01 Community description patterns

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${\bf Contents}$

1	Setting up the workspace					
	1.1	Loading Packages	2			
	1.2	Loading colorblind palette	2			
	1.3	Load OTU table and metadata	2			
2	Community patterns					
	2.1	Description of microcosms community composition at Genus level				
	2.2	Relative read number reads over the experiment (Figure 1b ms)				
	2.3	Patterns of diversity	6			
		2.3.1 Alpha and beta diversity over the incubation cycles	6			
		2.3.2 Calculation Raup–Crick (bRC) index	7			
	2.4	Get communities with common and abundant genera	7			
		2.4.1 Prepare RC figures	Ć			
		2.4.2 Calculate proportions of pairwise bRC	ć			
		2.4.3 Export Alpha and beta-diversity (Figure 2 ms)	10			
	2.5	Compare pairwise compositional dissimilarity (Bray Curtis-based) between C0 and C6 \ldots .	11			
		2.5.1 Comparison Bray Curtis dissimilarity at C0 and C6 cycles	12			
3	Des	scription of community types	12			
	3.1	Veen diagram	12			
	3.2	2 Heatmap Cycle 6 community types				
	3.3	Compare dominant genus from the community clusters	14			
	3.4	Alpha diversity patterns				
	3.5	Export figure 3 manuscript	16			

1 Setting up the workspace

1.1 Loading Packages

```
rm(list = ls())
library(cowplot)
library(vegan)
library(egg)
library(readxl)
library(FactoMineR)
library(factoextra)
library(RColorBrewer) #Expand color palette
library(dplyr)
library(stringr) # For editing string
library(reshape2) #For 'melt' function
library(ggsignif) #Significance values for gaplot
library(viridis) #Nice green and yellow color palette
library(rstatix)
library(olsrr)
library(nlme)
library(pheatmap)
library(NST)
library(tinytex)
library(kableExtra)
library(ggnewscale)
library(ggvenn) # Venn diagram
```

1.2 Loading colorblind palette

1.3 Load OTU table and metadata

```
levels(meta$time) = c(0, 1, 4, 6, 7) #Cycle number
meta$time = as.numeric(as.character(meta$time))
meta$Replicate = as.factor(sprintf("%02d", meta$Replicate)) # Add Microcosm ID
tibble(meta)
```

```
## # A tibble: 100 x 4
##
     Cycle Replicate M.ID
                            time
                     <chr> <dbl>
##
      <chr> <fct>
  1 CO
           01
                     C01
## 2 CO
                     C010
           10
                               0
## 3 CO
                     C011
           11
                               0
## 4 CO
                     C012
          12
                               0
## 5 CO
         13
                     C013
                               0
## 6 CO
           14
                     C014
                               0
## 7 CO
           15
                     C015
                               0
                     C016
                               0
## 8 CO
           16
## 9 CO
           17
                     C017
                               0
## 10 CO
           18
                     C018
                               0
## # i 90 more rows
```

2 Community patterns

2.1 Description of microcosms community composition at Genus level

```
# Overview at Genus level
df2 <- aggregate(. ~ Genus, data = df1[, c(6, 8:107)], sum, na.rm = TRUE)
genus <- df2[, 1] #Create a vector with Genus
rownames(df2) <- df2[, 1] #Add it as rownames
df2 <- df2[, -1] #remove column with orders and keep only abundance data
# New object with the aggregated reads
agg = df2
# Change format of agg matrix to dataframe
agg <- as.data.frame(agg)</pre>
rownames(agg) <- genus #add genus information
agg$Sum.agg <- rowSums(agg) # add column with counts across treatments for all genera
# Summarizing the most abundant genera Get the more
# abundant genera at Cycle 7
agg$Sum.agg10T <- rowSums(agg[, meta$M.ID[meta$Cycle == "C7"]])
## Get the more abundant genera from Cycle 0 to Cycle 6
agg$Sum.agg10C <- rowSums(agg[, meta$M.ID[meta$Cycle != "C7"]])
# select the most abundant genera from the full experiment
agg10T <- agg[with(agg, order(-Sum.agg10T)), ][1:11, 1:100]
agg10C <- agg[with(agg, order(-Sum.agg10C)), ][1:20, 1:100]
# Combine both dataframes avoiding duplicates
```

