Data Analysis

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Loading library and data

```
library(performance)
library(DHARMa)
library(mgcv)
library(fitdistrplus)
library(goft)
library(gamlss)
library(FSA)
library(fGarch)
library(LambertW)
library(ordinal)
library(ordinal)
library(here)
library(plotrix)
library(patchwork)
library(tidyverse)
```

```
health_data <- read.csv("Big_Data_Green.csv")
heat_data <- read.csv("Heat_Data.csv")
```

Health Data

Includes feeding time, PAM measurements, base measurements, and symbiont density.

Symbiont Data

PAM Threshold

Determining a minum threshold for aposymbiotic anemones that we will remove from the PAM data. From the data bellow, we decide that the threshold for PAM will be >1000 cells per mg (symbiont density).

```
Green_Threshold <- health_data %>%
  select(-c(Event, Field_Site, Acclimation_Period, Base_Width, Base_Length,
            Base_Diagonal, Base_Diameter_mm, Nb_Tentacles, Weight_Total_g,
            Weight_Tube_g, Weight_Tentacle_g, Div_Dino, Dino_Cells, Feeding_Time_Min,
            Feeding Time, Feeding Start, Feeding Stop, Photo ID, Removed, MI Dino,
            Fv_Fm_1, Fv_Fm_2, Fv_Fm_3, PAM_avg, Event_True, Dino_Density)) %>%
  drop na(Green Density) %>%
  drop_na(Time_Point) %>%
  filter(Green_Density != 0)
Dino_Threshold <- health_data %>%
  select(-c(Event, Field_Site, Acclimation_Period, Base_Width, Base_Length,
            Base_Diagonal, Base_Diameter_mm, Nb_Tentacles, Weight_Total_g,
            Weight_Tube_g, Weight_Tentacle_g, Green_Cells, Div_Green, Feeding_Time_Min,
            Feeding_Time, Feeding_Start, Feeding_Stop, Photo_ID, Removed, MI_Green,
            Fv_Fm_1, Fv_Fm_2, Fv_Fm_3,
            PAM_avg, Event_True, Green_Density)) %>%
  drop_na(Dino_Density) %>%
  drop_na(Time_Point) %>%
  filter(Dino_Density != 0)
```

Density Data

Data for models of symbiont density for both zoochlorellae and zooxanthellae. It filters out unwanted columns and anemones without symbiont count.

```
# Organizing the data
Green_Symbionts <- health_data %>%
  select(-c(Event, Base_Width, Base_Length, Base_Diagonal, Base_Diameter_mm,
            Nb_Tentacles, Weight_Total_g, Weight_Tube_g, Weight_Tentacle_g,
            Green_Cells, Dino_Cells, Feeding_Time_Min, Feeding_Time, Feeding_Start,
            Feeding_Stop, Photo_ID, Removed, Fv_Fm_1, Fv_Fm_2, Fv_Fm_3, PAM_avg,
            Event_True)) %>%
  filter(Date == "11/5/2021" | Date == "11/9/2021" | Date == "11/13/2021") %>%
  mutate(Date = fct relevel(Date, "11/5/2021", "11/9/2021", "11/13/2021")) %>%
  mutate(Green_Density_Log = log(Green_Density)) %>%
  drop na(Green Density) %>%
  drop_na(Dino_Density) %>%
  drop_na(MI_Green) %>%
  drop_na(MI_Dino) %>%
  filter(Green Density != 0) %>%
  mutate(Acclimation Period = as.factor(Acclimation Period)) %>%
```

```
group_by(Treatment, Date) %>%
  mutate(Mean_Green_Log = mean(Green_Density_Log),
         SE_Green_Log = std.error(Green_Density_Log))
Dino_Symbionts <- health_data %>%
  select(-c(Event, Base_Width, Base_Length, Base_Diagonal, Base_Diameter_mm,
            Nb_Tentacles, Weight_Total_g, Weight_Tube_g, Weight_Tentacle_g,
            Green_Cells, Dino_Cells, Feeding_Time_Min, Feeding_Time,
            Feeding_Start, Feeding_Stop, Photo_ID, Removed, Fv_Fm_1, Fv_Fm_2,
            Fv_Fm_3, PAM_avg, Event_True)) %>%
  filter(Date == "11/5/2021" | Date == "11/9/2021" | Date == "11/13/2021") %>%
  mutate(Date = fct_relevel(Date, "11/5/2021", "11/9/2021", "11/13/2021")) %>%
  drop na(Green Density) %>%
  drop_na(Dino_Density) %>%
  drop_na(MI_Green) %>%
  drop_na(MI_Dino) %>%
  filter(Dino_Density != 0) %>%
  mutate(Acclimation_Period = as.factor(Acclimation_Period))
kruskal.test(Green_Density ~ Treatment, data = Green_Symbionts) # p = 0.003
##
## Kruskal-Wallis rank sum test
## data: Green Density by Treatment
## Kruskal-Wallis chi-squared = 11.091, df = 2, p-value = 0.003905
dunnTest(Green_Density ~ Treatment, data = Green_Symbionts) # sig. difference between 25-30 and 30-cont
## Dunn (1964) Kruskal-Wallis multiple comparison
##
    p-values adjusted with the Holm method.
##
        Comparison
                            Z
                                  P.unadj
         25C - 30C -2.5463792 0.010884689 0.021769378
## 2 25C - Control 0.1437093 0.885729997 0.885729997
## 3 30C - Control 3.0094276 0.002617404 0.007852213
kruskal.test(Dino_Density ~ Treatment, data = Dino_Symbionts) # not significant
##
  Kruskal-Wallis rank sum test
##
## data: Dino_Density by Treatment
## Kruskal-Wallis chi-squared = 4.742, df = 2, p-value = 0.09339
dunnTest(Dino_Density ~ Treatment, data = Dino_Symbionts) # not significant
## Dunn (1964) Kruskal-Wallis multiple comparison
    p-values adjusted with the Holm method.
```

```
## Comparison Z P.unadj P.adj
## 1 25C - 30C 0.3672174 0.71345689 0.7134569
## 2 25C - Control -1.4749869 0.14021607 0.2804321
## 3 30C - Control -2.1053369 0.03526199 0.1057860
```

Checking Normality and Equal Variance:

```
# Green
shapiro.test(Green_Symbionts$Green_Density)
##
##
   Shapiro-Wilk normality test
##
## data: Green_Symbionts$Green_Density
## W = 0.64647, p-value = 3.82e-10
bartlett.test(Green_Density ~ Treatment, Green_Symbionts)
##
##
   Bartlett test of homogeneity of variances
##
## data: Green Density by Treatment
## Bartlett's K-squared = 23.091, df = 2, p-value = 9.681e-06
# Dino
shapiro.test(Dino_Symbionts$Dino_Density)
##
##
   Shapiro-Wilk normality test
## data: Dino_Symbionts$Dino_Density
## W = 0.80442, p-value = 5.198e-07
bartlett.test(Dino_Density ~ Treatment, Dino_Symbionts)
##
## Bartlett test of homogeneity of variances
##
## data: Dino_Density by Treatment
## Bartlett's K-squared = 3.8169, df = 2, p-value = 0.1483
```

