## Symbiodiniaceae ITS2 community analyses

# $Ryan\ Eckert-ryan.j.eckert@gmail.com\\10/15/2019$

## ${\bf Contents}$

About this document	2
Basic setup of R environment and data	2
Loading required packages	2
Creating a color palette to use for our data	3
Loading our data into R	3
Preparing data for analyses	3
Purging outlying OTUs	4
Samples with data for OTUs	5
Normalizing reads across all samples	6
Renaming OTU sequence columns	7
OTU $\alpha$ -diversity	7
Statistical tests of $\alpha$ -diversity	8
Assumption testing	8
Kruskal-Wallis Tests	12
K-W test on Shannon's indices	12
K-W test on Simpson's indices	13
K-W test on species richness	13
Pairwise comparisons with Dunn's test	13
Plotting $\alpha$ -diversity metrics	14
Plot species richness by site	14
Create pairwise comparison letters	15
Plot species richness by depth	15
Make legends grobs	16
Plot Shannon's index by site	17
Plot Shannon's index by depth	17
Plot Simpson's index by site	18
Plot Simpson's index by depth	19
Create a single figure panel	19
Save as high-res images	19
	•
Ordination with PCoA	20
Create distance matrix	
Perform PCoA	
Determine percent variation captured on each axis	
Plot the PCoA	
Save the generated PCoA plot	22
Ordination with nMDS	23
Prepare data to plot	23
Construct nMDS biplot	24
Saving nMDS plot	25
Remove outlying samples	25

nMDS without outlying samples	26
Relative abundance of Symbiodiniaceae OTUs	28
Caluculate OTU relative abundances	28
Add plot order to data frame	
Create OTU stack for plotting	
Consruct OTU barplot	
Save the OTU barplot	
ITS2 community differences	32
Cheking dispersion	32
Running PERMANOVA in R	
Pairwise PERMANOVA for multiple comparisons	
PERMANOVA without 35 m samples	
Subset our dataframe	
Normalize reads	34
Run PERMANOVA on subset of data	
Pairwise PERMANOVA	
Generalized linear mixed model of OTUs	36
Stack OTU data table	36
Creating the model	37
Fitting the model	
Calculate effect size and p-values	
Plotting the GLMM	
Save GLMM plot	

#### About this document

This is the code accompanies the publication Eckert RJ, Reaume A, Sturm AB, Studivan MS, and Voss JD.(in review) Symbiodiniaceae community variation among Montastraea cavernosa populations along a depth gradient on the Belize Barrier Reef. Here you will find all the code to repeat the analyses performed for this manuscript. All of the accompanying data can be found on my github. If you download my entire accompanying github directory you should be able to re-run these analyses by following along with the included code in R Studio. If you download the code separtely or you are using this pipeline on your own data, you may need to change the working directory to whereever the associated files are housed (ie. setwd("~/path/to/directory/with/data")).

The data used for this analysis are OTUs calculated with R packages dada2 and LULU. The raw Symbiodiniaceae ITS2 sequences obtained from  $Montastraea\ cavernosa$  samples can be found in the NCBI SRA under project number XXX. Hopefully you are able to follow along this file and find it useful to use with your own data!

## Basic setup of R environment and data

#### Loading required packages

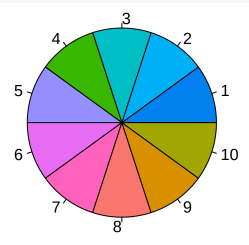
For the following analyses we will require the use of a number of different R packages. Most of these can be sourced from CRAN, but a couple need to be downloaded from GitHub or BioConducter. We can use the following code to quickly load in the packages and install any packages not previously installed in the R console.

```
if (!require("pacman")) install.packages("pacman") # for rapid install if not in library
#if(!require("pairwiseAdonis")) pacman::p_load_gh('pmartinezarbizu/pairwiseAdonis/
```

#### Creating a color palette to use for our data

We need 10 colors, I'm using mostly ggPlot colors, but in a better order for visualizing our OTU bar chart. I like to set a default palette at the beginning and call specific colors from it as needed throughout.

```
its2ColPal=c("#0081EF", "#00B0F6", "#00BFC4", "#39B600", "#9590FF", "#E76BF3", "#FF62BC", "#F8766D", "#D89000", "#A3A500")
```



#### Loading our data into R

First, we need to load in the data from dada2 and LULU analyses.

```
curated_result_97 = readRDS("curated_result_97.rds") #read in LULU output
ITS2data <- read.csv("ASVtable.csv") #read in dada2 output
head(ITS2data)</pre>
```

## Preparing data for analyses

Now that the data are loaded in, we can continue to make these data the way we need them for our analyses. We can combine our curated OTUs with the metadata we added to our ASV file.

```
ITS2data = cbind(ITS2data[, 1:3], data.frame(t(curated_result_97$curated_table)))
```

And we can order sites from north to south so later data can be plotted shallow to deep; north to south. To do this we can set "Depth" and "Site" as factors

```
ITS2data$Depth = factor(ITS2data$Depth, levels = c("10", "16", "25", "35"))
levels(ITS2data$Site)
```

```
## [1] "GR" "RW" "SR" "TR"

ITS2data$Site = factor(ITS2data$Site, levels(ITS2data$Site)[c(4, 2, 3, 1)])

ITS2data = ITS2data[order(ITS2data$Site, ITS2data$Depth), ]

ITS2data$newOrder = 1:nrow(ITS2data)
```

```
ITS2data = cbind(ITS2data[, length(ITS2data), drop = FALSE],
                   ITS2data[, c(1:length(ITS2data) - 1)])
row.names(ITS2data) = ITS2data$Sample
head(ITS2data)
##
       newOrder Sample Site Depth
                                         sq1 sq10 sq11 sq14 sq17 sq18 sq25 sq272
## 107
               1
                     107
                            TR
                                  10
                                        6456
                                                       0
                                                             0
                                                                  0
               2
                     108
                                   10 215412
                                                                  0
## 108
                            TR
                                                 0
                                                       0
                                                             0
                                                                        0
## 109
               3
                     109
                            TR
                                      16407
                                                       0
                                                             0
                                                                  0
                                                                        0
                                                                                    0
                                  10
                                                 0
## 110
               4
                     110
                            TR
                                  10 197896
                                                 0
                                                       0
                                                             6
                                                                  3
                                                                     618
                                                                             1
                                                                                    0
               5
                                                             0
                                                                  0
                                                                                    0
## 111
                     111
                            TR
                                   10 180119
                                                 0
                                                       0
                                                                        0
## 112
               6
                     112
                            TR
                                       24462
                                                             0
                                                                  0
                                                                        0
                                                                                    0
                                   10
                                                 0
                                                       0
       sq300 sq32 sq360 sq442 sq474 sq485
                                                 sq5 sq503 sq531 sq537 sq547 sq555
## 107
                  0
                        0
                               0
                                      0
                                             0
                                                 649
                                                          0
                                                                 0
                                                                        0
                                                                               0
            0
## 108
            0
                  0
                        0
                               0
                                      0
                                             0 17803
                                                          0
                                                                 0
                                                                        0
                                                                               0
                                                                                     0
## 109
                  0
                        0
                               0
                                      0
                                             0 1397
                                                                 0
                                                                        0
                                                                               0
                                                                                     0
            0
                                                          0
## 110
            0
                  2
                        0
                               0
                                      0
                                             0 19950
                                                          0
                                                                 0
                                                                        0
                                                                               0
                                                                                     0
## 111
                        0
                                                                               0
                                                                                     0
            0
                 0
                               6
                                      0
                                             0 16261
                                                          3
                                                                 0
                                                                        1
## 112
            0
                 0
                        0
                                      0
                                                2438
                                                          0
                                                                 0
                                                                        0
                                                                               0
                                                                                     0
       sq563 sq64
##
                     sq7 sq8
## 107
            0
                  0
                     225
## 108
            0
                  0 4736
                            0
```

Now we can call these data for the following analyses.

0 7923

0 8595

#### Purging outlying OTUs

## 109

## 110

## 111

## 112

```
goods = purgeOutliers(ITS2data, count.columns = 5:length(ITS2data), otu.cut = 0.001)
## [1] "samples with counts below z-score -2.5 :"
                    "212" "220" "67" "189" "186" "197"
## [1] "107" "44"
## [1] "zscores:"
##
         107
                              212
                                         220
## -3.289894 -2.977162 -2.594488 -2.840026 -2.895177 -4.272415 -3.211249
##
         197
## -4.032355
## [1] "OTUs passing frequency cutoff 0.001 : 10"
goods$newOrder = 1:nrow(goods)
row.names(goods) = goods$sample
head(goods)
##
       newOrder sample Site Depth
                                       sq1 sq10 sq11 sq14 sq17 sq18 sq25
## 108
                    108
                          TR
                                10 215412
                                                   0
                                                         0
                                                              0
                                                                   0
                                                                         0 17803
              1
                                              0
## 109
              2
                    109
                          TR
                                   16407
                                                   0
                                                         0
                                                              0
                                                                   0
                                                                           1397
## 110
              3
                    110
                          TR
                                10 197896
                                              0
                                                   0
                                                         6
                                                              3
                                                                 618
                                                                         1 19950
## 111
              4
                    111
                          TR
                                10 180119
                                              0
                                                   0
                                                         0
                                                              0
                                                                   0
                                                                         0 16261
## 112
              5
                    112
                          TR
                                    24462
                                                         0
                                                              0
                                                                   0
                                                                           2438
                                              0
                                                   0
## 113
              6
                    113
                          TR
                                10 195026
                                                        7
                                                             15
                                                                 826
                                                                       14 23224
        sq7 sq8
##
## 108 4736
```

```
## 109 266 0

## 110 7923 36

## 111 8595 0

## 112 862 0

## 113 6242 21

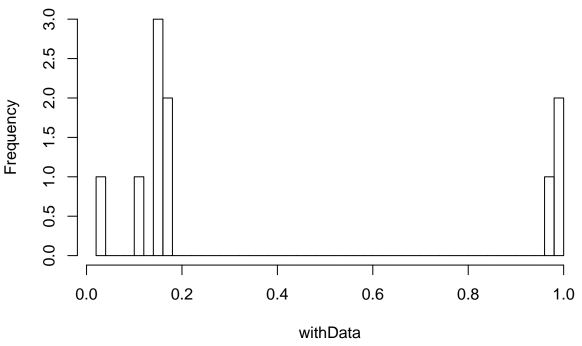
row.names(goods) = goods$sample
```

#### Samples with data for OTUs

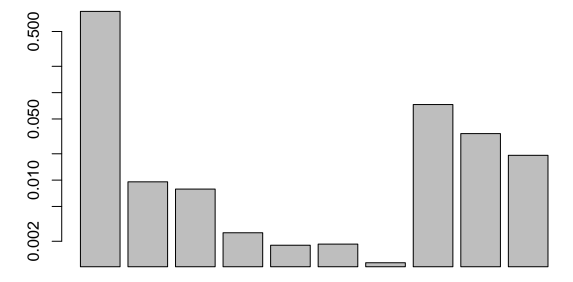
What is the proportion of samples with data for these OTUs?

```
withData = apply(goods[, 5:length(goods[1, ])], 2, function(x) {
    sum(x > 0)/length(x)
})
hist(withData, breaks = 50)
```

## Histogram of withData



```
props = apply(goods[, 5:length(goods[1, ])], 2, function(x) {
    sum(x)/sum(goods[, 5:length(goods[1, ])])
})
props
           sq1
                      sq10
                                   sq11
                                               sq14
                                                           sq17
## 0.848909686 0.009562495 0.007899518 0.002501895 0.001801993 0.001855771
          sq25
                       sq5
                                    sq7
                                                sq8
## 0.001136438 0.073101722 0.034023114 0.019207368
barplot(props, xaxt = "n", log = "y")
```



