# Blunt-nosed Leopard Lizard CEWL Measurement Triplicate Tests

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# Packages

```
if (!require("tidyverse")) install.packages("tidyverse")
library("tidyverse")
if (!require("UsingR")) install.packages("UsingR")
library("UsingR")
if (!require("MASS")) install.packages("MASS")
library("MASS")
if (!require("lme4")) install.packages("lme4")
library("lme4")
```

### Data Info

This data was collected during fieldwork with Blunt-nosed Leopard Lizards ( $Gambelia\ sila$ ) in the Elkhorn Plain of the Carrizo Plain National Monument in California, USA during the 2021 active season (April-August).

This is a preliminary analysis to decide whether CEWL measurements should be taken in triplicate or can be just one measurement.

## Load CEWL Data

Only the first weekend is loaded for now, but we can check with other replicates later.

```
# weekend 1
CEWL_April_23_25 <- read.csv("./data/BNLL CEWL 4-23-2021 to 4-25-2021.csv") %>%
  dplyr::mutate(weekend = "1") # add weekend index for throughout season
# weekend 2
# etc etc
# merge all CEWL datafiles & reformat
CEWL <- CEWL April 23 25 %>%
  # join
  #rbind(., CEWL_May_xx, CEWL_May_xx, etc
          ) %>%
  # reformat data
  dplyr::select(Date, Time,
                ID = Comments,
                TEWL_g_m2h = TEWL..g..m2h.., # rename
                CV = CV...., # rename
                SSWL_g_m2 = SSWL..g..m2.., # rename
                ambient_temp_C = AmbT..C., # rename
                ambient RH percent = AmbRH...., # rename
                weekend
                ) %>%
  dplyr::mutate(date_time = as.POSIXct(paste(Date, Time), format = "%m/%d/%y %I:%M:%S %p"),
                # date only
                Date = as.Date(Date, format = "%m/%d/%y"),
                # time with arbitrary date, need to figure out how to remove
                Time = as.POSIXct(Time, format = "%I:%M:%S %p"),
                # get length of current ID for editing
                length_ID = nchar((ID)),
                # extract CEWL replicate number
                replicate = as.numeric(substring(ID, length_ID, length_ID)),
                # extract group (male, female, widespread)
                ID_group = substring(ID, 1, 1),
                # extract ID number within group
                ID_number = substring(ID, 2, (length_ID-2)),
                # merge IDs to match format in other data files
                ID = paste(ID group, "-", ID number, sep = ""),
                weekend = as.numeric(weekend)
                ) %>%
  dplyr::select(-length_ID, # remove these since no longer need
                -ID group,
                -ID_number
summary(CEWL)
##
         Date
                              Time
                                                             ID
                                                        Length: 198
## Min.
          :2021-04-23
                        Min.
                                :2021-05-06 11:37:51
```

```
## Date Time ID

## Min. :2021-04-23 Min. :2021-05-06 11:37:51 Length:198

## 1st Qu.:2021-04-23 1st Qu.:2021-05-06 13:53:22 Class :character

## Median :2021-04-24 Median :2021-05-06 14:38:46 Mode :character

## Mean :2021-04-23 Mean :2021-05-06 15:01:37

## 3rd Qu.:2021-04-24 3rd Qu.:2021-05-06 15:58:40
```

```
:2021-05-06 19:56:00
##
   Max.
          :2021-04-25
                       Max.
##
     TEWL_g_m2h
                        CV
                                    SSWL_g_m2
                                                    ambient_temp_C
                   Min. : 0.1000 Min. :0.00000 Min.
## Min. : 0.140
                                                           :18.90
  1st Qu.: 8.213
                   1st Qu.: 0.4125
                                   1st Qu.:0.00900
                                                    1st Qu.:28.52
## Median :10.615
                   Median : 0.5900
                                   Median: 0.02439 Median: 30.50
## Mean
         :11.650
                   Mean
                        : 1.0383
                                  Mean
                                         :0.03701 Mean
                                                           :29.07
## 3rd Qu.:14.150
                   3rd Qu.: 0.8175
                                    3rd Qu.:0.05147
                                                    3rd Qu.:31.60
## Max.
          :65.310 Max.
                         :42.8500
                                   Max.
                                          :0.18780 Max.
                                                           :33.70
## ambient_RH_percent
                        weekend
                                  date time
                                                              replicate
## Min.
                                      :2021-04-23 12:57:55
                                                                   :1.000
         :11.60
                   Min. :1 Min.
                                                            Min.
## 1st Qu.:15.00
                    1st Qu.:1 1st Qu.:2021-04-23 14:55:45
                                                            1st Qu.:1.000
## Median :15.90
                     Median :1 Median :2021-04-24 12:57:24
                                                            Median :2.000
## Mean
          :19.31
                     Mean :1 Mean
                                      :2021-04-24 07:23:26
                                                            Mean
                                                                  :2.015
## 3rd Qu.:20.98
                     3rd Qu.:1
                                3rd Qu.:2021-04-24 15:36:36
                                                            3rd Qu.:3.000
## Max.
          :37.90
                     Max. :1 Max.
                                      :2021-04-25 14:41:17
                                                            Max.
                                                                  :4.000
```

