Friday, Feb 2

Marginal Means

A marginal mean is effectively an average of expected responses. The **emmeans** package is particularly useful for making inferences about marginal means.

```
library(trtools)
library(emmeans)
```

Attaching package: 'emmeans'

The following objects are masked from 'package:trtools':

```
contrast, neuralgia
```

Warning: The emmeans package contains a function called contrast which is not the same as the function of the same name in the trtools package, resulting in a namespace conflict if both packages are loaded. If you have both packages loaded in a given session, use trtools::contrast and emmeans::contrast to refer to a given function.

Example: Consider again the data from the platyfish study.

```
m <- lm(Percentage ~ Pair, data = Sleuth3::case0602)
summary(m)$coefficients</pre>
```

```
Estimate Std. Error t value Pr(>|t|)
(Intercept)
             56.406 3.864 14.5965 5.208e-24
PairPair2
              4.479
                         5.657 0.7919 4.308e-01
PairPair3
                         5.384 1.1187 2.667e-01
              6.023
PairPair4
             10.594
                         5.657 1.8727 6.485e-02
PairPair5
              7.805
                         6.441 1.2118 2.292e-01
              6.929
                         5.657 1.2250 2.243e-01
PairPair6
```

We see that there are indicator variables for male pairs 2-6. The model can be written as

```
E(Y_i) = \begin{cases} \beta_0, & \text{if the $i$-th observation was from the first male pair,} \\ \beta_0 + \beta_1, & \text{if the $i$-th observation was from the second male pair,} \\ \beta_0 + \beta_2, & \text{if the $i$-th observation was from the third male pair,} \\ \beta_0 + \beta_3, & \text{if the $i$-th observation was from the fourth male pair,} \\ \beta_0 + \beta_4, & \text{if the $i$-th observation was from the fifth male pair,} \\ \beta_0 + \beta_5, & \text{if the $i$-th observation was from the sixth male pair.} \end{cases}
```

We can use contrast to estimate the expected response for each pair.

```
contrast(m, a = list(Pair = paste("Pair", 1:6, sep = "")),
    cnames = paste("Pair", 1:6, sep = ""))
```

```
estimate se lower upper tvalue df pvalue
Pair1 56.41 3.864 48.71 64.10 14.60 78 5.208e-24
Pair2 60.89 4.131 52.66 69.11 14.74 78 2.990e-24
```

```
Pair3 62.43 3.749 54.97 69.89 16.65 78 2.114e-27
Pair4 67.00 4.131 58.78 75.22 16.22 78 1.052e-26
Pair5 64.21 5.152 53.95 74.47 12.46 78 3.039e-20
Pair6 63.34 4.131 55.11 71.56 15.33 78 3.006e-25
```

Note how I used a shortcut to specify the pairs.

```
paste("Pair", 1:6, sep = "")
```

[1] "Pair1" "Pair2" "Pair3" "Pair4" "Pair5" "Pair6"

This can also be done using the emmeans function from the package emmeans.

```
library(emmeans)
emmeans(m, ~ Pair)
```

```
Pair emmean
               SE df lower.CL upper.CL
                         48.7
        56.4 3.86 78
Pair1
                                   64.1
Pair2
        60.9 4.13 78
                         52.7
                                   69.1
       62.4 3.75 78
                         55.0
Pair3
                                   69.9
Pair4
        67.0 4.13 78
                         58.8
                                   75.2
        64.2 5.15 78
                         54.0
                                   74.5
Pair5
Pair6
        63.3 4.13 78
                         55.1
                                   71.6
```

Confidence level used: 0.95

Denote the six expected responses (one for each pair) as

$$\begin{split} \mu_1 &= \beta_0, \\ \mu_2 &= \beta_0 + \beta_1, \\ \mu_3 &= \beta_0 + \beta_2, \\ \mu_4 &= \beta_0 + \beta_3, \\ \mu_5 &= \beta_0 + \beta_4, \\ \mu_6 &= \beta_0 + \beta_5. \end{split}$$

One marginal mean would be the average expected response across the pairs. This could be written as

$$\mu = \frac{\mu_1 + \mu_2 + \mu_3 + \mu_4 + \mu_5 + \mu_6}{6} = \beta_0 + \frac{1}{6}\beta_1 + \frac{1}{6}\beta_2 + \frac{1}{6}\beta_3 + \frac{1}{6}\beta_4 + \frac{1}{6}\beta_5.$$

We can estimate this quantity with lincon.

```
lincon(m, a = c(1,1/6,1/6,1/6,1/6))
```

```
estimate se lower upper tvalue df pvalue (1,1/6,1/6,1/6,1/6,1/6),0 62.38 1.722 58.95 65.81 36.23 78 1.501e-50
```

