# Climate Water Loss Experiment - Plasma Osmolality Data Wrangling

## Savannah Weaver

## 2021

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`%nin%` = Negate(`%in%`)
<pre>if (!require("tidyverse")) install.packages("tidyverse")  ## Loading required package: tidyverse  ## Attaching packages</pre>

## **Background and Goals**

Blood was drawn from the postorbital sinus of adult male *Sceloporus occidentalis* between June - August 2021. After centrifuging and separating, plasma was run on a VAPRO vapor pressure osmometer in 1-4 replicates, when plasma volume allowed. In this R script, I check the distribution of replicates, omit outliers, and average remaining replicates. The final values will be more precise and accurate estimates of the true plasma osmolality for each lizard, and those values will be used in the capture\_analysis and experiment\_analysis R script files. Please refer to **doi:** for the published scientific journal article and full details.

#### Load Data

```
##
    date_blood_drawn
                          date_osmom_run
                                                time_osmom_run
##
    Min.
           :2021-06-16
                          Min.
                                 :2021-06-16
                                                Min.
                                                        :2021-11-07 01:00:00
##
    1st Qu.:2021-06-30
                          1st Qu.:2021-06-30
                                                1st Qu.:2021-11-07 10:38:00
##
   Median: 2021-07-24
                          Median :2021-07-24
                                                Median :2021-11-07 12:48:00
##
   Mean
           :2021-07-22
                          Mean
                                 :2021-07-22
                                                        :2021-11-07 12:57:52
                                                Mean
##
    3rd Qu.:2021-08-16
                          3rd Qu.:2021-08-16
                                                3rd Qu.:2021-11-07 15:14:00
           :2021-09-01
                                 :2021-09-02
                                                        :2021-11-07 21:22:00
##
    Max.
                          Max.
                                                Max.
##
                                                NA's
                                                        :211
##
    individual_ID
                   replicate_no osmolality_mmol_kg notes
    339
                    1:544
                                                     Mode:logical
##
           :
              13
                                 Min.
                                         :247.0
    206
                    2:531
                                 1st Qu.:336.0
                                                     NA's:1484
##
              12
##
    207
                    3:406
                                 Median :351.0
              12
##
    208
              12
                    4: 3
                                 Mean
                                         :357.3
##
    211
              12
                                 3rd Qu.:370.0
##
    214
                                 Max.
                                         :596.0
              12
    (Other):1411
```

#### Check Data

#### Dates

Blood was drawn on day 0, 4, 8, and 10 of the experiment. Create a list of the dates expected to have blood draw data, then determine whether I have dates outside those.

Trail 1: June 16-24 Trail 2: June 26 - July 4 Trial 3: July 20-28 Trial 4: August 8-16 Trial 5: August 22-30

```
# trial 1
expected_dates <- as.Date(c("2021-06-16", "2021-06-20",
                            "2021-06-24", "2021-06-26",
                            # trial 2
                            "2021-06-26", "2021-06-30",
                            "2021-07-04", "2021-07-06",
                            # trial 3
                            "2021-07-20", "2021-07-24",
                            "2021-07-28", "2021-07-30",
                            # trial 4
                            "2021-08-08", "2021-08-12",
                            "2021-08-16", "2021-08-18",
                            # trial 5
                            "2021-08-22", "2021-08-26",
                            "2021-08-30", "2021-09-01"))
length(osml_reps$date_blood_drawn[osml_reps$date_blood_drawn %nin% expected_dates]
```

## [1] 0

There are zero blood draw dates that are not in our expected list.

#### Number of Blood Draws

Each lizard should have had their blood drawn on 4 different dates, unless they were taken out of the experiment early.

```
# get ID's of the individuals that completed treatment
individuals <- read.csv("./data/tmt_assignments.csv") %>%
  dplyr::select(conclusion, individual_ID) %>%
  dplyr::filter(conclusion == "complete") %>%
  mutate(individual_ID = as.factor(individual_ID),
         conclusion = as.factor(conclusion))
summary(individuals)
##
       conclusion individual_ID
##
   complete:134
                  201
                        : 1
##
                   202
                          : 1
##
                   203
                        : 1
##
                   204
                        : 1
##
                   205
                          : 1
##
                   206
##
                   (Other):128
# calculate the number of dates for each individual
osml reps %>%
  dplyr::filter(individual_ID %in% individuals$individual_ID) %>%
  group_by(individual_ID, date_blood_drawn) %>%
  summarise(n()) %>%
  group_by(individual_ID) %>%
  summarise(n()) %>%
  arrange(n())
## `summarise()` regrouping output by 'individual_ID' (override with `.groups` argument)
## `summarise()` ungrouping output (override with `.groups` argument)
```

```
## # A tibble: 134 x 2
##
      individual_ID `n()`
##
    1 201
##
##
    2 202
                          4
##
    3 203
    4 204
##
    5 205
##
##
    6 206
##
   7 207
    8 208
    9 209
##
## 10 210
## # ... with 124 more rows
```

Wahoo, every lizard has blood draws from 4 different dates.

## Replicates

Now, I will try to identify outliers within the replicates for a given individual on a given date. There must be at least 3 replicates to do this, so the first thing I need to do is figure out which individuals/dates have enough replicates, then subset my data to be only those individuals.

#### Individuals w 3+ Replicates

```
# identify individuals with 3-4 reps
enuf reps <- osml reps %>%
  group_by(individual_ID, date_blood_drawn) %>%
  mutate(count = n()) %>%
  dplyr::filter(count > 2) %>%
  arrange(count)
enuf_reps
## # A tibble: 1,224 x 8
               individual_ID, date_blood_drawn [407]
  # Groups:
      date_blood_drawn date_osmom_run time_osmom_run
                                                           individual_ID
##
      <date>
                       <date>
                                       <dttm>
                                                           <fct>
   1 2021-06-16
                       2021-06-16
                                      2021-11-07 11:00:00 201
##
   2 2021-06-16
                       2021-06-16
                                      2021-11-07 11:00:00 201
##
   3 2021-06-16
                       2021-06-16
                                      2021-11-07 11:00:00 201
                                      2021-11-07 11:00:00 202
##
   4 2021-06-16
                       2021-06-16
##
   5 2021-06-16
                       2021-06-16
                                      2021-11-07 11:00:00 202
   6 2021-06-16
                       2021-06-16
                                      2021-11-07 11:00:00 202
##
##
   7 2021-06-16
                       2021-06-16
                                      2021-11-07 11:10:00 203
##
   8 2021-06-16
                       2021-06-16
                                      2021-11-07 11:10:00 203
##
  9 2021-06-16
                       2021-06-16
                                      2021-11-07 11:10:00 203
                                      2021-11-07 11:25:00 205
## 10 2021-06-16
                       2021-06-16
## # ... with 1,214 more rows, and 4 more variables: replicate_no <fct>,
       osmolality_mmol_kg <dbl>, notes <lgl>, count <int>
# identify individuals with 1-2 reps
not_reps <- osml_reps %>%
  group_by(individual_ID, date_blood_drawn) %>%
 mutate(count = n()) %>%
```

```
dplyr::filter(count < 3) %>%
  arrange(count)
not_reps
## # A tibble: 260 x 8
## # Groups:
               individual_ID, date_blood_drawn [136]
##
      date_blood_drawn date_osmom_run time_osmom_run
                                                           individual_ID
##
      <date>
                       <date>
                                      <dttm>
                                                           <fct>
   1 2021-06-24
                       2021-06-24
##
                                      2021-11-07 09:26:00 203
##
   2 2021-06-24
                       2021-06-26
                                      NA
                                                           204
  3 2021-06-24
##
                       2021-06-26
                                      NA
                                                           223
##
  4 2021-06-26
                       2021-06-26
                                      2021-11-07 09:47:00 201
                                      2021-11-07 11:40:00 230
## 5 2021-06-26
                       2021-06-26
##
   6 2021-07-04
                       2021-07-05
                                      2021-11-07 10:03:00 227
##
  7 2021-07-06
                       2021-07-06
                                      2021-11-07 09:59:00 239
## 8 2021-07-28
                       2021-07-28
                                      2021-11-07 10:32:00 259
## 9 2021-07-28
                       2021-07-28
                                      2021-11-07 12:07:00 274
## 10 2021-07-30
                       2021-07-31
                                      2021-11-07 13:55:00 291
## # ... with 250 more rows, and 4 more variables: replicate_no <fct>,
       osmolality_mmol_kg <dbl>, notes <lgl>, count <int>
# check total obs still add to original 1484
nrow(enuf_reps) + nrow(not_reps)
## [1] 1484
nrow(enuf_reps) + nrow(not_reps) == nrow(osml_reps)
## [1] TRUE
```

