# Mothur

Module 3: Project 1 by Team 5

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#### Abstract

This is the abstract. It consists of two paragraphs.

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#### 1 Introduction

#### 2 Problem Formulation

## 3 Materials and Experimental Configuration

#### 3.1 Experimental Protocols

Here . . . :

- **P1.** Analysis of microbial community structure along with depth and oxygen concentration (see Section 4.1).
- **P2.** Analysis of abundance information of [OTU\*\*\*\*] along with depth and/or oxygen concentration (see Section 4.2).
- P3. Estimate richness (number of OTUs/ASVs) for [OTU\*\*\*\*] (see Section 4.3).
- **P4.** Interpretation of abundance information of OTUs/ASVs of [OTU\*\*\*\*] along with depth and/or oxygen concentration (see Section 4.4).

#### 3.2 Dataset

#### 3.3 Parameters Configuration

```
## try http:// if https:// URLs are not supported
source("https://bioconductor.org/biocLite.R")
biocLite("phyloseq")
library("tidyverse")
library("gridExtra")
library("magrittr")
```

#### 3.4 Data Preporcessing

We use saanich inlet datasets that are propossed using mothur and QIIME2

```
load("data/mothur_phyloseq.RData")
load("data/qiime2_phyloseq.RData")
```

Samples are then rarefied/normalized to 100,000 sequences per sample to facilitate comparisons between samples. A random seed was set to ensure reproducibility.

```
set.seed(4832)
rarefied = rarefy_even_depth(mothur, sample.size = 1e+05)
```

Rarefied counts were converted to relative abundance percentages.

```
rarefiedPer = transform_sample_counts(rarefied, function(x) 100 * x/sum(x))
```

Next, we perform a series of filterings according to three rules: i)- exclude OTUs that are not observed for more than 4 samples; ii)- prune samples and OTUs with unknown values, such as unclassified value; and iii)- any phylum fail to have more than 5 OTUs should be trimmed. The codes used for applying the three rules are:

#### 4 Results

# 4.1 Analysis of microbial community structure along with depth and oxygen concentration

We first estimate the overall taxa diversity using Shannon's diversity index.

```
rarefiedRich <- estimate_richness(rarefied, measures = "Shannon")
rarefiedRichAlpha <- full_join(rownames_to_column(rarefiedRich), rownames_to_column(data.frame by = "rowname")

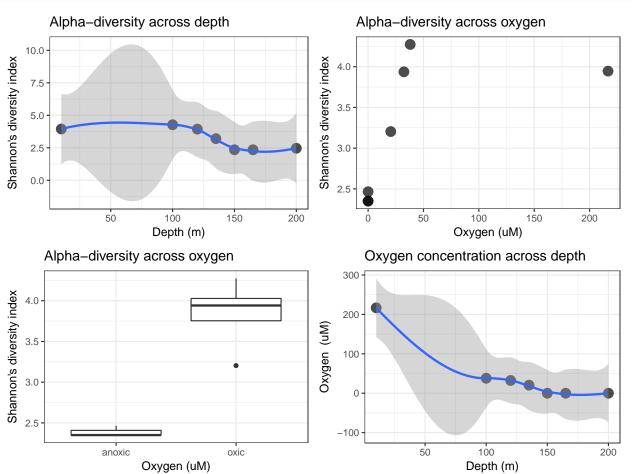
p1 <- rarefiedRichAlpha %>% ggplot() + geom_point(aes(x = Depth_m, y = Shannon), size = 4, alpha = 0.7) + geom_smooth(method = "loess", aes(x = as.numeric(Depth_m), y = Shannon)) + labs(title = "Alpha-diversity across depth", y = "Shannon's diversity index x = "Depth (m)")

p2 <- rarefiedRichAlpha %>% ggplot() + geom_point(aes(x = O2_uM, y = Shannon), size = 4, alpha = 0.7) + labs(title = "Alpha-diversity across oxygen", y = "Shannon's divers x = "Oxygen (uM)")

p3 <- rarefiedRichAlpha %>% mutate(O2_group = ifelse(O2_uM == 0, "anoxic", "oxic")) %>% ggplot() + geom_boxplot(aes(x = O2_group, y = Shannon)) + labs(title = "Alpha-diversity across oxygen")

p4 <- rarefiedRichAlpha %>% ggplot() + geom_point(aes(x = Depth_m, y = O2_uM),
```

```
size = 4, alpha = 0.7) + geom_smooth(method = "loess", aes(x = as.numeric(Depth_m),
    y = 02_uM)) + labs(title = "Oxygen concentration across depth", y = "Oxygen (uM)",
    x = "Depth (m)")
grid.arrange(p1, p2, p3, p4, ncol = 2)
```



