Interactions and post-hocs

KEY

Wed Apr 15 14:21:50 2020

## MANOVA interactions

Import the mouse data:

library(readxl)  
read\_excel("mouse.xlsx","mouse") -> mouse

Make a new column in mouse with combinations of parasite.treatment and caloric.treatment:

mouse$parasite.caloric <- with(mouse, interaction(parasite.treatment, caloric.treatment))

### Fat and RMR as responses - MANOVA

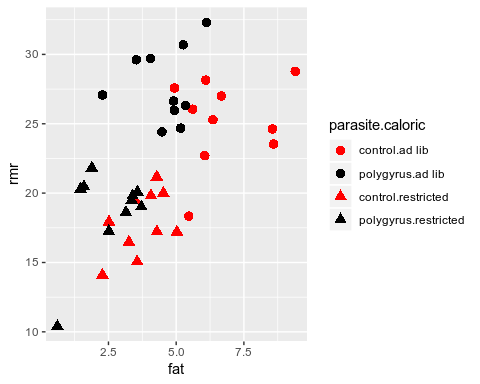
Load the ggplot2 library:

library(ggplot2)

## Registered S3 methods overwritten by 'ggplot2':  
## method from   
## [.quosures rlang  
## c.quosures rlang  
## print.quosures rlang

Graph the fat and rmr variables as a scatter plot, using color to indicate parasite treatment, and plot symbol to indicate caloric treatment:

ggplot(mouse, aes(x = fat, y = rmr, color = parasite.caloric, shape = parasite.caloric)) + geom\_point(size = 3) + scale\_color\_manual(values = c("red","black","red","black")) + scale\_shape\_manual(values = c(16,16,17,17))



**Question: ignore the plot symbol and just compare the colors - does it look like there is separation between control and polygyrus groups with respect to their fat and rmr values? How can you tell?**

Yes, the red points are slightly to the right of the black points.

**Question: ignore color and focus on the different plot symbols - does it look like there is separation between ad lib and restricted mice? How can you tell?**

Yes, the dots are to the upper right of the triangles.

To make a plot of the centroids we need to calculate them first. We can use the aggregate() function to calculate means for combinations of parasite.treatment and caloric.treatment:

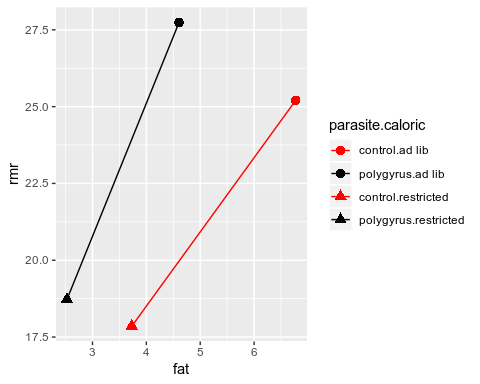
aggregate(cbind(fat, rmr, li, si) ~ parasite.treatment + caloric.treatment, data = mouse, FUN = mean) -> mouse.means

Add the combinations of parasite and caloric treatments to mouse.means:

mouse.means$parasite.caloric <- with(mouse.means, interaction(parasite.treatment, caloric.treatment))

Make the plot of the centroids:

ggplot(mouse.means, aes(x = fat, y = rmr, color = parasite.caloric, shape = parasite.caloric, group = parasite.treatment)) + geom\_point(size = 3) + scale\_color\_manual(values = c("red","black","red","black")) + scale\_shape\_manual(values = c(16,16,17,17)) + geom\_line()



**Question: why would you expect a main effect of parasite.treatment, based on this graph?**

Yes there should be a main effect of parasite.treatment because the red line is different on average from the black line.

**Question: why would you expect a main effect of caloric.treatment, based on this graph?**

Yes, caloric.treatment should be different because the circles are different on average from the triangles.

**Question: why does the fact that the lines are about the same length and pointed in the same direction indicate that we should not expect an interaction between parasite.treatment and caloric.treatment? Hint: is the amount of difference between ad lib and restricted mice about the same for both colors, and is the way they change similar?**

There should not be a significant interaction, because the amount of difference between the ad lib and restricted caloric treatments are the same for polygyrus and control parasite treatment groups.

