## MicrobiotaProcess: A tidy framework for microbiome or other related ecology data analysis

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# 1 Analysis of 16s rDNA dataset about 43 pediatric CD stool samples from iHMP

Here, we use the 43 pediatric IBD stool samples as example, which were obtained from the Integrative Human Microbiome Project Consortium (iHMP) (Research Network Consortium 2014).

### 1.1 Importing the output of dada2

The datasets were downloaded from web<sup>1</sup>. It contains ibd\_asv\_table.txt, which is feature table (row features X column samples), ibd\_meta.csv (metadata file of samples), and ibd\_taxa.txt (the taxonomic annotation of features). In the session, we use mp\_import\_dada2 of MicrobiotaProcess to import the dataset, and return a MPSE object.

```
library(MicrobiotaProcess)
otuda <- read.table("./data/IBD_data/ibd_asv_table.txt", header=T,
                    check.names=F, comment.char="", row.names=1, sep="\t")
# building the output format of removeBimeraDenovo of dada2
otuda <- data.frame(t(otuda), check.names=F)</pre>
sampleda <- read.csv("./data/IBD_data/ibd_meta.csv", row.names=1, comment.char="")</pre>
taxda <- read.table("./data/IBD_data/ibd_taxa.txt", header=T,</pre>
                    row.names=1, check.names=F, comment.char="")
# the feature names should be the same with rownames of taxda.
taxda <- taxda[match(colnames(otuda), rownames(taxda)),]</pre>
mpse <- mp_import_dada2(seqtab = otuda, taxatab = taxda, sampleda = sampleda)</pre>
# view the reads depth of samples and the prevalence of the OTUs. In this example,
# mpse %>% mp extract assay(.abundant=Abundance) %>% rowSums() %>% sort %>% head(100)
# mpse %>% mp_extract_assay(.abundant=Abundance) %>% colSums() %>% sort %>% head()
# head(sort(rowSums(assay(mpse, "Abundance"))), 100)
# head(sort(colSums(assay(mpse, "Abundance"))))
# In this example, we can find some OTUs have very low frequency in the samples.
# and some taxonomy are unreasonable, for example, the probability of chloroplasts
# in the intestine should be low. We can also remove the features.
mpse2 <- mpse %>%
         dplyr::filter(!Phylum %in% c("p_un_k_Bacteria", "p_Chloroflexi") &
                        !Class %in% "c__Chloroplast" &
                        !Family %in% "f__mitochondria"
         ) %>%
         mp_filter_taxa(.abundance = Abundance, min.abun = 1, min.prop = 0.1)
mpse2
```

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<sup>## #</sup> A MPSE-tibble (MPSE object) abstraction: 9,890 x 11 ## # OTU=230 | Samples=43 | Assays=Abundance | Taxanomy=Kingdom, Phylum, Class, Order, Family, Genus, Species Sample Abundance Group Kingdom Phylum Class Order Family Genus Species ## OTU ## <chr> <chr> <int> <chr> <chr> <chr> <chr> <chr> <chr> <chr> <chr> <chr> 1 OTU\_2~ S2067~ 0 CD k\_Bac~ p\_Ac~ c\_A~ o\_A~ f\_Ac~ g\_A~ s\_un\_~ 2 OTU\_5~ S2067~ 0 CD k\_Bac~ p\_Ac~ c\_A~ o\_A~ f\_Ac~ g\_A~ s\_un\_~ 3 OTU\_7~ S2067~ O CD k\_Bac~ p\_Ac~ c\_A~ o\_A~ f\_Mi~ g\_R~ s\_muc~ k\_Bac~ p\_Ac~ c\_A~ o\_B~ f\_Bi~ g\_B~ s\_ado~ 4 OTU\_42 S2067~ 0 CD

 $<sup>^{1}</sup> https://www.microbiomeanalyst.ca/MicrobiomeAnalyst/resources/data/ibd\_data.zip$ 

```
5 OTU_1~ S2067~
                         0 CD
                                k_Bac~ p_Ac~ c_A~ o_B~ f_Bi~ g_B~ s_un_~
  6 OTU_1~ S2067~
                                k_Bac~ p_Ac~ c_A~ o_B~ f_Bi~ g_B~ s_un_~
                         0 CD
   7 OTU_3~ S2067~
                         O CD
                                k_Bac~ p_Ac~ c_C~ o_C~ f_Co~ g_A~ s_un_~
## 8 OTU_1~ S2067~
                                k_Bac~ p_Ac~ c_C~ o_C~ f_Co~ g_C~ s_aer~
                         O CD
## 9 OTU 3~ S2067~
                         O CD
                                k_Bac~ p_Ac~ c_C~ o_C~ f_Co~ g_E~ s_len~
## 10 OTU_1~ S2067~
                         0 CD
                                k_Bac~ p_Ba~ c_B~ o_B~ f_[0~ g_0~ s_un_~
## # ... with 9,880 more rows
```

## 1.2 Other import functions

MicrobiotaProcess also presents some other import functions S1 to parse the output of the upstream pipelines. In addition, some common object of R can also be converted to MPSE object, such as phyloseq (McMurdie 2013), SummarizedExperiment (Morgan et al. 2021), TreeSummarizedExperiment (Huang et al. 2021), biom (McMurdie and Paulson 2021) (output of biomformat by read\_biom) 2.1.

Table S1: List of import functions provided by MicrobiotaProcess

Package	Import Function	Description
	$mp\_import\_qiime2$	Import function to load the output of qiime2
MicrobiotaProcess	$mp\_import\_qiime$	Import function to read the now legacy-format QIIME OTU table (tsv format)
	$mp\_import\_metaphlan$	Import function to read the output of MetaPhlAn