#### Normalizing reads across all samples

Here we normalize OTU counts with weighted trimmed mean of M-values (TMM; Robinson and Oshlack 2010; https://doi.org/10.1186/gb-2010-11-3-r25). This helps to account for disparity in sequencing depth across libraries. First we transpose the data to work with edgeR

```
itsGoodsTransposed = t(goods[, 5:length(goods[1, ])])
itsGoodsList = DGEList(counts = itsGoodsTransposed)
head(itsGoodsList$samples)
```

```
group lib.size norm.factors
## 108
            1
                 237951
## 109
            1
                  18070
                                     1
## 110
            1
                 226433
                                     1
## 111
            1
                 204975
                                     1
## 112
            1
                  27762
                                     1
## 113
            1
                 225375
                                     1
```

Now we can use TMM normalization in edgeR

```
its2Norm = calcNormFactors(itsGoodsList, method = "TMM")
head(its2Norm$samples)
```

```
##
       group lib.size norm.factors
## 108
                           1.090520
           1
                237951
## 109
           1
                 18070
                           1.095378
## 110
           1
                226433
                           1.057821
## 111
           1
                204975
                           1.063382
                           1.066040
## 112
           1
                 27762
                225375
                           1.048051
## 113
its2TMM = t(cpm(its2Norm, normalized.lib.sizes = TRUE))
its2Norm = cbind(goods[,c(2:4)], its2TMM)
head(its2Norm)
```

```
##
       sample Site Depth
                               sq1 sq10 sq11
                                                   sq14
                                                            sq17
                                                                      sq18
## 108
          108
                 TR
                       10 830134.8
                                       0
                                               0.00000
                                                         0.00000
                                                                     0.000
## 109
          109
                 TR
                       10 828909.5
                                       0
                                            0
                                               0.00000
                                                         0.00000
                                                                     0.000
## 110
          110
                 TR
                       10 826199.7
                                       0
                                            0 25.04951 12.52476 2580.100
                                              0.00000 0.00000
## 111
          111
                 TR
                       10 826360.2
                                       0
                                                                     0.000
```

```
## 112
          112
                TR
                       10 826546.9
                                           0 0.00000 0.00000
                                                                   0.000
## 113
          113
                       10 825665.6
                                           0 29.63533 63.50427 3496.969
                TR.
##
            sq25
                      sq5
                                          sq8
        0.000000 68607.55 18251.16
                                      0.00000
## 108
##
  109
        0.000000 70578.82 13438.77
                                      0.00000
        4.174918 83289.62 33077.88 150.29706
        0.000000 74603.14 39432.63
                                      0.00000
## 112 0.000000 82377.62 29126.13
                                      0.00000
## 113 59.270653 98321.55 26426.24
                                     88.90598
```

#### Renaming OTU sequence columns

Finally, I want to rename the columns with the Symbiodiniaceae sequences identified through our previose BLASTn query

```
colnames(its2Norm)[4:ncol(its2Norm)] = c("sq01_C3", "sq10_C3", "sq11_C3g", "sq14_C3g",
                                          "sq17_C3g", "sq18_C3g", "sq25_C3g", "sq05_C3z",
                                          "sq07_C3e", "sq08_C3g")
head(its2Norm)
##
       sample Site Depth sq01_C3 sq10_C3 sq11_C3g sq14_C3g sq17_C3g sq18_C3g
## 108
          108
                TR
                      10 830134.8
                                         0
                                                  0 0.00000 0.00000
                                                                          0.000
## 109
          109
                TR
                      10 828909.5
                                         0
                                                  0 0.00000 0.00000
                                                                          0.000
                                         0
                                                  0 25.04951 12.52476 2580.100
## 110
          110
                TR
                      10 826199.7
## 111
          111
                TR
                      10 826360.2
                                         0
                                                     0.00000
                                                              0.00000
                                                                          0.000
                                         0
## 112
          112
                TR
                      10 826546.9
                                                  0 0.00000 0.00000
                                                                          0.000
## 113
          113
                TR
                      10 825665.6
                                         0
                                                  0 29.63533 63.50427 3496.969
##
        sq25_C3g sq05_C3z sq07_C3e
                                     sq08_C3g
## 108
        0.000000 68607.55 18251.16
                                      0.00000
## 109
        0.000000 70578.82 13438.77
                                      0.00000
        4.174918 83289.62 33077.88 150.29706
## 110
## 111
        0.000000 74603.14 39432.63
                                      0.00000
## 112 0.000000 82377.62 29126.13
                                      0.00000
## 113 59.270653 98321.55 26426.24
                                     88.90598
levels(its2Norm$Site)
## [1] "TR" "RW" "SR" "GR"
levels(its2Norm$Depth)
```

```
## [1] "10" "16" "25" "35"
```

After all that everything looks good to go. On to the fun stuff!

## OTU $\alpha$ -diversity

Looking at  $\alpha$ -diversity of Symbiodiniaceae OTU sequences in each of our samples for comparison. We will look at species richness, Shannon's index and Simpson's index ##Calculate  $\alpha$ -diversity metrics

```
its2Richness = specnumber(its2Norm[,4:ncol(its2Norm)])
its2Shannon = diversity(its2Norm[,4:ncol(its2Norm)],index = "shannon")
its2Simpson = diversity(its2Norm[,4:ncol(its2Norm)],index = "simpson")
its2Div = cbind(its2Norm[,1:3], its2Richness, its2Shannon, its2Simpson)
colnames(its2Div)[4:6] = c("richness", "shannon", "simpson")
head(its2Div)
```

```
##
       sample Site Depth richness
                                     shannon
                                               simpson
## 108
          108
                TR
                      10
                                 3 0.3620252 0.1744764
                                 3 0.3476677 0.1693987
## 109
          109
                TR
                       10
## 110
                      10
          110
                TR
                                 8 0.4670561 0.2271793
## 111
          111
                TR
                      10
                                 3 0.4476261 0.2197705
## 112
          112
                TR
                      10
                                 3 0.4329306 0.2149295
## 113
          113
                      10
                                 8 0.4816535 0.2397877
```

#### Statistical tests of $\alpha$ -diversity

Now we can test whether or not there a significant differences in any of the calculated  $\alpha$ -diversity metrics across Site or Depth.

#### Assumption testing

First we need to see if our data are normally distributed. We can use Shapiro-Wilk's test.

```
tapply(its2Div$richness, its2Div$Site, shapiro.test)
```

```
## $TR
##
    Shapiro-Wilk normality test
##
##
## data: X[[i]]
## W = 0.49821, p-value = 8.713e-13
##
##
## $RW
##
##
   Shapiro-Wilk normality test
##
## data: X[[i]]
## W = 0.59852, p-value = 2.451e-11
##
##
## $SR
##
    Shapiro-Wilk normality test
##
##
## data: X[[i]]
## W = 0.51981, p-value = 1.024e-12
##
##
## $GR
##
##
   Shapiro-Wilk normality test
##
## data: X[[i]]
## W = 0.58742, p-value = 2.111e-11
tapply(its2Div$shannon, its2Div$Site, shapiro.test)
## $TR
##
##
    Shapiro-Wilk normality test
##
```

```
## data: X[[i]]
## W = 0.55244, p-value = 4.961e-12
##
##
## $RW
##
## Shapiro-Wilk normality test
##
## data: X[[i]]
## W = 0.61603, p-value = 4.648e-11
##
## $SR
##
## Shapiro-Wilk normality test
##
## data: X[[i]]
## W = 0.69458, p-value = 6.984e-10
##
##
## $GR
##
## Shapiro-Wilk normality test
## data: X[[i]]
## W = 0.51309, p-value = 1.8e-12
tapply(its2Div$simpson, its2Div$Site, shapiro.test)
## $TR
##
## Shapiro-Wilk normality test
##
## data: X[[i]]
## W = 0.56825, p-value = 8.468e-12
##
##
## $RW
##
## Shapiro-Wilk normality test
##
## data: X[[i]]
## W = 0.63812, p-value = 1.073e-10
##
##
## $SR
##
## Shapiro-Wilk normality test
## data: X[[i]]
## W = 0.73532, p-value = 4.536e-09
##
##
## $GR
##
```

```
## Shapiro-Wilk normality test
##
## data: X[[i]]
## W = 0.55991, p-value = 8.211e-12
tapply(its2Div$richness, its2Div$Depth, shapiro.test)
## $`10`
##
##
   Shapiro-Wilk normality test
##
## data: X[[i]]
## W = 0.7758, p-value = 3.534e-08
##
##
## $`16`
##
## Shapiro-Wilk normality test
## data: X[[i]]
## W = 0.67303, p-value = 4.331e-10
##
##
## $`25`
##
## Shapiro-Wilk normality test
##
## data: X[[i]]
## W = 0.2022, p-value = 3.019e-16
##
##
## $`35`
##
## Shapiro-Wilk normality test
##
## data: X[[i]]
## W = 0.17992, p-value = 4.725e-16
tapply(its2Div$shannon, its2Div$Depth, shapiro.test)
## $`10`
##
## Shapiro-Wilk normality test
## data: X[[i]]
## W = 0.67641, p-value = 3.197e-10
##
##
## $`16`
##
## Shapiro-Wilk normality test
##
## data: X[[i]]
## W = 0.68356, p-value = 6.73e-10
```

```
##
## $`25`
##
##
   Shapiro-Wilk normality test
##
## data: X[[i]]
## W = 0.71312, p-value = 1.977e-09
##
##
## $`35`
##
##
   Shapiro-Wilk normality test
## data: X[[i]]
## W = 0.77289, p-value = 6.555e-08
tapply(its2Div$simpson, its2Div$Depth, shapiro.test)
## $`10`
##
    Shapiro-Wilk normality test
##
##
## data: X[[i]]
## W = 0.70942, p-value = 1.353e-09
##
##
## $`16`
##
   Shapiro-Wilk normality test
##
##
## data: X[[i]]
## W = 0.70544, p-value = 1.734e-09
##
##
## $`25`
##
##
   Shapiro-Wilk normality test
##
## data: X[[i]]
## W = 0.74479, p-value = 8.794e-09
##
##
## $`35`
##
##
    Shapiro-Wilk normality test
##
## data: X[[i]]
## W = 0.83899, p-value = 2.827e-06
We see that our data are NOT normally distributed.
We can use Levene's test to test for equal variance among sites and depths.
leveneTest(its2Div$richness, its2Div$Depth)
## Levene's Test for Homogeneity of Variance (center = median)
```

```
Df F value
                       Pr(>F)
## group 3 25.117 4.368e-14 ***
##
        229
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
leveneTest(its2Div$shannon, its2Div$Depth)
## Levene's Test for Homogeneity of Variance (center = median)
        Df F value
                       Pr(>F)
         3 6.7836 0.0002115 ***
## group
##
        229
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
leveneTest(its2Div$simpson, its2Div$Depth)
## Levene's Test for Homogeneity of Variance (center = median)
        Df F value
                       Pr(>F)
## group 3 6.6744 0.0002442 ***
        229
##
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
leveneTest(its2Div$richness, its2Div$Site)
## Levene's Test for Homogeneity of Variance (center = median)
         Df F value Pr(>F)
## group 3 1.0165 0.3861
        229
leveneTest(its2Div$shannon, its2Div$Site)
## Levene's Test for Homogeneity of Variance (center = median)
##
         Df F value Pr(>F)
## group 3 0.4804 0.6962
        229
leveneTest(its2Div$simpson, its2Div$Site)
## Levene's Test for Homogeneity of Variance (center = median)
##
         Df F value Pr(>F)
## group
          3 0.5514 0.6477
        229
##
```

Data are non-normally distributed and also heteroschedastic. Using non-parametric tests moving forward.