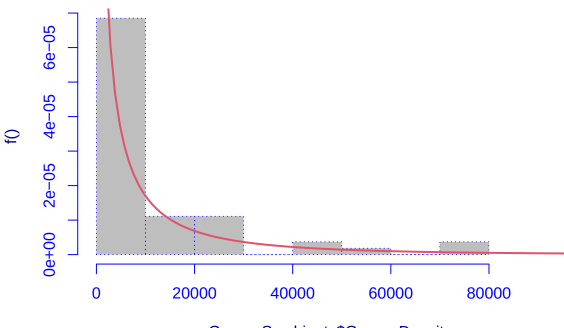
All p-values are <0.05 and therefore the distributions do not meet assumptions of normality and equal variance.

Checking Distributions of Symbiont Data: Determining if the data fits other distributions.

```
#Greens
exp_test(Green_Symbionts$Green_Density) #p-value < 0.05
```

```
##
## Test for exponentiality based on a transformation to uniformity
## data: Green_Symbionts$Green_Density
## T = 4.719, p-value < 2.2e-16
gamma_test(Green_Symbionts$Green_Density)
                                                #p = 0.123
##
  Test of fit for the Gamma distribution
##
## data: Green_Symbionts$Green_Density
## V = -0.12047, p-value = 0.9321
lnorm_test(Green_Symbionts$Green_Density)
                                                #p-value < 0.05
##
## Test for the lognormal distribution based on a transformation to
## normality
## data: Green_Symbionts$Green_Density
## p-value = 0.006495
normal_test(Green_Symbionts$Green_Density)
                                                #p-value < 0.05
##
## Correlation test for normality
## data: Green_Symbionts$Green_Density
## R = 0.89717, p-value = 1.019e-06
## alternative hypothesis: Green_Symbionts$Green_Density does not follow a normal distribution.
weibull_test(Green_Symbionts$Green_Density)
                                                #p = 0.44
##
## Test for the Weibull distribution
## data: Green_Symbionts$Green_Density
## p-value = 0.048
#Use Weibull since p value is larger than gamma
Green_Weibull <- histDist(Green_Symbionts$Green_Density, "WEI",</pre>
                          density = F, main = "Weibull")
```

Weibull

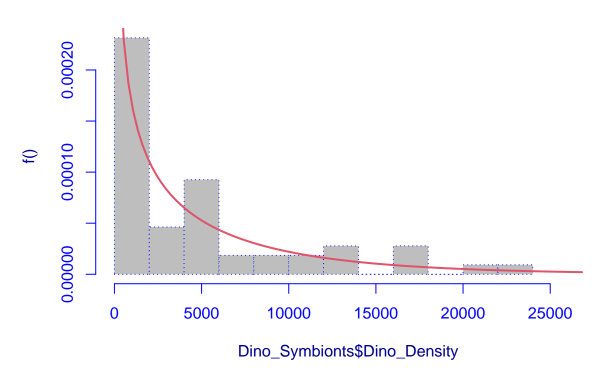


Green_Symbionts\$Green_Density

```
#Dinos
exp_test(Dino_Symbionts$Dino_Density)
                                                 #p-value > 0.05
##
##
   Test for exponentiality based on a transformation to uniformity
## data: Dino_Symbionts$Dino_Density
## T = 2.6262, p-value = 0.002
gamma_test(Dino_Symbionts$Dino_Density)
                                                 #p = 0.6709
##
    Test of fit for the Gamma distribution
##
## data: Dino_Symbionts$Dino_Density
## V = -1.131, p-value = 0.4239
lnorm_test(Dino_Symbionts$Dino_Density)
                                                 #p-value < 0.05
##
   Test for the lognormal distribution based on a transformation to
##
##
    normality
##
## data: Dino_Symbionts$Dino_Density
## p-value = 0.003319
```

```
normal_test(Dino_Symbionts$Dino_Density)
                                                 #p-value < 0.05
##
##
    Correlation test for normality
##
## data: Dino_Symbionts$Dino_Density
## R = 0.95707, p-value = 7.874e-05
## alternative hypothesis: Dino_Symbionts$Dino_Density does not follow a normal distribution.
weibull_test(Dino_Symbionts$Dino_Density)
                                                 #p = 0.774
##
##
    Test for the Weibull distribution
## data: Dino_Symbionts$Dino_Density
## p-value = 0.944
	t #Use Gamma since p value is larger than weibull
Dino_Gamma <- histDist(Dino_Symbionts$Dino_Density, "GA",</pre>
                       density = F, main = "Gamma")
```

Gamma



Dunn and Kruskal Wallis Test on Density Data: Used for box blot

```
kruskal.test(Green_Density ~ Treatment, data = Green_Symbionts) # p = 0.004
##
   Kruskal-Wallis rank sum test
##
##
## data: Green_Density by Treatment
## Kruskal-Wallis chi-squared = 11.091, df = 2, p-value = 0.003905
dunnTest (Green_Density ~ Treatment, data = Green_Symbionts) # sig. difference between 25-30 and between
## Dunn (1964) Kruskal-Wallis multiple comparison
     p-values adjusted with the Holm method.
##
##
        Comparison
                            Z
                                  P.unadj
         25C - 30C -2.5463792 0.010884689 0.021769378
## 1
## 2 25C - Control 0.1437093 0.885729997 0.885729997
## 3 30C - Control 3.0094276 0.002617404 0.007852213
kruskal.test(Dino_Density ~ Treatment, data = Dino_Symbionts) # not significant
##
  Kruskal-Wallis rank sum test
##
## data: Dino_Density by Treatment
## Kruskal-Wallis chi-squared = 4.742, df = 2, p-value = 0.09339
dunnTest(Dino_Density ~ Treatment, data = Dino_Symbionts) # not significant
## Dunn (1964) Kruskal-Wallis multiple comparison
     p-values adjusted with the Holm method.
##
##
        Comparison
                            Z
                                 P.unadj
                                             P.adj
         25C - 30C 0.3672174 0.71345689 0.7134569
## 2 25C - Control -1.4749869 0.14021607 0.2804321
## 3 30C - Control -2.1053369 0.03526199 0.1057860
```