## 1.3 alpha diversity analysis

#### 1.3.1 rarefaction visualization

Rarefaction, based on sampling technique, was used to compensate for the effect of sample size on the number of units observed in a sample. MicrobiotaProcess provided  $mp\_cal\_rarecurve$  and  $mp\_plot\_rarecurve$  to calculate and plot the curves.

```
library(MicrobiotaProcess)
library(patchwork)
cols <- c("orange", "deepskyblue")</pre>
mpse2 %<>%
    mp_rrarefy(.abundance=Abundance) %>%
    mp_cal_rarecurve(.abundance=RareAbundance, chunks=500)
p_rare <- mpse2 %>%
          mp_plot_rarecurve(
            .rare = RareAbundanceRarecurve,
            .alpha = c(Observe, Chao1, ACE),
          ) +
          theme(
            legend.key.width = unit(0.3, "cm"),
            legend.key.height = unit(0.3, "cm"),
            legend.spacing.y = unit(0.01, "cm"),
            legend.text = element_text(size=4)
          )
prare1 <- mpse2 %>%
          mp_plot_rarecurve(
            .rare = RareAbundanceRarecurve,
            .alpha = c(Observe, Chao1, ACE),
            .group = Group
          scale_fill_manual(values = cols)+
          scale_color_manual(values = cols)+
          theme bw()+
```

```
theme(
           axis.text=element_text(size=8), panel.grid=element_blank(),
           strip.background = element_rect(colour=NA,fill="grey"),
           strip.text.x = element_text(face="bold")
prare2 <- mpse2 %>%
         mp_plot_rarecurve(
           .rare = RareAbundanceRarecurve,
           .alpha = c(Observe, Chao1, ACE),
           .group = Group,
           plot.group = TRUE
         ) +
         scale_color_manual(values = cols)+
         scale_fill_manual(values = cols) +
         theme_bw()+
         theme(
           axis.text=element_text(size=8), panel.grid=element_blank(),
           strip.background = element_rect(colour=NA,fill="grey"),
           strip.text.x = element_text(face="bold")
         )
(p_rare / prare1 / prare2) + patchwork::plot_annotation(tag_levels="A")
```

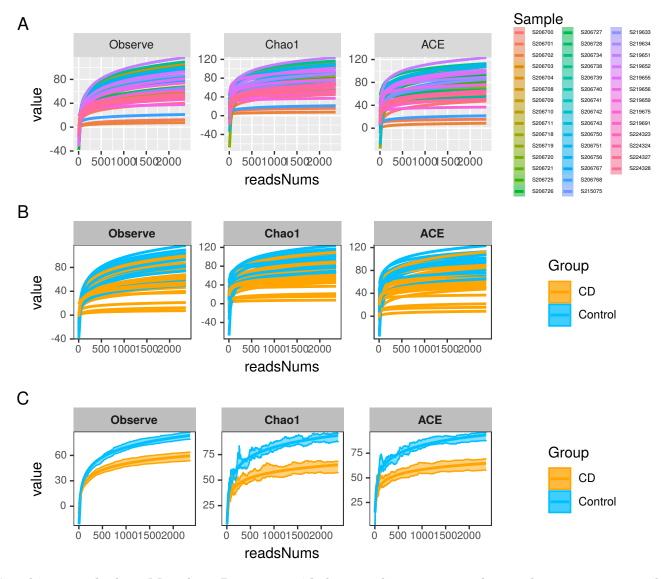


Fig. S1: This example show *MicrobiotaProcess* provided  $mp\_cal\_rarecurve$  and  $mp\_plot\_rarecurve$  to calculate and visualize the rarefaction curve. The horizontal coordinate represents the sequencing depth of samples, the vertical coordinate shows the Alpha diversity index (such as Observe OTU, Chao1 and ACE). The  $mp\_plot\_rarecurve$  provides three types of visualization. (A) the rarefaction curve for each sample. (B) the rarefaction curve for each sample with colored group (specified *.group* argument in  $mp\_plot\_rarecurve$ ). (C) the rarefaction curve for each group with standard error of the mean (specified *.group* argument and plot.group=TRUE in  $mp\_plot\_rarecurve$ )

Since the curves in each sample were near saturation, the sequencing data were great enough with very few new species undetected

#### 1.3.2 Calculation and different analysis of alpha index

Alpha index can evaluate the richness and abundance of microbial communities. MicrobiotaProcess provides  $mp\_cal\_alpha$  to calculate alpha index. Six common diversity measures (Observe, Chao1, ACE, Shannon, Simpson, Pielou) are supported. And the different groups of samples can be tested and visualize by  $mp\_plot\_alpha$ . This following example shows how to use  $mp\_cal\_alpha$  and  $mp\_plot\_alpha$  of MicrobiotaProcess to analysis the alpha diversity of the community. The RareAbundance is rarefied (default), which will be used to calculate the alpha diversity index, users can specified the force=TRUE of  $mp\_cal\_alpha$  to calculated the index if the abundance is not be rarefied (2.3.1).

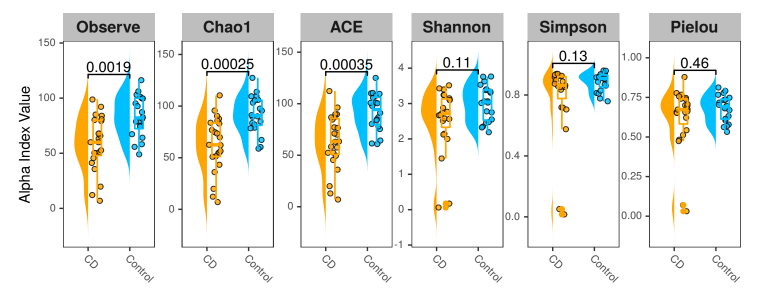


Fig. S2: The raincloud plot of alpha diversity index The horizontal coordinate represents each group (by .group argument of  $mp\_plot\_alpha$ ), the vertical coordinate represents the alpha diversity index.

## 1.4 Taxonomy composition analysis

### 1.4.1 Statistics and visualization of specific levels

MicrobiotaProcess presents the  $mp\_cal\_abundance$  and  $mp\_plot\_abundance$  for the calculation and visualization of composition of microbial communities. After the  $mp\_cal\_abundance$  done, you can get the abundance of specific levels of class by  $mp\_extract\_abundance$  1.5.4.

```
library(ggplot2)
library(MicrobiotaProcess)
# The relative abundance of all taxonomy for samples will be calculated
mpse2 %<>% mp_cal_abundance(.abundance = RareAbundance)
# The relative abundance of all taxonomy for group will be calculated
mpse2 %<>% mp_cal_abundance(.abundance = RareAbundance, .group = Group)
# The 30 most abundant taxonomy will be visualized.
pclass <- mpse2 %>%
          mp_plot_abundance(
             .abundance = RareAbundance,
             .group = Group,
             taxa.class = Class,
             topn = 30
          ) +
          xlab(NULL) +
          vlab("relative abundance (%)") +
             legend.key.width = unit(0.3, "cm"),
             legend.key.height = unit(0.3, "cm")
          ) +
          xlab(NULL) +
          ylab("relative abundance (%)") +
```

```
theme(
    legend.key.width = unit(0.3, "cm"),
    legend.key.height = unit(0.3, "cm"),
    legend.text = element_text(size=6)
)
pclass
```

