Wednesday, Feb 23

Iteratively Weighted Least Squares

Iteratively weighted least squares can be used when we assume that the variance is proportional to a function of the mean so that

$$Var(Y_i) \propto h[E(Y_i)],$$

where h is some specified function, implying that our weights should be

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

Because $E(Y_i)$ is unknown we can use the estimate \hat{y}_i to obtain weights

$$w_i = \frac{1}{h(\hat{y}_i)}.$$

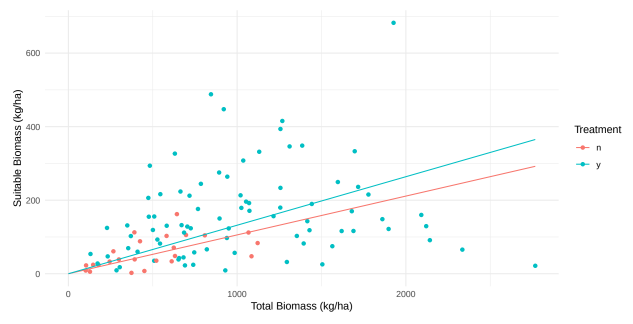
Because \hat{y}_i depends on the weights used in the weighted least squares algorithm, and w_i depends on \hat{y}_i , we can use the following algorithm known as iteratively weighted least squares.

- 1. Estimate the model using ordinary least squares where all $w_i = 1$.
- 2. Compute weights as $w_i = 1/h(\hat{y}_i)$.
- 3. Estimate the model using weighted least squares with the weights $w_i = 1/h(\hat{y}_i)$.

The second and third steps can be repeated until the estimates and thus the weights stop changing. Typically only a few iterations are necessary.

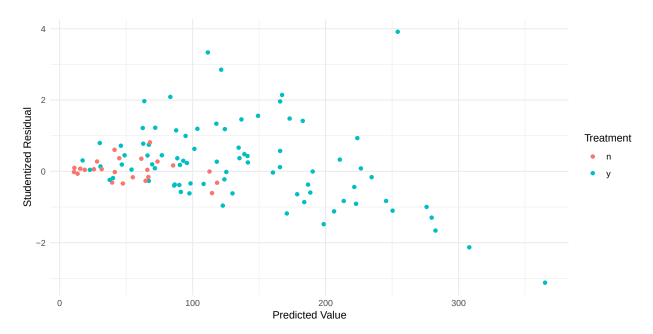
Example: Consider again following data from a study on the effects of fuel reduction on biomass.

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



```
biomass$yhat <- predict(m.ols)
biomass$rest <- rstudent(m.ols)

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



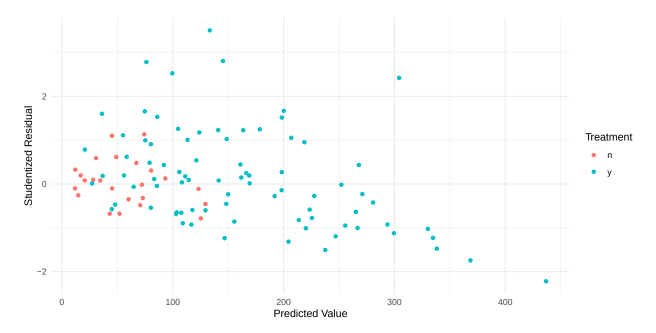
Assume that $Var(Y_i) \propto E(Y_i)$, which means the weights should be $w_i = 1/E(Y_i)$. We can program the iteratively weighted least squares algorithm as follows.