We can also use emmeans.

```
emmeans(m, ~ 1)
```

```
1 emmean SE df lower.CL upper.CL overall 62.4 1.72 78 59 65.8
```

Results are averaged over the levels of: Pair Confidence level used: 0.95

Note that we can use the confidence interval to test the null hypothesis that $\mu = 50$. For a test statistic and p-value for this test we could write this as

$$\mu = 50 \Leftrightarrow \beta_0 + \frac{1}{6}\beta_1 + \frac{1}{6}\beta_2 + \frac{1}{6}\beta_3 + \frac{1}{6}\beta_4 + \frac{1}{6}\beta_5 = 50 \Leftrightarrow \beta_0 + \frac{1}{6}\beta_1 + \frac{1}{6}\beta_2 + \frac{1}{6}\beta_3 + \frac{1}{6}\beta_4 + \frac{1}{6}\beta_5 - 50 = 0.$$

Here is how we can do that with lincon.

```
lincon(m, a = c(1,1/6,1/6,1/6,1/6), b = -50)
```

```
estimate se lower upper tvalue df pvalue (1,1/6,1/6,1/6,1/6,1/6),-50 12.38 1.722 8.95 15.81 7.189 78 3.439e-10
```

Here is how we do it with emmeans.

```
emmeans(m, ~ 1, offset = -50, infer = TRUE)
```

```
1 emmean SE df lower.CL upper.CL t.ratio p.value overall 12.4 1.72 78 8.95 15.8 7.189 <.0001
```

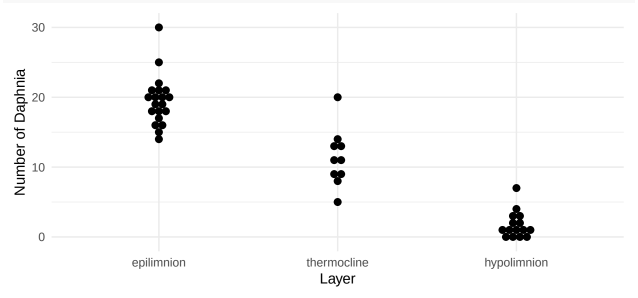
```
Results are averaged over the levels of: Pair Confidence level used: 0.95
```

By *not* listing an explanatory variable on the right-hand side of \sim , we are asking that emmeans average over that explanatory variable. Also note that the argument infer = TRUE makes the emmeans function provide both confidence intervals as well as tests.

Note: If we just want to know whether or not we would reject the null hypothesis that $\mu = 50$ we can also just look at the confidence interval for μ .

Example: Consider the following data from a survey of water fleas.

```
library(ggplot2)
p <- ggplot(daphniastrat, aes(x = layer, y = count)) +
  geom_dotplot(binaxis = "y", stackdir = "center") +
  labs(x = "Layer", y = "Number of Daphnia") + theme_minimal()
plot(p)</pre>
```



We might model these data using the following linear model.

```
m <- lm(count ~ layer, data = daphniastrat)
summary(m)$coefficients</pre>
```

```
Estimate Std. Error t value Pr(>|t|)
(Intercept) 19.50 0.7271 26.820 4.727e-28
layerthermocline -8.20 1.2593 -6.512 7.293e-08
layerhypolimnion -17.77 1.1106 -15.997 1.784e-19
```

So our model can be written as

$$E(Y_i) = \begin{cases} \beta_0, & \text{if the } i\text{-th observation is from the epilimnion layer,} \\ \beta_0 + \beta_1, & \text{if the } i\text{-th observation is from the thermocline layer,} \\ \beta_0 + \beta_2, & \text{if the } i\text{-th observation is from the hypolimnion layer.} \end{cases}$$

Let μ_e , μ_t , and μ_h denote the expected number of daphnia per liter for the epilimnion, thermocline, and hypolimnion layers, respectively (i.e., the density of daphnia in each layer). So

$$\mu_e = \beta_0, \ \mu_t = \beta_0 + \beta_1, \ \mu_h = \beta_0 + \beta_2.$$

It is known that the volumes of the epilimnion, thermocline, and hypolimnion layers are 100, 200, and 400 kL, respectively. The density for the entire lake is then

$$\mu = \frac{100}{700}\mu_e + \frac{200}{700}\mu_t + \frac{400}{700}\mu_h = \beta_0 + \frac{2}{7}\beta_1 + \frac{4}{7}\beta_2.$$

We can estimate this with lincon or emmeans using the weights option.

```
lincon(m, a = c(1, 2/7, 4/7))
```

```
estimate se lower upper tvalue df pvalue (1,2/7,4/7),0 7.005 0.572 5.85 8.159 12.25 42 1.907e-15 emmeans(m, ~ 1, weights = c(1/7, 2/7, 4/7))
```

```
1 emmean SE df lower.CL upper.CL overall 7 0.572 42 5.85 8.16
```

