# Chemo.03.ComunityComposition

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# 1 Load packages and formatting

# 1.1 Loading packages

```
rm(list = ls())
library(phyloseq) # Phyloseq object
library(ggplot2)
library(vegan)
library(dplyr)
library(ape)
library(iCAMP) # Null model
library(cowplot) # Mutiple plot arrangement
```

```
library(egg) # Tag edition
library(reshape) # Dataframe formation
library(stringr) # String manipulation
library(ggstatsplot)
library(microViz)
library(kableExtra) # To nicely print tables
```

## 1.2 Setting colorblind palette

### 1.3 Loading and formating datasets

In this section the object schema with information about the experimental setup (Sample.ID; Chemostat.ID: chemostats number from 1 to 12; sampling time T: 1:9, DOM: DOM regime; Sal: disturbance regime) is created. Metabarcoding sequence data (16s rRNA gene) were analyzed using the dada2-pipeline (Callahan et al. 2016, doi: 10.1038/nmeth.3869) using the code in script01\_dada2.R and the resulting phyloseq project containing the ASV count table is uploaded.

```
## # A tibble: 108 x 5
##
     sample.ID T
                     Chem.ID DOM
                                   Sal
      <chr>
               <fct> <chr>
                             <fct> <fct>
##
   1 C10-1-01 1
                             oDOM C
##
                     01
   2 C10-1-02 1
                     02
                             oDOM D
   3 C10-1-03 1
                     03
                             oDOM C
##
##
   4 C10-1-04 1
                     04
                             oDOM D
##
  5 C10-1-05 1
                     05
                             oDOM C
  6 C10-1-06 1
                     06
                             oDOM D
##
##
  7 C10-1-07 1
                     07
                             eDOM
                                  С
## 8 C10-1-08 1
                     80
                             eDOM D
## 9 C10-1-09 1
                     09
                             eDOM C
## 10 C10-1-10 1
                             eDOM D
                     10
## # ... with 98 more rows
```

```
# Loading phyloseg object from #dada2
ps <- readRDS("../data/dada2.output/chem.ps.rds")</pre>
# Loading phylogenetic tree
chem.tree = read tree("../data/dada2.output/dada-chem.GTR2")
phy_tree(ps) <- chem.tree #Adding phylo-tree to the phyloseq object</pre>
# Phyloseq object contain abundance table, sample information, taxonomic
# information and the phylogenetic tree
## phyloseq-class experiment-level object
## otu_table()
                OTU Table:
                              [ 1447 taxa and 110 samples ]
## sample_data() Sample Data:
                                   [ 110 samples by 3 sample variables ]
                Taxonomy Table: [ 1447 taxa by 7 taxonomic ranks ]
## tax_table()
## phy_tree()
                Phylogenetic Tree: [ 1447 tips and 1445 internal nodes ]
# Removing initial inoculum samples for downstream analysis
ps = subset_samples(ps, sample.ID != "C10-0-LL" & sample.ID != "C10-0-HH")
ps
## phyloseq-class experiment-level object
## otu_table()
                                    [ 1447 taxa and 108 samples ]
                OTU Table:
                                   [ 108 samples by 3 sample variables ]
## sample_data() Sample Data:
                Taxonomy Table: [ 1447 taxa by 7 taxonomic ranks ]
## tax_table()
                Phylogenetic Tree: [ 1447 tips and 1445 internal nodes ]
## phy_tree()
```