```
biomass$w <- 1 # initial weights are all equal to one
for (i in 1:5) {
  m.wls <- lm(suitable ~ -1 + treatment:total, weights = w, data = biomass)
  print(coef(m.wls)) # optional
  biomass$w <- 1 / predict(m.wls)
}</pre>
```

Now let's take a look at the residuals.

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

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



That may not be quite enough. Suppose we assume that $Var(Y_i) \propto E(Y_i)^p$ where p=2.

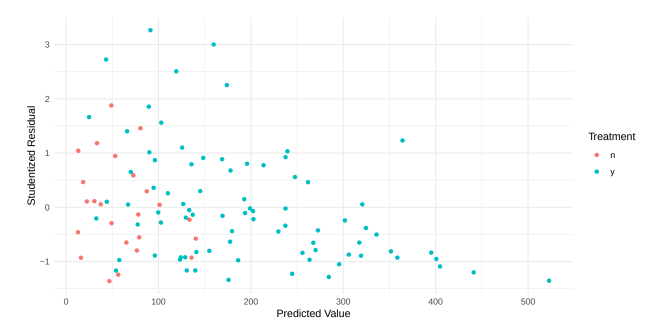
```
biomass$w <- 1 # initial weights are all equal to one
for (i in 1:5) {
  m.wls <- lm(suitable ~ -1 + treatment:total, weights = w, data = biomass)</pre>
```

```
biomass$w <- 1 / predict(m.wls)^2
}</pre>
```

Now let's take a look at the residuals.

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

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



Better. Maybe too much? We could try p = 1.5 or something like that. The residuals do get a little strange for higher predicted values, but we'll leave it here.

The model is $E(S_i) = \beta_1 n_i t_i + \beta_2 y_i t_i$, where n_i and y_i are indicator variables for if the *i*-th plot was treated or not by fuel reduction. We can also write the model as

$$E(S_i) = \begin{cases} \beta_1 t_i, & \text{if the i-th plot was not treated by fuel reduction,} \\ \beta_2 t_i, & \text{if the i-th plot was treated by fuel reduction.} \end{cases}$$

We can use $\beta_2 - \beta_1$ for inferences about the treatment effect.

(-1,1),0 0.06386 0.02359 0.01708 0.1106 2.707 104 0.007937

The contrast function from the **trtools** package can also do this. It can make inferences for a difference of differences.

```
contrast(m.wls,
   a = list(treatment = "y", total = 1),
   b = list(treatment = "y", total = 0),
   u = list(treatment = "n", total = 1),
   v = list(treatment = "n", total = 0))
```

```
estimate se lower upper tvalue df pvalue 0.06386 0.02359 0.01708 0.1106 2.707 104 0.007937
```

This estimates $E(Y_a) - E(Y_b) - [E(Y_u) - E(Y_v)]$. This can also be done using the emtrends function from the emmeans package.

```
library(emmeans)
emtrends(m.wls, ~treatment, var = "total") # estimate slopes
```

```
treatment total.trend SE df lower.CL upper.CL n 0.125 0.0183 104 0.0888 0.161 y 0.189 0.0149 104 0.1593 0.219
```

Confidence level used: 0.95

```
pairs(emtrends(m.wls, ~ treatment, var = "total")) # estimate difference between slopes
```

```
contrast estimate SE df t.ratio p.value n-y -0.0639 0.0236 104 -2.707 0.0079
```

Recall that both the **emmeans** and **trtools** packages have a **contrast** function. To avoid conflicts or having to use **trtools**::contrast to call te contrast function from the **trtools** package later, we can unload the **emmeans** package usin detach.

```
detach(package:emmeans)
```

Yet another approach to compare the slopes is to change the parameterization. Consider the following model.

```
m.wls <- lm(suitable ~ -1 + total + total:treatment, weights = w, data = biomass)
summary(m.wls)$coefficients</pre>
```

```
Estimate Std. Error t value Pr(>|t|) total 0.18892 0.01493 12.656 8.836e-23 total:treatmentn -0.06386 0.02359 -2.707 7.937e-03
```

From summary we can see that this model can be written as

$$E(S_i) = \beta_1 t_i + \beta_2 t_i n_i,$$

where n_i is an indicator variable where $n_i = 1$ if the treatment was not applied to the *i*-th plot, add $n_i = 0$ otherwise, so we can also write the model as

```
E(S_i) = \begin{cases} (\beta_1 + \beta_2)t_i, & \text{if the } i\text{-th plot was not treated by fuel reduction,} \\ \beta_1 t_i, & \text{if the } i\text{-th plot was treated by fuel reduction.} \end{cases}
```

Note that the meaning of β_1 and β_2 have changed here. The slopes of the lines with and without treatment are β_1 and $\beta_1 + \beta_2$, respectively, and the difference between the slopes is $\beta_1 - (\beta_1 + \beta_2) = -\beta_2$. So inferences for β_2 are for the difference in the slopes (after we reverse the sign). Although not necessary, we can change the reference category to avoid having to reverse the sign.

