

Chemo.04.ComunityAssembly

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1 Load packages and formatting

1.1 Loading packages

```
rm(list = ls())
library(phyloseq)
library(ggplot2)
library(vegan)
library(ecodist)
library(dplyr)
library(ape) #
library(iCAMP) #NULL model for microbial comm
library(cowplot) #Multiple panel formatting
library(egg) #Additional label for multiple panels
library(reshape) #Dataframe formatting
library(stringr) #Text manipulation
library(microViz) #Microbiome (re_order function for phyloseq-objects)
library(rstatix) #Versatile statistical package
```

1.2 Setting colorblind palette

```
# Colorblind palette
cbbPalette <- c("#000000", "#E69F00", "#56B4E9", "#009E73", "#F0E442", "#0072B2",
  "#D55E00", "#CC79A7", "#808080", "#7570B3")
```

1.3 Loading and formatting datasets

In this section the object schema with information about the experimental setup (Sample.ID; Chemostat.ID: chemostat 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.

```
# Create metadata for experimental setup
schema <- data.frame(sample.ID = paste0("C10-", rep(1:9, each = 12), "-", sprintf("%02d",
  1:12)), T = rep(1:9, each = 12), Chem.ID = sprintf("%02d", 1:12), DOM = rep(c("L",
  "H"), each = 6), Sal = c("C", "D"))
schema$T = as.factor(schema$T)
schema$DOM = as.factor(schema$DOM)
schema$Sal = as.factor(schema$Sal)
tibble(schema)
```

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

```
# Loading phyloseq object from #dada2
ps <- readRDS("../data/dada2.output/chem.ps.rds")
# Phyloseq object contain abundance table, sample information, taxonomic
# information and the phylogenetic tree

# Loadgin phylogenetic tree
chem.tree = read_tree("../data/dada2.output/dada-chem.GTR2")
phy_tree(ps) <- chem.tree #Adding phylo-tree to the phyloseq object

# Phyloseq object contain abundance table, sample information, taxonomic
# information and the phylogenetic tree
ps
```

```
## phyloseq-class experiment-level object
## otu_table() OTU Table: [ 1447 taxa and 110 samples ]
## sample_data() Sample Data: [ 110 samples by 3 sample variables ]
## tax_table() Taxonomy Table: [ 1447 taxa by 7 taxonomic ranks ]
## 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() OTU Table: [ 1447 taxa and 108 samples ]
## sample_data() Sample Data: [ 108 samples by 3 sample variables ]
## tax_table() Taxonomy Table: [ 1447 taxa by 7 taxonomic ranks ]
## phy_tree() Phylogenetic Tree: [ 1447 tips and 1445 internal nodes ]
```

