coralMicrobiomes

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This is an R Markdown document detailing the statistical and graphical steps for reproducting the results in: Neave, M.J., Rachmawati, R., Xun, L., Michell, C.T., Barber, P.H., Bourne, D.G., McCulloch, M.T., Apprill, A., Voolstra, C.R. A global microbiome analysis reveals that closely related corals exhibit fine-scale differences in their association with Endozoicomonas symbionts.

Load required libraries

```
library("phyloseq"); packageVersion("phyloseq")
## [1] '1.9.15'
library("ggplot2"); packageVersion("ggplot2")
## [1] '1.0.0'
library("plyr"); packageVersion("plyr")
## [1] '1.8.1'
library("vegan"); packageVersion("vegan")
## [1] '2.0.10'
library("grid"); packageVersion("grid")
## [1] '3.1.1'
library("knitr"); packageVersion("knitr")
## [1] '1.6'
library("clustsig"); packageVersion("clustsig")
## Warning: package 'clustsig' was built under R version 3.1.2
## [1] '1.1'
```

```
library('ape'); packageVersion("ape")

## [1] '3.1.4'

library('RColorBrewer'); packageVersion("RColorBrewer")

## [1] '1.0.5'

library("dunn.test"); packageVersion("dunn.test")

## Warning: package 'dunn.test' was built under R version 3.1.2

## [1] '1.2.3'

library("DESeq2"); packageVersion("DESeq2")

## Warning: package 'RcppArmadillo' was built under R version 3.1.2

## [1] '1.4.5'

setwd("./data")
opts_knit$set(root.dir = "./data")
#opts_chunk$set(tidy.opts=list(width.cutoff=80))
```

Import data

First the matrix percent file and count file generated by the minimum entropy decomposition (MED) pipeline, subsampled to 7974 reads per sample, and the accordated taxonony file

```
allShared = read.table("all.7974.matrixPercent.txt", header = T, row.names = 1)
allCounts = read.table("all.7974.matrixCount.txt", header = T, row.names = 1)
allTax = read.table("all.7974.nodeReps.nr_v119.knn.taxonomy", header = T, sep = "\t",
    row.names = 1)
```

Warning: number of items read is not a multiple of the number of columns

```
allTax = allTax[, 2:8]
allTax = as.matrix(allTax)
```

Import the shared and taxonomy files generated in mothur for 3% and 1% pairwise similarity, in order to calculate alpha diversity measures and to compare to the MED procedure. Also import the 3% OTU file without any subsampling for alpha diversity calculations.

```
all30TUshared = read.table("all.7974.0.03.pick.shared", header=T, row.names=2)
all30TUshared = all30TUshared[,3:length(all30TUshared)]
alpha30TUshared = read.table("all.7974.0.03.shared", header=T)
rownames(alpha30TUshared) = alpha30TUshared[,2]
alpha30TUshared = alpha30TUshared[,4:length(alpha30TUshared)]
all10TUshared = read.table("all.7974.0.01.pick.shared", header=T, row.names=2)
all10TUshared = all10TUshared[,3:length(all10TUshared)]
all30TUtax = read.table('all.7974.0.03.taxonomy', header=T, sep='\t', row.names=1)
all30TUtax = all30TUtax[,2:8]
all10TUtax = read.table('all.7974.0.01.taxonomy', header=T, sep='\t', row.names=1)
all10TUtax = all10TUtax[,2:8]
all10TUtax = all10TUtax[,2:8]
all10TUtax = as.matrix(all10TUtax)
```

Import Endozoicomonas phylogenetic tree (exported from ARB) using the APE package (Fig. 3). Also import a MED percent matrix that is slightly modified to accommodate the tree

```
endoTreeFile = read.tree(file='MEDNJ5.tree')
allSharedTree = read.table("all.7974.matrixPercent.tree.txt", header=T, row.names=1)
```

Import meta data for the samples, including metaData3.txt, which is slightly modified to accommodate heatmap sample ordering, and metaDataChem which contains additional columns of physiochemical data

```
metaFile = read.table('metaData2.MED', header=T, sep='\t', row.names=1)
metaFile3 = read.table('metaData3.txt', header=T, sep='\t', row.names=1)
metaFileChem = read.table('metaDataChem.txt', header=T, sep='\t', row.names=1)
```

Create phyloseq objects and add consistent coloring for sites

```
OTU = otu_table(allShared, taxa_are_rows = FALSE)
OTUcounts = otu_table(allCounts, taxa_are_rows = FALSE)
OTUs3 = otu_table(all30TUshared, taxa_are_rows = FALSE)
OTUs3alpha = otu_table(alpha30TUshared, taxa_are_rows = FALSE)
OTUs1 = otu_table(all10TUshared, taxa_are_rows = FALSE)
OTUtree = otu_table(allSharedTree, taxa_are_rows = FALSE)

TAX = tax_table(allTax)
TAX3 = tax_table(all30TUtax)
TAX1 = tax_table(all10TUtax)

META = sample_data(metaFile)
METAchem = sample_data(metaFileChem)
TREE = phy_tree(endoTreeFile)

allPhylo = phyloseq(OTU, TAX, META)
countPhylo = phyloseq(OTUcounts, TAX, META)
all30TUphylo = phyloseq(OTUcounts, TAX, META)
```

```
alpha30TUphylo = phyloseq(OTUs3alpha, META)
all10TUphylo = phyloseq(OTUs1, TAX1, META)
allPhyloChem = phyloseq(OTU, TAX, METAchem)
endoTree = phyloseq(OTUtree, META, TREE)

cols <- c(AmericanSamoa = "#D95F02", Indonesia = "#A6761D", MaggieIs = "#666666",
    Maldives = "#E6AB02", Micronesia = "#66A61E", Ningaloo = "#7570B3", RedSea = "#E7298A",
    other = "black")</pre>
```

Ordinations to compare MED vs pairwise OTUs

Subset samples for the two corals, remove taxa with 0s, create relative abundance and square-root sample counts

```
filter stylo data <- function(initial matrix){
  initial_coral <- subset_samples(initial_matrix, species=="Stylophora pistillata")</pre>
  coral_filt = filter_taxa(initial_coral, function(x) mean(x) > 0, TRUE)
  coral_filt_rel = transform_sample_counts(coral_filt, function(x) x / sum(x) )
  coral_filt_rel_sqrt = transform_sample_counts(coral_filt_rel, function(x) sqrt(x) )
  return(coral filt rel sqrt)
filter_pverr_data <- function(initial_matrix){</pre>
  initial_coral <- subset_samples(initial_matrix, species=="Pocillopora verrucosa")</pre>
  coral_filt = filter_taxa(initial_coral, function(x) mean(x) > 0, TRUE)
  coral_filt_rel = transform_sample_counts(coral_filt, function(x) x / sum(x) )
  coral_filt_rel_sqrt = transform_sample_counts(coral_filt_rel, function(x) sqrt(x) )
  return(coral_filt_rel_sqrt)
}
spistPhyloRelSqrt <- filter_stylo_data(allPhylo)</pre>
spist30TUphyloRelSqrt <- filter_stylo_data(all30TUphylo)</pre>
spist10TUphyloRelSqrt <- filter_stylo_data(all10TUphylo)</pre>
pverrPhyloRelSqrt <- filter_pverr_data(allPhylo)</pre>
pverr30TUphyloRelSqrt <- filter_pverr_data(all30TUphylo)</pre>
pverr10TUphyloRelSqrt <- filter_pverr_data(all10TUphylo)</pre>
```

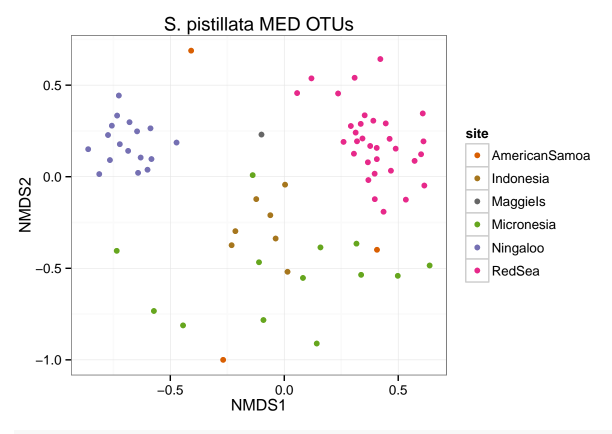
Now do ordinations for each

Run 1 stress 0.2459

```
compOrdinations <- function(sample_data, sample_name){
theme_set(theme_bw())
sample_dataOrd <- ordinate(sample_data, "NMDS", "bray")
plot_ordination(sample_data, sample_dataOrd, type = 'samples', color='site', title=sample_name) +
    geom_point(size=2) +
    scale_color_manual(values=cols)
}
compOrdinations(spistPhyloRelSqrt, "S. pistillata MED OTUs")

## Run O stress 0.2254</pre>
```

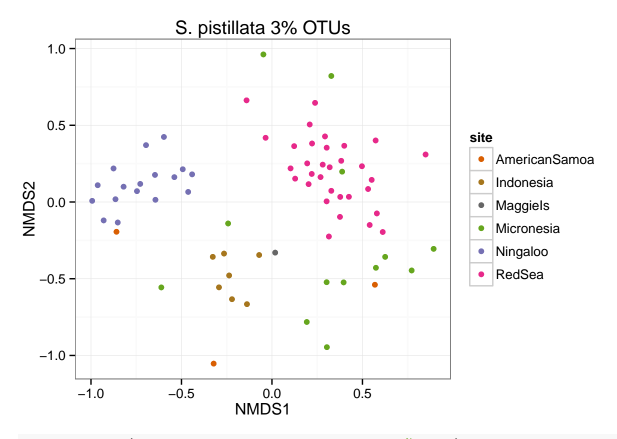
```
## Run 2 stress 0.2413
## Run 3 stress 0.2399
## Run 4 stress 0.2389
## Run 5 stress 0.2373
## Run 6 stress 0.2344
## Run 7 stress 0.2328
## Run 8 stress 0.2371
## Run 9 stress 0.229
## Run 10 stress 0.2273
## Run 11 stress 0.2341
## Run 12 stress 0.2302
## Run 13 stress 0.2352
## Run 14 stress 0.2288
## Run 15 stress 0.2503
## Run 16 stress 0.2381
## Run 17 stress 0.2486
## Run 18 stress 0.2409
## Run 19 stress 0.2242
## ... New best solution
## ... procrustes: rmse 0.02453 max resid 0.1974
## Run 20 stress 0.2292
```



compOrdinations(spist3OTUphyloRelSqrt, "S. pistillata 3% OTUs")

```
## Run 0 stress 0.2273
## Run 1 stress 0.235
## Run 2 stress 0.248
```