#### Kruskal-Wallis Tests

Since our data did not fit the assumptions of ANOVA, we will use single factor Krskal-Wallis tests on these data.

#### K-W test on Shannon's indices

```
kruskal.test(shannon ~ Site, data = its2Div)
##
## Kruskal-Wallis rank sum test
```

```
##
## data: shannon by Site
## Kruskal-Wallis chi-squared = 2.7884, df = 3, p-value = 0.4254
kruskal.test(shannon ~ Depth, data = its2Div)
##
## Kruskal-Wallis rank sum test
##
## data: shannon by Depth
## Kruskal-Wallis chi-squared = 3.0743, df = 3, p-value = 0.3803
Nothing significant
K-W test on Simpson's indices
kruskal.test(simpson ~ Site, data = its2Div)
##
## Kruskal-Wallis rank sum test
##
## data: simpson by Site
## Kruskal-Wallis chi-squared = 3.4937, df = 3, p-value = 0.3216
kruskal.test(simpson ~ Depth, data = its2Div)
##
##
   Kruskal-Wallis rank sum test
##
## data: simpson by Depth
## Kruskal-Wallis chi-squared = 3.8446, df = 3, p-value = 0.2787
Again, no significant tests
K-W test on species richness
kruskal.test(richness ~ Site, data = its2Div)
   Kruskal-Wallis rank sum test
##
##
## data: richness by Site
## Kruskal-Wallis chi-squared = 2.7628, df = 3, p-value = 0.4297
kruskal.test(richness ~ Depth, data = its2Div)
   Kruskal-Wallis rank sum test
##
## data: richness by Depth
```

## Pairwise comparisons with Dunn's test

Here we see that species richness changes significantly with Depth

## Kruskal-Wallis chi-squared = 38.993, df = 3, p-value = 1.741e-08

We now use Dunn's test to do multiple comparisons with bonferroni correction. This will reveal where the differences in species richness are across Depth

```
dunn.test(its2Div$richness, g = its2Div$Depth, method = "bonferroni", table = FALSE,
          list = TRUE)
     Kruskal-Wallis rank sum test
##
##
## data: x and group
## Kruskal-Wallis chi-squared = 38.9935, df = 3, p-value = 0
##
##
##
                              Comparison of x by group
##
                                    (Bonferroni)
##
## List of pairwise comparisons: Z statistic (adjusted p-value)
## 10 - 16 : 1.449108 (0.4419)
## 10 - 25 : 4.989355 (0.0000)*
## 16 - 25 : 3.504135 (0.0014)*
## 10 - 35 : 5.057011 (0.0000)*
## 16 - 35 : 3.591117 (0.0010)*
```

This shows us that species richness is different between our deepest sites and our shallowest sites

#### Plotting $\alpha$ -diversity metrics

## 25 - 35 : 0.133160 (1.0000)

## Reject Ho if p <= alpha/2

##

## alpha = 0.05

We can take all of our  $\alpha$ -diversity metrics and plot them all into a single figure panel to visualize the results of the previous tests. Here we make plots for richness, Simpson's and Shannon's indices by site and depth zone.

#### Plot species richness by site

```
its2RichPlotS = ggplot(data = its2Div, aes(x = Site, y = richness)) +
                geom point(aes(fill = Depth), shape = 21, color = "gray20", size = 2,
                           position = position_jitterdodge()) +
                geom boxplot(alpha = 0, outlier.shape = NA, color = "gray30") +
                scale_fill_manual(values = its2ColPal[c(1, 4, 6, 2)],
                                  labels = c("10 m", "16 m", "25 m", "35 m")) +
                xlab("Reef Site") +
                vlab("Richness") +
                stat_compare_means(size = 5, geom = "label", label.x = 1, label.y = 10.7) +
                coord_cartesian(ylim = c(1, 11)) +
                scale_y_continuous(breaks = seq(2, 10, 2)) +
                guides(fill = guide_legend(ncol = 2)) +
                theme_bw()
its2RichPlotSite = its2RichPlotS +
                   theme(axis.title.x = element_blank(),
                         axis.text.x = element_blank(),
                         axis.title.y = element_text(color = "black", size = 14),
                         axis.text.y = element text(color = "black", size = 12),
                         legend.title = element_text(color = "black", size = 14),
                         legend.text = element_text(color = "black", size = 12),
```

```
panel.border = element_rect(size = 1.1, color = "black"),
    panel.background = element_rect(fill = "white"),
    plot.background = element_rect(fill = "white"),
    legend.position = "bottom",
    legend.background = element_rect(color = "black"),
    plot.margin = unit(c(0.5, 0.1, 0.1, 0.61), "cm")
)
its2RichPlotSite
```

#### Create pairwise comparison letters

We know from our Kruskal-Wallis tests that there was a significant difference in the species diversity across depth zones. We can plot letters to denote similarity between groups on our plot.

```
richDkmc = kruskalmc(its2Div$richness ~ its2Div$Depth) # multiple-comparison test
print(richDkmc)
```

```
## Multiple comparison test after Kruskal-Wallis
## p.value: 0.05
## Comparisons
##
           obs.dif critical.dif difference
## 10-16 14.040805
                       32.74642
                                     FALSE
## 10-25 48.134463
                       32.60500
                                      TRUE
## 10-35 49.441667
                       33.04241
                                      TRUE
## 16-25 34.093659
                       32.88254
                                      TRUE
## 16-35 35.400862
                       33.31630
                                      TRUE
## 25-35 1.307203
                       33.17732
                                     FALSE
richDkmcDiff = richDkmc$dif.com$difference # select logical vector
names(richDkmcDiff) = row.names(richDkmc$dif.com)# add comparison names
# create a list with "homogenous groups" coded by letter
richDsigLetters = multcompLetters(richDkmcDiff, compare="<", threshold=0.05,
                                  Letters=c(letters, LETTERS, "."), reversed = FALSE)
richLetters = as.data.frame(richDsigLetters$Letters)
richLetters$depth = row.names(richLetters)
colnames(richLetters)[1] = "id"
richLettersy = c(1, 1, 1, 1)
head(richLetters)
      id depth y
##
## 10 a
           10 1
## 16 a
            16 1
            25 1
## 25 b
            35 1
## 35 b
```

These letters can now be added to the corresponding plot

#### Plot species richness by depth

```
scale_fill_manual(values = its2ColPal[c(8,5,3,9)],
                                  labels = c("Tobacco Reef", "Raph's Wall",
                                            "South Reef", "Gover's Reef")) +
                stat_compare_means(size = 5, geom = "label", label.x = 1.07,
                                  label.y = 10.7) +
                geom_text(data = richLetters, aes(x = depth, y = y, label = id),
                          size = 6) +
                coord cartesian( ylim = c(1, 11)) +
                scale_y_continuous(breaks = seq(2, 10, 2))+
                xlab("Depth (m)") +
                ylab("Richness") +
                guides(fill=guide_legend(ncol=2))+
                theme_bw()
its2RichPlotDepth = its2RichPlotD +
                     theme(axis.title.x = element_blank(),
                           axis.text.x = element_blank(),
                           axis.title.y = element_blank(),
                           axis.text.y = element_blank(),
                           legend.title = element_text(color = "black", size = 14),
                           legend.text = element_text(color = "black", size = 12),
                           panel.border = element_rect(size = 1.1, color = "black"),
                           panel.background = element rect(fill = "white"),
                           plot.background = element_rect(fill = "white"),
                           legend.position = "bottom",
                           legend.background = element_rect(color = "black"),
                           plot.margin = unit(c(0.5, 0.5, 0.1, 0.85), "cm")
                           )
its2RichPlotDepth
```

#### Make legends grobs

Now we want to take the Site and Depth legends and make them graphical objects (grobs). This will allow us to place them separately in our figure grid, so each plot won't have its own legend.

```
get_legend <- function(its2RichPlotSite) {</pre>
  tmp <- ggplot gtable(ggplot build(its2RichPlotSite))</pre>
  leg <-
  which(sapply(tmp$grobs, function(x)
  x$name) == "guide-box")
  legend <- tmp$grobs[[leg]]</pre>
  return(legend)
legend.site = get_legend(its2RichPlotSite)
get_legend = function(its2RichPlotDepth) {
  tmp <- ggplot_gtable(ggplot_build(its2RichPlotDepth))</pre>
  leg <-
  which(sapply(tmp$grobs, function(x)
  x$name) == "guide-box")
  legend <- tmp$grobs[[leg]]</pre>
  return(legend)
}
```

```
legend.depth = get_legend(its2RichPlotDepth)

# re-plot without legends
its2RichPlotSite = its2RichPlotSite + theme(legend.position = "none")
its2RichPlotDepth = its2RichPlotDepth + theme(legend.position = "none")
```

#### Plot Shannon's index by site

```
its2ShannonPlotS = ggplot(data = its2Div, aes(x = Site, y = shannon)) +
                   geom_point(aes(fill = Depth), shape = 21, color = "gray20",
                              size = 2, position = position_jitterdodge()) +
                   geom_boxplot(alpha = 0, outlier.shape = NA, color = "gray30") +
                   scale_fill_manual(values = its2ColPal[c(1, 4, 6, 2)],
                                     labels = c("10 m", "16 m", "25 m", "35 m")) +
                   stat_compare_means(size = 5, geom = "label", label.x = 1,
                                      label.y = 1.7) +
                   expand_limits(y = c(0, 1.75)) +
                   xlab("Reef Site") +
                   ylab("Shannon's index") +
                   theme_bw()
its2ShannonPlotSite = its2ShannonPlotS +
                      theme(axis.title.x = element_blank(),
                            axis.text.x = element_blank(),
                            axis.title.y = element_text(color = "black", size = 14),
                            axis.text.y = element text(color = "black", size = 12),
                            legend.title = element text(color = "black", size = 14),
                            legend.text = element_text(color = "black", size = 12),
                            panel.border = element_rect(size = 1.1, color = "black"),
                            panel.background = element_rect(fill = "white"),
                            plot.background = element rect(fill = "white"),
                            legend.position = "none",
                            legend.background = element_rect(color = "black"),
                            plot.margin = unit(c(0.1, 0.1, 0.1, 0.5), "cm")
its2ShannonPlotSite
```

#### Plot Shannon's index by depth

#### Plot Simpson's index by site

```
its2SimpsonPlotS = ggplot(data = its2Div, aes(x = Site, y = simpson)) +
                   geom_point(aes(fill = Depth), shape = 21, color = "gray20", size = 2,
                              position = position_jitterdodge()) +
                   geom_boxplot(alpha = 0, outlier.shape = NA) +
                   scale_fill_manual(values = its2ColPal[c(1,4,6,2)],
                                     labels = c("10 m", "16 m", "25 m", "35 m")) +
                   stat_compare_means(size = 5, geom = "label", label.x = 1,
                                      label.y = 0.77) +
                   expand_limits(y = c(0, 0.8)) +
                   xlab("Reef Site") +
                   ylab("Simpson's index") +
                   theme_bw()
its2SimpsonPlotSite = its2SimpsonPlotS +
                      theme(axis.title.x = element_text(color = "black", size = 14),
                            axis.text.x = element text(color = "black", size = 12),
                            axis.title.y = element text(color = "black", size = 14),
                            axis.text.y = element_text(color = "black", size = 12),
                            legend.title = element_text(color = "black", size = 14),
                            legend.text = element_text(color = "black", size = 12),
                            panel.border = element_rect(size = 1.1, color = "black"),
                            panel.background = element_rect(fill = "white"),
                            plot.background = element_rect(fill = "white"),
                            legend.position = "none",
                            legend.background = element_rect(color = "black"),
                            plot.margin = unit(c(0.11, 0.1, 0, 0.5), "cm")
its2SimpsonPlotSite
```