2 Microbial community dynamic

2.1 Preprocess phyloseq-object

```
# Rarefy by minimum readnumber and transform to relative data
ps = rarefy_even_depth(ps, min(rowSums(otu_table(ps))), rngseed = 1, replace = F,
  trimOTUs = F)
```

```
## 'set.seed(1)' was used to initialize repeatable random subsampling.
```

```
## Please record this for your records so others can reproduce.
```

```
## Try 'set.seed(1); .Random.seed' for the full vector
```

```
## ...
```

```
# Estimating relative abundance
rOTUdf.rar <- prop.table(otu_table(ps), 1)

# New phyloseq-project with rarefied ASV table
otu_table(ps) <- otu_table(rOTUdf.rar, taxa_are_rows = FALSE)
ps
```

```
## phyloseq-class experiment-level object
## otu_table() OTU Table: [ 1447 taxa and 108 samples ]
## sample_data() Sample Data: [ 108 samples by 3 sample variables ]
## tax_table() Taxonomy Table: [ 1447 taxa by 7 taxonomic ranks ]
## phy_tree() Phylogenetic Tree: [ 1447 tips and 1445 internal nodes ]
```

```
# Keep ASVs with prevalence equivalent to more 0 reads
ps <- prune_taxa(taxa_sums(ps) > 0, ps)
ps
```

```
## phyloseq-class experiment-level object
## otu_table() OTU Table: [ 973 taxa and 108 samples ]
## sample_data() Sample Data: [ 108 samples by 3 sample variables ]
## tax_table() Taxonomy Table: [ 973 taxa by 7 taxonomic ranks ]
## phy_tree() Phylogenetic Tree: [ 973 tips and 971 internal nodes ]
```

```
# Setting up metadata
head(sample_data(ps))
```

```
##          sample.ID Chem.ID T
## C10-1-09 C10-1-09      09 1
## C10-2-05 C10-2-05      05 2
## C10-3-01 C10-3-01      01 3
## C10-3-09 C10-3-09      09 3
## C10-4-05 C10-4-05      05 4
## C10-5-01 C10-5-01      01 5
```

```
# Some samples in phyloseq object are not included into this analyses
# (schema), so we proceed to reorder ps-data base in the schema$sample.ID
```

```
# Re-Order ps object by sample ID from schema-object
new_order <- schema$sample.ID
ps = ps %>%
  ps_reorder(new_order) #MicroViz package
```

```
# Vizualize ordered ps-object
head(sample_data(ps))
```

```
##          sample.ID Chem.ID T
## C10-1-01 C10-1-01      01 1
## C10-1-02 C10-1-02      02 1
## C10-1-03 C10-1-03      03 1
## C10-1-04 C10-1-04      04 1
## C10-1-05 C10-1-05      05 1
## C10-1-06 C10-1-06      06 1
```

```
tail(sample_data(ps))
```

```
##          sample.ID Chem.ID T
## C10-9-07 C10-9-07      07 9
## C10-9-08 C10-9-08      08 9
## C10-9-09 C10-9-09      09 9
## C10-9-10 C10-9-10      10 9
## C10-9-11 C10-9-11      11 9
## C10-9-12 C10-9-12      12 9
```

2.2 Subset data by treatment

```

# Create empty list-objects
LD.ps = list() # List to store LDOM results
LC.ps = list() # List to store LDOM results

HD.ps = list() # List to store HDOM results
HC.ps = list() # List to store HDOM results

# Sample ID (Control and disturbance treatments)
LDOM_C = c("01", "03", "05") #L-DOM Samples for control
LDOM_D = c("02", "04", "06") #L-DOM Samples for disturbed treatments
HDOM_C = c("07", "09", "11") #H-DOM Samples for control
HDOM_D = c("08", "10", "12") #H-DOM Samples for disturbed treatments

# LDOM level Filter by Sample
tmpL <- prune_samples(ps@sam_data[["Chem.ID"]] %in% LDOM_C, ps)
# Filter ASVs (taxa) to only those with abund equal to 0 in all the samples
LC.ps <- filter_taxa(tmpL, function(x) sum(x != 0) > 0, TRUE)
LC.ps

## phyloseq-class experiment-level object
## otu_table() OTU Table: [ 437 taxa and 27 samples ]
## sample_data() Sample Data: [ 27 samples by 3 sample variables ]
## tax_table() Taxonomy Table: [ 437 taxa by 7 taxonomic ranks ]
## phy_tree() Phylogenetic Tree: [ 437 tips and 435 internal nodes ]

tmpL <- prune_samples(ps@sam_data[["Chem.ID"]] %in% LDOM_D, ps)
LD.ps <- filter_taxa(tmpL, function(x) sum(x != 0) > 0, TRUE)
LD.ps

## phyloseq-class experiment-level object
## otu_table() OTU Table: [ 394 taxa and 27 samples ]
## sample_data() Sample Data: [ 27 samples by 3 sample variables ]
## tax_table() Taxonomy Table: [ 394 taxa by 7 taxonomic ranks ]
## phy_tree() Phylogenetic Tree: [ 394 tips and 392 internal nodes ]

# HDOM level Filter by Sample
tmpH <- prune_samples(ps@sam_data[["Chem.ID"]] %in% HDOM_C, ps)
# Filter ASVs (taxa) to only those with abund equal to 0 in all the samples
HC.ps <- filter_taxa(tmpH, function(x) sum(x != 0) > 0, TRUE)
HC.ps

## phyloseq-class experiment-level object
## otu_table() OTU Table: [ 576 taxa and 27 samples ]
## sample_data() Sample Data: [ 27 samples by 3 sample variables ]
## tax_table() Taxonomy Table: [ 576 taxa by 7 taxonomic ranks ]
## phy_tree() Phylogenetic Tree: [ 576 tips and 574 internal nodes ]

tmpH <- prune_samples(ps@sam_data[["Chem.ID"]] %in% HDOM_D, ps)
# Filter ASVs (taxa) to only those with abund equal to 0 in all the samples
HD.ps <- filter_taxa(tmpH, function(x) sum(x != 0) > 0, TRUE)
HD.ps