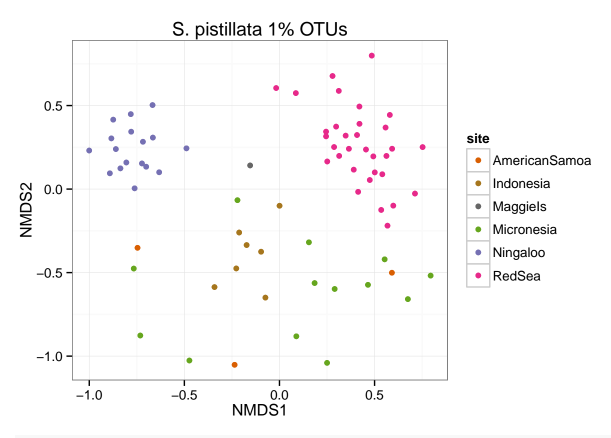
```
## Run 3 stress 0.2386
## Run 4 stress 0.2395
## Run 5 stress 0.2663
## Run 6 stress 0.2542
## Run 7 stress 0.2273
## ... New best solution
## ... procrustes: rmse 0.0004645 max resid 0.003106
## *** Solution reached
```



compOrdinations(spist10TUphyloRelSqrt, "S. pistillata 1% OTUs")

```
## Run 0 stress 0.2225
## Run 1 stress 0.2575
## Run 2 stress 0.2512
## Run 3 stress 0.2274
## Run 4 stress 0.2645
## Run 5 stress 0.2512
## Run 6 stress 0.2366
## Run 7 stress 0.2274
## Run 8 stress 0.2351
## Run 9 stress 0.2532
## Run 10 stress 0.2304
## Run 11 stress 0.2582
## Run 12 stress 0.2338
## Run 13 stress 0.2318
## Run 14 stress 0.2331
## Run 15 stress 0.2399
```

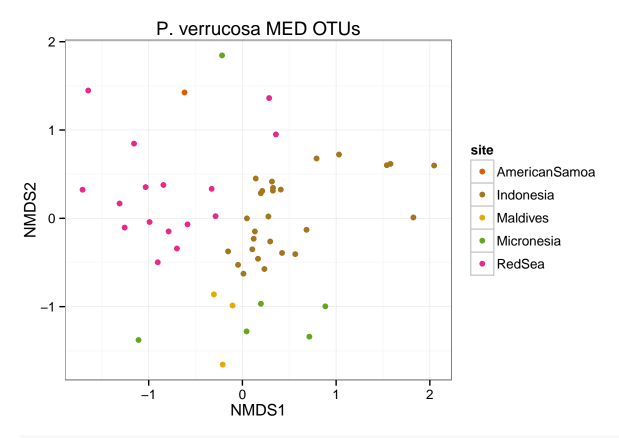
```
## Run 16 stress 0.2586
## Run 17 stress 0.2571
## Run 18 stress 0.2417
## Run 19 stress 0.2287
## Run 20 stress 0.2559
```



compOrdinations(pverrPhyloRelSqrt, "P. verrucosa MED OTUs")

```
## Run 0 stress 0.2437
## Run 1 stress 0.2485
## Run 2 stress 0.271
## Run 3 stress 0.2362
## ... New best solution
## ... procrustes: rmse 0.09863 max resid 0.3381
## Run 4 stress 0.2446
## Run 5 stress 0.2349
## ... New best solution
## ... procrustes: rmse 0.07606 max resid 0.353
## Run 6 stress 0.2133
## ... New best solution
## ... procrustes: rmse 0.08037 max resid 0.3474
## Run 7 stress 0.2259
## Run 8 stress 0.2227
## Run 9 stress 0.2438
## Run 10 stress 0.2136
## ... procrustes: rmse 0.02494 max resid 0.146
## Run 11 stress 0.2247
```

```
## Run 12 stress 0.2454
## Run 13 stress 0.2186
## Run 14 stress 0.2393
## Run 15 stress 0.2393
## Run 16 stress 0.2206
## Run 17 stress 0.2315
## Run 18 stress 0.2572
## Run 20 stress 0.239
```



compOrdinations(pverr30TUphyloRelSqrt, "P. verrucosa 3% OTUs")

```
## Run 0 stress 0.2409

## Run 1 stress 0.2354

## ... New best solution

## Run 2 stress 0.2482

## Run 3 stress 0.2598

## Run 4 stress 0.2349

## ... New best solution

## ... procrustes: rmse 0.09954 max resid 0.4044

## Run 5 stress 0.23

## ... New best solution

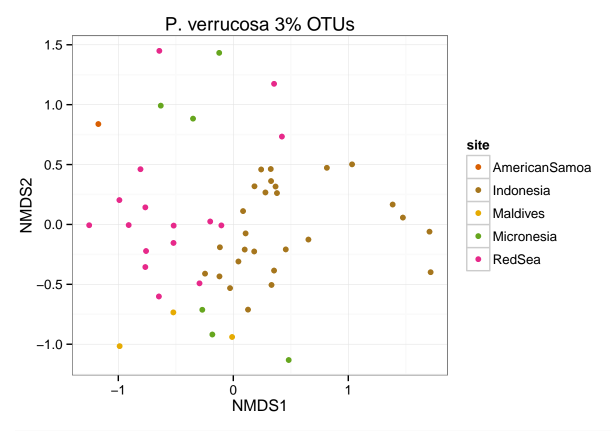
## ... procrustes: rmse 0.1085 max resid 0.3136

## Run 6 stress 0.2283

## ... New best solution

## ... procrustes: rmse 0.09872 max resid 0.2776
```

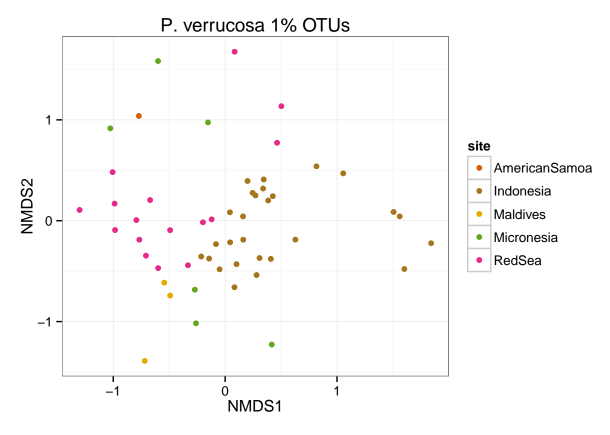
```
## Run 7 stress 0.224
## ... New best solution
## ... procrustes: rmse 0.0909 max resid 0.2662
## Run 8 stress 0.2289
## Run 9 stress 0.2407
## Run 10 stress 0.2288
## Run 11 stress 0.2238
## ... New best solution
## ... procrustes: rmse 0.04183 max resid 0.2614
## Run 12 stress 0.2593
## Run 13 stress 0.2852
## Run 14 stress 0.2277
## Run 15 stress 0.227
## Run 16 stress 0.2358
## Run 17 stress 0.2468
## Run 18 stress 0.2353
## Run 19 stress 0.2316
## Run 20 stress 0.4043
```



compOrdinations(pverr10TUphyloRelSqrt, "P. verrucosa 1% OTUs")

```
## Run 0 stress 0.234
## Run 1 stress 0.2289
## ... New best solution
## ... procrustes: rmse 0.08099 max resid 0.2296
## Run 2 stress 0.2458
## Run 3 stress 0.217
```

```
## ... New best solution
## ... procrustes: rmse 0.09259 max resid 0.4139
## Run 4 stress 0.2297
## Run 5 stress 0.2492
## Run 6 stress 0.2473
## Run 7 stress 0.228
## Run 8 stress 0.2341
## Run 9 stress 0.2263
## Run 10 stress 0.2306
## Run 11 stress 0.2187
## Run 12 stress 0.2263
## Run 13 stress 0.2331
## Run 14 stress 0.2356
## Run 15 stress 0.2355
## Run 16 stress 0.2172
  ... procrustes: rmse 0.04413 max resid 0.297
## Run 17 stress 0.2188
## Run 18 stress 0.2283
## Run 19 stress 0.2181
## Run 20 stress 0.2322
```