Now run the MANOVA to test for main effects and interaction between parasite.treatment and caloric.treatment:

lm(cbind(fat, rmr) ~ parasite.treatment \* caloric.treatment, data = mouse) -> fat.rmr.manova  
  
anova(fat.rmr.manova, test = "Wilks")

## Analysis of Variance Table  
##   
## Df Wilks approx F num Df den Df  
## (Intercept) 1 0.01373 1257.38 2 35  
## parasite.treatment 1 0.52434 15.88 2 35  
## caloric.treatment 1 0.25899 50.07 2 35  
## parasite.treatment:caloric.treatment 1 0.90863 1.76 2 35  
## Residuals 36   
## Pr(>F)   
## (Intercept) < 2.2e-16 \*\*\*  
## parasite.treatment 1.239e-05 \*\*\*  
## caloric.treatment 5.402e-11 \*\*\*  
## parasite.treatment:caloric.treatment 0.187   
## Residuals   
## ---  
## Signif. codes: 0 '\*\*\*' 0.001 '\*\*' 0.01 '\*' 0.05 '.' 0.1 ' ' 1

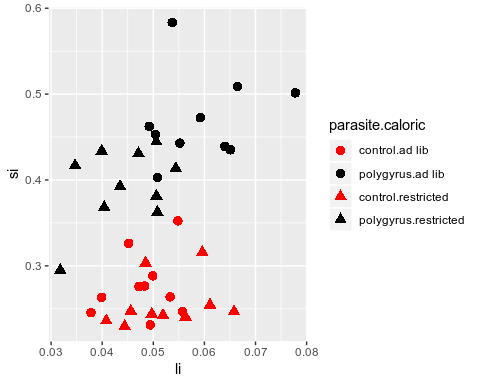
**Question: are there significant main effects for the predictors (if so, which)? Is there a significant interaction?**

The main effects are both significant, but there is not a significant interaction.

### Large intestine and small intestine as responses - MANOVA

Now do the graphs again using li and si. Start with the raw data in mouse:

ggplot(mouse, aes(x = li, y = si, color = parasite.caloric, shape = parasite.caloric)) + geom\_point(size = 3) + scale\_color\_manual(values = c("red","black","red","black")) + scale\_shape\_manual(values = c(16,16,17,17))



**Question: Looking just at color, does it appear that there will be a difference between control and polygyrus treatments in their large and small intestine lengths? How can you tell?**

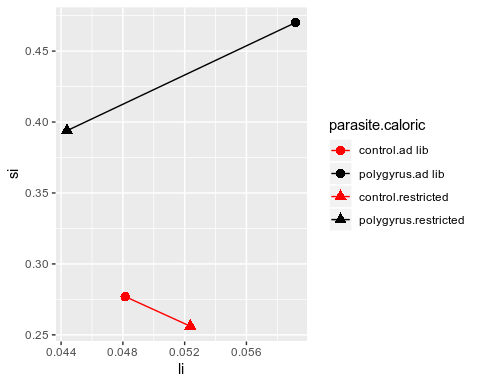
Yes, parasite.treatment should be significant becasue the red and black do not overlap.

**Now looking just at the circles and triangles, does it look like there will be a difference between ad lib and restricted animals? How can you tell?**

There may not be a main effect of caloric treatment, because the circles and triangles overlap a lot.

Now the centroids:

ggplot(mouse.means, aes(x = li, y = si, color = parasite.caloric, shape = parasite.caloric, group = parasite.treatment)) + geom\_point(size = 3) + scale\_color\_manual(values = c("red","black","red","black")) + scale\_shape\_manual(values = c(16,16,17,17)) + geom\_line()



**Question: why would you expect a main effect of parasite.treatment, based on this graph?**

Parasite treatment should be significant because the red line is different from the black line on average.