#### Plot Simpson's index by depth

```
its2SimpsonPlotD = ggplot(data = its2Div, aes(x = Depth, y = simpson)) +
                   geom point(aes(fill = Site), shape = 21, color = "gray20", size = 2,
                              position = position_jitterdodge()) +
                   geom_boxplot(alpha = 0, outlier.shape = NA) +
                   scale_fill_manual(values = its2ColPal[c(8,5,3,9)],
                                     labels = c("Tobacco Reef", "Raph's Wall",
                                     "South Reef", "Gover's Reef")) +
                   stat_compare_means(size = 5, geom = "label", label.x = 1,
                                      label.y = 0.77) +
                   expand_limits(y = c(0, 0.8)) +
                   xlab("Depth (m)") +
                   ylab("Simpson's index") +
                   theme_bw()
its2SimpsonPlotDepth = its2SimpsonPlotD +
                       theme(axis.title.x = element_text(color = "black", size = 14),
                             axis.text.x = element_text(color = "black", size = 12),
                             axis.title.y = element_blank(),
                             axis.text.y = element_blank(),
                             legend.title = element_text(color = "black", size = 14),
                             legend.text = element text(color = "black", size = 12),
                             panel.border = element_rect(size = 1.1, color = "black"),
                             panel.background = element_rect(fill = "white"),
                             plot.background = element_rect(fill = "white"),
                             legend.position = "none",
                             legend.background = element_rect(color = "black"),
                             plot.margin = unit(c(0.1, 0.5, 0, 0.85), "cm")
                             )
its2SimpsonPlotDepth
```

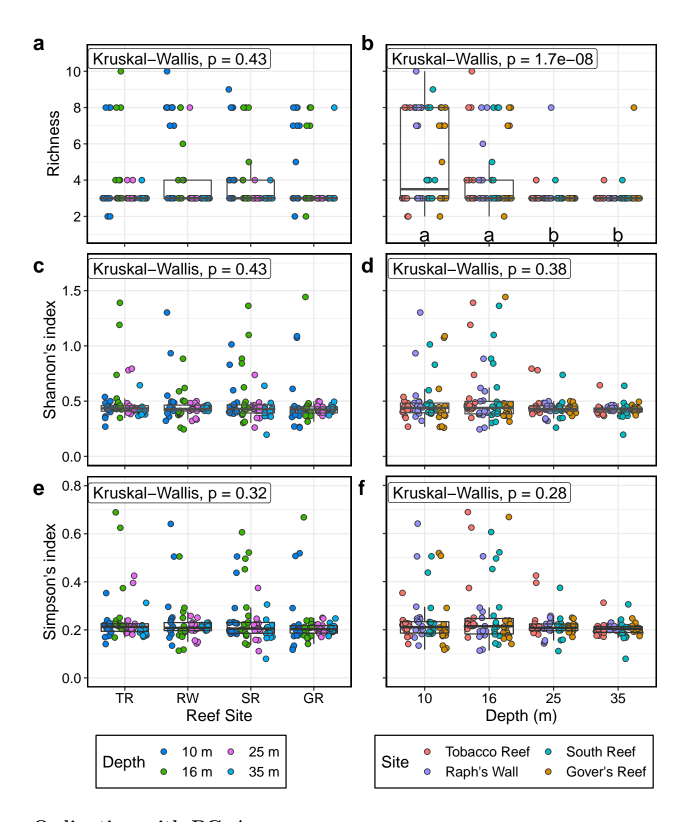
#### Create a single figure panel

Now we want to throw all those bad boys into a single figure grid.

#### Save as high-res images

Finally, we will save the resulting  $\alpha$ -diversity plot panel as a high resoultion .eps figure. Alternatively you could save it as a HQ .tiff by simply changing the extension in the code below.

```
ggsave("its2_diversityPlots.eps", plot = divPlots, width = 8.25, height = 10,
    unit = "in", dpi = 600)
```



### Ordination with PCoA

We can use principal coordinates analysis (PCoA) to visualize our samples based on significant OTUs. But, first we need to set up data for PCoA analysis in R.

#### Create distance matrix

```
We will create a distance matrix with Bray-Curtis similarity using the package vegan
```

```
its2Dist = vegdist(its2Norm[, c(4:ncol(its2Norm))], method = "bray")
```

#### Perform PCoA

```
This is the actual PCoA step
```

```
its2Mds = cmdscale(its2Dist, eig = TRUE, x.ret = TRUE)
```

#### Determine percent variation captured on each axis

Calculate the eigenvalues so later we can figure out % variation shown on each Principal Coordinate

```
its2Var = round(its2Mds$eig/sum(its2Mds$eig)*100, 1)
its2Var
```

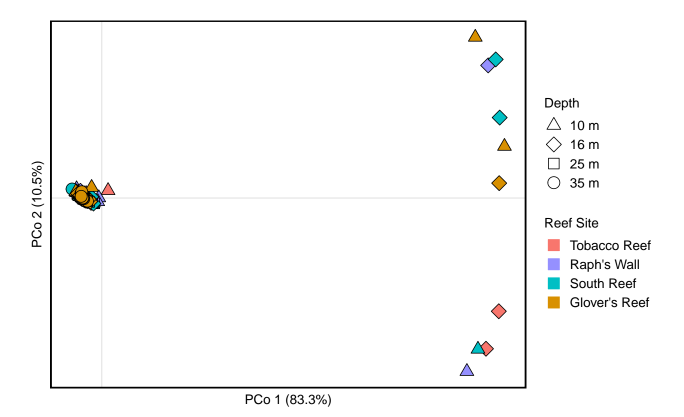
```
[1] 83.3 10.5
                    3.0
                                              0.3
                                                   0.2
##
                         1.8
                              1.3
                                   0.9
                                        0.4
                                                        0.2
                                                             0.1
                                                                   0.1
                                                                        0.1
                                                   0.0
##
    [15]
          0.0
               0.0
                    0.0
                         0.0
                              0.0
                                    0.0
                                        0.0
                                              0.0
                                                        0.0
                                                             0.0
                                                                   0.0
                                                                        0.0
##
    [29]
          0.0
               0.0
                    0.0
                         0.0
                              0.0
                                    0.0
                                        0.0
                                              0.0
                                                   0.0
                                                        0.0
                                                             0.0
                                                                  0.0
                                                                       0.0
                                                                             0.0
   [43]
          0.0
               0.0
                    0.0
                         0.0
                              0.0
                                   0.0
                                        0.0
                                              0.0
                                                   0.0
                                                        0.0
                                                             0.0
                                                                  0.0
                                                                        0.0
   [57]
               0.0
                    0.0
                         0.0
                              0.0
                                        0.0
                                              0.0
                                                   0.0
                                                        0.0
                                                             0.0
##
          0.0
                                   0.0
                                                                  0.0
                                                                        0.0
                                                                             0.0
##
    [71]
         0.0
               0.0
                    0.0
                         0.0
                              0.0
                                   0.0
                                        0.0
                                              0.0
                                                   0.0
                                                        0.0
                                                             0.0
                                                                  0.0
                                                                        0.0
                                                                             0.0
##
   [85]
         0.0
               0.0
                    0.0
                         0.0
                              0.0
                                   0.0
                                        0.0
                                              0.0
                                                   0.0
                                                        0.0
                                                             0.0
                                                                  0.0
                                                                        0.0
                                                                             0.0
   [99]
          0.0
               0.0
                    0.0
                         0.0
                              0.0
                                   0.0
                                        0.0
                                              0.0
                                                   0.0
                                                        0.0
                                                             0.0
                                                                  0.0
                                                                        0.0
                                                                             0.0
##
## [113]
          0.0
               0.0
                    0.0
                         0.0
                              0.0
                                   0.0
                                        0.0
                                              0.0
                                                   0.0
                                                        0.0
                                                             0.0
                                                                  0.0
                                                                        0.0
                                                                             0.0
## [127]
         0.0
               0.0
                    0.0
                         0.0
                              0.0
                                   0.0
                                        0.0
                                              0.0
                                                   0.0
                                                        0.0
                                                                  0.0
                                                                        0.0
                                                                             0.0
                                                             0.0
## [141]
         0.0
               0.0
                   0.0
                         0.0
                              0.0
                                   0.0
                                        0.0
                                              0.0
                                                   0.0
                                                        0.0
                                                             0.0
                                                                  0.0
                                                                        0.0
                                                                             0.0
                         0.0
## [155]
                              0.0
                                                   0.0
                                                        0.0
         0.0
               0.0
                    0.0
                                   0.0
                                        0.0
                                              0.0
                                                             0.0
                                                                  0.0
                                                                        0.0
                                                                             0.0
## [169]
         0.0
               0.0
                    0.0
                         0.0
                              0.0
                                   0.0
                                        0.0
                                              0.0
                                                   0.0
                                                        0.0
                                                             0.0
                                                                  0.0
## [183]
         0.0
               0.0 0.0
                         0.0
                              0.0
                                   0.0
                                        0.0
                                              0.0
                                                   0.0
                                                        0.0
                                                             0.0
                                                                  0.0
                                                                        0.0
                                                                             0.0
## [197]
               0.0
                   0.0
                         0.0
                              0.0
                                        0.0
                                              0.0
                                                   0.0
                                                        0.0
                                                                  0.0
         0.0
                                   0.0
                                                             0.0
                                                                        0.0
## [211]
               0.0 0.0 0.0 0.0 0.0 0.0 0.0 0.0
                                                        0.0 0.0 0.0 0.0 0.0
         0.0
## [225] 0.0 0.0 -0.1 -0.1 -0.1 -0.2 -0.3 -0.4 -0.6
```

```
##
     sample site depth
                               PCo1
                                            PCo2
## 1
        108
              TR
                    10 -0.04147518 0.004611102
## 2
        109
                    10 -0.04211077 0.005071409
              TR.
## 3
        110
              TR
                    10 -0.03039080 -0.004883738
## 4
        111
              TR
                    10 -0.03416250 -0.004766676
## 5
        112
              TR
                    10 -0.03216465 -0.003956257
## 6
        113
              TR.
                    10 -0.02452031 -0.006515757
```

#### Plot the PCoA

```
# plotting PCoA with ggPlot
its2PcoaA = ggplot(its2Pcoa, aes(x = PCo1, y = PCo2)) +
            geom_hline(yintercept = 0, color = "gray90", size = 0.5) +
            geom_vline(xintercept = 0, color = "gray90", size = 0.5) +
            geom_point(aes(shape = factor(depth), size = depth, fill = site),
                       color = "black") +
            scale_fill_manual(values = its2ColPal[c(8,5,3,9)], name = "Reef Site",
                                labels = c("Tobacco Reef", "Raph's Wall", "South Reef",
                                           "Glover's Reef")) +
            scale_shape_manual(values = c(24,23,22,21), name = "Depth",
                               labels = c("10 m", "16 m", "25 m", "35 m")) +
            scale_size_manual(values = c(4, 4.75, 4.75, 4.75)) +
            guides(shape = guide legend(override.aes = list(size = c(4, 4.75, 4.75, 4.75),
                   fill = "white")), fill = guide legend(override.aes =
                   list(shape = 22, size = 5.75, color = "white")), size = FALSE,
                   color = FALSE)+
             xlab(paste ("PCo 1 (", its2Var[1],"%)", sep = "")) +
             ylab(paste ("PCo 2 (", its2Var[2],"%)", sep = "")) +
             theme_bw()
its2Pcoa = its2PcoaA +
           theme(axis.title.x = element_text(color = "black", size = 12),
                 axis.text.x = element_blank(),
                 axis.ticks.x = element_blank(),
                 axis.line.x = element_blank(),
                 axis.title.y = element_text(color = "black", size = 12),
                 axis.text.y = element_blank(),
                 axis.ticks.y = element_blank(),
                 axis.line.y = element_blank(),
                 legend.position = "right",
                 legend.title = element text(color = "black", size = 12),
                 legend.text = element_text(color = "black", size = 12),
                 legend.key = element blank(),
                 panel.border = element_rect(color = "black", size = 1.2),
                 panel.background = element_rect(fill = "white"),
                 plot.background = element_rect(fill = "white"),
                 panel.grid.major = element blank(),
                 panel.grid.minor = element_blank()
its2PcoaA
```