Alpha diversity measures

First subset the corals, then plot using phyloseq and ggplot2

Note: I'll use unsubampled 3% pairwise OTUs for calculation of alpha diversity measures as this will make them more comparable to other studies, plus the MED pipeline is has not yet implemented alpha diversity

```
allAlphaTmp <- subset_samples(alpha30TUphylo, species == "seawater")</pre>
allAlphaTmp2 <- subset_samples(alpha3OTUphylo, species == "Stylophora pistillata")
allAlphaTmp3 <- subset_samples(alpha3OTUphylo, species == "Pocillopora verrucosa")
allAlpha2 <- merge_phyloseq(allAlphaTmp, allAlphaTmp2, allAlphaTmp3)
allAlphaPlot2 <- plot_richness(allAlpha2, x = "species", measures = c("Chao1", "Simpson",
    "observed"), color = "site", sortby = "Chao1")
ggplot(data = allAlphaPlot2$data) + geom_point(aes(x = species, y = value, color = site),
    position = position_jitter(width = 0.1, height = 0)) + geom_boxplot(aes(x = species,
    y = value, color = NULL), alpha = 0.1, outlier.shape = NA) + scale_color_manual(values = cols) +
    theme(axis.text.x = element_text(angle = 90)) + facet_wrap(~variable, scales = "free_y") +
    scale_x_discrete(limits = c("Stylophora pistillata", "Pocillopora verrucosa",
         "seawater"))
## Warning: Removed 3 rows containing missing values (geom_point).
## Warning: Removed 7 rows containing missing values (geom point).
## Warning: Removed 4 rows containing missing values (geom_point).
## Warning: Removed 6 rows containing missing values (geom point).
## Warning: Removed 4 rows containing missing values (geom_point).
## Warning: Removed 7 rows containing missing values (geom_point).
## Warning: Removed 4 rows containing missing values (geom_point).
## Warning: Removed 4 rows containing missing values (geom_point).
            Observed
                                    Chao1
                                                        Simpson
                                               1.00
                                                                       site
  2000
                                                                           AmericanSamoa
                         4000
                                               0.75
                                                                           Indonesia
  1500
                                                                           Maggiels
                                                                           Maldives
                                               0.50
  1000
                         2000
                                                                           Micronesia
                                                                           Ningaloo
   500
                                               0.25
                                                                           RedSea
                            0
      0
                Pocillopora verrucosa
                                      Pocillopora verrucosa
                                                            Pocillopora verrucosa
           Stylophora pistillata
                                  Stylophora pistillata
                                                       Stylophora pistillata
                     seawater
                                           seawater
                                                                 seawater
```

Check for significant differences in the alpha diversity measures using a kruskal-wallis test and a dunn post-hoc test to check which specific groups were different

species

```
alphaObserved = estimate_richness(allAlpha2, measures="Observed")
alphaSimpson = estimate_richness(allAlpha2, measures="Simpson")
alphaChao = estimate_richness(allAlpha2, measures="Chao1")
alpha.stats <- cbind(alphaObserved, sample_data(allAlpha2))</pre>
alpha.stats2 <- cbind(alpha.stats, alphaSimpson)</pre>
alpha.stats3 <- cbind(alpha.stats2, alphaChao)</pre>
kruskal.test(Observed~species, data = alpha.stats3)
##
## Kruskal-Wallis rank sum test
##
## data: Observed by species
## Kruskal-Wallis chi-squared = 61.88, df = 2, p-value = 3.662e-14
dunn.test(alpha.stats3$Observed, alpha.stats3$species, method="bonferroni")
     Kruskal-Wallis rank sum test
##
##
## data: x and group
## Kruskal-Wallis chi-squared = 61.8764, df = 2, p-value = 0
##
##
##
                              Comparison of x by group
                                     (Bonferroni)
##
## Col Mean-|
## Row Mean |
                Pocillop
                           seawater
## seawater |
                7.510384
##
                  0.0000
##
## Stylopho |
                1.357184
                          -6.783011
##
            1
                  0.2621
                             0.0000
kruskal.test(Simpson~species, data = alpha.stats3)
##
## Kruskal-Wallis rank sum test
## data: Simpson by species
## Kruskal-Wallis chi-squared = 12.25, df = 2, p-value = 0.002193
dunn.test(alpha.stats3$Simpson, alpha.stats3$species, method="bonferroni")
##
     Kruskal-Wallis rank sum test
##
## data: x and group
## Kruskal-Wallis chi-squared = 12.2453, df = 2, p-value = 0
##
```

```
##
                              Comparison of x by group
##
                                     (Bonferroni)
## Col Mean-|
## Row Mean |
                Pocillop
                           seawater
## -----
## seawater |
                3.397898
##
          - 1
                  0.0010
##
            ## Stylopho |
                0.811204
                          -2.904738
            ١
                  0.6259
                             0.0055
kruskal.test(Chao1~species, data = alpha.stats3)
##
##
   Kruskal-Wallis rank sum test
##
## data: Chao1 by species
## Kruskal-Wallis chi-squared = 64.31, df = 2, p-value = 1.086e-14
dunn.test(alpha.stats3$Chao1, alpha.stats3$species, method="bonferroni")
     Kruskal-Wallis rank sum test
##
##
## data: x and group
## Kruskal-Wallis chi-squared = 64.3067, df = 2, p-value = 0
##
##
##
                              Comparison of x by group
##
                                     (Bonferroni)
## Col Mean-|
## Row Mean |
                Pocillop
                           seawater
## ----+
## seawater |
                7.581725
##
                  0.0000
## Stylopho |
                1.146749
                          -7.033279
                  0.3772
                             0.0000
```

In each case, the seawater was significantly different to the corals, while the corals were not different to each other. This suggests the corals have a more 'selective' community of microbes compared to the surrounding seawater.

Similarity Profile Analysis (SIMPROF)

This will show how the samples cluster without any a priori assumptions regarding sample origin

Need to import the shared file containing just spist OTUs, then calcualte the simprof clusters based on the braycurtis metric.

```
# spist <- subset_samples(allPhylo, species=='Stylophora pistillata') spistShared
# = otu_table(spist) class(spistShared) <- 'numeric' spistSIMPROF <-
# simprof(spistShared, num.expected=1000, num.simulated=999,</pre>
```

```
# method.cluster='average', method.distance='braycurtis',
# method.transform='squareroot', alpha=0.05, sample.orientation='row',
# silent=FALSE) simprof.plot(spistSIMPROF, leafcolors=NA, plot=TRUE, fill=TRUE,
# leaflab='perpendicular', siglinetype=1) pVerr <- subset_samples(allPhylo,
# species=='Pocillopora verrucosa') pVerrShared = otu_table(pVerr)
# class(pVerrShared) <- 'numeric' pVerrSIMPROF <- simprof(pVerrShared,
# num.expected=1000, num.simulated=999, method.cluster='average',
# method.distance='braycurtis', method.transform='squareroot', alpha=0.05,
# sample.orientation='row', silent=FALSE) simprof.plot(pVerrSIMPROF,
# leafcolors=NA, plot=TRUE, fill=TRUE, leaflab='perpendicular', siglinetype=1)</pre>
```

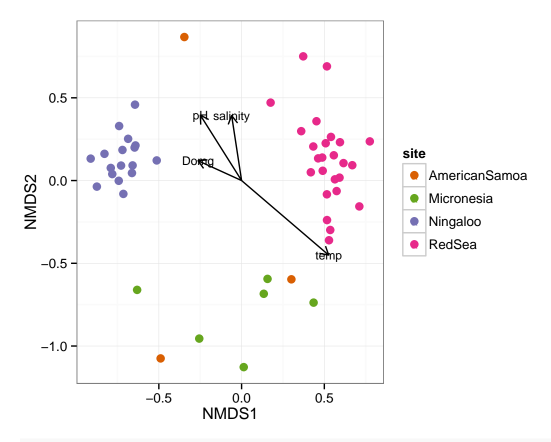
Chemical and biological correlations

Use the envfit function from the Vegan package to test if any environmental variables are significantly correlated with microbiome differences in the corals

```
draw_envfit_ord <- function(coral_chem, env_data) {</pre>
    chemNoNA <- na.omit(metaFileChem[sample_names(coral_chem), env_data])</pre>
    coralNoNA <- prune_samples(rownames(chemNoNA), coral_chem)</pre>
    theme_set(theme_bw())
    coralNoNAOrd <- ordinate(coralNoNA, "NMDS", "bray")</pre>
    coralNoNAOrdPlot <- plot_ordination(coralNoNA, coralNoNAOrd, type = "samples",</pre>
        color = "site") + geom_point(size = 3) + scale_color_manual(values = c(cols))
    # get points for ggplot
    pointsNoNA <- coralNoNAOrd$points[rownames(chemNoNA), ]</pre>
    chemFit <- envfit(pointsNoNA, env = chemNoNA, na.rm = TRUE)</pre>
    print(chemFit)
    chemFit.scores <- as.data.frame(scores(chemFit, display = "vectors"))</pre>
    chemFit.scores <- cbind(chemFit.scores, Species = rownames(chemFit.scores))</pre>
    # create arrow info
    chemNames <- rownames(chemFit.scores)</pre>
    arrowmap <- aes(xend = MDS1, yend = MDS2, x = 0, y = 0, shape = NULL, color = NULL)
    labelmap <- aes(x = MDS1, y = MDS2 + 0.04, shape = NULL, color = NULL, size = 1.5,
        label = chemNames)
    arrowhead = arrow(length = unit(0.25, "cm"))
    # note: had to use aes_string to get applot to recognize variables
    coralNoNAOrdPlot + coord_fixed() + geom_segment(arrowmap, size = 0.5, data = chemFit.scores,
        color = "black", arrow = arrowhead, show guide = FALSE) + geom text(aes string(x = "MDS1",
        y = "MDS2", shape = NULL, color = NULL, size = 1.5, label = "Species"), size = 3,
        data = chemFit.scores)
}
waterQual <- c("temp", "salinity", "Domg", "pH")</pre>
nutrients <- c("PO4", "N.N", "silicate", "NO2", "NH4")
FCM <- c("prok", "syn", "peuk", "pe.peuk", "Hbact")</pre>
spistChem <- subset_samples(allPhyloChem, species == "Stylophora pistillata")</pre>
pverrChem <- subset_samples(allPhyloChem, species == "Pocillopora verrucosa")</pre>
```

draw_envfit_ord(spistChem, waterQual)