Results are averaged over the levels of: layer Confidence level used: 0.95

Note that when using emmeans it is important to put the weights in the correct order. We can verify the order using level (if the variable is a factor) or by using weights = "slow.levels".

```
levels(daphniastrat$layer)
```

```
[1] "epilimnion" "thermocline" "hypolimnion"
emmeans(m, ~ 1, weights = "show.levels")
```

emmeans are obtained by averaging over these factor combinations

```
layer
1 epilimnion
2 thermocline
3 hypolimnion
```

We can estimate the expected number of daphnia per liter for each layer.

```
emmeans(m, ~ layer)
```

```
layer emmean SE df lower.CL upper.CL epilimnion 19.50 0.727 42 18.033 20.97 thermocline 11.30 1.028 42 9.225 13.38 hypolimnion 1.73 0.840 42 0.039 3.43
```

Confidence level used: 0.95

```
trtools::contrast(m, a = list(layer = c("epilimnion","thermocline","hypolimnion")),
    cnames = c("epilimnion","thermocline","hypolimnion"))
```

```
estimate se lower upper tvalue df pvalue epilimnion 19.500 0.7271 18.03274 20.967 26.820 42 4.727e-28 thermocline 11.300 1.0282 9.22498 13.375 10.990 42 6.221e-14 hypolimnion 1.733 0.8395 0.03909 3.428 2.065 42 4.517e-02
```

We can also do inferences concerning the differences between pairs of layers.

```
pairs(emmeans(m, ~ layer), adjust = "none")
 contrast
                           estimate
                                       SE df t.ratio p.value
 epilimnion - thermocline
                               8.20 1.26 42
                                               6.512 < .0001
                              17.77 1.11 42 15.997
 epilimnion - hypolimnion
                                                     <.0001
 thermocline - hypolimnion
                               9.57 1.33 42
                                               7.207 < .0001
trtools::contrast(m,
  a = list(layer = c("epilimnion", "epilimnion", "thermocline")),
  b = list(layer = c("thermocline", "hypolimnion", "hypolimnion")),
  cnames = c("E-T","E-H", "T-H"))
```

```
estimate se lower upper tvalue df pvalue
E-T 8.200 1.259 5.659 10.74 6.512 42 7.293e-08
E-H 17.767 1.111 15.525 20.01 15.997 42 1.784e-19
T-H 9.567 1.327 6.888 12.25 7.207 42 7.363e-09
```

The adjust = "none" option for pairs specifies that no adjustment be made to confidence intervals or tests for the family-wise Type I error rate.¹

Something to note when using the weights argument with the emmeans function is that the weights that are used must sum to one, and if they do not they will be normalized so that they do. For example, the following provide the same result.

```
emmeans(m, ~ 1, weights = c(1/7, 2/7, 4/7))
                   SE df lower.CL upper.CL
 1
         emmean
              7 0.572 42
                             5.85
 overall
                                       8.16
Results are averaged over the levels of: layer
Confidence level used: 0.95
emmeans(m, ~ 1, weights = c(1, 2, 4)) # original weights multiplied by 7
         emmean
                   SE df lower.CL upper.CL
              7 0.572 42
                             5.85
                                       8.16
 overall
```

Results are averaged over the levels of: layer Confidence level used: 0.95

If you want to use weights that do not sum to one, you can use the contrast function from the emmeans package (different from the function of the same name from **trtools**).

¹The family-wise Type I error rate is the probability of making at least one Type I error. If it is desired that the family-wise Type I error rate be no greater than α (default is 0.05), then some adjustment can be made. This adjustment is seen in the p-values and confidence intervals. The most general method is to use adjust = "mvt". Some special cases are more widely known such as Tukey (adjust = "tukey") and Bonferroni (adjust = "bonferroni"), but the adjustment based on the multivariate t-distribution (adjust = "mvt") is the most general and accurate. Note that an adjustment will produce "simultaneous" confidence intervals. A method of producing simultaneous confidence intervals has the property that the probability that all of the confidence intervals will contain the quantities being estimated is equal to the specified confidence level (95% by default). The multivariate t-distribution adjustment is perhaps not as well known, so a reference that you can cite is Edwards, D. & Berry, J. T. (1987). The efficiency of simulation-based multiple comparisons. Biometrics, 43(4), 913–928.

```
emmeans(m, ~1, weights = c(1/7, 2/7, 4/7))
                   SE df lower.CL upper.CL
         emmean
              7 0.572 42
                             5.85
                                      8.16
overall
Results are averaged over the levels of: layer
Confidence level used: 0.95
emmeans::contrast(emmeans(m, ~layer), method = list(layer = c(1/7, 2/7, 4/7)), infer = TRUE)
                      SE df lower.CL upper.CL t.ratio p.value
 contrast estimate
                                         8.16 12.245 < .0001
                 7 0.572 42
                                5.85
layer
```