Running Models on Symbiont Data: We chose to use our Weibull distribution for green density and Gamma distribution for dino density. Here we run our models.

```
## GAMLSS-RS iteration 1: Global Deviance = 927.0726
## GAMLSS-RS iteration 2: Global Deviance = 910.9055
## GAMLSS-RS iteration 3: Global Deviance = 908.0624
## GAMLSS-RS iteration 4: Global Deviance = 907.6404
## GAMLSS-RS iteration 5: Global Deviance = 907.5682
## GAMLSS-RS iteration 6: Global Deviance = 907.5528
## GAMLSS-RS iteration 7: Global Deviance = 907.549
## GAMLSS-RS iteration 8: Global Deviance = 907.5483
summary(Green_Weibull_model) # significance in the 30 degree treatment
## Family: c("WEI", "Weibull")
##
## Call: gamlss(formula = Green_Density ~ Date * Treatment +
     random(Species_ID) + random(Field_Site) + random(Acclimation_Period),
##
     family = WEI(), data = Green_Symbionts, control = gamlss.control(n.cyc = 10))
##
##
## Fitting method: RS()
## -----
## Mu link function: log
## Mu Coefficients:
                         Estimate Std. Error t value Pr(>|t|)
                          ## (Intercept)
## Date11/9/2021
                         -0.17523 0.27804 -0.630 0.5354
                          0.28886 0.28856
## Date11/13/2021
                                            1.001
                                                   0.3283
                         -0.12147 0.32291 -0.376
## Treatment25C
                                                   0.7106
## Treatment30C
                          ## Date11/9/2021:Treatment25C 0.05102 0.45057
                                           0.113
                                                   0.9109
## Date11/13/2021:Treatment25C -0.83396 0.44295 -1.883
                                                   0.0738 .
## Date11/9/2021:Treatment30C 0.13530 0.38302 0.353
                                                   0.7275
## Date11/13/2021:Treatment30C 0.32660 0.38314 0.852
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
##
## -----
## Sigma link function: log
## Sigma Coefficients:
            Estimate Std. Error t value Pr(>|t|)
## (Intercept) 0.6939
                    0.1161 5.977 6.48e-06 ***
## ---
## Signif. codes: 0 '*** 0.001 '** 0.01 '* 0.05 '.' 0.1 ' 1
##
## NOTE: Additive smoothing terms exist in the formulas:
## i) Std. Error for smoothers are for the linear effect only.
## ii) Std. Error for the linear terms maybe are not accurate.
## No. of observations in the fit: 54
## Degrees of Freedom for the fit: 33.20121
       Residual Deg. of Freedom: 20.79879
```

at cycle: 8

##

```
##
## Global Deviance: 907.5483
                 973.9507
##
           AIC:
##
           SBC:
                  1039.987
## ***********************
Dino_Symbionts\( \frac{1}{3}\) orderTreatment = ordered(Dino_Symbionts\( \frac{1}{3}\) Treatment,
                               levels = c("Control", "25C", "30C"))
Dino_Gamma_model <- gamlss(formula = Dino_Density ~ Date*Treatment +</pre>
                       random(Species_ID) + random(Field_Site) +
                       random(Acclimation_Period),
                     family = GA(), data = Dino_Symbionts,
                     control = gamlss.control(n.cyc = 3))
## GAMLSS-RS iteration 1: Global Deviance = 908.9065
## GAMLSS-RS iteration 2: Global Deviance = 908.8828
## GAMLSS-RS iteration 3: Global Deviance = 908.8819
summary (Dino_Gamma_model) # significance in both the 25 and 30 degree treatment
## Family: c("GA", "Gamma")
##
## Call: gamlss(formula = Dino_Density ~ Date * Treatment +
     random(Species_ID) + random(Field_Site) + random(Acclimation_Period),
##
     family = GA(), data = Dino_Symbionts, control = gamlss.control(n.cyc = 3))
##
##
##
## Fitting method: RS()
##
## -----
## Mu link function: log
## Mu Coefficients:
                     Estimate Std. Error t value Pr(>|t|)
##
## (Intercept)
                       8.06079 0.25027 32.208 < 2e-16 ***
                       ## Date11/9/2021
## Date11/13/2021
                        -0.23477 0.35394 -0.663 0.51405
## Treatment25C
                       -1.32556 0.39574 -3.350 0.00291 **
                    -0.74465 0.32308 -2.305 0.03104 *
## Treatment30C
## Date11/13/2021:Treatment25C 1.03874 0.54261 1.914 0.06874 .
## Date11/9/2021:Treatment30C -0.23333 0.46976 -0.497 0.62435
## Date11/13/2021:Treatment30C 0.03936 0.46977 0.084 0.93399
## Signif. codes: 0 '*** 0.001 '** 0.01 '* 0.05 '.' 0.1 ' 1
## -----
## Sigma link function: log
## Sigma Coefficients:
           Estimate Std. Error t value Pr(>|t|)
```

```
## ---
## Signif. codes: 0 '*** 0.001 '** 0.01 '* 0.05 '.' 0.1 ' 1
## -----
## NOTE: Additive smoothing terms exist in the formulas:
## i) Std. Error for smoothers are for the linear effect only.
## ii) Std. Error for the linear terms maybe are not accurate.
## No. of observations in the fit: 54
## Degrees of Freedom for the fit: 32.10774
##
      Residual Deg. of Freedom: 21.89226
                   at cycle:
##
##
## Global Deviance:
                  908.8819
##
           AIC:
                  973.0974
##
           SBC:
                  1036.959
```

Figure 1: ZC symbiont density, using gamlss with Weibull distribution. Significance compared to pre-heat of control.

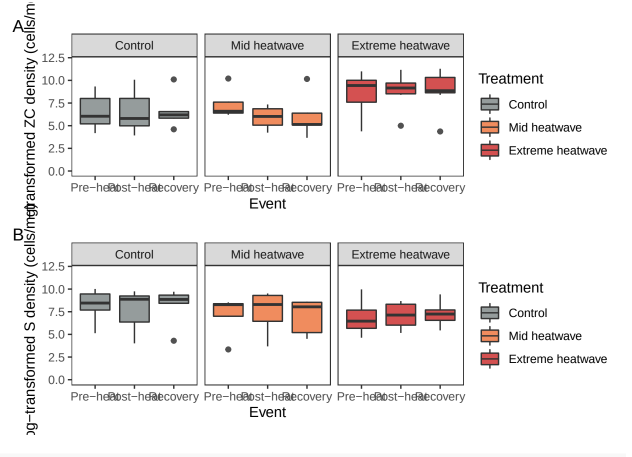
Interaction [ask sara]	Standard Error	t-value	p-value
Post-heat	0.27804	-0.630	0.5354
Recovery	0.28856	1.001	0.3283
Mid heatwave	0.32291	-0.376	0.7106
Extreme heatwave	0.26344	5.600	< 0.001
Post-heat: Mid heatwave	0.45057	0.113	0.9109
Post-heat : Extreme heatwave	0.44295	-1.883	0.0738
Recovery: Mid heatwave	0.38302	0.353	0.7275
Recovery: Extreme heatwave	0.38314	0.852	0.4037

Figure 2: ZX symbiont density, using gamlss with Gamma distribution. Significance compared to pre-heat of control.