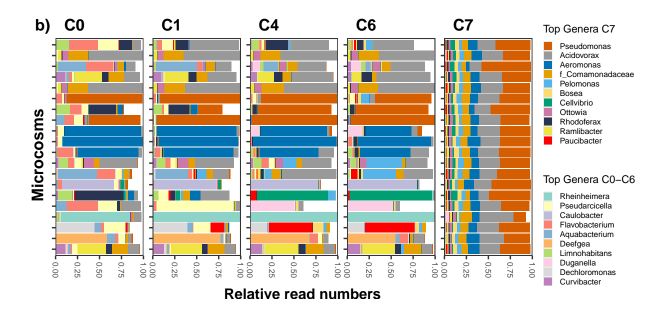
```
duprows <- which(!is.na(match(rownames(agg10C), rownames(agg10T))))</pre>
agg10combined <- rbind(agg10T, agg10C[-duprows, ])
agg10combined$genus <- rownames(agg10combined)</pre>
# convert from wide to long format
agg10.long <- melt(agg10combined, id.vars = "genus", variable.name = "treat")</pre>
agg10.long$M.ID = str remove(agg10.long$treat, "V1.")
# Combine metadata with results
agg10.long = merge(agg10.long, meta, by = "M.ID")
# Reformat Replicate with 02 digits
agg10.long$Replicate = sprintf("%02d", as.numeric(agg10.long$Replicate))
agg10.long$Replicate = as.factor(agg10.long$Replicate)
## Set Genus as factor
agg10.long$order = factor(agg10.long$genus, row.names(agg10combined))
# Reorder factor levels according C6 microcosms
agg10.long$Replicate = factor(agg10.long$Replicate, c(c("04",
    "05", "08", "11", "12", "16", "17", "10", "18"), c("01",
    "03", "07"), c("02", "13", "14"), c("06", "09", "15", "19",
    "20")))
# Classify them
agg10.long$Class = "NA"
agg10.long$Class[which(agg10.long$order %in% rownames(agg10T))] = "Top10"
agg10.long$Class[(agg10.long$Class) == "NA"] = "Others"
# set column Class as factor
agg10.long$Class <- as.factor(agg10.long$Class)</pre>
tibble(agg10.long)
## # A tibble: 2,100 x 9
##
     M.ID genus
                                      value Cycle Replicate time order
                                                                              Class
                            treat
      <chr> <chr>
                                      <dbl> <chr> <fct>
                                                            <dbl> <fct>
                                                                              <fct>
##
                            <fct>
## 1 CO1
          Pseudomonas
                            CO1 0
                                            CO
                                                  01
                                                                0 Pseudomonas Top10
## 2 CO1
                                            CO
                                                  01
          Acidovorax
                            CO1 0.0405
                                                                0 Acidovorax Top10
## 3 CO1 Aeromonas
                            CO1 0.695
                                            CO
                                                  01
                                                                O Aeromonas
                                                                              Top10
## 4 CO1
          f_Comamonadaceae CO1 0.00756 CO
                                                                0 f_Comamona~ Top10
                                                  01
## 5 CO1
           Pelomonas
                            C01
                                  Ω
                                            CO
                                                  01
                                                                O Pelomonas
                                                                              Top10
## 6 CO1
           Bosea
                            C01 0
                                            CO
                                                  01
                                                                0 Bosea
                                                                              Top10
## 7 CO1
           Cellvibrio
                            C01
                                            CO
                                                  01
                                                                0 Cellvibrio Top10
                                  Ω
## 8 CO1
           Ottowia
                            C01
                                  0.000344 CO
                                                  01
                                                                O Ottowia
                                                                              Top10
## 9 CO1
           Rhodoferax
                            C01
                                  0.0113
                                           CO
                                                  01
                                                                0 Rhodoferax Top10
## 10 CO1
           Ramlibacter
                            C01
                                  0.00550 CO
                                                  01
                                                                O Ramlibacter Top10
## # i 2,090 more rows
# loop to plot microbial relative abundance per cycle
list.plots <- list()</pre>
cycles.index <- c("CO", "C1", "C4", "C6", "C7")
```

```
for (i in 1:length(cycles.index)) {
    list.plots[[i]] <- ggplot(agg10.long %>%
        subset(Cycle == cycles.index[i]), aes(x = Replicate,
        y = value, fill = order)) + geom_bar(stat = "identity") +
        scale_fill_manual(values = c(cbp1, mycolors2)) + scale_y_continuous(limits = c(0,
        1), expand = c(0, 0)) + guides(fill = guide_legend(override.aes = list(size = 1))) +
        theme(legend.text = element_text(size = 5)) + guides(fill = guide_legend(ncol = 1)) +
        labs(title = "", x = NULL, y = "") + my_theme + coord_flip() +
        theme(legend.position = "none")
}
# Temporal plot to create legends
plot_tmp <- ggplot() + geom_bar(data = agg10.long %>%
    subset(Cycle == "C7"), aes(x = Replicate, y = value, fill = order),
    stat = "identity") + scale_fill_manual(name = "Top Genera C7",
   values = c(cbp1, mycolors2), breaks = levels(agg10.long$order)[1:11]) +
   new_scale_fill() + geom_bar(data = agg10.long %>%
    subset(Cycle == "C7"), aes(x = Replicate, y = value, fill = order),
    stat = "identity") + scale_fill_manual(name = "Top Genera CO-C6",
   values = c(mycolors2, mycolors2), breaks = levels(agg10.long$order)[12:22]) +
    theme(legend.key.width = unit(0.25, "cm"), legend.key.height = unit(0.25,
        "cm"), legend.key = element_rect(fill = NULL, colour = NULL),
        legend.text = element_text(size = 6), legend.title = element_text(size = 8)) +
    guides(fill = guide_legend(ncol = 1))
# Extract legend
legend.plot <- get_legend(plot_tmp)</pre>
rm(plot_tmp)
```

