Lingering land use legacies after different wetland restoration strategies

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Project Description: Anthropogenic legacy effects often occur as a consequence of land use change and can leave behind long-lasting changes to ecosystem structure and function. These physical, chemical, and biological changes can linger and interact with contemporary management to impact restoration outcomes. In the present study, we evaluated how restoration strategy can overcome the lingering effects of land-use in wetlands.

Initial Setup

```
rm(list=ls())
setwd("~/GitHub/IL_Wetlands/analyses")
se <- function(x, ...){sd(x, na.rm = TRUE)/sqrt(length(na.omit(x)))}</pre>
ci \leftarrow function(x, ...)\{1.96 * sd(x,na.rm = TRUE)\}
# Code Dependencies
source("../bin/DiversityFunctions.R")
source("../bin/MothurTools.R")
require("vegan")
require("reshape")
require("ggplot2")
require("nlme")
require("ade4")
require("grid"); require("png")
require("ape"); require("picante")
library("agricolae")
myColors <- c("#448844", "#33CC33", "#CCFF00", "#FFF000", "#FF9933", "#A9A9A9")
names(myColors) <- c("BalledBurlapped", "Bareroot", "Seedling",</pre>
                      "Acorn", "Seedbank", "Reference")
```

Import Data Files

Experimental Design File

```
# "Seedbank", "Reference")

# Note: The sample names in the microbial part were incorrect (skipped 72)

# They are correct in this design file and will be corrected for each below
```

Microbial Data

```
# Import OTU data
# Import Raw Data
WLdata.in <- read.otu("../data/WL.final.shared")</pre>
# Remove Mock Community
WLdata.in2 <- WLdata.in[which(rownames(WLdata.in) != "Mock"), ]</pre>
# Correct Sample IDs
all.equal(rownames(WLdata.in2), rownames(design))
## [1] "19 string mismatches"
rownames(WLdata.in2) <- rownames(design)</pre>
# Remove OTUs with less than two occurences across all sites
WLdata <- WLdata.in2[, which(colSums(WLdata.in2) >= 2)]
# Make Presence Absence Matrix
WLdataPA <- (WLdata > 0) * 1
# Make Relative Abundence Matrices
WLdataREL <- WLdata
for(i in 1:dim(WLdata)[1]){
 WLdataREL[i,] <- WLdata[i,]/sum(WLdata[i,])</pre>
}
# Log Transform Relative Abundances
WLdataREL.log <- decostand(WLdataREL, method="log")</pre>
## Warning: non-integer data: divided by smallest positive value
# Import Taxonomy File
WL.tax <- read.tax(taxonomy = "../data/WL.final.0.03.taxonomy",
                   format = "rdp", tax.levels = 6, col.tax = 3)
```

Plant Data

Phylogenetic Tree and UniFrac Distance Matrix

Notes: Phylogenetic Analysis

```
# The following was done outside of R
python ../bin/name_change.py WL.final.0.03.rep.fasta WL.final.0.03.rep.rename.fasta
FastTree -gtr -nt -gamma -fastest WL.final.0.03.rep.rename.fasta > WL.bac.tree
Output:
ML-NNI round 11: LogLk = -1017514.896 NNIs 4825 max delta 3.21 Time 626.69 (final)
Optimize all lengths: LogLk = -1017490.876 Time 645.88
Gamma(20) LogLk = -1017848.611 alpha = 2.130 rescaling lengths by 1.471
Total time: 733.31 seconds Unique: 56413/56413
Bad splits: 16/56410 Worst delta-LogLk 3.347
# Weighted UniFrac was done using Mothur
This caused an error because of the names. Mothur actually crashed
FastTree -gtr -nt -gamma -fastest WL.final.0.03.rep.fasta > WL.bac.tree
ML-NNI round 12: LogLk = -1017096.874 NNIs 4848 max delta 3.23
Time 633.03 (final)x delta 3.226)
Optimize all lengths: LogLk = -1017090.230 Time 650.74
Gamma(20) LogLk = -1017412.647 alpha = 2.058 rescaling lengths by 1.484
Total time: 735.70 seconds Unique: 56413/56413
Bad splits: 13/56410 Worst delta-LogLk 1.738
Mothur (v 1.38)
unifrac.weighted(tree=WL.bac.tree, count=WL.final.rep.count_table, distance=square)
Output File Names:
WL.bac.treewsummary
WL.bac.tree1.weighted.phylip.dist
```

Soil Data

```
WL.soil <- read.csv("../data/WL_plant_soil.csv")
row.names(WL.soil) <- WL.soil$Sample_Code</pre>
```

Soil Factors

```
# Organize Data
WL.trts <- WL.soil[, "Treatment"]</pre>
WL.soil.phys <- WL.soil[, which(colnames(WL.soil) %in% c("Moisture",
                                                      "Temp", "pH"))]
WL.soil.nuts <- WL.soil[, which(colnames(WL.soil) %in% c("TOC", "TN",
                                                      "OM", "NH4.N", "NO3.N"))]
# Linear Models
moisture.lm <- lm(Moisture ~ Treatment, data=WL.soil)</pre>
anova(moisture.lm)
## Analysis of Variance Table
## Response: Moisture
            Df Sum Sq Mean Sq F value
## Treatment 5 255.28 51.055 4.8689 0.0005844 ***
## Residuals 84 880.83 10.486
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
summary(moisture.lm)
##
## Call:
## lm(formula = Moisture ~ Treatment, data = WL.soil)
## Residuals:
##
      Min
               1Q Median
                              3Q
                                    Max
## -7.0653 -2.0458 -0.0647 1.8072 9.7347
## Coefficients:
##
                    Estimate Std. Error t value Pr(>|t|)
## (Intercept)
                    ## TreatmentBareroot -2.68733 1.18243 -2.273 0.0256 *
## TreatmentSeedling -2.82333 1.18243 -2.388 0.0192 *
## TreatmentAcorn
                    -0.78333 1.18243 -0.662 0.5095
## TreatmentSeedbank -0.07667 1.18243 -0.065 0.9485
## TreatmentReference 2.07067
                               1.18243
                                        1.751 0.0836 .
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## Residual standard error: 3.238 on 84 degrees of freedom
## Multiple R-squared: 0.2247, Adjusted R-squared: 0.1785
## F-statistic: 4.869 on 5 and 84 DF, p-value: 0.0005844
```

```
HSD <- HSD.test(moisture.lm, "Treatment", console=TRUE)</pre>
## Study: moisture.lm ~ "Treatment"
## HSD Test for Moisture
## Mean Square Error: 10.48604
## Treatment, means
##
                   Moisture
                                 std r
                                          Min
## Acorn
                   31.72133 2.359243 15 26.57 35.57
## BalledBurlapped 32.50467 2.623563 15 27.52 36.71
                  29.81733 2.715018 15 26.70 34.69
## Bareroot
## Reference
                  34.57533 5.581024 15 27.51 44.31
                  32.42800 2.095302 15 29.06 37.33
## Seedbank
## Seedling
                   29.68133 2.749122 15 24.98 34.28
##
## alpha: 0.05; Df Error: 84
## Critical Value of Studentized Range: 4.124617
## Honestly Significant Difference: 3.448607
## Means with the same letter are not significantly different.
## Groups, Treatments and means
         Reference
                             34.58
         {\tt BalledBurlapped}
                             32.5
## ab
         Seedbank
                             32.43
## ab
## ab
         Acorn
                             31.72
## b
        Bareroot
                             29.82
## b
         Seedling
                             29.68
pH.lm <- lm(pH ~ Treatment, data=WL.soil)
anova(pH.lm)
## Analysis of Variance Table
## Response: pH
             Df
                  Sum Sq Mean Sq F value
## Treatment 5 0.055259 0.011052 3.7149 0.004362 **
## Residuals 84 0.249896 0.002975
## Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
summary(pH.lm)
##
## lm(formula = pH ~ Treatment, data = WL.soil)
## Residuals:
         Min
                    1Q
                          Median
                                        3Q
                                                 Max
## -0.135000 -0.024792 0.008333 0.025625 0.125833
```

```
##
## Coefficients:
##
                     Estimate Std. Error t value Pr(>|t|)
                      7.52417 0.01408 534.274 < 2e-16 ***
## (Intercept)
## TreatmentBareroot -0.00750
                                 0.01992 -0.377 0.70744
## TreatmentSeedling 0.05917 0.01992
                                          2.971 0.00387 **
## TreatmentAcorn
                      0.05083
                               0.01992
                                          2.552 0.01251 *
## TreatmentSeedbank
                      0.01750
                                 0.01992
                                           0.879 0.38208
## TreatmentReference 0.03583
                                 0.01992
                                           1.799 0.07558 .
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## Residual standard error: 0.05454 on 84 degrees of freedom
## Multiple R-squared: 0.1811, Adjusted R-squared: 0.1323
## F-statistic: 3.715 on 5 and 84 DF, p-value: 0.004362
HSD <- HSD.test(pH.lm, "Treatment", console=TRUE)</pre>
##
## Study: pH.lm ~ "Treatment"
## HSD Test for pH
##
## Mean Square Error: 0.00297495
## Treatment, means
##
##
                                  std r
                        рΗ
                  7.575000 0.05088502 15 7.450 7.65
## Acorn
## BalledBurlapped 7.524167 0.04735567 15 7.450 7.65
## Bareroot
               7.516667 0.06315137 15 7.400 7.60
## Reference
                  7.560000 0.05809475 15 7.425 7.65
## Seedbank
                  7.541667 0.05643159 15 7.450 7.65
## Seedling
                  7.583333 0.04970149 15 7.450 7.65
##
## alpha: 0.05; Df Error: 84
## Critical Value of Studentized Range: 4.124617
##
## Honestly Significant Difference: 0.05808685
## Means with the same letter are not significantly different.
##
## Groups, Treatments and means
## a
        Seedling
                            7.583
## ab
         Acorn
                            7.575
        Reference
## abc
                            7.56
## abc
        Seedbank
                            7.542
## bc
        BalledBurlapped
                            7.524
## c
        Bareroot
                            7.517
temp.lm <- lm(Temp ~ Treatment, data=WL.soil)</pre>
anova(temp.lm)
## Analysis of Variance Table
```