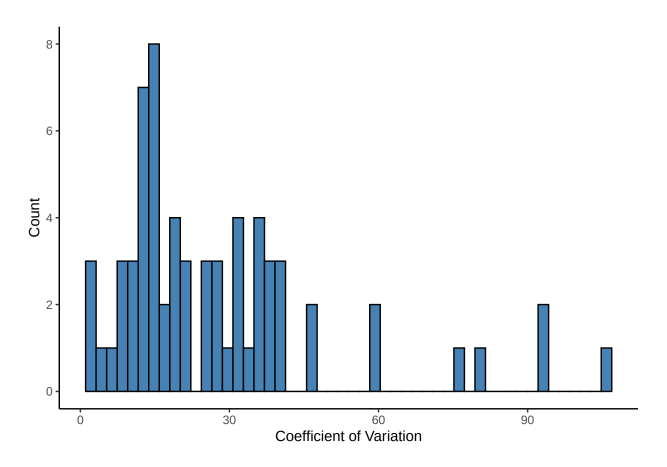
### **Basic Calculations**

Get mean, sd, and CV values for each lizard's CEWL:

## `summarise()` regrouping output by 'Date' (override with `.groups` argument)

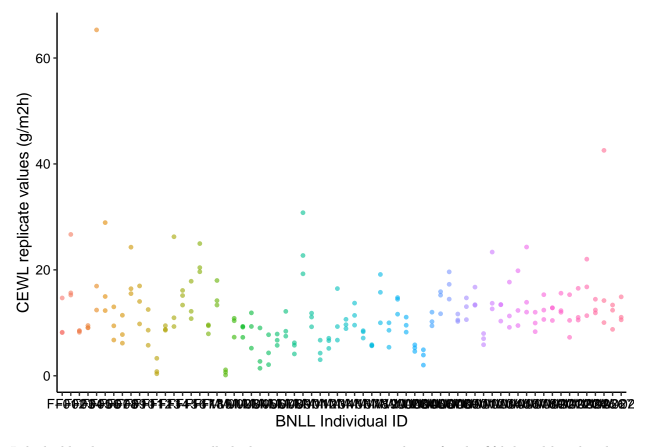
# Plot CV Histogram

```
CEWL_rep_avg %>%
  ggplot(., aes(x = TEWL_CV)) +
  geom_histogram(color = "black", fill="steelblue", bins=50) +
  theme_classic() +
  xlab("Coefficient of Variation") +
  ylab("Count")
```