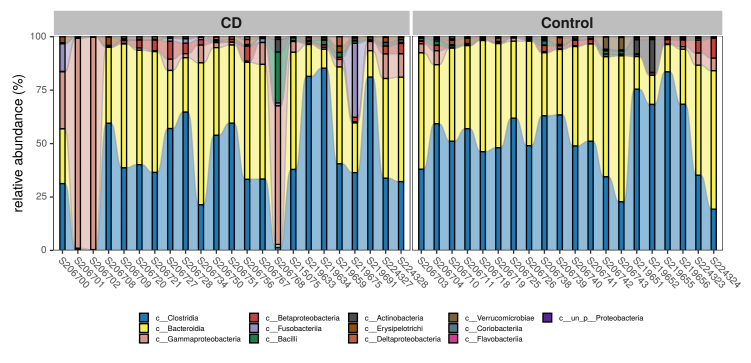


Fig. S3: The relative abundance of each sample in *class* level

The relative abundance of groups also can be visualized by providing .group argument and setting plot.group = TRUE in the  $mp\_plot\_abundance$ . If you want to view the raw abundance (count or others) of taxa, you can set the relative parameter of  $mp\_plot\_abundance$  to FALSE.

```
# Show the abundance in different groups.
fclass <- mpse2 %>%
          mp_plot_abundance(
             .abundance = RareAbundance,
             .group = Group,
             taxa.class = Class,
             topn = 30,
             plot.group = TRUE
          ) +
          xlab(NULL) +
          ylab("relative abundance (%)") +
          theme(legend.position = "none")
pclass2 <- mpse2 %>%
          mp_plot_abundance(
             .abundance = RareAbundance,
             .group = Group,
             relative = FALSE,
             taxa.class = Class,
             topn = 30
          ) +
          xlab(NULL) +
          ylab("count reads") +
          theme(legend.key.width = unit(0.3, "cm"),
                 legend.key.height = unit(0.3, "cm"),
```



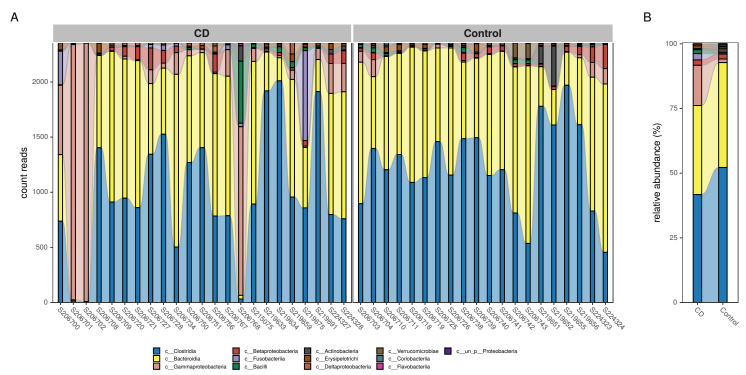


Fig. S4: This example show how to displayed the abundance (count or other) of sample and the relative abundance of groups. The Abundance (count by rarefied) of each sample (A) and the relative abundance of group (B), these results show the *Gammaproteobacteria* of *CD* group might be more abundant than the *control* group.

The abundance of features also can be visualized by mp\_plot\_abundance with heatmap plot by setting geom="heatmap".

```
hclass1 <- mpse2 %>%
          mp_plot_abundance(
             .abundance = RareAbundance,
             .group = Group,
             taxa.class = Class,
             topn = 30,
             geom = "heatmap"
          ) %>%
          set_scale_theme(
            x = list(scale_fill_viridis_c(option = "H"),
                     theme(
                       axis.text.x = element_text(size = 6),
                       axis.text.y = element_text(size = 7),
                       legend.title = element_text(size = 7),
                       legend.text = element_text(size = 5),
                       legend.key.width = unit(0.3, "cm"),
                       legend.key.height = unit(0.3, "cm")
                ),
            aes_var = RelRareAbundance
          ) %>%
          set_scale_theme(
            x = list(scale_fill_manual(values = cols),
                     theme (
                       legend.key.height = unit(0.3, "cm"),
                       legend.key.width = unit(0.3, "cm"),
```

```
legend.spacing.y = unit(0.02, "cm"),
                       legend.text = element_text(size = 7),
                       legend.title = element_text(size = 9)
                ),
            aes_var = Group
hclass2 <- mpse2 %>%
          mp_plot_abundance(
             .abundance = RareAbundance,
             .group = Group,
             taxa.class = Class,
             topn = 30,
             geom = 'heatmap',
             relative = FALSE
          ) %>%
          set_scale_theme(
            x = list(scale_fill_viridis_c(option = "H"),
                     theme(
                       axis.text.x = element_text(size = 6),
                       axis.text.y = element_text(size = 7),
                       legend.title = element_text(size = 7),
                       legend.text = element_text(size = 5),
                       legend.key.width = unit(0.3, "cm"),
                       legend.key.height = unit(0.3, "cm")
                ),
            aes_var = RareAbundance
          ) %>%
          set_scale_theme(
            x = list(scale fill manual(values = cols),
                     theme(
                       legend.key.height = unit(0.3, "cm"),
                       legend.key.width = unit(0.3, "cm"),
                       legend.spacing.y = unit(0.02, "cm"),
                       legend.text = element_text(size = 7),
                       legend.title = element_text(size = 9)
                ),
            aes_var = Group
p <- aplot::plot_list(hclass1, hclass2, nrow = 1, tag_levels = "A")</pre>
p
```

#### 1.4.2 Venn or Upset plot

The Venn or UpSet plot can help us to obtain the difference between groups in overview. MicrobiotaProcess provides mp\_cal\_venn (mp\_plot\_venn) and mp\_cal\_upset (mp\_plot\_upset) to perform the Venn and Upset analysis.

```
mpse2 %<>%
    mp_cal_venn(
        .abundance = RareAbundance,
        .group = Group
)

venn_p <- mpse2 %>%
    mp_plot_venn(
        .group = Group,
```