##

```
## Response: Temp
##
            Df Sum Sq Mean Sq F value
                                         Pr(>F)
## Treatment 5 42.674 8.5348 7.1405 1.308e-05 ***
## Residuals 84 100.403 1.1953
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
summary(temp.lm)
##
## Call:
## lm(formula = Temp ~ Treatment, data = WL.soil)
## Residuals:
##
      Min
               1Q Median
                              ЗQ
                                     Max
## -2.4133 -0.7467 -0.2133 0.7867 2.3200
## Coefficients:
                     Estimate Std. Error t value Pr(>|t|)
## (Intercept)
                     ## TreatmentBareroot
                    -1.9667
                                 0.3992 -4.926 4.15e-06 ***
                                 0.3992 -3.006 0.00349 **
## TreatmentSeedling
                      -1.2000
## TreatmentAcorn
                      -1.6333
                                 0.3992 -4.091 9.80e-05 ***
## TreatmentSeedbank -1.4000
                                 0.3992 -3.507 0.00073 ***
## TreatmentReference -2.0867
                                 0.3992 -5.227 1.24e-06 ***
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## Residual standard error: 1.093 on 84 degrees of freedom
## Multiple R-squared: 0.2983, Adjusted R-squared: 0.2565
## F-statistic: 7.14 on 5 and 84 DF, p-value: 1.308e-05
HSD <- HSD.test(temp.lm, "Treatment", console=TRUE)</pre>
##
## Study: temp.lm ~ "Treatment"
##
## HSD Test for Temp
## Mean Square Error: 1.19527
## Treatment, means
##
##
                      Temp
                                std r Min Max
## Acorn
                  21.08000 0.9630309 15 19.9 23.4
## BalledBurlapped 22.71333 0.7818172 15 21.5 24.3
## Bareroot
                 20.74667 1.2374321 15 19.7 22.6
## Reference
                  20.62667 0.6943308 15 19.7 21.9
                  21.31333 1.3314153 15 18.9 23.2
## Seedbank
## Seedling
                  21.51333 1.3590263 15 19.7 23.6
##
## alpha: 0.05; Df Error: 84
## Critical Value of Studentized Range: 4.124617
## Honestly Significant Difference: 1.164316
```

```
##
## Means with the same letter are not significantly different.
## Groups, Treatments and means
## a
        BalledBurlapped
                             22.71
## b
        Seedling
                            21.51
        Seedbank
                            21.31
## b
        Acorn
                            21.08
## b
        Bareroot
                             20.75
        Reference
                            20.63
TOC.lm <- lm(TOC ~ Treatment, data=WL.soil)
anova (TOC.lm)
## Analysis of Variance Table
##
## Response: TOC
            Df Sum Sq Mean Sq F value
## Treatment 5 2.5744 0.51488 5.8751 0.0001054 ***
## Residuals 84 7.3615 0.08764
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
summary(TOC.lm)
##
## Call:
## lm(formula = TOC ~ Treatment, data = WL.soil)
##
## Residuals:
##
      Min
               1Q Median
                                3Q
                                      Max
## -0.9659 -0.1943 0.0587 0.2202 0.5948
##
## Coefficients:
##
                      Estimate Std. Error t value Pr(>|t|)
## (Intercept)
                      5.116667 0.076436 66.941
                                                    <2e-16 ***
## TreatmentBareroot -0.258933 0.108097 -2.395
                                                     0.0188 *
## TreatmentSeedling -0.236300 0.108097 -2.186
                                                    0.0316 *
## TreatmentAcorn
                       0.007333
                                 0.108097
                                            0.068
                                                     0.9461
## TreatmentSeedbank
                      0.178767
                                 0.108097
                                             1.654
                                                     0.1019
## TreatmentReference 0.145567
                                 0.108097
                                            1.347
                                                     0.1817
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
##
## Residual standard error: 0.296 on 84 degrees of freedom
## Multiple R-squared: 0.2591, Adjusted R-squared: 0.215
## F-statistic: 5.875 on 5 and 84 DF, p-value: 0.0001054
HSD <- HSD.test(TOC.lm,"Treatment", console=TRUE)</pre>
## Study: TOC.lm ~ "Treatment"
##
## HSD Test for TOC
## Mean Square Error: 0.08763679
```

```
##
## Treatment, means
##
##
                                 std r
                       TOC
                                           Min
                                                 Max
## Acorn
                  5.124000 0.2037942 15 4.8010 5.384
## BalledBurlapped 5.116667 0.2281678 15 4.6815 5.426
## Bareroot
                 4.857733 0.3032293 15 4.4155 5.394
                 5.262233 0.2957362 15 4.8190 5.857
## Reference
## Seedbank
                  5.295433 0.3555548 15 4.3295 5.834
## Seedling
                  4.880367 0.3555291 15 4.1775 5.312
## alpha: 0.05; Df Error: 84
## Critical Value of Studentized Range: 4.124617
## Honestly Significant Difference: 0.315269
## Means with the same letter are not significantly different.
## Groups, Treatments and means
        Seedbank
## a
        Reference
                            5.262
        Acorn
                            5.124
        BalledBurlapped
## ab
                            5.117
## b
        Seedling
                             4.88
## b
                            4.858
        Bareroot
NH4.lm <- lm(NH4.N ~ Treatment, data=WL.soil)
anova(NH4.lm)
## Analysis of Variance Table
##
## Response: NH4.N
            Df Sum Sq Mean Sq F value Pr(>F)
## Treatment 5 8.607 1.7213 1.4738 0.207
## Residuals 84 98.108 1.1680
summary(NH4.lm)
##
## lm(formula = NH4.N ~ Treatment, data = WL.soil)
##
## Residuals:
##
                 1Q
       Min
                     Median
                                   3Q
                                            Max
## -2.01667 -0.75542 -0.01167 0.67708 2.50833
##
## Coefficients:
                     Estimate Std. Error t value Pr(>|t|)
##
## (Intercept)
                       9.9867
                                  0.2790 35.789
                                                  <2e-16 ***
                                          0.325
                                                    0.746
## TreatmentBareroot
                       0.1283
                                  0.3946
## TreatmentSeedling
                                          1.204
                       0.4750
                                  0.3946
                                                    0.232
## TreatmentAcorn
                                  0.3946
                                          0.266
                                                    0.791
                       0.1050
## TreatmentSeedbank
                      -0.3583
                                  0.3946 -0.908
                                                    0.366
## TreatmentReference 0.5800
                                  0.3946 1.470
                                                    0.145
```

```
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
##
## Residual standard error: 1.081 on 84 degrees of freedom
## Multiple R-squared: 0.08065,
                                Adjusted R-squared: 0.02593
## F-statistic: 1.474 on 5 and 84 DF, p-value: 0.207
NO3.lm <- lm(NO3.N ~ Treatment, data=WL.soil)
anova(NO3.lm)
## Analysis of Variance Table
##
## Response: NO3.N
##
            Df Sum Sq Mean Sq F value
                                        Pr(>F)
## Treatment 5 3316.9 663.37 7.1093 1.376e-05 ***
## Residuals 84 7838.1
                       93.31
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
summary(NO3.lm)
##
## Call:
## lm(formula = NO3.N ~ Treatment, data = WL.soil)
## Residuals:
               1Q Median
                               ЗQ
                                      Max
## -16.400 -5.818 -1.142
                            5.270 57.375
## Coefficients:
##
                     Estimate Std. Error t value Pr(>|t|)
                       21.447
                                   2.494
                                         8.599 3.81e-13 ***
## (Intercept)
## TreatmentBareroot -10.495
                                   3.527 -2.975 0.00382 **
## TreatmentSeedling
                        1.428
                                   3.527
                                          0.405 0.68655
## TreatmentAcorn
                        4.470
                                   3.527
                                          1.267 0.20856
## TreatmentSeedbank
                      3.968
                                   3.527
                                          1.125 0.26377
## TreatmentReference -9.504
                                   3.527 -2.694 0.00852 **
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## Residual standard error: 9.66 on 84 degrees of freedom
## Multiple R-squared: 0.2973, Adjusted R-squared: 0.2555
## F-statistic: 7.109 on 5 and 84 DF, p-value: 1.376e-05
HSD <- HSD.test(NO3.lm, "Treatment", console=TRUE)</pre>
## Study: NO3.lm ~ "Treatment"
##
## HSD Test for NO3.N
## Mean Square Error: 93.31097
## Treatment, means
##
##
                     NO3.N
                                 std r
                                          Min
                                                 Max
## Acorn
                  25.91667 7.176535 15 16.750 39.350
```

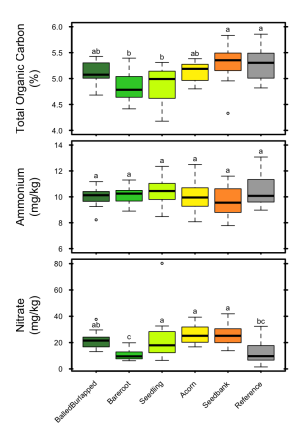
```
## BalledBurlapped 21.44667 6.560344 15 13.175 37.675
## Bareroot
                   10.95167 4.180081 15 6.250 19.850
## Reference
                   11.94300 8.227502 15 1.510 32.375
## Seedbank
                   25.41500 7.739381 15 13.825 41.875
## Seedling
                   22.87500 17.895869 15 6.475 80.250
##
## alpha: 0.05; Df Error: 84
## Critical Value of Studentized Range: 4.124617
##
## Honestly Significant Difference: 10.28737
## Means with the same letter are not significantly different.
## Groups, Treatments and means
         Acorn
## a
                              25.92
## a
         Seedbank
                              25.42
## a
                             22.88
         Seedling
         BalledBurlapped
                              21.45
## bc
         Reference
                              11.94
## c
         Bareroot
                              10.95
WL.phys.mean <- apply(WL.soil.phys, 2,
                      FUN = function(avg) aggregate(avg ~ WL.trts, WL.soil.phys, mean))
WL.phys.sem <- apply(WL.soil.phys, 2,
                     FUN = function(sem) aggregate(sem ~ WL.trts, WL.soil.phys, se))
WL.phys.95 <- apply(WL.soil.phys, 2,
                    FUN = function(ci_95) aggregate(ci_95 ~ WL.trts, WL.soil.phys, ci))
WL.nuts.mean <- apply(WL.soil.nuts, 2,
                      FUN = function(avg) aggregate(avg ~ WL.trts, WL.soil.nuts, mean))
WL.nuts.sem <- apply(WL.soil.nuts, 2,
                     FUN = function(sem) aggregate(sem ~ WL.trts, WL.soil.nuts, se))
WL.nuts.95 <- apply(WL.soil.nuts, 2,
                    FUN = function(ci_95) aggregate(ci_95 ~ WL.trts, WL.soil.nuts, ci))
phys.means.table <- data.frame(trt = WL.phys.mean$Moisture$WL.trts,</pre>
                               moisture = WL.phys.mean$Moisture$avg,
                               temp = WL.phys.mean$Temp$avg,
                               pH = WL.phys.mean$pH$avg)
phys.sem.table <- data.frame(trt = WL.phys.sem$Moisture$WL.trts,</pre>
                               moisture = WL.phys.sem$Moisture$sem,
                               temp = WL.phys.sem$Temp$sem,
                               pH = WL.phys.sem$pH$sem)
phys.ci.table <- data.frame(trt = WL.phys.95$Moisture$WL.trts,</pre>
                               moisture = WL.phys.95$Moisture$ci_95,
                               temp = WL.phys.95$Temp$ci_95,
                               pH = WL.phys.95\$pH\$ci_95)
nuts.means.table <- data.frame(trt = WL.nuts.mean$TOC$WL.trts,</pre>
                               toc = WL.nuts.mean$TOC$avg,
                               tn = WL.nuts.mean$TN$avg,
                               om = WL.nuts.mean$OM$avg,
                               nh4 = WL.nuts.mean$NH4.N$avg,
```

Plot: WL Soil Nuts

```
png(filename="../figures/soil.chem.png",
    width = 800, height = 1200, res = 96*2)
layout(matrix(1:3, 3, byrow = T))
par(mar = (c(0.5, 5, 0, 1) + 0.1), oma = c(7, 1, 1.5, 1))
# TOC
toc <- boxplot(WL.soil.nuts$TOC ~ WL.trts,</pre>
        col = myColors, xaxt = "n", yaxt = "n",
        xlab = "", ylab = "", ylim = c(4,6))
  # Axes with Tick Marks
  axis(side = 1, labels = F, tck = -0.01, lwd = 2)
  axis(side = 2, labels = T, tck = -0.02, lwd = 2, las = 1)
  axis(side = 4, labels = F, tck = -0.02, lwd = 2)
  axis(side = 2, labels = F, tck = 0.01, lwd = 2)
  axis(side = 4, labels = F, tck = 0.01, lwd = 2)
  box(lwd = 2)
  # Lables
  mtext("Total Organic Carbon\n(%)", side = 2, cex = 1, line = 3.5)
  # PostHoc Test
  tuk <- HSD.test(aov(WL.soil.nuts$TOC ~ WL.trts), "WL.trts")</pre>
  grp <- tuk$groups[c(match(levels(WL.trts), gsub(" ", "", tuk$groups$trt))), ]</pre>
  text(x = seq_along(levels(WL.trts)),
       y = toc\$stats[5, ] + ((par("usr")[4] - par("usr")[3]) * 0.05),
       labels = grp$M)
  #par("usr")
  \#text(x = seq\_along(levels(WL.trts)), y = 5.5, labels = grp$trt)
# # Total Nitrogen
# boxplot(WL.soil.nuts$TN ~ WL.trts,
          col = myColors, xaxt = "n", yaxt = "n",
#
#
          xlab = "", ylab = "", ylim = c(0.25, 0.4))
```

```
# Axes with Tick Marks
   axis(side = 1, labels = F, tck = -0.01, lwd = 2)
#
   axis(side = 2, labels = T, tck = -0.02, lwd = 2, las = 1)
#
   axis(side = 4, labels = F, tck = -0.02, lwd = 2)
   axis(side = 2, labels = F, tck = 0.01, lwd = 2)
#
   axis(side = 4, labels = F, tck = 0.01, lwd = 2)
#
   box(lwd = 2)
#
   # Lables
   mtext("Total\ Nitrogen\n(?)",\ side = 2,\ cex = 1,\ line = 3.5)
# # Organic Matter
# boxplot(WL.soil.nuts$OM ~ WL.trts,
          col = myColors, xaxt = "n", yaxt = "n",
          xlab = "", ylab = "", ylim = c(7, 11))
#
#
#
   # Axes with Tick Marks
#
   axis(side = 1, labels = F, tck = -0.01, lwd = 2)
   axis(side = 2, labels = T, tck = -0.02, lwd = 2, las = 1)
#
   axis(side = 4, labels = F, tck = -0.02, lwd = 2)
   axis(side = 2, labels = F, tck = 0.01, lwd = 2)
#
#
   axis(side = 4, labels = F, tck = 0.01, lwd = 2)
#
   box(lwd = 2)
#
  # Lables
  mtext("Organic Matter \n(?)", side = 2, cex = 1, line = 3.5)
# Ammonium
nh4 <- boxplot(WL.soil.nuts$NH4.N ~ WL.trts,
        col = myColors, xaxt = "n", yaxt = "n",
        xlab = "", ylab = "", ylim = c(6, 14))
  # Axes with Tick Marks
  axis(side = 1, labels = F, tck = -0.01, lwd = 2)
  axis(side = 2, labels = T, tck = -0.02, lwd = 2, las = 1)
  axis(side = 4, labels = F, tck = -0.02, lwd = 2)
  axis(side = 2, labels = F, tck = 0.01, lwd = 2)
  axis(side = 4, labels = F, tck = 0.01, lwd = 2)
  box(lwd = 2)
  # PostHoc Test
  tuk <- HSD.test(aov(WL.soil.nuts$NH4.N ~ WL.trts), "WL.trts")</pre>
  grp <- tuk$groups[c(match(levels(WL.trts), gsub(" ", "", tuk$groups$trt))), ]</pre>
  text(x = seq_along(levels(WL.trts)),
       y = nh4\$stats[5, ] + ((par("usr")[4] - par("usr")[3]) * 0.05),
       labels = grp$M)
  # Labels
  mtext("Ammonium\n(mg/kg)", side = 2, cex = 1, line = 3.5)
# Nitrate
no3 <- boxplot(WL.soil.nuts$NO3.N ~ WL.trts,</pre>
        col = myColors, xaxt = "n", yaxt = "n",
```

```
xlab = "", ylab = "")
  # Axes with Tick Marks
  axis(side = 1, labels = F, tck = -0.01, lwd = 2)
  axis(side = 2, labels = T, tck = -0.02, lwd = 2, las = 1)
  axis(side = 4, labels = F, tck = -0.02, lwd = 2)
  axis(side = 2, labels = F, tck = 0.01, lwd = 2)
  axis(side = 4, labels = F, tck = 0.01, lwd = 2)
  box(lwd = 2)
  # PostHoc Test
  tuk <- HSD.test(aov(WL.soil.nuts$NO3.N ~ WL.trts), "WL.trts")</pre>
  grp <- tuk$groups[c(match(levels(WL.trts), gsub(" ", "", tuk$groups$trt))), ]</pre>
  text(x = seq_along(levels(WL.trts)),
       y = no3$stats[5,] + ((par("usr")[4] - par("usr")[3]) * 0.05),
       labels = grp$M)
  # Lables
  mtext("Nitrate \setminus n(mg/kg)", side = 2, cex = 1, line = 3.5)
# Plot X labs at default X position
par(xpd = NA)
text(x = seq_along(levels(WL.trts)),
     y = par("usr")[3] - 0.1 * (par("usr")[4] - par("usr")[3]),
     srt = 45, adj = 1, labels = levels(WL.trts), xpd=NA)
# Close Plot Device
dev.off()
## pdf
##
   2
graphics.off()
```