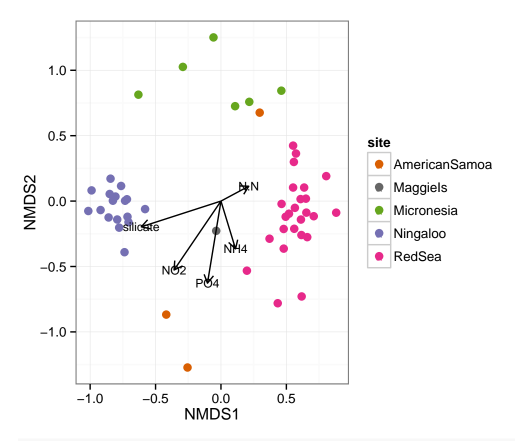
```
## Square root transformation
## Wisconsin double standardization
## Run 0 stress 0.189
## Run 1 stress 0.1729
## ... New best solution
## ... procrustes: rmse 0.04878 max resid 0.2672
## Run 2 stress 0.2191
## Run 3 stress 0.2
## Run 4 stress 0.2133
## Run 5 stress 0.2092
## Run 6 stress 0.2003
## Run 7 stress 0.1853
## Run 8 stress 0.1848
## Run 9 stress 0.2156
## Run 10 stress 0.2121
## Run 11 stress 0.1828
## Run 12 stress 0.2202
## Run 13 stress 0.2
## Run 14 stress 0.1931
## Run 15 stress 0.1848
## Run 16 stress 0.2178
## Run 17 stress 0.2018
## Run 18 stress 0.2141
## Run 19 stress 0.1885
## Run 20 stress 0.1941
##
## ***VECTORS
##
##
             MDS1
                    MDS2 r2 Pr(>r)
           0.760 -0.650 0.48 0.001 ***
## temp
## salinity -0.151 0.989 0.16 0.020 *
           -0.910 0.415 0.08 0.122
## Domg
## pH
           -0.533 0.846 0.22 0.004 **
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
## P values based on 999 permutations.
```



draw_envfit_ord(spistChem, nutrients)

```
## Square root transformation
## Wisconsin double standardization
## Run 0 stress 0.1927
## Run 1 stress 0.1979
## Run 2 stress 0.1848
## ... New best solution
## ... procrustes: rmse 0.0305 max resid 0.1973
## Run 3 stress 0.1837
## ... New best solution
## ... procrustes: rmse 0.01026 max resid 0.06491
## Run 4 stress 0.2098
## Run 5 stress 0.2
## Run 6 stress 0.2092
## Run 7 stress 0.1963
## Run 8 stress 0.1874
## Run 9 stress 0.1957
## Run 10 stress 0.2218
## Run 11 stress 0.2114
## Run 12 stress 0.1888
## Run 13 stress 0.1955
## Run 14 stress 0.2078
## Run 15 stress 0.1821
## ... New best solution
## ... procrustes: rmse 0.07277 max resid 0.4077
## Run 16 stress 0.2004
```

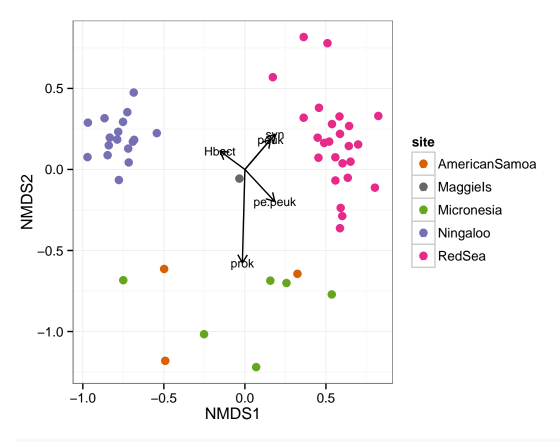
```
## Run 17 stress 0.1837
## Run 18 stress 0.2134
## Run 19 stress 0.2139
## Run 20 stress 0.1955
## ***VECTORS
##
##
              MDS1
                     MDS2
                            r2 Pr(>r)
## P04
            -0.162 -0.987 0.40 0.001 ***
## N.N
            0.882 0.472 0.06 0.197
## silicate -0.952 -0.307 0.41 0.001 ***
            -0.563 -0.827 0.41 0.001 ***
## NO2
## NH4
             0.292 -0.957 0.15 0.016 *
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
## P values based on 999 permutations.
```



draw_envfit_ord(spistChem, FCM)

```
## Square root transformation
## Wisconsin double standardization
## Run 0 stress 0.1927
## Run 1 stress 0.2191
## Run 2 stress 0.2193
## Run 3 stress 0.1971
## Run 4 stress 0.2141
## Run 5 stress 0.1848
```

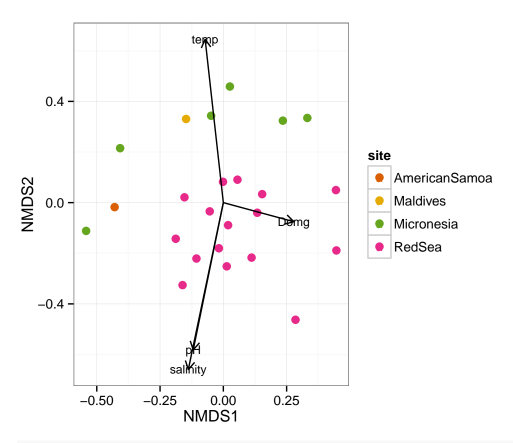
```
## ... New best solution
## ... procrustes: rmse 0.03064 max resid 0.1973
## Run 6 stress 0.2075
## Run 7 stress 0.2075
## Run 8 stress 0.1936
## Run 9 stress 0.1922
## Run 10 stress 0.2043
## Run 11 stress 0.216
## Run 12 stress 0.196
## Run 13 stress 0.206
## Run 14 stress 0.2108
## Run 15 stress 0.1943
## Run 16 stress 0.1949
## Run 17 stress 0.2147
## Run 18 stress 0.2017
## Run 19 stress 0.2089
## Run 20 stress 0.2179
##
## ***VECTORS
##
##
           MDS1 MDS2 r2 Pr(>r)
## prok
        -0.027 -1.000 0.33 0.002 **
## syn
          0.650 0.760 0.08 0.092 .
         0.650 0.760 0.06 0.214
## peuk
## pe.peuk 0.682 -0.731 0.07 0.163
## Hbact -0.805 0.593 0.04 0.426
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## P values based on 999 permutations.
```



draw_envfit_ord(pverrChem, waterQual)

```
## Square root transformation
## Wisconsin double standardization
## Run 0 stress 0.2599
## Run 1 stress 0.2639
## Run 2 stress 0.2739
## Run 3 stress 0.2583
## ... New best solution
## ... procrustes: rmse 0.1098 \max resid 0.2726
## Run 4 stress 0.263
## Run 5 stress 0.2699
## Run 6 stress 0.2583
## ... New best solution
## ... procrustes: rmse 0.1761 max resid 0.4596
## Run 7 stress 0.259
## Run 8 stress 0.2535
## ... New best solution
## ... procrustes: rmse 0.1793 max resid 0.436
## Run 9 stress 0.2579
## Run 10 stress 0.2545
## Run 11 stress 0.247
## ... New best solution
## ... procrustes: rmse 0.1713 max resid 0.384
## Run 12 stress 0.2708
## Run 13 stress 0.2762
## Run 14 stress 0.2536
```

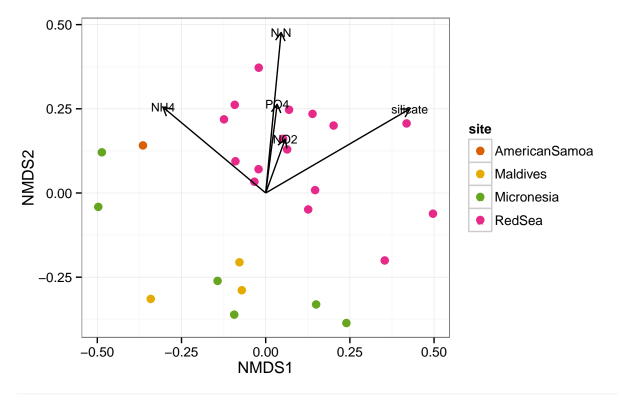
```
## Run 15 stress 0.2559
## Run 16 stress 0.2555
## Run 17 stress 0.263
## Run 18 stress 0.2692
## Run 19 stress 0.2582
## Run 20 stress 0.2581
## Warning: skipping half-change scaling: too few points below threshold
##
## ***VECTORS
##
##
             MDS1
                    MDS2 r2 Pr(>r)
            -0.111 0.994 0.42 0.006 **
## temp
## salinity -0.205 -0.979 0.45 0.007 **
## Domg
           0.965 -0.261 0.08 0.369
           -0.200 -0.980 0.35 0.010 **
## pH
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## P values based on 999 permutations.
```



draw_envfit_ord(pverrChem, nutrients)

```
## Square root transformation
## Wisconsin double standardization
## Run 0 stress 0.262
```

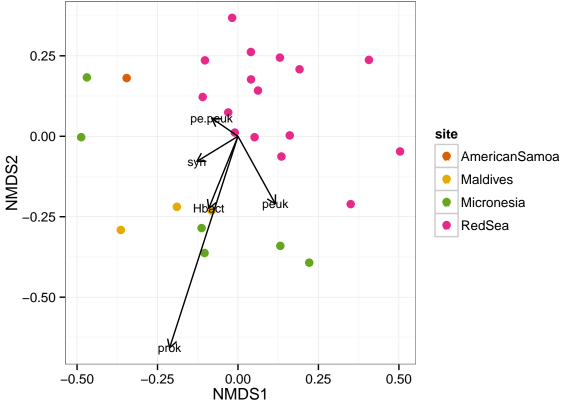
```
## Run 1 stress 0.2529
## ... New best solution
## ... procrustes: rmse 0.1795 max resid 0.3817
## Run 2 stress 0.2605
## Run 3 stress 0.276
## Run 4 stress 0.2585
## Run 5 stress 0.2703
## Run 6 stress 0.282
## Run 7 stress 0.2666
## Run 8 stress 0.2939
## Run 9 stress 0.2439
## ... New best solution
## ... procrustes: rmse 0.08096 max resid 0.2958
## Run 10 stress 0.2542
## Run 11 stress 0.2778
## Run 12 stress 0.2645
## Run 13 stress 0.2743
## Run 14 stress 0.2654
## Run 15 stress 0.2766
## Run 16 stress 0.2719
## Run 17 stress 0.2748
## Run 18 stress 0.2627
## Run 19 stress 0.2657
## Run 20 stress 0.2907
## Warning: skipping half-change scaling: too few points below threshold
##
## ***VECTORS
            MDS1 MDS2 r2 Pr(>r)
##
## P04
           0.128 0.992 0.07 0.437
        0.096 0.995 0.23 0.038 *
## N.N
## silicate 0.864 0.504 0.25 0.027 *
## NO2
       0.340 0.940 0.03 0.711
## NH4
          -0.768 0.641 0.16 0.135
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## P values based on 999 permutations.
```