#### Save the generated PCoA plot



#### Ordination with nMDS

Now we can view the data with nMDS using the package *vegan* again. First we have to run the calculations for the nMDS. This is similar to PCoA, but can be scaled freely and uses an iterative process. We'll run it with 50 iterations, which should be enough to arrive at a solution.

```
its2Nmds = metaMDS(its2Norm[4:ncol(its2Norm)], try = 50)
its2Nmds
```

#### Prepare data to plot

```
its2Scores = as.data.frame(scores(its2Nmds))
its2Scores$site = factor(its2Norm$Site)
its2Scores$depth = as.factor(its2Norm$Depth)
its2Scores$sample = row.names(its2Scores)
head(its2Scores)
##
              NMDS1
                           NMDS2 site depth sample
## 108 -0.052295737 -0.058459256
                                   TR
                                          10
## 109 -0.048164717 -0.091740894
                                   TR
                                          10
                                                109
## 110 -0.022926842 0.009377931
                                          10
                                                110
## 111 -0.069887493 0.011330805
                                   TR
                                          10
                                                111
## 112 -0.056906218 -0.016534503
                                   TR
                                          10
                                                112
## 113 0.002797965 -0.011414792
                                   TR
                                          10
                                                113
its2Clades = as.data.frame(scores(its2Nmds, "species"))
its2Clades$seq = row.names(its2Clades)
its2Clades
```

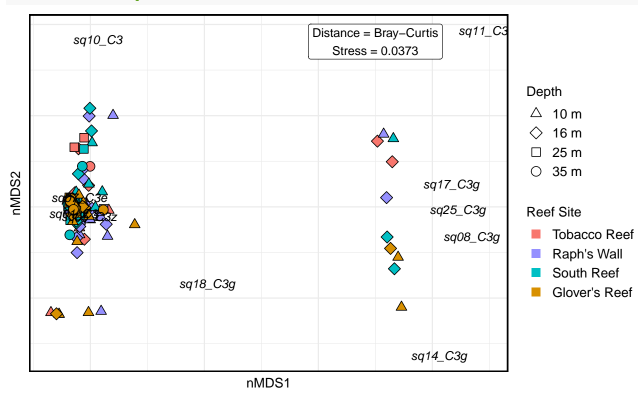
```
##
                  NMDS1
                              NMDS2
## sq01_C3 -0.074715881 -0.03014023 sq01_C3
## sq10 C3 0.031219752 0.73387836 sq10 C3
## sq11_C3g 1.736981629 0.77272569 sq11_C3g
## sq14_C3g 1.528327383 -0.65265579 sq14_C3g
## sq17 C3g 1.581939252 0.10004674 sq17 C3g
## sq18 C3g 0.511145818 -0.33556718 sq18 C3g
## sq25_C3g 1.610315632 -0.01124022 sq25_C3g
## sq05_C3z -0.006323011 -0.04149856 sq05_C3z
## sq07_C3e -0.049387470 0.03985393 sq07_C3e
## sq08_C3g 1.670623995 -0.13028495 sq08_C3g
its2Clades$seq
## [1] "sq01_C3" "sq10_C3" "sq11_C3g" "sq14_C3g" "sq17_C3g" "sq18_C3g"
## [7] "sq25_C3g" "sq05_C3z" "sq07_C3e" "sq08_C3g"
```

#### Construct nMDS biplot

```
its2NmdsPlotA = ggplot() +
                geom_point(data = its2Scores, aes(x = NMDS1, y = NMDS2,
                           shape = depth, size = depth, fill = site),
                           color = "black") + # add the site points
                scale_fill_manual(values = its2ColPal[c(8,5,3,9)],
                                  name = "Reef Site", labels = c("Tobacco Reef",
                                  "Raph's Wall", "South Reef", "Glover's Reef")) +
                scale shape manual (values = c(24, 23, 22, 21), name = "Depth",
                                   labels = c("10 m", "16 m", "25 m", "35 m")) +
                scale_size_manual(values = c(3, 3.75, 3.75, 3.75)) +
                guides(shape = guide legend( override.aes = list(size =
                       c(3, 3.75, 3.75, 3.75), fill = "white")), fill =
                         guide_legend(override.aes = list(shape = 22, size = 3.75,
                         color = NA)), size = FALSE)+
                geom_text(data = its2Clades, aes(x = NMDS1, y = NMDS2,
                                                 label = seq),
                            color = "#000000", size = 4, fontface = "italic") +
                          # add seq labels
                annotate("label", x = 1.25, y = 0.725, label = paste
                            ("Distance = Bray-Curtis\nStress = ",
                            round(its2Nmds$stress, 4), sep = ""), size = 4) +
                labs(x = "nMDS1", y = "nMDS2") +
                coord equal() +
                theme bw()
its2NmdsPlot = its2NmdsPlotA +
               theme(axis.title.x = element_text(color = "black", size = 12),
                     axis.text.x = element_blank(),
                     axis.ticks.x = element_blank(),
                     axis.title.y = element_text(color = "black", size = 12),
                     axis.text.y = element_blank(),
                     axis.ticks.y = element_blank(),
                     legend.position = "right",
                     legend.title = element_text(color = "black", size = 12),
                     legend.text = element_text(color = "black", size = 12),
```

```
legend.key = element_blank(),
legend.background = element_blank(),
panel.border = element_rect(color = "black", size = 1.2),
panel.background = element_rect(fill = "white"),
plot.background = element_blank()
)
its2NmdsPlot
```

#### Saving nMDS plot



#### Remove outlying samples

Samples with the C3g types skewing the data: 42, 80, 104, 98, 20, 57, 149, 152, 58, 163 let's look at these samples specifically

## sample Site Depth sq01\_C3 sq10\_C3 sq11\_C3g sq14\_C3g sq17\_C3g

```
## 98
           98
                TR
                       16
                            5962.282 2442.380
                                                1152540
                                                               0.0 107105.56
## 104
          104
                TR
                       16
                           14559.120 3065.078
                                                1047363
                                                              0.0 306539.72
## 42
           42
                RW
                       10
                            3327.416 3104.598
                                                1241319
                                                               0.0 200848.17
                           34844.366
                                                               0.0 219056.79
##
  20
           20
                RW
                       16
                                        0.000
                                                      0
##
  80
           80
                SR
                       10
                            7562.987 9943.468
                                                2905229
                                                               0.0 187462.89
## 57
           57
                SR
                       16
                           59249.888
                                        0.000
                                                         438911.4 361382.30
                                                      0
## 58
           58
                SR
                       16
                           69160.226
                                        0.000
                                                      0
                                                         704896.7 155133.96
## 152
          152
                GR
                       10
                            6045.766
                                        0.000
                                                      0
                                                         560943.6 79987.07
## 163
          163
                GR
                       10 171477.140
                                        0.000
                                                      0 1164666.6 101868.88
## 149
          149
                GR
                       16
                          14587.886
                                        0.000
                                                      0
                                                         499950.3 168854.14
##
        sq18_C3g
                  sq25_C3g sq05_C3z sq07_C3e
                                                sq08_C3g
                                                            sums
## 98
            0.00
                  61370.79 169218.7 13145.75
                                                884093.8 2395879
##
  104
            0.00 143260.47 165801.6 61908.19 1215750.4 2958247
## 42
            0.00 129932.62 130690.2 38369.26
                                                596691.8 2344283
## 20
            0.00 122446.05 383743.7 22680.39 5925715.7 6708487
## 80
            0.00 107791.16 164873.1 31417.39
                                                947977.0 4362257
## 57
       184315.48 189620.65 346779.9 47641.53 2467484.7 4095386
       312760.62 116130.53 474786.2 20674.75 4183582.7 6037126
## 152 62068.54 53954.48 187677.3 15253.63 1879099.6 2845030
## 163 158405.47 281910.22 621065.3
                                         0.00 5294413.0 7793807
## 149 169574.53
                  92441.40 243388.7 29973.35 1338843.8 2557614
max(itsOuts$sum)
## [1] 7793807
max(itsNoOut$sum)
## [1] 1372236
mean(itsNoOut$sum)
## [1] 942949
summary(itsOuts$sum)
##
      Min. 1st Qu. Median
                               Mean 3rd Qu.
                                                Max.
## 2344283 2629468 3526817 4209812 5618408 7793807
summary(itsNoOut$sum)
##
      Min. 1st Qu.
                               Mean 3rd Qu.
                    Median
                                                Max.
    871864 923180
                    932594
                             942949
                                     943096 1372236
```

Nothing unusual here, neither the highest nor the lowest sequencing depth of all samples