Plot: WL Soil Phys

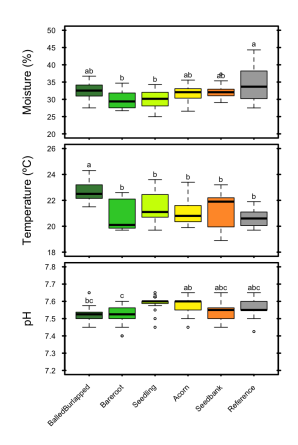
```
# PostHoc Test
  tuk <- HSD.test(aov(WL.soil.phys$Moisture ~ WL.trts), "WL.trts")</pre>
  grp <- tuk$groups[c(match(levels(WL.trts), gsub(" ", "", tuk$groups$trt))), ]</pre>
  text(x = seq_along(levels(WL.trts)),
       y = moisture stats[5, ] + ((par("usr")[4] - par("usr")[3]) * 0.05),
       labels = grp$M)
  # Labels
  mtext("Moisture (%)", side = 2, cex = 1, line = 3.5)
# Temperature
temp <- boxplot(WL.soil.phys$Temp ~ WL.trts,</pre>
        col = myColors, xaxt = "n", yaxt = "n",
        xlab = "", ylab = "", ylim = c(18, 26))
  # Axes with Tick Marks
  axis(side = 1, labels = F, tck = -0.01, lwd = 2)
  axis(side = 2, labels = T, tck = -0.02, lwd = 2, las = 1)
  axis(side = 4, labels = F, tck = -0.02, lwd = 2)
  axis(side = 2, labels = F, tck = 0.01, lwd = 2)
  axis(side = 4, labels = F, tck = 0.01, lwd = 2)
  box(lwd = 2)
  # PostHoc Test
  tuk <- HSD.test(aov(WL.soil.phys$Temp ~ WL.trts), "WL.trts")</pre>
  grp <- tuk$groups[c(match(levels(WL.trts), gsub(" ", "", tuk$groups$trt))), ]</pre>
  text(x = seq_along(levels(WL.trts)),
       y = temp\$stats[5, ] + ((par("usr")[4] - par("usr")[3]) * 0.05),
       labels = grp$M)
  # Lables
  mtext("Temperature (°C)", side = 2, cex = 1, line = 3.5)
# pH
  pH <- boxplot(WL.soil.phys$pH ~ WL.trts,
        col = myColors, xaxt = "n", yaxt = "n",
        xlab = "", ylab = "", ylim = c(7.2, 7.8))
  # Axes with Tick Marks
  axis(side = 1, labels = F, tck = -0.01, lwd = 2)
  axis(side = 2, labels = T, tck = -0.02, lwd = 2, las = 1)
  axis(side = 4, labels = F, tck = -0.02, lwd = 2)
  axis(side = 2, labels = F, tck = 0.01, lwd = 2)
  axis(side = 4, labels = F, tck = 0.01, lwd = 2)
  box(lwd = 2)
  # PostHoc Test
  tuk <- HSD.test(aov(WL.soil.phys$pH ~ WL.trts), "WL.trts")</pre>
  grp <- tuk$groups[c(match(levels(WL.trts), gsub(" ", "", tuk$groups$trt))), ]</pre>
  text(x = seq_along(levels(WL.trts)),
       y = pH$stats[5, ] + ((par("usr")[4] - par("usr")[3]) * 0.05),
       labels = grp$M)
```

```
# Lables
mtext("pH", side = 2, cex = 1, line = 3.5)

# Plot x labs at default x position
par(xpd = NA)
text(x = seq_along(levels(WL.trts)),
    y = par("usr")[3] - 0.1 * (par("usr")[4] - par("usr")[3]),
    srt = 45, adj = 1, labels = levels(WL.trts), xpd=NA)

# Close Plot Device
dev.off()

## pdf
## 2
graphics.off()
```



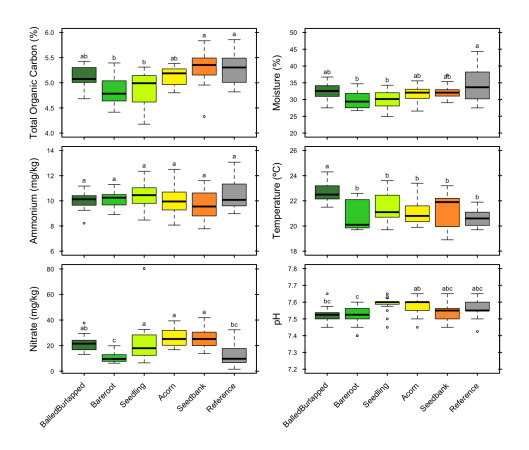
Joint Plot: Wetland Soil

```
png(filename="../figures/WL.soil.png",
    width = 1600, height = 1400, res = 96*2)
layout(matrix(1:6, 3, byrow = F))
par(mar = (c(1, 4, 0, 2) + 0.1), oma = c(8, 1, 2, 0))
# TOC
toc <- boxplot(WL.soil.nuts$TOC ~ WL.trts,</pre>
        col = myColors, xaxt = "n", yaxt = "n",
        xlab = "", ylab = "", ylim = c(4,6))
  # Axes with Tick Marks
  axis(side = 1, labels = F, tck = -0.02, lwd = 1.5)
  axis(side = 2, labels = T, tck = -0.02, lwd = 1.5, las = 1)
  axis(side = 4, labels = F, tck = -0.02, lwd = 1.5)
  axis(side = 2, labels = F, tck = 0.01, lwd = 1.5)
  axis(side = 4, labels = F, tck = 0.01, lwd = 1.5)
  box(lwd = 1.5)
  # Lables
  mtext("Total Organic Carbon (%)", side = 2, cex = 1, line = 3)
  # PostHoc Test
 tuk <- HSD.test(aov(WL.soil.nuts$TOC ~ WL.trts), "WL.trts")</pre>
  grp <- tuk$groups[c(match(levels(WL.trts), gsub(" ", "", tuk$groups$trt))), ]</pre>
 text(x = seq_along(levels(WL.trts)),
       y = toc\$stats[5, ] + ((par("usr")[4] - par("usr")[3]) * 0.05),
       labels = grp$M)
# Ammonium
nh4 <- boxplot(WL.soil.nuts$NH4.N ~ WL.trts,
        col = myColors, xaxt = "n", yaxt = "n",
        xlab = "", ylab = "", ylim = c(6, 14))
  # Axes with Tick Marks
  axis(side = 1, labels = F, tck = -0.02, lwd = 1.5)
  axis(side = 2, labels = T, tck = -0.02, lwd = 1.5, las = 1)
  axis(side = 4, labels = F, tck = -0.02, lwd = 1.5)
  axis(side = 2, labels = F, tck = 0.01, lwd = 1.5)
  axis(side = 4, labels = F, tck = 0.01, lwd = 1.5)
  box(lwd = 1.5)
  # PostHoc Test
  tuk <- HSD.test(aov(WL.soil.nuts$NH4.N ~ WL.trts), "WL.trts")</pre>
  grp <- tuk$groups[c(match(levels(WL.trts), gsub(" ", "", tuk$groups$trt))), ]</pre>
 text(x = seq_along(levels(WL.trts)),
       y = nh4\$stats[5, ] + ((par("usr")[4] - par("usr")[3]) * 0.05),
       labels = grp$M)
  mtext("Ammonium (mg/kg)", side = 2, cex = 1, line = 3)
# Nitrate
```

```
no3 <- boxplot(WL.soil.nuts$NO3.N ~ WL.trts,</pre>
        col = myColors, xaxt = "n", yaxt = "n",
        xlab = "", ylab = "")
  # Axes with Tick Marks
  axis(side = 1, labels = F, tck = -0.02, lwd = 1.5)
  axis(side = 2, labels = T, tck = -0.02, lwd = 1.5, las = 1)
  axis(side = 4, labels = F, tck = -0.02, lwd = 1.5)
  axis(side = 2, labels = F, tck = 0.01, lwd = 1.5)
  axis(side = 4, labels = F, tck = 0.01, lwd = 1.5)
  box(lwd = 1.5)
  # PostHoc Test
  tuk <- HSD.test(aov(WL.soil.nuts$NO3.N ~ WL.trts), "WL.trts")</pre>
  grp <- tuk$groups[c(match(levels(WL.trts), gsub(" ", "", tuk$groups$trt))), ]</pre>
 text(x = seq_along(levels(WL.trts)),
       y = no3$stats[5, ] + ((par("usr")[4] - par("usr")[3]) * 0.05),
       labels = grp$M)
  # Lables
  mtext("Nitrate (mg/kg)", side = 2, cex = 1, line = 3)
# Plot X labs at default X position
par(xpd = NA)
text(x = seq along(levels(WL.trts)),
     y = par("usr")[3] - 0.1 * (par("usr")[4] - par("usr")[3]),
     srt = 45, adj = 1, labels = levels(WL.trts), xpd=NA, cex = 1.25)
# Moisture
moisture <- boxplot(WL.soil.phys$Moisture ~ WL.trts,</pre>
        col = myColors, xaxt = "n", yaxt = "n",
        xlab = "", ylab = "", ylim = c(20, 50))
  # Axes with Tick Marks
  axis(side = 1, labels = F, tck = -0.02, lwd = 1.5)
  axis(side = 2, labels = T, tck = -0.02, lwd = 1.5, las = 1)
  axis(side = 4, labels = F, tck = -0.02, lwd = 1.5)
  axis(side = 2, labels = F, tck = 0.01, lwd = 1.5)
  axis(side = 4, labels = F, tck = 0.01, lwd = 1.5)
  box(lwd = 1.5)
  # PostHoc Test
 tuk <- HSD.test(aov(WL.soil.phys$Moisture ~ WL.trts), "WL.trts")</pre>
  grp <- tuk$groups[c(match(levels(WL.trts), gsub(" ", "", tuk$groups$trt))), ]</pre>
  text(x = seq_along(levels(WL.trts)),
       y = moisture stats[5, ] + ((par("usr")[4] - par("usr")[3]) * 0.05),
       labels = grp$M)
  # Labels
  mtext("Moisture (%)", side = 2, cex = 1, line = 3)
# Temperature
temp <- boxplot(WL.soil.phys$Temp ~ WL.trts,</pre>
```

```
col = myColors, xaxt = "n", yaxt = "n",
        xlab = "", ylab = "", ylim = c(18, 26))
  # Axes with Tick Marks
  axis(side = 1, labels = F, tck = -0.02, lwd = 1.5)
  axis(side = 2, labels = T, tck = -0.02, lwd = 1.5, las = 1)
  axis(side = 4, labels = F, tck = -0.02, lwd = 1.5)
  axis(side = 2, labels = F, tck = 0.01, lwd = 1.5)
  axis(side = 4, labels = F, tck = 0.01, lwd = 1.5)
  box(lwd = 1.5)
  # PostHoc Test
  tuk <- HSD.test(aov(WL.soil.phys$Temp ~ WL.trts), "WL.trts")</pre>
  grp <- tuk$groups[c(match(levels(WL.trts), gsub(" ", "", tuk$groups$trt))), ]</pre>
  text(x = seq_along(levels(WL.trts)),
       y = temp\$stats[5, ] + ((par("usr")[4] - par("usr")[3]) * 0.05),
       labels = grp$M)
  # Lables
  mtext("Temperature (°C)", side = 2, cex = 1, line = 3)
# pH
 pH <- boxplot(WL.soil.phys$pH ~ WL.trts,</pre>
        col = myColors, xaxt = "n", yaxt = "n",
        xlab = "", ylab = "", ylim = c(7.2, 7.8))
  # Axes with Tick Marks
  axis(side = 1, labels = F, tck = -0.02, lwd = 1.5)
  axis(side = 2, labels = T, tck = -0.02, lwd = 1.5, las = 1)
  axis(side = 4, labels = F, tck = -0.02, lwd = 1.5)
  axis(side = 2, labels = F, tck = 0.01, lwd = 1.5)
  axis(side = 4, labels = F, tck = 0.01, lwd = 1.5)
  box(lwd = 1.5)
  # PostHoc Test
  tuk <- HSD.test(aov(WL.soil.phys$pH ~ WL.trts), "WL.trts")</pre>
  grp <- tuk$groups[c(match(levels(WL.trts), gsub(" ", "", tuk$groups$trt))), ]</pre>
  text(x = seq along(levels(WL.trts)),
       y = pH$stats[5, ] + ((par("usr")[4] - par("usr")[3]) * 0.05),
       labels = grp$M)
  # Lables
  mtext("pH", side = 2, cex = 1, line = 3)
# Plot X labs at default X position
par(xpd = NA)
text(x = seq_along(levels(WL.trts)),
     y = par("usr")[3] - 0.1 * (par("usr")[4] - par("usr")[3]),
     srt = 45, adj = 1, labels = levels(WL.trts), xpd=NA, cex = 1.25)
# Close Plot Device
dev.off()
```

```
## pdf
## 2
graphics.off()
```