Confidence level used: 0.95

But suppose we wanted to estimate the number of daphnia in the lake (τ) . It can be shown that this is

```
\tau = 700000\mu = 700000 \left(\frac{1}{7}\mu_e + \frac{2}{7}\mu_t + \frac{4}{7}\mu_h\right) = 100000\mu_e + 200000\mu_t + 400000\mu_h.
```

Note that there are 700kL in the lake, which is 700000L (which is the scale used for the observations). This can be estimated as follows.

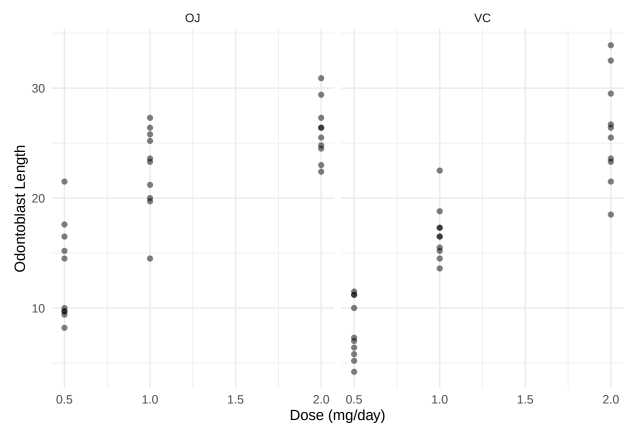
Confidence level used: 0.95

Another approach is to use lincon but your weights/coefficients will be different since they are applied to β_0 , β_1 , and β_2 .

Marginal Means and "Main Effects"

Consider data from a randomized experiment with guinea pigs administered one of three doses of vitamin C (0.5, 1, or 2 mg/day) via one of two supplement methods: orange juice (OJ) or ascorbic acid (VC).

```
p <- ggplot(ToothGrowth, aes(x = dose, y = len)) +
    geom_point(alpha = 0.5) + facet_wrap(~supp) +
    labs(x = "Dose (mg/day)", y = "Odontoblast Length") + theme_minimal()
plot(p)</pre>
```



Here we are going to model dose as a categorical variable so we need to coerce it to a factor. Perhaps the safest approach is to create a new variable.

```
ToothGrowth$dosef <- factor(ToothGrowth$dose)</pre>
```

Note: Whether a variable is a numeric, a factor, or something else can be seen use str (for "structure").

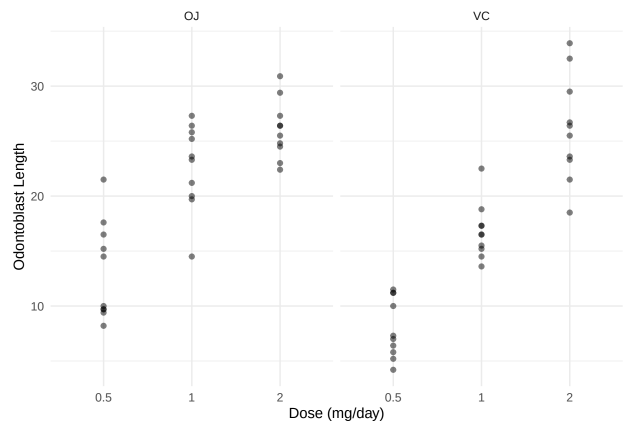
str(ToothGrowth)

```
'data.frame': 60 obs. of 4 variables:
$ len : num 4.2 11.5 7.3 5.8 6.4 10 11.2 11.2 5.2 7 ...
$ supp : Factor w/ 2 levels "OJ","VC": 2 2 2 2 2 2 2 2 2 2 2 ...
$ dose : num 0.5 0.5 0.5 0.5 0.5 0.5 0.5 0.5 ...
$ dosef: Factor w/ 3 levels "0.5","1","2": 1 1 1 1 1 1 1 1 1 ...
```

Notice that ggplot responds differently.

summary(ToothGrowth)

```
len
                supp
                              dose
                                        dosef
Min.
        : 4.2
                OJ:30
                        Min.
                                :0.50
                                        0.5:20
 1st Qu.:13.1
                VC:30
                        1st Qu.:0.50
                                        1 :20
Median:19.2
                        Median:1.00
                                        2 :20
Mean
       :18.8
                        Mean
                               :1.17
 3rd Qu.:25.3
                         3rd Qu.:2.00
Max.
        :33.9
                        Max.
                                :2.00
p <- ggplot(ToothGrowth, aes(x = dosef, y = len)) +</pre>
  geom_point(alpha = 0.5) + facet_wrap(~supp) +
  labs(x = "Dose (mg/day)", y = "Odontoblast Length") + theme_minimal()
plot(p)
```