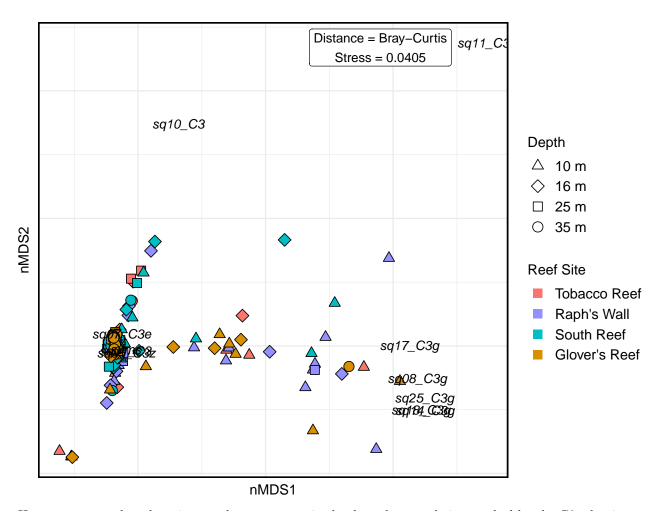
#### nMDS without outlying samples

Same idea as above, just with the new data lacking 10 C3g dominated samples. This way we can see if these samples are skewing our plot and obscuring a larger pattern in rest of the data.

```
its2Nmds2 = metaMDS(itsNoOut[4:ncol(itsNoOut)], try = 50)
its2Nmds2

its2Scores2 = as.data.frame(scores(its2Nmds2))
its2Scores2$site = factor(itsNoOut$Site)
its2Scores2$depth = as.factor(itsNoOut$Depth)
its2Scores2$sample = row.names(its2Scores2)
head(its2Scores2)
```

```
its2Clades2 = as.data.frame(scores(its2Nmds2, "species"))
its2Clades2$seq = row.names(its2Clades2)
its2Clades2
its2Clades2$seq
## Plot the data
its2NmdsPlotB = ggplot() +
                geom_point(data = its2Scores2, aes(x = NMDS1, y = NMDS2,
                           shape = depth, size = depth, fill = site),
                           color = "black") + # add the site points
                scale_fill_manual(values = its2ColPal[c(8,5,3,9)], name =
                                  "Reef Site", labels = c("Tobacco Reef",
                                  "Raph's Wall", "South Reef", "Glover's Reef")) +
                scale_shape_manual(values = c(24, 23, 22, 21), name = "Depth",
                                   labels = c("10 m", "16 m", "25 m", "35 m")) +
                scale_size_manual(values = c(3, 3.75, 3.75, 3.75)) +
                guides(shape = guide_legend( override.aes = list(size =
                       c(3, 3.75, 3.75, 3.75), fill = "white")), fill =
                       guide_legend(override.aes = list(shape = 22, size = 3.75,
                       color = NA)), size = FALSE)+
                geom text(data = its2Clades2, aes(x = NMDS1, y = NMDS2,
                          label = seq), color = "#000000", size = 4,
                          fontface = "italic") +
                          # add seq labels
                annotate("label", x = 0.95, y = 1.17, label = paste
                         ("Distance = Bray-Curtis\nStress = ",
                         round(its2Nmds2$stress, 4), sep = ""), size = 4) +
                labs(x = "nMDS1", y = "nMDS2") +
                coord_equal() +
                theme_bw()
its2NmdsPlot2 = its2NmdsPlotB +
                theme(axis.title.x = element_text(color = "black", size = 12),
                      axis.text.x = element_blank(),
                      axis.ticks.x = element_blank(),
                      axis.title.y = element_text(color = "black", size = 12),
                      axis.text.y = element_blank(),
                      axis.ticks.y = element blank(),
                      legend.position = "right",
                      legend.title = element text(color = "black", size = 12),
                      legend.text = element_text(color = "black", size = 12),
                      legend.key = element_blank(),
                      legend.background = element_blank(),
                      panel.border = element_rect(color = "black", size = 1.2),
                      panel.background = element_rect(fill = "white"),
                      plot.background = element_blank()
its2NmdsPlot2
# saving new nMDS plot
ggsave("nMDS_ASVs_noOut.eps", plot = its2NmdsPlot2, width = 9.2, height = 5.75,
      dpi = 600, device = "eps")
```



Here we can see that there is not a larger pattern in the data that was being masked by the C3g dominant samples

## Relative abundance of Symbiodiniaceae OTUs

Now we can calculate the relative abundance of each unique OTU per sample. This will allow us to view the community structure in a faceted barplot.

#### Caluculate OTU relative abundances

#### head(its2Norm) sample Site Depth sq01\_C3 sq10\_C3 sq11\_C3g sq14\_C3g sq17\_C3g sq18\_C3g ## ## 108 108 TR 10 830134.8 0 0.00000 0.00000 0.000 ## 109 109 TR 10 828909.5 0 0.00000 0.00000 0.000 TR 0 0 25.04951 12.52476 2580.100 ## 110 110 10 826199.7 ## 111 111 TR 10 826360.2 0 0.00000 0.00000 0.000 0.00000 0.00000 ## 112 112 TR 10 826546.9 0 0.000 ## 113 113 TR 10 825665.6 0 0 29.63533 63.50427 3496.969 sq25\_C3g sq05\_C3z sq07\_C3e ## sq08\_C3g ## 108 0.000000 68607.55 18251.16 0.00000 0.000000 70578.82 13438.77 0.00000 ## 109

```
4.174918 83289.62 33077.88 150.29706
## 111
       0.000000 74603.14 39432.63
                                         0.00000
## 112 0.000000 82377.62 29126.13
                                         0.00000
## 113 59.270653 98321.55 26426.24
                                        88.90598
its2NormPerc = its2Norm
its2NormPerc$sum = apply(its2NormPerc[, c(4:length(its2NormPerc[1,]))], 1, function(x) {
  sum(x, na.rm = T)
})
its2NormPerc = cbind(its2NormPerc[, c(1:3)], (its2NormPerc[,
                       c(4:(ncol(its2NormPerc)-1))] / its2NormPerc$sum))
head(its2NormPerc)
##
       sample Site Depth
                              sq01_C3 sq10_C3 sq11_C3g
                                                              sq14_C3g
                                                                             sq17_C3g
## 108
           108
                 TR
                        10 0.9052788
                                             0
                                                       0 0.000000e+00 0.000000e+00
## 109
           109
                 TR
                        10 0.9079690
                                             0
                                                       0 0.000000e+00 0.000000e+00
                 TR
                                             0
                                                       0 2.649790e-05 1.324895e-05
## 110
           110
                        10 0.8739716
## 111
           111
                 TR.
                        10 0.8787364
                                             0
                                                       0 0.000000e+00 0.000000e+00
## 112
           112
                 TR
                        10 0.8811325
                                             0
                                                       0 0.000000e+00 0.000000e+00
##
   113
           113
                 TR
                        10 0.8653400
                                             0
                                                       0 3.105935e-05 6.655574e-05
##
           sq18_C3g
                         sq25_C3g
                                      sq05_C3z
                                                  sq07_C3e
                                                                 sq08 C3g
## 108 0.000000000 0.000000e+00 0.07481792 0.01990326 0.000000e+00
## 109 0.000000000 0.000000e+00 0.07731046 0.01472053 0.000000e+00
## 110 0.002729284 4.416317e-06 0.08810553 0.03499048 1.589874e-04
   111 0.000000000 0.000000e+00 0.07933163 0.04193194 0.000000e+00
## 112 0.000000000 0.000000e+00 0.08781788 0.03104964 0.000000e+00
## 113 0.003665003 6.211869e-05 0.10304603 0.02769606 9.317804e-05
Now a quick sanity check. If this worked the sum of each row should = 100\% (i.e. "1")
# test that all are now 100% = 1
apply(its2NormPerc[, c(4:(ncol(its2NormPerc)))], 1, function(x) {
  sum(x, na.rm = T)
})
                                                                            95
   108 109 110 111 112 113 114 116 117 118 119 120 121 156
                                                                                96
##
                                                                   93
                                                                       94
                            1
                                1
                                                           1
                                                                    1
                                                                        1
                                                                                 1
##
        98
             99 100 101 102 103 104 105 106 137 122 123 124
                                                                 125 126 127
                                                                               128
     1
          1
              1
                       1
                            1
                                1
                                     1
                                         1
                                             1
                                                  1
                                                       1
                                                           1
                                                                1
                                                                    1
                                                                        1
   129 130
           131 132 133 134 135
                                  136 226
                                           227
                                                228
                                                    229
                                                        230 231
                                                                  232 233
##
                                                                          234
                                                                               236
##
     1
          1
              1
                   1
                       1
                           1
                                1
                                     1
                                         1
                                             1
                                                  1
                                                      1
                                                           1
                                                               1
                                                                    1
                                                                        1
                                                                             1
                                                                                 1
##
   237 238
            239
                240
                      29
                           30
                               31
                                   32
                                        33
                                            34
                                                 35
                                                     36
                                                          37
                                                              38
                                                                   39
                                                                       40
                                                                            41
                                                                                42
##
     1
          1
              1
                   1
                           1
                                1
                                     1
                                         1
                                             1
                                                  1
                                                      1
                                                           1
                                                               1
                                                                    1
                                                                        1
                                                                             1
                                                                                 1
                       1
##
    43
        17
             18
                 19
                      20
                          21
                               22
                                   23
                                        24
                                            25
                                                 26
                                                     27
                                                          28
                                                              45
                                                                   46
                                                                                 3
##
          1
              1
                   1
                           1
                                1
                                         1
                                             1
                                                  1
                                                           1
                                                               1
                                                                    1
                                                                        1
                                                                                 1
     1
                       1
                                    1
                                                      1
                   7
##
     4
          5
              6
                       8
                           9
                               10
                                   11
                                        12
                                            13
                                                 14
                                                     15
                                                          16 211
                                                                  213 214
                                                                          215
                                                                               216
##
     1
          1
              1
                   1
                       1
                           1
                                1
                                     1
                                         1
                                             1
                                                  1
                                                      1
                                                           1
                                                               1
                                                                    1
                                                                        1
                                                                             1
                                                                                 1
##
  217 218 219 221 222 223 224
                                  225
                                        76
                                            77
                                                 78
                                                     79
                                                          80
                                                              81
                                                                   82
                                                                       83
                                                                            84
                                                                                85
##
                           1
                                1
                                         1
                                                  1
                                                           1
                                                               1
                                                                    1
                                                                             1
                                                                                 1
     1
          1
              1
                   1
                       1
                                     1
                                             1
                                                      1
                                                                        1
             88
                 89
                      90
                          91
                                   48
                                        49
                                            50
                                                     52
                                                          53
                                                              54
                                                                   55
                                                                            57
##
    86
        87
                               47
                                                 51
                                                                       56
                                                                                58
##
              1
                   1
                            1
                                1
                                     1
                                         1
                                             1
                                                  1
                                                           1
                                                                1
                                                                    1
                                                                        1
                                                                             1
                                                                                 1
     1
          1
                       1
                                                      1
    59
        60
             92
                 61
                      62
                          63
                               64
                                   65
                                        66
                                            68
                                                 69
                                                     70
                                                          71
                                                              72
                                                                   73
                                                                       74
                                                                               198
##
     1
          1
              1
                   1
                       1
                            1
                                1
                                     1
                                         1
                                             1
                                                  1
                                                      1
                                                           1
                                                                1
                                                                    1
                                                                        1
                                                                             1
                                                                                 1
   199
       200
            201
                202 203
                         204
                              205
                                  206
                                      207
                                           208
                                                209
                                                    210 241
                                                             242
                                                                  115
                                                                      152
                                                                           153
                                                  1
     1
          1
              1
                   1
                       1
                           1
                                1
                                     1
                                         1
                                             1
                                                      1
                                                           1
                                                                1
```

```
## 155 157 158 159 160 161 162 163 164 165 166 148 149 150 151 176 177 178
                           1
                                1
                                     1
                                         1
                                             1
                                                  1
                                                      1
                                                           1
                                                               1
                                                                    1
                                                                        1
          1
              1
                   1
                       1
  179 180 181 182 183 187 188 145 146 147 167 168 169 170 171 172 173 174
                   1
                       1
                            1
                                1
                                     1
                                         1
                                             1
                                                  1
                                                      1
                                                           1
                                                               1
                                                                    1
                                                                        1
## 175 184
           185 138 139 140 141 142 143
                                          144
                                               190 191 192 193 194 195
##
                   1
                       1
                           1
                                1
                                     1
                                         1
                                             1
                                                  1
```

Everything adds up to 1, this is good! The code works.