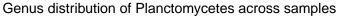
The following function will assist us to understand the unique Phylum rank:

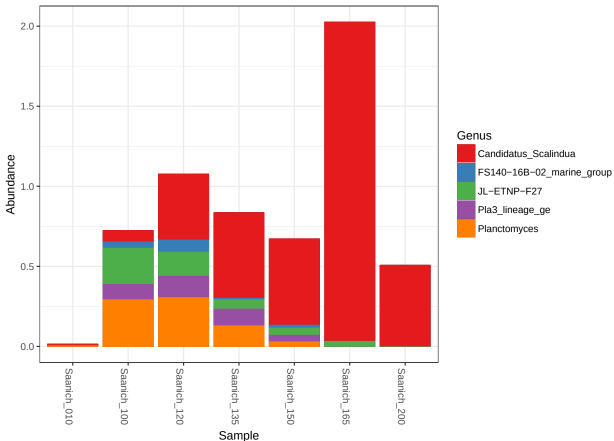
```
get_taxa_unique(physeq = thirdTaxa, taxonomic.rank = "Phylum")
```

```
## [1] "Proteobacteria" "Bacteroidetes"
## [3] "Thaumarchaeota" "Actinobacteria"
## [5] "Marinimicrobia_(SAR406_clade)" "Planctomycetes"
## [7] "Verrucomicrobia"
```

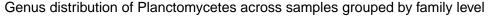
We choose the *Planctomycetes* phylum, and explored the distribution of genera of this phylum.

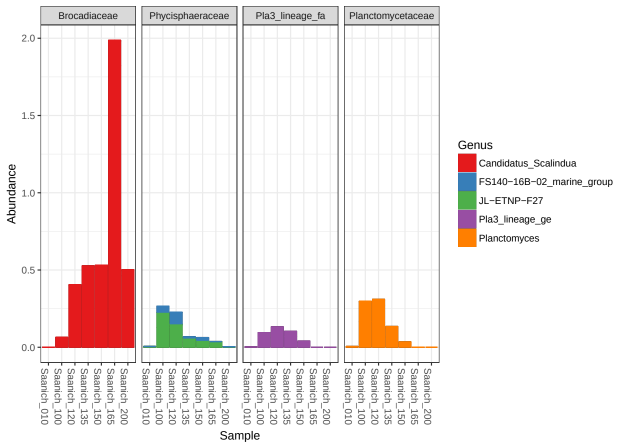
```
subTaxa <- subset_taxa(thirdTaxa, Phylum == "Planctomycetes")
plot_bar(subTaxa, fill = "Genus") + geom_bar(aes(color = Genus, fill = Genus),
    stat = "identity", position = "stack") + labs(title = "Genus distribution of Planctomycetes")</pre>
```





We further investigated the genus distribution of this phylum across samples grouped by family level.





Finally, we settled on performing experimental analysis at *Planctomyces* genus level and it's associated OTUs.

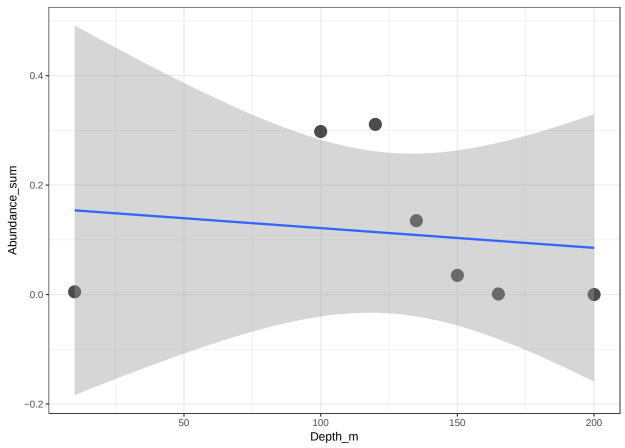
```
workingTaxa = subset_taxa(thirdTaxa, Genus == "Planctomyces")
(suggestedOTUs <- colnames(otu_table(workingTaxa)))</pre>
```

