PS: Indpendent and Dependent t-test for TAs

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R practice.

Install packages.

```
library(car)
## Loading required package: carData
library(ggplot2)
library(dplyr)
## Attaching package: 'dplyr'
## The following object is masked from 'package:car':
##
##
       recode
## The following objects are masked from 'package:stats':
##
##
       filter, lag
## The following objects are masked from 'package:base':
##
##
       intersect, setdiff, setequal, union
```

Load the Bird Migration data into R.

```
##
     Birds
                Species Gender Treatment value
          1 Hylocichla
                              M Pre_weight
## 2
          2 Hylocichla
                              M Pre_weight
                                                 44
         3 Hylocichla
4 Hylocichla
5 Hylocichla
                              M Pre_weight
                                                 48
                              M Pre_weight
                                                 42
          5 Hylocichla
## 5
                              M Pre_weight
                                                 51
          6 Hylocichla
                              {\tt M\ Pre\_weight}
                                                 51
## 6
```

Make a subgroup of Weight before migration (Pre_weight).

```
df_bird_Pre_weight <- df_bird_all %>%
  filter(Treatment == 'Pre_weight')
summary(df_bird_Pre_weight)
```

```
##
       Birds
                    Species
                                      Gender
                                                      {\tt Treatment}
## Min. : 1.00
                  Length:28
                                   Length:28
                                                     Length:28
## 1st Qu.: 7.75 Class :character
                                   Class :character
                                                     Class :character
## Median :14.50 Mode :character
                                   Mode :character
                                                     Mode :character
## Mean :14.50
   3rd Qu.:21.25
##
## Max. :28.00
##
       value
## Min. :41.00
## 1st Qu.:46.75
## Median :49.00
## Mean :48.68
## 3rd Qu.:52.00
## Max. :55.00
```

Part I.

2. Analyze data distribution. Is your pre-weight data normally distributed?

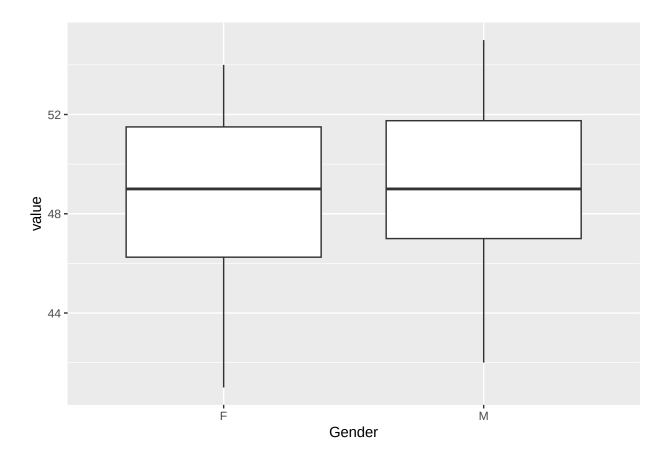
```
shapiro.test(df_bird_Pre_weight$value)
##
##
  Shapiro-Wilk normality test
##
## data: df_bird_Pre_weight$value
## W = 0.95557, p-value = 0.2725
3. Test to see if you meet the assumption of equal variance for this test. Based on the results
of this test for unequal variance, which test should you use?
# Perform Levene's test
result <- leveneTest(value~Gender, data=df_bird_Pre_weight)</pre>
## Warning in leveneTest.default(y = y, group = group, \dots): group coerced to
## factor.
print(result)
## Levene's Test for Homogeneity of Variance (center = median)
        Df F value Pr(>F)
## group 1 0.0061 0.9381
         26
##
p value <- result$Pr[1]</pre>
# Significance level
alpha <- 0.05
# Interpret the result
if (p_value < alpha) {</pre>
 cat("Reject null hypothesis: Variances are significantly different.\n")
} else {
  cat("Fail to reject null hypothesis: Variances are not significantly different.\n")
```

Fail to reject null hypothesis: Variances are not significantly different.