Simple Community Hypothesis Testing

```
# Check that Bac and Plant data have same structure as design
all.equal(row.names(WLdataREL), row.names(design))

## [1] TRUE
all.equal(row.names(WL.unifrac), row.names(design))

## [1] TRUE
all.equal(row.names(WL.plant), row.names(design))

## [1] "90 string mismatches"

# Same info seem to be inside so I'm going to rename the plant data
row.names(WL.plant) <- row.names(design)</pre>
```

```
WL.bac.adonis <- adonis(WLdataREL ~ treatments, method = "bray", perm=999)
## Call:
## adonis(formula = WLdataREL ~ treatments, permutations = 999, method = "bray")
## Permutation: free
## Number of permutations: 999
## Terms added sequentially (first to last)
##
             Df SumsOfSqs MeanSqs F.Model
                                               R2 Pr(>F)
                 0.4235 0.084706 1.8661 0.09997 0.004 **
## treatments 5
## Residuals 84
                   3.8130 0.045392
                                          0.90003
## Total
           89
                 4.2365
                                           1.00000
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
# Odd sites in bacterial composition data (explore more)
odd.sites <- c("HC_A3_ss1_S11", "HC_A3_ss2_S12", "HC_A3_ss3_S13",
              "HC_C3_ss5_S45", "HC_D2_ss3_S53", "HC_E1_ss1_S61")
WLdataREL.2 <- WLdataREL[setdiff(rownames(WLdataREL), odd.sites), ]</pre>
treatments.2 <- design[setdiff(rownames(design), odd.sites), ]$Treatment</pre>
odd.sites
## [1] "HC_A3_ss1_S11" "HC_A3_ss2_S12" "HC_A3_ss3_S13" "HC_C3_ss5_S45"
## [5] "HC_D2_ss3_S53" "HC_E1_ss1_S61"
WL.bac.adonis <- adonis(WLdataREL.2 ~ treatments.2, method = "bray", perm=999)
WL.bac.adonis
## Call:
## adonis(formula = WLdataREL.2 ~ treatments.2, permutations = 999,
                                                                   method = "bray")
## Permutation: free
## Number of permutations: 999
## Terms added sequentially (first to last)
##
              Df SumsOfSqs MeanSqs F.Model
                                                 R2 Pr(>F)
## treatments.2 5 0.34744 0.069488 2.3362 0.13025 0.001 ***
## Residuals 78 2.31999 0.029743
                                             0.86975
## Total
               83 2.66743
                                             1.00000
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
WL.unifrac.adonis <- adonis(WL.unifrac.dist ~ treatments, perm = 999)
WL.unifrac.adonis
##
## Call:
## adonis(formula = WL.unifrac.dist ~ treatments, permutations = 999)
```

```
##
## Permutation: free
## Number of permutations: 999
## Terms added sequentially (first to last)
##
             Df SumsOfSqs MeanSqs F.Model
                                                R2 Pr(>F)
                  0.05081 0.0101624 1.6563 0.08974 0.062 .
## treatments 5
## Residuals 84
                  0.51538 0.0061355
                                           0.91026
## Total
             89
                 0.56619
                                            1.00000
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
WL.unifrac.dist2 <- as.matrix(WL.unifrac.dist)[setdiff(labels(WL.unifrac.dist),
                             odd.sites), setdiff(labels(WL.unifrac.dist),
                             odd.sites)]
WL.unifrac.adonis <- adonis(as.dist(WL.unifrac.dist2) ~ treatments.2, perm = 999)
WL.unifrac.adonis
## Call:
## adonis(formula = as.dist(WL.unifrac.dist2) ~ treatments.2, permutations = 999)
##
## Permutation: free
## Number of permutations: 999
## Terms added sequentially (first to last)
##
               Df SumsOfSqs MeanSqs F.Model
## treatments.2 5 0.036942 0.0073883 2.6616 0.14575 0.001 ***
## Residuals
               78 0.216521 0.0027759
                                             0.85425
## Total
               83 0.253462
                                             1.00000
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
WL.plant.adonis <- adonis(WL.plant ~ treatments, method = "bray", perm=999)
WL.plant.adonis
##
## Call:
## adonis(formula = WL.plant ~ treatments, permutations = 999, method = "bray")
## Permutation: free
## Number of permutations: 999
## Terms added sequentially (first to last)
##
             Df SumsOfSqs MeanSqs F.Model
                                             R2 Pr(>F)
## treatments 5
                10.3524 2.07047 18.608 0.52553 0.001 ***
                 9.3466 0.11127
## Residuals 84
                                          0.47447
## Total
             89 19.6990
                                          1.00000
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
```

```
# Simper Analysis for Species Responses
#WL.bac.simper <- simper(WLdataREL, group = treatments, permutations = 999)
#bac.sum <- summary(WL.bac.simper, ordered = T)

# Which plants influenced plant communities between trts
#WL.plant.simper <- simper(WL.plant, group = treatments, permutations = 999)
#plant.sum <- summary(WL.plant.simper, ordered = T)
#plant.sum</pre>
```

Indicator Species Analysis

```
# Plant Communities
library("labdsv")
## Loading required package: mgcv
## This is mgcv 1.8-16. For overview type 'help("mgcv-package")'.
## Loading required package: MASS
## Loading required package: cluster
## Attaching package: 'labdsv'
## The following object is masked from 'package:stats':
##
##
       density
plant.ind <- indval(WL.plant, treatments)</pre>
summary(plant.ind)
##
                        cluster indicator_value probability
## BARE
                              2
                                         0.3201
                                                       0.001
## Phalaris.arundinacea
                              4
                                          0.3148
                                                       0.001
## Lemna.minor
                              6
                                         0.4000
                                                       0.001
## Acer.saccharinum
                              6
                                         0.2000
                                                       0.024
## Bidens.frondosa
                                         0.2000
                                                       0.017
## Sum of probabilities
                                         = 11.671
## Sum of Indicator Values
                                         = 2.51
## Sum of Significant Indicator Values = 1.43
## Number of Significant Indicators
                                        = 5
## Significant Indicator Distribution
##
## 2 4 6
## 1 1 3
plant.ind$indval
```

```
##
                                  BalledBurlapped
                                                     Bareroot Seedling
## Acer.saccharinum
                                       0.0000000 0.00000000 0.0000000
                                       0.11001710 0.000000000 0.0000000
## Aster.lanceolatus.var..simplex
                                       0.12122001 0.320139686 0.1304718
## BARE
## Bidens.frondosa
                                       0.0000000 0.00000000 0.0000000
## Carex.sp.
                                       0.00000000 0.000000000 0.0000000
## Elymus.virginicus
                                       0.0000000 0.00000000 0.0000000
## Impatiens.capensis
                                       0.00000000 0.000000000 0.0000000
## Leersia.virginica
                                       0.05807388 0.000000000 0.0000000
## Lemna.minor
                                       0.00000000 0.000000000 0.0000000
## Lycopus.virginicus
                                       0.0000000 0.00000000 0.0000000
                                       0.05678233 0.000000000 0.0000000
## Morus.alba
                                       0.0000000 0.000000000 0.0000000
## Persicaria.pensylvanica
## Phalaris.arundinacea
                                       0.16332556 0.006775177 0.1310697
## Physostegia.virginiana
                                       0.0000000 0.00000000 0.0000000
## Pilea.pumila
                                       0.03255814 0.000000000 0.0000000
## Sicyos.angulatus
                                       0.0000000 0.06666667 0.0000000
## Stachys.palustris
                                       0.0000000 0.00000000 0.0000000
## Ulmus.americana
                                       0.0666667 0.00000000 0.0000000
                                       0.0666667 0.00000000 0.0000000
## Vitis.riparia
##
                                        Acorn
                                                Seedbank
                                                           Reference
## Acer.saccharinum
                                  0.00000000 0.00000000 0.200000000
## Aster.lanceolatus.var..simplex 0.000000000 0.00000000 0.156649562
## BARE
                                  0.004877485 0.01486021 0.262185693
                                  0.00000000 0.00000000 0.200000000
## Bidens.frondosa
## Carex.sp.
                                  0.00000000 0.00000000 0.066666667
## Elymus.virginicus
                                  0.00000000 0.00000000 0.066666667
## Impatiens.capensis
                                  0.00000000 0.00000000 0.066666667
## Leersia.virginica
                                  0.00000000 0.00000000 0.017185568
                                  0.00000000 0.00000000 0.400000000
## Lemna.minor
                                  0.00000000 0.00000000 0.066666667
## Lycopus.virginicus
## Morus.alba
                                  0.00000000 0.00000000 0.009884332
## Persicaria.pensylvanica
                                  0.00000000 0.00000000 0.066666667
## Phalaris.arundinacea
                                  0.314842939 0.30629608 0.007130337
## Physostegia.virginiana
                                  0.00000000 0.00000000 0.066666667
## Pilea.pumila
                                  0.00000000 0.00000000 0.068217054
## Sicyos.angulatus
                                  0.000000000 0.00000000 0.000000000
## Stachys.palustris
                                  0.00000000 0.00000000 0.133333333
                                  0.000000000 0.00000000 0.000000000
## Ulmus.americana
                                  0.00000000 0.00000000 0.000000000
## Vitis.riparia
levels(treatments)
## [1] "BalledBurlapped" "Bareroot"
                                           "Seedling"
                                                              "Acorn"
## [5] "Seedbank"
                         "Reference"
plant.groups <- rep(NA, length(treatments))</pre>
for (i in 1:length(treatments)){
  if (treatments[i] == "Reference" |
      treatments[i] == "Bareroot"){
   plant.groups[i] <- "Group 1"</pre>
  if (treatments[i] == "Seedling" |
      treatments[i] == "BalledBurlapped"){
  plant.groups[i] <- "Group 2"</pre>
```

```
if (treatments[i] == "Seedbank" |
      treatments[i] == "Acorn"){
  plant.groups[i] <- "Group 3"</pre>
}
plant.ind2 <- indval(WL.plant, plant.groups)</pre>
summary(plant.ind2)
                         cluster indicator_value probability
##
## BARE
                                         0.5810
## Lemna.minor
                                          0.2000
                                                       0.003
                               1
                               3
## Phalaris.arundinacea
                                         0.6211
                                                       0.001
## Sum of probabilities
                                         = 12.706
## Sum of Indicator Values
                                         = 2.14
## Sum of Significant Indicator Values = 1.4
## Number of Significant Indicators
## Significant Indicator Distribution
##
## 1 3
## 2 1
WLdataREL.2 <- WLdataREL.2[, colSums(WLdataREL.2) > 0]
bac.ind <- indval(WLdataREL.2, treatments.2)</pre>
summary(bac.ind)
##
## Sum of probabilities
                                         = 17132.223
## Sum of Indicator Values
                                            2897.24
## Sum of Significant Indicator Values =
                                            464.74
## Number of Significant Indicators
                                         = 1911
## Significant Indicator Distribution
##
       2 3 4 5
## 417 93 172 518 240 471
#bac.ind
WLdataREL.2 <- WLdataREL.2[, colSums(WLdataREL.2) > 0.05]
bac.ind <- indval(WLdataREL.2, treatments.2)</pre>
levels(treatments.2)
## [1] "BalledBurlapped" "Bareroot"
                                            "Seedling"
                                                               "Acorn"
## [5] "Seedbank"
                         "Reference"
```

шш		-1	:	
##	Otu000173	cruster 1	indicator_value 0.2498	0.001
##	Otu000173	1	0.2439	0.001
##	Otu000133	1	0.2439	0.001
##	Otu000130	1	0.2372	0.001
##	Otu000070	1	0.2372	0.003
##	Otu000047	1	0.2249	0.024
##	Otu000230	1	0.2249	0.014
##	Otu000176	1	0.2187	0.005
##	Otu000244	1	0.2179	0.043
##	Otu000133	1	0.2173	0.045
##	Otu000100	1	0.2148	0.001
##	Otu000092	1	0.2141	0.001
##	Otu000192	1	0.2134	0.021
##	Otu000057	1	0.2126	0.005
##	Otu000090	1	0.2118	0.002
##	Otu000089	1	0.2105	0.002
##	Otu000170	1	0.2105	0.047
##	Otu000020	1	0.2095	0.001
##	Otu000204	1	0.2086	0.003
##	Otu000097	1	0.2065	0.003
##	Otu000260	1	0.2046	0.022
##	Otu000049	1	0.2005	0.023
##	Otu000245	1	0.2002	0.044
##	Otu000258	1	0.1992	0.040
##	Otu000105	1	0.1986	0.048
##	Otu000014	1	0.1949	0.001
##	Otu000005	1	0.1938	0.001
##	Otu000021	1	0.1936	0.010
##	Otu000100	1	0.1931	0.045
##	Otu000037	1	0.1910	0.011
##	Otu000033	1	0.1909	0.039
##	Otu000008	1	0.1794	0.025
##	Otu000030	2	0.2209	0.002
##	Otu000189	2	0.2206	0.002
##	Otu000207	2	0.2151	0.009
##	Otu000056	2	0.2119	0.009
##	Otu000211	2	0.2107	0.031
##	Otu000177	2	0.2082	0.037
##	Otu000205	2	0.2072	0.028
##	Otu000140	2	0.2053	0.013
##	Otu000144	2	0.2035	0.022
##	Otu000143	2	0.1982	0.033
##	Otu000059	2	0.1938	0.012
##	Otu000044	2	0.1920	0.005
##	Otu000093	2	0.1890	0.035
##	Otu000193	3	0.2910	0.017
##	Otu000058	3	0.2382	0.001
##	Otu000186	3	0.2335	0.033
##	Otu000220	3	0.1986	0.040
##	Otu000078	3	0.1965	0.006
##	Otu000040	3	0.1951	0.001