Interaction [ask sara]	Standard Error	t-value	p-value
Post-heat	0.34106	-0.496	0.62449
Recovery	0.35394	-0.663	0.51405
Mid heatwave	0.39574	-3.350	0.00291
Treatment30C	0.32308	-2.305	0.03104
Extreme heatwave	0.55161	0.788	0.43886
Post-heat: Mid heatwave	0.54261	1.914	0.06874
Post-heat : Extreme heatwave	0.46976	-0.497	0.62435
Recovery : Extreme heatwave	0.46977	0.084	0.93399

Plots for symbiont data:

```
xlab("Event") +
  ylab("Log-transformed ZC density (cells/mg)") +
  facet_grid(. ~ Treatment,
             labeller = labeller(Treatment = treatment_labels)) +
  scale_fill_manual(limits = c("Control", "25C", "30C"),
                    labels = c("Control", "Mid heatwave", "Extreme heatwave"),
                    values = cols_for_heat) +
 theme test()
green_plot <- green_plot + ylim(0,12) +</pre>
  scale_x_discrete(labels=c("11/5/2021" = "Pre-heat", "11/9/2021" = "Post-heat",
                             "11/13/2021" = "Recovery"))
dino_plot <- ggplot(Dino_Symbionts, aes(x = Date, y = log(Dino_Density),</pre>
                                        fill = Treatment)) +
  geom_boxplot(alpha = 0.85) +
  xlab("Event") +
 ylab("Log-transformed S density (cells/mg)") +
 facet_grid(. ~ Treatment,
             labeller = labeller(Treatment = treatment_labels)) +
 scale_fill_manual(limits = c("Control", "25C", "30C"),
                    labels = c("Control", "Mid heatwave", "Extreme heatwave"),
                    values = cols_for_heat) +
 theme_test()
dino_plot <- dino_plot + ylim(0,12) +</pre>
  scale_x_discrete(labels=c("11/5/2021" = "Pre-heat", "11/9/2021" = "Post-heat",
                            "11/13/2021" = "Recovery"))
density_plot = (green_plot + dino_plot) + plot_layout(ncol=1)
density_plot <- density_plot + plot_annotation(tag_levels = 'A')</pre>
density_plot
```



ggsave(here("./images/density_plot.png"), density_plot, width = 10, height = 9)

Mitotic Index Data

MI is the amount of cell division in the symbionts, calculated as a %.

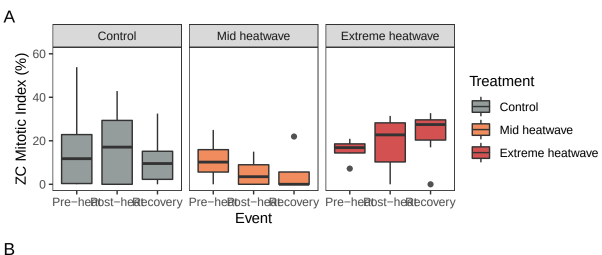
```
# Organizing the data
MI_Data <- health_data %>%
  select(-c(Event, Base_Width, Base_Length, Base_Diagonal, Base_Diameter_mm,
            Nb_Tentacles, Weight_Total_g, Weight_Tube_g, Weight_Tentacle_g,
            Green_Cells, Dino_Cells, Feeding_Time_Min, Feeding_Time, Feeding_Start,
            Feeding_Stop, Photo_ID, Removed, Fv_Fm_1, Fv_Fm_2, Fv_Fm_3, PAM_avg,
            Event_True)) %>%
  filter(Date == "11/5/2021" | Date == "11/9/2021" | Date == "11/13/2021") %>%
  mutate(Date = fct_relevel(Date, "11/5/2021", "11/9/2021", "11/13/2021")) %>%
  drop na(Green Density) %>%
  drop_na(Dino_Density) %>%
  drop_na(MI_Green) %>%
  drop_na(MI_Dino) %>%
  mutate(Acclimation_Period = as.factor(Acclimation_Period)) %>%
  mutate(MI Dino scaled = MI Dino + 0.01) %>%
  mutate(MI_Green_scaled = MI_Green + 0.01)
```

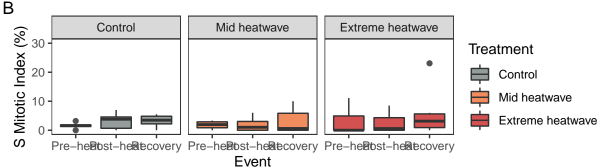
Checking Normality and Equal Variance:

```
shapiro.test(MI_Data$Green_Density)
##
##
    Shapiro-Wilk normality test
## data: MI_Data$Green_Density
## W = 0.64647, p-value = 3.82e-10
bartlett.test(Green_Density ~ Treatment, MI_Data)
##
##
   Bartlett test of homogeneity of variances
##
## data: Green_Density by Treatment
## Bartlett's K-squared = 23.091, df = 2, p-value = 9.681e-06
shapiro.test(MI_Data$Dino_Density)
##
##
    Shapiro-Wilk normality test
## data: MI_Data$Dino_Density
## W = 0.80442, p-value = 5.198e-07
bartlett.test(Dino_Density ~ Treatment, MI_Data) # p-value = 0.1483
##
## Bartlett test of homogeneity of variances
##
## data: Dino_Density by Treatment
## Bartlett's K-squared = 3.8169, df = 2, p-value = 0.1483
ZC (green) symbiont density does not meet assumptions of normality or equal variance, while ZC (dino)
does not meet assumption of normality but meets the assumption of equal variance. We will run Dunn and
Kruskal Wallis tests on the MI data to test for significant differences.
Dunn and Kruskal Wallis Test on MI Data:
kruskal.test(MI_Green ~ Treatment, data = MI_Data) # p = 0.009
##
##
    Kruskal-Wallis rank sum test
##
## data: MI_Green by Treatment
## Kruskal-Wallis chi-squared = 9.492, df = 2, p-value = 0.008686
```

```
dunnTest(MI_Green ~ Treatment, data = MI_Data) # sig. difference between 25-30
## Dunn (1964) Kruskal-Wallis multiple comparison
     p-values adjusted with the Holm method.
##
##
        Comparison
                           Z
                                 P.unadj
## 1
         25C - 30C -3.036394 0.002394262 0.007182786
## 2 25C - Control -1.441840 0.149347449 0.149347449
## 3 30C - Control 1.734539 0.082822412 0.165644824
kruskal.test(MI Dino ~ Treatment, data = MI Data) # not significant
##
##
   Kruskal-Wallis rank sum test
## data: MI_Dino by Treatment
## Kruskal-Wallis chi-squared = 0.51372, df = 2, p-value = 0.7735
dunnTest(MI_Dino ~ Treatment, data = MI_Data) # not significant
## Dunn (1964) Kruskal-Wallis multiple comparison
     p-values adjusted with the Holm method.
##
        Comparison
                                 P.unadj
                             Z
                                             P.adj
         25C - 30C -0.07275603 0.9420003 0.9420003
## 2 25C - Control -0.60896106 0.5425503 1.0000000
## 3 30C - Control -0.61859436 0.5361836 1.0000000
Plot for MI data:
green_MI_plot <- ggplot(Green_Symbionts, aes(x = Date, y = MI_Green,</pre>
                                             fill = Treatment)) +
  geom_boxplot(alpha = 0.85) +
  xlab("Event") +
  ylab("ZC Mitotic Index (%)") +
  facet_grid(. ~ Treatment,
             labeller = labeller(Treatment = treatment_labels)) +
  scale_fill_manual(limits = c("Control", "25C", "30C"),
                    labels = c("Control", "Mid heatwave", "Extreme heatwave"),
                    values = cols_for_heat) +
  theme test()
green_MI_plot <- green_MI_plot + ylim(0,60) +</pre>
  scale_x_discrete(labels=c("11/5/2021" = "Pre-heat", "11/9/2021" = "Post-heat",
                            "11/13/2021" = "Recovery"))
dino_MI_plot <- ggplot(Dino_Symbionts, aes(x = Date, y = MI_Dino,
                                           fill = Treatment)) +
  geom_boxplot(alpha = 0.85) +
```