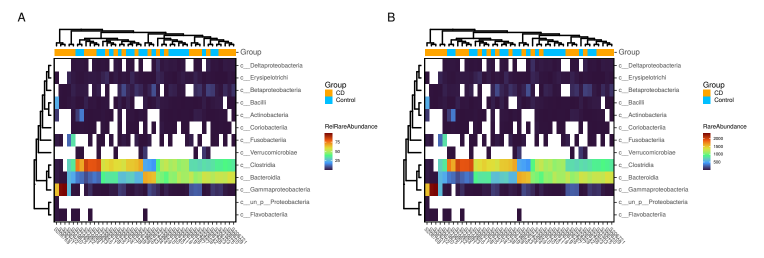


Fig. S5: The heatmap of abundance for each sample in *class* level. The color (continuous) of heatmap represents the abundance of taxon, the color of bar represents the group name of sample, the horizontal coordinate represents the sample, and the vertical coordinate represents the taxon.

```
set_size = 2.5,
      label_size = 2,
      edge_size = 2.5
    ) +
    scale colour manual(values = cols) +
    scale_fill_viridis_c(guide = guide_colorbar(barwidth=.3, barheight=2)) +
      legend.title = element_text(size = 8),
      legend.text = element_text(size = 6)
    )
mpse2 %<>%
    mp_cal_upset(
      .abundance = RareAbundance,
      .group = Group
    )
upset_p <- mpse2 %>%
    mp_plot_upset(
      .group = Group
    theme bw() +
    theme(
      plot.background = element_blank(),
      panel.border = element_blank(),
      panel.grid = element_blank(),
      axis.line.x.bottom = element_line(size = .5),
      axis.line.y.left = element_line(size = .5)
    ) +
    ggupset::theme_combmatrix(
      combmatrix.label.extra_spacing = 40
    )
library(ggpp)
p.up.venn <- upset_p +</pre>
             ggpp::annotate(
                "plot_npc",
               npcx = "right",
               npcy = "top",
               label = venn_p,
```

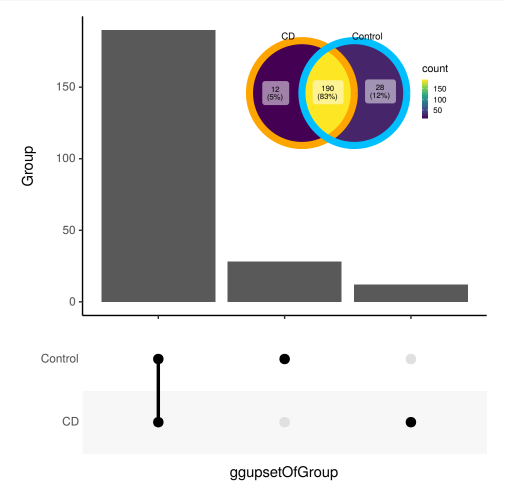


Fig. S6: The venn diagram and upset plot for groups in OTU/ASV level

## 1.5 beta analysis

#### 1.5.1 PCA analysis

PCA (Principal component analysis) and PCoA (Principal Coordinate Analysis) are general statistical procedures to compare dissimilarity of samples. And PCoA can based on the phylogenetic or count-based distance metrics, such as Bray-Curtis, Jaccard, Unweighted-UniFrac and weighted-UniFrac. MicrobiotaProcess presents the mp\_cal\_dist, mp\_cal\_pca, mp\_cal\_pca, mp\_cal\_dca, mp\_cal\_nmds, mp\_cal\_cca, mp\_cal\_rda, mp\_adonis, mp\_anosim, mp\_mrpp, mp\_envfit and mp\_mantel for the analysis.

```
library(MicrobiotaProcess)
library(patchwork)
# hellinger transform
mpse2 %<>%
    mp_decostand(
        .abundance = Abundance,
        method = "hellinger"
    )

mpse2 %<>% mp_cal_pca(.abundance = hellinger)
# Visulizing the result
pcaplot1 <- mpse2 %<%</pre>
```

```
mp_plot_ord(
              .ord = pca,
              .group = Group,
              .starshape = Group,
              .size = Observe
            scale_fill_manual(values = cols) +
            scale_size_continuous(
             range = c(1, 3),
              guide = guide_legend(override.aes = list(starshape = 15))
            ) +
            theme(
              legend.key.width = unit(0.3, "cm"),
              legend.key.height = unit(0.3, "cm"),
              legend.text = element_text(size = 6),
              legend.title = element_text(size = 7)
\# .dim = c(1, 3) to show the first and third principal components.
pcaplot2 <- mpse2 %>%
            mp_plot_ord(
              .ord = pca,
              .dim = c(1, 3),
              .group = Group,
              .starshape = Group,
              .size = Observe
            ) +
            scale_fill_manual(values = cols) +
            scale_size_continuous(
              range = c(1, 3),
              guide = guide_legend(override.aes = list(starshape = 15))
            theme (
              legend.key.width = unit(0.3, "cm"),
              legend.key.height = unit(0.3, "cm"),
              legend.text = element_text(size = 6),
              legend.title = element_text(size = 7)
(pcaplot1 | pcaplot2) + plot_annotation(tag_levels = "A")
```

## 1.5.2 PCoA analysis

```
# distmethod
# "unifrac",
             "wunifrac", "manhattan", "euclidean", "canberra", "bray", "kulczynski" ... (vegdist, dist)
mpse2 %<>%
   mp_cal_dist(
      .abundance = hellinger,
      distmethod = "bray"
    )
# PCoA analysis
mpse2 %<>%
    mp_cal_pcoa(
      .abundance = hellinger,
     distmethod = "bray"
    )
pcoaplot1 <- mpse2 %>%
             mp_plot_ord(
               .ord = pcoa,
```