#### **Assess Variation**

We want the Coefficient of Variation (CV) among our technical replicates to be small. We need to calculate it to identify whether there may be outliers.

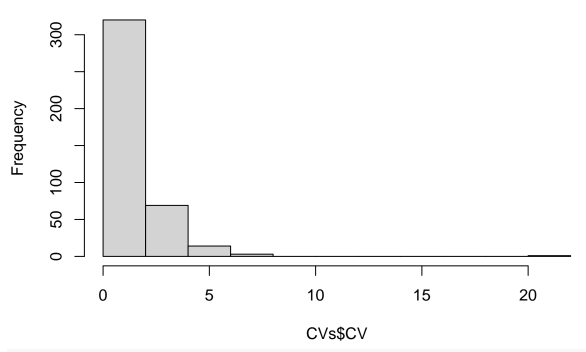
## `summarise()` regrouping output by 'individual\_ID' (override with `.groups` argument)
summary(CVs)

```
##
    individual_ID date_blood_drawn
                                             mean
                                                               SD
##
    206
                  Min.
                          :2021-06-16
                                               :295.3
                                                                : 0.000
           : 4
                                        Min.
                                                        Min.
   207
           : 4
                  1st Qu.:2021-06-30
                                        1st Qu.:335.0
                                                        1st Qu.: 2.581
    208
                  Median :2021-07-24
                                        Median :350.0
                                                        Median: 4.359
##
           :
              4
                                                        Mean
##
   211
             4
                  Mean
                          :2021-07-23
                                        Mean
                                               :356.2
                                                                : 5.596
           :
## 214
             4
                  3rd Qu.:2021-08-16
                                        3rd Qu.:368.0
                                                         3rd Qu.: 6.658
## 220
           :
             4
                  Max.
                         :2021-09-01
                                        Max.
                                               :576.0
                                                        Max.
                                                                :69.816
## (Other):383
```

```
##
                          min
                                          max
                                                         range
         : 0.0000
                            :247.0
                                            :300.0
                                                     Min. : 0.00
##
   Min.
                     Min.
                                    Min.
   1st Qu.: 0.7278
                     1st Qu.:330.5
                                     1st Qu.:339.0
                                                     1st Qu.: 5.00
   Median : 1.2021
                     Median :345.0
                                     Median :356.0
                                                     Median: 8.00
                            :351.1
         : 1.5627
                                            :361.7
                                                           : 10.66
   Mean
                     Mean
                                     Mean
                                                     Mean
##
   3rd Qu.: 1.8746
                     3rd Qu.:364.0
                                     3rd Qu.:374.0
                                                     3rd Qu.: 12.50
   Max.
           :21.4600
                     Max.
                            :566.0
                                     Max.
                                            :596.0
                                                     Max.
                                                            :134.00
##
```

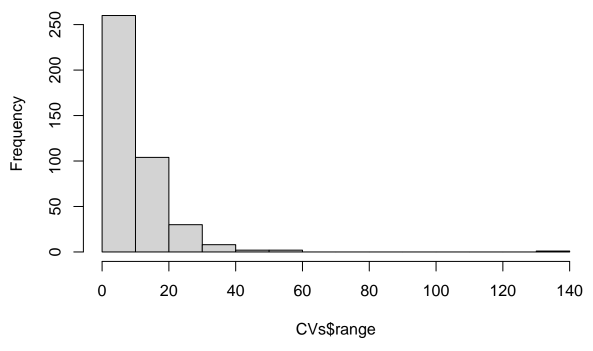
hist(CVs\$CV)

## **Histogram of CVs\$CV**



hist(CVs\$range)

## **Histogram of CVs\$range**



Ideally, CV would be 10-15%. If it's larger, and one of the replicates is very different than the others, we can assume that the replicates that are closer together are more reliable.

The CV is >10 for only one lizard on one date, so our replicates are already likely to accurately represent the true value. We don't want to lose accuracy by searching for precision, so we will only remove the one point driving the enormous CV value.

#### Find Outlier

```
# write function to find outliers for each individual on each date
find_outliers <- function(df) {</pre>
  # initiate dataframe to compile info and list to compile plots
  outliers <- data.frame()</pre>
  #boxplots <- list()</pre>
  # initiate a for loop to go through every who in df
  for(indiv_ch in unique(df$individual_ID)) {
    # select data for only the individual of interest
    df_sub <- df %>%
      dplyr::filter(individual_ID == as.numeric(indiv_ch))
    # make a boxplot
    df_sub %>%
      ggplot(.) +
      geom_boxplot(aes(x = as.factor(date_blood_drawn),
                        y = osmolality_mmol_kg,
                        fill = as.factor(date_blood_drawn))) +
      ggtitle(paste("Individual", indiv_ch)) +
```

```
theme_classic() -> plot

# print/save
print(plot)
#boxplots[[indiv_ch]] <- plot

# extract outliers
outs <- df_sub %>%
    group_by(individual_ID, date_blood_drawn) %>%
    summarise(outs = boxplot.stats(osmolality_mmol_kg)$out)

# add to running dataframe of outliers
outliers <- outliers %>%
    rbind(outs)
}
#return(boxplots)
return(outliers)
}
```

Now apply the function to the data:

```
par(mfrow = c(71, 2))
outliers_found <- find_outliers(enuf_reps)</pre>
```