draw_envfit_ord(pverrChem, FCM)

```
## Square root transformation
## Wisconsin double standardization
## Run 0 stress 0.262
## Run 1 stress 0.2789
## Run 2 stress 0.2602
## ... New best solution
## ... procrustes: rmse 0.1728 max resid 0.4041
## Run 3 stress 0.2598
## ... New best solution
## ... procrustes: rmse 0.15 max resid 0.4096
## Run 4 stress 0.2633
## Run 5 stress 0.2603
## Run 6 stress 0.2684
## Run 7 stress 0.2588
## ... New best solution
## ... procrustes: rmse 0.1693 max resid 0.3954
## Run 8 stress 0.2437
## ... New best solution
## ... procrustes: rmse 0.1479 max resid 0.3598
## Run 9 stress 0.244
## ... procrustes: rmse 0.01929 max resid 0.08613
## Run 10 stress 0.2873
## Run 11 stress 0.2627
## Run 12 stress 0.2963
## Run 13 stress 0.255
## Run 14 stress 0.2608
## Run 15 stress 0.2957
## Run 16 stress 0.2736
```

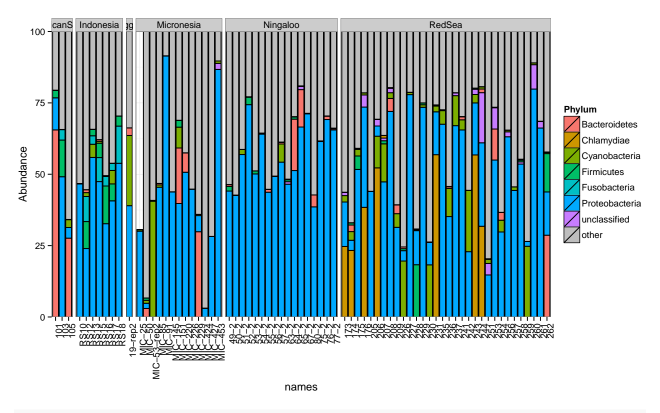
```
## Run 17 stress 0.2652
## Run 18 stress 0.2667
## Run 19 stress 0.2906
## Run 20 stress 0.2621
## Warning: skipping half-change scaling: too few points below threshold
##
## ***VECTORS
##
##
            MDS1
                   MDS2
                          r2 Pr(>r)
## prok
          -0.308 -0.952 0.48 0.001 ***
          -0.853 -0.522 0.02 0.785
## syn
           0.484 -0.875 0.06 0.511
## peuk
## pe.peuk -0.823 0.568 0.01 0.886
## Hbact
          -0.375 -0.927 0.06 0.492
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## P values based on 999 permutations.
```



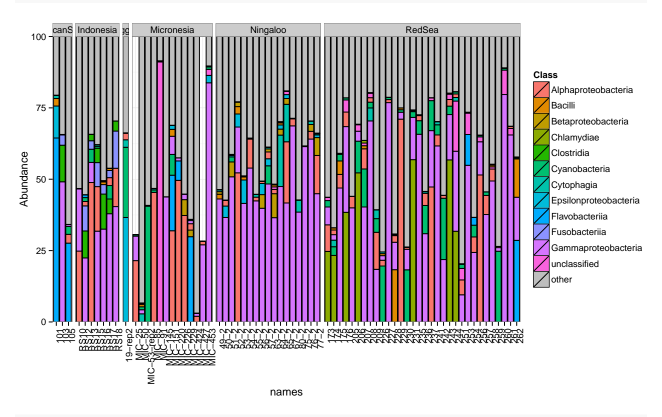
Taxonomic barcharts of bacteria in the corals and seawaters, and core microbiome members

```
# define a function to draw barcharts at a specific taxonomic level also need to # create my own ggplot colors then replace the last one ('other' column) with # gray
```

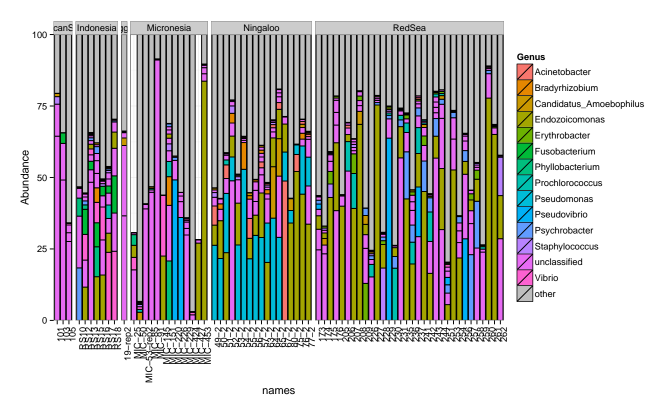
```
gg_color_hue <- function(n) {</pre>
    hues = seq(15, 375, length = n + 1)
    hcl(h = hues, l = 65, c = 100)[1:n]
}
draw_barcharts <- function(coral_species, tax_level) {</pre>
    coralFiltGlom <- tax glom(coral species, taxrank = tax level)</pre>
    physeqdf <- psmelt(coralFiltGlom)</pre>
    # get total abundance so can make an 'other' column had to add ^ and $ characters
    # to make sure grep matches whole word
    physeqdfOther <- physeqdf</pre>
    for (j in unique(physeqdf$Sample)) {
        jFirst = paste("^", j, sep = "")
        jBoth = paste(jFirst, "$", sep = "")
        rowNumbers = grep(jBoth, physeqdf$Sample)
        otherValue = 100 - sum(physeqdf[rowNumbers, "Abundance"])
        newRow = (physeqdf[rowNumbers, ])[1, ]
        newRow[, tax_level] = "other"
        newRow[, "Abundance"] = otherValue
        physeqdfOther <- rbind(physeqdfOther, newRow)</pre>
    }
    ggCols <- gg_color_hue(length(unique(physeqdf0ther[, tax_level])))</pre>
    ggCols \leftarrow head(ggCols, n = -1)
    physeqdf0ther$names <- factor(physeqdf0ther$Sample, levels = rownames(metaFile),</pre>
        ordered = TRUE)
    theme_set(theme_bw())
    ggplot(physeqdfOther, aes_string(x = "names", y = "Abundance", fill = tax_level,
        order = as.factor(tax_level))) + geom_bar(stat = "identity", colour = "black") +
        scale_fill_manual(values = c(ggCols, "gray")) + scale_y_continuous(expand = c(0,
        0), limits = c(0, 100)) + facet_grid(~site, scales = "free", space = "free_x") +
        theme(axis.text.x = element text(angle = 90, hjust = 1))
}
# subset coral samples, create names factor for label ordering and filter so the
# graph isn't too full
spist <- subset_samples(allPhylo, species == "Stylophora pistillata")</pre>
sample_data(spist)$names <- factor(sample_names(spist), levels = rownames(metaFile),</pre>
    ordered = TRUE)
spistFilt = filter_taxa(spist, function(x) mean(x) > 0.5, TRUE)
draw_barcharts(spistFilt, "Phylum")
```



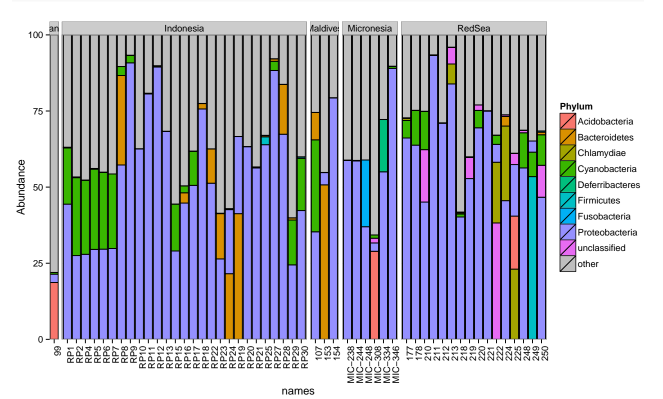
draw_barcharts(spistFilt, "Class")



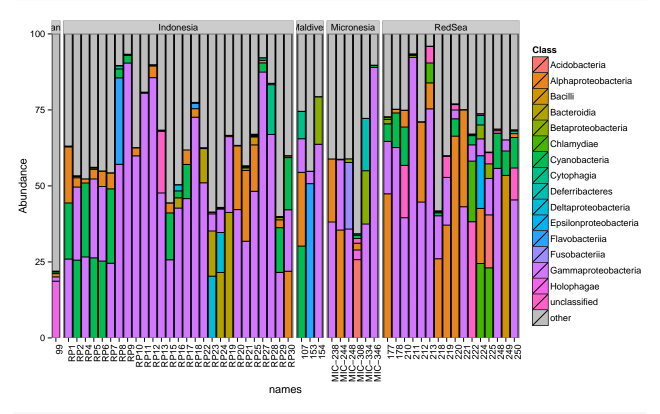
draw_barcharts(spistFilt, "Genus")



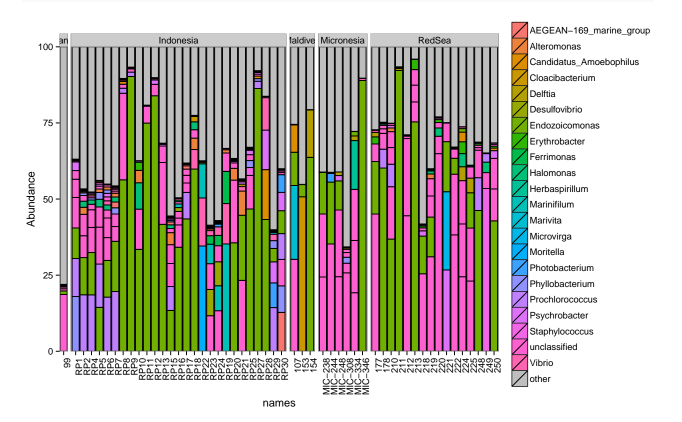
pVerr <- subset_samples(allPhylo, species == "Pocillopora verrucosa")
sample_data(pVerr)\$names <- factor(sample_names(pVerr), levels = unique(sample_names(pVerr)))
pVerrFilt = filter_taxa(pVerr, function(x) mean(x) > 0.3, TRUE)
draw_barcharts(pVerrFilt, "Phylum")

