warmXtrophic Project: Plant Composition Data Analyses

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Load in packages & data

```
# Clear all existing data
rm(list = ls())
# Load packages
library(tidyverse)
library(ggplot2)
library(lme4)
library(olsrr)
library(predictmeans)
library(car)
library(fitdistrplus)
library(ggpubr)
library(rstatix)
library(vegan)
library(interactions)
library(sjPlot)
library(effects)
library(glmmTMB)
# Set working directory
Sys.getenv("L1DIR")
```

[1] "/Volumes/GoogleDrive/Shared drives/SpaCE_Lab_warmXtrophic/data/L1/"

```
L1_dir <- Sys.getenv("L1DIR")
L2_dir <- Sys.getenv("L2DIR")</pre>
list.files(L1_dir)
                                                  "CN"
## [1] "ANPP"
                             "climate_data"
## [4] "greenup"
                             "herbivory"
                                                  "HOBO data"
## [7] "PAR"
                             "phenology"
                                                  "plant_composition"
## [10] "SLA"
# read in plant comp data
comp <- read.csv(file.path(L1_dir, "plant_composition/final_plantcomp_L1.csv"))</pre>
comp <- comp %>% select(-X) # get rid of 'X' column that shows up
```

```
# adding sequential year variable starting at 1: this is because 2016... are
# large numbers compare with other values in the dataset. We can always label
# axes with these real years.
comp$year_factor[comp$year == 2015] <- 1</pre>
comp$year_factor[comp$year == 2016] <- 1</pre>
comp$year_factor[comp$year == 2017] <- 2</pre>
comp$year_factor[comp$year == 2018] <- 3</pre>
comp$year_factor[comp$year == 2019] <- 4</pre>
comp$year_factor[comp$year == 2020] <- 5</pre>
# Remove non-plant data - MY: not sure if I should remove these?
comp <- comp[!(comp$species == "Bare_Ground" | comp$species == "Unknown" | comp$species ==
    "Brown" | comp$species == "Litter" | comp$species == "Vert_Litter" | comp$species ==
    "Animal_Disturbance"), ]
# calculating total composition sums - proxy for most common species
comp yearly totals <- comp %>% group by(species, site, year, plot) %>% summarize(comp sum = sum(cover,
   na.rm = T)
## 'summarise()' has grouped output by 'species', 'site', 'year'. You can override using the '.groups'
Relative abundance
# Calculating relative abundance
comp1 <- comp %>% select(species, site, year, plot, cover, state)
# calculate plot cover mean by species-plot per year
plot_mean <- aggregate(cover ~ plot * species * year * site * state, data = comp1,</pre>
   FUN = mean, na.rm = T)
names(plot_mean) [names(plot_mean) == "cover"] <- "plot_mean" # change 'cover' column name to 'plot_mea</pre>
View(plot mean)
# convert cover to relative abundance first get summed cover for all plants per
# plot
plot_cover_total = aggregate(plot_mean ~ plot * year * site * state, data = plot_mean,
   FUN = sum, na.rm = T)
names(plot_cover_total) [names(plot_cover_total) == "plot_mean"] <- "plot_cover_total"</pre>
View(plot_cover_total)
comp2 <- merge(plot_mean, plot_cover_total, by = c("plot", "year", "site", "state"))</pre>
# calculate relative percent cover per species in each quadrat (='relative
# abundance')
comp2$relabun <- comp2$plot_mean/comp2$plot_cover_total</pre>
summary(comp2)
##
                            year
                                           site
                                                             state
       plot
## Length:2472
                       Min. :2015
                                      Length:2472
                                                          Length:2472
                                     Class :character
## Class:character 1st Qu.:2016
                                                          Class :character
## Mode :character Median :2017
                                      Mode :character Mode :character
##
                       Mean :2017
##
                       3rd Qu.:2019
                              :2020
##
                       Max.
```