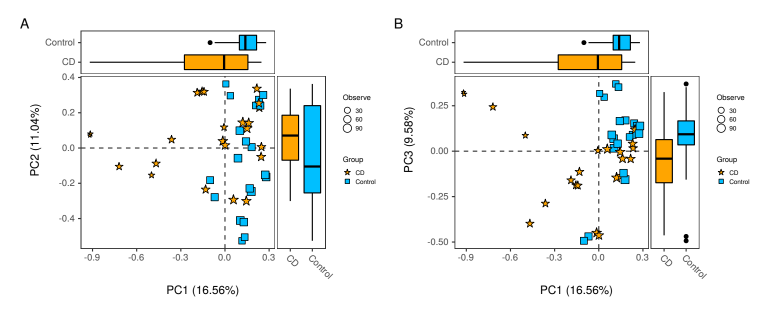


Fig. S7: The PCA plot of the community. Each point represents one sample, the size of point represents the observe OTU of the sample. The color of point represents the group name of the sample, based on the first and second component (A), based on the first and third component (B).

```
.group = Group,
               .starshape = Group,
               .color = Group,
               .size = Observe,
               ellipse = TRUE,
               show.legend = FALSE
            ) +
            scale_color_manual(
               values = cols
            ) +
            scale_fill_manual(values = cols) +
            scale_size_continuous(
               range = c(1, 3),
               guide = guide_legend(override.aes = list(starshape = 15))
            ) +
            theme(
               legend.key.width = unit(0.3, "cm"),
               legend.key.height = unit(0.3, "cm"),
               legend.text = element_text(size=6),
               legend.title = element_text(size=7)
# first and third principal co-ordinates
pcoaplot2 <- mpse2 %>%
             mp_plot_ord(
               .ord = pcoa,
               .group = Group,
               .starshape = Group,
               .color = Group,
               .size = Observe,
               ellipse = TRUE,
                .dim = c(1, 3),
               show.legend = FALSE
             ) +
             scale_color_manual(
               values = cols
             ) +
             scale_fill_manual(
```

```
values = cols
) +
scale_size_continuous(
    range = c(1, 3),
    guide = guide_legend(override.aes = list(starshape = 15))
) +
theme(
    legend.key.width = unit(0.3, "cm"),
    legend.key.height = unit(0.3, "cm"),
    legend.text = element_text(size = 6),
    legend.title = element_text(size = 7)
)
(pcoaplot1 | pcoaplot2) + plot_annotation(tag_levels = "A")
```

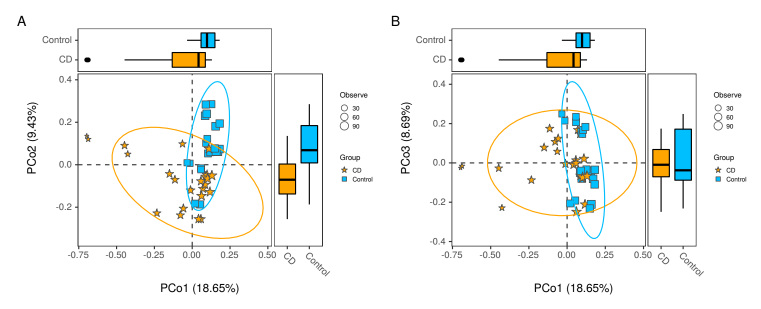


Fig. S8: The PCoA plot based on Bray-Curtis distance.

The result of distance between the samples also can be visualized by mp\_plot\_dist with heatmap or boxplot.

```
pdist1 <- mpse2 %>%
          mp_plot_dist(
            .distmethod = bray,
            .group = Group
          ) %>%
          set_scale_theme(
            x = scale_fill_manual(
                  values=cols,
                  guide = guide_legend(
                              keywidth = 0.5,
                              keyheight = 0.5,
                              label.theme=element_text(size=6)
                ),
            aes_var = Group
          ) %>%
          set_scale_theme(
            x = list(scale_size_continuous(range = c(1, 3)),
                     scale_color_viridis_c(option = "H"),
                     theme(
                       legend.key.width = unit(0.3, "cm"),
                       legend.text = element_text(size = 6),
                       legend.title = element_text(size = 7)
```

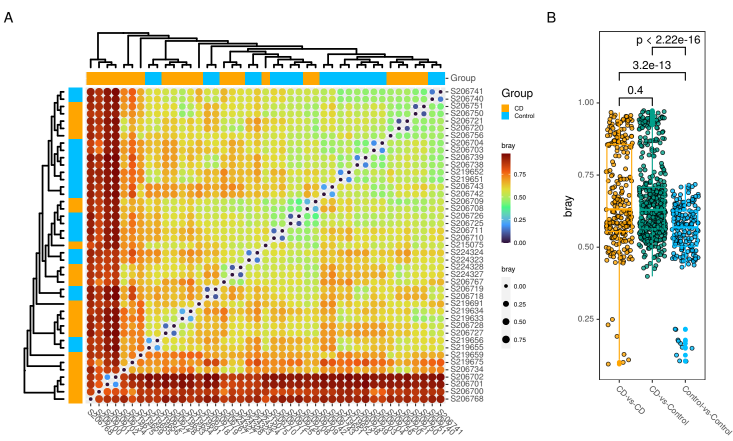


Fig. S9: The distance heatmap and the boxplot for each sample. The size and color of the heatmap represent the distance of each sample, the color of bar represent the group of sample (A). The boxplot represent the distance pairs of sample among the group, it show the dissimilarity of sample between the *control* and *CD* is significant, which is consistent with the result of Permutational Multivariate Analysis of Variance 1.5.3.

#### 1.5.3 Permutational Multivariate Analysis of Variance

We also can perform the Permutational Multivariate Analysis of Variance using mp\_adonis wrapping the adonis of vegan (Oksanen et al. 2020).

```
mpse2 %<>% mp_adonis(
            .abundance = hellinger,
            distmethod = "bray",
            .formula = ~Group,
            permutation = 9999,
            action = "add")
mpse2 %>%
    mp_extract_internal_attr(name=adonis) %>%
    mp_fortify()
## # A tibble: 3 x 7
##
                  Df SumsOfSqs MeanSqs F.Model
                                                     R2 `Pr(>F)`
     factors
                                           <dbl> <dbl>
##
     <chr>
               <dbl>
                          <dbl>
                                  <dbl>
                                                           <dbl>
                          0.789
                                  0.789
                                           3.88 0.0864
                                                          0.0001
## 1 Group
                   1
## 2 Residuals
                  41
                          8.34
                                  0.203
                                          NA
                                                 0.914
                                                         NA
```

From the result, we found the pvalue of the analysis of adonis is smaller than 0.05 for the Group, meaning the dissimilarity of samples between the Group is significant, which is consistent with the 1.5.2.