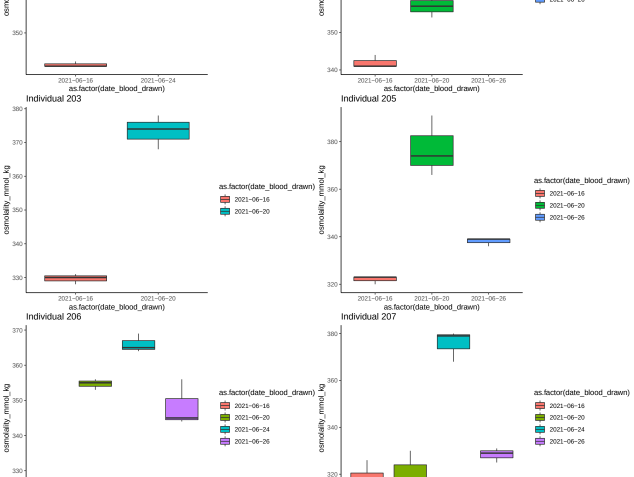
```
## `summarise()` regrouping output by 'individual_ID', 'date_blood_drawn' (override with `.groups` argu
## `summarise()` regrouping output by 'individual_ID', 'date_blood_drawn' (override with `.groups` argu
## `summarise()` regrouping output by 'individual_ID', 'date_blood_drawn' (override with `.groups` argu
## `summarise()` regrouping output by 'individual_ID', 'date_blood_drawn' (override with `.groups` argu
## `summarise()` regrouping output by 'individual_ID', 'date_blood_drawn' (override with `.groups` argu
## `summarise()` regrouping output by 'individual_ID', 'date_blood_drawn' (override with `.groups` argu
## `summarise()` regrouping