# 2 Microbial community dynamic

### 2.1 Preprocess phyloseq-object

```
# Rarefy by minimum readnumber and transform to relative data set.seed(1)
# # Set seed for rarefaction
ps = rarefy_even_depth(ps, min(rowSums(otu_table(ps))), rngseed = 1, replace = F,
   trimOTUs = F)
# Estimating relative abundance
rOTUdf.rar <- prop.table(otu_table(ps), 1)</pre>
# New phyloseq-project with rarefied ASV table
otu_table(ps) <- otu_table(rOTUdf.rar, taxa_are_rows = FALSE)</pre>
## phyloseq-class experiment-level object
## otu_table()
               OTU Table:
                               [ 1447 taxa and 108 samples ]
## sample_data() Sample Data: [ 108 samples by 3 sample variables ]
                Taxonomy Table: [ 1447 taxa by 7 taxonomic ranks ]
## tax_table()
## phy_tree()
                Phylogenetic Tree: [ 1447 tips and 1445 internal nodes ]
```

```
# Keep ASVs with prevalence equivalent to more O reads
ps <- prune_taxa(taxa_sums(ps) > 0, ps)
ps
## phyloseq-class experiment-level object
                OTU Table: [ 973 taxa and 108 samples ]
## otu table()
## sample_data() Sample Data:
                                   [ 108 samples by 3 sample variables ]
                Taxonomy Table: [ 973 taxa by 7 taxonomic ranks ]
## tax table()
## phy_tree()
                Phylogenetic Tree: [ 973 tips and 971 internal nodes ]
# Setting up metadata Creating the additional identifier 'sample.names'
# for downstream analysis
schema$sample.names = str_replace_all(schema$sample.ID, "-", ".")
head(schema)
    sample.ID T Chem.ID DOM Sal sample.names
## 1 C10-1-01 1
                     O1 oDOM
                              С
                                     C10.1.01
## 2 C10-1-02 1
                     02 oDOM
                               D
                                     C10.1.02
## 3 C10-1-03 1
                     03 oDOM C
                                     C10.1.03
## 4 C10-1-04 1
                     04 oDOM D
                                     C10.1.04
## 5 C10-1-05 1
                     O5 oDOM
                              С
                                     C10.1.05
## 6 C10-1-06 1
                     06 oDOM D
                                     C10.1.06
# Re-Order ps object by sample ID from schema-object
new_order <- schema$sample.ID</pre>
ps = ps %>%
   ps_reorder(new_order) #From microViz
```

