## preliminary analyses blending groups

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## Data Grouping and Exclusion Rationale

The following groups were combined because there were no significant differences between them:

- 10°C and 15°C temperature treatments
- Lemna minor strains from a US lab and field collection
- Preparation methods with and without repeated first-daughter separation

Exception: At 27°C, there were significant differences in performance between preparation methods. Therefore, the six replicates that did not include first-daughter separation were excluded from the analysis.

```
library(lme4)
library(performance)
library(sjPlot)
## Testing for differences in preparation methods, average temperatures,
## and strains to justify merging of groups performed during data analysis
### This dataset contains replicates for which a preparation technique was performed (repeated first bo
original_dataset_2 <- read.csv("https://raw.githubusercontent.com/Cuddington-Lab/thermal-experiments/ma
original_dataset_2$prep <- rep("yes",times=length(original_dataset_2$Experiment_Number))
### This dataset contains replicates of experiments performed without a preparation technique
original dataset 1 <- read.csv("https://raw.githubusercontent.com/Cuddington-Lab/thermal-experiments/ma
original_dataset_1$prep <- rep("no",times=length(original_dataset_1$Experiment_Number))
datin <- rbind(original_dataset_1,original_dataset_2)</pre>
datin <- datin[!(datin$Treatment == 0 & (datin$Obs_sd <= 2.1 | datin$Obs_sd >= 2.9))
               &!(datin$Treatment == 0.95 & (datin$Obs_sd <= 2.1 | datin$Obs_sd >= 2.9)),]
datin <- datin[!(datin$Treatment == 0 & (datin$Obs_ac <= -0.2 | datin$Obs_ac >= 0.2))
               &!(datin$Treatment == 0.95 & (datin$0bs_ac <= 0.92 | datin$0bs_ac >= 0.98)),]
datin <- subset(datin, !Errors == "y"|is.na(Errors))</pre>
datin <- subset(datin, !Treatment == "constant")</pre>
colnames(datin)[6] <- "Label"</pre>
table(datin$Label, datin$cat_1)
##
##
                  N N/A P
##
   0
              66 0 16 0
```

```
0 78 0 73
##
     0.95
##
     constant 0 0 0 0
levels(datin$cat_1) = c("no autocorrelation", "hot-cold", "no autocorrelation", "cold-hot")
datin$Treatment<-paste0(datin$cat_1)</pre>
### Relabel groups to simplify visualization of results
library(stringr)
datin$Label <- str_replace(datin$Label, "0.95", "strong autocorrelation")
datin$Label <- str_replace(datin$Label, "0", "no autocorrelation")</pre>
datin$Exp_run <- str_sub(datin$Profile_name,-2,-1)</pre>
datin$Exp_run <- gsub('_','',datin$Exp_run)</pre>
datin$Exp_run <- as.numeric(datin$Exp_run)</pre>
# Subset data to keep only those Exp_run that have at least 3 occurrences within each Mean_Temperature
datin <- datin[</pre>
  ave(seq_along(datin$Exp_run),
      interaction(datin$Exp_run, datin$Mean_temperature, datin$Species),
      FUN = length) >= 3,
]
datin$total_living_fronds <- (datin$Frond_count_1 + datin$Frond_count_2 + datin$Frond_count_3)</pre>
# Datasets and variables to iterate over
datin1015 <- subset(datin, (Species == "Lab_LM" | Species == "Field_LM") &</pre>
                       (Mean_temperature == 10 | Mean_temperature == 15))
datin27 <- subset(datin, (Species == "Lab_LM" | Species == "Field_LM") &</pre>
                     (Mean_temperature == 27))
datasets <- list(datin1015 = datin1015, datin27 = datin27)</pre>
predictors_list <- list(</pre>
 datin1015 = c("Mean_temperature*Treatment", "prep*Treatment"),
  datin27 = c("prep*Treatment")
)
# Initialize lists to save results
best models <- list()</pre>
model_diagnostics <- list()</pre>
# Loop over datasets and predictors
for (data_name in names(datasets)) {
  data <- datasets[[data name]]</pre>
  predictors <- predictors_list[[data_name]]</pre>
 for (pred in predictors) {
    # Fit models
    simple <- glm(as.formula(paste("total_living_fronds ~", pred)), data = data, family="poisson")</pre>
    exp_number <- glmer(as.formula(paste("total_living_fronds ~", pred, "+ (1|Exp_run)")), data = data,
    # Store candidate models
    Cand.modsF <- list("no random effects" = simple,</pre>
                        "experiment number" = exp_number)
```

```
# Compare models
    compareF <- compare_performance(Cand.modsF)</pre>
    # Select best model and save it
   best_model_index <- which.min(compareF$AIC)</pre>
   best_model <- Cand.modsF[[best_model_index]]</pre>
    # Save best model and its diagnostics
   best_models[[paste(data_name, pred)]] <- best_model</pre>
}}
#at 10-15C, no differences between average temperature and preparation method
summary(best_models[[1]])
##
## Call:
## glm(formula = as.formula(paste("total_living_fronds ~", pred)),
       family = "poisson", data = data)
##
## Coefficients:
##
                                                 Estimate Std. Error z value
## (Intercept)
                                                 2.987619 0.309239 9.661
                                                 -0.008150 0.023784 -0.343
## Mean_temperature
## Treatmenthot-cold
                                                 0.153296   0.434862   0.353
                                                 -0.179723
## Treatmentno autocorrelation
                                                             0.445021 -0.404
## Mean_temperature:Treatmenthot-cold
                                                -0.011498
                                                             0.033533 -0.343
## Mean_temperature:Treatmentno autocorrelation 0.009838 0.034148 0.288
                                                Pr(>|z|)
## (Intercept)
                                                   <2e-16 ***