output by 'individual_ID', 'date_blood_drawn' (override with `.groups` argu
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```

## `summarise()` regrouping output by 'individual\_ID', 'date\_blood\_drawn' (override with `.groups` argument outliers\_found

## # A tibble: 0 x 3 ## # Groups: individual\_ID, date\_blood\_drawn [0] ## # ... with 3 variables: individual\_ID <fct>, date\_blood\_drawn <date>, ## # outs <dbl> par(mfrow = c(1, 1))Individual 201 Individual 202 380 osmolality\_mmol\_kg as.factor(date\_blood\_drawn) as.factor(date blood drawn) 2021-06-16 2021-06-16 osmolality 2021-06-20 2021-06-24 2021-06-26 350 350 2021-06-20 2021-06-26 as.factor(date\_blood\_drawn) as.factor(date\_blood\_drawn)

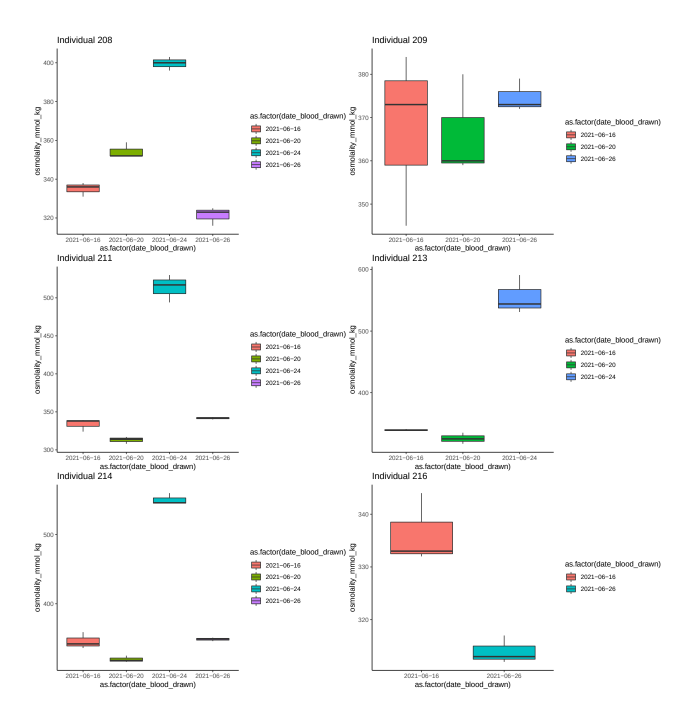