```
biomass$treatment <- relevel(biomass$treatment, ref = "y")
m.wls <- lm(suitable ~ -1 + total + total:treatment, weights = w, data = biomass)
summary(m.wls)$coefficients</pre>
```

```
Estimate Std. Error t value Pr(>|t|) total 0.12506 0.01827 6.847 5.428e-10 total:treatmenty 0.06386 0.02359 2.707 7.937e-03
```

Now the model can be written as

$$E(S_i) = \beta_1 t_i + \beta_2 t_i n_i,$$

or

$$E(S_i) = \begin{cases} \beta_1 t_i, & \text{if the } i\text{-th plot was not treated by fuel reduction,} \\ (\beta_1 + \beta_2) t_i, & \text{if the } i\text{-th plot was treated by fuel reduction.} \end{cases}$$

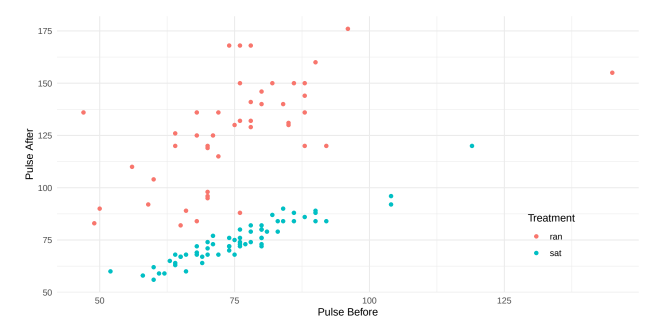
Note: For some reason the reference category (y) is getting an indicator variable here, where normally it does not. I am not sure if this is a bug or intentional, but it appears to be due to the somewhat unusual parameterization I am using.

Parametric Models for Heteroscedasticity

Example: Consider the following data where variability appears to vary by treatment.

```
library(trtools) # for pulse data

p <- ggplot(pulse, aes(x = pulse1, y = pulse2, color = treatment)) +
   geom_point() + theme_minimal() +
   labs(x = "Pulse Before", y = "Pulse After", color = "Treatment") +
   theme(legend.position = c(0.85,0.2))
plot(p)</pre>
```



There is one case with missing values on pulse1 and pulse2.

```
subset(pulse, !complete.cases(pulse)) # show observations with missing data
```

height weight age gender smokes alcohol exercise treatment pulse1 pulse2 year 76 173 64 20 female no yes moderate sat NA NA 97

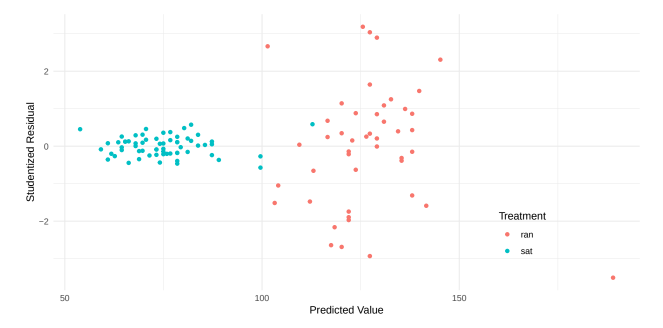
This will cause problems so we are going to remove it.

pulse <- subset(pulse, complete.cases(pulse)) # overwrite pulse with only complete cases</pre>

Let's consider a simple linear model.