```

```
## phyloseq-class experiment-level object
## otu_table() OTU Table: [ 537 taxa and 27 samples ]
## sample_data() Sample Data: [ 27 samples by 3 sample variables ]
## tax_table() Taxonomy Table: [ 537 taxa by 7 taxonomic ranks ]
## phy_tree() Phylogenetic Tree: [ 537 tips and 535 internal nodes ]
```

2.3 Compute the beta-Nearest Taxon Index (bNTI)

The Influence of deterministic versus stochastic processes on microbial community dynamics was quantified during the course of the continuous culture experiment via null model analyses using the beta nearest taxon indices (bNTI) between sample pairs. For this purpose we applied bNTIs between two temporally succeeding samples separately for disturbance and DOM regimes by applying a sliding window setup in the continuous cultures.

```
knitr::opts_chunk$set(cache = T)

NTI.out.LC = list() #Loop for LDOM for each sampling point
NTI.out.LD = list() #Loop for LDOM for each sampling point

# L-DOM x Control
comm = otu_table(LC.ps)
dist = cophenetic(phy_tree(LC.ps)) #Standard distance calculation for tree used in the manual
set.seed(1)
NTI.out.LC = bNTIn.p(comm@.Data, dist, nworker = 2, memo.size.GB = 50, weighted = TRUE,
  exclude.consp = FALSE, rand = 1000, output.bMNTD = FALSE, sig.index = "SES",
  unit.sum = NULL, correct.special = FALSE, detail.null = FALSE, special.method = "MNTD")

## Now calculating observed betaMNTD. Begin at Sat Jan 14 12:44:56 2023. Please wait...

## Now randomizing by parallel computing. Begin at Sat Jan 14 12:44:57 2023. Please wait...

# L-DOM x Disturbance
comm = otu_table(LD.ps)
dist = cophenetic(phy_tree(LD.ps)) #Standard distance calculation for tree used in the manual
NTI.out.LD = bNTIn.p(comm@.Data, dist, nworker = 2, memo.size.GB = 50, weighted = TRUE,
  exclude.consp = FALSE, rand = 1000, output.bMNTD = FALSE, sig.index = "SES",
  unit.sum = NULL, correct.special = FALSE, detail.null = FALSE, special.method = "MNTD")

## Now calculating observed betaMNTD. Begin at Sat Jan 14 12:46:33 2023. Please wait...

## Now randomizing by parallel computing. Begin at Sat Jan 14 12:46:34 2023. Please wait...

NTI.out.HC = list() #Loop for HDOM for each sampling point
NTI.out.HD = list() #Loop for LDOM for each sampling point

# H-DOM x Control
comm = otu_table(HC.ps)
dist = cophenetic(phy_tree(HC.ps)) #Standard distance calculation for tree used in the manual
NTI.out.HC = bNTIn.p(comm@.Data, dist, nworker = 2, memo.size.GB = 50, weighted = TRUE,
  exclude.consp = FALSE, rand = 100, output.bMNTD = FALSE, sig.index = "SES",
  unit.sum = NULL, correct.special = FALSE, detail.null = FALSE, special.method = "MNTD")
```

```
## Now calculating observed betaMNTD. Begin at Sat Jan 14 12:47:58 2023. Please wait...
```

```
## Now randomizing by parallel computing. Begin at Sat Jan 14 12:47:59 2023. Please wait...
```

```
# H-DOM x Disturbance
comm = otu_table(HD.ps)
dist = cophenetic(phy_tree(HD.ps)) #Standard distance calculation for tree used in the manual
NTI.out.HD = bNTIn.p(comm@.Data, dist, nworker = 2, memo.size.GB = 50, weighted = TRUE,
  exclude.consp = FALSE, rand = 1000, output.bMNTD = FALSE, sig.index = "SES",
  unit.sum = NULL, correct.special = FALSE, detail.null = FALSE, special.method = "MNTD")
```

```
## Now calculating observed betaMNTD. Begin at Sat Jan 14 12:48:12 2023. Please wait...
```

```
## Now randomizing by parallel computing. Begin at Sat Jan 14 12:48:13 2023. Please wait...
```