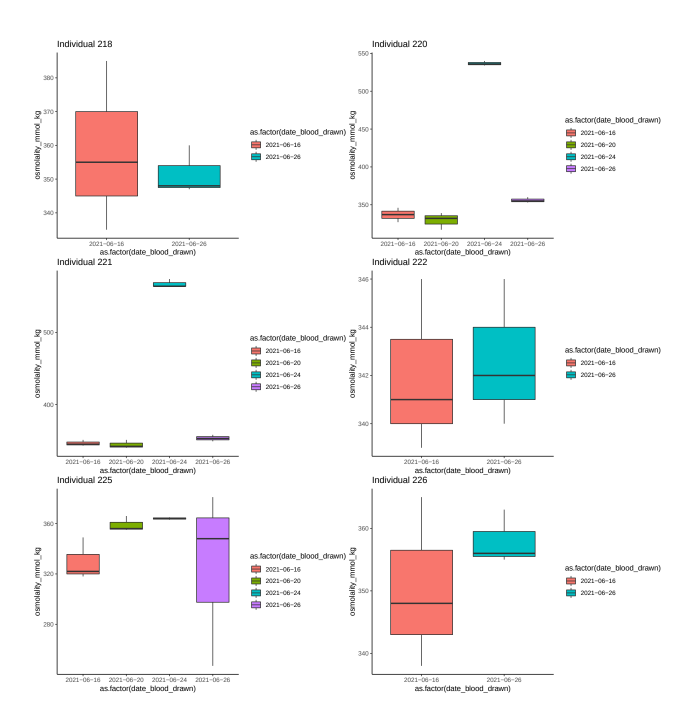
2021-06-16 2021-06-20 2021-06-24 2021-06-26

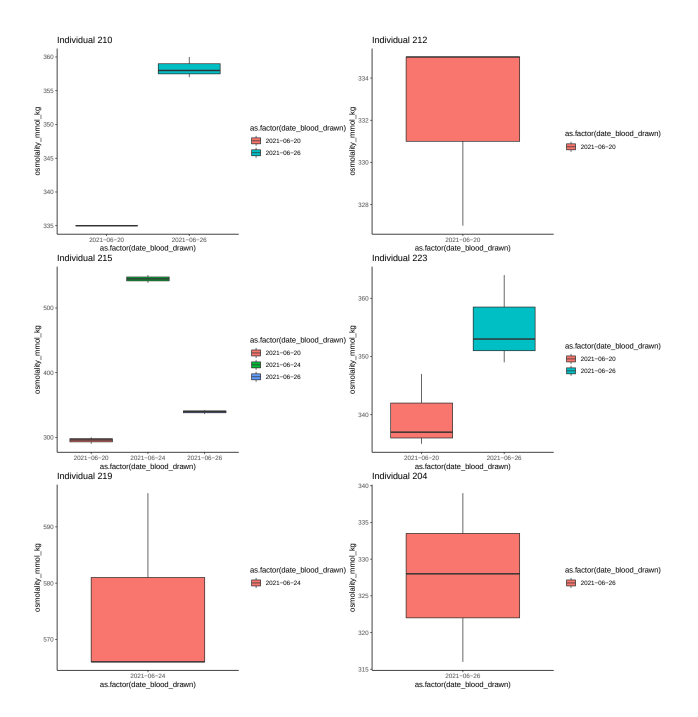
as.factor(date\_blood\_drawn)

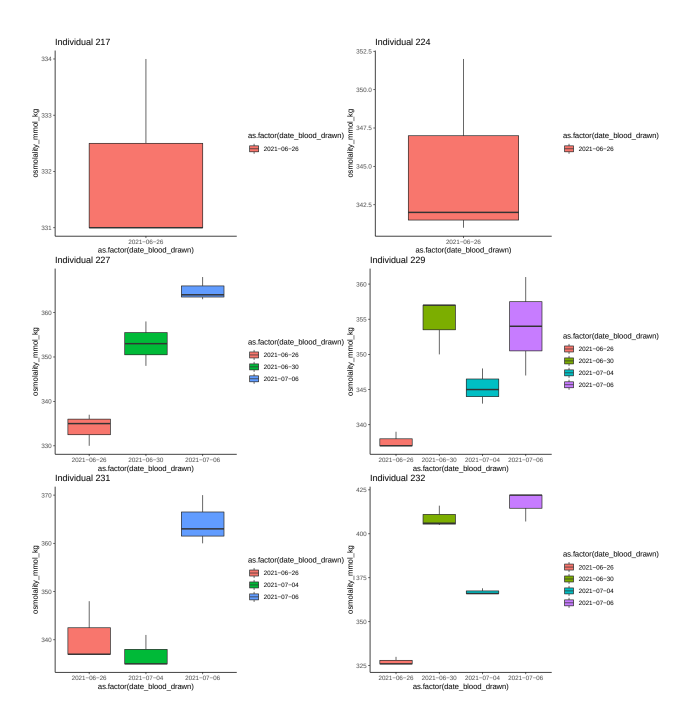
2021-06-16 2021-06-20 2021-06-24 2021-06-26

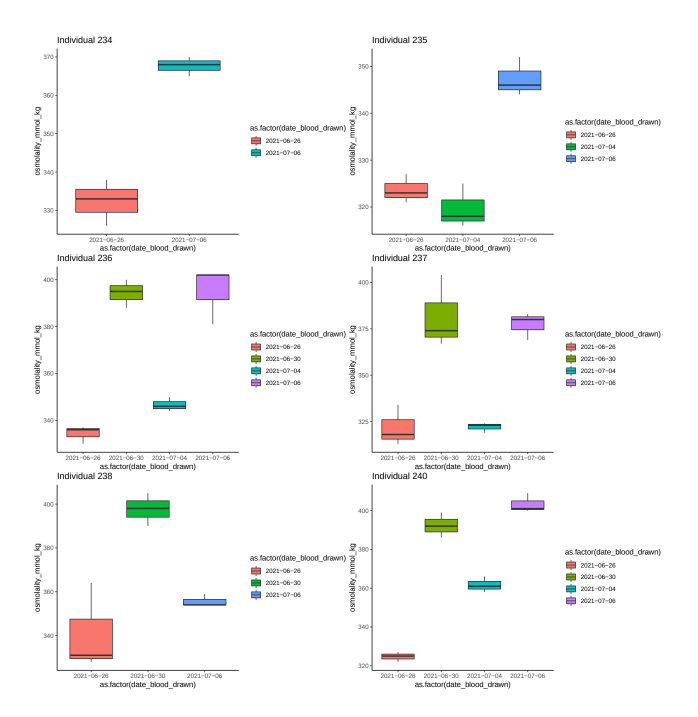
as.factor(date\_blood\_drawn)

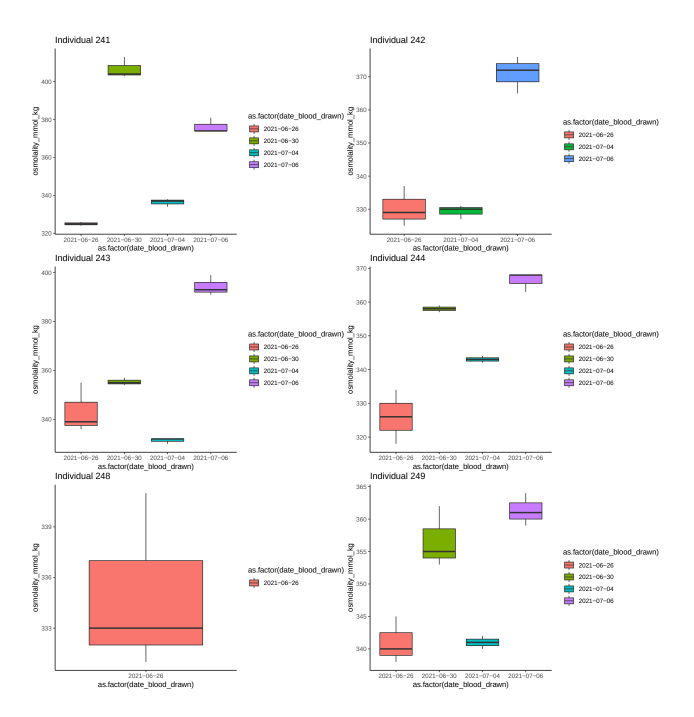


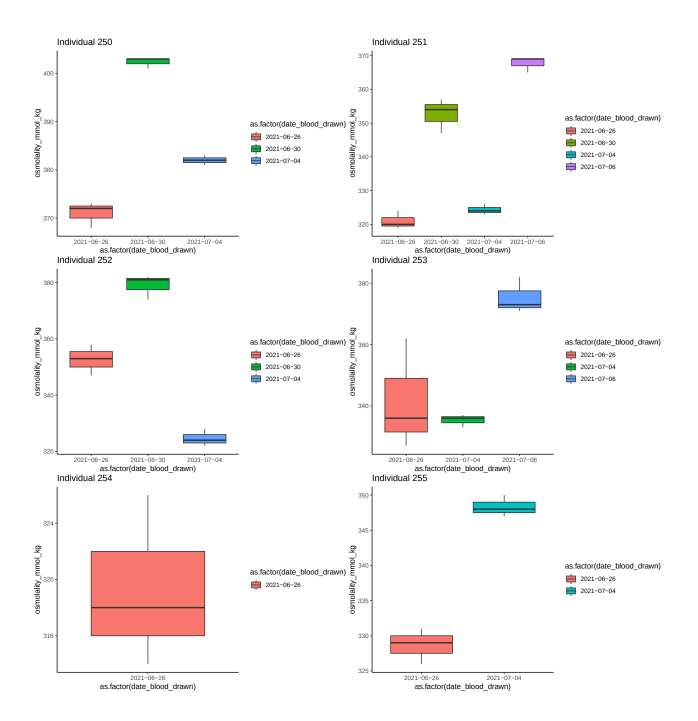


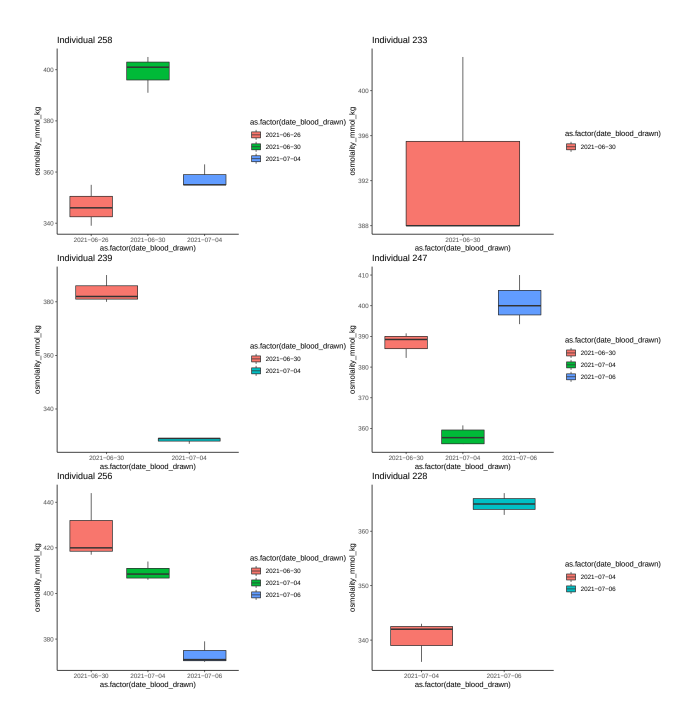


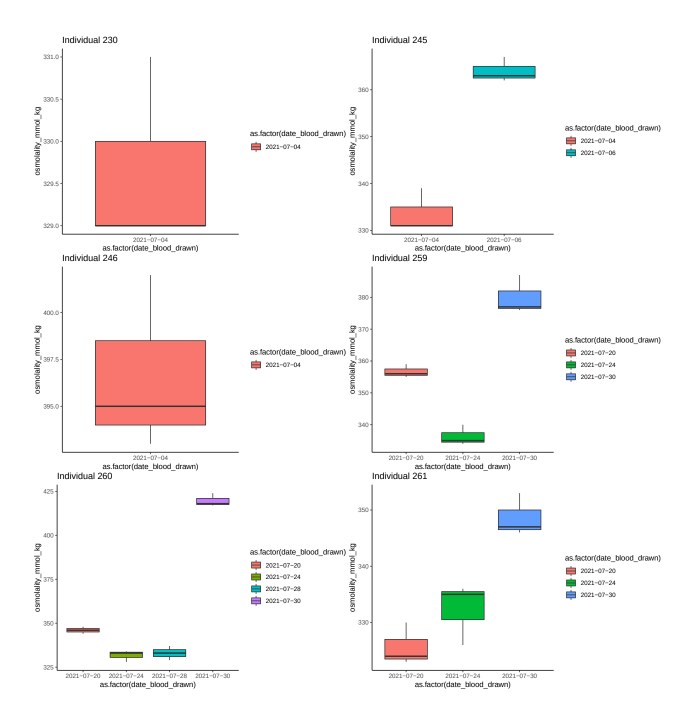


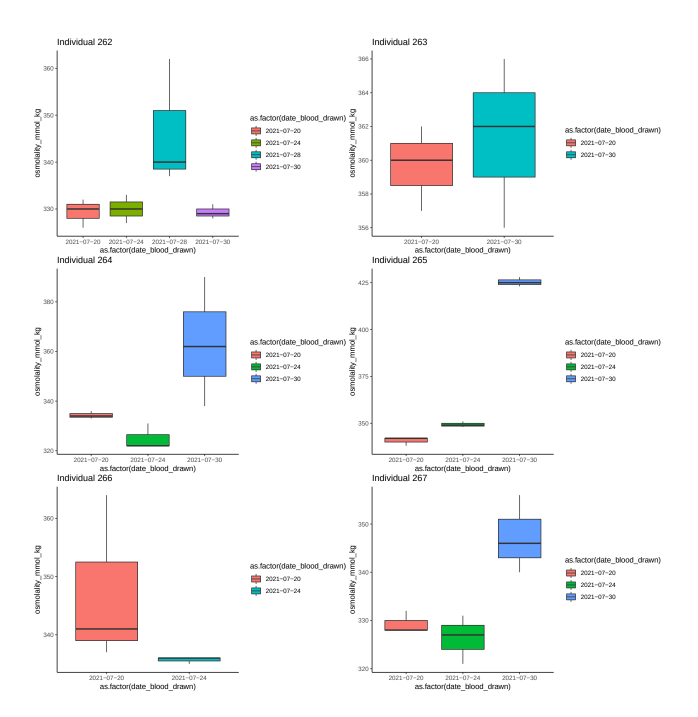


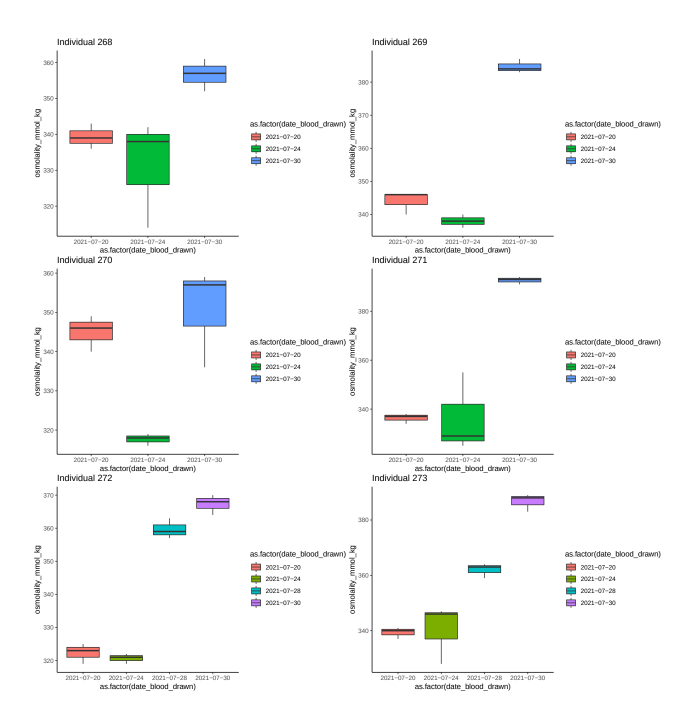


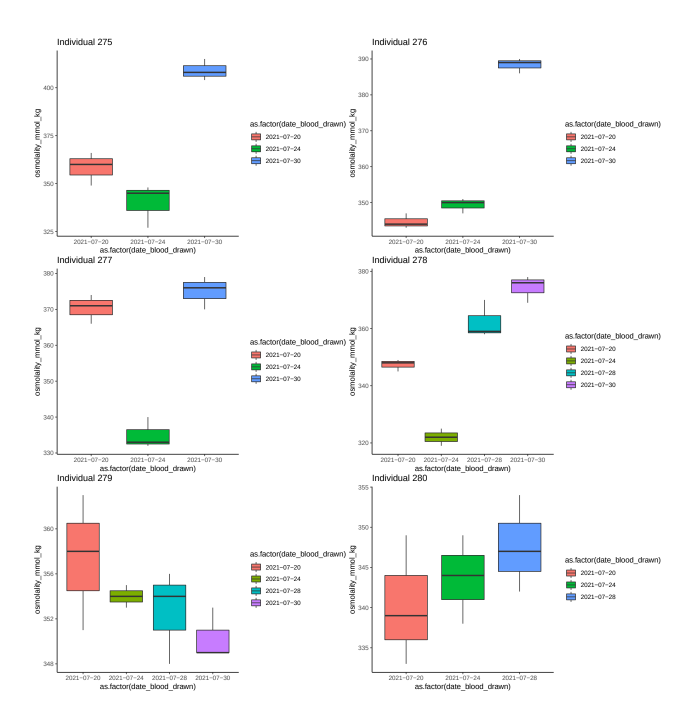


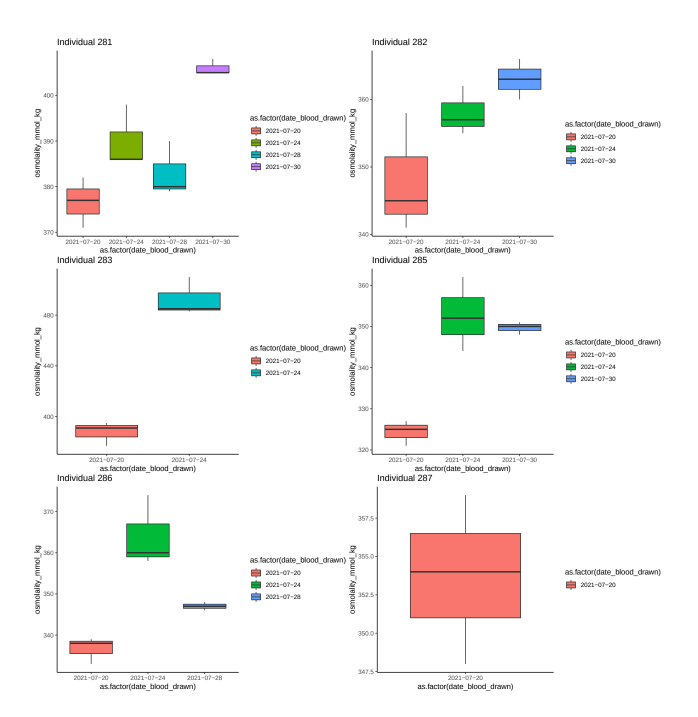


