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
                                        Gender
                    Species
                                                        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?

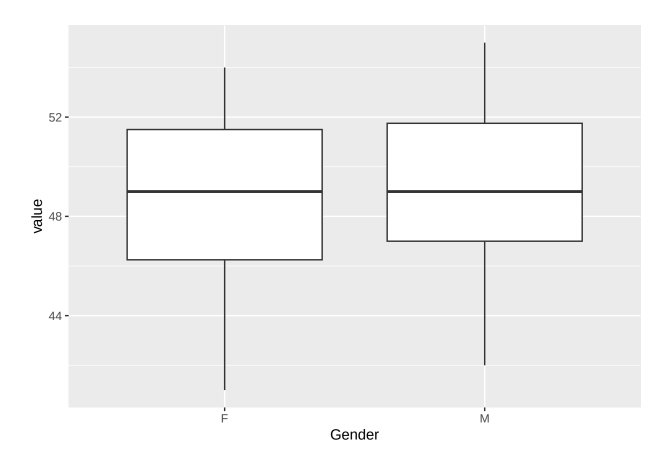
```
##
## Shapiro-Wilk normality test
##
## data: df_bird_Pre_weight$value
## W = 0.95557, p-value = 0.2725
```

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)
## Warning in leveneTest.default(y = y, group = group, ...): group coerced to
## factor.</pre>
```

```
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 0
## 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



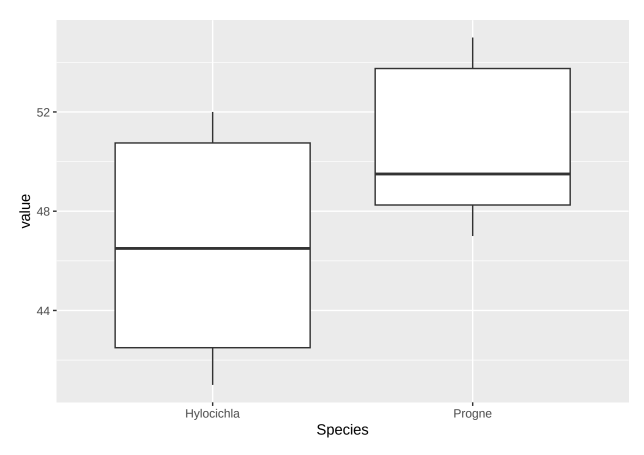
Part II. Analyze data distribution. Is your pre-weight data normally distributed?

```
##
## Shapiro-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, ...): 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 \leftarrow 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.
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"),
# Perform t-test
t_test_result <- t.test(value~Species, alternative = "two.sided",</pre>
                          var.equal=TRUE, data=df_bird_Pre_weight)
print(t_test_result)
##
## Two Sample t-test
##
## 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
                                              50.64286
##
                   46.71429
```

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



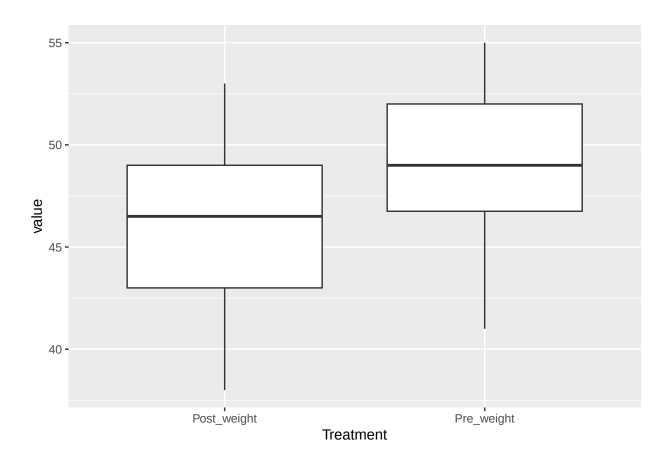
Part III. 12. Analyze data distribution. Is your pre-weight data normally distributed?

```
shapiro.test(df_bird_all$value)

##
## 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)</pre>
## 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]</pre>
# Significance level
alpha \leftarrow 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.
Perform and Print the paired t-test result.
t_paired_result <- t.test(value ~ Treatment, paired = TRUE,</pre>
                          data = df_bird_all)
print(t_paired_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$value
## W = 0.95617, p-value = 0.282
leveneTest(value~Treatment, data=df_bird_female)
## Warning in leveneTest.default(y = y, group = group, ...): group coerced to
## factor.
## Levene's Test for Homogeneity of Variance (center = median)
        Df F value Pr(>F)
## group 1 0.0053 0.9424
         26
##
Perform and Print the paired t-test result
t_paired_female <- t.test(value ~ Treatment, paired = TRUE,</pre>
                          alternative= 'less',
                          data = df_bird_female)
print(t_paired_female)
##
## Paired t-test
## data: value by Treatment
## t = -1.9868, df = 13, p-value = 0.03422
## alternative hypothesis: true difference in means is less than 0
## 95 percent confidence interval:
##
         -Inf -0.1164104
## sample estimates:
## mean of the differences
##
                 -1.071429
Assumptions for Males
shapiro.test(df_bird_male$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

```
##
## 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
```