Now consider the following linear model.

```
m <- lm(len ~ dosef + supp + dosef:supp, data = ToothGrowth)
summary(m)$coefficients</pre>
```

	Estimate Std.	Error	t value	Pr(> t)
(Intercept)	13.23	1.148	11.5208	3.603e-16
dosef1	9.47	1.624	5.8312	3.176e-07
dosef2	12.83	1.624	7.9002	1.430e-10
suppVC	-5.25	1.624	-3.2327	2.092e-03
dosef1:suppVC	-0.68	2.297	-0.2961	7.683e-01
dosef2:suppVC	5.33	2.297	2.3207	2.411e-02

The model is

$$E(Y_i) = \beta_0 + \beta_1 x_{i1} + \beta_2 x_{i2} + \beta_3 x_{i3} + \beta_4 x_{i4} + \beta_5 x_{i5},$$

where

$$x_{i1} = \begin{cases} 1, & \text{if dose is 1 mg/day,} \\ 0, & \text{otherwise,} \end{cases}$$

$$x_{i2} = \begin{cases} 1, & \text{if dose is 2 mg/day,} \\ 0, & \text{otherwise,} \end{cases}$$

$$x_{i3} = \begin{cases} 1, & \text{if supplement type is VC} \\ 0, & \text{otherwise,} \end{cases}$$

$$x_{i4} = x_{i1}x_{i3} = \begin{cases} 1, & \text{if dose is 1 mg/day and supplement type is VC,} \\ 0, & \text{otherwise,} \end{cases}$$

$$x_{i5} = x_{i2}x_{i3} = \begin{cases} 1, & \text{if dose is 2 mg/day and supplement type is VC,} \\ 0, & \text{otherwise.} \end{cases}$$

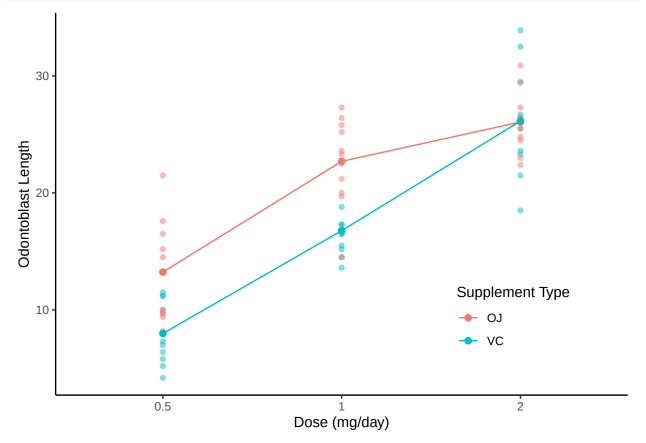
We can write this model case-wise.

$$E(Y_i) = \begin{cases} \beta_0, & \text{if dose is } 0.5 \text{ mg/day and supplement type is OJ,} \\ \beta_0 + \beta_1, & \text{if dose is } 1 \text{ mg/day and supplement type is OJ,} \\ \beta_0 + \beta_2, & \text{if dose is } 2 \text{ mg/day and supplement type is OJ,} \\ \beta_0 + \beta_3, & \text{if dose is } 0.5 \text{ mg/day and supplement type is VC,} \\ \beta_0 + \beta_1 + \beta_3 + \beta_4, & \text{if dose is } 1 \text{ mg/day and supplement type is VC,} \\ \beta_0 + \beta_2 + \beta_3 + \beta_5, & \text{if dose is } 2 \text{ mg/day and supplement type is VC.} \end{cases}$$

And we can visualize it.

```
d <- expand.grid(dosef = levels(ToothGrowth$dosef), supp = levels(ToothGrowth$supp))
d$yhat <- predict(m, newdata = d)

p <- ggplot(ToothGrowth, aes(x = dosef, y = len, color = supp)) +
    geom_point(alpha = 0.5) + theme_classic() +
    theme(legend.position = c(0.8,0.2)) +
    geom_point(aes(y = yhat), size = 2, data = d) +
    geom_line(aes(y = yhat, group = supp), data = d) +
    labs(x = "Dose (mg/day)", y = "Odontoblast Length", color = "Supplement Type")
plot(p)</pre>
```



