820 -213.70 705.3
              672.75
                          317.9 2.1160 0.0385749
OrganOvary
                                                    36.56 1308.9
                          259.1 4.5713 0.0000253 665.91 1702.7
OrganBreast
             1184.32
```

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

	Estimate	Std.	Error	t value	Pr(> t )	2.5 %	97.5 %
(Intercept)	211.59		50.9	4.1571	0.0001057	109.74	313.4
OrganStomach	74.41		108.7	0.6846	0.4963078	-143.10	291.9
OrganColon	245.82		115.4	2.1296	0.0373858	14.85	476.8
OrganOvary	672.75		451.4	1.4904	0.1414343	-230.45	1575.9
OrganBreast	1184.32		377.0	3.1413	0.0026291	429.92	1938.7

```
organs <- unique(CancerSurvival$Organ)</pre>
trtools::contrast(m.ols, a = list(Organ = organs), cnames = organs)
                         lower upper tvalue df
        estimate
Stomach
           286.0 185.7 -85.57 657.6 1.540 59 1.289e-01
Bronchus
           211.6 162.4 -113.34 536.5 1.303 59 1.976e-01
Colon
           457.4 162.4 132.48 782.3 2.817 59 6.587e-03
           884.3 273.3 337.39 1431.3 3.235 59 1.993e-03
Ovary
Breast
          1395.9 201.9 991.96 1799.9 6.915 59 3.770e-09
trtools::contrast(m.wls, a = list(Organ = organs), cnames = organs)
        estimate
                     se lower upper tvalue df
                                                   pvalue
           286.0 96.05 93.81 478.2 2.978 59 0.0042091
Stomach
Bronchus
           211.6 50.90 109.74 313.4 4.157 59 0.0001057
           457.4 103.60 250.10 664.7 4.415 59 0.0000437
Colon
           884.3 448.49 -13.10 1781.8 1.972 59 0.0533281
Ovary
          1395.9 373.56 648.41 2143.4 3.737 59 0.0004228
Breast
trtools::contrast(m.ols,
  a = list(Organ = "Breast"),
  b = list(Organ = c("Bronchus", "Stomach", "Colon", "Ovary")),
  cnames = c("Breast vs Bronchus", "Breast vs Stomach",
     "Breast vs Colon", "Breast vs Ovary"))
                              se lower upper tvalue df
                  estimate
                   1184.3 259.1 665.9 1703 4.571 59 0.0000253
Breast vs Bronchus
Breast vs Stomach
                  1109.9 274.3 561.1 1659 4.046 59 0.0001533
Breast vs Colon
                    938.5 259.1 420.1 1457 3.622 59 0.0006083
                     511.6 339.8 -168.4 1192 1.506 59 0.1375263
Breast vs Ovary
trtools::contrast(m.wls,
  a = list(Organ = "Breast"),
  b = list(Organ = c("Bronchus", "Stomach", "Colon", "Ovary")),
  cnames = c("Breast vs Bronchus", "Breast vs Stomach",
     "Breast vs Colon", "Breast vs Ovary"))
                              se lower upper tvalue df
                  estimate
Breast vs Bronchus 1184.3 377.0 429.9 1939 3.1413 59 0.002629
                    1109.9 385.7 338.1 1882 2.8776 59 0.005572
Breast vs Stomach
                    938.5 387.7 162.8 1714 2.4209 59 0.018577
Breast vs Colon
                     511.6 583.7 -656.4 1680 0.8765 59 0.384340
Breast vs Ovary
Here's how you can do the comparison of one level with all others using the contrast function from the
emmeans package.
library(emmeans)
contrast(emmeans(m.wls, ~ Organ), "trt.vs.ctrl", ref = "Breast",
  reverse = TRUE, adjust = "none", infer = TRUE)
contrast
                  estimate SE df lower.CL upper.CL t.ratio p.value
Breast - Bronchus
                      1184 377 59
                                       430
                                               1939
                                                      3.141 0.0026
Breast - Stomach
                      1110 386 59
                                       338
                                               1882
                                                      2.878 0.0056
Breast - Colon
                       938 388 59
                                       163
                                               1714
                                                      2.421 0.0186
Breast - Ovary
                       512 584 59
                                      -656
                                               1680
                                                      0.876 0.3843
```

Confidence level used: 0.95

Another approach is to assume that the variance of the response variable is some function of its expected response, and thus the weights are a function of the expected response. With right-skewed response variables one common functional relationship is that

$$Var(Y_i) \propto E(Y_i)$$
,

or, more generally,

$$Var(Y_i) \propto E(Y_i)^p$$
,

where p is some power (usually  $p \ge 1$ ). So the weights would then be

$$w_i \propto \frac{1}{E(Y_i)^p}.$$

We do not know  $E(Y_i)$ , but  $\hat{y}_i$  is an estimate of  $E(Y_i)$ . But we need the weights to compute  $\hat{y}_i$ !

Two situations:

- 1. Estimates of the model parameters and thus  $\hat{y}_i$  do not depend on the weights.<sup>1</sup> Here we can compute the weights with an initial regression model without weights.
- 2. Estimates of the model parameters and thus  $\hat{y}_i$  do depend on the weights. An approach we can use here is iteratively weighted least squares.

It can be shown that  $\hat{y}_i$  does not depend on the weights for the model for the CancerSurvival model. We can use the estimates from ordinary least squares to obtain weights of  $w_i = 1/\hat{y}_i^p$ .

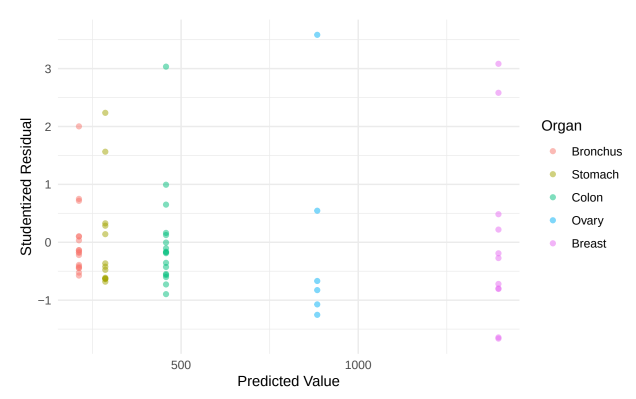
```
m.ols <- lm(Survival ~ Organ, data = CancerSurvival)