##	Otu000119	4	0.2595		0.035
##	Otu000125	4	0.2389		0.001
##	Otu000221	4	0.2298		0.004
##	Otu000067	4	0.2295		0.002
##	Otu000212	4	0.2272		0.001
##	Otu000181	4	0.2181		0.005
##	Otu000096	4	0.2178		0.003
##	Otu000266	4	0.2178		0.015
##	Otu000188	4	0.2110		0.015
##	Otu000128	4	0.2080		0.014
##	Otu000122	4	0.2049		0.035
##	Otu000118	4	0.2031		0.046
##	Otu000230	4	0.1999		0.021
##	Otu000130	4	0.1977		0.017
##	Otu000017	4	0.1932		0.037
##	Otu000066	4	0.1918		0.041
##	Otu000110	4	0.1914		0.044
##	Otu000237	5	0.2656		0.029
##	Otu000279	5	0.2631		0.001
##	Otu000201	5	0.2576		0.001
##	Otu000268	5	0.2473		0.001
##	Otu000274	5	0.2435		0.008
##	Otu000011	5	0.2295		0.004
##	Otu000155	5	0.2259		0.009
##	Otu000142	5	0.2246		0.002
##	Otu000142	5	0.2239		0.002
##	Otu000242	5	0.2219		0.001
##	Otu000120	5	0.2213		0.050
##	Otu000200	5	0.2154		0.001
##	Otu00001	5	0.2101		0.001
##	Otu000129	5	0.2019		0.003
##	Otu000138	5	0.2019		0.006
##	Otu000084	5	0.1992		0.000
##	Otu000074	5	0.1947		0.009
	Otu000029 Otu000041	5	0.1946		0.026
## ##	Otu000041	5	0.1944		0.036
		5			
##	Otu000080		0.1843		0.039
	Otu000200	6	0.2881		0.017
##		6	0.2607		0.035
	Otu000232	6	0.2550		0.009
	Otu000275	6	0.2332		0.024
	Otu000117	6	0.2260		0.005
	Otu000082	6	0.2157		0.004
	Otu000055	6	0.2130		0.001
	Otu000183	6	0.2093		0.016
##	Otu000087	6	0.2081		0.005
##		6	0.2046		0.044
##	Otu000111	6	0.2034		0.022
##	Otu000031	6	0.1927		0.017
##					
##	Sum of probabil	ities		=	65.58
##					
##	Sum of Indicato	r Values		=	55.48
##					

```
## Sum of Significant Indicator Values = 21.54
##
## Number of Significant Indicators
                                             100
## Significant Indicator Distribution
##
## 1 2 3 4 5 6
## 32 13 6 17 20 12
inds <- which(bac.ind$pval <= 0.05)</pre>
bac.indicators <- as.data.frame(matrix(NA, nrow = length(inds), ncol = 4))</pre>
colnames(bac.indicators) <- c("OTU", "Cluster", "IndVal", "Prob")</pre>
bac.indicators$OTU <- names(inds)</pre>
bac.indicators$Cluster <- bac.ind$maxcls[inds]</pre>
bac.indicators$IndVal <- bac.ind$indcls[inds]</pre>
bac.indicators$Prob <- bac.ind$pval[inds]</pre>
ind.tax <- WL.tax[which(as.character(WL.tax$OTU) %in% bac.indicators$OTU), ]</pre>
ind.tax <- ind.tax[match(ind.tax$OTU, bac.indicators$OTU), ]</pre>
indicator.bac <- cbind(bac.indicators, ind.tax[, -c(1)])
indicator.bac <- indicator.bac[order(as.numeric(indicator.bac$Cluster)), ]</pre>
table(indicator.bac$Cluster)
## 1 2 3 4 5 6
## 32 13 6 17 20 12
table(indicator.bac$Phylum)
##
##
           Acidobacteria
                                 Actinobacteria Bacteria_unclassified
##
                                                                     17
##
           Bacteroidetes
                                    Chloroflexi
                                                             Firmicutes
##
##
          Planctomycetes
                                 Proteobacteria
                                                       Verrucomicrobia
                                              39
                                                                      7
table(indicator.bac$Cluster)
## 1 2 3 4 5 6
## 32 13 6 17 20 12
levels(treatments.2)
## [1] "BalledBurlapped" "Bareroot"
                                             "Seedling"
                                                                "Acorn"
## [5] "Seedbank"
                          "Reference"
# Export Bacteria Indicator Table
write.table(indicator.bac, "../data/BacterialIndicators.txt",
            sep="\t", row.names = F, quote = F)
split(indicator.bac$Phylum, indicator.bac$Cluster)
```

```
## $`1`
                                 "Proteobacteria"
    [1] "Acidobacteria"
    [3] "Acidobacteria"
                                 "Acidobacteria"
   [5] "Verrucomicrobia"
##
                                 "Bacteroidetes"
    [7] "Proteobacteria"
                                 "Proteobacteria"
##
  [9] "Bacteria unclassified" "Bacteroidetes"
       "Bacteria unclassified" "Acidobacteria"
## [13] "Bacteria unclassified" "Verrucomicrobia"
## [15] "Verrucomicrobia"
                                 "Proteobacteria"
## [17] "Bacteria_unclassified" "Acidobacteria"
## [19] "Acidobacteria"
                                 "Bacteria_unclassified"
## [21] "Bacteroidetes"
                                 "Proteobacteria"
## [23] "Planctomycetes"
                                 "Verrucomicrobia"
## [25] "Proteobacteria"
                                 "Bacteria_unclassified"
## [27] "Proteobacteria"
                                 "Bacteroidetes"
## [29] "Actinobacteria"
                                 "Bacteria_unclassified"
## [31] "Acidobacteria"
                                 "Acidobacteria"
##
## $`2`
##
   [1] "Proteobacteria"
                                 "Proteobacteria"
##
   [3] "Acidobacteria"
                                 "Bacteria_unclassified"
  [5] "Firmicutes"
                                 "Bacteria unclassified"
##
  [7] "Proteobacteria"
                                 "Chloroflexi"
   [9] "Verrucomicrobia"
                                 "Bacteria unclassified"
                                 "Proteobacteria"
## [11] "Acidobacteria"
## [13] "Chloroflexi"
##
## $`3`
## [1] "Bacteria_unclassified" "Proteobacteria"
                                                         "Proteobacteria"
## [4] "Proteobacteria"
                                "Proteobacteria"
                                                         "Proteobacteria"
##
## $`4`
   [1] "Proteobacteria"
                                 "Proteobacteria"
##
   [3] "Acidobacteria"
                                 "Verrucomicrobia"
    [5] "Proteobacteria"
                                 "Acidobacteria"
   [7] "Proteobacteria"
                                 "Acidobacteria"
## [9] "Proteobacteria"
                                 "Firmicutes"
## [11] "Firmicutes"
                                 "Bacteria_unclassified"
## [13] "Proteobacteria"
                                 "Acidobacteria"
## [15] "Bacteria_unclassified" "Verrucomicrobia"
## [17] "Acidobacteria"
##
## $\5\
   [1] "Proteobacteria"
                                 "Actinobacteria"
##
   [3] "Proteobacteria"
                                 "Bacteria_unclassified"
##
   [5] "Proteobacteria"
                                 "Bacteria_unclassified"
##
    [7] "Proteobacteria"
                                 "Actinobacteria"
##
   [9] "Bacteroidetes"
                                 "Bacteria_unclassified"
## [11] "Proteobacteria"
                                 "Bacteroidetes"
## [13] "Proteobacteria"
                                 "Proteobacteria"
## [15] "Acidobacteria"
                                 "Proteobacteria"
## [17] "Planctomycetes"
                                 "Bacteria_unclassified"
## [19] "Proteobacteria"
                                 "Bacteroidetes"
##
```

```
## $`6`
    [1] "Actinobacteria" "Acidobacteria"
                                           "Proteobacteria" "Proteobacteria"
    [5] "Proteobacteria" "Acidobacteria"
                                           "Proteobacteria" "Bacteroidetes"
    [9] "Proteobacteria" "Proteobacteria" "Proteobacteria" "Proteobacteria"
##
split(indicator.bac$Class, indicator.bac$Cluster)
## $`1`
   [1] "Acidobacteria_Gp6"
##
                                        "Alphaproteobacteria"
    [3] "Acidobacteria_Gp6"
                                        "Acidobacteria_Gp6"
##
    [5] "Subdivision3"
                                         "Sphingobacteria"
##
   [7] "Alphaproteobacteria"
                                        "Gammaproteobacteria"
  [9] "Bacteria_unclassified"
                                        "Sphingobacteria"
## [11] "Bacteria_unclassified"
                                         "Acidobacteria_Gp5"
## [13] "Bacteria_unclassified"
                                         "Subdivision3"
                                        "Proteobacteria_unclassified"
  [15] "Spartobacteria"
##
  [17] "Bacteria_unclassified"
                                        "Acidobacteria_Gp4"
  [19] "Acidobacteria_Gp6"
                                        "Bacteria_unclassified"
  [21] "Flavobacteria"
                                        "Deltaproteobacteria"
## [23] "Planctomycetacia"
                                        "Verrucomicrobia_unclassified"
## [25] "Gammaproteobacteria"
                                        "Bacteria_unclassified"
## [27] "Gammaproteobacteria"
                                         "Sphingobacteria"
## [29] "Actinobacteria"
                                        "Bacteria_unclassified"
  [31] "Acidobacteria_Gp17"
                                        "Acidobacteria_Gp22"
##
## $`2`
   [1] "Alphaproteobacteria"
##
                                    "Alphaproteobacteria"
    [3] "Acidobacteria_Gp6"
                                    "Bacteria_unclassified"
   [5] "Bacilli"
##
                                    "Bacteria_unclassified"
##
    [7] "Alphaproteobacteria"
                                    "Chloroflexi_unclassified"
##
   [9] "Subdivision3"
                                    "Bacteria_unclassified"
  [11] "Acidobacteria_Gp4"
                                    "Deltaproteobacteria"
  [13] "Chloroflexi_unclassified"
##
## $`3`
## [1] "Bacteria_unclassified"
                                      "Deltaproteobacteria"
  [3] "Deltaproteobacteria"
                                      "Proteobacteria_unclassified"
  [5] "Proteobacteria_unclassified" "Betaproteobacteria"
##
## $`4`
##
    [1] "Betaproteobacteria"
                                   "Alphaproteobacteria"
##
    [3] "Acidobacteria_Gp4"
                                   "Spartobacteria"
##
       "Betaproteobacteria"
                                   "Acidobacteria_Gp6"
##
   [7] "Gammaproteobacteria"
                                   "Acidobacteria_Gp4"
##
   [9] "Alphaproteobacteria"
                                   "Bacilli"
                                   "Bacteria_unclassified"
## [11] "Firmicutes_unclassified"
  [13] "Deltaproteobacteria"
                                   "Acidobacteria_Gp3"
   [15] "Bacteria_unclassified"
                                   "Subdivision3"
##
   [17] "Acidobacteria_Gp25"
##
## $\5\
                                       "Actinobacteria"
##
    [1] "Deltaproteobacteria"
##
    [3] "Alphaproteobacteria"
                                       "Bacteria_unclassified"
##
    [5]
       "Deltaproteobacteria"
                                       "Bacteria_unclassified"
```

"Actinobacteria"

[7] "Betaproteobacteria"

```
[9] "Sphingobacteria"
                                         "Bacteria_unclassified"
  [11] "Proteobacteria_unclassified" "Sphingobacteria"
                                         "Deltaproteobacteria"
  [13] "Deltaproteobacteria"
  [15] "Acidobacteria_Gp22"
                                         "Gammaproteobacteria"
  [17] "Planctomycetacia"
                                         "Bacteria_unclassified"
##
   [19] "Deltaproteobacteria"
                                         "Flavobacteria"
##
## $`6`
    [1] "Actinobacteria"
##
                                         "Acidobacteria_Gp10"
                                         "Betaproteobacteria"
##
    [3] "Alphaproteobacteria"
    [5] "Deltaproteobacteria"
                                         "Acidobacteria_Gp7"
                                         "Sphingobacteria"
    [7] "Alphaproteobacteria"
##
                                         "Betaproteobacteria"
##
   [9] "Betaproteobacteria"
## [11] "Proteobacteria_unclassified" "Proteobacteria_unclassified"
plot(as.factor(indicator.bac$Cluster), as.factor(indicator.bac$Phylum))
     Chloroflexi Verrucomicrobia
                                                                                             \infty
                                                                                             9.0
     Acidobacteria
                      1
                                       2
                                               3
                                                                        5
                                                         4
                                                                                     6
```

Beta Diversity Analysis

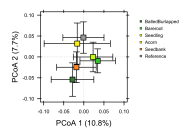
Bray Curtis Ordination (PCoA)

```
png(filename="../figures/WL.bac.PCoA.png",
    width = 1200, height = 800, res = 96*2)
# Create Distance Matrix
sampleREL.dist <- vegdist(WLdataREL, method="bray")
# Principal Coordinates Analysis
WL_pcoa <- cmdscale(sampleREL.dist, k=3, eig=TRUE, add=FALSE)</pre>
```

Χ

```
# Classical (Metric) Multidimensional Scaling; returns PCoA coordinates
  \# eig=TRUE returns eigenvalues; k = \# of dimensions to calculate
# Remove Odd Sites
odd.sites <- row.names(WLdataREL[c(abs(WL_pcoa$points[, 1]) > 0.3), ])
odd.sbys <- WLdataREL[c(abs(WL_pcoa$points[, 1]) > 0.3), ]
mean(rowSums((WLdataREL > 0) * 1))
## [1] 3605.033
rowSums((odd.sbys > 0) * 1)
## HC_A3_ss1_S11 HC_A3_ss2_S12 HC_A3_ss3_S13 HC_C3_ss5_S45 HC_D2_ss3_S53
                           3415
                                          3883
                                                         3340
## HC E1 ss1 S61
            3673
WLdataREL.2 <- WLdataREL[c(abs(WL_pcoaspoints[, 1]) < 0.3), ]</pre>
design2 <- design[c(abs(WL_pcoa$points[, 1]) < 0.3), ]</pre>
treatments <- design2$Treatment</pre>
# Create Distance Matrix
sampleREL.dist2 <- vegdist(WLdataREL.2, method="bray")</pre>
# Principal Coordinates Analysis
WL_pcoa <- cmdscale(sampleREL.dist2, k=2, eig=TRUE, add=FALSE)</pre>
# Classical (Metric) Multidimensional Scaling; returns PCoA coordinates
\# eig=TRUE returns eigenvalues; k = \# of dimensions to calculate
explainvar1 <- round(WL_pcoa$eig[1] / sum(WL_pcoa$eig), 3) * 100
explainvar2 <- round(WL_pcoa$eig[2] / sum(WL_pcoa$eig), 3) * 100</pre>
sum.eig <- sum(explainvar1, explainvar2)</pre>
# Plot
points <- cbind(as.data.frame(WL_pcoa$points), treatments)</pre>
L.centroids <- melt(points, id="treatments", measure.vars = c("V1", "V2"))</pre>
centroids <- cast(L.centroids, variable ~ treatments, mean)</pre>
centroids.se <- cast(L.centroids, variable ~ treatments, se)</pre>
centroids.sd <- cast(L.centroids, variable ~ treatments, sd)</pre>
cent.dataframe <- t(data.frame(rbind(centroids[1,-1], centroids[2,-1],</pre>
                              centroids.sd[1,-1], centroids.sd[2,-1]))
colnames(cent.dataframe) <- c("V1", "V2", "V1e", "V2e")</pre>
cent.treats <- rownames(cent.dataframe)</pre>
# Define Plot Parameters
par(mar = c(5, 5.5, 1, 1) + 0.1)
layout(matrix(1:2, 1, 2), widths = c(4,2))
plot(cent.dataframe[,1], cent.dataframe[,2], type = 'n', las = 1,
     xlim = c(-0.1, 0.1), ylim = c(-0.1, 0.1),
     xaxt = "n", xlab = "", yaxt = "n", ylab="")
abline(h = 0, lty = 3, col = "gray")
abline(v = 0, lty = 3, col = "gray")
```

```
arrows(x0 = cent.dataframe[,1],
       y1 = cent.dataframe[,2] - cent.dataframe[,4],
       y0 = cent.dataframe[,2] + cent.dataframe[,4],
       angle = 90,
       length=0.1, lwd = 2, code = 3)
arrows(y0 = cent.dataframe[,2],
       x1 = cent.dataframe[,1] - cent.dataframe[,3],
       x0 = cent.dataframe[,1] + cent.dataframe[,3],
       angle = 90,
       length=0.1, lwd = 2, code = 3)
points(cent.dataframe[,1], cent.dataframe[,2],
       cex = 2.5, bg = myColors, col = "black", pch = 22, lwd = 2)
# text(cent.dataframe[,1], cent.dataframe[,2], rownames(cent.dataframe))
axis(side = 1, labels = T, las = 1, lwd.ticks = 2)
axis(side = 2, labels = T, las = 1, lwd.ticks = 2)
axis(side=1, lwd.ticks = 2, tck = -0.02, labels=F, cex.axis=1)
axis(side=3, lwd.ticks = 2, tck = -0.02, labels=F, cex.axis=1)
axis(side=1, lwd.ticks = 2, tck = 0.01, labels=F, cex.axis=1)
axis(side=3, lwd.ticks = 2, tck = 0.01, labels=F, cex.axis=1)
axis(side = 2, lwd.ticks = 2, tck = -0.02, labels=F, cex.axis=1)
axis(side = 4, lwd.ticks = 2, tck = -0.02, labels=F, cex.axis=1)
axis(side = 2, lwd.ticks = 2, tck = 0.01, labels=F, cex.axis=1)
axis(side = 4, lwd.ticks = 2, tck = 0.01, labels=F, cex.axis=1)
mtext(paste("PCoA 1 (", explainvar1, "%)", sep = ""), side = 1, line = 3, cex = 1.5)
mtext(paste("PCoA 2 (", explainvar2, "%)", sep = ""), side = 2, line = 3.5, cex = 1.5)
box(lwd = 2)
par(mar = c(5, 0, 1, 1) + 0.1)
plot.new()
legend(0, 1, legend = levels(treatments), pt.bg = myColors, col = "black",
       pch = 22, cex = 0.9, bty = 'n', inset = c(0.1, 0.05),
       y.intersp = 1.25)
# Close Plot Device
dev.off()
## pdf
##
    2
graphics.off()
```