2.2 Relative read number reads over the experiment (Figure 1b ms)

Proportions of reads of 16S rRNA genes affiliated with the most abundant genera in metagenomes from the experimental communities at C0, C1, C4, C6 and C7.

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



2.3 Patterns of diversity

2.3.1 Alpha and beta diversity over the incubation cycles

```
### Shannon diversity index
meta$shannon = vegan::diversity(t(df1[, 8:107]))
### Richness
meta$Richness = colSums(df1[, 8:107] != 0)

## Evenness (Pielou)
meta$Even = meta$shannon/log(meta$Richness)
tibble(meta)
```

```
## # A tibble: 100 x 7
##
      Cycle Replicate M.ID
                               time shannon Richness Even
      <chr> <fct>
                        <chr> <dbl>
                                                 <dbl> <dbl>
##
                                       <dbl>
##
    1 CO
             01
                        C01
                                        2.97
                                                   116 0.626
                                   0
##
    2 CO
             10
                        C010
                                   0
                                        3.57
                                                   176 0.690
##
    3 CO
             11
                        C011
                                   0
                                        2.71
                                                    72 0.633
##
    4 CO
             12
                        C012
                                   0
                                        3.47
                                                   125 0.719
##
                                        3.43
    5 CO
             13
                        C013
                                   0
                                                   107 0.735
##
    6 CO
             14
                        C014
                                   0
                                        3.06
                                                   113 0.648
##
    7
      CO
             15
                        C015
                                   0
                                        3.82
                                                   149 0.763
##
    8 CO
             16
                        C016
                                  0
                                        3.02
                                                   122 0.629
##
   9 CO
             17
                        C017
                                   0
                                        3.27
                                                   123 0.680
## 10 CO
             18
                        C018
                                   0
                                        3.18
                                                    85 0.716
## # i 90 more rows
```

```
sum(rowSums(df1[, 68:87] != 0) > 0), sum(rowSums(df1[,
            88:107] != 0) > 0)))
tibble(df.rich)
## # A tibble: 5 x 2
##
     Cycle nOTUS
##
     <chr> <int>
## 1 CO
             722
## 2 C1
             654
## 3 C4
             709
## 4 C6
             737
## 5 C7
             469
```

