## Week 6 Assignment

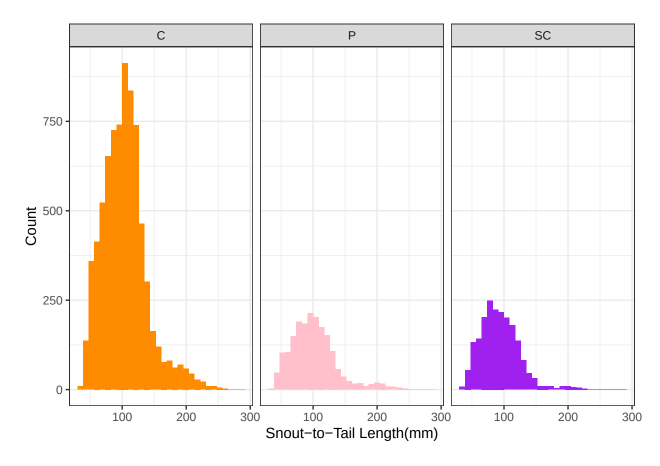
## Jessalyn Chuang

```
# Load necessary packages.
library(here)
## here() starts at /home/guest/Statistical_Modeling_Sp25
library(tidyverse)
## -- Attaching core tidyverse packages ----- tidyverse 2.0.0 --
## v dplyr
          1.1.4
                       v readr
                                   2.1.5
## v forcats 1.0.0
                                   1.5.1
                       v stringr
## v ggplot2 3.5.1
                       v tibble
                                   3.2.1
## v lubridate 1.9.3
                       v tidyr
                                   1.3.1
## v purrr
              1.0.2
## -- Conflicts ----- tidyverse_conflicts() --
## x dplyr::filter() masks stats::filter()
                   masks stats::lag()
## x dplyr::lag()
## i Use the conflicted package (<a href="http://conflicted.r-lib.org/">http://conflicted.r-lib.org/</a>) to force all conflicts to become error
library(moments)
library(dplyr)
```

#(1) Coastal Giant Salamander- a. Filter the dataset only for coastal giant salamanders in cascades (C), pools (P), and side channels (SC). Create a figure that helps you to evaluate whether their snout-to-tail length (length\_2\_mm) by habitat type (unittype) is normally distributed. You may also calculate skew and kurtosis values to help with your decision-making. Are these data normally distributed? Why or why not? If they are not, apply a log-transform (log10()) to the length data and re-evaluate if the data now meets the criteria to be considered normally-distributed.

```
theme(legend.position = "none")
salamander_fig1
```

## 'stat\_bin()' using 'bins = 30'. Pick better value with 'binwidth'.
## Warning: Removed 220 rows containing non-finite outside the scale range
## ('stat\_bin()').



```
# Separate data by unittype
C_dat <- salamander_data %>%
filter(unittype == "C")

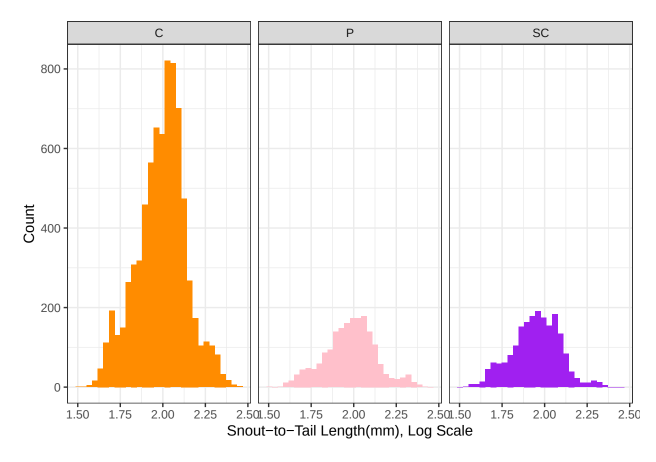
P_dat <- salamander_data %>%
filter(unittype == "P")

SC_dat <- salamander_data %>%
filter(unittype == "SC")

# Calculate skew.
skewness(C_dat$length_2_mm, na.rm = TRUE)
```