2.4 Reshape bNTI index into a dataframe

```
# Function to transform distance matrix into dataframe
dist2df_AR <- function(m) {
  xy <- t(combn(colnames(m$index), 2))
  tmp = data.frame(xy, dist = m$index[xy])
  tmp$t1 = str_split_fixed(as.character(tmp$X1), "-", 3)[, 2] # Extract the time point from sample 'x'
  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)
}

# LDOM control regime
df.NTI.LC <- dist2df_AR(NTI.out.LC)
df.NTI.LC = df.NTI.LC[df.NTI.LC$dif == 1, ] # Get data space by only 1 time unit
df.NTI.LC$DOM = "LDM"
df.NTI.LC$Treatment = "Control"
tibble(df.NTI.LC)
```

```
## # A tibble: 72 x 8
##   X1      X2      dist t1    t2    dif DOM   Treatment
##   <chr>   <chr>   <dbl> <chr> <chr> <dbl> <chr>   <chr>
## 1 C10-1-01 C10-2-01 -0.950 1     2     1 LDM     Control
## 2 C10-1-01 C10-2-03  0.709 1     2     1 LDM     Control
## 3 C10-1-01 C10-2-05  1.48  1     2     1 LDM     Control
## 4 C10-1-03 C10-2-01  0.540 1     2     1 LDM     Control
## 5 C10-1-03 C10-2-03  2.39  1     2     1 LDM     Control
## 6 C10-1-03 C10-2-05  1.21  1     2     1 LDM     Control
## 7 C10-1-05 C10-2-01  0.420 1     2     1 LDM     Control
## 8 C10-1-05 C10-2-03  0.547 1     2     1 LDM     Control
## 9 C10-1-05 C10-2-05  1.04  1     2     1 LDM     Control
## 10 C10-2-01 C10-3-01  0.505 2     3     1 LDM     Control
## # ... with 62 more rows
```

```
# LDOM disturbance regime
```

```
df.NTI.LD <- dist2df_AR(NTI.out.LD)
df.NTI.LD = df.NTI.LD[df.NTI.LD$dif == 1, ] # Get data space by only 1 time unit
df.NTI.LD$DOM = "LDOM"
df.NTI.LD$Treatment = "Disturbance"
tibble(df.NTI.LD)
```

```
## # A tibble: 72 x 8
##   X1      X2      dist t1    t2      dif DOM    Treatment
##   <chr>   <chr>   <dbl> <chr> <chr> <dbl> <chr> <chr>
## 1 C10-1-02 C10-2-02 -1.07  1    2      1 LDOM Disturbance
## 2 C10-1-02 C10-2-04  1.21  1    2      1 LDOM Disturbance
## 3 C10-1-02 C10-2-06  0.0468 1    2      1 LDOM Disturbance
## 4 C10-1-04 C10-2-02  1.85  1    2      1 LDOM Disturbance
## 5 C10-1-04 C10-2-04  2.81  1    2      1 LDOM Disturbance
## 6 C10-1-04 C10-2-06  2.03  1    2      1 LDOM Disturbance
## 7 C10-1-06 C10-2-02  2.93  1    2      1 LDOM Disturbance
## 8 C10-1-06 C10-2-04  1.98  1    2      1 LDOM Disturbance
## 9 C10-1-06 C10-2-06  1.80  1    2      1 LDOM Disturbance
## 10 C10-2-02 C10-3-02 -0.315 2    3      1 LDOM Disturbance
## # ... with 62 more rows
```

```
# HDOM control regime
```

```
df.NTI.HC <- dist2df_AR(NTI.out.HC)
df.NTI.HC = df.NTI.HC[df.NTI.HC$dif == 1, ] # Get data space by only 1 time unit
df.NTI.HC$DOM = "HDOM"
df.NTI.HC$Treatment = "Control"
tibble(df.NTI.HC)
```

```