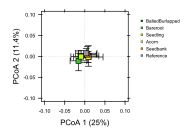


Plot: Microbial Phylogenetic Ordination

```
png(filename="../figures/WL.unifrac.PCoA.png",
    width = 1200, height = 800, res = 96*2)
# Principal Coordinates Analysis
WL_pcoa <- cmdscale(WL.unifrac.dist, k=3, eig=TRUE, add=FALSE)
  # Classical (Metric) Multidimensional Scaling; returns PCoA coordinates
  \# eig=TRUE returns eigenvalues; k = \# of dimensions to calculate
# Remove Odd Sites
odd.sites <- row.names(WL.unifrac[c(abs(WL_pcoa$points[, 1]) > 0.1), ])
# Create Distance Matrix W/O Odd Sites
WL.unifrac.2 <- WL.unifrac[-c(which(row.names(WL.unifrac) %in% odd.sites)),
                            -c(which(row.names(WL.unifrac) %in% odd.sites))]
WL.unifrac.dist2 <- as.dist(WL.unifrac.2, upper = T, diag = T)</pre>
treatments <- design[-c(which(row.names(design) %in% odd.sites)), ]$Treatment
# Principal Coordinates Analysis
WL_pcoa <- cmdscale(WL.unifrac.dist2, k=2, eig=TRUE, add=FALSE)
# Classical (Metric) Multidimensional Scaling; returns PCoA coordinates
\# eig=TRUE returns eigenvalues; k = \# of dimensions to calculate
explainvar1 <- round(WL pcoa$eig[1] / sum(WL pcoa$eig), 3) * 100
explainvar2 <- round(WL_pcoa$eig[2] / sum(WL_pcoa$eig), 3) * 100
sum.eig <- sum(explainvar1, explainvar2)</pre>
# Plot
points <- cbind(as.data.frame(WL_pcoa$points), treatments)</pre>
L.centroids <- melt(points, id="treatments", measure.vars = c("V1", "V2"))
centroids <- cast(L.centroids, variable ~ treatments, mean)</pre>
centroids.se <- cast(L.centroids, variable ~ treatments, se)</pre>
centroids.sd <- cast(L.centroids, variable ~ treatments, sd)</pre>
cent.dataframe <- t(data.frame(rbind(centroids[1,-1], centroids[2,-1],</pre>
                              centroids.sd[1,-1],centroids.sd[2,-1])))
colnames(cent.dataframe) <- c("V1", "V2", "V1e", "V2e")</pre>
```

```
cent.treats <- rownames(cent.dataframe)</pre>
# Define Plot Parameters
par(mar = c(5, 5.5, 1, 1) + 0.1)
layout(matrix(1:2, 1, 2), widths = c(4,2))
plot(cent.dataframe[,1], cent.dataframe[,2], type = 'n', las = 1,
     xlim = c(-0.1, 0.1), ylim = c(-0.1, 0.1),
     xaxt = "n", xlab = "", yaxt = "n", ylab="")
abline(h = 0, lty = 3, col = "gray")
abline(v = 0, lty = 3, col = "gray")
arrows(x0 = cent.dataframe[,1],
       y1 = cent.dataframe[,2] - cent.dataframe[,4],
       y0 = cent.dataframe[,2] + cent.dataframe[,4],
       angle = 90,
       length=0.1, lwd = 2, code = 3)
arrows(y0 = cent.dataframe[,2],
       x1 = cent.dataframe[,1] - cent.dataframe[,3],
       x0 = cent.dataframe[,1] + cent.dataframe[,3],
       angle = 90,
       length=0.1, lwd = 2, code = 3)
points(cent.dataframe[,1], cent.dataframe[,2],
       cex = 2.5, bg = myColors, col = "black", pch = 22, lwd = 2)
#text(cent.dataframe[,1], cent.dataframe[,2], rownames(cent.dataframe))
axis(side = 1, labels = T, las = 1, lwd.ticks = 2)
axis(side = 2, labels = T, las = 1, lwd.ticks = 2)
axis(side=1, lwd.ticks = 2, tck = -0.02, labels=F, cex.axis=1)
axis(side=3, lwd.ticks = 2, tck = -0.02, labels=F, cex.axis=1)
axis(side=1, lwd.ticks = 2, tck = 0.01, labels=F, cex.axis=1)
axis(side=3, lwd.ticks = 2, tck = 0.01, labels=F, cex.axis=1)
axis(side = 2, lwd.ticks = 2, tck = -0.02, labels=F, cex.axis=1)
axis(side = 4, lwd.ticks = 2, tck = -0.02, labels=F, cex.axis=1)
axis(side = 2, lwd.ticks = 2, tck = 0.01, labels=F, cex.axis=1)
axis(side = 4, lwd.ticks = 2, tck = 0.01, labels=F, cex.axis=1)
mtext(paste("PCoA 1 (", explainvar1, "%)", sep = ""), side = 1, line = 3, cex = 1.5)
mtext(paste("PCoA 2 (", explainvar2, "%)", sep = ""), side = 2, line = 3.5, cex = 1.5)
box(1wd = 2)
par(mar = c(5, 0, 1, 1) + 0.1)
plot.new()
legend(0, 1, legend = cent.treats, pt.bg = myColors, col = "black",
       pch = 22, cex = 0.9, bty = 'n', inset = c(0.1, 0.05),
       y.intersp = 1.25)
# Close Plot Device
```

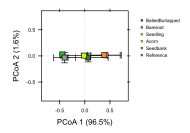
```
dev.off()
## pdf
## 2
graphics.off()
```



Plot: Plant Principal Coordinates Ordination

```
png(filename="../figures/WL.plant.PCoA.png",
    width = 1200, height = 800, res = 96*2)
# Create Distance Matrix
treatments <- design$Treatment</pre>
samplePlant.dist <- vegdist(WL.plant, method="bray")</pre>
# Principal Coordinates Analysis
WLplant_pcoa <- cmdscale(samplePlant.dist, k=3, eig=TRUE, add=FALSE)</pre>
  # Classical (Metric) Multidimensional Scaling; returns PCoA coordinates
  \# eig=TRUE returns eigenvalues; k = \# of dimensions to calculate
# # Remove Odd Sites
# odd.sites <- row.names(WLdataREL[c(abs(WL_pcoa$points[, 1]) > 0.3), ])
# odd.sbys <- WLdataREL[c(abs(WL_pcoa$points[, 1]) > 0.3), ]
# mean(rowSums((WLdataREL > 0) * 1))
# rowSums((odd.sbys > 0) * 1)
# WLdataREL.2 <- WLdataREL[c(abs(WL_pcoa$points[, 1]) < 0.3), ]</pre>
# design2 \leftarrow design[c(abs(WL_pcoa$points[, 1]) < 0.3), ]
# treatments <- design2$Treatment</pre>
# Create Distance Matrix
#sampleREL.dist2 <- vegdist(WLdataREL.2, method="bray")</pre>
# Principal Coordinates Analysis
#WL_pcoa <- cmdscale(sampleREL.dist2, k=2, eig=TRUE, add=FALSE)</pre>
# Classical (Metric) Multidimensional Scaling; returns PCoA coordinates
\# eig=TRUE returns eigenvalues; k = \# of dimensions to calculate
explainvar1 <- round(WLplant_pcoa$eig[1] / sum(WLplant_pcoa$eig), 3) * 100
explainvar2 <- round(WLplant_pcoa$eig[2] / sum(WLplant_pcoa$eig), 3) * 100
sum.eig <- sum(explainvar1, explainvar2)</pre>
```

```
# Plot
points <- cbind(as.data.frame(WLplant_pcoa$points), treatments)</pre>
L.centroids <- melt(points, id="treatments", measure.vars = c("V1", "V2"))
centroids <- cast(L.centroids, variable ~ treatments, mean)</pre>
centroids.se <- cast(L.centroids, variable ~ treatments, se)</pre>
centroids.sd <- cast(L.centroids, variable ~ treatments, sd)</pre>
cent.dataframe <- t(data.frame(rbind(centroids[1,-1], centroids[2,-1],</pre>
                              centroids.sd[1,-1], centroids.sd[2,-1]))
colnames(cent.dataframe) <- c("V1", "V2", "V1e", "V2e")</pre>
cent.treats <- rownames(cent.dataframe)</pre>
# Define Plot Parameters
par(mar = c(5, 5.5, 1, 1) + 0.1)
layout(matrix(1:2, 1, 2), widths = c(4,2))
plot(cent.dataframe[,1], cent.dataframe[,2], type = 'n', las = 1,
     xlim = c(-0.8, 0.8), ylim = c(-0.8, 0.8),
     xaxt = "n", xlab = "", yaxt = "n", ylab="")
abline(h = 0, lty = 3, col = "gray")
abline(v = 0, lty = 3, col = "gray")
arrows(x0 = cent.dataframe[,1],
       y1 = cent.dataframe[,2] - cent.dataframe[,4],
       y0 = cent.dataframe[,2] + cent.dataframe[,4],
       angle = 90,
       length=0.1, lwd = 2, code = 3)
arrows(y0 = cent.dataframe[,2],
       x1 = cent.dataframe[,1] - cent.dataframe[,3],
       x0 = cent.dataframe[,1] + cent.dataframe[,3],
       angle = 90,
       length=0.1, lwd = 2, code = 3)
points(cent.dataframe[,1], cent.dataframe[,2],
       cex = 2.5, bg = myColors, col = "black", pch = 22, lwd = 2)
axis(side = 1, labels = T, las = 1, lwd.ticks = 2)
axis(side = 2, labels = T, las = 1, lwd.ticks = 2)
axis(side=1, lwd.ticks = 2, tck = -0.02, labels=F, cex.axis=1)
axis(side=3, lwd.ticks = 2, tck = -0.02, labels=F, cex.axis=1)
axis(side=1, lwd.ticks = 2, tck = 0.01, labels=F, cex.axis=1)
axis(side=3, lwd.ticks = 2, tck = 0.01, labels=F, cex.axis=1)
axis(side = 2, lwd.ticks = 2, tck = -0.02, labels=F, cex.axis=1)
axis(side = 4, lwd.ticks = 2, tck = -0.02, labels=F, cex.axis=1)
axis(side = 2, lwd.ticks = 2, tck = 0.01, labels=F, cex.axis=1)
axis(side = 4, lwd.ticks = 2, tck = 0.01, labels=F, cex.axis=1)
mtext(paste("PCoA 1 (", explainvar1, "%)", sep = ""), side = 1, line = 3, cex = 1.5)
mtext(paste("PCoA 2 (", explainvar2, "%)", sep = ""), side = 2, line = 3.5, cex = 1.5)
box(lwd = 2)
par(mar = c(5, 0, 1, 1) + 0.1)
```



Joint Plot: Microbial and Plant Bray Curtis Ordination (PCoA)

```
png(filename="../figures/WL.PCoA.png",
    width = 1800, height = 800, res = 96*2)
layout(matrix(1:3, 1, 3, byrow = T), widths = c(4, 4, 1.5))
# Define Plot Parameters
par(mar = c(5, 6, 1, 2), oma = c(1, 1, 1, 1))
# Plant PcoA
# Create Distance Matrix
treatments <- design$Treatment</pre>
samplePlant.dist <- vegdist(WL.plant, method="bray")</pre>
# Principal Coordinates Analysis
WLplant_pcoa <- cmdscale(samplePlant.dist, k=3, eig=TRUE, add=FALSE)
 # Classical (Metric) Multidimensional Scaling; returns PCoA coordinates
  \# eig=TRUE returns eigenvalues; k = \# of dimensions to calculate
explainvar1 <- round(WLplant_pcoa$eig[1] / sum(WLplant_pcoa$eig), 3) * 100
explainvar2 <- round(WLplant_pcoa$eig[2] / sum(WLplant_pcoa$eig), 3) * 100
sum.eig <- sum(explainvar1, explainvar2)</pre>
# Plot
```