We might want to compare the two supplement types at each dose level. These are sometimes called simple effects. We can do this using contrast.

```
trtools::contrast(m,
    a = list(supp = "OJ", dosef = c("0.5","1","2")),
    b = list(supp = "VC", dosef = c("0.5","1","2")),
    cnames = c("OJ-VC @ 0.5 mg/day","OJ-VC @ 1.0 mg/day","OJ-VC @ 2.0 mg/day"))
```

```
estimate se lower upper tvalue df pvalue OJ-VC @ 0.5 mg/day 5.25 1.624 1.994 8.506 3.23273 54 0.0020925 OJ-VC @ 1.0 mg/day 5.93 1.624 2.674 9.186 3.65144 54 0.0005897 OJ-VC @ 2.0 mg/day -0.08 1.624 -3.336 3.176 -0.04926 54 0.9608934
```

We can also do this use emmeans. First note that we can use emmeans to also estimate the expected response for each level of supp within each level of dosef.

```
emmeans(m, ~ supp | dosef)
```

```
dosef = 0.5:
               SE df lower.CL upper.CL
 supp emmean
OJ
       13.23 1.15 54
                        10.93
                                  15.5
VC
       7.98 1.15 54
                         5.68
                                  10.3
dosef = 1:
               SE df lower.CL upper.CL
 supp emmean
       22.70 1.15 54
                        20.40
                                  25.0
VC
       16.77 1.15 54
                        14.47
                                  19.1
dosef = 2:
 supp emmean
               SE df lower.CL upper.CL
       26.06 1.15 54
                        23.76
                                  28.4
VC
       26.14 1.15 54
                        23.84
                                  28.4
```

Confidence level used: 0.95

We can use pairs to make inferences about the differences between levels of supp within each level of dosef.

```
pairs(emmeans(m, ~ supp | dosef), adjust = "none", infer = TRUE)
```

```
dosef = 0.5:
 contrast estimate
                    SE df lower.CL upper.CL t.ratio p.value
OJ - VC
             5.25 1.62 54
                               1.99
                                       8.51
                                              3.233 0.0021
dosef = 1:
 contrast estimate
                    SE df lower.CL upper.CL t.ratio p.value
             5.93 1.62 54
                               2.67
                                        9.19
                                               3.651 0.0006
dosef = 2:
                    SE df lower.CL upper.CL t.ratio p.value
 contrast estimate
OJ - VC
            -0.08 1.62 54
                              -3.34
                                       3.18 -0.049 0.9609
```

Confidence level used: 0.95

The "main effect" of supplement method concerns μ_{OJ} and μ_{VC} , defined as

$$\mu_{\rm OJ} = \frac{\mu_{\rm OJ,0.5} + \mu_{\rm OJ,1.0} + \mu_{\rm OJ,2.0}}{3}, \quad \mu_{\rm VC} = \frac{\mu_{\rm VC,0.5} + \mu_{\rm VC,1.0} + \mu_{\rm OJ,2.0}}{3}.$$

So μ_{OJ} and μ_{VC} are the marginal means for supp, averaging over levels of dose.

emmeans(m, ~ supp)

```
      supp
      emmean
      SE df lower.CL upper.CL
      upper.CL upper.CL

      OJ
      20.7 0.663 54
      19.3 22.0

      VC
      17.0 0.663 54
      15.6 18.3
```

Results are averaged over the levels of: dosef Confidence level used: 0.95

Inference for the main effect $\mu_{OJ} - \mu_{VC}$ can then be obtained as follows.

```
pairs(emmeans(m, ~ supp), infer = TRUE)
```

```
contrast estimate SE df lower.CL upper.CL t.ratio p.value OJ - VC 3.7 0.938 54 1.82 5.58 3.946 0.0002
```

Results are averaged over the levels of: dosef Confidence level used: 0.95

This main effect is simply a single linear function of $\beta_0, \beta_1, \dots, \beta_5$. From the case-wise representation of the model,

$$E(Y_i) = \begin{cases} \beta_0, & \text{if dose is } 0.5 \text{ mg/day and supplement type is OJ,} \\ \beta_0 + \beta_1, & \text{if dose is } 1 \text{ mg/day and supplement type is OJ,} \\ \beta_0 + \beta_2, & \text{if dose is } 2 \text{ mg/day and supplement type is OJ,} \\ \beta_0 + \beta_3, & \text{if dose is } 0.5 \text{ mg/day and supplement type is VC,} \\ \beta_0 + \beta_1 + \beta_3 + \beta_4, & \text{if dose is } 1 \text{ mg/day and supplement type is VC,} \\ \beta_0 + \beta_2 + \beta_3 + \beta_5, & \text{if dose is } 2 \text{ mg/day and supplement type is VC,} \end{cases}$$