**Question: why would you expect a main effect of caloric.treatment, based on this graph?**

The main effect of caloric.treatment should be significant becasue the mean of the triangles is diferent than the mean of the circles.

**Question: why does the fact that the lines are different lengths and pointed in the different directions indicate that we should expect an interaction between parasite.treatment and caloric.treatment? Hint: is the amount of difference between ad lib and restricted mice about the same for both colors, and is the way they change similar?**

The lines are different lengths and point in different directions, so the difference between caloric treatments is not the same for polygyrus and control groups.

Run your MANOVA using li and si as responses:

lm(cbind(li, si) ~ parasite.treatment \* caloric.treatment, data = mouse) -> li.si.manova  
  
anova(li.si.manova, test = "Wilks")

## Analysis of Variance Table  
##   
## Df Wilks approx F num Df den Df  
## (Intercept) 1 0.00976 1775.53 2 35  
## parasite.treatment 1 0.17551 82.21 2 35  
## caloric.treatment 1 0.70233 7.42 2 35  
## parasite.treatment:caloric.treatment 1 0.68917 7.89 2 35  
## Residuals 36   
## Pr(>F)   
## (Intercept) < 2.2e-16 \*\*\*  
## parasite.treatment 5.963e-14 \*\*\*  
## caloric.treatment 0.002063 \*\*   
## parasite.treatment:caloric.treatment 0.001482 \*\*   
## Residuals   
## ---  
## Signif. codes: 0 '\*\*\*' 0.001 '\*\*' 0.01 '\*' 0.05 '.' 0.1 ' ' 1

**Question: is there a significant main effect of parasite.treatment? How do you know?**

The parasite.treatment effect is statistically significant because its p-value is less than 0.05.

**Question: is there a significant main effect of caloric.treatment? How do you know?**

The caloric.treatment effect is statistically significant because its p-value is also less than 0.05.

**Question: is there a significant interaction between parasite.treatment and caloric.treatment? How do yo know?**

Yes, the interaction term also has a p-value less than 0.05.

## Post-hoc procedures, with and without interactions

The procedure for following up after a significant MANOVA depends on whether there is a significant interaction or not.

### Post-hoc procedures with a non-significant interaction - rmr and fat

Since the interaction was not significant we can focus on main effects. With only two groups we already know that the groups are different in a multivariate sense, but we still need to test main effects of each predictor on fat, and on rmr separately. With only two groups it isn’t necessary to use multiple comparison procedures, and we can just do a univariate factorial ANOVA.

First rmr:

anova(lm(rmr ~ parasite.treatment + caloric.treatment, data = mouse))

## Analysis of Variance Table  
##   
## Response: rmr  
## Df Sum Sq Mean Sq F value Pr(>F)   
## parasite.treatment 1 29.08 29.08 3.6421 0.06412 .   
## caloric.treatment 1 669.75 669.75 83.8736 4.765e-11 \*\*\*  
## Residuals 37 295.46 7.99   
## ---  
## Signif. codes: 0 '\*\*\*' 0.001 '\*\*' 0.01 '\*' 0.05 '.' 0.1 ' ' 1

**Question: is rmr different between parasite treatments? What about between caloric treatments? How can you tell?**

RMR is not different between parasite treatment groups, but is different between caloric treatment groups.

Now fat:

anova(lm(fat ~ parasite.treatment + caloric.treatment, data = mouse))

## Analysis of Variance Table  
##   
## Response: fat  
## Df Sum Sq Mean Sq F value Pr(>F)   
## parasite.treatment 1 28.365 28.365 20.474 6.057e-05 \*\*\*  
## caloric.treatment 1 65.656 65.656 47.391 4.048e-08 \*\*\*  
## Residuals 37 51.261 1.385   
## ---  
## Signif. codes: 0 '\*\*\*' 0.001 '\*\*' 0.01 '\*' 0.05 '.' 0.1 ' ' 1

**Question: is fat different between parasite treatments? What about between caloric treatments? How can you tell?**

Fat is different between both parasite treatments and caloric treatments.