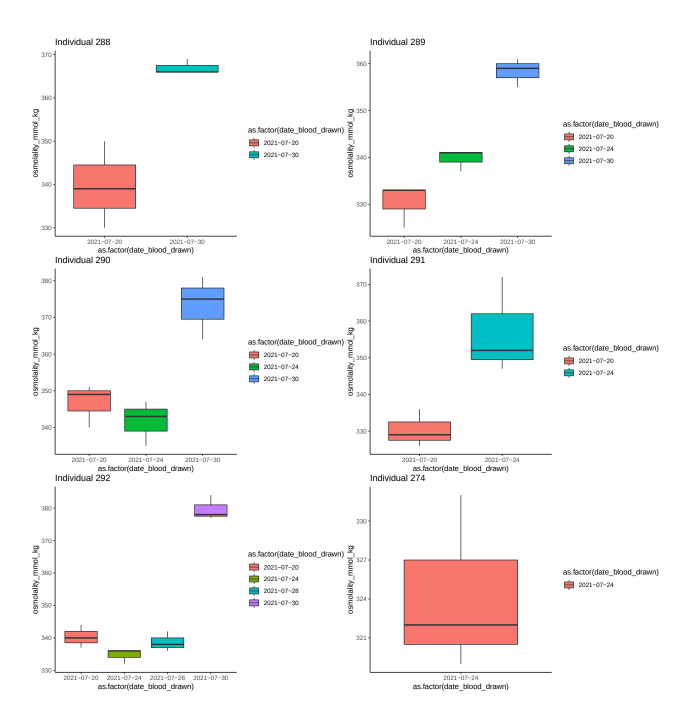


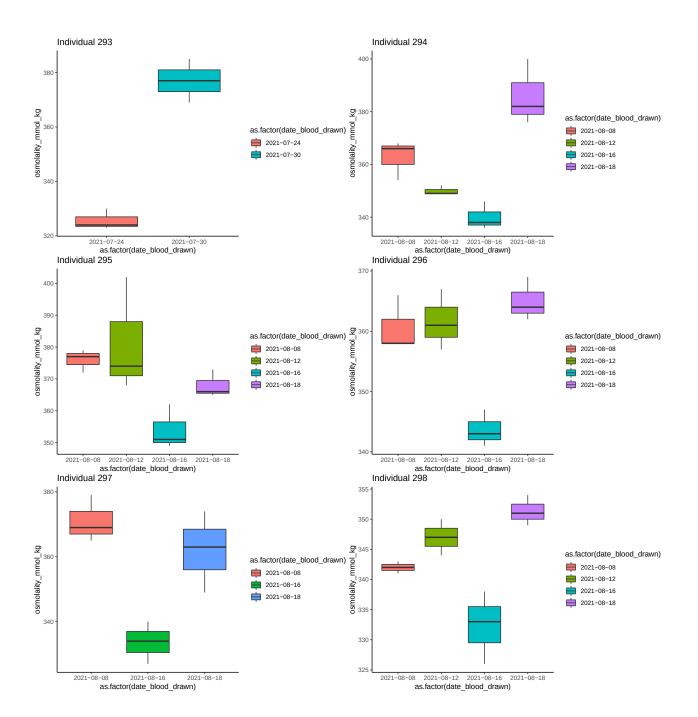


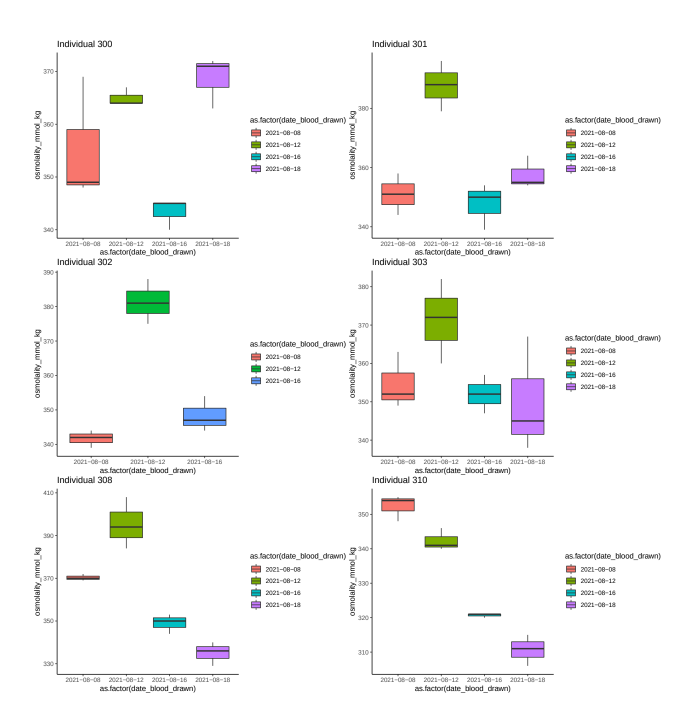


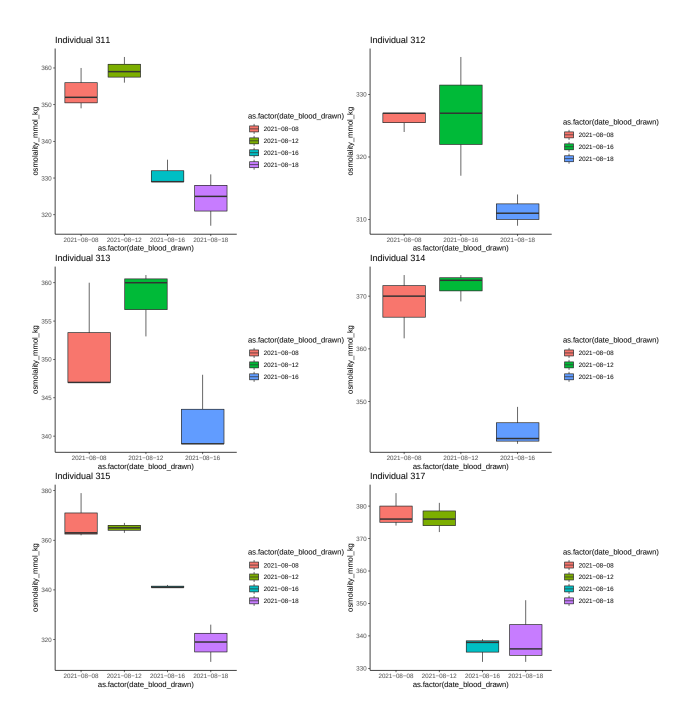


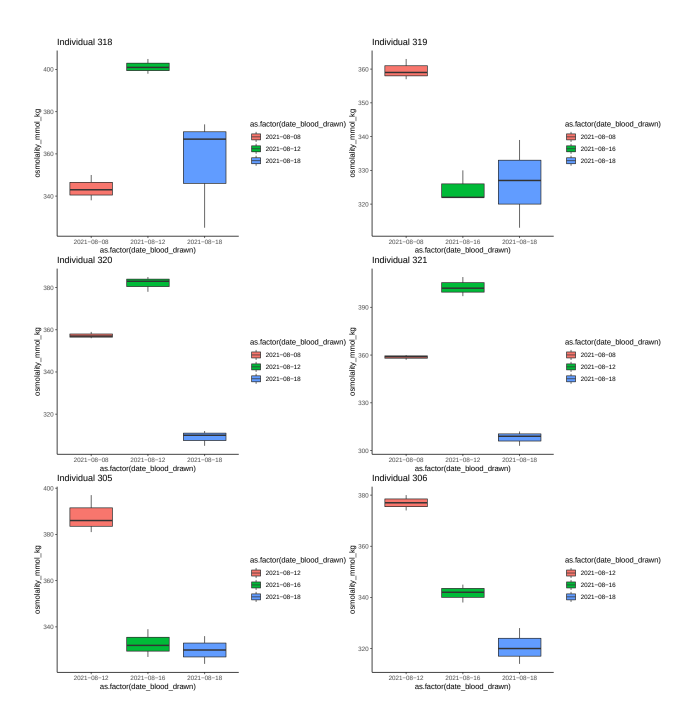


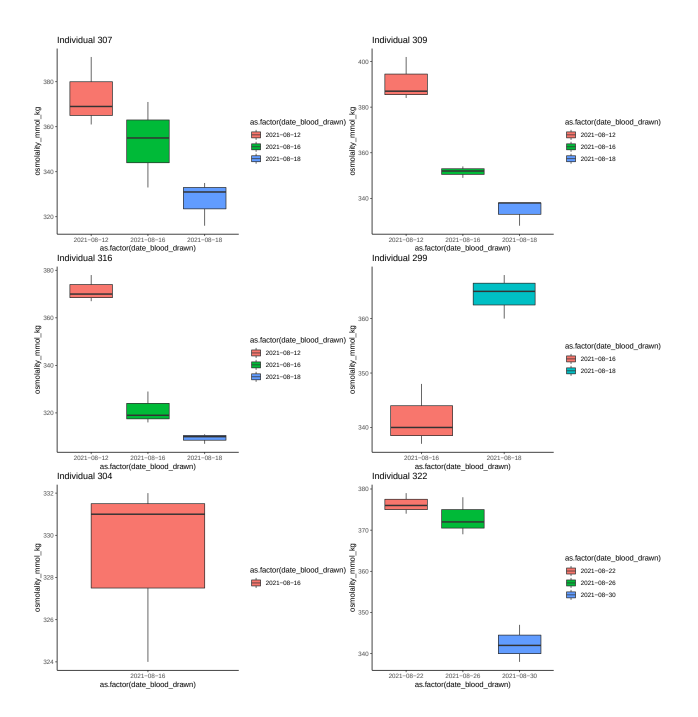


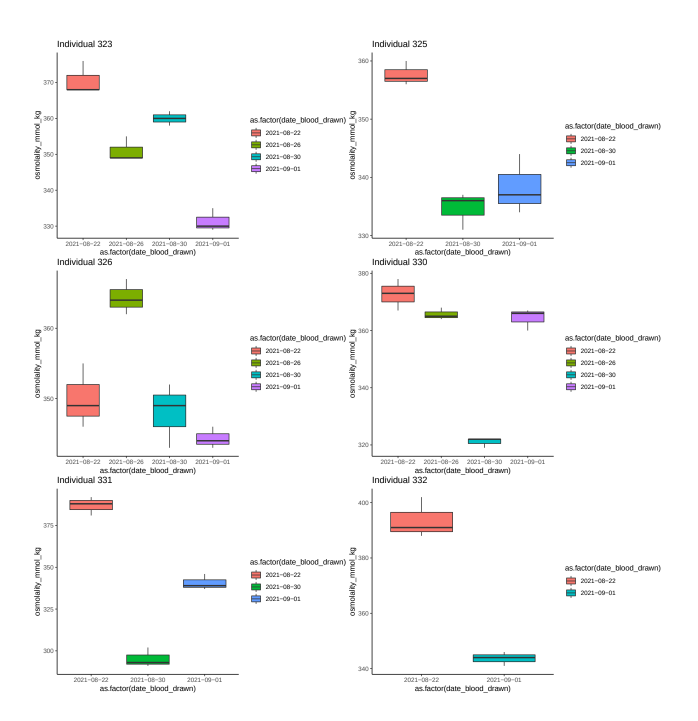


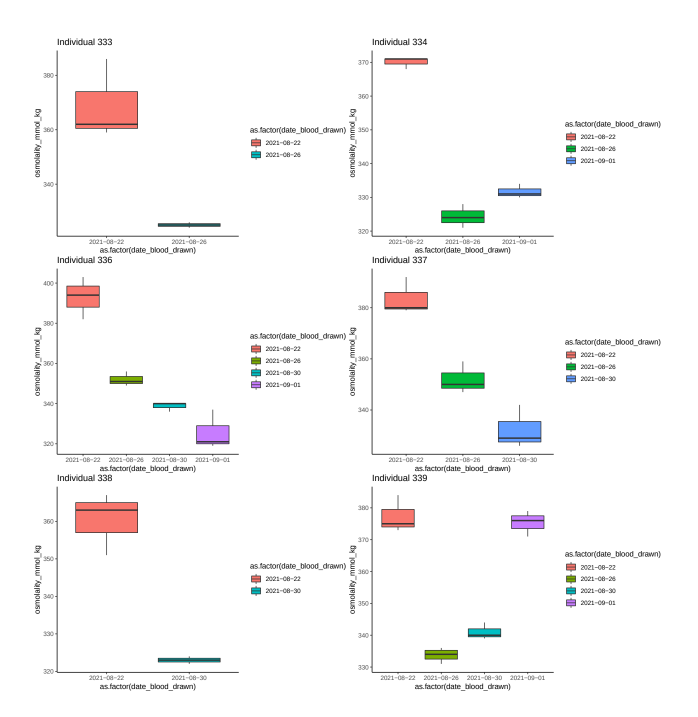


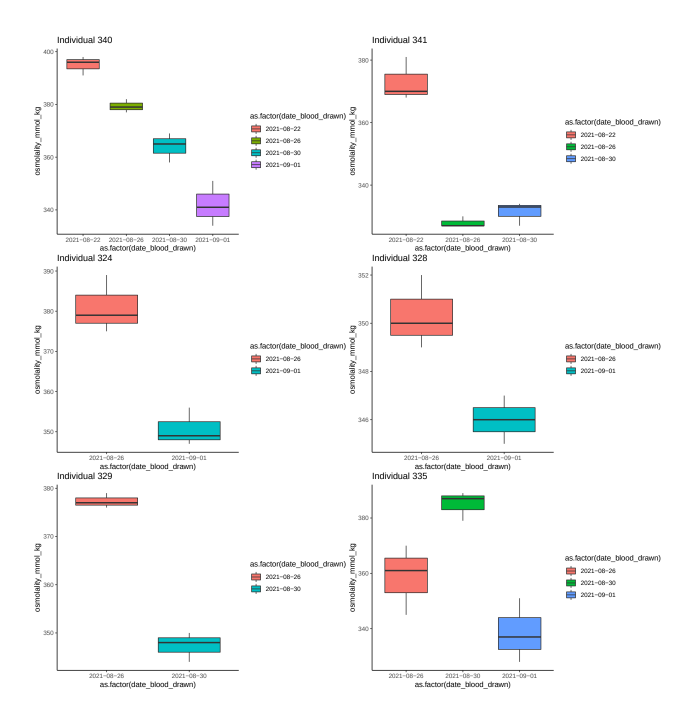


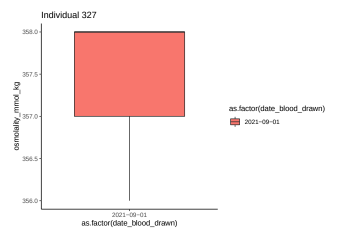












Based on boxplots, there are no outliers, but we still want to find and omit the point severely increasing CV for one individual on one date.

Determine which replicate is an outlier and pulls the CV above our acceptable range (10-15%).