## [1] 0.9428883

```
skewness(P_dat$length_2_mm, na.rm = TRUE)
## [1] 1.106651
skewness(SC_dat$length_2_mm, na.rm = TRUE)
## [1] 0.9912964
# Calculate kurtosis.
kurtosis(C_dat$length_2_mm, na.rm = TRUE)
## [1] 4.691787
kurtosis(P_dat$length_2_mm, na.rm = TRUE)
## [1] 4.894066
kurtosis(SC_dat$length_2_mm, na.rm = TRUE)
## [1] 4.937146
Skew results: Cascades: 0.94 Pool: 1.11 Side Channels: 0.99
Kurtosis results: Cascades: 4.69 Pool: 4.89 Side Channels: 4.94
Visually, the three data from the three habitats all seem to be positive skewed, with the majority of points
concentrated at lower values. From the skew results, all three have values that suggest positive skewing as
they are all greater than 0. Lastly for kurtosis, all three have values greater than 3, suggesting that the data
is leptokurtic. Therefore, this data is not normally distributed!
#log-transformation of data
salamander_data_transformed <- mutate(salamander_data, length_2_mm = log10(length_2_mm))</pre>
salamander_fig2 <- ggplot(salamander_data_transformed, aes(x = length_2_mm, fill = unittype)) +</pre>
  geom_histogram() +
  scale_fill_manual(values = c("darkorange", "pink", "purple")) +
  labs(x = "Snout-to-Tail Length(mm), Log Scale", y = "Count") +
  facet_grid(.~unittype) +
  theme_bw() +
  theme(legend.position = "none")
salamander_fig2
## 'stat_bin()' using 'bins = 30'. Pick better value with 'binwidth'.
## Warning: Removed 220 rows containing non-finite outside the scale range
## ('stat_bin()').
```



```
# Separate data by unittype
C_dat_trans <- salamander_data_transformed %>%
filter(unittype == "C")

P_dat_trans <- salamander_data_transformed %>%
filter(unittype == "P")

SC_dat_trans <- salamander_data_transformed %>%
filter(unittype == "SC")

# Calculate skew.
skewness(C_dat_trans$length_2_mm, na.rm = TRUE)
```

## [1] -0.1713156

```
skewness(P_dat_trans$length_2_mm, na.rm = TRUE)
```

## [1] -0.04432009

```
skewness(SC_dat_trans$length_2_mm, na.rm = TRUE)
```

## [1] -0.07959806

```
# Calculate kurtosis.
kurtosis(C_dat_trans$length_2_mm, na.rm = TRUE)
## [1] 3.08717
kurtosis(P_dat_trans$length_2_mm, na.rm = TRUE)
## [1] 3.083047
kurtosis(SC_dat_trans$length_2_mm, na.rm = TRUE)
```

## [1] 2.981829

Now, all three habitat types appear to be normally distributed from the visual inspection. Their skewness values are now all around 0 and kurtosis values are all around 3 (mesokurtic).

b. If you found the data from part a were not normally distributed and a log-transform did not change this finding, you may stop here and proceed to question 2. If you found the data from part a were normally distributed, conduct a Bartlett's test for equal variance to determine if these data also satisfy the need for homogeneity of variances across groups. Do these data have approximately equal variances? Why or why not? Remember, the data may not pass the Bartlett's test, but if they adhere to the rule of thumb mentioned above, you may proceed with a one-way ANOVA.

```
# Perform Bartlett test.
salamander_var <- bartlett.test(salamander_data_transformed$length_2_mm,</pre>
                                 salamander_data_transformed$unittype)
# Examine results.
salamander_var
##
```

Bartlett test of homogeneity of variances ## ## data: salamander\_data\_transformed\$length\_2\_mm and salamander\_data\_transformed\$unittype ## Bartlett's K-squared = 11.213, df = 2, p-value = 0.003674

Since p-value is less than 0.05, I reject the null hypothesis stating that the variance between all groups included in the data are the same in favor of the alternate hypothesis that states that the variance between all groups included in the data are not the same. Thus, this data does not pass the Bartlett's test. However, if the largest sample variance is less than 4 times the smallest sample variance, we may still assume variances are equal across samples and conduct an ANOVA. Checking this:

```
C_var <- var(C_dat_trans$length_2_mm, na.rm = TRUE)</pre>
P_var <- var(P_dat_trans$length_2_mm, na.rm = TRUE)
SC_var <- var(SC_dat_trans$length_2_mm, na.rm = TRUE)
C_var
```

## [1] 0.0208925

##

```
P_var

## [1] 0.02270146

SC_var

## [1] 0.01949329
```

Variance results: Cascades: 0.02 Pool: 0.02 Side Channels: 0.02

When finding their variances, they actually all turned out equal, so we may assume variances are approximately equal across samples and conduct an ANOVA.