```
m <- lm(pulse2 ~ treatment + pulse1 + treatment:pulse1, data = pulse)
summary(m)$coefficients</pre>
```

```
t value Pr(>|t|)
                      Estimate Std. Error
(Intercept)
                      59.41757
                                   10.4467
                                            5.68767 1.171e-07
treatmentsat
                     -51.25896
                                   15.7451 -3.25554 1.524e-03
                       0.89363
                                    0.1357 6.58544 1.841e-09
pulse1
treatmentsat:pulse1
                      -0.01437
                                    0.2049 -0.07011 9.442e-01
pulse$yhat <- predict(m)</pre>
pulse$rest <- rstudent(m)</pre>
p \leftarrow ggplot(pulse, aes(x = yhat, y = rest, color = treatment)) +
  geom_point() + theme_minimal() +
  labs(x = "Predicted Value", y = "Studentized Residual",
    color = "Treatment") +
  theme(legend.position = c(0.8,0.2))
plot(p)
```



Consider that the model assumed by 1m is

$$E(Y_i) = \beta_0 + \beta_1 t_i + \beta_2 x_i + \beta_3 t_i x_i, \tag{1}$$

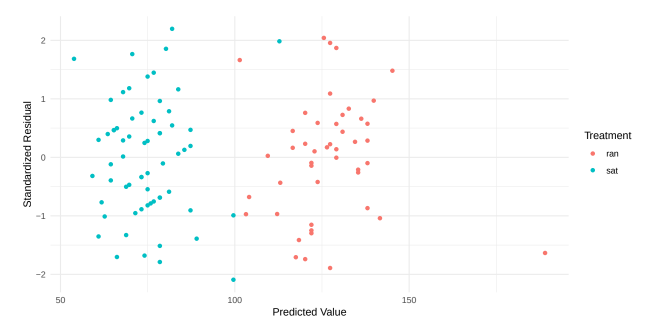
$$Var(Y_i) = \sigma^2, \tag{2}$$

where Y_i is the second pulse measurement, t_i is an indicator variable for the treatment (i.e., $t_i = 1$ if the *i*-th observation was from the sitting treatment condition, and $t_i = 0$ otherwise), and x_i is the first pulse measurement. Maybe it would make sense to have something like

$$\operatorname{Var}(Y_i) = \begin{cases} \sigma_s^2, & \text{if the i-th observation is from the sitting treatment,} \\ \sigma_r^2, & \text{if the i-th observation is from the running treatment.} \end{cases}$$

We can estimate such a model using the gls function from the nlme package.

```
library(nlme) # should come with R
m <- gls(pulse2 ~ treatment + pulse1 + treatment:pulse1, data = pulse,
 method = "ML", weights = varIdent(form = ~ 1|treatment))
summary(m)
Generalized least squares fit by maximum likelihood
  Model: pulse2 ~ treatment + pulse1 + treatment:pulse1
  Data: pulse
    AIC BIC logLik
  763.1 779.3 -375.6
Variance function:
 Structure: Different standard deviations per stratum
 Formula: ~1 | treatment
Parameter estimates:
  sat
      ran
1.000 5.723
Coefficients:
                    Value Std.Error t-value p-value
                    59.42 15.755 3.771 0.0003
(Intercept)
                             16.058 -3.192 0.0019
treatmentsat
                    -51.26
pulse1
                     0.89 0.205 4.367 0.0000
                             0.209 -0.069 0.9452
treatmentsat:pulse1 -0.01
 Correlation:
                    (Intr) trtmnt pulse1
treatmentsat
                    -0.981
pulse1
                    -0.980 0.962
treatmentsat:pulse1 0.962 -0.980 -0.981
Standardized residuals:
            Q1
                   Med
                             QЗ
-2.0920 -0.7688 0.1026 0.5886 2.1968
Residual standard error: 3.634
Degrees of freedom: 109 total; 105 residual
Note the different syntax for extracting standardized residuals.
pulse$yhat <- predict(m)</pre>
pulse$resz <- residuals(m, type = "p") # note different syntax</pre>
p \leftarrow ggplot(pulse, aes(x = yhat, y = resz, color = treatment)) +
  geom_point() + theme_minimal() +
  labs(x = "Predicted Value", y = "Standardized Residual",
    color = "Treatment")
plot(p)
```



Here is an example with the ${\tt CancerSurvival}$ data.