CancerSurvival$w <- 1/predict(m.ols)
m.wls <- lm(Survival ~ Organ, data = CancerSurvival, weights = w)

CancerSurvival$yhat <- predict(m.wls)
CancerSurvival$rest <- rstudent(m.wls)

p <- ggplot(CancerSurvival, aes(x = yhat, y = rest, color = Organ)) +
    geom_point(alpha = 0.5) + theme_minimal() +
    labs(x = "Predicted Value", y = "Studentized Residual")
plot(p)</pre>
```

<sup>&</sup>lt;sup>1</sup>In practice the most common situation where this will happen is with models with only a single categorical explanatory variable (or models with multiple categorical explanatory variables that include interactions).



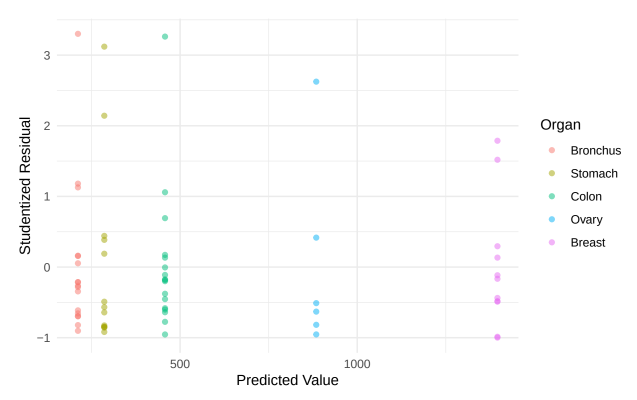
Maybe we could do better. Let's try p = 2 — i.e.,  $Var(Y_i) \propto E(Y_i)^2$ .

```
m.ols <- lm(Survival ~ Organ, data = CancerSurvival)

CancerSurvival$w <- 1/predict(m.ols)^2
m.wls <- lm(Survival ~ Organ, data = CancerSurvival, weights = w)

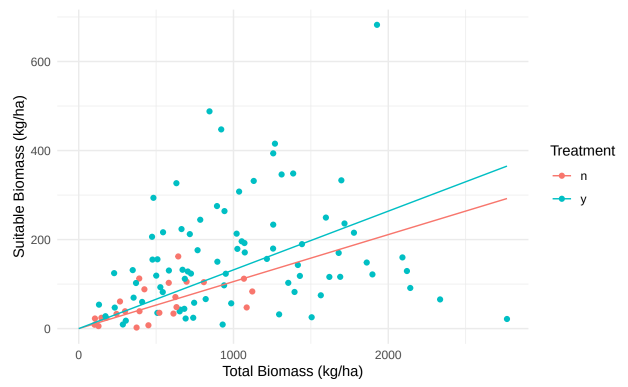
CancerSurvival$yhat <- predict(m.wls)
CancerSurvival$rest <- rstudent(m.wls)