4. Run this analysis. What is the test statistic (obtained) value for the test you selected? Use alternative to determine your hypothesis. Default= "two.sided". alternative = c("two.sided", "less", "greater"),

```
# Perform t-test
t_test_result <- t.test(value~Gender,alternative = "two.sided",</pre>
                         var.equal=TRUE, data=df_bird_Pre_weight)
print(t_test_result)
## Two Sample t-test
##
## data: value by Gender
## t = -0.78654, df = 26, p-value = 0.4387
## alternative hypothesis: true difference in means between group F and group M is not equal to O
## 95 percent confidence interval:
## -4.387692 1.959120
## sample estimates:
## mean in group F mean in group M
##
          48.07143
                          49.28571
```

```
# Create a boxplot plot using ggplot2
ggplot(df_bird_Pre_weight, aes(x = Gender, y = value)) +
geom_boxplot() # Add jitter for better visualization
```



Part II.

Analyze data distribution. Is your pre-weight data normally distributed?

```
shapiro.test(df_bird_Pre_weight$value)
##
##
  Shapiro-Wilk normality test
## data: df_bird_Pre_weight$value
## W = 0.95557, p-value = 0.2725
Test to see if you meet the assumption of equal variance for this test. Based on the results of
this test for unequal variance, which test should you use?
# Perform Levene's test
result <- leveneTest(value~Species, data=df_bird_Pre_weight)</pre>
## Warning in leveneTest.default(y = y, group = group, \dots): group coerced to
## factor.
print(result)
## Levene's Test for Homogeneity of Variance (center = median)
        Df F value Pr(>F)
## group 1 1.2128 0.2809
         26
##
p value <- result$Pr[1]</pre>
# Significance level
alpha <- 0.05
# Interpret the result
if (p_value < alpha) {</pre>
 cat("Reject null hypothesis: Variances are significantly different.\n")
  cat("Fail to reject null hypothesis: Variances are not significantly different.\n")
```

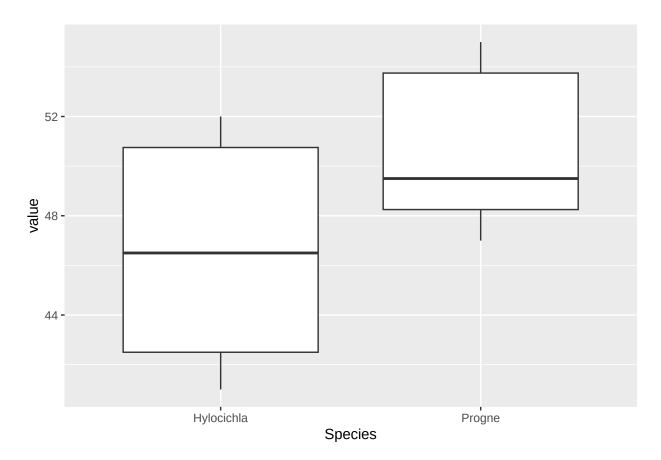
Fail to reject null hypothesis: Variances are not significantly different.

8. Run this analysis. What is the test statistic (obtained) value for the test you selected? Use alternative to determine your hypothesis. Default= "two.sided". alternative = c("two.sided", "less", "greater"),

```
## data: value by Species
## t = -2.891, df = 26, p-value = 0.007655
## alternative hypothesis: true difference in means between group Hylocichla and group Progne is not eq
## 95 percent confidence interval:
## -6.721783 -1.135360
## sample estimates:
## mean in group Hylocichla mean in group Progne
## 46.71429 50.64286
```

Perform t-test

```
# Create a boxplot plot using ggplot2
ggplot(df_bird_Pre_weight, aes(x = Species, y = value)) +
  geom_boxplot() # Add jitter for better visualization
```



Part III.