plot_cover_total

relabun

plot_mean

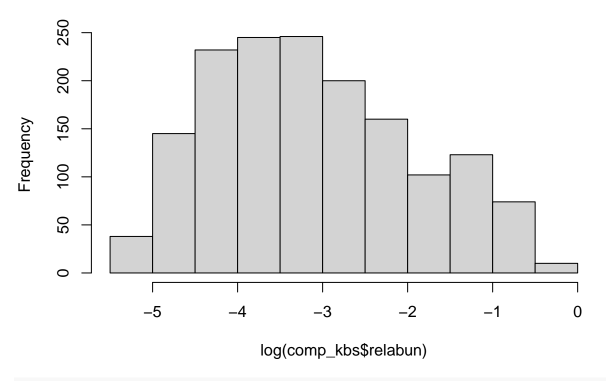
##

species

```
## Length:2472
                      Min. : 0.500
                                       Min. : 15.88
                                                          Min.
                                                                 :0.005242
## Class:character 1st Qu.: 1.816 1st Qu.: 57.75
                                                         1st Qu.:0.023009
                                                          Median :0.054084
## Mode :character Median : 4.445 Median : 85.83
##
                      Mean
                            : 9.515 Mean : 90.90
                                                          Mean
                                                                 :0.116505
##
                       3rd Qu.: 11.500
                                        3rd Qu.:124.18
                                                          3rd Qu.:0.148558
##
                      Max.
                             :110.000
                                        Max. :190.75
                                                                 :0.809170
                                                          Max.
# create dataframes for kbs and umbs - remember that these contain species within
# plots
comp kbs <- subset(comp2, site == "kbs")</pre>
comp_umbs <- subset(comp2, site == "umbs")</pre>
Relative Abundance for comparisons between native and exotic species
# getting relative % cover for comparisons between native & exotic most code from
# Kileigh's old script average sub-quadrats for plots - MY: what are sub
# quadrats? Just plots?
comp_org <- subset(comp, origin == "Exotic" | origin == "Native")</pre>
plot_mean_org <- aggregate(cover ~ plot * origin * species * year * site * state,</pre>
    data = comp_org, FUN = mean, na.rm = T)
names(plot_mean_org)[names(plot_mean_org) == "cover"] <- "plot_mean_org"</pre>
head(plot_mean_org)
    plot origin species year site
                                    state plot mean org
                 Acmi 2015 kbs ambient
## 1
      A1 Native
                                               13.625000
## 2
      A3 Native
                   Acmi 2015 kbs ambient
                                               9.111111
## 3
      A6 Native Acmi 2015 kbs ambient
                                               6.333333
## 4
      B1 Native Acmi 2015 kbs ambient
                                               6.777778
                   Acmi 2015 kbs ambient
## 5
      B4 Native
                                               15.44444
                   Acmi 2015 kbs ambient
                                               6.250000
## 6
      B5 Native
# convert cover to relative abundance first get summed cover for all plants per
# plot
cover_sum = aggregate(plot_mean_org ~ plot * origin * year * site * state, data = plot_mean_org,
    FUN = sum, na.rm = T)
names(cover sum) [names(cover sum) == "plot mean org"] <- "cover sum"</pre>
head(cover sum)
##
    plot origin year site
                            state cover_sum
      A1 Exotic 2015 kbs ambient 22.38333
      A3 Exotic 2015 kbs ambient 44.24762
## 2
## 3
      A6 Exotic 2015 kbs ambient
                                    24.02222
## 4
      B1 Exotic 2015 kbs ambient 13.00000
## 5
      B4 Exotic 2015 kbs ambient
## 6
      B5 Exotic 2015 kbs ambient 55.80000
comp2 <- merge(plot_mean_org, cover_sum, by = c("plot", "origin", "year", "site",</pre>
   "state"))
# calculate relative percent cover per species in each quadrat (='relative
# abundance')
comp2$relabun <- comp2$plot mean org/comp2$cover sum</pre>
summary(comp2)
```