#### 2.2 Setting-up matrix for barplot

```
# Combine relative count-table with taxonomic information in a new
# dataframe
df1 <- data.frame(tax_table(ps), t(otu_table(ps)))</pre>
# Aggregate relative counts by taxonomic Order
df2 \leftarrow aggregate(. \sim Order, data = df1[, c(4, 8:115)], sum, na.rm = TRUE)
tibble(df2)
## # A tibble: 28 x 109
##
      Order C10.1~1 C10.1~2 C10.1~3 C10.1~4 C10.1~5 C10.1~6 C10.1~7 C10.1~8 C10.1~9
##
              <dbl>
                      <dbl>
                              <dbl>
                                      <dbl>
                                              <dbl>
                                                      <dbl>
                                                              <dbl>
  1 Acho~ 5.31e-4 4.25e-4 3.19e-4 2.12e-4 1.06e-4 1.06e-4 0
                                                                    0
   2 Acti~ 1.81e-3 1.81e-3 1.49e-3 1.49e-3 2.02e-3 2.34e-3 8.50e-4 6.37e-4 4.25e-4
                            0
                                    0
## 3 Bact~ 0
                    0
                                            0
                                                    0
                                                            0
                                                                    0
## 4 Baln~ 0
                    0
                            0
                                    0
## 5 Beta~ 8.07e-3 1.59e-3 2.01e-2 2.23e-3 2.23e-3 2.76e-3 6.69e-3 2.23e-3 1.59e-3
## 6 Caul~ 1.06e-4 0
                            2.12e-4 2.12e-4 0
                                                    0
                                                            0
                                                                     1.06e-4 0
## 7 Chit~ 0
                    0
                            0
                                    0
                                            0
                                                    0
                                                            0
                                                                    Ω
## 8 Cyto~ 6.37e-4 7.44e-4 4.25e-4 4.25e-4 3.19e-4 4.25e-4 1.27e-3 1.49e-3 2.12e-4
## 9 Ente~ 2.88e-1 3.86e-1 2.76e-1 3.57e-1 2.46e-1 3.37e-1 2.03e-1 2.74e-1 1.40e-1
```

```
## 10 Flav~ 6.69e-1 5.94e-1 6.97e-1 6.30e-1 7.34e-1 6.40e-1 5.90e-1 5.46e-1 7.41e-1
## # ... with 18 more rows, 99 more variables: C10.1.10 <dbl>, C10.1.11 <dbl>,
      C10.1.12 <dbl>, C10.2.01 <dbl>, C10.2.02 <dbl>, C10.2.03 <dbl>,
      C10.2.04 <dbl>, C10.2.05 <dbl>, C10.2.06 <dbl>, C10.2.07 <dbl>,
      C10.2.08 <dbl>, C10.2.09 <dbl>, C10.2.10 <dbl>, C10.2.11 <dbl>,
## #
      C10.2.12 <dbl>, C10.3.01 <dbl>, C10.3.02 <dbl>, C10.3.03 <dbl>,
      C10.3.04 <dbl>, C10.3.05 <dbl>, C10.3.06 <dbl>, C10.3.07 <dbl>,
## # C10.3.08 <dbl>, C10.3.09 <dbl>, C10.3.10 <dbl>, C10.3.11 <dbl>, ...
order <- df2[, 1] #Create a vector with only taxonomic Orders
rownames(df2) <- df2[, 1] # Set taxonomic orders as rownames
df2 <- df2[, -1] #remove column with orders and keep only abundance data
# tcolSums to see representatively of the top orders ()values close to 1
# are reached)
summary(colSums(df2))
     Min. 1st Qu. Median
                             Mean 3rd Qu.
## 0.9686 0.9996 0.9998 0.9990 0.9999 1.0000
# Create a new object for the aggregated relative abundance
agg = df2
agg <- as.data.frame(agg) #Formatting it as dataframe</pre>
rownames(agg) <- order #add order information as rownames</pre>
agg$Sum.agg <- rowSums(agg) # add an extra column with counts across treatments for all orders
# select the 10 most abundand orders
agg10 <- agg[with(agg, order(-Sum.agg)), ][1:10, 1:108]
agg10$order <- rownames(agg10)</pre>
tibble(agg10)
## # A tibble: 10 x 109
     C10.1.01 C10.1.02 C10.1.03 C10.1.04 C10.1.05 C10.1.06 C10.1~1 C10.1~2 C10.1~3
##
        <dbl>
                 <dbl>
                          <dbl>
                                   <dbl>
                                           <dbl>
                                                    <dbl>
                                                            <dbl>
                                                                    <dbl>
1.89e-1 1.59e-1 1.11e-1
                       0.697
## 2 0.669
              0.594
                                                 0.640
                                0.630
                                        0.734
                                                          5.90e-1 5.46e-1 7.41e-1
## 3 0.288
              0.386
                       0.276
                                0.357
                                         0.246
                                                 0.337
                                                          2.03e-1 2.74e-1 1.40e-1
## 4 0
              Ω
                       Ω
                                0
                                        0.000106 0.000106 3.19e-4 0
## 5 0.000106 0
                       0.000212 0.000212 0
                                                 0
                                                          0
                                                                  1.06e-4 0
## 6 0.000956 0.000106 0
                                0
                                                 0.000106 1.06e-4 2.12e-4 2.12e-4
                                        0
             0.00106 0.000319 0.000425 0.000212 0.000106 8.39e-3 1.48e-2 5.21e-3
## 7 0.0119
                                0.000212 0.000106 0
## 8 0.000212 0
                                                          0
                       0
## 9 0.00807 0.00159 0.0201
                                0.00223 0.00223 0.00276 6.69e-3 2.23e-3 1.59e-3
## 10 0.000637 0.000744 0.000425 0.000425 0.000319 0.000425 1.27e-3 1.49e-3 2.12e-4
## # ... with 100 more variables: C10.1.10 <dbl>, C10.1.11 <dbl>, C10.1.12 <dbl>,
      C10.2.01 <dbl>, C10.2.02 <dbl>, C10.2.03 <dbl>, C10.2.04 <dbl>,
      C10.2.05 <dbl>, C10.2.06 <dbl>, C10.2.07 <dbl>, C10.2.08 <dbl>,
      C10.2.09 <dbl>, C10.2.10 <dbl>, C10.2.11 <dbl>, C10.2.12 <dbl>,
## #
## #
      C10.3.01 <dbl>, C10.3.02 <dbl>, C10.3.03 <dbl>, C10.3.04 <dbl>,
## #
      C10.3.05 <dbl>, C10.3.06 <dbl>, C10.3.07 <dbl>, C10.3.08 <dbl>,
## #
      C10.3.09 <dbl>, C10.3.10 <dbl>, C10.3.11 <dbl>, C10.3.12 <dbl>, ...
```

```
# Check accumulated abundance after pooling by 10 most abundant orders
summary(colSums(agg10[, 1:108])) # All above or around 0.95
##
      Min. 1st Qu.
                    Median
                              Mean 3rd Qu.
                                              Max.
   0.9674 0.9975 0.9985
                            0.9974 0.9993
                                            1.0000
# convert dataframe from wide to long format
agg10.long <- melt(agg10, id.vars = "order", variable.name = "variable")</pre>
# Combine aggregated data with metadata
agg10.long = merge(agg10.long, schema, by.x = "variable", by.y = "sample.names")
# Add experimental time (sampling day) as a new column
agg10.long$Time = as.data.frame(str_split_fixed(as.vector(agg10.long$variable),
    "", 7))[, 5]
# Formatting columns as factors
agg10.long$Time = as.factor(agg10.long$Time)
levels(agg10.long$Time) = c("4", "8", "15", "18", "22", "29", "36", "39", "41")
agg10.long$Sal = as.factor(agg10.long$Sal)
# Rename treatments
levels(agg10.long$Sal) = c("Control", "Disturbance")
agg10.long$DOM = as.factor(agg10.long$DOM)
# Add replicate identifier
agg10.long$Rep = rep(c(1, 2, 3), each = 20)
tibble(agg10.long)
## # A tibble: 1,080 x 10
##
      variable order
                                value sampl~1 T
                                                    Chem.ID DOM
                                                                   Sal
                                                                         Time
##
      <fct>
               <chr>>
                                <dbl> <chr>
                                              <fct> <chr>
                                                             <fct> <fct> <fct> <dbl>
## 1 C10.1.01 Rhodobacteral~ 1.73e-2 C10-1-~ 1
                                                    01
                                                             oDOM Cont~ 4
## 2 C10.1.01 Flavobacteria~ 6.69e-1 C10-1-~ 1
                                                             oDOM Cont~ 4
                                                                                   1
                                                    01
## 3 C10.1.01 Enterobactera~ 2.88e-1 C10-1-~ 1
                                                    01
                                                             oDOM Cont~ 4
                                                                                   1
## 4 C10.1.01 Sphingomonada~ 0
                                      C10-1-~ 1
                                                    01
                                                             oDOM Cont~ 4
                                                                                   1
## 5 C10.1.01 Caulobacteral~ 1.06e-4 C10-1-~ 1
                                                             oDOM Cont~ 4
                                                    01
                                                                                   1
## 6 C10.1.01 Rhodospirilla~ 9.56e-4 C10-1-~ 1
                                                    01
                                                             oDOM Cont~ 4
                                                                                   1
## 7 C10.1.01 Pseudomonadal~ 1.19e-2 C10-1-~ 1
                                                    01
                                                             oDOM Cont~ 4
                                                                                   1
## 8 C10.1.01 Rhizobiales
                              2.12e-4 C10-1-~ 1
                                                    01
                                                             oDOM Cont~ 4
                                                                                   1
## 9 C10.1.01 Betaproteobac~ 8.07e-3 C10-1-~ 1
                                                    01
                                                             oDOM
                                                                  Cont~ 4
                                                                                   1
## 10 C10.1.01 Cytophagales
                              6.37e-4 C10-1-~ 1
                                                    01
                                                             oDOM
                                                                  Cont~ 4
                                                                                   1
```