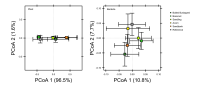
```
points <- cbind(as.data.frame(WLplant_pcoa$points), treatments)</pre>
L.centroids <- melt(points, id="treatments", measure.vars = c("V1", "V2"))
centroids <- cast(L.centroids, variable ~ treatments, mean)</pre>
centroids.se <- cast(L.centroids, variable ~ treatments, se)</pre>
centroids.sd <- cast(L.centroids, variable ~ treatments, sd)</pre>
cent.dataframe <- t(data.frame(rbind(centroids[1,-1], centroids[2,-1],</pre>
                             centroids.sd[1,-1],centroids.sd[2,-1])))
colnames(cent.dataframe) <- c("V1", "V2", "V1e", "V2e")</pre>
cent.treats <- rownames(cent.dataframe)</pre>
# Initiate Plot
plot(cent.dataframe[,1], cent.dataframe[,2], type = 'n', las = 1,
     xlim = c(-0.8, 0.8), ylim = c(-0.8, 0.8),
     xaxt = "n", xlab = "", yaxt = "n", ylab="")
abline(h = 0, lty = 3, col = "gray")
abline(v = 0, lty = 3, col = "gray")
arrows(x0 = cent.dataframe[,1],
       y1 = cent.dataframe[,2] - cent.dataframe[,4],
       y0 = cent.dataframe[,2] + cent.dataframe[,4],
       angle = 90,
       length=0.1, lwd = 2, code = 3)
arrows(y0 = cent.dataframe[,2],
       x1 = cent.dataframe[,1] - cent.dataframe[,3],
       x0 = cent.dataframe[,1] + cent.dataframe[,3],
       angle = 90,
       length=0.1, lwd = 2, code = 3)
points(cent.dataframe[,1], cent.dataframe[,2],
       cex = 2.5, bg = myColors, col = "black", pch = 22, lwd = 2)
range.x <- par("usr")[2] - par("usr")[1]</pre>
range.y <- par("usr")[4] - par("usr")[3]
text(par("usr")[1] + 0.1 * range.x, par("usr")[4] - 0.05 * range.y, "Plant")
axis(side = 1, labels = T, las = 1, lwd.ticks = 2)
axis(side = 2, labels = T, las = 1, lwd.ticks = 2)
axis(side=1, lwd.ticks = 2, tck = -0.02, labels=F, cex.axis=1)
axis(side=3, lwd.ticks = 2, tck = -0.02, labels=F, cex.axis=1)
axis(side=1, lwd.ticks = 2, tck = 0.01, labels=F, cex.axis=1)
axis(side=3, lwd.ticks = 2, tck = 0.01, labels=F, cex.axis=1)
axis(side = 2, lwd.ticks = 2, tck = -0.02, labels=F, cex.axis=1)
axis(side = 4, lwd.ticks = 2, tck = -0.02, labels=F, cex.axis=1)
axis(side = 2, lwd.ticks = 2, tck = 0.01, labels=F, cex.axis=1)
axis(side = 4, lwd.ticks = 2, tck = 0.01, labels=F, cex.axis=1)
mtext(paste("PCoA 1 (", explainvar1, "%)", sep = ""), side = 1,
      line = 3.5, cex = 1.5)
mtext(paste("PCoA 2 (", explainvar2, "%)", sep = ""), side = 2,
      line = 3.5, cex = 1.5)
box(lwd = 2)
```

```
# Bacterial Principal Coordinates Analysis
# Create Distance Matrix
sampleREL.dist <- vegdist(WLdataREL, method="bray")</pre>
WL_pcoa <- cmdscale(sampleREL.dist, k=3, eig=TRUE, add=FALSE)</pre>
  # Classical (Metric) Multidimensional Scaling; returns PCoA coordinates
  \# eig=TRUE returns eigenvalues; k = \# of dimensions to calculate
# Remove Odd Sites
odd.sites <- row.names(WLdataREL[c(abs(WL_pcoa$points[, 1]) > 0.3), ])
odd.sbys <- WLdataREL[c(abs(WL_pcoa$points[, 1]) > 0.3), ]
mean(rowSums((WLdataREL > 0) * 1))
## [1] 3605.033
rowSums((odd.sbys > 0) * 1)
## HC_A3_ss1_S11 HC_A3_ss2_S12 HC_A3_ss3_S13 HC_C3_ss5_S45 HC_D2_ss3_S53
            3907
                           3415
                                          3883
                                                         3340
## HC_E1_ss1_S61
            3673
WLdataREL.2 <- WLdataREL[c(abs(WL_pcoa$points[, 1]) < 0.3), ]</pre>
design2 <- design[c(abs(WL_pcoa$points[, 1]) < 0.3), ]</pre>
treatments <- design2$Treatment</pre>
# Create Distance Matrix
sampleREL.dist2 <- vegdist(WLdataREL.2, method="bray")</pre>
# Principal Coordinates Analysis
WL_pcoa <- cmdscale(sampleREL.dist2, k=2, eig=TRUE, add=FALSE)</pre>
# Classical (Metric) Multidimensional Scaling; returns PCoA coordinates
\# eig=TRUE returns eigenvalues; k = \# of dimensions to calculate
explainvar1 <- round(WL_pcoa$eig[1] / sum(WL_pcoa$eig), 3) * 100
explainvar2 <- round(WL_pcoa$eig[2] / sum(WL_pcoa$eig), 3) * 100
sum.eig <- sum(explainvar1, explainvar2)</pre>
points <- cbind(as.data.frame(WL_pcoa$points), treatments)</pre>
L.centroids <- melt(points, id="treatments", measure.vars = c("V1", "V2"))
centroids <- cast(L.centroids, variable ~ treatments, mean)</pre>
centroids.se <- cast(L.centroids, variable ~ treatments, se)</pre>
centroids.sd <- cast(L.centroids, variable ~ treatments, sd)</pre>
cent.dataframe <- t(data.frame(rbind(centroids[1,-1], centroids[2,-1],
                              centroids.sd[1,-1],centroids.sd[2,-1])))
colnames(cent.dataframe) <- c("V1", "V2", "V1e", "V2e")</pre>
cent.treats <- rownames(cent.dataframe)</pre>
# Bacterial Plot
plot(cent.dataframe[,1], cent.dataframe[,2], type = 'n', las = 1,
     xlim = c(-0.1, 0.1), ylim = c(-0.1, 0.1),
     xaxt = "n", xlab = "", yaxt = "n", ylab="")
```

```
abline(h = 0, lty = 3, col = "gray")
abline(v = 0, lty = 3, col = "gray")
arrows(x0 = cent.dataframe[,1],
       y1 = cent.dataframe[,2] - cent.dataframe[,4],
       y0 = cent.dataframe[,2] + cent.dataframe[,4],
       angle = 90,
       length=0.1, lwd = 2, code = 3)
arrows(y0 = cent.dataframe[,2],
       x1 = cent.dataframe[,1] - cent.dataframe[,3],
       x0 = cent.dataframe[,1] + cent.dataframe[,3],
       angle = 90,
       length=0.1, lwd = 2, code = 3)
points(cent.dataframe[,1], cent.dataframe[,2],
       cex = 2.5, bg = myColors, col = "black", pch = 22, lwd = 2)
range.x <- par("usr")[2] - par("usr")[1]</pre>
range.y <- par("usr")[4] - par("usr")[3]</pre>
text(par("usr")[1] + 0.13 * range.x, par("usr")[4] - 0.05 * range.y, "Bacteria")
# text(cent.dataframe[,1], cent.dataframe[,2], rownames(cent.dataframe))
axis(side = 1, labels = T, las = 1, lwd.ticks = 2)
axis(side = 2, labels = T, las = 1, lwd.ticks = 2)
axis(side=1, lwd.ticks = 2, tck = -0.02, labels=F, cex.axis=1)
axis(side=3, lwd.ticks = 2, tck = -0.02, labels=F, cex.axis=1)
axis(side=1, lwd.ticks = 2, tck = 0.01, labels=F, cex.axis=1)
axis(side=3, lwd.ticks = 2, tck = 0.01, labels=F, cex.axis=1)
axis(side = 2, lwd.ticks = 2, tck = -0.02, labels=F, cex.axis=1)
axis(side = 4, lwd.ticks = 2, tck = -0.02, labels=F, cex.axis=1)
axis(side = 2, lwd.ticks = 2, tck = 0.01, labels=F, cex.axis=1)
axis(side = 4, lwd.ticks = 2, tck = 0.01, labels=F, cex.axis=1)
mtext(paste("PCoA 1 (", explainvar1, "%)", sep = ""), side = 1,
      line = 3.5, cex = 1.5)
mtext(paste("PCoA 2 (", explainvar2, "%)", sep = ""), side = 2,
      line = 3.5, cex = 1.5)
box(1wd = 2)
par(mar = c(5, 0, 1, 0) + 0.5)
plot.new()
legend(0, 1, legend = c("Balled Burlapped", cent.treats[-1]),
       pt.bg = myColors, col = "black",
       pch = 22, cex = 1, bty = 'n', inset = c(0.1, 0.05),
       y.intersp = 1.25)
# Close Plot Device
dev.off()
## pdf
```

2

```
graphics.off()
```



Mantel Tests

Similarity between microbe and plant communities

```
# Name check
all.equal(rownames(WL.plant), rownames(WLdataREL))
## [1] TRUE
# Remove Odd Sites from Bac Community
bac.comm <- WLdataREL[-which(rownames(WLdataREL) %in% odd.sites),]
plant.comm <- WL.plant[-which(rownames(WL.plant) %in% odd.sites), ]</pre>
# Remove Bare Treatment
bare.sites <- rownames(design[design$Treatment == "Bareroot", ])</pre>
#bac.comm <- bac.comm[-which(rownames(bac.comm) %in% bare.sites),]</pre>
#plant.comm <- plant.comm[-which(rownames(plant.comm) %in% bare.sites), ]</pre>
all.equal(rownames(bac.comm), rownames(plant.comm))
## [1] TRUE
#rowSums(plant.comm)
bac.comm <- bac.comm[, colSums(bac.comm) > 0]
plant.comm <- plant.comm[, colSums(plant.comm) > 0]
dist.bac <- vegdist(bac.comm, method = "bray")</pre>
dist.plant <- vegdist(plant.comm, method = "bray")</pre>
mantel.rtest(dist.bac, dist.plant, nrepet = 999)
## Warning in is.euclid(m2): Zero distance(s)
## Monte-Carlo test
## Observation: 0.09849538
## Call: mantelnoneuclid(m1 = m1, m2 = m2, nrepet = nrepet)
## Based on 999 replicates
## Simulated p-value: 0.001
WL.plant_points <- cmdscale(dist.plant, k=3, eig=TRUE, add=FALSE)$points
```

```
dbRDA <- dbrda(bac.comm ~ WL.plant_points[, 1], distance = "bray")</pre>
anova(dbRDA)
## Permutation test for dbrda under reduced model
## Permutation: free
## Number of permutations: 999
## Model: dbrda(formula = bac.comm ~ WL.plant_points[, 1], distance = "bray")
           Df SumOfSqs
                            F Pr(>F)
##
            1 0.09609 3.0644 0.001 ***
## Model
## Residual 82 2.57134
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
RsquareAdj(dbRDA)
## $r.squared
## [1] 0.03602451
##
## $adj.r.squared
## [1] 0.02426871
```

Similarity between microbes and soil

```
# Remove Odd Sites from Bac Community
bac.comm <- WLdataREL[-which(rownames(WLdataREL) %in% odd.sites),]</pre>
plant.comm <- WL.plant[-which(rownames(WL.plant) %in% odd.sites), ]</pre>
# Remove Zero Sum Columns
bac.comm <- bac.comm[, colSums(bac.comm) > 0]
plant.comm <- plant.comm[, colSums(plant.comm) > 0]
# Remove Odd Sites from Soil
WL.soil2 <- WL.soil[-which(rownames(WLdataREL) %in% odd.sites),]</pre>
WL.soil.phys <- WL.soil2[, which(colnames(WL.soil2) %in%
                                     c("Moisture", "Temp", "pH"))]
WL.soil.nuts <- WL.soil2[, which(colnames(WL.soil2) %in%
                                     c("TOC", "TN", "OM", "NH4.N", "NO3.N"))]
WL.soil.all <- WL.soil2[, which(colnames(WL.soil2) %in%
                               c("Moisture", "Temp", "pH", "TOC", "TN",
                                  "OM", "NH4.N", "NO3.N"))]
all.equal(grep("[0-9][0-9]$", rownames(WL.soil2)),
          grep("S*$", rownames(bac.comm)))
## [1] TRUE
dist.bac <- vegdist(bac.comm, method = "bray")</pre>
dist.plant <- vegdist(plant.comm, method = "bray")</pre>
dist.nuts <- vegdist(WL.soil.nuts, method = "euclidean")</pre>
dist.phys <- vegdist(WL.soil.phys, method = "euclidean")</pre>
dist.soil <- vegdist(WL.soil.all, method = "euclidean")</pre>
```

```
mantel.rtest(dist.bac, dist.plant, nrepet = 999)
## Warning in is.euclid(m2): Zero distance(s)
## Monte-Carlo test
## Observation: 0.09849538
## Call: mantelnoneuclid(m1 = m1, m2 = m2, nrepet = nrepet)
## Based on 999 replicates
## Simulated p-value: 0.001
mantel.rtest(dist.bac, dist.nuts, nrepet = 999)
## Monte-Carlo test
## Observation: -0.04139427
## Call: mantel.rtest(m1 = dist.bac, m2 = dist.nuts, nrepet = 999)
## Based on 999 replicates
## Simulated p-value: 0.688
mantel.rtest(dist.bac, dist.phys, nrepet = 999)
## Monte-Carlo test
## Observation: 0.1637996
## Call: mantel.rtest(m1 = dist.bac, m2 = dist.phys, nrepet = 999)
## Based on 999 replicates
## Simulated p-value: 0.021
mantel.rtest(dist.bac, dist.soil, nrepet = 999)
## Monte-Carlo test
## Observation: -0.01610846
## Call: mantel.rtest(m1 = dist.bac, m2 = dist.soil, nrepet = 999)
## Based on 999 replicates
## Simulated p-value: 0.538
mantel.rtest(dist.plant, dist.nuts, nrepet = 999)
## Warning in is.euclid(m1): Zero distance(s)
## Monte-Carlo test
## Observation: 0.265419
## Call: mantelnoneuclid(m1 = m1, m2 = m2, nrepet = nrepet)
## Based on 999 replicates
## Simulated p-value: 0.001
mantel.rtest(dist.plant, dist.phys, nrepet = 999)
## Warning in is.euclid(m1): Zero distance(s)
## Monte-Carlo test
## Observation: 0.04439773
## Call: mantelnoneuclid(m1 = m1, m2 = m2, nrepet = nrepet)
## Based on 999 replicates
## Simulated p-value: 0.023
mantel.rtest(dist.plant, dist.soil, nrepet = 999)
## Warning in is.euclid(m1): Zero distance(s)
## Monte-Carlo test
## Observation: 0.2481641
```

```
## Call: mantelnoneuclid(m1 = m1, m2 = m2, nrepet = nrepet)
## Based on 999 replicates
## Simulated p-value: 0.001
```