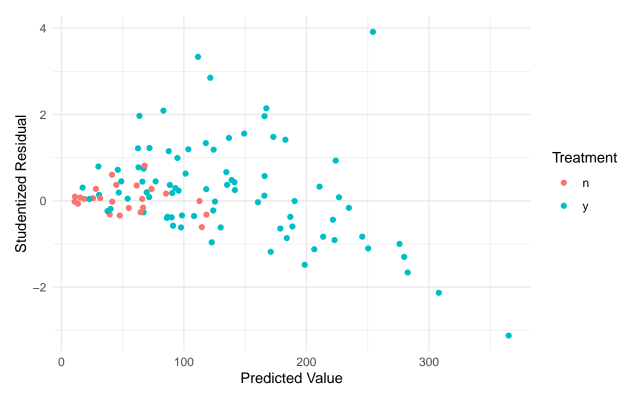
p <- ggplot(CancerSurvival, aes(x = yhat, y = rest, color = Organ)) +
    geom_point(alpha = 0.5) + theme_minimal() +
    labs(x = "Predicted Value", y = "Studentized Residual")
plot(p)</pre>
```



**Example**: Consider again following data from a study on the effects of fuel reduction on biomass.

```
library(trtools) # for biomass data
m <- lm(suitable ~ -1 + treatment:total, data = biomass)</pre>
summary(m)$coefficients
                  Estimate Std. Error t value Pr(>|t|)
treatmentn:total
                    0.1056
                              0.04183
                                         2.524 1.31e-02
                    0.1319
                              0.01121 11.773 7.61e-21
treatmenty:total
d \leftarrow expand.grid(treatment = c("n","y"), total = seq(0, 2767, length = 10))
d$yhat <- predict(m, newdata = d)</pre>
p <- ggplot(biomass, aes(x = total, y = suitable, color = treatment)) +</pre>
   geom_point() + geom_line(aes(y = yhat), data = d) + theme_minimal() +
   labs(x = "Total Biomass (kg/ha)",
      y = "Suitable Biomass (kg/ha)",
      color = "Treatment")
plot(p)
```





Here we might also assume that  $\operatorname{Var}(Y_i) \propto E(Y_i)^p$ , with weights of  $w_i = 1/\hat{y}_i$ . But here things are a bit more complicated for this model: the  $w_i$  depend on the  $\hat{y}_i$ , the  $\hat{y}_i$  depend on the  $w_i$ . In the model for the CancerSurvival data this was not an issue because there the estimates of the model parameters, and thus  $\hat{y}_i$ , did not depend on the weights so we could use ordinary least squares where all  $w_i = 1$  to get the  $\hat{y}_i$ . But that is not true for this model. But we can solve this problem using iteratively weighted least squares.

```
biomass$w <- 1 # initial weights are all equal to one
for (i in 1:5) {
 m.wls <- lm(suitable ~ -1 + treatment:total, weights = w, data = biomass)</pre>
 print(coef(m.wls)) # optional
 print(biomass$w)
                 # optional
 biomass$w <- 1 / predict(m.wls)</pre>
}
treatmentn:total treatmenty:total
        0.1056
                      0.1319
 treatmentn:total treatmenty:total
        0.1155
                      0.1578
 [1] 0.007064 0.032550 0.008232 0.011412 0.014898 0.074071 0.006703 0.005470
 [9] 0.007314 0.010754 0.021027 0.024910 0.035501 0.016278 0.008970 0.015132
 [17] 0.022310 0.033109 0.021716 0.010972 0.014911 0.015997 0.092343 0.011300
[25] 0.013910 0.008150 0.007978 0.011553 0.006235 0.005782 0.018173 0.025335
[33] 0.010453 0.014369 0.011716 0.009869 0.008049 0.005353 0.003933 0.024211
[41] 0.015701 0.005305 0.064582 0.011616 0.011111 0.005036 0.007079 0.005436
[49] 0.004746 0.009239 0.026575 0.015901 0.021376 0.008873 0.012010 0.024113
[57] 0.014990 0.008727 0.008070 0.014726 0.008485 0.020475 0.038766 0.005255
[65] 0.006031 0.013957 0.007197 0.006031 0.010244 0.010559 0.005978 0.009652
[73] 0.057874 0.018494 0.013587 0.007693 0.053023 0.004846 0.010163 0.007394
```