#### 2.3 Microbial community relative abundance

Barplots indicate the change of the 10 most abundant order along the chemostat experiment by DOM level (HDOM and LDOM) by disturbance regime (Disturbed and undisturbed Controls). Early stages during the succession were dominated by members of Flavobacteriales and Enterobacterales orders at LDOM, as well as Rhodobacterales under the HDOM regime. At intermediate succession stages, Rhodobacterales and

## # ... with 1,070 more rows, and abbreviated variable name 1: sample.ID

Caulobacterales increased in abundance. At the experiment end diverse taxonomic structures were observed and depending on the culture vessel for instance members of the Caulobacterales, Sphingomonadales, Rhodospirillales, or Enterobacterales dominated the community.

```
# Create an individual plot for oDOM treatment
plot0 <- agg10.long[agg10.long$DOM == "oDOM", ] %>%
    ggplot(aes(x = Time, y = value, fill = order)) + geom_bar(stat = "identity",
    width = 1) + scale_fill_manual(values = cbbPalette) + facet_grid(Sal ~ Rep) +
   xlab("Sampling day") + ylab("Relative abundance") + theme_bw() + theme(strip.placement.y = "outside
    axis.text.x = element_text(size = 7), strip.text.y = element_text(angle = 270),
    strip.background = element_blank()) + theme(plot.margin = margin(t = 2,
   r = -12, b = 0, l = 0, unit = "pt"))
plot0 <- plot0 + theme(legend.position = "none")</pre>
# Create an individual plot for eDOM treatment
plot1 <- agg10.long[agg10.long$DOM == "eDOM", ] %>%
    ggplot(aes(x = Time, y = value, fill = order)) + geom_bar(stat = "identity",
    width = 1) + scale_fill_manual(values = cbbPalette) + facet_grid(Sal ~ Rep) +
   xlab("Sampling day") + ylab("Relative abundance") + theme bw() + theme(strip.placement.y = "outside
    axis.text.x = element_text(size = 7), strip.text.y = element_text(angle = 270),
    strip.background = element_blank()) + theme(plot.margin = margin(t = 2,
    r = 0, b = 0, l = 0, unit = "pt"))
```