2.3.2 Calculation Raup-Crick (bRC) index

2.4 Get communities with common and abundant genera

#Common dominant taxa are here operationally defined as the genus with highest relative abundance in a community and high co-occurrency (>3) following Avolio et al., 2019.

```
df.order <- df2[order(rowSums(df2[, 61:80]), decreasing = TRUE),
      61:80]
df.dominance <- data.frame(sample.ID = colnames(df.order), most_abundant = rownames(df.order)[max.col(t tibble(df.dominance)</pre>
```

```
## # A tibble: 20 x 2
##
      sample.ID most_abundant
##
      <chr>
                <chr>
##
   1 C61
                Aeromonas
##
    2 C610
                Pelomonas
    3 C611
##
                Rheinheimera
##
  4 C612
                Duganella
##
  5 C613
                Pseudomonas
## 6 C614
                Pseudomonas
##
  7 C615
                Acidovorax
  8 C616
##
                Cellvibrio
  9 C617
                Caulobacter
## 10 C618
                Pelomonas
## 11 C619
                Acidovorax
## 12 C62
                Pseudomonas
## 13 C620
                Acidovorax
## 14 C63
                Aeromonas
## 15 C64
                Ramlibacter
## 16 C65
                Deefgea
                Acidovorax
## 17 C66
## 18 C67
                Aeromonas
## 19 C68
                Paucibacter
## 20 C69
                Acidovorax
```

```
# Create a dataframe with community type labels
labels2 <- data.frame(labels = levels(meta$Replicate), cluster = NA)</pre>
# Indicate clusters identity
labels2$cluster[labels2$labels %in% c("01", "03", "07")] = "Aeromonas"
labels2$cluster[labels2$labels %in% c("06", "09", "15", "19",
    "20")] = "Acidovorax"
labels2$cluster[labels2$labels %in% c("02", "13", "14")] = "Pseudomonas"
labels2$cluster[labels2$labels %in% c("04", "05", "08", "10",
    "11", "12", "16", "17", "18")] = "Others"
# Combine cluster labels with metadata
meta = merge(meta, labels2, by.x = "Replicate", by.y = "labels")
meta$cluster = as.factor(meta$cluster)
meta$cluster = factor(meta$cluster, c("Acidovorax", "Aeromonas",
    "Pseudomonas", "Others"))
tibble(meta)
## # A tibble: 100 x 8
     Replicate Cycle M.ID
##
                          time shannon Richness Even cluster
##
               <chr> <chr> <dbl>
                                   <dbl> <dbl> <fct>
     <fct>
## 1 01
               CO
                     C01
                              0 2.97
                                            116 0.626 Aeromonas
## 2 01
               C4
                     C41
                               4 2.80
                                             92 0.620 Aeromonas
## 3 01
              C7
                    C71
                              7
                                   4.21
                                             219 0.782 Aeromonas
              C1 C11
                                            93 0.570 Aeromonas
## 4 01
                              1 2.58
## 5 01
              C6
                              6 2.99
                     C61
                                             91 0.663 Aeromonas
              C7
                                            226 0.775 Pseudomonas
## 6 02
                    C72
                              7 4.20
## 7 02
               C6
                   C62
                              6 2.95
                                             58 0.727 Pseudomonas
## 8 02
               C4 C42
                              4 2.80
                                             48 0.724 Pseudomonas
## 9 02
               C1
                    C12
                             1 2.93
                                             63 0.706 Pseudomonas
                                            143 0.780 Pseudomonas
## 10 02
               CO
                               0 3.87
                     C02
## # i 90 more rows
dist.method = "bray" # Select distance method
comm <- df1[, 8:107]
TaxonID <- row.names(comm)</pre>
comm \leftarrow comm[-1,]
meta <- meta[order(meta$M.ID), ]</pre>
meta.groupi <- data.frame(cluster = meta[, c("cluster")])</pre>
row.names(meta.groupi) <- meta$M.ID</pre>
tnst = tNST(comm = t(comm), meta.com = NULL, group = meta.groupi,
   dist.method = "bray", abundance.weighted = TRUE, rand = 1000,
   output.rand = TRUE, nworker = 8, LB = FALSE, null.model = "PF",
   between.group = TRUE, SES = TRUE, RC = TRUE)
## All match very well.
## Now randomizing by parallel computing. Begin at Thu Jul 4 09:51:54 2024. Please wait...
```