## # A tibble: 72 x 8
##   X1      X2      dist t1    t2      dif DOM    Treatment
##   <chr>   <chr>   <dbl> <chr> <chr> <dbl> <chr> <chr>
## 1 C10-1-07 C10-2-07 -0.330 1    2      1 HDOM Control
## 2 C10-1-07 C10-2-09  1.69  1    2      1 HDOM Control
## 3 C10-1-07 C10-2-11 -0.865 1    2      1 HDOM Control
## 4 C10-1-09 C10-2-07 -0.697 1    2      1 HDOM Control
## 5 C10-1-09 C10-2-09  0.429 1    2      1 HDOM Control
## 6 C10-1-09 C10-2-11 -1.66  1    2      1 HDOM Control
## 7 C10-1-11 C10-2-07  0.271 1    2      1 HDOM Control
## 8 C10-1-11 C10-2-09  1.04  1    2      1 HDOM Control
## 9 C10-1-11 C10-2-11 -1.03  1    2      1 HDOM Control
## 10 C10-2-07 C10-3-07 -1.24  2    3      1 HDOM Control
## # ... with 62 more rows
```

```
# HDOM disturbance regime
```

```
df.NTI.HD <- dist2df_AR(NTI.out.HD)
df.NTI.HD = df.NTI.HD[df.NTI.HD$dif == 1, ] # Get data space by only 1 time unit
df.NTI.HD$DOM = "HDOM"
df.NTI.HD$Treatment = "Disturbance"
tibble(df.NTI.HD)
```

```
## # A tibble: 72 x 8
```



```
##      X1      X2      dist t1      t2      dif DOM      Treatment
##      <chr>    <chr>    <dbl> <chr> <chr> <dbl> <chr> <chr>
## 1 C10-1-08 C10-2-08 0.590 1      2      1 HDOM Disturbance
## 2 C10-1-08 C10-2-10 -0.200 1      2      1 HDOM Disturbance
## 3 C10-1-08 C10-2-12 -1.39 1      2      1 HDOM Disturbance
## 4 C10-1-10 C10-2-08 -1.02 1      2      1 HDOM Disturbance
## 5 C10-1-10 C10-2-10 0.907 1      2      1 HDOM Disturbance
## 6 C10-1-10 C10-2-12 -0.408 1      2      1 HDOM Disturbance
## 7 C10-1-12 C10-2-08 -0.611 1      2      1 HDOM Disturbance
## 8 C10-1-12 C10-2-10 -0.775 1      2      1 HDOM Disturbance
## 9 C10-1-12 C10-2-12 -1.62 1      2      1 HDOM Disturbance
## 10 C10-2-08 C10-3-08 -0.658 2      3      1 HDOM Disturbance
## # ... with 62 more rows
```

```
# Pooling the dataframes together
df.NTI.all = rbind(df.NTI.LC, df.NTI.LD, df.NTI.HC, df.NTI.HD)
tibble(df.NTI.all)
```

```
## # A tibble: 288 x 8
##      X1      X2      dist t1      t2      dif DOM      Treatment
##      <chr>    <chr>    <dbl> <chr> <chr> <dbl> <chr> <chr>
## 1 C10-1-01 C10-2-01 -0.950 1      2      1 LDOM Control
## 2 C10-1-01 C10-2-03 0.709 1      2      1 LDOM Control
## 3 C10-1-01 C10-2-05 1.48 1      2      1 LDOM Control
## 4 C10-1-03 C10-2-01 0.540 1      2      1 LDOM Control
## 5 C10-1-03 C10-2-03 2.39 1      2      1 LDOM Control
## 6 C10-1-03 C10-2-05 1.21 1      2      1 LDOM Control
## 7 C10-1-05 C10-2-01 0.420 1      2      1 LDOM Control
## 8 C10-1-05 C10-2-03 0.547 1      2      1 LDOM Control
## 9 C10-1-05 C10-2-05 1.04 1      2      1 LDOM Control
## 10 C10-2-01 C10-3-01 0.505 2      3      1 LDOM Control
## # ... with 278 more rows
```