NΑ

1

#### 1.5.4 hierarchical cluster analysis of samples

9.12

NA

NA

42

## 3 Total

beta diversity metrics can assess the differences between microbial communities. It can be visualized with PCA or PCoA, this can also be visualized with hierarchical clustering based on ggplot2 (Wickham 2011), ggtree (Yu et al. 2017) and ggtreeExtra (Xu et al. 2021)

```
library(ggplot2)
library(MicrobiotaProcess)
library(ggtree)
library(ggtreeExtra)
mpse2 %<>%
    mp_cal_clust(
      .abundance = hellinger,
      distmethod = "bray",
      action = "add"
    )
hcsample <- mpse2 %>%
            mp_extract_internal_attr(name=SampleClust)
# rectangular layout + relative abundance of phyla
phy.tb <- mpse2 %>%
          mp_extract_abundance(
            taxa.class = Phylum,
            topn = 30
          ) %>%
          tidyr::unnest(cols=RareAbundanceBySample) %>%
          dplyr::rename(Phyla="label")
cplot1 <- ggtree(</pre>
            hcsample,
            layout = "rectangular"
          ) +
          geom_treescale(fontsize = 2) +
          geom_tippoint(mapping=aes(color=Group)) +
          geom_fruit(
            data = phy.tb,
            geom = geom_col,
```

```
mapping = aes(x = RelRareAbundanceBySample, y = Sample, fill = Phyla),
            orientation = "y",
            offset = 0.08,
            pwidth = 3,
           width = .6,
            axis.params = list(
             axis = x,
             title = "The relative abundance of phyla (%)",
             title.size = 3,
             title.height = 0.04,
             text.size = 2,
             vjust = 1
            )
          ) +
          geom_tiplab(as_ylab = TRUE) +
          scale_color_manual(
           values = cols,
            guide = guide_legend(
             keywidth = .5,
             keyheight = .5,
             title.theme = element_text(size = 8),
             label.theme = element_text(size = 6)
           )
          ) +
          scale_fill_manual(
            values=c(colorRampPalette(RColorBrewer::brewer.pal(12, "Set2"))(6)),
            guide = guide_legend(
             keywidth = .5,
             keyheight = .5,
             title.theme = element_text(size = 8),
             label.theme = element_text(size = 6)
           )
         ) +
          scale_x_continuous(expand = c(0, 0.01))
cplot1
```

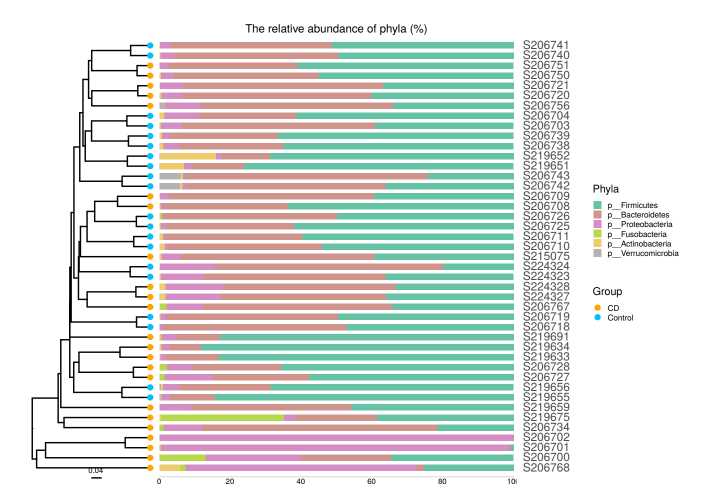


Fig.~S10: The hierarchical clustering plot of samples based on Bray-Curtis distance calculated with abundance of OTU/ASV and the relative Abundance of phyla for samples

## 1.6 biomarker discovery

This package provides mp\_diff\_analysis to detect the biomarker. And the result (with action = "get") can be visualized by ggdiffbox, ggdiffclade, ggeffectsize, ggdifftaxbar and mp\_plot\_diff\_res, or displayed manually using ggtree (Yu et al. 2017) and ggtreeExtra (Xu et al. 2021).

```
# for the kruskal_test and wilcox_test
library(coin)
library(MicrobiotaProcess)
# get result (diffAnalysisClass) of the different analysis with action = 'get'.
deres <- mpse2 %>%
         mp_diff_analysis(
            .abundance = RareAundance,
            .group = Group,
            first.test.method = "kruskal_test",
            filter.p = "pvalue",
            first.test.alpha = 0.05,
            strict = TRUE,
            second.test.method = "wilcox_test",
            second.test.alpha = 0.05,
            subcl.min = 3,
            subcl.test = TRUE,
            ml.method = "lda",
            ldascore = 3,
            action = "get"
```

#### 1.6.1 visualization of different results by ggdiffclade

The color of discriminative taxa represent the taxa is more abundant in the corresponding group. The point size shows the negative logarithms (base 10) of pvalue. The bigger size of point shows more significant (lower pvalue), the *pvalue* was calculated in the first step test (default is *kruskal.test*).

```
diffclade_p <- ggdiffclade(</pre>
                   obj=deres,
                   alpha=0.3,
                   linewd=0.15,
                   skpointsize=0.6,
                   layout="radial",
                   taxlevel=3,
                   removeUnkown = TRUE,
                   reduce = FALSE # This argument is to remove the branch of unknown taxonomy.
               ) +
               scale_fill_manual(
                   values = cols
               guides(color = guide_legend(
                                   keywidth = 0.1,
                                   keyheight = 0.2,
                                   order = 3,
                                   ncol=1)
               ) +
               theme(
                   panel.background = element_rect(fill=NA),
                   legend.position = "right",
                   plot.margin = margin(0,0,0,0),
                   legend.key.width = unit(0.2, "cm"),
                   legend.key.height = unit(0.2, "cm"),
                   legend.spacing.y = unit(0.02, "cm"),
                   legend.title = element_text(size=7),
                   legend.text = element_text(size=6),
                   legend.box.spacing = unit(0.02, "cm")
               )
diffclade_p
```

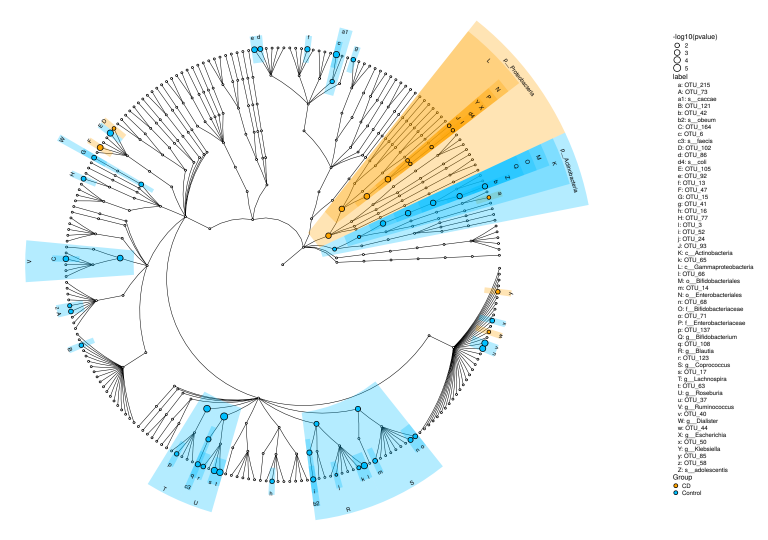


Fig. S11: The taxa tree clade plot of different analysis result.