Organize Raup-Crick (bRC) data

2.4.1 Prepare RC figures

```
set.seed(666)
plot_RC <- nst.temp[nst.temp$Cycle1 == nst.temp$Cycle2 & nst.temp$group !=</pre>
    "Others" & nst.temp$Cycle1 != "C7", ] %>%
    ggplot(aes(x = Cycle1, y = RC.ij.bray)) + geom_violin(aes(fill = group),
   alpha = 0.3, size = 0.5) + geom_jitter(aes(fill = group),
   width = 0.2, alpha = 0.7, size = 1.5, shape = 21) + geom_hline(yintercept = c(-0.95,
    0.95), linetype = "dashed") + theme_bw() + theme(panel.grid.minor = element_blank(),
    panel.grid.major = element_blank()) + labs(y = expression(beta *
    "RC"), x = \text{"Cycle"}) + ylim(-1.05, 1.1) + scale_fill_manual(values = c("#999999",
   "#D55E00", "#0072B2")) + theme(plot.margin = unit(c(0.1,
   0.2, 0, 0), "cm")) + theme(plot.title = element_text(vjust = -1,
    size = 8), text = element_text(size = 8.5), axis.title.y = element_text(size = 8.5,
    vjust = -2)) + facet_wrap(~group, nrow = 1) + theme(legend.position = "none")
plot RC <- tag facet(plot RC, tag pool = c("Acidovorax-type",</pre>
    "Pseudomonas-type", "Aeromonas-type"), hjust = -0.1, open = "",
    close = "", fontface = 3, size = 2.8)
```

2.4.2 Calculate proportions of pairwise bRC

```
data.bRC = nst.temp[nst.temp$Cycle1 == nst.temp$Cycle2, ]
data.bRC$category <- "bRC<|0.95|"
data.bRC$category[data.bRC$RC.ij.bray > 0.95] <- "bRC>0.95"
data.bRC$category[data.bRC$RC.ij.bray < (-0.95)] <- "bRC<-0.95"

agg.bRC <- aggregate(RC.ij.bray ~ category + Cycle1, data = data.bRC,
    length)
agg.bRC$category <- as.factor(agg.bRC$category) #Set it as factor
agg.bRC$category <- factor(agg.bRC$category, c("bRC>0.95", "bRC<|0.95|",
    "bRC<-0.95")) #Reorder factor levels

agg.bRC$norm.count <- agg.bRC$RC.ij.bray/52

prop.bRC <- agg.bRC %>%
    ggplot(aes(x = Cycle1, y = norm.count, fill = category)) +
    geom_bar(stat = "identity") + scale_fill_manual(values = c(cbp1[6:9]),
    name = "", labels = c(expression(beta * "RC >0.95"), expression(beta *
```

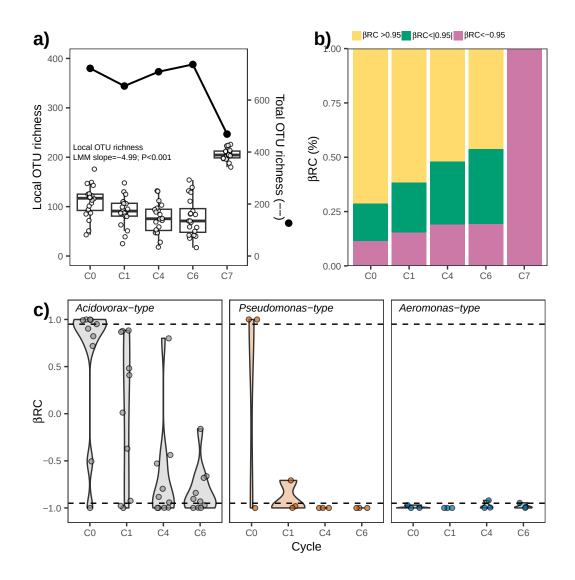
```
"RC<|0.95|"), expression(beta * "RC<-0.95"))) + scale_y_continuous(limits = c(0,
1), expand = c(0, 0)) + labs(title = "", x = NULL, y = expression(beta *
"RC (%) ")) + theme_bw() + theme(text = element_text(size = 8.5),
axis.title.y = element_text(size = 8.5, vjust = -0.05)) +
theme(panel.grid.minor = element_blank(), panel.grid.major = element_blank(),
    legend.position = c(0.35, 1.06), legend.direction = "horizontal",
    legend.title = element_blank(), legend.text = element_text(size = 5.5,
        margin = margin(0, -0.2, 0, 0.01, unit = "cm")),
    legend.margin = margin(-10, -20, -10, -10), legend.key.size = unit(0.3,
        "cm")) + theme(plot.margin = unit(c(0.1, 0.15, 0.1,
0), "cm"))</pre>
```

```
## Warning: A numeric 'legend.position' argument in 'theme()' was deprecated in ggplot2
## 3.5.0.
## i Please use the 'legend.position.inside' argument of 'theme()' instead.
## This warning is displayed once every 8 hours.
## Call 'lifecycle::last_lifecycle_warnings()' to see where this warning was
## generated.
```

2.4.3 Export Alpha and beta-diversity (Figure 2 ms)

OTU richness (a) per microcosms (left axis) and total OTU richness across microcosms (right axis) over cycles. b) Percentage of Raup-Crick (bRC) pairwise dissimilarity distances among all microcosms. c) Per identified community-common genera cluster during cycles.