                                                    0.732
## Mean_temperature
## Treatmenthot-cold
                                                    0.724
                                                    0.686
## Treatmentno autocorrelation
## Mean temperature:Treatmenthot-cold
                                                    0.732
## Mean_temperature:Treatmentno autocorrelation
                                                   0.773
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## (Dispersion parameter for poisson family taken to be 1)
##
       Null deviance: 15.024 on 47 degrees of freedom
## Residual deviance: 13.597 on 42 degrees of freedom
## AIC: 251.52
## Number of Fisher Scoring iterations: 4
summary(best_models[[2]])
##
## glm(formula = as.formula(paste("total_living_fronds ~", pred)),
       family = "poisson", data = data)
##
##
## Coefficients:
##
                                       Estimate Std. Error z value Pr(>|z|)
```

```
## (Intercept)
                                         2.86479
                                                    0.07198 39.799
                                                                       <2e-16 ***
                                         0.05837
                                                    0.12623 0.462
                                                                        0.644
## prepyes
## Treatmenthot-cold
                                        -0.02094
                                                    0.10233 -0.205
                                                                        0.838
## Treatmentno autocorrelation
                                        -0.05868
                                                    0.10332 -0.568
                                                                        0.570
## prepyes:Treatmenthot-cold
                                         0.08346
                                                    0.17699
                                                              0.472
                                                                        0.637
## prepyes:Treatmentno autocorrelation 0.01472
                                                    0.18073
                                                              0.081
                                                                        0.935
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
##
## (Dispersion parameter for poisson family taken to be 1)
##
##
       Null deviance: 15.024 on 47 degrees of freedom
## Residual deviance: 12.592 on 42 degrees of freedom
## AIC: 250.52
##
## Number of Fisher Scoring iterations: 4
#at 10-15C, no differences between duckweed strains (lab and field LM)
exp_number <- glmer(total_living_fronds ~ Species * Treatment + (1|Exp_run), data = datin1015, family="
simple <- glm(total_living_fronds ~ Species * Treatment, data = datin1015, family="poisson")</pre>
tab_model(simple)
total_living_fronds
Predictors
Incidence Rate Ratios
CI
р
(Intercept)
17.55
15.19 - 20.14
< 0.001
Species [Field_LM]
1.06
0.82 - 1.35
0.644
Treatment [hot-cold]
0.98
0.80 - 1.20
0.838
Treatment [noautocorrelation]
0.94
0.77 - 1.15
0.570
```

```
Species [Field LM] ×Treatment [hot-cold]
1.09
0.77 - 1.54
0.637
Species [Field_LM] ×Treatment [noautocorrelation]
1.01
0.71 - 1.45
0.935
Observations
48
R2 Nagelkerke
0.184
#at 27C, preparation method makes a difference (excluding older replicates)
summary(best_models[[3]])
## Generalized linear mixed model fit by maximum likelihood (Laplace
     Approximation) [glmerMod]
  Family: poisson (log)
## Formula: total_living_fronds ~ prep * Treatment + (1 | Exp_run)
##
     Data: data
##
##
        AIC
                       logLik deviance df.resid
                 BIC
##
      358.9
               371.0
                       -172.4
                                 344.9
## Scaled residuals:
                10 Median
       Min
                                3Q
                                       Max
## -1.9587 -0.7471 -0.0553 0.5738 3.6317
##
## Random effects:
## Groups Name
                        Variance Std.Dev.
## Exp_run (Intercept) 0.06156 0.2481
## Number of obs: 42, groups: Exp_run, 14
##
## Fixed effects:
##
                                         Estimate Std. Error z value Pr(>|z|)
## (Intercept)
                                        3.345e+00 2.192e-01 15.259 < 2e-16 ***
## prepyes
                                        9.872e-01 2.329e-01
                                                              4.239 2.25e-05 ***
## Treatmenthot-cold
                                        2.413e-05 1.838e-01
                                                               0.000
                                                                         1.000
## Treatmentno autocorrelation
                                        3.335e-02 1.822e-01
                                                               0.183
                                                                         0.855
## prepyes:Treatmenthot-cold
                                        6.349e-03 1.894e-01
                                                               0.034
                                                                         0.973
## prepyes:Treatmentno autocorrelation -3.122e-02 1.880e-01 -0.166
                                                                         0.868
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
##
## Correlation of Fixed Effects:
               (Intr) prepys Trtmn- Trtmna prp:T-
## prepyes
               -0.941
## Trtmntht-cl -0.419 0.394
```