#### 2.3.1 Figure relative abundance

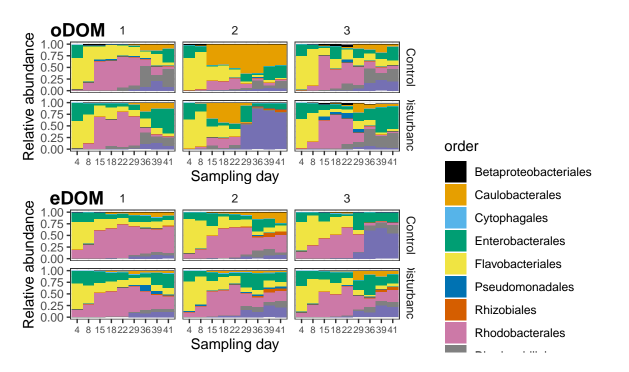


Figure 1: Community structure. Relative abundance of the 10 most abundant orders in the initial communities as well as in the resuscitated communities at LDOM and HDOM levels at each sampling day.

## 3 Betadiversity

#### 3.1 Bray-Curtis ordination visualization

```
# Bray-curtis (for relative abundance)
ord.pcoa.bray <- ordinate(ps, method = "PCoA", distance = "bray")</pre>
# Setting dataframes with ordination estimations
df.ord.pcoa.bray = data.frame(pco1 = ord.pcoa.bray$vectors[, 1], pco2 = ord.pcoa.bray$vectors[,
   2], Treatment = schema$Sal, DOM = schema$DOM, day = as.factor(schema$T),
   Chem.ID = schema$Chem.ID)
tibble(df.ord.pcoa.bray)
## # A tibble: 108 x 6
       pco1
             pco2 Treatment DOM
                                    day
                                          Chem. ID
##
      <dbl>
              <dbl> <fct>
                              <fct> <fct> <chr>
  1 -0.487 -0.237 C
                              oDOM 1
## 2 -0.428 -0.298 D
                              oDOM 1
                                          02
## 3 -0.534 -0.220 C
                              oDOM 1
                                          03
## 4 -0.472 -0.270 D
                              oDOM 1
                                          04
## 5 -0.472 -0.267 C
                              oDOM 1
                                          05
## 6 -0.448 -0.288 D
                                          06
                              oDOM 1
## 7 -0.440 -0.0472 C
                              eDOM 1
                                          07
## 8 -0.433 -0.0623 D
                              eDOM 1
                                          08
## 9 -0.459 -0.130 C
                                          09
                              eDOM 1
## 10 -0.457 -0.134 D
                              eDOM 1
                                          10
## # ... with 98 more rows
# Replace sampling ID by sampling day
levels(df.ord.pcoa.bray$day) = c("4", "8", "15", "18", "22", "29", "36", "39",
    "41")
# Visualize dataframe for Bray-curtis distance ordination
tibble(df.ord.pcoa.bray)
## # A tibble: 108 x 6
##
              pco2 Treatment DOM
                                    day
                                          Chem. ID
      <dbl>
              <dbl> <fct>
                              <fct> <fct> <chr>
  1 -0.487 -0.237 C
                              oDOM 4
##
                                          01
   2 -0.428 -0.298 D
                              oDOM 4
                                          02
## 3 -0.534 -0.220 C
                              oDOM 4
                                          03
## 4 -0.472 -0.270 D
                              oDOM 4
                                          04
## 5 -0.472 -0.267 C
                              oDOM 4
                                          05
## 6 -0.448 -0.288 D
                              oDOM 4
                                          06
## 7 -0.440 -0.0472 C
                              eDOM 4
                                          07
## 8 -0.433 -0.0623 D
                                          80
                              eDOM 4
## 9 -0.459 -0.130 C
                              eDOM 4
                                          09
## 10 -0.457 -0.134 D
                              eDOM 4
                                          10
## # ... with 98 more rows
```

```
# Create a label vector for DNA-sampling days
labels = c("d4", "d8", "d15", "d18", "d22", "d29", "d36", "d39", "d41")
# Figure all data points
plot.ord.pcoa.bray = ggplot(df.ord.pcoa.bray, aes(x = pco1, y = pco2)) + geom_point(aes(shape = interac
    DOM), colour = factor(day)), size = 5, alpha = 0.9) + scale_shape_manual(values = c(1,
    20, 2, 17), name = "Treatments", labels = c("C-eDOM", "D-eDOM", "C-oDOM",
    "D-oDOM")) + scale_color_manual(values = cbbPalette, name = "Sampling day") +
    theme_bw() + labs(title = "") + theme(text = element_text(size = 10, family = "ArialMT"),
    panel.grid.minor = element_blank()) + scale_x_continuous(breaks = seq(-0.5,
    0.5, 0.125), expand = c(0.02, 0.02)) + scale_y_continuous(breaks = seq(-0.5,
    0.5, 0.125), expand = c(0.02, 0.02)) + xlab(paste("PCOA1 [", round(ord.pcoa.bray$values[3][1,
    ] * 100, 0), "%]")) + ylab(paste("PCOA2 [", round(ord.pcoa.bray$values[3][2,
    ] * 100, 0), "%]")) + theme(legend.position = "none")
# Panels
plot.ord.pcoa.bray.b = ggplot(df.ord.pcoa.bray, aes(x = pco1, y = pco2)) + geom_point(aes(shape = inter
    DOM), colour = factor(day)), size = 3, alpha = 0.9) + scale_shape_manual(values = c(1,
    20, 2, 17), name = "Treatments", labels = c("C-eDOM", "D-eDOM", "C-oDOM",
    "D-oDOM")) + scale_color_manual(values = cbbPalette, name = "Sampling day") +
    theme_bw() + labs(title = "") + theme_bw() + labs(title = "", x = "", y = "") +
    theme(text = element_text(size = 10, family = "ArialMT"), panel.grid.minor = element_blank()) +
    scale_x_{continuous}(breaks = seq(-0.5, 0.5, 0.125), expand = c(0.02, 0.02)) +
    scale_y = continuous(breaks = seq(-0.5, 0.5, 0.125), expand = c(0.02, 0.02)) +
    facet_wrap(~day, ncol = 3, scales = "free") + theme(axis.text.x = element_blank(),
    axis.text.y = element_blank(), axis.ticks = element_blank(), legend.key.size = unit(0.8,
        "lines"))
plot.ord.pcoa.bray.b = tag_facet(plot.ord.pcoa.bray.b, open = "", close = ")",
    tag pool = labels, hjust = -0.1, size = 3)
plot_grid(plot.ord.pcoa.bray, plot.ord.pcoa.bray.b, rel_widths = c(0.9, 1),
    ncol = 2, labels = c("A", "B"))
                                            В
Α
                                                                                Sampling day
                                                                     d15)
   0.375
                                                                                   8
                                                                                 15
   0.250
                                                                                   18
                                                                                   22
   0.125
PCOA2 [ 9 %]
                                                          d22)
   0.000
                                                                                   36
                                                                                   39
   -0.125
                                                                                 41
  -0.250
                                                          d39)
                                                                    d41)
                                                d36)
                                                                                Treatments
  -0.375
                                                                                 O C-eDOM
                                                                                   D-eDOM
  -0.500
                                                                                 △ C-oDOM
        -0.500 -0.375 -0.250 -0.125 0.000 0.125 0.250 0.375
                                                                                 ▲ D-oDOM
                   PCOA1 [ 14 %]
```