c. If you found the data from part b did not display approximately equal variances, you may stop here. If you found the data from part b did display approximately equal variances, conduct a one-way ANOVA to see if there is a significant difference between coastal giant salamander lengths across cascades, pools, and side channels of Mack Creek. If you find evidence of significant differences, perform a post-hoc Tukey's HSD test to determine which are significant from which habitats. Communicate your findings as a figure (with an appropriate caption) and in a sentence, as it might appear in a final report.

```
salamander_ANOVA <- aov(length_2_mm ~ unittype, data = salamander_data_transformed)</pre>
summary(salamander_ANOVA)
                  Df Sum Sq Mean Sq F value Pr(>F)
##
## unittype
                   2
                        3.2 1.5993
                                      76.35 <2e-16 ***
                      239.0 0.0209
## Residuals
               11411
## ---
## Signif. codes: 0 '*** 0.001 '** 0.01 '* 0.05 '.' 0.1 ' 1
## 220 observations deleted due to missingness
#Perform a post-hoc Tukey's HSD test since p < 0.001
salamander_Tukey <- TukeyHSD(salamander_ANOVA)</pre>
salamander_Tukey
##
     Tukey multiple comparisons of means
##
       95% family-wise confidence level
##
## Fit: aov(formula = length_2_mm ~ unittype, data = salamander_data_transformed)
##
## $unittype
##
               diff
                            lwr
                                         upr
                                                  p adj
## P-C -0.01310767 -0.02187368 -0.004341656 0.0013319
## SC-C -0.04502145 -0.05359132 -0.036451579 0.0000000
## SC-P -0.03191378 -0.04286293 -0.020964629 0.0000000
salamander_summary <- salamander_data %>%
  group_by(unittype) %>%
  summarize(mean = mean(length_2_mm, na.rm = TRUE),
  sd = sd(length_2_mm, na.rm = TRUE)) %>%
  ungroup()
```

salamander\_summary

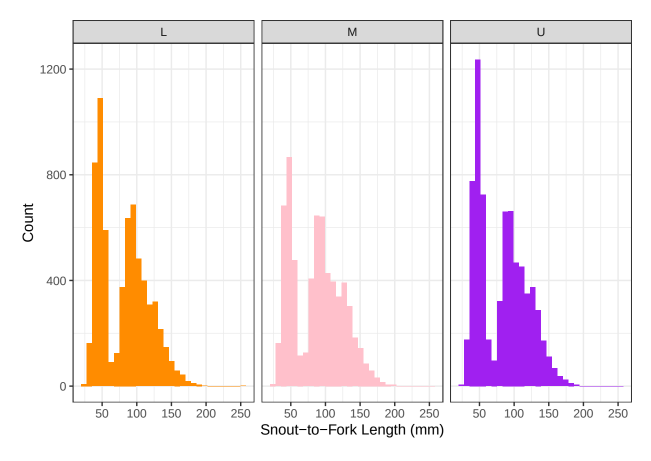
```
## # A tibble: 3 x 3
##
     unittype mean
                        sd
##
     <fct>
               <dbl> <dbl>
## 1 C
               104.
                      34.7
## 2 P
               101.
                      36.1
## 3 SC
               93.4
                      30.5
```

Salamander species displayed significant differences in Snout-to-Tail length as determined by one-way ANOVA (F(2, 11411) = 76.35, p < 0.001). Post-hoc testing by Tukey's HSD revealed that mean Snout-to-Tail length for salamanders in the Cascades habitat (mean = 104 mm, s.d. = 34.7 mm), Pool habitat (mean = 101 mm, s.d. = 36.1 mm), and Side Channels habitat (mean = 93.4 mm, s.d. = 30.5 mm) all differed significantly from each other.

## #(2) Cutthroat Trout-

a. Filter the dataset only for cutthroat trout and create a figure that helps you to evaluate whether their snout-to-fork length (length\_1\_mm) by reach (reach) is normally distributed. You may also calculate skew and kurtosis values to help with your decision-making. Are these data normally distributed? Why or why not? If they are not, apply a log-transform (log10()) to the length data and re-evaluate if the data now meets the criteria to be considered normally- distributed.

```
## 'stat_bin()' using 'bins = 30'. Pick better value with 'binwidth'.
## Warning: Removed 5 rows containing non-finite outside the scale range
## ('stat_bin()').
```



```
# Separate data by unittype
L_dat <- trout_data %>%
filter(reach == "L")

M_dat <- trout_data %>%
filter(reach == "M")

U_dat <- trout_data %>%
filter(reach == "U")

# Calculate skew.
skewness(L_dat$length_1_mm, na.rm = TRUE)
```