### Significant interaction - li and si

First, re-run the MANOVA using parasite.caloric as the only predictor:

lm(cbind(li, si) ~ parasite.caloric, data = mouse) -> parasite.caloric.manova  
anova(parasite.caloric.manova, test = "Wilks")

## Analysis of Variance Table  
##   
## Df Wilks approx F num Df den Df Pr(>F)   
## (Intercept) 1 0.00976 1775.53 2 35 < 2.2e-16 \*\*\*  
## parasite.caloric 3 0.11022 23.47 6 70 5.307e-15 \*\*\*  
## Residuals 36   
## ---  
## Signif. codes: 0 '\*\*\*' 0.001 '\*\*' 0.01 '\*' 0.05 '.' 0.1 ' ' 1

Source the hotelling\_t2\_posthoc.R file:

source("hotelling\_t2\_posthoc.R")

Run the multivariate Hotelling T2 comparisons for each pair of groups in parasite.caloric:

tsquare.ph(parasite.caloric.manova)

## $`T2 Post-hoc comparisons`  
## Comparison P.value  
## 1 control.ad lib - polygyrus.ad lib 1.57934295596724e-07  
## 2 control.ad lib - control.restricted 0.128494924022069  
## 3 control.ad lib - polygyrus.restricted 8.08486847414539e-06  
## 4 polygyrus.ad lib - control.restricted 1.28325950279748e-08  
## 5 polygyrus.ad lib - polygyrus.restricted 0.0011191292978385  
## 6 control.restricted - polygyrus.restricted 1.44957574190861e-07

**Quetion: which groups are significantly different according to these Hotelling T2 tests?**

All of them are, except for control.ad lib and control.restricted

Lastly, do Tukey post-hocs on li and si separately:

TukeyHSD(aov(li ~ parasite.caloric, data = mouse))

## Tukey multiple comparisons of means  
## 95% family-wise confidence level  
##   
## Fit: aov(formula = li ~ parasite.caloric, data = mouse)  
##   
## $parasite.caloric  
## diff lwr upr  
## polygyrus.ad lib-control.ad lib 0.01104 0.001681469 0.020398531  
## control.restricted-control.ad lib 0.00422 -0.005138531 0.013578531  
## polygyrus.restricted-control.ad lib -0.00377 -0.013128531 0.005588531  
## control.restricted-polygyrus.ad lib -0.00682 -0.016178531 0.002538531  
## polygyrus.restricted-polygyrus.ad lib -0.01481 -0.024168531 -0.005451469  
## polygyrus.restricted-control.restricted -0.00799 -0.017348531 0.001368531  
## p adj  
## polygyrus.ad lib-control.ad lib 0.0154259  
## control.restricted-control.ad lib 0.6218967  
## polygyrus.restricted-control.ad lib 0.7008222  
## control.restricted-polygyrus.ad lib 0.2209573  
## polygyrus.restricted-polygyrus.ad lib 0.0007724  
## polygyrus.restricted-control.restricted 0.1171011

TukeyHSD(aov(si ~ parasite.caloric, data = mouse))

## Tukey multiple comparisons of means  
## 95% family-wise confidence level  
##   
## Fit: aov(formula = si ~ parasite.caloric, data = mouse)  
##   
## $parasite.caloric  
## diff lwr  
## polygyrus.ad lib-control.ad lib 0.19318542 0.14344444  
## control.restricted-control.ad lib -0.02090357 -0.07064455  
## polygyrus.restricted-control.ad lib 0.11695700 0.06721602  
## control.restricted-polygyrus.ad lib -0.21408899 -0.26382997  
## polygyrus.restricted-polygyrus.ad lib -0.07622842 -0.12596940  
## polygyrus.restricted-control.restricted 0.13786057 0.08811959  
## upr p adj  
## polygyrus.ad lib-control.ad lib 0.24292641 0.0000000  
## control.restricted-control.ad lib 0.02883742 0.6725941  
## polygyrus.restricted-control.ad lib 0.16669799 0.0000015  
## control.restricted-polygyrus.ad lib -0.16434801 0.0000000  
## polygyrus.restricted-polygyrus.ad lib -0.02648744 0.0011418  
## polygyrus.restricted-control.restricted 0.18760155 0.0000000

**Question: which groups had different large intestine lengths?**

For large intestine, only polygyrus.ad lib and control.ad lib, and polygyrus.restricted and polygyrus.ad lib are different.