## [1] "Otu0125" "Otu0144" "Otu0401" "Otu0592"

# 4.2 Analysis of abundance information of $[OTU^{****}]$ along with depth and/or oxygen concentration

```
##
## Coefficients:
                 Estimate Std. Error t value Pr(>|t|)
##
## (Intercept) 0.1575152 0.1406836
                                       1.120
                                                0.314
## Depth m
               -0.0003609
                          0.0010227 -0.353
                                                0.739
## Residual standard error: 0.1511 on 5 degrees of freedom
## Multiple R-squared: 0.0243, Adjusted R-squared: -0.1708
## F-statistic: 0.1245 on 1 and 5 DF, p-value: 0.7386
workingTaxa %>% psmelt() %>% group_by(Sample) %>% summarize(Abundance_sum = sum(Abundance),
   Depth_m = mean(Depth_m)) %>% ggplot() + geom_point(aes(x = Depth_m, y = Abundance_sum),
    size = 5, alpha = 0.7) + geom_smooth(method = "lm", aes(x = as.numeric(Depth_m),
   y = Abundance_sum)) + labs(title = "Abundance of Plantomyces across depth")
```

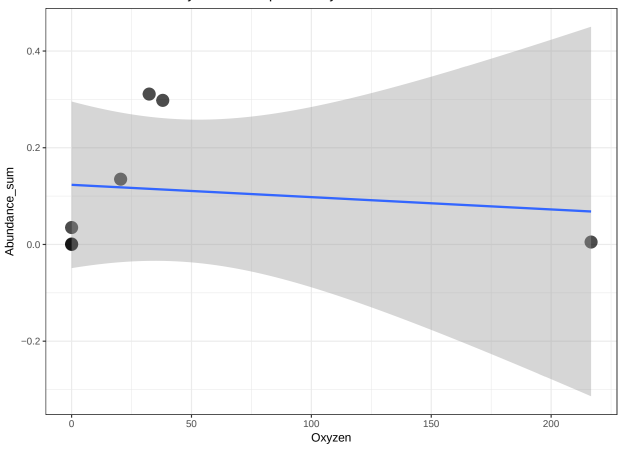
### Abundance of Plantomyces across depth



```
##
## Call:
## lm(formula = Abundance ~ 02_uM, data = .)
##
```

```
## Residuals:
##
          3
                   2
                                      5
   0.19591 \quad 0.18435 \quad 0.01688 \ -0.08832 \ -0.06319 \ -0.12232 \ -0.12332
##
##
## Coefficients:
                 Estimate Std. Error t value Pr(>|t|)
## (Intercept) 0.1233192 0.0670191
                                         1.84
## 02_uM
               -0.0002544 0.0007941
                                        -0.32
                                                 0.762
## Residual standard error: 0.1514 on 5 degrees of freedom
## Multiple R-squared: 0.02012,
                                     Adjusted R-squared: -0.1759
## F-statistic: 0.1027 on 1 and 5 DF, p-value: 0.7616
workingTaxa %>% psmelt() %>% group_by(Sample) %>% summarize(Abundance_sum = sum(Abundance),
    Oxyzen = mean(O2_uM)) %>% ggplot() + geom_point(aes(x = Oxyzen, y = Abundance_sum),
    size = 5, alpha = 0.7) + geom_smooth(method = "lm", aes(x = as.numeric(Oxyzen),
    y = Abundance sum)) + labs(title = "Abundance of Plantomyces with respect to oxygen concen
```