2.5 Plot boxplot bNTI per DOM regime

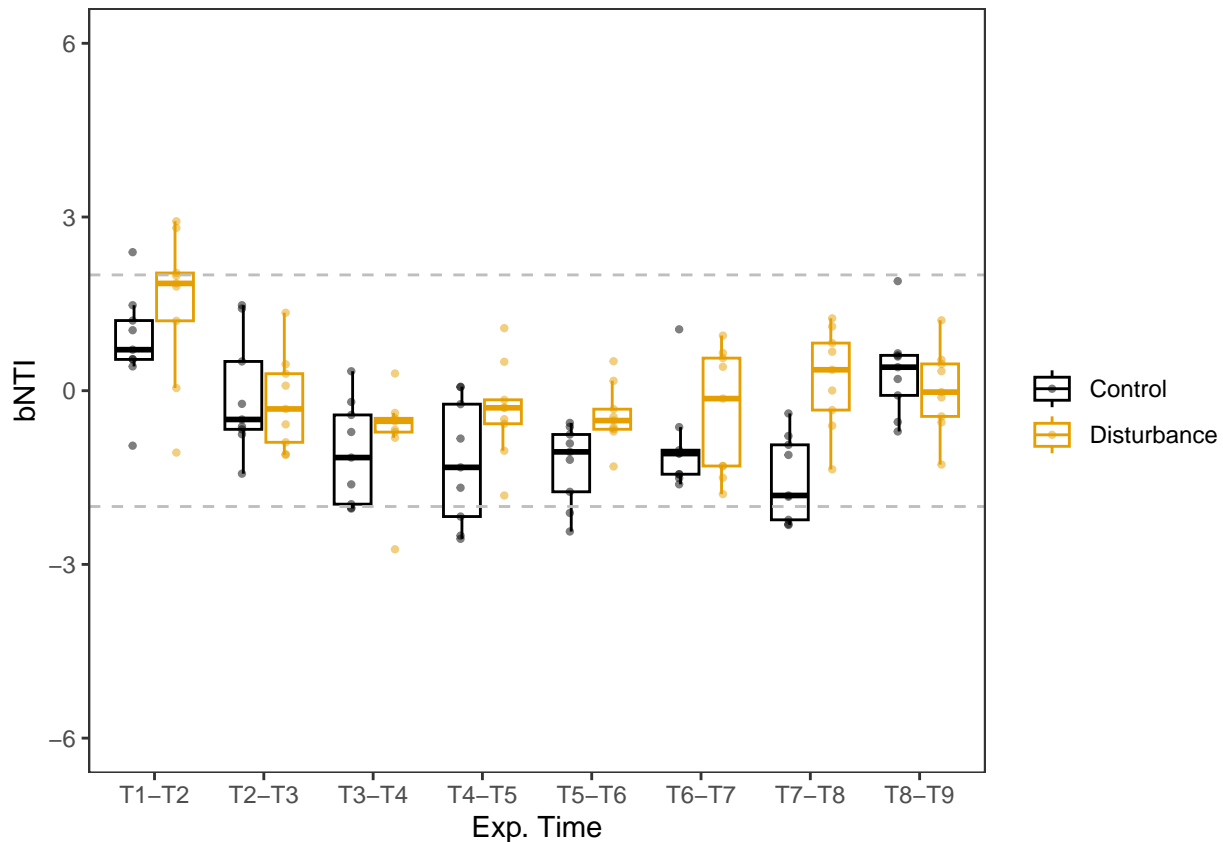
The bNTI evaluates whether the phylogenetic similarity between a pair of sample is significantly lower or higher than expected by chance relative to a reference species pool. Phylogenetic similarity surpassing the theoretical expectation ($bNTI > 2$) indicates the prevalence of variable deterministic assembly processes during community succession. Phylogenetic similarity below the theoretical expectation ($bNTI < -2$), indicates the prevalence of homogeneous deterministic assembly processes during community succession. bNTIs between -2 and 2 indicate that community assembly is driven by stochastic rather than a deterministic processes.

```
# LDOM
plot.NTI.LDOM = df.NTI.all[df.NTI.all$DOM == "LDOM", ] %>%
  ggplot(aes((t2), dist, colour = (Treatment))) + geom_boxplot(outlier.size = -1,
    alpha = 0.5, size = 0.5) + geom_jitter(aes(group = interaction(t2, Treatment)),
    position = position_dodge(0.8), alpha = 0.5, size = 0.8) + ylab(" bNTI ") +
  xlab("Exp. Time") + scale_color_manual(values = cbbPalette, name = "") +
  theme_bw() + ylim(-6, 6) + geom_hline(aes(yintercept = -2), color = "grey",
    linetype = "dashed", size = 0.5) + geom_hline(aes(yintercept = 2), color = "grey",
    linetype = "dashed", size = 0.5) + theme(panel.grid.major = element_blank(),
    panel.grid.minor = element_blank()) + scale_x_discrete(labels = c(`2` = "T1-T2",
```

```
`3` = "T2-T3", `4` = "T3-T4", `5` = "T4-T5", `6` = "T5-T6", `7` = "T6-T7",  
`8` = "T7-T8", `9` = "T8-T9"))
```