we have that $\mu_{\text{OJ},0.5} = \beta_0$, $\mu_{\text{OJ},1.0} = \beta_0 + \beta_1$, $\mu_{\text{OJ},2.0} = \beta_0 + \beta_3$, $\mu_{\text{VC},0.5} = \beta_0 + \beta_3$, $\mu_{\text{VC},1.0} = \beta_0 + \beta_1 + \beta_3 + \beta_4$, $\mu_{\text{VC},2.0} = \beta_0 + \beta_1 + \beta_3 + \beta_5$. So, we can write this as

$$\mu_{\rm OJ} - \mu_{\rm VC} = \frac{\mu_{\rm OJ,0.5} + \mu_{\rm OJ,1.0} + \mu_{\rm OJ,2.0}}{3} - \frac{\mu_{\rm VC,0.5} + \mu_{\rm VC,1.0} + \mu_{\rm OJ,2.0}}{3} = -\beta_3 - \frac{1}{3}\beta_4 - \frac{1}{3}\beta_5.$$

lincon(m, a =
$$c(0,0,0,-1,-1/3,-1/3)$$
)

```
estimate se lower upper tvalue df pvalue (0,0,0,-1,-1/3,-1/3),0 3.7 0.9376 1.82 5.58 3.946 54 0.0002312
```

Clearly using emmeans is much less tedious!

The "main effect" of dose concerns differences among the marginal means of dose defined as $\mu_{0.5}$, μ_1 and μ_2 where

$$\mu_{0.5} = \frac{\mu_{\text{OJ},0.5} + \mu_{\text{VC},0.5}}{2}, \quad \mu_1 = \frac{\mu_{\text{OJ},1} + \mu_{\text{VC},1}}{2}, \quad \mu_2 = \frac{\mu_{\text{OJ},2} + \mu_{\text{VC},2}}{2}.$$

emmeans(m, ~ dosef)

```
    dosef
    emmean
    SE
    df
    lower.CL
    upper.CL

    0.5
    10.6
    0.812
    54
    8.98
    12.2

    1
    19.7
    0.812
    54
    18.11
    21.4

    2
    26.1
    0.812
    54
    24.47
    27.7
```

Results are averaged over the levels of: supp Confidence level used: 0.95

```
pairs(emmeans(m, ~ dosef), adjust = "none")
```

```
contrast estimate SE df t.ratio p.value
dosef0.5 - dosef1 -9.13 1.15 54 -7.951 <.0001
dosef0.5 - dosef2 -15.49 1.15 54 -13.493 <.0001
dosef1 - dosef2 -6.37 1.15 54 -5.543 <.0001
```

Results are averaged over the levels of: supp

```
pairs(emmeans(m, ~ dosef), reverse = TRUE, adjust = "none")
```

```
      contrast
      estimate
      SE df t.ratio p.value

      dosef1 - dosef0.5
      9.13 1.15 54 7.951 <.0001</td>

      dosef2 - dosef0.5
      15.49 1.15 54 13.493 <.0001</td>

      dosef2 - dosef1
      6.37 1.15 54 5.543 <.0001</td>
```

Results are averaged over the levels of: supp

In ANOVA tables the test of the "main effect" is the (joint) null hypothesis that all pairwise differences are zero. For the variable dose the null hypothesis is $\mu_{0.5} = \mu_1 = \mu_2$. This can be done using the test function.

```
test(pairs(emmeans(m, ~ dosef)), joint = TRUE)
```

```
df1 df2 F.ratio p.value note
2 54 92.000 <.0001 d
```

d: df1 reduced due to linear dependence

This is the traditional main effect that is sometimes reported in an "ANOVA table" such as that produced by Anova from the car package.

```
library(car)
m <- lm(len ~ dosef + supp + dosef:supp, data = ToothGrowth,
    contrast = list(dosef = contr.sum, supp = contr.sum))
Anova(m, type = 3)</pre>
```

Anova Table (Type III tests)

Response: len

```
Sum Sq Df F value Pr(>F)

(Intercept) 21236 1 1610.39 < 2e-16 ***
dosef 2426 2 92.00 < 2e-16 ***
supp 205 1 15.57 0.00023 ***
dosef:supp 108 2 4.11 0.02186 *
Residuals 712 54
```

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

The option contrast = list(dosef = contr.sum, supp = contr.sum) is necessary here for the Anova function to do the correct calculations.²