```
library(Stat2Data)
data(CancerSurvival)
m <- gls(Survival ~ Organ, data = CancerSurvival,
    method = "ML", weights = varIdent(form = ~ 1|Organ))
summary(m)</pre>
```

Generalized least squares fit by maximum likelihood

Model: Survival ~ Organ Data: CancerSurvival AIC BIC logLik 976.8 998.4 -478.4

Variance function:

Structure: Different standard deviations per stratum

Formula: ~1 | Organ Parameter estimates:

Stomach Bronchus Colon Ovary Breast 1.0000 0.6119 1.2455 3.0141 3.5504

Coefficients:

Value Std.Error t-value p-value (Intercept) 1395.9 371.0 3.763 0.0004 OrganBronchus -1184.3 374.5 -3.162 0.0025 OrganColon -938.5 385.5 -2.435 0.0179 OrganOvary -511.6 565.2 -0.905 0.3691 OrganStomach -1109.9 383.2 -2.896 0.0053

Correlation:

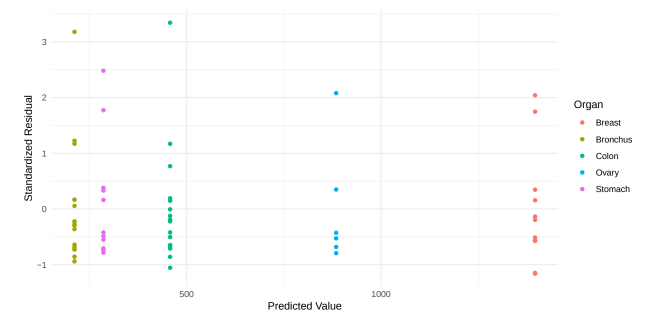
(Intr) OrgnBr OrgnCl OrgnOv

OrganBronchus -0.991

OrganColon -0.962 0.953

OrganOvary -0.656 0.650 0.632

```
OrganStomach -0.968 0.959 0.932 0.635
Standardized residuals:
    Min
             Q1
                              Q3
                                     Max
-1.1613 -0.6824 -0.2878
                         0.1748
                                 3.3435
Residual standard error: 332.7
Degrees of freedom: 64 total; 59 residual
CancerSurvival$yhat <- predict(m)</pre>
CancerSurvival$resz <- residuals(m, type = "p")</pre>
p <- ggplot(CancerSurvival, aes(x = yhat, y = resz, color = Organ)) +
  geom_point() + theme_minimal() +
  labs(x = "Predicted Value", y = "Standardized Residual", color = "Organ")
plot(p)
```



Comments about parametric models for heteroscedasticity.

Advantages: Potentially very effective if we can specify an accurate model for the variance.

Disadvantages: If we do not specify an accurate model for the variance, it may bias estimation of parameters concerning the expected response.

Heteroscedastic Consistent Standard Errors

The idea is to estimate the model parameters using ordinary least squares, but estimate the standard errors in such a way that we do not assume heteroscedasticity. This is sometimes called *heteroscedastic consistent standard errors*, robust standard errors, or sandwich estimators.

Example: Consider again the cancer survival data.

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

The **sandwich** package provides resources for using heteroscedastic-consistent standard errors. Technically, what is being estimated is the *covariance matrix* of the parameter estimators.