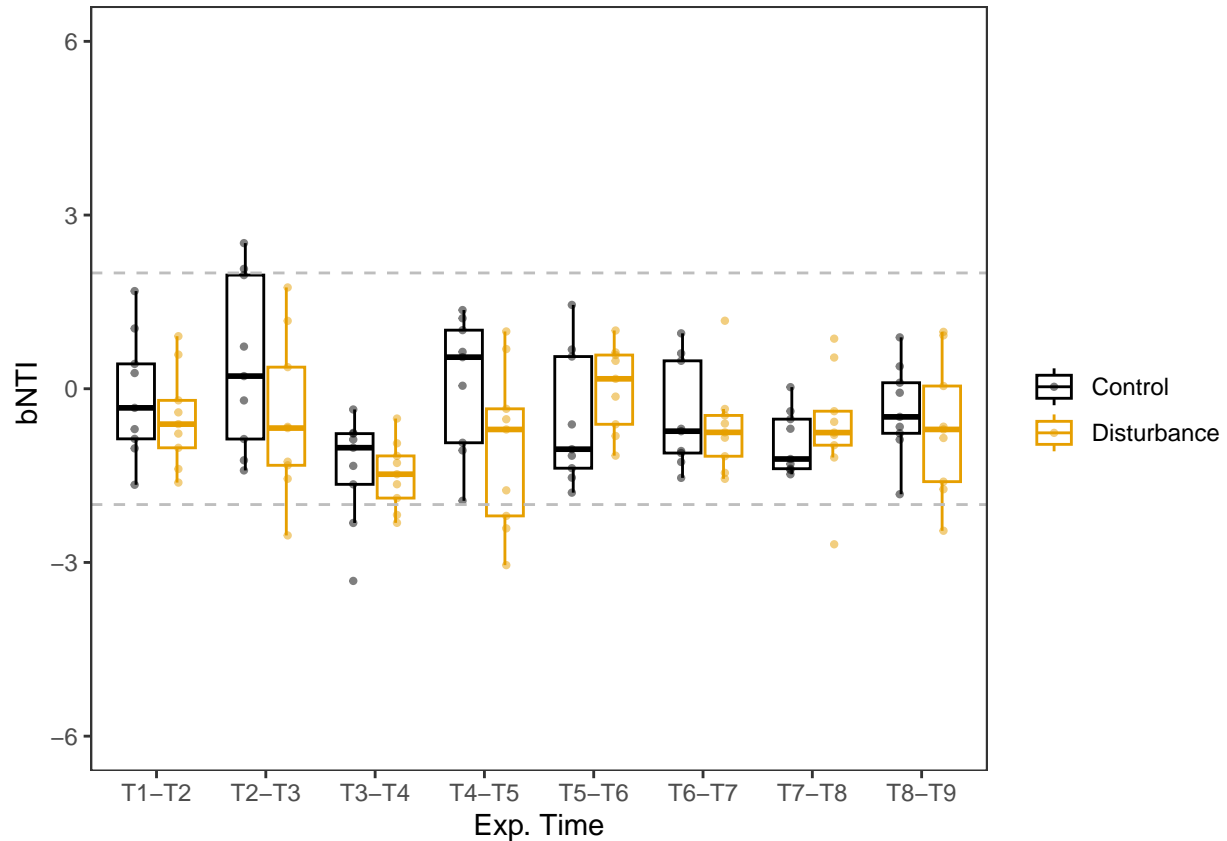
```
## Warning: Using 'size' aesthetic for lines was deprecated in ggplot2 3.4.0.  
## i Please use 'linewidth' instead.
```

```
plot.NTI.LDOM
```



```
# HDOM  
plot.NTI.HDOM = df.NTI.all[df.NTI.all$DOM == "HDOM", ] %>%  
  ggplot(aes((t2), dist, colour = (Treatment))) + geom_boxplot(outlier.size = -1,  
    alpha = 0.5, size = 0.5) + geom_jitter(aes(group = interaction(t2, Treatment)),  
    position = position_dodge(0.8), alpha = 0.5, size = 0.8) + ylab("bNTI ") +  
  xlab("Exp. Time") + scale_color_manual(values = cbbPalette, name = "") +  
  theme_bw() + ylim(-6, 6) + geom_hline(aes(yintercept = -2), color = "grey",  
    linetype = "dashed", size = 0.5) + geom_hline(aes(yintercept = 2), color = "grey",  
    linetype = "dashed", size = 0.5) + theme(panel.grid.major = element_blank(),  
    panel.grid.minor = element_blank()) + scale_x_discrete(labels = c(`2` = "T1-T2",  
    `3` = "T2-T3", `4` = "T3-T4", `5` = "T4-T5", `6` = "T5-T6", `7` = "T6-T7",  
    `8` = "T7-T8", `9` = "T8-T9"))
```

```
plot.NTI.HDOM
```