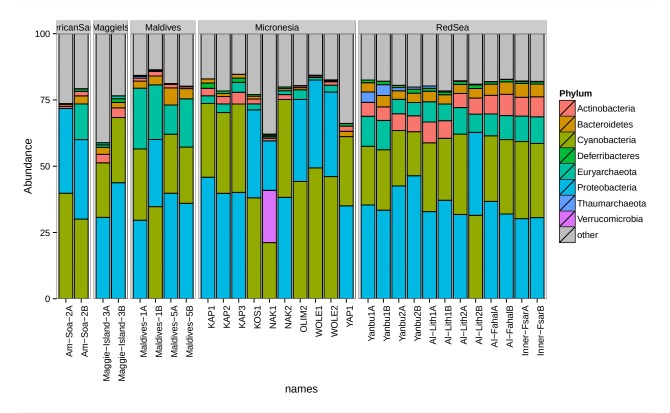


draw_barcharts(pVerrFilt, "Class")

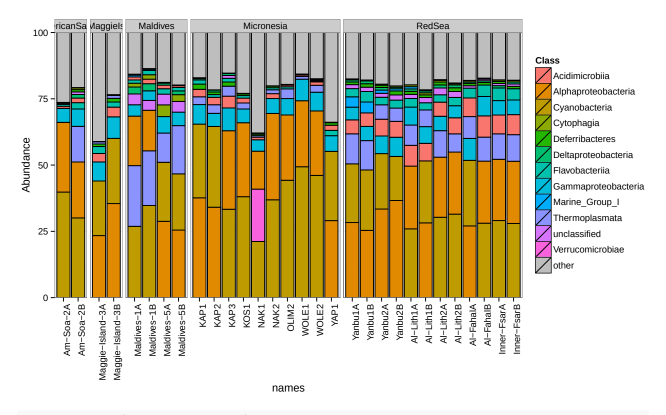


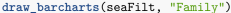
draw_barcharts(pVerrFilt, "Genus")

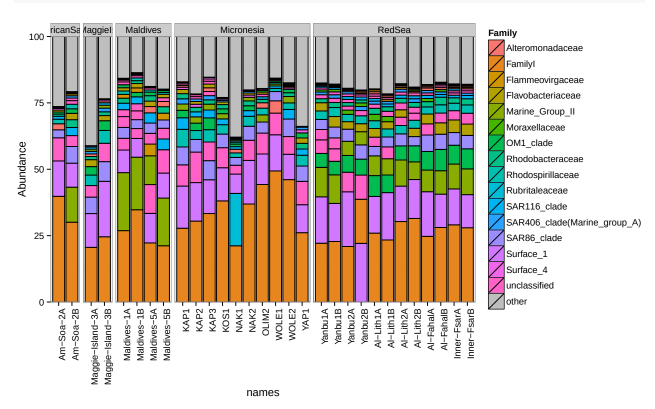




draw_barcharts(seaFilt, "Class")







The two corals are both dominated by Gammaproteobacteria at the higher taxonomic levels. At the genus level, there is more variability but Endozoicomonas seem to be fairly prevalent. Let's check which bacterial genera are most consistently associated with the corals and may be considered a 'core' microbiome member.

Core coral microbiome members

```
# check for 'core' microbiome members at the genus level which taxa are present
# at 1% overall abundance and at least 50% of samples in Stylophora pistillata?
spistGenusGlom <- tax_glom(spistFilt, taxrank = "Genus")</pre>
coreTaxa = filter_taxa(spistGenusGlom, function(x) sum(x > 1) > (0.5 * length(x)),
    TRUE)
tax_table(coreTaxa)
                       [1 taxa by 7 taxonomic ranks]:
## Taxonomy Table:
##
                Domain
                           Phylum
                                            Class
## MED000008661 "Bacteria" "Proteobacteria" "Gammaproteobacteria"
                Order
                                    Family
                                                   Genus
## MED000008661 "Oceanospirillales" "Hahellaceae" "Endozoicomonas" NA
sum(otu_table(coreTaxa) > 1)/nsamples(spist)
## [1] 0.7671
# which taxa are present at 1% overall abundance and at least 50% of samples in
# Pocillopora verrucosa?
pVerrGenusGlom <- tax_glom(pVerrFilt, taxrank = "Genus")</pre>
coreTaxa = filter_taxa(pVerrGenusGlom, function(x) sum(x > 1) > (0.5 * length(x)),
   TRUE)
tax table(coreTaxa)
## Taxonomy Table:
                       [1 taxa by 7 taxonomic ranks]:
                Domain
                           Phylum
## MED000008683 "Bacteria" "Proteobacteria" "Gammaproteobacteria"
                                    Family
                                                                    Species
## MED000008683 "Oceanospirillales" "Hahellaceae" "Endozoicomonas" NA
sum(otu_table(coreTaxa) > 1)/nsamples(pVerr)
```

[1] 0.8491

Indeed Endozoicomonas were the most prevalent bacteria in the corals and were they only bacterial genera to occur in more than 50% of the colonies sampled. In fact, for both coral species Endozoicomonas occurred in more than 75% of colonies.

Let's check if these Endozoicomonas OTUs show any patterns across the coral species or at different geographic areas.

Heatmap of different Endozoicomonas MED OTUs across the coral species

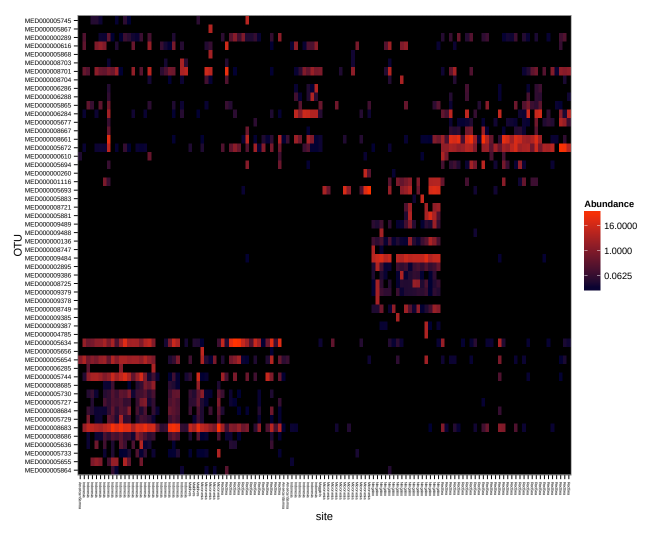
```
allPhyloEndo <- subset_taxa(allPhylo, Genus == "Endozoicomonas")
spistPhyloEndo <- subset_samples(allPhyloEndo, species == "Stylophora pistillata")
pVerrPhyloEndo <- subset_samples(allPhyloEndo, species == "Pocillopora verrucosa")
```

```
spistPverrEndo <- merge_phyloseq(spistPhyloEndo, pVerrPhyloEndo)

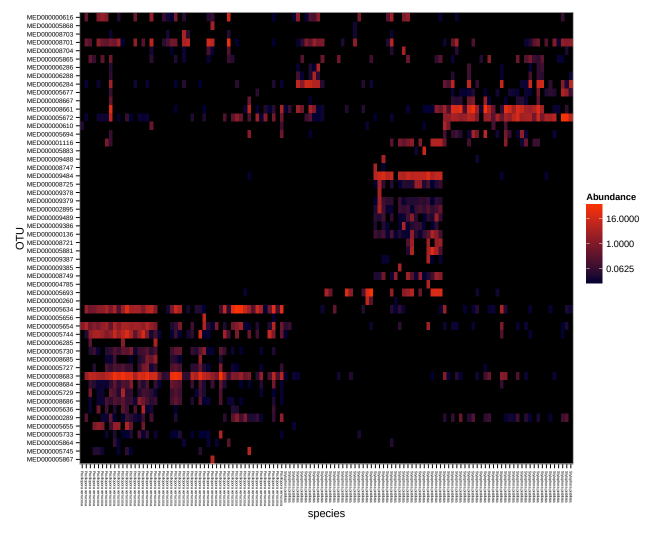
spistPverrEndoFilt = filter_taxa(spistPverrEndo, function(x) mean(x) > 0, TRUE)

spistPverrEndoFiltPrune = prune_samples(sample_sums(spistPverrEndoFilt) > 0, spistPverrEndoFilt)

plot_heatmap(spistPverrEndoFiltPrune, "NMDS", "bray", "site", low = "#000033", high = "#FF3300", sample.order = rownames(metaFile3))
```



```
plot_heatmap(spistPverrEndoFiltPrune, "NMDS", "bray", "species", low = "#000033",
    high = "#FF3300", sample.order = rownames(metaFile3))
```



Pretty cool. Looks like the two corals have different Endozoicomonas types, and the types also seem to partition differently across sites for the coral species, i.e., Pocillopora verrucosa has similar Endozoicomonas types across large geographic areas but Stylophora pistillata seems to have different Endozoicomonas types at each area. Let's do some significance testing to see if more of the Endozoicomonas OTUs are different across sites for S. pistillata compared to P. verrucosa.