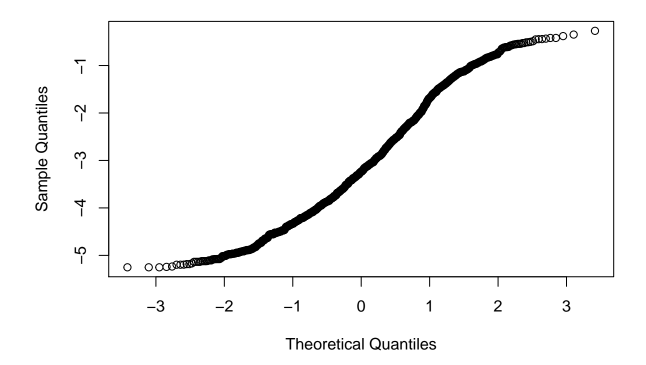
```
year
##
       plot
                         origin
                                                          site
## Length:1914
                     Length: 1914
                                        Min. :2015 Length:1914
                                        1st Qu.:2016 Class:character
## Class :character Class :character
## Mode :character Mode :character
                                        Median: 2017 Mode: character
##
                                        Mean :2017
##
                                        3rd Qu.:2019
##
                                        Max.
                                              :2020
##
                                        plot_mean_org
                                                           cover_sum
      state
                        species
## Length:1914
                      Length: 1914
                                        Min. : 0.500
                                                         Min. : 1.00
## Class :character
                      Class : character
                                        1st Qu.: 1.667
                                                        1st Qu.: 19.35
## Mode :character Mode :character
                                        Median: 4.333 Median: 30.62
                                        Mean : 9.553
##
                                                        Mean : 33.95
##
                                        3rd Qu.: 11.000 3rd Qu.: 45.46
                                        Max. :110.000 Max. :126.00
##
##
      relabun
## Min.
          :0.009009
## 1st Qu.:0.065864
## Median :0.160139
## Mean :0.299373
## 3rd Qu.:0.457948
## Max. :1.000000
# create dataframes for kbs and umbs - remember that these contain species within
# plots
comp_org_kbs <- subset(comp2, site == "kbs")</pre>
comp_org_umbs <- subset(comp2, site == "umbs")</pre>
# https://rfunctions.blogspot.com/2016/08/comparing-and-estimating-species.html
# rarefired species richness rarecurve(comp2$species)
# specpool(comp2)
# checking for normality in raw data
hist(log(comp_kbs$relabun))
```

Histogram of log(comp_kbs\$relabun)



qqnorm(log(comp_kbs\$relabun))

Normal Q-Q Plot



```
shapiro.test(log(comp_kbs$relabun))

##

## Shapiro-Wilk normality test

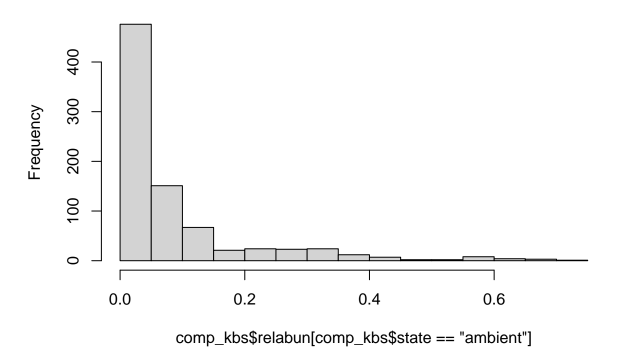
##

## data: log(comp_kbs$relabun)

## W = 0.9707, p-value < 2.2e-16

hist(comp_kbs$relabun[comp_kbs$state == "ambient"])</pre>
```

Histogram of comp_kbs\$relabun[comp_kbs\$state == "ambient"]



hist(comp_kbs\$relabun[comp_kbs\$state == "warmed"])

Histogram of comp_kbs\$relabun[comp_kbs\$state == "warmed"]