2.6 Export figure

```
pdf("../figures/FigureS3_betaNTI.pdf", width = 7, height = 5)
plot_grid(plot.NTI.LDOM, plot.NTI.HDOM, ncol = 1, labels = c("LDOM", "HDOM"),
  label_x = 0.07, label_y = 0.97, label_size = 12)
dev.off()
```

```
## pdf
## 2
```

2.7 Statistical analysis

```
df.NTI.all$Rep = rep(c(1, 2, 3), each = 3)
df.NTI.all$DOM = as.factor(df.NTI.all$DOM)
df.NTI.all$Treatment = as.factor(df.NTI.all$Treatment)
df.NTI.all$t2 = as.factor(df.NTI.all$t2)

# Testing assumptions Normality
df.NTI.all %>%
  group_by(DOM, Treatment, t2) %>%
  shapiro_test(dist)
```

```
## # A tibble: 32 x 6
##   t2    DOM Treatment variable statistic    p
##   <fct> <fct> <fct>    <chr>      <dbl> <dbl>
## 1 2    HDOM Control    dist      0.972 0.911
## 2 3    HDOM Control    dist      0.915 0.354
## 3 4    HDOM Control    dist      0.881 0.162
## 4 5    HDOM Control    dist      0.908 0.304
## 5 6    HDOM Control    dist      0.889 0.194
## 6 7    HDOM Control    dist      0.871 0.126
## 7 8    HDOM Control    dist      0.876 0.143
## 8 9    HDOM Control    dist      0.981 0.967
## 9 2    HDOM Disturbance dist      0.955 0.746
## 10 3   HDOM Disturbance dist      0.951 0.702
## # ... with 22 more rows
```

Homogeneity of variances

```
df.NTI.all %>%
  group_by(t2) %>%
  levene_test(dist ~ DOM * Treatment)
```

```
## # A tibble: 8 x 5
##   t2    df1 df2 statistic    p
##   <fct> <int> <int>      <dbl> <dbl>
## 1 2      3  32    0.325 0.807
## 2 3      3  32    1.19 0.331
## 3 4      3  32    0.606 0.616
## 4 5      3  32    1.07 0.375
## 5 6      3  32    1.39 0.265
## 6 7      3  32    1.13 0.351
## 7 8      3  32    0.389 0.762
## 8 9      3  32    0.784 0.512
```

Repeated measurement ANOVA

```
summary(aov(dist ~ DOM * Treatment * t2 + Error(Rep), data = df.NTI.all))
```

```
##
## Error: Rep
##           Df Sum Sq Mean Sq F value Pr(>F)
## Residuals  1 0.3936  0.3936
##
## Error: Within
##           Df Sum Sq Mean Sq F value    Pr(>F)
## DOM         1   3.39   3.393   3.659 0.056902 .
## Treatment   1   1.68   1.681   1.812 0.179434
## t2          7  58.90   8.415   9.073 5.32e-10 ***
## DOM:Treatment 1  13.41  13.412  14.461 0.000179 ***
## DOM:t2       7  27.63   3.947   4.255 0.000183 ***
## Treatment:t2  7  15.91   2.272   2.450 0.019007 *
## DOM:Treatment:t2 7   6.88   0.983   1.060 0.390186
## Residuals    255 236.50   0.927
## ---
## Signif. codes:  0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
```