```
library(sandwich) # for vcovHC used below
vcov(m)
          # bad estimate if there is heteroscedasticity
              (Intercept) OrganBronchus OrganColon OrganOvary OrganStomach
(Intercept)
                    40752
                                  -40752
                                             -40752
                                                        -40752
                                                                      -40752
OrganBronchus
                   -40752
                                   67121
                                              40752
                                                         40752
                                                                       40752
OrganColon
                   -40752
                                   40752
                                              67121
                                                         40752
                                                                       40752
                                   40752
OrganOvary
                   -40752
                                              40752
                                                        115464
                                                                       40752
OrganStomach
                   -40752
                                   40752
                                              40752
                                                         40752
                                                                       75235
vcovHC(m) # better estimate if there is heteroscedasticity
              (Intercept) OrganBronchus OrganColon OrganOvary OrganStomach
                   153504
                                -153504
                                            -153504
                                                       -153504
                                                                     -153504
(Intercept)
OrganBronchus
                  -153504
                                 156256
                                             153504
                                                        153504
                                                                      153504
OrganColon
                  -153504
                                                        153504
                                                                      153504
                                  153504
                                             164908
OrganOvary
                  -153504
                                  153504
                                             153504
                                                        394879
                                                                      153504
                                  153504
OrganStomach
                  -153504
                                             153504
                                                        153504
                                                                      163498
The square root of the diagonal elements are the standard errors.
                      # bad estimates of the standard errors
sqrt(diag(vcov(m)))
  (Intercept) OrganBronchus
                                OrganColon
                                              OrganOvary OrganStomach
        201.9
                      259.1
                                     259.1
                                                   339.8
                                                                 274.3
sqrt(diag(vcovHC(m))) # better estimates of the standard errors
  (Intercept) OrganBronchus
                                OrganColon
                                              OrganOvary
                                                          OrganStomach
        391.8
                      395.3
                                     406.1
                                                   628.4
But the usual way to interface with the functions in the sandwich package is through other functions.
              Estimate Std. Error t value Pr(>|t|) 2.5 % 97.5 %
(Intercept)
                1395.9
                            201.9
                                    6.915 3.770e-09
                                                       992 1799.9
OrganBronchus
               -1184.3
                            259.1 -4.571 2.530e-05 -1703 -665.9
OrganColon
                -938.5
                            259.1 -3.622 6.083e-04 -1457 -420.1
OrganOvary
                -511.6
                            339.8 -1.506 1.375e-01 -1192 168.4
OrganStomach
               -1109.9
                            274.3 -4.046 1.533e-04 -1659 -561.1
confint(m) # bad confidence intervals due to bad standard error estimates
              2.5 % 97.5 %
(Intercept)
                992 1799.9
OrganBronchus -1703 -665.9
OrganColon
              -1457 -420.1
OrganOvary
              -1192 168.4
OrganStomach -1659 -561.1
library(lmtest) # for coeftest and coefci used below
coeftest(m, vcov = vcovHC) # better standard error estimates
t test of coefficients:
              Estimate Std. Error t value Pr(>|t|)
(Intercept)
                  1396
                              392
                                      3.56 0.00073 ***
                              395
OrganBronchus
                 -1184
                                     -3.00 0.00400 **
                              406
OrganColon
                  -938
                                     -2.31 0.02434 *
OrganOvary
                  -512
                              628
                                     -0.81 0.41886
```