**Question: which groups had different small intestine lengths?**

For small intestine, all are different except for control.restricted and control.ad lib.

**Question: were there any groups that were only different when both variables were used? Or, were all of the groups that were different in the Hotelling’s T2 tests also different in either their li or si measurements?**

No, the only groups that were never different using li and si separately were control.ad lib and control.restricted, and that was the comparison that was not different using Hotelling’s T2 either.

## Abledu paper - post-hocs following significant multivariate analysis

Last time you confirmed that you had multivariate differences between the sexes for left feet and for right feet. You also confirmed that left and right feet different, using a paired multivariate analysis. Now you can follow up with comparisons of each variable to determine if the differences are present for every variable, or if only some of the variables differ.

Import the food measurments data:

data.frame(read\_excel("footprint\_dimensions.xlsx")) -> foot  
foot[complete.cases(foot),] -> foot.complete

Make a list of variable names for the left foot (columns 2 to 9) and the right foot (columns 10 to 17):

left.var <- colnames(foot.complete)[2:9]  
right.var <- colnames(foot.complete)[10:17]

Calculate the differences between right and left:

foot.complete[left.var] - foot.complete[right.var] -> foot.complete.diffs

Conduct the paired t-tests for each variable, using sapply():

apply(foot.complete.diffs, MARGIN = 2, FUN = function(x) t.test(x)) -> foot.paired.t  
foot.paired.t

## $T1.l  
##   
## One Sample t-test  
##   
## data: x  
## t = 1.5507, df = 121, p-value = 0.1236  
## alternative hypothesis: true mean is not equal to 0  
## 95 percent confidence interval:  
## -0.02290628 0.18848005  
## sample estimates:  
## mean of x   
## 0.08278689   
##   
##   
## $T2.l  
##   
## One Sample t-test  
##   
## data: x  
## t = 3.5317, df = 121, p-value = 0.0005851  
## alternative hypothesis: true mean is not equal to 0  
## 95 percent confidence interval:  
## 0.07600102 0.26990062  
## sample estimates:  
## mean of x   
## 0.1729508   
##   
##   
## $T3.l  
##   
## One Sample t-test  
##   
## data: x  
## t = 3.1668, df = 121, p-value = 0.001951  
## alternative hypothesis: true mean is not equal to 0  
## 95 percent confidence interval:  
## 0.04700772 0.20381196  
## sample estimates:  
## mean of x   
## 0.1254098   
##   
##   
## $T4.l  
##   
## One Sample t-test  
##   
## data: x  
## t = 2.4085, df = 121, p-value = 0.01753  
## alternative hypothesis: true mean is not equal to 0  
## 95 percent confidence interval:  
## 0.01984308 0.20310774  
## sample estimates:  
## mean of x   
## 0.1114754   
##   
##   
## $T5.l  
##   
## One Sample t-test  
##   
## data: x  
## t = 3.3873, df = 121, p-value = 0.0009528  
## alternative hypothesis: true mean is not equal to 0  
## 95 percent confidence interval:  
## 0.1035428 0.3948178  
## sample estimates:  
## mean of x   
## 0.2491803   
##   
##   
## $BAB.l  
##   
## One Sample t-test  
##   
## data: x  
## t = -0.32045, df = 121, p-value = 0.7492  
## alternative hypothesis: true mean is not equal to 0  
## 95 percent confidence interval:  
## -0.08825530 0.06366513  
## sample estimates:  
## mean of x   
## -0.01229508   
##   
##   
## $BAH.l  
##   
## One Sample t-test  
##   
## data: x  
## t = 0.48796, df = 121, p-value = 0.6265  
## alternative hypothesis: true mean is not equal to 0  
## 95 percent confidence interval:  
## -0.05512989 0.09119546  
## sample estimates:  
## mean of x   
## 0.01803279   
##   
##   
## $Index.l  
##   
## One Sample t-test  
##   
## data: x  
## t = 0.68422, df = 121, p-value = 0.4951  
## alternative hypothesis: true mean is not equal to 0  
## 95 percent confidence interval:  
## -0.6367858 1.3094088  
## sample estimates:  
## mean of x   
## 0.3363115