We also can visualized the result with ggtree (Yu et al. 2017) and ggtreeExtra (Xu et al. 2021).

```
taxa.tree <- mpse2 %>% mp_extract_tree(type='taxatree')
p1 <- ggtree(
        taxa.tree,
        layout="radial",
        size = 0.3
      ) +
      geom_point(
        data = td_filter(!isTip),
        fill="white",
        size=1,
        shape=21
# display the high light of phylum clade.
p2 <- p1 +
      geom_hilight(
        data = td_filter(nodeClass == "Phylum"),
        mapping = aes(node = node, fill = label)
# display the relative abundance of features(OTU)
p3 <- p2 +
      ggnewscale::new_scale("fill") +
      geom_fruit(
         data = td_unnest(RareAbundanceBySample),
         geom = geom_star,
```

```
mapping = aes(
                       x = fct_reorder(Sample, Group, .fun=min),
                       size = RelRareAbundanceBySample,
                       fill = Group,
                       subset = RelRareAbundanceBySample > 0
                   ),
         starshape = 13,
         starstroke = 0.25,
         offset = 0.04,
         pwidth = 1.5,
         grid.params = list(vline = TRUE, size = 0.01, linetype = 1)
      ) +
      scale_size_continuous(
         name="Relative Abundance (%)",
         range = c(0.5, 3)
      ) +
      scale fill manual(values=cols)
# display the tip labels of taxa tree
p4 <- p3 + geom_tiplab(size=2, offset=12.8)
# display the LDA of significant OTU.
p5 <- p4 +
      ggnewscale::new_scale("fill") +
      geom_fruit(
         geom = geom_col,
         mapping = aes(
                       x = LDAmean,
                       fill = Sign_Group,
                       subset = !is.na(LDAmean)
         orientation = "y",
         offset = 0.5,
         pwidth = 1,
         axis.params = list(axis = "x",
                            title = "Log10(LDA)",
                            title.height = 0.005,
                            title.size = 2,
                            text.size = 1.8,
                            vjust = 1),
         grid.params = list(linetype = 3)
      )
# display the significant (FDR) taxonomy after kruskal.test (default)
p6 <- p5 +
      ggnewscale::new_scale("size") +
      geom_point(
         data=td_filter(!is.na(fdr)),
         mapping = aes(size = -log10(fdr),
                       fill = Sign_Group,
         shape = 21,
      ) +
      scale_size_continuous(range=c(1, 3)) +
      scale_fill_manual(values=cols)
p6 <- p6 + theme(
           legend.key.height = unit(0.3, "cm"),
           legend.key.width = unit(0.3, "cm"),
           legend.spacing.y = unit(0.02, "cm"),
           legend.text = element_text(size = 7),
           legend.title = element_text(size = 9),
```



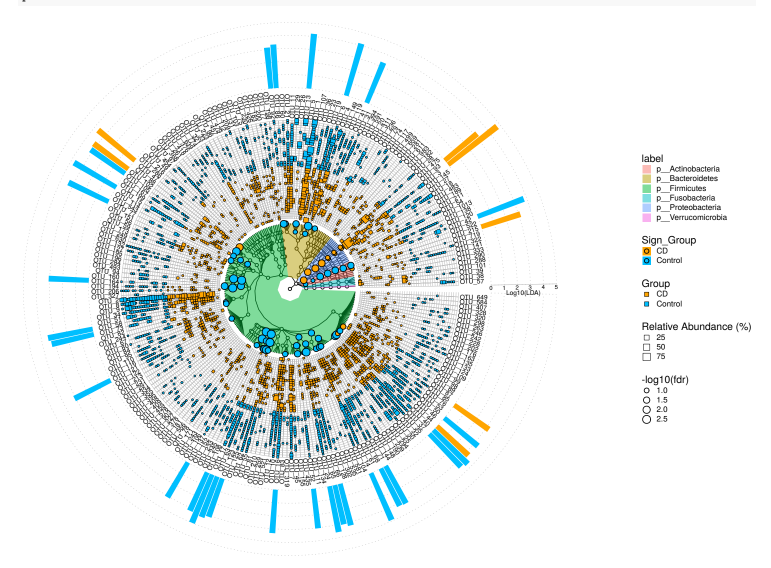


Fig. S12: The taxa tree of the community with the relative abundance of each OTU/ASV on sample and the LDA of different OTU/ASV. The taxa tree is built with the taxa of all samples. The high light color of taxa tree represents the phyla of the clade. The external point layer represents the relative abundance of each OTU on sample. The external bar layer represents the LDA of the different OTU. The colored points represent the different taxa, the size represents the pvalue or fdr.

To decreases coding burden, we also developed mp\_plot\_diff\_res to visualize the result of different analysis.

```
library(ggplot2)

pp <- mpse2 %>%
    mp_plot_diff_res() +
    scale_fill_manual(
        values = cols
    ) +
    scale_fill_manual(
        aesthetics = "fill_new",
        values = cols
    )

pp
```

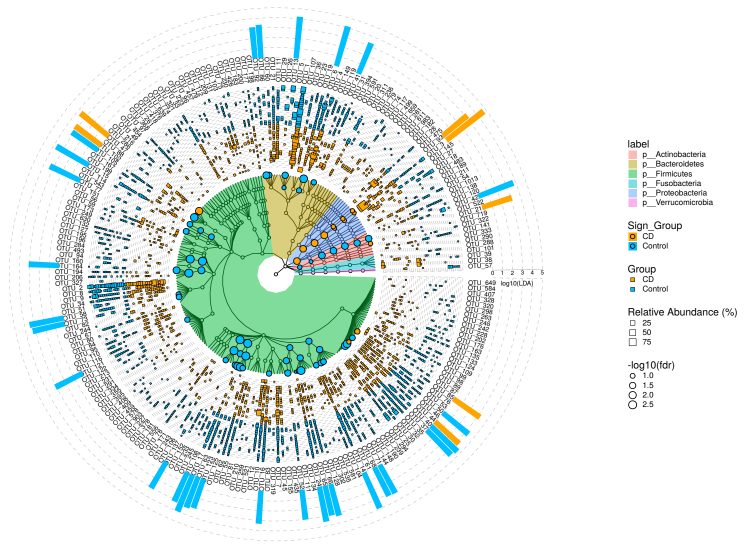


Fig. S13: seel also Fig. S12

## 1.6.2 visualization of different results by ggdiffbox

The left panel represents the relative abundance or abundance (according the standard\_method) of biomarker, the right panel represents the confident interval of effect size (LDA or MDA) of biomarker. The bigger confident interval shows that the biomarker is more fluctuant, owing to the influence of samples number.