Figure 2: Community structure. A) Overview PCoA Bray-Curtis based biplot including all data points. B) PCoAs for individual sampling days; axes of the individual plots are differently scaled as indicated by the grid lines

#### 3.2 Bray-Curtis distance between T1-T9

```
# Bray-Curtis distance
dist.bray <- phyloseq::distance(ps, method = "bray")</pre>
# Reshape Bray-curtis distance into dataframe Function to transform
# distance matrix into dataframe
dist2df_AR <- function(m) {</pre>
    xy <- t(combn(colnames(m), 2))</pre>
    tmp = data.frame(xy, dist = m[xy])
   tmp$t1 = str_split_fixed(as.character(tmp$X1), "-", 3)[, 2]  # Extract the time point from sample '
   tmp$t2 = str_split_fixed(as.character(tmp$X2), "-", 3)[, 2] # Extract time point from sample 'y'
   tmp$dif = as.numeric(tmp$t2) - as.numeric(tmp$t1) # Calculate the difference in time units
   return(tmp)
}
df.dist.bray <- dist2df_AR(as.matrix(dist.bray))</pre>
# Summarizing only Bray-Curtis distances between T1 and T9
summary(df.dist.bray[df.dist.bray$t1 == 1 & df.dist.bray$t2 == 9, "dist"])
      Min. 1st Qu. Median
                              Mean 3rd Qu.
## 0.7182 0.8164 0.8540 0.8560 0.8985 0.9503
```