The test of the main effect of supplement method was given by

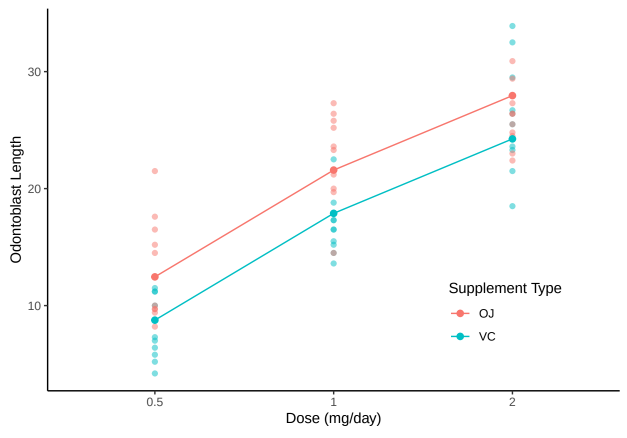
```
pairs(emmeans(m, ~ supp), infer = TRUE)
```

²I am demonstrating here what is sometimes called inferences based on Type III sums of squares. Another common approach is to use what is called Type II sums of squares. This can be done with the Anova function with type = 2. For inferences based on Type II sums of squares with the functions from the emmeans package an extra step is needed (email me for an example if you really want to know how to do it).

```
Results are averaged over the levels of: dosef
Confidence level used: 0.95
```

We do not need a joint test here since there are only two marginal means, but here it is anyway.

```
test(pairs(emmeans(m, ~ supp)), joint = TRUE)
df1 df2 F.ratio p.value
  1 54 15.572 0.0002
Now consider a model without the interaction.
m <- lm(len ~ dosef + supp, data = ToothGrowth)
summary(m)$coefficients
           Estimate Std. Error t value Pr(>|t|)
                         0.9883 12.603 5.490e-18
(Intercept) 12.45
dosef1
               9.13
                        1.2104 7.543 4.383e-10
dosef2
               15.50
                        1.2104 12.802 2.852e-18
suppVC
               -3.70
                         0.9883 -3.744 4.293e-04
d <- expand.grid(dosef = levels(ToothGrowth$dosef), supp = levels(ToothGrowth$supp))</pre>
d$yhat <- predict(m, newdata = d)</pre>
p <- ggplot(ToothGrowth, aes(x = dosef, y = len, color = supp)) +</pre>
  geom_point(alpha = 0.5) + theme_classic() +
 theme(legend.position = c(0.8,0.2)) +
  geom_point(aes(y = yhat), size = 2, data = d) +
 geom_line(aes(y = yhat, group = supp), data = d) +
 labs(x = "Dose (mg/day)", y = "Odontoblast Length", color = "Supplement Type")
plot(p)
```



Note that without the interaction the "simple effects" and "main effects" are equivalent. Here are the simple and main effects for supplement.

```
pairs(emmeans(m, ~ supp | dosef)) # simple
dosef = 0.5:
 contrast estimate
                      SE df t.ratio p.value
OJ - VC
               3.7 0.988 56
                              3.744 0.0004
dosef = 1:
 contrast estimate
                      SE df t.ratio p.value
OJ - VC
               3.7 0.988 56
                              3.744 0.0004
dosef = 2:
 contrast estimate
                      SE df t.ratio p.value
               3.7 0.988 56
                              3.744 0.0004
                                   # main
pairs(emmeans(m, ~ supp))
contrast estimate
                      SE df t.ratio p.value
OJ - VC
               3.7 0.988 56
                              3.744 0.0004
Results are averaged over the levels of: dosef
And here are the simple and main effects for dose.
pairs(emmeans(m, ~ dosef | supp), adjust = "none") # simple
supp = OJ:
                              SE df t.ratio p.value
contrast
                   estimate
```

```
dosef0.5 - dosef1 -9.13 1.21 56 -7.543 <.0001
dosef0.5 - dosef2 -15.49 1.21 56 -12.802 <.0001
dosef1 - dosef2 -6.37 1.21 56 -5.259 <.0001
supp = VC:
contrast
                 estimate
                            SE df t.ratio p.value
dosef0.5 - dosef1 -9.13 1.21 56 -7.543 <.0001
dosef0.5 - dosef2 -15.49 1.21 56 -12.802 <.0001
dosef1 - dosef2
                    -6.37 1.21 56 -5.259 <.0001
pairs(emmeans(m, ~ dosef), adjust = "none")
                                                 # main
contrast
                  estimate
                             SE df t.ratio p.value
dosef0.5 - dosef1
                    -9.13 1.21 56 -7.543 <.0001
dosef0.5 - dosef2 -15.49 1.21 56 -12.802 <.0001
dosef1 - dosef2
                  -6.37 1.21 56 -5.259 <.0001
Results are averaged over the levels of: supp
The joint test of the overall main effect for dose can be be obtained as follows.
test(pairs(emmeans(m, ~ dosef)), joint = TRUE)
df1 df2 F.ratio p.value note
```

d: df1 reduced due to linear dependence

2 56 82.810 <.0001 d