```
CV_all <- enuf_reps %>%
    group_by(individual_ID, date_blood_drawn) %>%
    summarise(mean = mean(osmolality_mmol_kg),
              SD = sd(osmolality_mmol_kg),
              CV \ all = (SD/mean) *100) %>%
    dplyr::select(-mean, -SD)
## `summarise()` regrouping output by 'individual_ID' (override with `.groups` argument)
CV 12 <- enuf reps %>%
    dplyr::filter(replicate_no != 3) %>%
    group_by(individual_ID, date_blood_drawn) %>%
    summarise(mean = mean(osmolality_mmol_kg),
              SD = sd(osmolality_mmol_kg),
              CV_12 = (SD/mean) *100) %>%
   dplyr::select(-mean, -SD)
## `summarise()` regrouping output by 'individual_ID' (override with `.groups` argument)
CV_23 <- enuf_reps %>%
   dplyr::filter(replicate_no != 1) %>%
    group_by(individual_ID, date_blood_drawn) %>%
    summarise(mean = mean(osmolality_mmol_kg),
              SD = sd(osmolality_mmol_kg),
              CV_23 = (SD/mean) *100) %>%
   dplyr::select(-mean, -SD)
## `summarise()` regrouping output by 'individual_ID' (override with `.groups` argument)
CV 31 <- enuf reps %>%
   dplyr::filter(replicate_no != 2) %>%
    group_by(individual_ID, date_blood_drawn) %>%
    summarise(mean = mean(osmolality mmol kg),
              SD = sd(osmolality_mmol_kg),
              CV_31 = (SD/mean) *100) %>%
    dplyr::select(-mean, -SD)
```

## `summarise()` regrouping output by 'individual\_ID' (override with `.groups` argument)