#### 3.3 Unifrac ordination visualization

```
# Unifrac (Weighted=T for considering relative abundances)
ord.pcoa.unif <- ordinate(ps, "PCoA", "unifrac", weighted = T)</pre>
## Warning in UniFrac(physeq, ...): Randomly assigning root as --
## SV_738_Rhodobacterales -- in the phylogenetic tree in the data you provided.
# Setting dataframes with ordination estimations
df.ord.pcoa.unif = data.frame(pco1 = ord.pcoa.unif$vectors[, 1], pco2 = ord.pcoa.unif$vectors[,
    2], Treatment = schema$Sal, DOM = schema$DOM, day = as.factor(schema$T),
    Chem.ID = schema$Chem.ID)
tibble(df.ord.pcoa.unif)
## # A tibble: 108 x 6
##
               pco2 Treatment DOM
                                           Chem. ID
       pco1
                                     day
             <dbl> <fct>
                               <fct> <fct> <chr>
##
       <dbl>
## 1 -0.327 -0.0980 C
                               oDOM 1
                                           01
## 2 -0.319 -0.111 D
                               oDOM 1
                                           02
## 3 -0.335 -0.0973 C
                                           03
                               oDOM 1
## 4 -0.326 -0.109 D
                               oDOM 1
                                           04
## 5 -0.337 -0.0966 C
                               oDOM 1
                                           05
## 6 -0.326 -0.108 D
                               oDOM 1
                                           06
                              eDOM 1
## 7 -0.285 -0.0385 C
                                           07
## 8 -0.281 -0.0572 D
                               eDOM 1
                                           80
## 9 -0.327 -0.0473 C
                               eDOM 1
                                           09
## 10 -0.308 -0.0643 D
                               eDOM 1
                                           10
## # ... with 98 more rows
```

```
# Replace sampling ID by sampling day
levels(df.ord.pcoa.unif$day) = c("4", "8", "15", "18", "22", "29", "36", "39",
    "41")
# Visualize dataframe for Unifrac distance ordination
tibble(df.ord.pcoa.unif)
## # A tibble: 108 x 6
##
       pco1
              pco2 Treatment DOM
                                   day
                                          Chem. ID
##
       <dbl>
              <dbl> <fct>
                              <fct> <fct> <chr>
                               oDOM 4
## 1 -0.327 -0.0980 C
                                          01
## 2 -0.319 -0.111 D
                               oDOM 4
                                          02
## 3 -0.335 -0.0973 C
                                          03
                               oDOM 4
## 4 -0.326 -0.109 D
                               oDOM 4
                                          04
## 5 -0.337 -0.0966 C
                               oDOM 4
                                          05
## 6 -0.326 -0.108 D
                               oDOM 4
                                          06
## 7 -0.285 -0.0385 C
                               eDOM 4
                                          07
## 8 -0.281 -0.0572 D
                                          80
                               eDOM 4
## 9 -0.327 -0.0473 C
                               eDOM 4
                                          09
## 10 -0.308 -0.0643 D
                               eDOM 4
                                          10
## # ... with 98 more rows
# Figure all data points
plot.ord.pcoa.unif = ggplot(df.ord.pcoa.unif, aes(x = pco1, y = pco2)) + geom_point(aes(shape = interac
   DOM), colour = factor(day)), size = 5, alpha = 0.9) + scale_shape_manual(values = c(1,
    20, 2, 17), name = "Treatments", labels = c("C-eDOM", "D-eDOM", "C-oDOM",
    "D-oDOM")) + scale_color_manual(values = cbbPalette, name = "Sampling day") +
   theme bw() + labs(title = "") + theme(text = element text(size = 10, family = "ArialMT"),
   panel.grid.minor = element_blank()) + scale_x_continuous(breaks = seq(-0.5,
   0.5, 0.125), expand = c(0.02, 0.02)) + scale_y_continuous(breaks = seq(-0.5, 0.125))
   0.5, 0.125), expand = c(0.02, 0.02)) + xlab(paste("PCOA1 [", round(ord.pcoa.unif$values[3][1,
   ] * 100, 0), "%]")) + ylab(paste("PCOA2 [", round(ord.pcoa.unif$values[3][2,
   ] * 100, 0), "%]")) + theme(legend.position = "none")
# Panels
plot.ord.pcoa.unif.b = ggplot(df.ord.pcoa.unif, aes(x = pco1, y = pco2)) + geom_point(aes(shape = inter
    DOM), colour = factor(day)), size = 3, alpha = 0.9) + scale_shape_manual(values = c(1,
    20, 2, 17), name = "Treatments", labels = c("C-eDOM", "D-eDOM", "C-oDOM",
    "D-oDOM")) + scale_color_manual(values = cbbPalette, name = "Sampling day") +
   theme_bw() + labs(title = "") + theme_bw() + labs(title = "", x = "", y = "") +
    theme(text = element_text(size = 10, family = "ArialMT"), panel.grid.minor = element_blank()) +
    scale_x_{continuous}(breaks = seq(-0.5, 0.5, 0.125), expand = c(0.02, 0.02)) +
    scale_y = scale_y = seq(-0.5, 0.5, 0.125), expand = c(0.02, 0.02)) +
   facet_wrap(~day, ncol = 3, scales = "free") + theme(axis.text.x = element_blank(),
    axis.text.y = element blank(), axis.ticks = element blank(), legend.key.size = unit(0.8,
        "lines"))
plot.ord.pcoa.unif.b = tag_facet(plot.ord.pcoa.unif.b, open = "", close = ")",
    tag_pool = labels, hjust = -0.1, size = 3)
plot grid(plot.ord.pcoa.unif, plot.ord.pcoa.unif.b, rel widths = c(0.9, 1),
   ncol = 2, labels = c("A", "B"))
```