```
OrganStomach
                 -1110
                              404
                                    -2.74 0.00801 **
Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
coefci(m, vcov = vcovHC)
                           # better confidence intervals
                2.5 % 97.5 %
(Intercept)
                611.9 2179.9
OrganBronchus -1975.3 -393.3
OrganColon
              -1751.1 -125.9
              -1769.0 745.8
OrganOvary
OrganStomach -1919.0 -300.8
Both lincon and contrast will accept a fcov argument to provide a function to estimate standard errors.
lincon(m, fcov = vcovHC)
              estimate
                          se
                               lower upper tvalue df
                                                          pvalue
(Intercept)
                1395.9 391.8
                               611.9 2179.9 3.5628 59 0.0007337
OrganBronchus -1184.3 395.3 -1975.3 -393.3 -2.9961 59 0.0039950
OrganColon
                -938.5 406.1 -1751.1 -125.9 -2.3111 59 0.0243421
                -511.6 628.4 -1769.0 745.8 -0.8141 59 0.4188611
OrganOvary
OrganStomach
              -1109.9 404.3 -1919.0 -300.8 -2.7449 59 0.0080080
organs <- sort(unique(CancerSurvival$Organ)) # sorted organ names</pre>
contrast(m, a = list(Organ = organs),
 cnames = organs, fcov = vcovHC)
                                                    pvalue
         estimate
                      se lower upper tvalue df
Breast
           1395.9 391.80 611.93 2179.9 3.563 59 7.337e-04
Bronchus
           211.6 52.46 106.61 316.6 4.033 59 1.604e-04
            457.4 106.79 243.72 671.1 4.283 59 6.884e-05
Colon
            884.3 491.30 -98.75 1867.4 1.800 59 7.698e-02
Ovary
           286.0 99.97 85.96 486.0 2.861 59 5.836e-03
Stomach
lincon(m, a = c(1,0,0,0,1), fcov = vcovHC)
                          se lower upper tvalue df
              estimate
                                                     pvalue
(1,0,0,0,1),0
                   286 99.97 85.96
                                     486 2.861 59 0.005836
You can use a similar approach with the emmeans function from the emmeans package, but there the
argument is vcov.
library(emmeans)
emmeans(m, ~Organ, vcov = vcovHC)
                    SE df lower.CL upper.CL
Organ
          emmean
 Breast
            1396 391.8 59
                             611.9
                                       2180
Bronchus
             212 52.5 59
                             106.6
                                        317
Colon
             457 106.8 59
                             243.7
                                        671
Ovary
             884 491.3 59
                             -98.8
                                       1867
Stomach
             286 100.0 59
                              86.0
                                        486
Confidence level used: 0.95
pairs(emmeans(m, ~Organ, vcov = vcovHC), adjust = "none", infer = TRUE)
                    estimate SE df lower.CL upper.CL t.ratio p.value
                                         393
                                               1975.3
                                                        2.996 0.0040
 Breast - Bronchus
                      1184.3 395 59
```

```
Breast - Colon
                      938.5 406 59
                                        126
                                              1751.1
                                                       2.311 0.0243
                     511.6 628 59
Breast - Ovary
                                              1769.0
                                                       0.814 0.4189
                                       -746
Breast - Stomach
                     1109.9 404 59
                                        301
                                              1919.0
                                                       2.745 0.0080
                                                     -2.066 0.0432
Bronchus - Colon
                     -245.8 119 59
                                       -484
                                                -7.7
Bronchus - Ovary
                     -672.7 494 59
                                      -1661
                                               315.9
                                                     -1.362 0.1785
                                                     -0.659 0.5124
Bronchus - Stomach
                      -74.4 113 59
                                       -300
                                               151.5
Colon - Ovary
                     -426.9 503 59
                                                     -0.849 0.3992
                                      -1433
                                               579.1
Colon - Stomach
                      171.4 146 59
                                       -121
                                               464.1
                                                       1.172 0.2460
Ovary - Stomach
                      598.3 501 59
                                       -405
                                              1601.6
                                                       1.193 0.2375
```

Confidence level used: 0.95

Use the function waldtest in place of anova when using heteroscedastic-consistent standard errors.

```
m.full <- lm(Survival ~ Organ, data = CancerSurvival)
m.null <- lm(Survival ~ 1, data = CancerSurvival)
waldtest(m.null, m.full, vcov = vcovHC)</pre>
```

Wald test

```
Model 1: Survival ~ 1
Model 2: Survival ~ Organ
  Res.Df Df F Pr(>F)
1 63
2 59 4 3.52 0.012 *
---
Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
```

Comments about heteroscedastic-consistent standard errors:

Advantages: Does not require us to specify a variance structure/function. We let the data inform the estimator.

Disadvantages: Highly dependent on the data to help produce better estimates of the standard errors, and tends to work well only if n is relatively large.

Note: There are a variety of variations of the "sandwich" estimator. Different estimators can be specified through the type argument to vcovHC so instead of writing vcov = vcovHC or fcov = vcovHC we write vcov = function(m) vcovHC(m, type = "HCO") or vcov = function(m) vcovHC(m, type = "HCO") if we wanted to use that particular type of estimator (sometimes called "White's estimator").