#### Abundance of Plantomyces with respect to oxyzen concentration

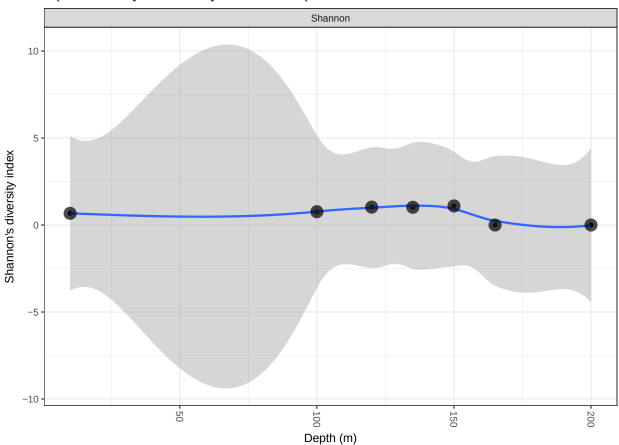


### 4.3 Estimate richness (number of OTUs/ASVs) for [OTU\*\*\*\*]

We explore the diversity of *Planctomyces* across depth.

```
workingTaxa %>% plot_richness(x = "Depth_m", measures = "Shannon") + geom_smooth(method = "loe
    aes(x = as.numeric(Depth_m))) + geom_point(size = 5, alpha = 0.7) + labs(title = "Alpha-di
    y = "Shannon's diversity index", x = "Depth (m)")
```

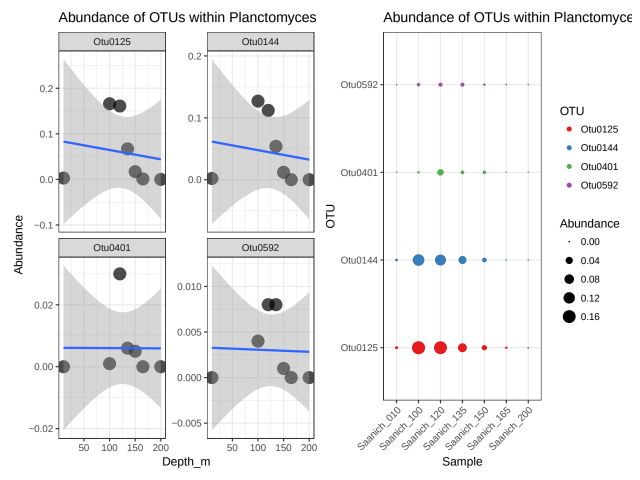
#### Alpha-diversity of Plantomyces across depth