#### Add plot order to data frame

I added an additional column to sort better for the stacked barplot. This was just a work around to get the facet\_grid() function to play nice with our data. I added a coulumn "barPlotOrder" and for each population I filled in a series 1:n foreach sample at each Site:Depth combo, so now there's no large blank expanses on the plot.

```
its2ra = its2NormPerc
sampleCounts = count(its2ra, c('Site', 'Depth'))
meltedList = melt(lapply(sampleCounts$freq,function(x){c(1:x)}))
its2ra$barPlotOrder = meltedList$value
colnames(its2ra)[c(5:ncol(its2ra)-1)] = c("sq01_C3", "sq10_C3", "sq11_C3g", "sq14_C3g", "sq17_C3g",
                                           "sq18_C3g", "sq25_C3g", "sq05_C3z", "sq07_C3e",
                                           "sq08_C3g")
its2ra=its2ra[c(1,ncol(its2ra),2:(ncol(its2ra)-1))]
head(its2ra)
       sample barPlotOrder Site Depth
                                         sq01_C3 sq10_C3 sq11_C3g
##
                                                                       sq14 C3g
                                    10 0.9052788
## 108
          108
                         1
                             TR
                                                       0
                                                                 0 0.00000e+00
## 109
          109
                         2
                             TR
                                    10 0.9079690
                                                       0
                                                                 0 0.000000e+00
                         3
                             TR
## 110
          110
                                    10 0.8739716
                                                       0
                                                                 0 2.649790e-05
                                                       0
## 111
          111
                         4
                             TR
                                    10 0.8787364
                                                                 0 0.00000e+00
## 112
          112
                         5
                             TR
                                    10 0.8811325
                                                       0
                                                                 0 0.00000e+00
                             TR
                                                                 0 3.105935e-05
##
  113
          113
                         6
                                    10 0.8653400
                                                       0
                                     sq25_C3g
##
           sq17_C3g
                       sq18_C3g
                                                sq05_C3z
                                                           sq07_C3e
## 108 0.000000e+00 0.000000000 0.000000e+00 0.07481792 0.01990326
## 109 0.000000e+00 0.000000000 0.000000e+00 0.07731046 0.01472053
## 110 1.324895e-05 0.002729284 4.416317e-06 0.08810553 0.03499048
## 111 0.000000e+00 0.000000000 0.000000e+00 0.07933163 0.04193194
## 112 0.000000e+00 0.000000000 0.000000e+00 0.08781788 0.03104964
## 113 6.655574e-05 0.003665003 6.211869e-05 0.10304603 0.02769606
           sq08 C3g
## 108 0.000000e+00
## 109 0.000000e+00
## 110 1.589874e-04
## 111 0.00000e+00
## 112 0.00000e+00
## 113 9.317804e-05
```

#### Create OTU stack for plotting

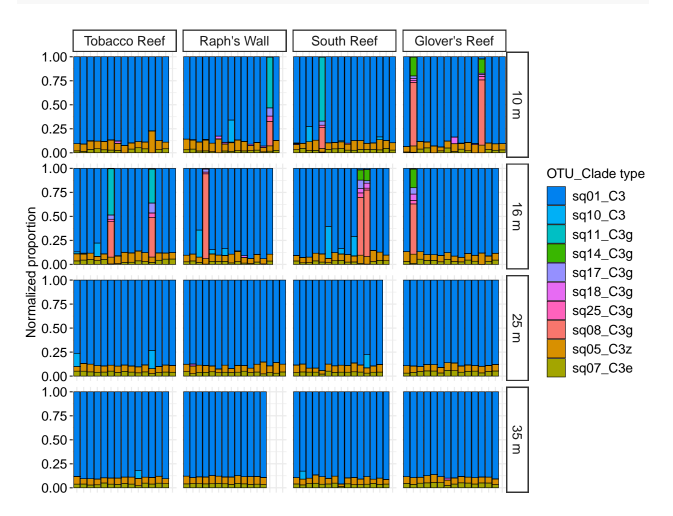
Stack the OTU data and adjest our columns and factor names for plotting

```
## [1] "summ"
## [7] "sq18_C3g" "sq25_C3g" "sq05_C3z" "sq07_C3e" "sq08_C3g"
gss\$otu = factor(gss\$otu, levels(gss\$otu)[c(1:8, 11, 9, 10)])
levels(gss$otu)
## [1] "summ"
                ## [7] "sq18_C3g" "sq25_C3g" "sq08_C3g" "sq05_C3z" "sq07_C3e"
levels(gss$Depth)
## [1] "10" "16" "25" "35"
levels(gss$Depth) = c("10 m", "16 m", "25 m", "35 m")
levels(gss$Site)
## [1] "TR" "RW" "SR" "GR"
levels(gss$Site) = c("Tobacco Reef", "Raph's Wall", "South Reef", "Glover's Reef")
levels(gss$Depth)
## [1] "10 m" "16 m" "25 m" "35 m"
levels(gss$Site)
## [1] "Tobacco Reef" "Raph's Wall"
                                 "South Reef"
                                              "Glover's Reef"
```

#### Consruct OTU barplot

```
OTUplotA = ggplot(gss, aes(x = barPlotOrder, y = count, fill = factor(otu))) +
          geom_bar(position = "stack", stat = "identity", color = "black",
                   size = 0.25) +
            ylab("Normalized proportion") +
            scale fill manual(values=its2ColPal)+
            labs(fill = "OTU_Clade type") +
            facet_grid(Depth ~ Site, scales = "free_x") + #faceting plots by Depth and Site
            theme_bw()
OTUplot = OTUplotA +
          theme(axis.title.x = element_blank(),
                axis.text.x = element_blank(),
                axis.ticks.x = element_blank(),
                axis.title.y = element_text(color = "black", size = 12),
                axis.text.y = element text(color = "black", size = 12),
                legend.position = "right",
                legend.title = element text(color = "black", size = 12),
                legend.text = element_text(color = "black", size = 12),
                legend.key = element_blank(),
                legend.background = element_blank(),
                panel.border = element_blank(),
                panel.background = element rect(fill = "white"),
                plot.background = element_blank(),
                strip.text.x = element_text(size = 12),
                strip.text.y = element_text(size = 12),
                strip.background = element_rect(fill = "white", size = 0.9)
                )
```

#### Save the OTU barplot



## ITS2 community differences

Now we can use permutational multivariate analysis of variance (PERMANOVA) to test for differences in Symbiodiniaceae  $\beta$ -diversity across Site and Depth based on our OTUs.

#### Cheking dispersion

Using betadisper() in *vegan* to look at multivariate homogeneity of dispersion between sites and depths. This is using Bray-Curtis similarity.

```
anova(betadisper(its2Dist, its2Norm$Depth))
```

```
## Warning in betadisper(its2Dist, its2Norm$Depth): some squared distances are
## negative and changed to zero
```

## Analysis of Variance Table

```
##
## Response: Distances
             Df Sum Sq Mean Sq F value
##
              3 0.5242 0.174742 5.0941 0.001971 **
## Groups
## Residuals 229 7.8553 0.034303
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
```

Dispersion is heteroschedastic, but PERMANOVA is robust to deviations in homgeneity of variance (Anderson and Walsh, 2013; https://esajournals.onlinelibrary.wiley.com/doi/10.1890/12-2010.1).

#### Running PERMANOVA in R

Now let's see how different communities are from each other with PERMANOVA. We will utilize the adonis() function in vegan. We will use Bray-Curtis similarity for our distance matrix and run a total 0f 9,999 permutations, and test the effects of Site, Depth, and the interaction between Site and Depth.

```
its2Adonis = adonis(its2Norm[, c(4:ncol(its2Norm))] ~ Depth*Site,
                     data = its2Norm, permutations = 9999, method = "bray")
its2Adonis
##
## Call:
## adonis(formula = its2Norm[, c(4:ncol(its2Norm))] ~ Depth * Site,
                                                                         data = its2Norm, permutations
##
## Permutation: free
## Number of permutations: 9999
## Terms added sequentially (first to last)
##
##
               Df SumsOfSqs MeanSqs F.Model
                                                  R2 Pr(>F)
## Depth
                3
                     0.3946 0.131523 3.4741 0.04443 0.0082 **
## Site
                3
                     0.0349 0.011633 0.3073 0.00393 0.9114
## Depth:Site
                9
                     0.2364 0.026268
                                      0.6938 0.02662 0.7949
## Residuals
              217
                     8.2153 0.037859
                                             0.92502
                                             1.00000
## Total
              232
                     8.8812
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
```

We see that  $\mathbf{Depth}$  has a significant effect on Symbiodiniaceae community structure in our M. cavernosasamples.

#### Pairwise PERMANOVA for multiple comparisons

Since we found that Depth was a significant factor in our PERMANOVA we can now use pairwise PER-MANOVA to reveal where differences occur across depth. This utilizes the package pairwiseAdonis, where we will again use Bray-Curtis similarity and 9,999 permutations. We also have added false discovery rate (FDR) corrections since we are perforing multiple comparisons.

```
its2PWAdonis = pairwise.adonis(its2Norm[, c(4:ncol(its2Norm))],
                  factors = its2Norm$Depth, sim.method = "bray",
                  p.adjust.m = "BH", perm = 9999)
its2PWAdonis
        pairs Df
                    {\tt SumsOfSqs}
                                F.Model
                                                  R2 p.value p.adjusted sig
              1 3.791992e-02 0.5212379 0.004473330
                                                     0.5278
                                                                 0.63336
## 2 10 vs 25  1 1.215016e-01 4.0807976 0.033703094 0.0003
```

0.00180

```
## 3 10 vs 35
               1 1.209040e-01 3.9840789 0.033767936
                                                      0.0026
                                                                0.00580
                                                      0.0029
## 4 16 vs 25
               1 2.553940e-01 5.8424234 0.048347453
                                                                0.00580
                                                                           *
## 5 16 vs 35
               1 2.551471e-01 5.7118059 0.048523645
                                                      0.0095
                                                                0.01425
## 6 25 vs 35
               1 7.599081e-05 0.1802355 0.001592465
                                                      0.8247
                                                                0.82470
```

We see that again see differences between our deeper (25 + 35 m) and shallower (10 + 16 m) samples.

#### PERMANOVA without 35 m samples

#### Subset our dataframe

First we need to remove the deep samples from the dataframe. We will use our dataframe of good OTUs, which haven't been normalized yet. This way we can calculate the normalization based on only the samples we are keeping in the analysis.

```
goods2 = subset(goods, !Depth=="35")
goods2[] = lapply(goods2, function(x) if(is.factor(x)) factor(x) else x)
summary(goods2)
```

```
##
       new0rder
                          sample
                                        Site
                                                 Depth
                                                                sq1
##
           : 1.0
                             : 1.00
                                        TR:44
                                                 10:60
                                                                  :
                                                                      224
    Min.
                     Min.
                                                          Min.
##
    1st Qu.: 59.0
                     1st Qu.: 46.00
                                        RW:45
                                                 16:58
                                                          1st Qu.: 90703
                                                          Median :132450
    Median :103.0
                     Median: 91.00
                                        SR:45
                                                 25:59
##
##
    Mean
            :109.6
                     Mean
                             : 92.51
                                        GR:43
                                                          Mean
                                                                  :132460
##
    3rd Qu.:160.0
                     3rd Qu.:136.00
                                                          3rd Qu.:176321
##
    Max.
            :219.0
                     Max.
                             :188.00
                                                          Max.
                                                                  :430489
##
         sq10
                                              sq14
                                                               sq17
                           sq11
##
                 0
                                    0
                                                      0
                                                                       0.0
    Min.
                     Min.
                                        Min.
                                                          Min.
                                    0
##
    1st Qu.:
                 0
                     1st Qu.:
                                        1st Qu.:
                                                          1st Qu.:
                                                                       0.0
                                                      0
    Median:
                     Median:
                                    0
                                        Median:
                                                      0
                                                          Median:
                                                                       0.0
                                 1592
##
    Mean
            : 1739
                     Mean
                                        Mean
                                                   504
                                                          Mean
                                                                     363.1
##
    3rd Qu.:
                 0
                     3rd Qu.:
                                    0
                                        3rd Qu.:
                                                      0
                                                          3rd Qu.:
                                                                       0.0
##
                                                                  :13521.0
    Max.
            :53422
                     Max.
                             :117162
                                        Max.
                                                :28206
                                                          Max.
                                             sq5
##
         sq18
                            sq25
                                                              sq7
##
    Min.
            :
                0.0
                       Min.
                              :
                                   0
                                       Min.
                                               :
                                                  935
                                                         Min.
##
    1st Qu.:
                0.0
                       1st Qu.:
                                   0
                                       1st Qu.: 7391
                                                         1st Qu.: 2389
                0.0
##
    Median:
                       Median:
                                   0
                                       Median :11115
                                                         Median: 4938
##
    Mean
            : 370.1
                       Mean
                               : 229
                                               :11759
                                                                 : 5134
                                       Mean
                                                         Mean
##
    3rd Qu.:
                0.0
                       3rd Qu.:
                                   0
                                       3rd Qu.:14646
                                                         3rd Qu.: 7149
            :9958.0
##
    Max.
                              :8747
                                               :56300
                                                                 :15419
                       Max.
                                       Max.
                                                         Max.
##
         sq8
##
                  0
    Min.
##
    1st Qu.:
                  0
##
                  0
    Median:
##
    Mean
               3870
##
    3rd Qu.:
                  0
    Max.
            :169042
```