xlab("Event") +





```
ggsave(here("./images/MI_plot.png"), MI_plot, width = 10, height = 9)
```

PAM data

We filter out aposymbiotic anemones.

```
# Organizing the data
PAM_individuals <- health_data %>%
filter(Green_Density > 1000 | Dino_Density > 1000) %>%
```

```
distinct(Species_ID)
PAM_data <- health_data %>%
  select(-c(Base_Width, Base_Length, Base_Diagonal, Base_Diameter_mm, Nb_Tentacles,
            Weight_Total_g, Weight_Tube_g, Weight_Tentacle_g, Green_Cells, Dino_Cells,
            Feeding_Time_Min, Feeding_Time, Feeding_Start, Feeding_Stop, Photo_ID,
            Removed, Div_Green, Div_Dino, MI_Green, MI_Dino, Acclimation_Period,
            Green Density, Dino Density)) %>%
  drop_na(Time_Point) %>%
  drop_na(PAM_avg) %>%
  group_by(Treatment, Date) %>%
  mutate(Mean_PAM = mean(PAM_avg), SE_PAM = std.error(PAM_avg))
filtered_PAM_data = PAM_individuals %>%
  left_join(PAM_data) %>%
  filter(Date == "11/6/2021" | Date == "11/9/2021" | Date == "11/13/2021") %>%
  filter(Time_Point != 4) %>%
 mutate(Date = fct_relevel(Date, "11/6/2021", "11/9/2021", "11/13/2021"))
```

Checking Normality and Equal Variance:

```
##
## Shapiro-Wilk normality test
##
## data: filtered_PAM_data$PAM_avg
## W = 0.84753, p-value = 3.465e-08

bartlett.test(PAM_avg ~ Treatment, data = filtered_PAM_data)

##
## Bartlett test of homogeneity of variances
##
## data: PAM_avg by Treatment
## Bartlett's K-squared = 21.408, df = 2, p-value = 2.245e-05
```

Our data does not meet the assumptions of normality and equal variance (p < 0.05) and therefore we will use Dunn and Kruskal Wallis tests to test for significant differences.

```
### ## Kruskal-Wallis rank sum test
## ## data: PAM_avg by Treatment
## Kruskal-Wallis chi-squared = 8.0406, df = 2, p-value = 0.01795

dunnTest(PAM_avg ~ Treatment, data = filtered_PAM_data)
```

```
## Comparison Z P.unadj P.adj

## 1 25C - 30C -2.6353649 0.008404688 0.02521406

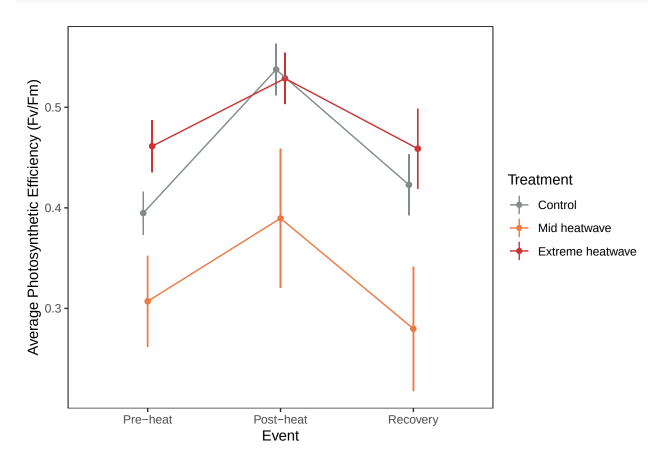
## 2 25C - Control -0.6723531 0.501358906 0.50135891

## 3 30C - Control 2.1391425 0.032424133 0.06484827

# significance only between 25 and 30
```

Plot for PAM:

```
PAM_over_time <- ggplot(data = filtered_PAM_data, aes(x=Date, y = Mean_PAM,
                                                       colour = Treatment,
                                                       group = Treatment)) +
  geom point(position = position dodge(width = 0.1)) +
  geom_linerange(aes(ymin = Mean_PAM - SE_PAM,
                     ymax = Mean_PAM + SE_PAM), position = position_dodge(width = 0.1)) +
  geom_line(position = position_dodge(width = 0.1)) +
  labs(x="Event",
       y = "Average Photosynthetic Efficiency (Fv/Fm)") +
  scale_colour_manual(limits = c("Control", "25C", "30C"),
                    labels = c("Control", "Mid heatwave", "Extreme heatwave"),
                    values = cols_for_heat) +
  theme_test()
PAM_over_time <- PAM_over_time +</pre>
  scale_x_discrete(labels=c("11/6/2021" = "Pre-heat", "11/9/2021" = "Post-heat",
                            "11/13/2021" = "Recovery"))
PAM_over_time
```



```
ggsave(here("./images/PAM_over_time.png"), PAM_over_time)
```

Size Data

Measuring the size of the base as a proxy for weight.