```
compare <- CV_all %>%
    left_join(CV_12, by = c('individual_ID', 'date_blood_drawn')) %>%
   left_join(CV_23, by = c('individual_ID', 'date_blood_drawn')) %>%
   left_join(CV_31, by = c('individual_ID', 'date_blood_drawn')) %>%
  arrange(desc(CV_all)) %>%
  dplyr::filter(CV_all>10)
compare
## # A tibble: 1 x 6
## # Groups: individual ID [1]
   individual_ID date_blood_drawn CV_all CV_12 CV_23 CV_31
##
     <fct>
                   <date>
                                    <dbl> <dbl> <dbl> <dbl>
## 1 225
                   2021-06-26
                                     21.5 30.2 24.0 6.40
```

We should remove replicate 2 for individual 225 on June 26.

#### Remove Outlier

```
enuf_reps[199:201, 5:6]
## # A tibble: 3 x 2
   replicate_no osmolality_mmol_kg
     <fct>
                                <dbl>
## 1 1
                                  381
## 2 2
                                  247
## 3 3
                                  348
enuf_reps[200, "osmolality_mmol_kg"] = NA
enuf_reps[199:201, 5:6]
## # A tibble: 3 x 2
##
    replicate_no osmolality_mmol_kg
## 1 1
                                  381
## 2 2
                                   NA
## 3 3
                                  348
# check
enuf_reps %>%
 dplyr::filter(individual_ID == 225,
                date_blood_drawn == as.Date("2021-06-26",
                                               format= "%Y-%m-%d")
                #replicate_no == 2
## # A tibble: 3 x 8
## # Groups: individual_ID, date_blood_drawn [1]
     date_blood_drawn date_osmom_run time_osmom_run
                                                          individual_ID replicate_no
     <date>
                      <date>
                                     <dttm>
                                                                         <fct>
                                                           \langle fct. \rangle
## 1 2021-06-26
                      2021-06-26
                                      2021-11-07 09:42:00 225
                                                                         1
## 2 2021-06-26
                      2021-06-26
                                      2021-11-07 09:42:00 225
                                                                         2
## 3 2021-06-26
                      2021-06-26
                                     2021-11-07 09:42:00 225
## # ... with 3 more variables: osmolality_mmol_kg <dbl>, notes <lgl>, count <int>
# filter out the one NA
enuf_reps_trimmed <- enuf_reps %>%
```

```
dplyr::filter(complete.cases(osmolality_mmol_kg))
```

### Average Remaining Replicates

Now that the outliers are removed from the technical replicates when there were enough replicates to identify them, I will average the remaining replicates of the rejoined data for lizards with 1-2 and 3-4 replicates.

```
osml_means <- not_reps %>%
  rbind(enuf_reps_trimmed) %>%
  group_by(date_blood_drawn, individual_ID) %>%
  summarise(osmolality_mmol_kg_mean = mean(osmolality_mmol_kg))
```

## `summarise()` regrouping output by 'date\_blood\_drawn' (override with `.groups` argument)

## **Export**

```
write.csv(osml_means, "./data/osml_means_clean.csv")
```