Distance Based RDA Models

```
# Nutrients
chem.dist <- dist(apply(WL.soil.nuts, 2, scale),</pre>
                  method = "euclidean")
chem.pcoa <- cmdscale(chem.dist, k = 3, eig = T)</pre>
explainvar1 <- round(chem.pcoa$eig[1] / sum(chem.pcoa$eig), 3) * 100
explainvar2 <- round(chem.pcoa$eig[2] / sum(chem.pcoa$eig), 3) * 100
# Physical Factors
phys.dist <- dist(apply(WL.soil.phys, 2, scale),</pre>
                  method = "euclidean")
phys.pcoa <- cmdscale(phys.dist, k = 3, eig = T)</pre>
explainvar1 <- round(phys.pcoa$eig[1] / sum(phys.pcoa$eig), 3) * 100
explainvar2 <- round(phys.pcoa$eig[2] / sum(phys.pcoa$eig), 3) * 100
# Bac dbRDA
dbRDA.c <- dbrda(bac.comm ~ chem.pcoa$points[,1], distance = "bray")
anova(dbRDA.c, by = "terms", model = "direct")
## Permutation test for dbrda under direct model
## Terms added sequentially (first to last)
## Permutation: free
## Number of permutations: 999
## Model: dbrda(formula = bac.comm ~ chem.pcoa$points[, 1], distance = "bray")
                         Df SumOfSqs
                                          F Pr(>F)
## chem.pcoa$points[, 1] 1 0.08661 2.7518 0.001 ***
## Residual
                         82 2.58082
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
RsquareAdj(dbRDA.c)
## $r.squared
## [1] 0.03246874
##
## $adj.r.squared
## [1] 0.02066957
dbRDA.p <- dbrda(bac.comm ~ phys.pcoa$points[,1], distance = "bray")
anova(dbRDA.p)
## Permutation test for dbrda under reduced model
## Permutation: free
## Number of permutations: 999
##
## Model: dbrda(formula = bac.comm ~ phys.pcoa$points[, 1], distance = "bray")
```

```
Df SumOfSqs
                           F Pr(>F)
            1 0.07508 2.375 0.001 ***
## Model
## Residual 82 2.59235
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
RsquareAdj(dbRDA.p)
## $r.squared
## [1] 0.02814833
## $adj.r.squared
## [1] 0.01629648
#error here ???
dbRDA.cp <- dbrda(bac.comm ~ chem.pcoa$points[,1] * phys.pcoa$points[,],
              distance = "bray", add = T)
anova(dbRDA.cp, by = "terms", model = "direct")
## Permutation test for dbrda under direct model
## Terms added sequentially (first to last)
## Permutation: free
## Number of permutations: 999
## Model: dbrda(formula = bac.comm ~ chem.pcoa$points[, 1] * phys.pcoa$points[, ], distance = "bray", a
                                             Df SumOfSqs
                                                              F Pr(>F)
                                              1 0.08661 2.8960 0.001 ***
## chem.pcoa$points[, 1]
## phys.pcoa$points[, ]
                                              3 0.19182 2.1380 0.001 ***
## chem.pcoa$points[, 1]:phys.pcoa$points[, ] 3 0.11616 1.2947 0.018 *
## Residual
                                             76 2.27285
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
RsquareAdj(dbRDA.cp)
## $r.squared
## [1] 0.1479263
## $adj.r.squared
## [1] 0.0694458
dbRDA.P.cp <- dbrda(plant.comm ~ phys.pcoa$points[,1] * chem.pcoa$points[,1],
              distance = "bray", add = T)
anova(dbRDA.P.cp, by = "terms", model = "direct")
## Permutation test for dbrda under direct model
## Terms added sequentially (first to last)
## Permutation: free
## Number of permutations: 999
## Model: dbrda(formula = plant.comm ~ phys.pcoa$points[, 1] * chem.pcoa$points[, 1], distance = "bray"
                                              Df SumOfSqs
                                                                F Pr(>F)
## phys.pcoa$points[, 1]
                                                   0.2752 1.1953 0.282
## chem.pcoa$points[, 1]
                                               1
                                                  3.0843 13.3955 0.001 ***
## phys.pcoa$points[, 1]:chem.pcoa$points[, 1] 1
                                                  1.0717 4.6545 0.024 *
## Residual
                                              80 18.4197
## ---
```

```
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
RsquareAdj(dbRDA.P.cp)

## $r.squared
## [1] 0.1939162
##
## $adj.r.squared
## [1] 0.163688
```

PostHoc Tests

```
phys.mod <- lm(phys.pcoa$points[,1] ~ WL.soil2$Treatment)</pre>
chem.mod <- lm(chem.pcoa$points[,1] ~ WL.soil2$Treatment)</pre>
anova(phys.mod)
## Analysis of Variance Table
## Response: phys.pcoa$points[, 1]
                     Df Sum Sq Mean Sq F value Pr(>F)
## WL.soil2$Treatment 5 14.534 2.9068 2.1447 0.06879 .
## Residuals
                      78 105.717 1.3554
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
anova (chem.mod)
## Analysis of Variance Table
##
## Response: chem.pcoa$points[, 1]
                      Df Sum Sq Mean Sq F value
## WL.soil2$Treatment 5 73.39 14.678 6.7206 2.959e-05 ***
## Residuals
                     78 170.35
                                  2.184
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
TukeyHSD(aov(phys.mod))
##
     Tukey multiple comparisons of means
##
       95% family-wise confidence level
##
## Fit: aov(formula = phys.mod)
##
## $`WL.soil2$Treatment`
##
                                   diff
                                                lwr
                                                          upr
                                                                  p adj
## Bareroot-BalledBurlapped
                              0.9960375 -0.32135712 2.3134322 0.2454950
## Seedling-BalledBurlapped
                              1.3681678 0.03002546 2.7063100 0.0421120
## Acorn-BalledBurlapped
                              1.1715954 -0.16654691 2.5097377 0.1203240
## Seedbank-BalledBurlapped
                              0.6934262 -0.64471608 2.0315685 0.6563663
## Reference-BalledBurlapped 0.8241642 -0.49323048 2.1415588 0.4542536
## Seedling-Bareroot
                              0.3721302 -0.89190627 1.6361667 0.9547234
## Acorn-Bareroot
                              0.1755579 -1.08847864 1.4395944 0.9985335
## Seedbank-Bareroot
                             -0.3026113 -1.56664780 0.9614252 0.9814775
## Reference-Bareroot
                             -0.1718734 -1.41392493 1.0701782 0.9985591
```

```
## Acorn-Seedling
                            -0.1965724 -1.48221791 1.0890732 0.9976763
## Seedbank-Seedling
                            -0.6747415 -1.96038707 0.6109040 0.6440150
## Reference-Seedling
                            -0.5440036 -1.80804009 0.7200329 0.8068369
## Seedbank-Acorn
                            -0.4781692 -1.76381469 0.8074764 0.8852677
## Reference-Acorn
                            -0.3474312 -1.61146771 0.9166053 0.9661433
## Reference-Seedbank
                             0.1307379 -1.13329855 1.3947745 0.9996494
TukeyHSD(aov(chem.mod))
##
     Tukey multiple comparisons of means
##
       95% family-wise confidence level
##
## Fit: aov(formula = chem.mod)
##
## $`WL.soil2$Treatment`
##
                                   diff
                                               lwr
                                                          upr
                                                                  p adj
## Bareroot-BalledBurlapped
                              1.4379873 -0.2343365 3.1103111 0.1331238
## Seedling-BalledBurlapped
                              0.8114554 -0.8872059 2.5101166 0.7294419
## Acorn-BalledBurlapped
                             -0.4487753 -2.1474366
                                                   1.2498859 0.9714488
## Seedbank-BalledBurlapped -1.3101913 -3.0088525 0.3884700 0.2258638
## Reference-BalledBurlapped -0.6494620 -2.3217858 1.0228618 0.8653924
## Seedling-Bareroot
                             -0.6265319 -2.2311220 0.9780582 0.8627205
## Acorn-Bareroot
                            -1.8867626 -3.4913527 -0.2821726 0.0118037
## Seedbank-Bareroot
                            -2.7481786 -4.3527686 -1.1435885 0.0000489
## Reference-Bareroot
                            -2.0874493 -3.6641313 -0.5107672 0.0029948
                            -1.2602307 -2.8922516 0.3717902 0.2247611
## Acorn-Seedling
## Seedbank-Seedling
                            -2.1216467 -3.7536676 -0.4896257 0.0037703
## Reference-Seedling
                            -1.4609174 -3.0655074 0.1436727 0.0952960
## Seedbank-Acorn
                            -0.8614159 -2.4934369 0.7706050 0.6384076
## Reference-Acorn
                             -0.2006866 -1.8052767
                                                   1.4039034 0.9991156
## Reference-Seedbank
                             0.6607293 -0.9438608 2.2653194 0.8340697
```

Invasive Species Biomass

```
##
                   Median
                1Q
                                3Q
                             7.307 122.081
## -67.006 -17.864
                   -0.514
##
## Coefficients:
##
                     Estimate Std. Error t value Pr(>|t|)
                                  8.113
                                           1.019
                                                    0.312
## (Intercept)
                        8.269
## TreatmentBareroot
                       -8.190
                                  11.474 -0.714
                                                    0.478
## TreatmentSeedling
                       13.460
                                  11.474
                                           1.173
                                                    0.245
## TreatmentAcorn
                       48.649
                                  11.677
                                           4.166 8.82e-05 ***
## TreatmentSeedbank
                       58.737
                                  11.474
                                           5.119 2.64e-06 ***
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## Residual standard error: 31.42 on 69 degrees of freedom
     (1 observation deleted due to missingness)
## Multiple R-squared: 0.4339, Adjusted R-squared: 0.401
## F-statistic: 13.22 on 4 and 69 DF, p-value: 4.775e-08
anova(RC.mod)
## Analysis of Variance Table
##
## Response: RCGbiomass
##
             Df Sum Sq Mean Sq F value
                                          Pr(>F)
## Treatment 4 52209 13052.2
                                 13.22 4.775e-08 ***
## Residuals 69
                68125
                         987.3
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
TukeyHSD(aov(RC.mod))
##
     Tukey multiple comparisons of means
       95% family-wise confidence level
##
##
## Fit: aov(formula = RC.mod)
##
## $Treatment
##
                                diff
                                           lwr
                                                    upr
                                                            p adj
## Bareroot-BalledBurlapped -8.19000 -40.33004 23.95004 0.9526582
## Seedling-BalledBurlapped 13.46000 -18.68004 45.60004 0.7665404
## Acorn-BalledBurlapped
                            48.64852 15.93959 81.35746 0.0008175
## Seedbank-BalledBurlapped 58.73667 26.59663 90.87671 0.0000256
## Seedling-Bareroot
                            21.65000 -10.49004 53.79004 0.3340189
## Acorn-Bareroot
                            56.83852 24.12959 89.54746 0.0000662
## Seedbank-Bareroot
                            66.92667
                                      34.78663 99.06671 0.0000016
## Acorn-Seedling
                            35.18852
                                       2.47959 67.89746 0.0287522
## Seedbank-Seedling
                            45.27667 13.13663 77.41671 0.0017170
## Seedbank-Acorn
                            10.08814 -22.62079 42.79708 0.9090114
tuk <- HSD.test(lm(WL.soil.RCGbiomass ~ WL.soil.Treatment,</pre>
                   data = WL.rc), "WL.soil.Treatment")
```

Plot: Invasive Species

```
png(filename="../figures/reed.biomass.png",
    width = 1200, height = 800, res = 96*2)
par(mar = (c(6.5,5,1,1) + 0.1))
rcg <- boxplot(WL.rc$WL.soil.RCGbiomass ~ WL.rc$WL.soil.Treatment,</pre>
        col = myColors, xaxt = "n", yaxt = "n",
        xlab = "", ylab = "")
# Lables
mtext("Reed Canary Biomass (g)", side = 2, cex = 1.25, line = 3)
# PostHoc Test
tuk <- HSD.test(lm(WL.soil.RCGbiomass ~ WL.soil.Treatment,</pre>
                   data = WL.rc), "WL.soil.Treatment")
grp <- tuk$groups[c(match(gsub(" ", "", tuk$groups$trt),</pre>
                          levels(WL.rc$WL.soil.Treatment))), ]
text(x = seq_along(levels(WL.rc$WL.soil.Treatment)),
     y = rcg\$stats[5, ] + ((par("usr")[4] - par("usr")[3]) * 0.05),
     labels = grp$M)
# Plot x labs at default x position
text(x = seq_along(levels(WL.rc$WL.soil.Treatment)),
     y = par("usr")[3] - 10, srt = 45, adj = 1,
     labels = levels(WL.rc$WL.soil.Treatment),
     xpd = TRUE)
# Axes with Tick Marks
axis(side = 1, labels = F, tck = -0.01, lwd = 2)
axis(side = 2, labels = T, tck = -0.02, lwd = 2, las = 1)
\#axis(side = 3, labels = F, tck = -0.02, lwd = 2)
axis(side = 4, labels = F, tck = -0.02, lwd = 2)
\#axis(side = 1, labels = F, tck = 0.01, lwd = 2)
axis(side = 2, labels = F, tck = 0.01, lwd = 2)
\#axis(side = 3, labels = F, tck = 0.01, lwd = 2)
axis(side = 4, labels = F, tck = 0.01, lwd = 2)
box(lwd = 2)
# Close Plot Device
dev.off()
## pdf
##
   2
graphics.off()
```