Extract the p-values:

sapply(foot.paired.t, FUN = function(x) x$p.value)

## T1.l T2.l T3.l T4.l T5.l   
## 0.1235850701 0.0005851031 0.0019510352 0.0175275494 0.0009528220   
## BAB.l BAH.l Index.l   
## 0.7491808978 0.6264592813 0.4951416351

Conduct two-sample t-tests for every variable between the sexes:

apply(foot.complete[,-1], MARGIN = 2, FUN = function(x) t.test(x ~ sex, data = foot.complete)) -> foot.sex.t  
foot.sex.t

## $T1.l  
##   
## Welch Two Sample t-test  
##   
## data: x by sex  
## t = -7.908, df = 119.9, p-value = 1.442e-12  
## alternative hypothesis: true difference in means is not equal to 0  
## 95 percent confidence interval:  
## -2.042505 -1.224530  
## sample estimates:  
## mean in group female mean in group male   
## 23.44679 25.08030   
##   
##   
## $T2.l  
##   
## Welch Two Sample t-test  
##   
## data: x by sex  
## t = -7.9203, df = 119.97, p-value = 1.348e-12  
## alternative hypothesis: true difference in means is not equal to 0  
## 95 percent confidence interval:  
## -2.144816 -1.286937  
## sample estimates:  
## mean in group female mean in group male   
## 23.14321 24.85909   
##   
##   
## $T3.l  
##   
## Welch Two Sample t-test  
##   
## data: x by sex  
## t = -8.2902, df = 119.82, p-value = 1.901e-13  
## alternative hypothesis: true difference in means is not equal to 0  
## 95 percent confidence interval:  
## -2.120853 -1.303108  
## sample estimates:  
## mean in group female mean in group male   
## 22.22893 23.94091   
##   
##   
## $T4.l  
##   
## Welch Two Sample t-test  
##   
## data: x by sex  
## t = -8.6907, df = 118.57, p-value = 2.364e-14  
## alternative hypothesis: true difference in means is not equal to 0  
## 95 percent confidence interval:  
## -1.971495 -1.239804  
## sample estimates:  
## mean in group female mean in group male   
## 21.13071 22.73636   
##   
##   
## $T5.l  
##   
## Welch Two Sample t-test  
##   
## data: x by sex  
## t = -9.4395, df = 119.98, p-value = 3.73e-16  
## alternative hypothesis: true difference in means is not equal to 0  
## 95 percent confidence interval:  
## -1.890261 -1.234783  
## sample estimates:  
## mean in group female mean in group male   
## 19.67536 21.23788   
##   
##   
## $BAB.l  
##   
## Welch Two Sample t-test  
##   
## data: x by sex  
## t = -7.2405, df = 97.265, p-value = 1.05e-10  
## alternative hypothesis: true difference in means is not equal to 0  
## 95 percent confidence interval:  
## -1.0089012 -0.5748001  
## sample estimates:  
## mean in group female mean in group male   
## 8.871786 9.663636   
##   
##   
## $BAH.l  
##   
## Welch Two Sample t-test  
##   
## data: x by sex  
## t = -6.5872, df = 116.27, p-value = 1.364e-09  
## alternative hypothesis: true difference in means is not equal to 0  
## 95 percent confidence interval:  
## -0.8288528 -0.4456494  
## sample estimates:  
## mean in group female mean in group male   
## 5.052143 5.689394   
##   
##   
## $Index.l  
##   
## Welch Two Sample t-test  
##   
## data: x by sex  
## t = -1.6382, df = 109.25, p-value = 0.1043  
## alternative hypothesis: true difference in means is not equal to 0  
## 95 percent confidence interval:  
## -3.9737259 0.3772973  
## sample estimates:  
## mean in group female mean in group male   
## 57.15679 58.95500   
##   