#### Normalize reads

We will normalize samples again, as we did with the entire dataframe originally

```
itsgoods2Transposed = t(goods2[, 5:length(goods2[1, ])])
itsgoods2List = DGEList(counts = itsgoods2Transposed)
head(itsgoods2List$samples)
```

```
## group lib.size norm.factors
```

```
## 108
           1
               237951
## 109
               18070
           1
## 110
              226433
## 111
              204975
                                 1
## 112
                27762
                                 1
## 113
               225375
           1
                                 1
its2Norm2 = calcNormFactors(itsgoods2List, method = "TMM")
head(its2Norm2$samples)
       group lib.size norm.factors
## 108
           1
              237951
                         1.080109
## 109
               18070
                          1.084610
## 110
              226433
                          1.047130
           1
## 111
               204975
                          1.052500
## 112
               27762
                          1.055361
           1
## 113
               225375
                          1.037930
its2TMM = t(cpm(its2Norm2, Normalized.lib.sizes = TRUE))
its2Norm2 = cbind(goods2[,c(2:4)], its2TMM)
head(its2Norm2)
##
       sample Site Depth
                              sq1 sq10 sq11
                                                sq14
                                                         sq17
                                                                  sq18
## 108
          108
                      10 838136.9
                                     0
                                          0 0.00000 0.00000
                                                                 0.000
## 109
                      10 837139.0
                                          0 0.00000 0.00000
                                                                 0.000
          109
                TR
                                     0
## 110
         110
               TR
                      10 834635.2
                                     0
                                          0 25.30527 12.65263 2606.443
## 111
               TR
                      10 834904.0
                                     0
                                          0 0.00000 0.00000
                                                                 0.000
         111
## 112
         112
                TR
                      10 834911.1
                                          0 0.00000 0.00000
                                                                 0.000
## 113
                                          0 29.92431 64.12352 3531.068
          113
                TR
                      10 833716.8
##
            sq25
                      sq5
                               sq7
                                         sq8
## 108 0.000000 69268.90 18427.09
                                     0.00000
       0.000000 71279.53 13572.19
                                     0.00000
## 110 4.217545 84140.02 33415.61 151.83161
## 111 0.000000 75374.47 39840.33
                                     0.00000
## 112 0.000000 83211.23 29420.87
                                     0.00000
## 113 59.848614 99280.30 26683.93 89.77292
Run PERMANOVA on subset of data
its2Dist2 = vegdist(its2Norm2[, c(4:ncol(its2Norm2))], method = "bray")
anova(betadisper(its2Dist2, its2Norm2$Depth))
## Warning in betadisper(its2Dist2, its2Norm2$Depth): some squared distances
## are negative and changed to zero
## Analysis of Variance Table
##
## Response: Distances
              Df Sum Sq Mean Sq F value Pr(>F)
               2 0.3443 0.172163 3.9313 0.02138 *
## Groups
## Residuals 174 7.6199 0.043793
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
its2Adonis2 <- adonis(its2Norm2[, c(4:ncol(its2Norm2))] ~ Depth*Site,
                       data = its2Norm2, permutations = 9999, method = "bray")
```

#### its2Adonis2

```
##
## Call:
                                                                     Site, data = its2Norm2, permutation
## adonis(formula = its2Norm2[, c(4:ncol(its2Norm2))] ~ Depth *
##
## Permutation: free
## Number of permutations: 9999
##
## Terms added sequentially (first to last)
##
               Df SumsOfSqs MeanSqs F.Model
##
                                                  R2 Pr(>F)
## Depth
                2
                     0.2828 0.141396 2.92686 0.03326 0.0368 *
                     0.0367 0.012230 0.25316 0.00432 0.9386
## Site
                3
## Depth:Site
                6
                     0.2111 0.035186 0.72835 0.02483 0.6873
## Residuals
              165
                     7.9711 0.048310
                                             0.93759
## Total
              176
                     8.5017
                                             1.00000
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
```

Depth is still has a significant effect on community structure

#### Pairwise PERMANOVA

Let' see where differences lie across depth using pairwise permanova again.

The differences are still between 25 m and the shallower sites (10 + 16 m). This gives us confidence that our deeper samples aren't changing our interpretation.

0.00345

#### Generalized linear mixed model of OTUs

## 3 16 vs 25 1 0.26179196 6.1925677 0.051096926 0.0023

To see how significantly changing OTUs across depth we will use the package MCMC.OTU, which performs Bayesian analysis of multivariate count data. This will create a MCMC-based GLMM analysis of our OTU data.

#### Stack OTU data table

```
## [1] "summ"
  [7] "sq18_C3g" "sq25_C3g" "sq05_C3z" "sq07_C3e" "sq08_C3g"
glmStack$otu = factor(glmStack$otu, levels(glmStack$otu)[c(1:8, 11, 9, 10)])
levels(glmStack$otu)
## [1] "summ"
               ## [7] "sq18_C3g" "sq25_C3g" "sq08_C3g" "sq05_C3z" "sq07_C3e"
glmStack$count = round(glmStack$count, 0)
head(glmStack)
##
    count
            otu sample Site Depth
## 1 830135 sq01 C3
                108
                       TR
                            10
                109
## 2 828910 sq01_C3
                       TR
                            10
## 3 826200 sq01_C3 110 TR
                           10
## 4 826360 sq01_C3
                  111 TR
                            10
## 5 826547 sq01_C3
                112 TR
                           10
## 6 825666 sq01_C3
                  113 TR
                           10
```

#### Creating the model

Now that we have our data setup correctly we can create the GLMM

#### Fitting the model

Here we are using Depth as a fixed effect and running 10<sup>6</sup> iterations with a 5000 burn-in period.

#### Calculate effect size and p-values

```
ssD = OTUsummary(its2MmD, glmStack, summ.plot = FALSE)
ssD = padjustOTU(ssD)
sigsD = signifOTU(ssD)
ss2D = OTUsummary(its2MmD, glmStack, otus = sigsD, whiskers = "sd",
                  ptype = "mcmc")
its2MCMCglmD = ss2D$summary
levels(its2MCMCglmD$otu)
## [1] "sq01_C3" "sq07_C3e" "sq08_C3g" "sq10_C3" "sq14_C3g" "sq17_C3g"
## [7] "sq18 C3g" "sq25 C3g"
its2MCMCglmD$Depth = as.factor(its2MCMCglmD$Depth)
its2MCMCglmD$otu = factor(its2MCMCglmD$otu, levels = c("sq01_C3", "sq10_C3",
                          "sq14_C3g", "sq17_C3g", "sq18_C3g", "sq25_C3g",
                          "sq08_C3g", "sq07_C3e"))
ssD$otuWise[sigsD]
## $sq01_C3
        difference
## pvalue
                                         25
                                                    35
                10
                               16
##
                 NA -0.061342676 0.1662458 0.16790133
```

```
16 0.49373133 NA 0.2275885 0.22924401
     25 0.03691520 0.005191352 NA 0.00165554
##
     35 0.03737995 0.005401985 0.9983354 NA
##
## $sq10 C3
## difference
## pvalue 10 16 25 35
     10 NA 2.09427370 -3.6073627 -5.064233
     16 0.3776807 NA -5.7016364 -7.158507
     25 0.2228695 0.03645170 NA -1.456871
     35 0.1137648 0.02472841 0.7422496 NA
##
## $sq14_C3g
## difference
## pvalue 10 16 25 35
## 10 NA -2.23508519 -7.0628806 -6.8949807
##
     16 0.040969131 NA -4.8277954 -4.6598955
     25 0.001607853 0.02499062 NA 0.1678999
##
     35 0.001178064 0.02313775 0.9983354
##
## $sq17_C3g
## difference
## pvalue 10 16 25 35
          NA -1.388122781 -7.5474321 -7.4377685
     16 0.206788675 NA -6.1593093 -6.0496457
     25 0.001245135 0.006000357 NA 0.1096636
##
     35 0.001241094 0.006041373 0.9983354
## $sq18_C3g
## difference
## pvalue 10 16 25 35
## 10 NA -3.16142817 -9.626732 -9.632088918
     16 0.024865893 NA -6.465303 -6.470660752
     25 0.000949371 0.01900819 NA -0.005357278
     35 0.001178064 0.02169629 0.998571
##
##
## $sq25 C3g
## difference
## pvalue 10 16 25
              NA -1.272909432 -7.0865630 -7.01764595
     16 0.236396410 NA -5.8136536 -5.74473652
     25 0.001245135 0.006000357 NA 0.06891705
     35 0.001477129 0.007169117 0.9983354 NA
##
## $sq08_C3g
## difference
## pvalue 10 16 25 ## 10 NA -1 700710114 0 7007101
              NA -1.799719114 -8.7903141 -8.69496991
     16 0.152251158 NA -6.9905950 -6.89525079
     25 0.000949371 0.004565093
                             NA 0.09534424
##
##
     35 0.000949371 0.004776074 0.9983354 NA
## $sq07_C3e
## difference
```

```
## pvalue
                    10
                               16
                                         25
                   NA 0.34735180 0.6075924 0.59452579
##
      10
                               NA 0.2602406 0.24717399
##
       16 0.0190342629
       25 0.0001186316 0.07833303
                                         NA -0.01306663
##
##
       35 0.0001186316 0.10555832 0.9983354
                                                     NA
```

#### Plotting the GLMM

We are now plotting the GLMM using our color pallete we created for the OTUs

```
pd = position_dodge(0.3)
its2glmPlotA = ggplot(its2MCMCglmD, aes(x = Depth, y = mean, group = otu,
                      colour =otu))+
                      geom errorbar(aes(ymin = mean-sd, ymax = mean+sd),
                                    lwd = 0.6, width = 0.5,
                                                                position = pd) +
                      geom_line(aes(group = otu), lwd = 0.6, position = pd)+
                      geom_point(aes(group = otu), position = pd, size = 2.5)+
                      scale color manual(values = its2ColPal[c(1,2,4:8,10)],
                                         name = "OTU_Clade type") +
                      xlab("Depth")+
                      ylab("log10(proportion)") +
                      theme_bw()
its2glmPlotD = its2glmPlotA +
               theme(axis.title.x = element_text(color = "black", size = 12),
                     axis.text.x = element_text(color = "black", size = 12),
                     axis.title.y = element_text(color = "black", size = 12),
                     axis.text.y = element_text(color = "black", size = 12),
                     legend.position = "right",
                     legend.title = element_text(color = "black", size = 12),
                     legend.text = element_text(color = "black", size = 12),
                     legend.key = element_blank(),
                     legend.background = element_blank(),
                     panel.border = element_rect(color = "black", size = 1.2),
                     panel.background = element_rect(fill = "white"),
                     plot.background = element_blank()
                     )
its2glmPlotD
```

#### Save GLMM plot

```
ggsave("its2glmDepth.eps", plot=its2glmPlotD, width = 8, height = 6, dpi = 600)
```