```
[81] 0.013009 0.007429 0.006036 0.004073 0.043850 0.003994 0.003538 0.004487
  [89] 0.090312 0.005598 0.003575 0.008461 0.003624 0.008445 0.004467 0.004414
  [97] 0.005854 0.002740 0.031633 0.011055 0.015161 0.004513 0.015473 0.004263
[105] 0.004681 0.003246
treatmentn:total treatmenty:total
                  0.1155
                                                 0.1578
   [1] 0.005904 0.027206 0.006880 0.009538 0.012453 0.067738 0.005602 0.004572
   [9] 0.006114 0.008988 0.019229 0.020820 0.032465 0.014886 0.007498 0.012647
  [17] 0.020403 0.027673 0.018151 0.009171 0.012463 0.013371 0.084448 0.009445
  [25] 0.011626 0.006812 0.006668 0.009656 0.005211 0.004833 0.016620 0.023169
  [33] 0.008737 0.012010 0.010714 0.008249 0.006727 0.004474 0.003288 0.022141
  [41] 0.013123 0.004434 0.059060 0.009709 0.009287 0.004209 0.005917 0.004544
  [49] 0.003967 0.007722 0.022212 0.013290 0.017867 0.008114 0.010038 0.022052
  [57] 0.013709 0.007981 0.006745 0.013467 0.007092 0.017114 0.035451 0.004392
  [65] 0.005041 0.011666 0.006016 0.005041 0.008562 0.008826 0.004997 0.008068
  [73] \quad 0.048373 \quad 0.015458 \quad 0.012425 \quad 0.006430 \quad 0.048490 \quad 0.004050 \quad 0.008494 \quad 0.006180
  [81] 0.010873 0.006210 0.005045 0.003405 0.036651 0.003339 0.002957 0.003750
  [89] 0.082590 0.004679 0.002988 0.007072 0.003029 0.007723 0.003734 0.003690
  [97] 0.004893 0.002290 0.028928 0.009240 0.013865 0.003772 0.014150 0.003563
[105] 0.003912 0.002713
treatmentn:total treatmenty:total
                                                 0.1578
   [1] 0.005904 0.027206 0.006880 0.009538 0.012453 0.067738 0.005602 0.004572
   [9] 0.006114 0.008988 0.019229 0.020820 0.032465 0.014886 0.007498 0.012647
  [17] 0.020403 0.027673 0.018151 0.009171 0.012463 0.013371 0.084448 0.009445
  [25] 0.011626 0.006812 0.006668 0.009656 0.005211 0.004833 0.016620 0.023169
  [33] \quad 0.008737 \quad 0.012010 \quad 0.010714 \quad 0.008249 \quad 0.006727 \quad 0.004474 \quad 0.003288 \quad 0.022141 \quad 0.008737 \quad 0.
  [41] 0.013123 0.004434 0.059060 0.009709 0.009287 0.004209 0.005917 0.004544
  [49] \ \ 0.003967 \ \ 0.007722 \ \ 0.022212 \ \ 0.013290 \ \ 0.017867 \ \ 0.008114 \ \ 0.010038 \ \ 0.022052
  [57] 0.013709 0.007981 0.006745 0.013467 0.007092 0.017114 0.035451 0.004392
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  [73] 0.048373 0.015458 0.012425 0.006430 0.048490 0.004050 0.008494 0.006180
  [81] 0.010873 0.006210 0.005045 0.003405 0.036651 0.003339 0.002957 0.003750
  [89] 0.082590 0.004679 0.002988 0.007072 0.003029 0.007723 0.003734 0.003690
  [97] 0.004893 0.002290 0.028928 0.009240 0.013865 0.003772 0.014150 0.003563
[105] 0.003912 0.002713
treatmentn:total treatmenty:total
                  0.1155
                                                 0.1578
    [1] 0.005904 0.027206 0.006880 0.009538 0.012453 0.067738 0.005602 0.004572
   [9] 0.006114 0.008988 0.019229 0.020820 0.032465 0.014886 0.007498 0.012647
  [17] 0.020403 0.027673 0.018151 0.009171 0.012463 0.013371 0.084448 0.009445
  [25] 0.011626 0.006812 0.006668 0.009656 0.005211 0.004833 0.016620 0.023169
  [33] 0.008737 0.012010 0.010714 0.008249 0.006727 0.004474 0.003288 0.022141
  [41] 0.013123 0.004434 0.059060 0.009709 0.009287 0.004209 0.005917 0.004544
  [49] 0.003967 0.007722 0.022212 0.013290 0.017867 0.008114 0.010038 0.022052
  [57] \ \ 0.013709 \ \ 0.007981 \ \ 0.006745 \ \ 0.013467 \ \ 0.007092 \ \ 0.017114 \ \ 0.035451 \ \ 0.004392
  [65] 0.005041 0.011666 0.006016 0.005041 0.008562 0.008826 0.004997 0.008068
  [73] 0.048373 0.015458 0.012425 0.006430 0.048490 0.004050 0.008494 0.006180
  [81] 0.010873 0.006210 0.005045 0.003405 0.036651 0.003339 0.002957 0.003750
  [89] 0.082590 0.004679 0.002988 0.007072 0.003029 0.007723 0.003734 0.003690
  [97] 0.004893 0.002290 0.028928 0.009240 0.013865 0.003772 0.014150 0.003563
[105] 0.003912 0.002713
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