DESeq2 significance testing for Endozoicomonas MED OTUs

```
# subset endozoicomonas OTUs from the count data as required by DESeq2
countEndos <- subset_taxa(countPhylo, Genus=='Endozoicomonas')
spistCountEndo <- subset_samples(countEndos, species=='Stylophora pistillata')
pVerrCountEndo <- subset_samples(countEndos, species=='Pocillopora verrucosa')
coralCountEndo <- merge_phyloseq(spistCountEndo, pVerrCountEndo)

# do some filtering for Os
spistCountEndoFilt = filter_taxa(spistCountEndo, function(x) mean(x) > 0.0, TRUE)
spistCountEndoFiltPrune = prune_samples(sample_sums(spistCountEndoFilt) > 0, spistCountEndoFilt)
pVerrCountEndoFilt = filter_taxa(pVerrCountEndo, function(x) mean(x) > 0.0, TRUE)
pVerrCountEndoFiltPrune = prune_samples(sample_sums(pVerrCountEndoFilt) > 0, pVerrCountEndoFilt)
```

```
# convert phyloseq object to DESeq object
spistDeseq <- phyloseq_to_deseq2(spistCountEndoFiltPrune, ~ site)</pre>
pverrDeseq <- phyloseq to deseq2(pVerrCountEndoFiltPrune, ~ site)</pre>
# need to calculate geometric means separately because there are zeros in the data
gm_mean = function(x, na.rm=TRUE){
  exp(sum(log(x[x > 0]), na.rm=na.rm) / length(x))
spistMeans <- apply(counts(spistDeseq), 1, gm_mean)</pre>
spistDeseq <- estimateSizeFactors(spistDeseq, geoMeans = spistMeans)</pre>
pverrMeans <- apply(counts(pverrDeseq), 1, gm_mean)</pre>
pverrDeseq <- estimateSizeFactors(pverrDeseq, geoMeans = pverrMeans)</pre>
# now can run the DESeq tests
spistDeseq <- DESeq(spistDeseq, fitType="local")</pre>
## using pre-existing size factors
## estimating dispersions
## gene-wise dispersion estimates
## mean-dispersion relationship
## final dispersion estimates
## fitting model and testing
## -- replacing outliers and refitting for 38 genes
## -- DESeq argument 'minReplicatesForReplace' = 7
## -- original counts are preserved in counts(dds))
## estimating dispersions
## fitting model and testing
pverrDeseq <- DESeq(pverrDeseq, fitType="local")</pre>
## using pre-existing size factors
## estimating dispersions
## gene-wise dispersion estimates
## mean-dispersion relationship
## final dispersion estimates
## fitting model and testing
## -- replacing outliers and refitting for 29 genes
## -- DESeq argument 'minReplicatesForReplace' = 7
## -- original counts are preserved in counts(dds))
## estimating dispersions
## fitting model and testing
# create a function to check the results for each of the site comparisons
# output significant p-value (FDR adjusted) ordered OTUs
check site results <- function(coral deseq, site comparison){</pre>
 res <- results(coral_deseq, contrast = site_comparison)</pre>
 res = res[order(res$padj, na.last=NA), ]
 sigtab = as.data.frame(res[(res$padj < 0.05), ])</pre>
 print(sigtab)
}
```

```
# Stylophora pistillata
check_site_results(spistDeseq, c("site", "AmericanSamoa", "RedSea"))
                baseMean log2FoldChange lfcSE
                                               stat
                                                        pvalue
                                                                    padj
## MED00000289
                 0.4453
                                -6.767 1.515 -4.466 7.983e-06 0.0002954
## MED00005672
                45.9003
                                -8.229 1.987 -4.140 3.468e-05 0.0006417
                                -5.288 1.521 -3.478 5.060e-04 0.0062405
## MED00005677
                 0.2435
## MED00005694
                 0.8396
                                -5.769 1.825 -3.160 1.576e-03 0.0145776
                                -5.025 1.663 -3.021 2.518e-03 0.0186321
## MED000008667
                 0.2736
## MED000008661 23.8289
                                -4.508 1.696 -2.658 7.855e-03 0.0484376
check_site_results(spistDeseq, c("site", "AmericanSamoa", "Ningaloo"))
                baseMean log2FoldChange lfcSE
                                                        pvalue
                                                stat
## MED000002895 0.33262
                                -7.194 1.520 -4.732 2.227e-06 6.905e-05
## MED000009484 24.74728
                                -9.485 2.124 -4.466 7.965e-06 8.231e-05
## MED000009379 0.22244
                                -6.706 1.493 -4.490 7.119e-06 8.231e-05
## MED000008725
                0.30197
                                -6.708 1.614 -4.157 3.230e-05 2.503e-04
                                -6.297 1.539 -4.090 4.309e-05 2.671e-04
## MED000009489
                0.17044
## MED000008749 0.58961
                                -6.953 1.738 -4.000 6.340e-05 3.275e-04
                                -6.343 1.618 -3.921 8.816e-05 3.904e-04
## MED000000136 0.31981
## MED000009386 0.09981
                                -5.449 1.566 -3.480 5.006e-04 1.940e-03
                                -5.531 1.915 -2.888 3.879e-03 1.336e-02
## MED000005881
                0.46854
## MED000008721 0.10896
                                -4.545 1.797 -2.529 1.142e-02 3.541e-02
## MED000005693 1.79057
                                -5.378 2.162 -2.487 1.287e-02 3.628e-02
check_site_results(spistDeseq, c("site", "AmericanSamoa", "Micronesia"))
## [1] baseMean
                     log2FoldChange lfcSE
                                                    stat
## [5] pvalue
                     padi
## <0 rows> (or 0-length row.names)
check_site_results(spistDeseq, c("site", "AmericanSamoa", "Indonesia"))
## [1] baseMean
                     log2FoldChange lfcSE
                                                    stat
## [5] pvalue
                     padj
## <0 rows> (or 0-length row.names)
check_site_results(spistDeseq, c("site", "RedSea", "Ningaloo"))
                baseMean log2FoldChange lfcSE
                                                stat
                                                         pvalue
## MED000005672 45.90025
                                11.032 1.0441 10.566 4.273e-26 1.795e-24
## MED000009484 24.74728
                               -11.528 1.3365 -8.625 6.399e-18 1.344e-16
## MED000008661 23.82895
                                 6.504 0.8044 8.085 6.212e-16 8.696e-15
## MED000006284
                                 9.652 1.6475
                                              5.858 4.671e-09 4.905e-08
                4.66332
                                8.958 1.8933 4.731 2.231e-06 1.561e-05
## MED000005694
                0.83957
## MED00000289
                                 9.426 1.9822 4.755 1.982e-06 1.561e-05
                0.44531
## MED000005693
               1.79057
                                -6.795 1.4752 -4.606 4.102e-06 2.154e-05
                                -9.394 2.0396 -4.606 4.103e-06 2.154e-05
## MED000002895 0.33262
                                -8.944 1.9658 -4.550 5.368e-06 2.255e-05
## MED000000136 0.31981
```

```
## MED000008749 0.58961
                                 -9.471 2.0788 -4.556 5.213e-06 2.255e-05
                                 8.322 1.8948 4.392 1.124e-05 4.293e-05
## MED000005677
                0.24346
## MED000008725
                0.30197
                                 -9.002 2.0704 -4.348 1.373e-05 4.805e-05
                                 -8.767 2.0752 -4.224 2.396e-05 7.739e-05
## MED000009379
                0.22244
## MED000008667
                0.27364
                                  7.956 1.9383 4.105 4.047e-05 1.214e-04
                                 -8.376 2.1012 -3.986 6.715e-05 1.880e-04
## MED000009489
                0.17044
                                 4.573 1.1844 3.861 1.128e-04 2.961e-04
## MED000008701
                4.18358
                0.18381
                                 6.537 1.9231 3.399 6.762e-04 1.590e-03
## MED000006288
## MED000009386
                0.09981
                                 -7.387 2.1746 -3.397 6.816e-04 1.590e-03
## MED00005881
                0.46854
                                 -7.859 2.3362 -3.364 7.689e-04 1.700e-03
## MED000008683
                0.32075
                                 4.732 1.4268 3.316 9.126e-04 1.916e-03
                                 -5.296 1.6137 -3.282 1.031e-03 2.061e-03
## MED00000616
                0.94550
## MED000008721
                0.10896
                                 -6.484 2.3768 -2.728 6.373e-03 1.217e-02
## MED000006286 0.07270
                                 5.374 2.0396 2.635 8.416e-03 1.537e-02
check_site_results(spistDeseq, c("site", "RedSea", "Micronesia"))
                baseMean log2FoldChange lfcSE stat
                                                        pvalue
                                                                    padj
## MED000008661
                23.8289
                                9.196 1.0638 8.644 5.405e-18 2.270e-16
## MED00005672
                45.9003
                                  6.468 0.9063 7.137 9.569e-13 2.009e-11
## MED00000289
                 0.4453
                                  9.307 2.0013 4.650 3.312e-06 4.637e-05
## MED00005694
                 0.8396
                                 7.736 1.8558 4.168 3.069e-05 3.222e-04
## MED000006284
                 4.6633
                                  5.816 1.4129 4.116 3.849e-05 3.233e-04
## MED000005677
                 0.2435
                                  5.543 1.4749 3.758 1.714e-04 1.028e-03
## MED000008667
                 0.2736
                                  7.529 1.9983 3.768 1.649e-04 1.028e-03
## MED000001116
                 1.2828
                                  6.022 1.8659 3.228 1.248e-03 6.555e-03
## MED000008701
                                  4.117 1.3038 3.158 1.589e-03 7.417e-03
                  4.1836
## MED00006288
                  0.1838
                                  6.060 1.9856 3.052 2.272e-03 9.541e-03
check_site_results(spistDeseq, c("site", "RedSea", "Indonesia"))
##
                baseMean log2FoldChange lfcSE stat
                                                       pvalue
                                                                   padj
## MED00005672
                45.9003
                                 6.244 1.084 5.761 8.368e-09 2.092e-07
## MED00000289
                  0.4453
                                 7.879 2.219 3.551 3.834e-04 4.793e-03
## MED00005677
                  0.2435
                                 6.634 2.146 3.092 1.990e-03 1.658e-02
## MED00005694
                 0.8396
                                  4.940 1.766 2.797 5.163e-03 3.227e-02
## MED000001116
                  1.2828
                                  5.369 2.028 2.647 8.115e-03 4.058e-02
                                  4.922 1.936 2.542 1.102e-02 4.591e-02
## MED000005634
                  0.4262
check_site_results(spistDeseq, c("site", "Ningaloo", "Micronesia"))
                                                        pvalue
                baseMean log2FoldChange lfcSE
                                                stat
                                                                    padj
## MED000009484 24.74728
                                 10.236 1.473
                                              6.949 3.672e-12 1.359e-10
                0.33262
                                 10.466 1.870
                                              5.596 2.193e-08 4.056e-07
## MED000002895
## MED000009379
                0.22244
                                 9.810 1.906
                                              5.148 2.632e-07 3.246e-06
## MED00000136
                0.31981
                                 7.785 1.580
                                              4.926 8.387e-07 7.758e-06
## MED000008725
                                 9.297 2.025
                                              4.592 4.388e-06 3.247e-05
                0.30197
                                 8.964 2.006 4.469 7.867e-06 4.851e-05
## MED000009489
                0.17044
## MED000008749
                                 9.274 2.107 4.401 1.076e-05 5.685e-05
                0.58961
## MED000005672 45.90025
                                 -4.565 1.209 -3.775 1.603e-04 6.699e-04
## MED000001116
                                 7.406 1.964 3.770 1.630e-04 6.699e-04
                1.28283
## MED000000616 0.94550
                                 7.190 2.040 3.524 4.253e-04 1.431e-03
```