# Plot Replicates

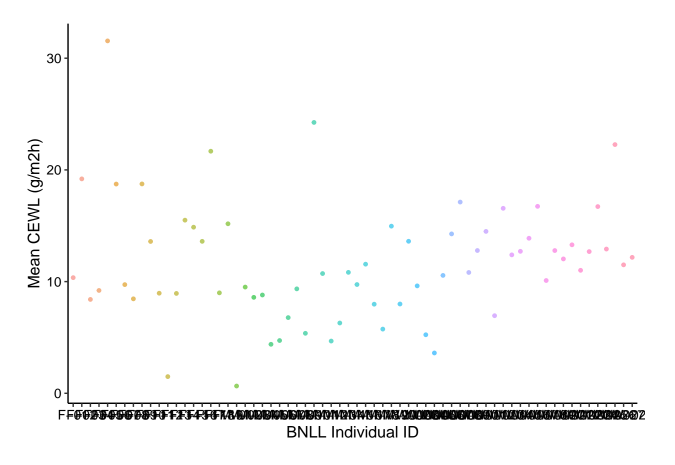
```
CEWL %>%
  ggplot(data = .) +
  geom_point(aes(x = ID,
                 y = TEWL_g_m2h,
                 color = ID
                 ),
             size = 1,
             alpha = 0.6) +
  theme_classic() +
  xlab("BNLL Individual ID") +
  ylab("CEWL replicate values (g/m2h)") +
  theme(text = element_text(color = "black",
                            family = "sans",
                            size = 12),
        axis.text = element_text(color = "black",
                                 family = "sans",
                                 size = 10),
        legend.text.align = 0,
        legend.position = "none"
)
```



It looks like there are two unusually high measurements - one replicate for the fifth lizard listed and one replicate for the third-to-last lizard listed. I think they're both above 40, so I could see how filtering them out affects replicability.

Plot means:

```
CEWL_rep_avg %>%
  ggplot(data = .) +
  geom_point(aes(x = ID,
                 y = mean_TEWL_g_m2h,
                 color = ID
                 ),
             size = 1,
             alpha = 0.6) +
  theme_classic() +
  xlab("BNLL Individual ID") +
  ylab("Mean CEWL (g/m2h)") +
  theme(text = element_text(color = "black",
                            family = "sans",
                            size = 12),
        axis.text = element_text(color = "black",
                                  family = "sans",
                                  size = 10),
        legend.text.align = 0,
        legend.position = "none"
```



## **Model Replicates**

make a GLMM to figure out whether replicates have significantly different CEWL values:

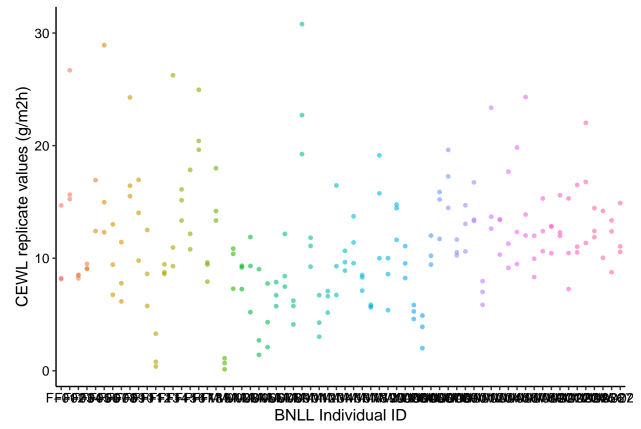
```
mod <- lme4::lmer(data = CEWL, TEWL_g_m2h ~ replicate + (1|ID))</pre>
mod2 <- nlme::lme(data = CEWL, TEWL_g_m2h ~ replicate, random = ~ 1|ID) # for p-values</pre>
summary(mod)
## Linear mixed model fit by REML ['lmerMod']
## Formula: TEWL_g_m2h ~ replicate + (1 | ID)
      Data: CEWL
##
##
  REML criterion at convergence: 1270.7
##
## Scaled residuals:
##
       Min
                1Q Median
  -2.3402 -0.3939 -0.0861 0.2671
                                    7.7263
##
## Random effects:
##
    Groups
             Name
                         Variance Std.Dev.
##
             (Intercept) 22.56
                                   4.750
                          23.09
                                   4.805
    Residual
## Number of obs: 198, groups: ID, 66
##
## Fixed effects:
               Estimate Std. Error t value
                              1.079 15.028
## (Intercept) 16.217
```

```
-2.267
## replicate
                        0.417 -5.436
##
## Correlation of Fixed Effects:
##
            (Intr)
## replicate -0.779
summary(mod2)
## Linear mixed-effects model fit by REML
##
    Data: CEWL
##
         AIC
                  BIC
                         logLik
    1278.673 1291.785 -635.3363
##
##
## Random effects:
## Formula: ~1 | ID
##
           (Intercept) Residual
## StdDev:
             4.749645 4.804734
##
## Fixed effects: TEWL_g_m2h ~ replicate
                 Value Std.Error DF
                                      t-value p-value
## (Intercept) 16.21740 1.0791558 131 15.027857
## replicate -2.26668 0.4170131 131 -5.435512
## Correlation:
##
             (Intr)
## replicate -0.779
## Standardized Within-Group Residuals:
## -2.34017065 -0.39388613 -0.08605866 0.26705911 7.72628368
## Number of Observations: 198
## Number of Groups: 66
```