Checking for Normality and Equal Variance:

```
shapiro.test(Size_Data$Base_Diameter_mm)

##

## Shapiro-Wilk normality test

##

## data: Size_Data$Base_Diameter_mm

## W = 0.94517, p-value = 0.0002429

bartlett.test(Base_Diameter_mm ~ Treatment, data = Size_Data)

##

## Bartlett test of homogeneity of variances

##

## data: Base_Diameter_mm by Treatment

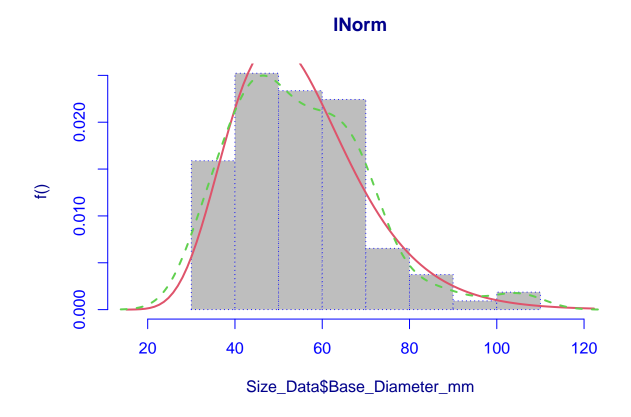
## Bartlett's K-squared = 3.8061, df = 2, p-value = 0.1491
```

The distributions do not meet assumptions of normality but does meet the assumption of equal variance.

Checking Distributions of Symbiont Data: Determining if the data fits other distributions.

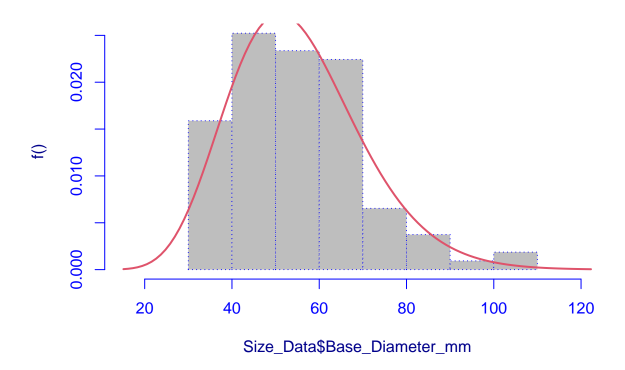
```
##
##
## Test for exponentiality based on a transformation to uniformity
##
## data: Size_Data$Base_Diameter_mm
## T = -11.277, p-value < 2.2e-16</pre>
```

```
gamma_test(Size_Data$Base_Diameter_mm)
                                                #p = 0.1874
##
##
   Test of fit for the Gamma distribution
## data: Size_Data$Base_Diameter_mm
## V = 1.8642, p-value = 0.1874
lnorm_test(Size_Data$Base_Diameter_mm)
                                                #p = 0.4117
##
## Test for the lognormal distribution based on a transformation to
## normality
## data: Size_Data$Base_Diameter_mm
## p-value = 0.4117
normal_test(Size_Data$Base_Diameter_mm)
                                                #p-value < 0.05
##
## Correlation test for normality
##
## data: Size_Data$Base_Diameter_mm
## R = 0.98889, p-value = 0.001424
## alternative hypothesis: Size_Data$Base_Diameter_mm does not follow a normal distribution.
weibull_test(Size_Data$Base_Diameter_mm)
                                                \#p-value < 0.05
##
## Test for the Weibull distribution
##
## data: Size_Data$Base_Diameter_mm
## p-value < 2.2e-16
# Run lnorm since it has the higest p value
Size_lnorm <- histDist(Size_Data$Base_Diameter_mm, "LOGNO", density = T, main = "lNorm")
# Choosing the best distribution by comparing AIC values
Size_lnorm <- histDist(Size_Data$Base_Diameter_mm, "LOGNO", density = T, main = "lNorm")
```



#AIC = 880.625
Feeding_Gamma <- histDist(Size_Data\$Base_Diameter_mm, "GA", density = F, main = "Gamma")</pre>

Gamma



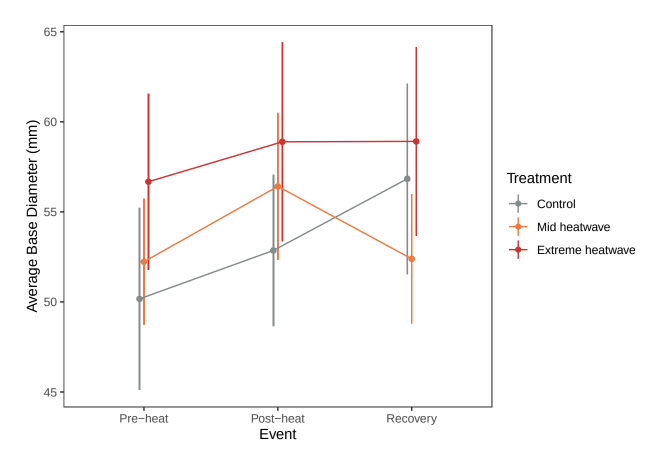
```
#AIC = 883.161
```

Running Models: We are going to run a lNorm model on our size data because the p value is larger and AIC value is smaller.

```
##
     random(Species_ID) + random(Field_Site) + random(Acclimation_Period),
##
     family = LOGNO(), data = Size_Data, control = gamlss.control(n.cyc = 4))
##
## Fitting method: RS()
## -----
## Mu link function: identity
## Mu Coefficients:
                     Estimate Std. Error t value Pr(>|t|)
##
## (Intercept)
                     ## Date11/9/2021
                     0.034371 0.026852 1.280 0.205309
                     ## Date11/13/2021
                     0.039133 0.026852 1.457 0.150054
## Treatment25C
## Treatment30C
                     ## Date11/9/2021:Treatment25C 0.038217 0.037560 1.018 0.312859
## Date11/9/2021:Treatment30C -0.004105 0.037560 -0.109 0.913320
## Date11/13/2021:Treatment30C -0.055277 0.037560 -1.472 0.146148
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
##
## Sigma link function: log
## Sigma Coefficients:
          Estimate Std. Error t value Pr(>|t|)
## Signif. codes: 0 '*** 0.001 '** 0.01 '* 0.05 '.' 0.1 ' 1
## -----
## NOTE: Additive smoothing terms exist in the formulas:
## i) Std. Error for smoothers are for the linear effect only.
## ii) Std. Error for the linear terms maybe are not accurate.
## No. of observations in the fit: 107
## Degrees of Freedom for the fit: 44.92919
##
      Residual Deg. of Freedom: 62.07081
##
                 at cycle: 4
## Global Deviance:
                 566.2585
          AIC:
                 656.1169
          SBC:
                 776.2049
##
```

The control and 25 did have significant values, but the 30 did not.

Plot for Size:



```
ggsave(here("./images/size_plot.png"), size_plot)
```

Saving 6.5×4.5 in image

Feeding Data

This is the feeding time before and after heatwaves.