## [1] 0.4042481

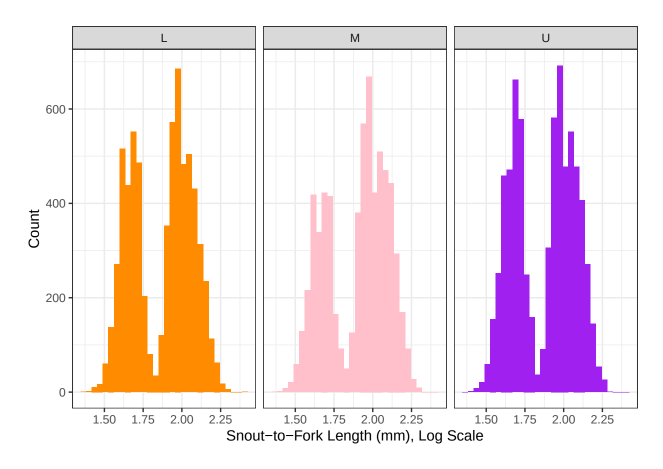
```
skewness(M_dat$length_1_mm, na.rm = TRUE)
```

## [1] 0.2626566

```
skewness(U_dat$length_1_mm, na.rm = TRUE)
```

## [1] 0.3832519

```
# Calculate kurtosis.
kurtosis(L_dat$length_1_mm, na.rm = TRUE)
## [1] 2.286412
kurtosis(M_dat$length_1_mm, na.rm = TRUE)
## [1] 2.158395
kurtosis(U_dat$length_1_mm, na.rm = TRUE)
## [1] 2.104812
Skew results: L: 0.40 M: 0.26 U: 0.38
Kurtosis results: L: 2.28 M: 2.16 U: 2.10
Visually, the three data sets appear bi-modal. From the skew results, all three have values slightly above 0
indicating slight positive skewing. All three have kurtosis values less than 3, indicating that they may be
platykurtic. Overall, they do not seem normally distributed.
#log-transformation of data
trout_data_transformed <- mutate(trout_data, length_1_mm = log10(length_1_mm))</pre>
trout_fig2 <- ggplot(trout_data_transformed, aes(x = length_1_mm, fill = reach)) +</pre>
  geom_histogram() +
  scale_fill_manual(values = c("darkorange", "pink", "purple")) +
  labs(x = "Snout-to-Fork Length (mm), Log Scale", y = "Count") +
  facet_grid(.~reach) +
  theme_bw() +
  theme(legend.position = "none")
trout_fig2
## 'stat_bin()' using 'bins = 30'. Pick better value with 'binwidth'.
## Warning: Removed 5 rows containing non-finite outside the scale range
## ('stat_bin()').
```



```
# Separate data by unittype
L_dat_trans <- trout_data_transformed %>%
filter(reach == "L")

M_dat_trans <- trout_data_transformed %>%
filter(reach == "M")

U_dat_trans <- trout_data_transformed %>%
filter(reach == "U")

# Calculate skew.
skewness(L_dat_trans$length_1_mm, na.rm = TRUE)
```

```
skewness(M_dat_trans$length_1_mm, na.rm = TRUE)
## [1] -0.3168654
skewness(U_dat_trans$length_1_mm, na.rm = TRUE)
```

## [1] -0.1169573

## [1] -0.1397749

```
# Calculate kurtosis.
kurtosis(L_dat_trans$length_1_mm, na.rm = TRUE)

## [1] 1.73097

kurtosis(M_dat_trans$length_1_mm, na.rm = TRUE)

## [1] 1.882288

kurtosis(U_dat_trans$length_1_mm, na.rm = TRUE)
```

## [1] 1.697403

After applying the log 10 transformation, the data is still bi-modal as apparent from a visual inspection. While skewness was brought closer to 0, kurtosis values are still less than 3, and so these distributions are considered platykurtic. The data are still not normally distributed.

b. If you found the data from part a were not normally distributed and a log-transform did not change this finding, you may stop here. If you found the data from part a were normally distributed, conduct a Bartlett's test for equal variance to determine if these data also satisfy the need for homogeneity of variances across groups. Do these data have approximately similar variances? Why or why not? Remember, the data may not pass the Bartlett's test, but if they adhere to the rule of thumb mentioned above, you may proceed with a one-way ANOVA.

Since the data was not normally distributed, I did not conduct a Bartlett's test for equal variance.

c. If you found the data from part b did not display equal variances, you may stop here. If you found the data from part b did display equal variances, conduct a one-way ANOVA to see if there is a significant difference between cutthroat trout lengths across lower, middle, and upper reaches of Mack Creek. If you find evidence of significant differences, perform a post- hoc Tukey's HSD test to determine which are significant from which reaches. Communicate your findings as a figure (with an appropriate caption) and a sentence, as it might appear in a final report.

Since the data was not normally distributed, I did not conduct a one-way ANOVA.