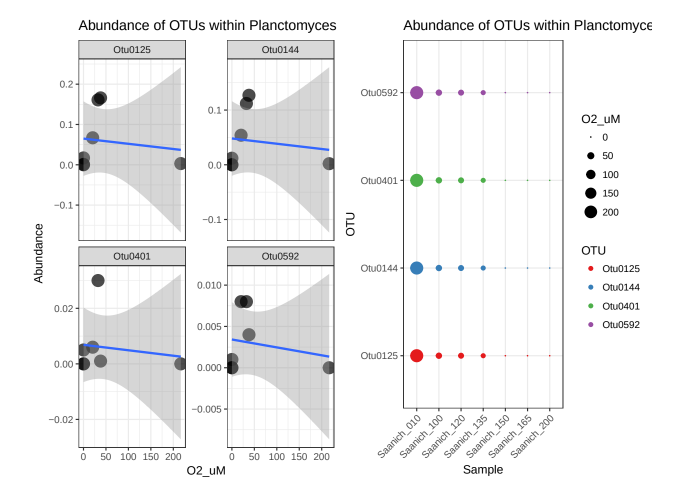
# 4.4 Interpretation of abundance information of OTUs/ASVs of [OTU\*\*\*\*] along with depth and/or oxygen concentration

```
## 0.101457 0.100546 0.009613 -0.037320 -0.079944 -0.050254 -0.044098
##
## Coefficients:
                Estimate Std. Error t value Pr(>|t|)
##
## (Intercept) 0.0849882 0.0753618
                                      1.128
                                               0.311
## Depth m
              -0.0002045 0.0005479 -0.373
                                               0.724
##
## Residual standard error: 0.08093 on 5 degrees of freedom
## Multiple R-squared: 0.0271, Adjusted R-squared: -0.1675
## F-statistic: 0.1393 on 1 and 5 DF, p-value: 0.7243
##
## ### General linear model for OtuO144
## Call:
## lm(formula = Abundance ~ Depth_m, data = .)
## Residuals:
                  2
                           3
                                    4
                                             5
   0.07920 0.06727 0.01157 -0.02814 -0.05959 -0.03784 -0.03247
##
## Coefficients:
##
                Estimate Std. Error t value Pr(>|t|)
## (Intercept) 0.0631237 0.0554978
                                      1.137
                                              0.307
## Depth_m
              -0.0001533 0.0004035 -0.380
                                               0.720
## Residual standard error: 0.0596 on 5 degrees of freedom
## Multiple R-squared: 0.02805,
                                   Adjusted R-squared: -0.1663
## F-statistic: 0.1443 on 1 and 5 DF, p-value: 0.7197
##
## ### General linear model for OtuO401
## Call:
## lm(formula = Abundance ~ Depth_m, data = .)
##
## Residuals:
           1
                                 3
## 2.399e-02 8.511e-06 -9.777e-04 -5.024e-03 -6.106e-03 -5.964e-03
##
## -5.932e-03
## Coefficients:
                Estimate Std. Error t value Pr(>|t|)
## (Intercept) 6.115e-03 1.110e-02
                                     0.551
                                               0.605
## Depth_m
              -9.165e-07 8.067e-05 -0.011
                                               0.991
##
## Residual standard error: 0.01192 on 5 degrees of freedom
## Multiple R-squared: 2.582e-05, Adjusted R-squared:
## F-statistic: 0.0001291 on 1 and 5 DF, p-value: 0.9914
##
## ### General linear model for Otu0592
```

```
## Call:
## lm(formula = Abundance ~ Depth_m, data = .)
##
## Residuals:
##
                                                                6
   0.0049869
             ##
## -0.0028298
##
## Coefficients:
##
                Estimate Std. Error t value Pr(>|t|)
## (Intercept) 3.288e-03 3.768e-03
                                     0.873
                                              0.423
## Depth_m
              -2.291e-06 2.740e-05 -0.084
                                              0.937
##
## Residual standard error: 0.004047 on 5 degrees of freedom
## Multiple R-squared: 0.001397,
                                 Adjusted R-squared: -0.1983
## F-statistic: 0.006996 on 1 and 5 DF, p-value: 0.9366
p.adjust(runif(length(suggestedOTUs), min = 0.005, max = 0.85), method = "fdr")
## [1] 0.5484101 0.5484101 0.5484101 0.5484101
# Abundance of OTUs within unclassified domain across depth
p5 <- workingTaxa %>% psmelt() %>% ggplot() + geom_point(aes(x = Depth_m, y = Abundance),
   size = 5, alpha = 0.7) + geom_smooth(method = "lm", aes(x = Depth_m, y = Abundance)) +
   facet_wrap(~OTU, scales = "free_y") + labs(title = "Abundance of OTUs within Planctomyces )
# Abundance of OTUs within unclassified depth by colour
p6 <- workingTaxa %>% psmelt() %>% ggplot() + geom_point(aes(x = Sample, y = OTU,
   size = Abundance, color = OTU)) + scale_size_continuous(range = c(0, 5)) +
   theme(axis.text.x = element_text(angle = 45, hjust = 1)) + labs(title = "Abundance of OTUs")
grid.arrange(p5, p6, ncol = 2)
```



```
# Abundance of OTUs within Planctomyces genus across depth
p7 <- workingTaxa %>% psmelt() %>% ggplot() + geom_point(aes(x = O2_uM, y = Abundance),
    size = 5, alpha = 0.7) + geom_smooth(method = "lm", aes(x = O2_uM, y = Abundance)) +
    facet_wrap(~OTU, scales = "free_y") + labs(title = "Abundance of OTUs within Planctomyces genus by colour
p8 <- workingTaxa %>% psmelt() %>% ggplot() + geom_point(aes(x = Sample, y = OTU,
    size = O2_uM, color = OTU)) + scale_size_continuous(range = c(0, 5)) + theme(axis.text.x =
    hjust = 1)) + labs(title = "Abundance of OTUs within Planctomyces genus across oxygen concergid.arrange(p7, p8, ncol = 2)
```



#### 5 Discussion

(Hawley et al. 2017; Torres-Beltrán et al. 2017)

## References

Hawley, Alyse K, Mónica Torres-Beltrán, Elena Zaikova, David A Walsh, Andreas Mueller, Melanie Scofield, Sam Kheirandish, et al. 2017. "A Compendium of Multi-Omic Sequence Information from the Saanich Inlet Water Column." *Scientific Data* 4. Nature Publishing Group: 170160.

Torres-Beltrán, Mónica, Alyse K Hawley, David Capelle, Elena Zaikova, David A Walsh, Andreas Mueller, Melanie Scofield, et al. 2017. "A Compendium of Geochemical Information from the Saanich Inlet Water Column." *Scientific Data* 4. Nature Publishing Group: 170159.