Checking for Normality and Equal Variance:

```
shapiro.test(Feeding_Data$Feeding_Time_Min)

##

## Shapiro-Wilk normality test

##

## data: Feeding_Data$Feeding_Time_Min

## W = 0.62812, p-value = 1.973e-05

bartlett.test(Feeding_Time_Min ~ Treatment, data = Feeding_Data)

##

## Bartlett test of homogeneity of variances

##

## data: Feeding_Time_Min by Treatment

## Bartlett's K-squared = 9.0434, df = 2, p-value = 0.01087
```

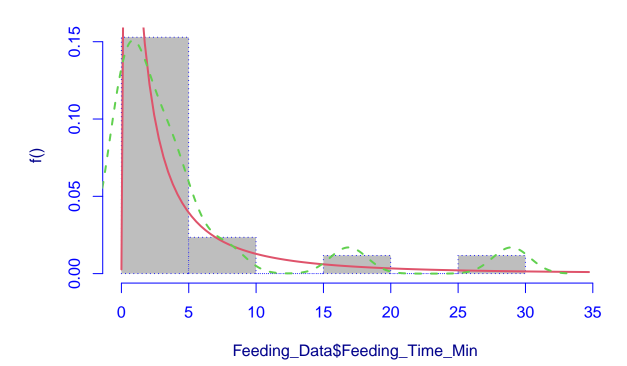
Distribution is not normal but meets the assumption of equal variance. We will check if it fits other distributions.

Checking Distributions: Determining if the data fits other distributions.

```
exp_test(Feeding_Data$Feeding_Time_Min)
                                                   \#p-value < 0.05
##
   Test for exponentiality based on a transformation to uniformity
##
## data: Feeding_Data$Feeding_Time_Min
## T = 2.048, p-value = 0.004
gamma_test(Feeding_Data$Feeding_Time_Min)
                                                   #p = 0.3717
##
## Test of fit for the Gamma distribution
##
## data: Feeding_Data$Feeding_Time_Min
## V = 0.86695, p-value = 0.5399
lnorm_test(Feeding_Data$Feeding_Time_Min)
                                                   #p = 0.6937
```

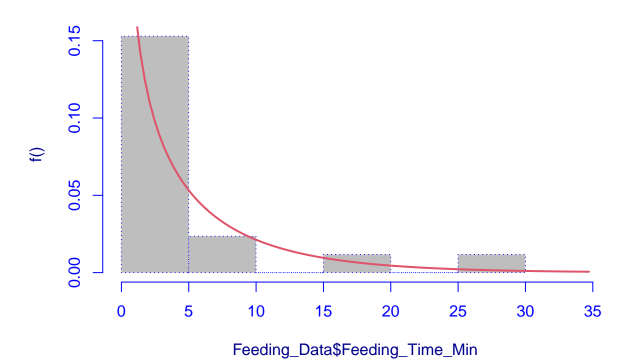
```
##
## Test for the lognormal distribution based on a transformation to
## normality
##
## data: Feeding_Data$Feeding_Time_Min
## p-value = 0.7746
normal_test(Feeding_Data$Feeding_Time_Min)
                                                   #p-value < 0.05
##
##
   Correlation test for normality
## data: Feeding_Data$Feeding_Time_Min
## R = 0.88533, p-value = 0.0001004
## alternative hypothesis: Feeding_Data$Feeding_Time_Min does not follow a normal distribution.
weibull_test(Feeding_Data$Feeding_Time_Min)
                                                   #p = 0.088
##
## Test for the Weibull distribution
## data: Feeding_Data$Feeding_Time_Min
## p-value = 0.13
# Choosing the best distribution by comparing AIC value
Feeding_lnorm <- histDist(Feeding_Data$Feeding_Time_Min, "LOGNO",</pre>
                          density = T, main = "lNorm") #AIC = 112.738
```

INorm

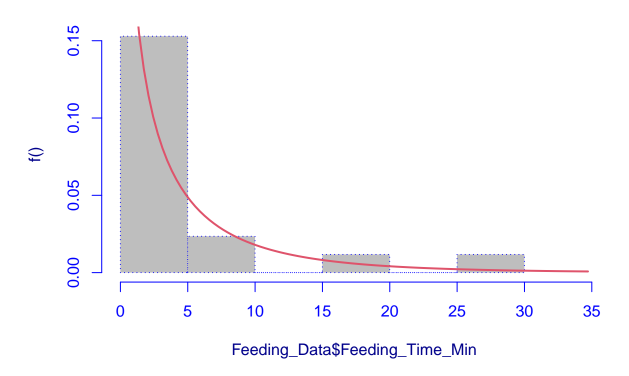


Feeding_Gamma <- histDist(Feeding_Data\$Feeding_Time_Min, "GA", density = F, main = "Gamma") #AIC = 116.789

Gamma



Weibull

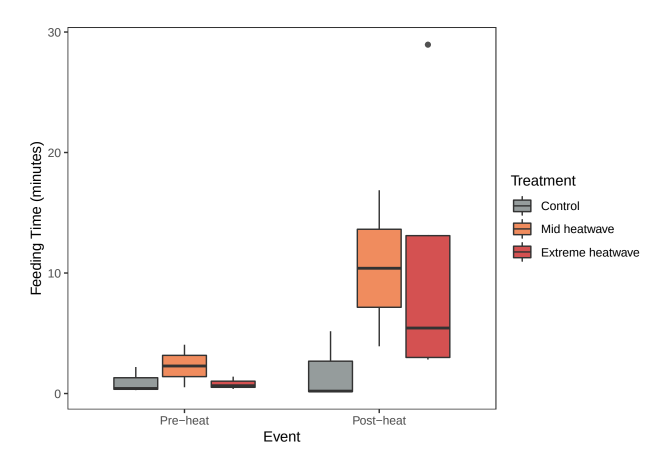


Running model: We are going to use the lnorm distribution because the p value is larger and AIC value is smaller.

```
Feeding_Data$orderTreatment = ordered(Feeding_Data$Treatment,
                                   levels = c("Control", "25C", "30C"))
Feeding_lNorm_model <- gamlss(formula = Feeding_Time_Min ~ Date*Treatment +
                             random(Species_ID) + random(Field_Site) +
                             random(Acclimation_Period),
                           family = LOGNO(), data = Feeding_Data,
                           control = gamlss.control(n.cyc = 3))
## GAMLSS-RS iteration 1: Global Deviance = 60.3695
## GAMLSS-RS iteration 2: Global Deviance = 60.3691
summary(Feeding_lNorm_model) # significance is only in post-heat 30C
## Family: c("LOGNO", "Log Normal")
##
## Call: gamlss(formula = Feeding_Time_Min ~ Date * Treatment +
##
      random(Species_ID) + random(Field_Site) + random(Acclimation_Period),
##
      family = LOGNO(), data = Feeding_Data, control = gamlss.control(n.cyc = 3))
##
##
```

```
## Fitting method: RS()
##
## -----
## Mu link function: identity
## Mu Coefficients:
                    Estimate Std. Error t value Pr(>|t|)
##
## (Intercept)
                     -0.5298 0.5149 -1.029 0.3330
                      -0.2449 0.7281 -0.336 0.7451
## Date11/9/2021
## Treatment25C
                      0.8233
                            0.8141 1.011 0.3409
## Treatment30C
                      0.3369 0.7281 0.463 0.6557
## Date11/9/2021:Treatment25C 1.9684
                             1.1513 1.710 0.1249
## Date11/9/2021:Treatment30C 2.3532
                            0.9970 2.360 0.0453 *
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## -----
## Sigma link function: log
## Sigma Coefficients:
          Estimate Std. Error t value Pr(>|t|)
## (Intercept) -0.1145 0.1715 -0.668 0.523
##
## -----
## NOTE: Additive smoothing terms exist in the formulas:
## i) Std. Error for smoothers are for the linear effect only.
## ii) Std. Error for the linear terms maybe are not accurate.
## -----
## No. of observations in the fit: 17
## Degrees of Freedom for the fit: 8.831933
      Residual Deg. of Freedom: 8.168067
##
                 at cycle: 2
##
## Global Deviance:
                 60.36911
##
          AIC:
                78.03298
          SBC:
##
                 85.39186
```