##   
## $T1.r  
##   
## Welch Two Sample t-test  
##   
## data: x by sex  
## t = -6.6602, df = 113.97, p-value = 1.01e-09  
## alternative hypothesis: true difference in means is not equal to 0  
## 95 percent confidence interval:  
## -1.912276 -1.035494  
## sample estimates:  
## mean in group female mean in group male   
## 23.45036 24.92424   
##   
##   
## $T2.r  
##   
## Welch Two Sample t-test  
##   
## data: x by sex  
## t = -6.5279, df = 119.44, p-value = 1.696e-09  
## alternative hypothesis: true difference in means is not equal to 0  
## 95 percent confidence interval:  
## -1.974537 -1.055484  
## sample estimates:  
## mean in group female mean in group male   
## 23.07893 24.59394   
##   
##   
## $T3.r  
##   
## Welch Two Sample t-test  
##   
## data: x by sex  
## t = -6.5556, df = 119.1, p-value = 1.491e-09  
## alternative hypothesis: true difference in means is not equal to 0  
## 95 percent confidence interval:  
## -1.879957 -1.007748  
## sample estimates:  
## mean in group female mean in group male   
## 22.24857 23.69242   
##   
##   
## $T4.r  
##   
## Welch Two Sample t-test  
##   
## data: x by sex  
## t = -6.4709, df = 115.96, p-value = 2.423e-09  
## alternative hypothesis: true difference in means is not equal to 0  
## 95 percent confidence interval:  
## -1.7288214 -0.9185162  
## sample estimates:  
## mean in group female mean in group male   
## 21.17179 22.49545   
##   
##   
## $T5.r  
##   
## Welch Two Sample t-test  
##   
## data: x by sex  
## t = -6.1192, df = 115.13, p-value = 1.333e-08  
## alternative hypothesis: true difference in means is not equal to 0  
## 95 percent confidence interval:  
## -1.8911753 -0.9662273  
## sample estimates:  
## mean in group female mean in group male   
## 19.49857 20.92727   
##   
##   
## $BAB.r  
##   
## Welch Two Sample t-test  
##   
## data: x by sex  
## t = -6.9999, df = 112.87, p-value = 1.945e-10  
## alternative hypothesis: true difference in means is not equal to 0  
## 95 percent confidence interval:  
## -0.8841950 -0.4940951  
## sample estimates:  
## mean in group female mean in group male   
## 8.939643 9.628788   
##   
##   
## $BAH.r  
##   
## Welch Two Sample t-test  
##   
## data: x by sex  
## t = -5.8434, df = 111.73, p-value = 5.137e-08  
## alternative hypothesis: true difference in means is not equal to 0  
## 95 percent confidence interval:  
## -0.7158761 -0.3533230  
## sample estimates:  
## mean in group female mean in group male   
## 5.089643 5.624242   
##   
##   
## $Index.r  
##   
## Welch Two Sample t-test  
##   
## data: x by sex  
## t = -1.6489, df = 111.71, p-value = 0.102  
## alternative hypothesis: true difference in means is not equal to 0  
## 95 percent confidence interval:  
## -3.3084032 0.3030569  
## sample estimates:  
## mean in group female mean in group male   
## 56.98036 58.48303

Extract the p-values for the t-tests:

sapply(foot.sex.t, FUN = function(x) x$p.value)

## T1.l T2.l T3.l T4.l T5.l   
## 1.442207e-12 1.348046e-12 1.901345e-13 2.363915e-14 3.730286e-16   
## BAB.l BAH.l Index.l T1.r T2.r   
## 1.050028e-10 1.363798e-09 1.042582e-01 1.009614e-09 1.696248e-09   
## T3.r T4.r T5.r BAB.r BAH.r   
## 1.491302e-09 2.423427e-09 1.333400e-08 1.945204e-10 5.136988e-08   
## Index.r   
## 1.019810e-01