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



2.5 Compare pairwise compositional dissimilarity (Bray Curtis-based) between C0 and C6

```
set.seed(666)
mantel(data.dist0, data.dist6, permutations = 1000)

##

## Mantel statistic based on Pearson's product-moment correlation

##

## Call:
## mantel(xdis = data.dist0, ydis = data.dist6, permutations = 1000)

##

## Mantel statistic r: 0.5542

## Significance: 0.000999

##

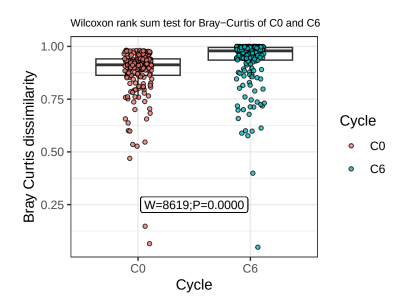
## Upper quantiles of permutations (null model):
## 90% 95% 97.5% 99%

## 0.105 0.140 0.189 0.252

## Permutation: free
```

2.5.1 Comparison Bray Curtis dissimilarity at C0 and C6 cycles

Number of permutations: 1000



3 Description of community types

3.1 Veen diagram

```
# Subset data from C6
df.C6 <- df2[, c(61:80)]

# Genus taxonomic classification
my_genus_col = data.frame(df1[, c(5, 6)])
my_genus_col <- unique(my_genus_col)</pre>
```

```
row.names(my_genus_col) <- my_genus_col$Genus</pre>
# Remove not present taxa
df.C6 <- df.C6[rowSums(df.C6) > 0, ] # Reduced from 119 to 85 Genera
# Average by community type
acido = rowMeans(df.C6[, meta$M.ID[meta$Cycle == "C6" & meta$cluster ==
    "Acidovorax"]])
pseu = rowMeans(df.C6[, meta$M.ID[meta$Cycle == "C6" & meta$cluster ==
    "Pseudomonas"]])
aero = rowMeans(df.C6[, meta$M.ID[meta$Cycle == "C6" & meta$cluster ==
    "Aeromonas"]])
othe = rowMeans(df.C6[, meta$M.ID[meta$Cycle == "C6" & meta$cluster ==
    "Others"]])
df.simple <- data.frame(acido, pseu, aero, othe)</pre>
# Combining samples per cluster Get only data from Cycle 6
# for Venn diagram and Heatmap
df.simple.cluster <- df.simple[, c("acido", "pseu", "aero")]</pre>
# Remove the uniques uniques taxa from the. 'Others
# communities
df.simple.cluster <- df.simple.cluster[rowSums(df.simple.cluster) >
    0, ]
df.simple.cluster <- as.data.frame(df.simple.cluster > 0)
names(df.simple.cluster) <- c("Acidovorax-type", "Pseudomonas-type",</pre>
    "Aeromonas-type")
# veen diagram plot (save for later)
venn_plot <- ggvenn(df.simple.cluster, fill_color = c("#868686FF",</pre>
    "#fc6f00", "#0073C2FF", "#f0f2f2"), stroke_size = 0.25, set_name_size = 1.7,
    show_percentage = F, text_size = 2)
```

3.2 Heatmap Cycle 6 community types

```
# Rename Allorhizobium-Neorhizobium-Pararhizobium-Rhizobium
# for ANPR
row.names(df.simple.log10)[27] <- "ANPR"</pre>
col.for.labels$genus[27] <- "ANPR"</pre>
names(df.simple.log10) <- c("Acidovorax-type", "Pseudomonas-type",</pre>
    "Aeromonas-type")
## create the breaks to state 0 as white
bk2 = seq(-4, -0.09119, length = 70)
# set different color vectors for each interval
col1 = "white" #set the order of greys
col2 = viridis(69)
colors2 <- c(col1, col2)</pre>
my_heatmap <- pheatmap::pheatmap(t(df.simple.log10), color = colors2,</pre>
    breaks = bk2, annotation_colors = NA, angle_col = 45, fontsize = 4.7,
    legend = TRUE, cluster_rows = F, cluster_cols = F, cellwidth = 5.4,
    cellheight = 7, margin = c(0, 0, 0, 0), labels_col = paste0("f",
        c(1:70)))
```