Plot for Feeding:

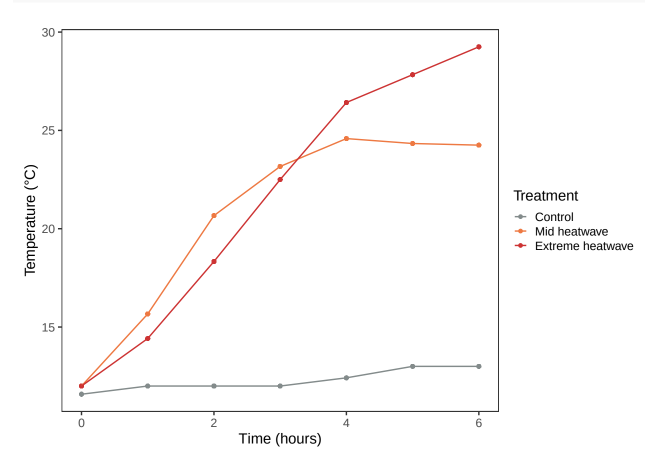


```
ggsave(here("./images/feeding_plot.png"), feeding_plot)
```

Heat Data

Includes temperature during heatwave and behavioral responses (open vs closed)

Temperature over Time Graph:



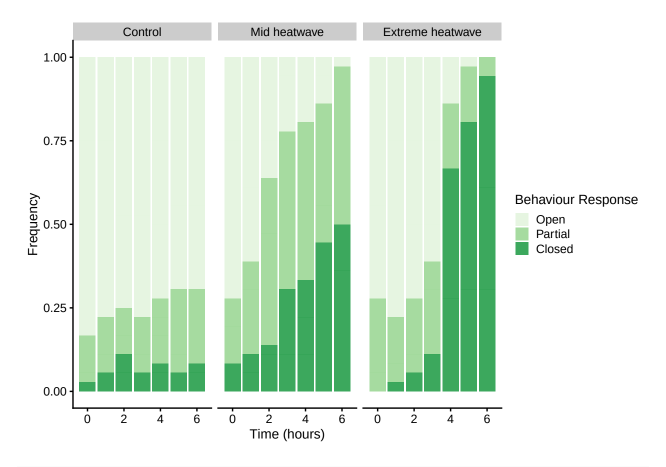
```
ggsave(here("./images/heat_plot.png"), heat_plot)
```

Ordinal Regression

Model to test significance on the open/partial/closed data.

```
## Cumulative Link Mixed Model fitted with the Laplace approximation
##
## formula: Open_Closed ~ Treatment + (1 | Bucket) + (1 | Species_ID)
## data: heat_data
##
## link threshold nobs logLik AIC niter max.grad cond.H
## logit flexible 756 -682.12 1376.25 220(930) 2.31e-05 1.1e+02
```

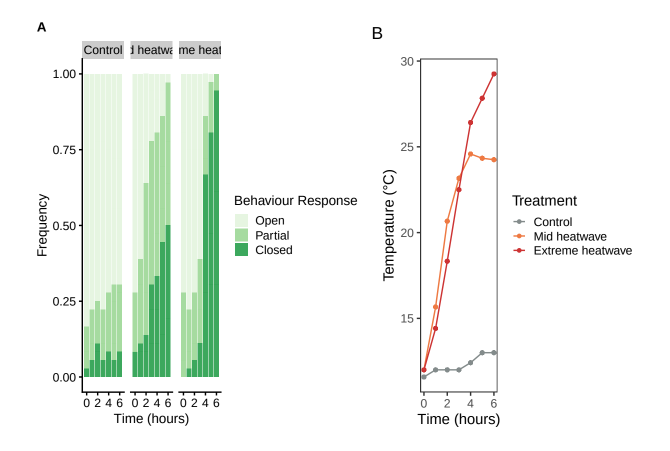
```
##
## Random effects:
## Groups
                          Variance Std.Dev.
## Species_ID (Intercept) 1.069e+00 1.034e+00
               (Intercept) 1.190e-11 3.450e-06
## Number of groups: Species_ID 36, Bucket 12
## Coefficients:
##
               Estimate Std. Error z value Pr(>|z|)
## Treatment25C 2.0438
                            0.4768 4.286 1.82e-05 ***
## Treatment30C 1.9778
                            0.4777 4.140 3.47e-05 ***
## Signif. codes: 0 '*** 0.001 '** 0.01 '* 0.05 '.' 0.1 ' ' 1
##
## Threshold coefficients:
##
                 Estimate Std. Error z value
## Open|Partial
                   1.3765
                              0.3492
                                       3.941
## Partial|Closed
                   2.9049
                              0.3613 8.040
Open/Closed Graph:
cols_for_behaviour = c("Open" = "olivedrab3", "Partial" = "forestgreen", "Closed" = "darkgreen")
behaviour_data <- heat_data %>%
  group_by(Day, Treatment, Time_Block) %>%
  count(Open Closed)
behaviour_plot <- ggplot(data = behaviour_data, aes(x = Time_Block, y = n, fill = Open_Closed)) +
  geom_bar(alpha = 0.95, position="fill", stat="identity") +
  facet_grid(. ~ Treatment,
            labeller = labeller(Treatment = treatment_labels)) +
 xlab("Time (hours)") +
 ylab("Frequency") +
  scale_fill_brewer(palette = "Greens") +
  theme_cowplot(10)
behaviour_plot <- behaviour_plot + labs(fill = "Behaviour Response")
behaviour_plot
```



```
ggsave(here("./images/behaviour_plot.png"), behaviour_plot, width = 10, height = 6)
```

Combining plots together for comparision:

```
behaviour_heat <- behaviour_plot + heat_plot
behaviour_heat <- behaviour_heat + plot_layout(ncol=2, widths = c(1.8,1))
behaviour_heat <- behaviour_heat + plot_annotation(tag_levels = 'A')
behaviour_heat</pre>
```



ggsave(here("./images/behaviour_heat.png"), behaviour_heat, width = 10, height = 4)