12. Analyze data distribution. Is your pre-weight data normally distributed?

```
##
## Shapiro-Wilk normality test
##
## data: df_bird_all$value
## W = 0.97528, p-value = 0.3028
```

Test to see if you meet the assumption of equal variance for this test. Based on the results of this test for unequal variance, which test should you use?

```
# Perform Levene's test
result_paired_t_test <- leveneTest(value~Treatment, data=df_bird_all)

## Warning in leveneTest.default(y = y, group = group, ...): group coerced to
## factor.

print(result_paired_t_test)

## Levene's Test for Homogeneity of Variance (center = median)
## Df F value Pr(>F)
## group 1 0.0035 0.9529
## 54
```

```
p_value <- result_paired_t_test$Pr[1]
# Significance level
alpha <- 0.05

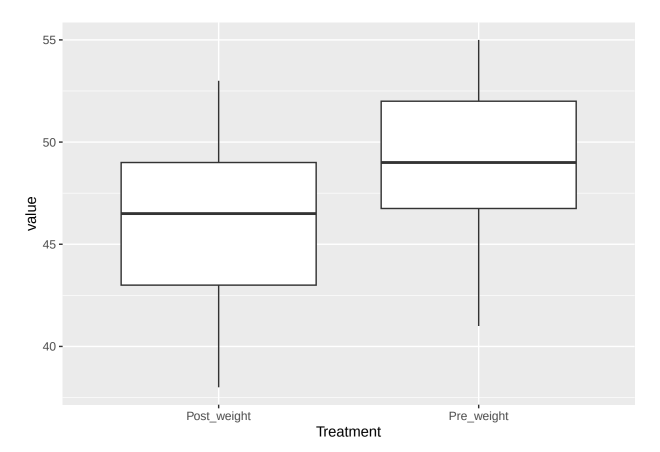
# Interpret the result
if (p_value < alpha) {
   cat("Reject null hypothesis: Variances are significantly different.\n")
} else {
   cat("Fail to reject null hypothesis: Variances are not significantly different.\n")
}</pre>
```

Fail to reject null hypothesis: Variances are not significantly different.

Perform and Print the paired t-test result.

```
##
## Paired t-test
##
## data: value by Treatment
## t = -3.4184, df = 27, p-value = 0.002013
## alternative hypothesis: true difference in means is not equal to 0
## 95 percent confidence interval:
## -4.400616 -1.099384
## sample estimates:
## mean of the differences
## -2.75
```

```
# Create a boxplot plot using ggplot2
ggplot(df_bird_all, aes(x = Treatment, y = value)) +
geom_boxplot() # Add jitter for better visualization
```



Part IV.

From the data.frame make two groups (females and males)

```
df_bird_female <- df_bird_all %>%
filter(Gender == 'F')
```

```
df_bird_male <- df_bird_all %>%
  filter(Gender == 'M')
```

Assumptions for Females

```
##
## Shapiro.test(df_bird_female$value)

##
## Shapiro-Wilk normality test
##
## data: df_bird_female$value
## W = 0.95617, p-value = 0.282

leveneTest(value~Treatment, data=df_bird_female)

## Warning in leveneTest.default(y = y, group = group, ...): group coerced to
## factor.

## Levene's Test for Homogeneity of Variance (center = median)
## Df F value Pr(>F)
## group 1 0.0053 0.9424
## 26
```

Perform and Print the paired t-test result

Assumptions for Males

```
shapiro.test(df_bird_male$value)
##
## Shapiro-Wilk normality test
##
## data: df_bird_male$value
## W = 0.97278, p-value = 0.6567
leveneTest(value~Treatment, data=df_bird_male)
## Warning in leveneTest.default(y = y, group = group, ...): group coerced to
## factor.
## Levene's Test for Homogeneity of Variance (center = median)
       Df F value Pr(>F)
## group 1 0.334 0.5683
##
         26
Perform and Print the paired t-test result
t_paired_male <- t.test(value ~ Treatment, paired = TRUE,</pre>
                        alternative= 'less',
                        data = df_bird_male)
print(t_paired_male)
##
## Paired t-test
##
## data: value by Treatment
## t = -3.1601, df = 13, p-value = 0.003762
\#\# alternative hypothesis: true difference in means is less than 0
## 95 percent confidence interval:
##
        -Inf -1.946804
## sample estimates:
## mean of the differences
                 -4.428571
```