### Remove Unusual Points

```
CEWL_sub <- CEWL %>%
dplyr::filter(TEWL_g_m2h <= 40)</pre>
```

### Re-Plot



This looks a lot better....

## Re-Model

```
mod3 <- lme4::lmer(data = CEWL_sub, TEWL_g_m2h ~ replicate + (1|ID))</pre>
mod4 <- nlme::lme(data = CEWL_sub, TEWL_g_m2h ~ replicate, random = ~ 1|ID) # for p-values</pre>
summary(mod3)
## Linear mixed model fit by REML ['lmerMod']
## Formula: TEWL_g_m2h ~ replicate + (1 | ID)
##
      Data: CEWL_sub
##
## REML criterion at convergence: 1097.5
##
## Scaled residuals:
       Min
                1Q Median
##
                                 ЗQ
                                        Max
## -2.0156 -0.5504 -0.0698 0.4158 3.3646
```

```
##
## Random effects:
##
    Groups
                          Variance Std.Dev.
    ID
              (Intercept) 18.687
                                   4.323
##
##
    Residual
                           7.935
                                   2.817
                                 ID, 66
## Number of obs: 196, groups:
##
## Fixed effects:
##
               Estimate Std. Error t value
##
   (Intercept)
                14.8771
                             0.7584
                                       19.61
   replicate
                -1.7947
                             0.2472
                                       -7.26
##
  Correlation of Fixed Effects:
##
             (Intr)
##
## replicate -0.661
summary(mod4)
## Linear mixed-effects model fit by REML
##
     Data: CEWL_sub
##
          AIC
                    BIC
                           logLik
##
     1105.523 1118.594 -548.7614
##
## Random effects:
##
    Formula: ~1 | ID
##
           (Intercept) Residual
##
  StdDev:
              4.322866 2.816841
##
## Fixed effects:
                    TEWL_g_m2h ~ replicate
##
                    Value Std.Error DF
                                           t-value p-value
                                                          0
##
   (Intercept) 14.877077 0.7584422 129 19.615307
   replicate
               -1.794696 0.2471906 129 -7.260373
                                                          0
##
    Correlation:
##
             (Intr)
   replicate -0.661
##
##
##
  Standardized Within-Group Residuals:
##
           Min
                         Q1
                                     Med
                                                  QЗ
                                                              Max
##
   -2.01563852 -0.55038247 -0.06984615 0.41576743
                                                     3.36460933
## Number of Observations: 196
## Number of Groups: 66
```

The p-values are still 0 even after removing the two unusually high points...

### Conclusion

There's definitely a lot of variation among triplicate measurements for a given lizard, and the models show that variation is significant. I would conclude that it is important to take these measurements in triplicate to even out the variation in measurements. Taking measurements in triplicate will also allow us to exclude unusual points like the two in the first weekend's dataset, while still having the other replicates for that lizard for us to analyse.