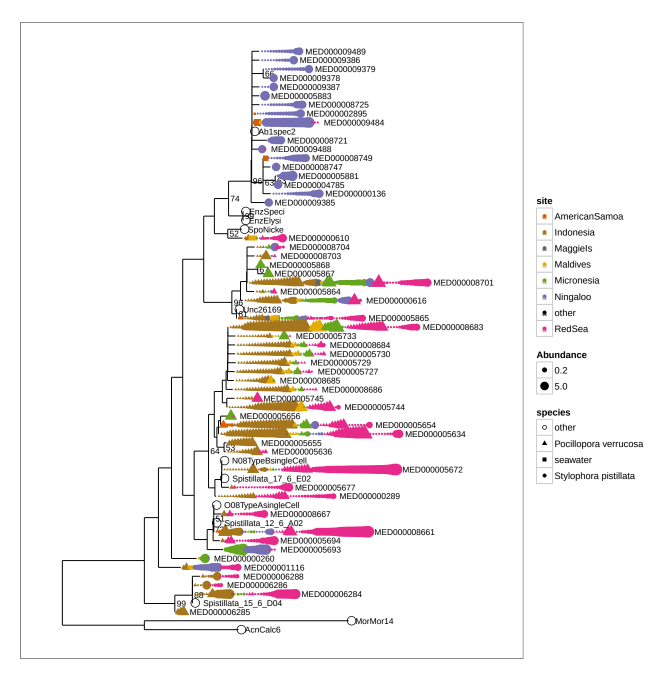
```
## MED000009386 0.09981
                                7.593 2.142 3.545 3.925e-04 1.431e-03
## MED000005881 0.46854
                                7.028 2.416 2.909 3.623e-03 1.117e-02
                                5.866 2.434 2.410 1.596e-02 4.543e-02
## MED000008721 0.10896
check_site_results(spistDeseq, c("site", "Ningaloo", "Indonesia"))
               baseMean log2FoldChange lfcSE stat
                                                    pvalue
                                                                  padj
## MED000009484 24.74728
                             12.009 1.936 6.203 5.529e-10 1.493e-08
## MED000006284 4.66332
                                -9.180 1.960 -4.683 2.822e-06 3.810e-05
## MED000008661 23.82895
                               -4.134 1.134 -3.645 2.674e-04 1.031e-03
## MED000005693 1.79057
                                7.649 2.050 3.731 1.905e-04 1.031e-03
                                7.841 2.134 3.675 2.380e-04 1.031e-03
## MED00000136 0.31981
## MED000008749 0.58961
                                8.285 2.246 3.689 2.250e-04 1.031e-03
## MED000002895 0.33262
                                8.414 2.192 3.839 1.237e-04 1.031e-03
## MED000005672 45.90025
                               -4.788 1.341 -3.570 3.576e-04 1.073e-03
                                7.950 2.227 3.570 3.573e-04 1.073e-03
## MED000008725 0.30197
## MED000009379
                                7.823 2.215 3.532 4.130e-04 1.115e-03
               0.22244
## MED000009489 0.17044
                                7.405 2.241 3.305 9.495e-04 2.330e-03
## MED000001116 1.28283
                                6.753 2.109 3.201 1.368e-03 3.079e-03
## MED000006288 0.18381
                               -6.595 2.160 -3.054 2.260e-03 4.694e-03
## MED000009386 0.09981
                                6.434 2.293 2.806 5.016e-03 9.675e-03
## MED000005881 0.46854
                                6.451 2.442 2.642 8.245e-03 1.484e-02
## MED000008701 4.18358
                               -4.162 1.609 -2.586 9.710e-03 1.639e-02
## MED000006286 0.07270
                               -5.611 2.284 -2.457 1.401e-02 2.225e-02
## MED000008721 0.10896
                                5.337 2.439 2.188 2.864e-02 4.297e-02
check_site_results(spistDeseq, c("site", "Micronesia", "Indonesia"))
               baseMean log2FoldChange lfcSE stat
                                                      pvalue
## MED000008661 23.8289
                               -6.826 1.323 -5.161 2.463e-07 2.709e-06
## MED00006284
                4.6633
                                -5.344 1.793 -2.981 2.875e-03 1.581e-02
## MED00000616
                 0.9455
                               -6.324 2.259 -2.799 5.126e-03 1.879e-02
# Pocillopora verrucosa
check_site_results(pverrDeseq, c("site", "Maldives", "RedSea"))
               baseMean log2FoldChange lfcSE stat
                                                      pvalue
                               -9.037 1.267 -7.132 9.904e-13 1.783e-11
## MED000005634 255.716
## MED000000289
                  1.384
                                -5.780 1.929 -2.995 2.740e-03 2.466e-02
                                -4.019 1.514 -2.654 7.949e-03 4.769e-02
## MED000005672
                  3.650
check_site_results(pverrDeseq, c("site", "Maldives", "Micronesia"))
## [1] baseMean
                     log2FoldChange lfcSE
                                                   stat
## [5] pvalue
                     padj
## <0 rows> (or 0-length row.names)
check_site_results(pverrDeseq, c("site", "Maldives", "Indonesia"))
               baseMean log2FoldChange lfcSE stat
                                                      pvalue
## MED000005634
                  255.7
                                -5.488 1.231 -4.459 8.238e-06 0.0001977
```

```
check_site_results(pverrDeseq, c("site", "RedSea", "Micronesia"))
##
                baseMean log2FoldChange lfcSE
                                                stat
                                                        pvalue
                                                                    padj
## MED00005634
                255.716
                                10.107 1.1443 8.832 1.028e-18 1.542e-17
## MED00005672
                  3.650
                                 4.675 1.3549 3.450 5.603e-04 3.279e-03
## MED00000289
                  1.384
                                 6.330 1.8576 3.407 6.557e-04 3.279e-03
## MED000008683 359.232
                                -2.339 0.9037 -2.588 9.660e-03 3.623e-02
check_site_results(pverrDeseq, c("site", "RedSea", "Indonesia"))
##
                baseMean log2FoldChange lfcSE
                                                stat
                                                        pvalue
                                                                    padj
## MED00005672
                  3.650
                                 5.503 0.8747 6.291 3.157e-10 7.578e-09
## MED000005634 255.716
                                 3.549 0.6308 5.626 1.846e-08 2.215e-07
## MED000008683 359.232
                                -2.506 0.5847 -4.286 1.817e-05 1.453e-04
## MED000008686
                                -2.844 0.7519 -3.782 1.557e-04 9.344e-04
                  1.164
## MED00005655
                  3.205
                                -7.518 2.0199 -3.722 1.975e-04 9.482e-04
## MED000005729
                  1.074
                                -3.505 0.9725 -3.604 3.130e-04 1.252e-03
## MED000008684
                  1.428
                                -2.267 0.7219 -3.141 1.685e-03 5.779e-03
## MED00000289
                                 2.103 0.8030 2.618 8.833e-03 2.650e-02
                  1.384
check_site_results(pverrDeseq, c("site", "Micronesia", "Indonesia"))
                baseMean log2FoldChange lfcSE
                                                                   padj
                                               stat
                                                       pvalue
                                -6.558 1.103 -5.947 2.730e-09 4.914e-08
## MED000005634 255.716
## MED000005655
                   3.205
                                -6.374 2.164 -2.945 3.225e-03 2.902e-02
```

There is quite a few more significant differences across sites for Stylophora pistillata (90) compared to Pocillopora verrucosa (18), supporting what is shown in the heatmaps. I'll add an asterisk next to each of the significantly different OTUs in the heatmap to visually display these results.

Endozoicomonas seem to be displaying strain-specific relationships with the corals and across sites. I'll do a phylogenetic analysis of the Endozoicomonas sequences to further explore these relationships. The sequences were aligned using the SINA web service and imported into ARB for manual refinement, before being exported for use here.

Endozoicomonas phylogenetic tree with meta-data



Some interesting things coming up here. Several abundant but phylogenetically diverse Endozoicomonas OTUs appear to co-inhabit the same coral colonies. There are also clear host and site groupings of Endozoicomonas strains. I also included single cell 16S sequences in the tree and they are identical to the abundant MED OTUs from the Red Sea, suggesting that the MED procedure produces biologically relevant OTUs. Also in the tree are sequences from an earlier study of Red Sea Stylophora pistillata, named 'Spistillata_17_6_E02, Spistillata_12_6_A02, Spistillata_15_6_D04', and these are also the same as the abundant MED nodes in this study, suggesting that the most abundant MED OTUs in Red Sea S. pistillata have not changed for ~ 4 years.