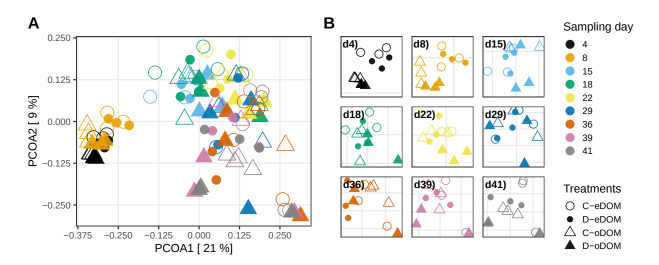


Figure 3: Community structure. A) Overview PCoA weighted Unifrac based biplot including all data points. B) PCoAs for individual sampling days; axes of the individual plots are differently scaled as indicated by the grid lines

## 4 Statistical analysis

#### 4.1 Permanova

# Bray-Curtis distance

We performed a Permutational Multivariate Analysis of Variance (Permanova) usgin the difference distance matrix estimated (Bray-curtis and Unifrac distances). Factor considered for the analysis were: 1) DOM level availability, 2) Disturbance treatment (as "Sal" treatment) and 3) Time.

```
dist.bray <- phyloseq::distance(ps, method = "bray")</pre>
# Unifrac distance
dist.wunif <- phyloseq::distance(ps, method = "wunifrac")</pre>
## Warning in UniFrac(physeq, weighted = TRUE, ...): Randomly assigning root as --
## SV_358_Enterobacterales -- in the phylogenetic tree in the data you provided.
# Permanova all factors: 1) DOM level availability, 2) Disturbance
# treatment (Sal) and 3) Time.
# Bray Curtis distance
set.seed(1)
\# permanova.bray <- adonis2(dist.bray ~ DOM*Sal*T, data=schema, permutations
permanova.bray <- adonis2(dist.bray ~ DOM * Sal, data = schema, permutations = 1000,
    strata = schema$T)
# Unifrac distance
set.seed(1)
permanova.wunifrac <- adonis2(dist.wunif ~ DOM * Sal * T, data = schema, permutations = 1000)
permanova.wunifrac
```

```
## Permutation test for adonis under reduced model
## Terms added sequentially (first to last)
## Permutation: free
## Number of permutations: 1000
## adonis2(formula = dist.wunif ~ DOM * Sal * T, data = schema, permutations = 1000)
             Df SumOfSqs
                          R2
                                      F Pr(>F)
             1 0.1902 0.03137 12.8421 0.000999 ***
## DOM
             1 0.0867 0.01430 5.8545 0.008991 **
## Sal
## T
              8 4.3430 0.71646 36.6611 0.000999 ***
## DOM:Sal
             1 0.0406 0.00669 2.7393 0.053946 .
## DOM:T
              8 0.2405 0.03967 2.0301 0.020979 *
              8 0.0445 0.00733 0.3753 0.989011
## Sal:T
## DOM:Sal:T 8 0.0502 0.00828 0.4239 0.981019
## Residual
            72 1.0662 0.17588
            107 6.0617 1.00000
## Total
## ---
## Signif. codes: 0 '*** 0.001 '** 0.01 '* 0.05 '.' 0.1 ' ' 1
```