3.3 Compare dominant genus from the community clusters

3.4 Alpha diversity patterns

```
df.rmANOVA <- data.frame(Variable = c("Richness", "Pielou evenness"),
    Normality = NA, HOV = NA, Comm.type.pvalue = NA, Cycle.pvalue = NA)
# ANOVAs
ref.anova.rich = aov(Richness ~ cluster * Cycle + Error(Replicate),
    data = meta[meta$Cycle != "C7", ])
ref.anova.even = aov(Even ~ cluster * Cycle + Error(Replicate),
    data = meta[meta$Cycle != "C7", ])</pre>
```

	Assumptions		${ m rmANOVA}$	
Variable	Normality	HOV	Comm.type.pvalue	Cycle.pvalue
Richness	TRUE	TRUE	0.7938670	0.0000002
Pielou evenness	TRUE	TRUE	0.0070899	0.0446801

```
# Alpha-diversity plots for final panel
plot.Richness = meta[meta$Cycle != "C7", ] %>%
    ggplot(aes(cluster, Richness, fill = Cycle)) + geom_boxplot(outlier.shape = NA,
    alpha = 0.5, size = 0.2) + xlab("") + ylab("OTU Richness") +
    geom point(position = position jitterdodge(jitter.width = 0.15),
        shape = 21, size = 0.5, stroke = 0.1) + theme_bw() +
    scale fill manual(values = c(cbp1[c(6, 7, 8, 10)]), name = "") +
    theme(axis.title.y = element_text(size = 6), text = element_text(size = 6),
        axis.text.x = element_blank(), panel.grid.minor = element_blank(),
       panel.grid.major = element_blank()) + theme(legend.position = "top",
    legend.text = element_text(size = 5.5, margin = margin(0,
        -0.2, 0, 0.01, unit = "cm")), legend.key.size = unit(1,
        "lines"), legend.box = "horizontal") + theme(plot.margin = unit(c(-0.5,
    0.1, -0.25, 0.15), "cm"), axis.text.y = element_text(size = 5),
    legend.box = "horizontal", legend.margin = margin(0, 0, -10,
        -10))
plot.Evennes = meta[meta$Cycle != "C7", ] %>%
    ggplot(aes(cluster, Even, fill = Cycle)) + geom_boxplot(outlier.shape = NA,
    alpha = 0.5, size = 0.2) + xlab("") + ylab("Pielou Evennes") +
    geom_point(position = position_jitterdodge(jitter.width = 0.15),
        shape = 21, size = 0.5, stroke = 0.1) + ylim(0.25, 1) +
    theme_bw() + theme(legend.position = "none") + theme(axis.title.y = element_text(size = 6),
```

```
text = element_text(size = 6), axis.text.x = element_text(angle = 13,
    hjust = 0.5), panel.grid.minor = element_blank(), panel.grid.major = element_blank()) +
scale_fill_manual(values = c(cbp1[c(6, 7, 8, 10)]), name = "") +
theme(legend.position = "none") + geom_signif(comparisons = list(c("Acidovorax",
    "Aeromonas"), c("Pseudomonas", "Aeromonas"), c("Others",
    "Aeromonas")), map_signif_level = TRUE, annotations = c("***",
    "***", "***"), y_position = c(0.78, 0.82, 0.87), textsize = 2.5,
    vjust = 0.5, col = "black", step_increase = 0.05, tip_length = 0.01,
    orientation = "x", size = 0.17) + theme(plot.margin = unit(c(-0.05,
    0.1, -0.45, 0.15), "cm"))
```

3.5 Export figure 3 manuscript

a) Community specific and shared genera in community types dominated by Acidovorax, Pseudomonas, or Aeromonas at cycle 6. b) Relative abundance of the three representative genera per community type (percentage of reads number). c) Shannon diversity (H) and Pielou Evenness indices per community type in the experimental cycles. d) Relative abundances of genera in the three community types.