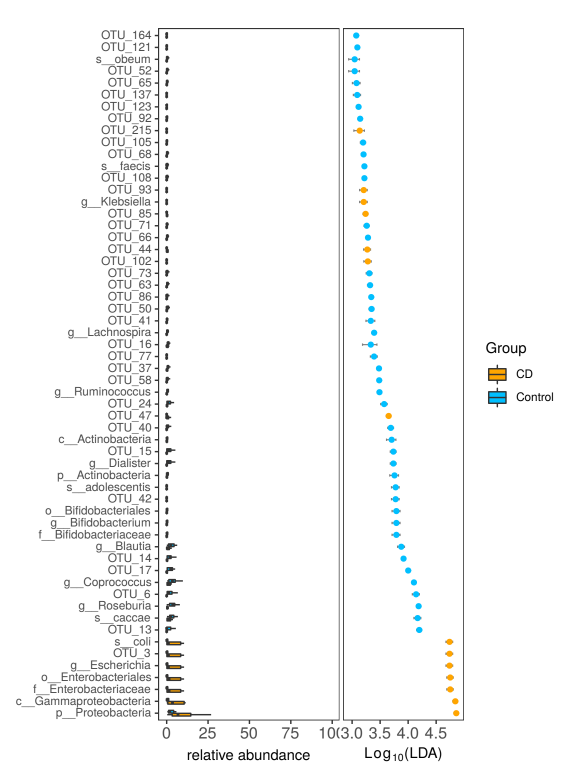


Fig. S14: The boxplot and the LDA score of different taxa. The left panel represents the relative abundance of the different taxa, the right panel represents the LDA effect size (95% confidence interval) of different taxa.

#### 1.6.3 visualization of different results by ggdifftaxbar

ggdifftaxbar can visualize the abundance of biomarker in each samples of groups, the mean and median abundance of groups or subgroups are also showed. output parameter is the directory of output.

### 1.7 Interoperable with the existing computing ecosystem

Because the MPSE object MicrobiotaProcess inherits the SummarizedExperiment object (Morgan et al. 2021), The related inherited methods for signature SummarizedExperiment can also be applied to the MPSE. For example, the tidybulk (Mangiola et al. 2021) provides an R tidy framework for modular transcriptomic data analysis. It provides a test\_differential\_abundance to perform differential transcription testing using edgeR quasi-likelihood edgeR likelihood-ratio (LR), limma-voom, limma-voom-with-quality-weights or DESeq2. It can also be compatible with MPSE.

```
library(tidybulk)
library(edgeR)
library(aplot)
library(shadowtext)
library(ggrepel)
mpse2 %<>% test_differential_abundance(.abundance = Abundance, .formula = ~Group)
# extract the different OTUs from the MPSE class
res <- mpse2 %>% dplyr::filter(FDR <= .05 & abs(logFC) >= 2)
pp <- res %>%
      mp_plot_abundance(
        .abundance = RareAbundance,
        force = TRUE,
       relative = TRUE,
       feature.dist = "bray",
        geom = "heatmap",
        topn = "all",
        .group = Group
      ) %>%
      set scale theme(
        x = list(scale_fill_viridis_c(option = "H"),
                 theme(
                   axis.text.x = element_text(size = 6),
                   axis.text.y = element_text(size = 6),
                   legend.title = element_text(size = 7),
                   legend.text = element_text(size = 5),
                   legend.key.width = unit(0.3, "cm"),
                   legend.key.height = unit(0.3, "cm")
            ),
        aes var = RelRareAbundance
      ) %>%
      set_scale_theme(
        x = list(scale_fill_manual(values = cols),
                   legend.key.height = unit(0.3, "cm"),
                   legend.key.width = unit(0.3, "cm"),
                   legend.spacing.y = unit(0.02, "cm"),
                   legend.text = element_text(size = 7),
                   legend.title = element_text(size = 9)
                 )
            ),
        aes_var = Group
f <- res %>%
     mp_extract_taxonomy %>%
     ggplot() +
     geom text(
       mapping = aes(y=OTU, x=0, label=Genus, color=Phylum),
      hjust = 0,
       size = 2
     scale x continuous(expand=c(0, 0, 0, 0.1)) +
```

```
theme_bw() +
     theme(
       legend.text = element_text(size = 5),
       legend.title = element_text(size = 7),
       legend.key.width = unit(0.3, "cm"),
       legend.key.height = unit(0.3, "cm"),
       panel.background = element_blank(),
       panel.grid = element_blank(),
       axis.text = element_blank(),
       axis.ticks = element_blank(),
       panel.border = element_blank()
     ) +
     labs(x = NULL, y = NULL)
pp <- pp %>% insert_right(f, width = 0.2)
sample.tree <- res %>%
      select(-bray) %>% # remove the bray, Because it was the result of all OTU,
      mp_cal_clust(.abundance = RelRareAbundanceBySample, distmethod = "bray") %>%
      ggtree(layout = igraph::layout_with_kk, color = "#afb7b8") +
      geom_nodepoint(color = "#afb7b8", size = .5) +
      geom_tippoint(aes(fill = Group), shape = 21, size=3) +
      geom_text_repel(
        data = td_filter(isTip),
        mapping = aes(label = label),
        size = 2,
        max.overlaps = 30,
        colour = "black",
        bg.colour = "white"
      ) +
      scale_fill_manual(
        values = cols,
        guide = guide_legend(
           title.theme = element_text(size = 7),
           label.theme = element_text(size = 5),
        )
      )
## # Note: MPSE object is converted to a tibble data (tbl_mpse object) for independent data analysis.
## #
        A new MPSE object can be returned by setting keep.mpse = TRUE.
p <- mpse2 %>%
      mp_cal_dist(
         .abundance = RelRareAbundanceBySample,
         distmethod = "bray",
         cal.feature.dist = T
      ) %>%
      hclust() %>%
      ggtree(layout = igraph::layout_with_kk, color = "#bed0d1") +
      geom_nodepoint(color = "#bed0d1", size = .5)
{\it \# The \ data.frame \ contained \ results \ of \ test\_differential\_abundance}
otu.tab <- mpse2 %>% mp_extract_feature()
p <- p %<+% otu.tab +
     geom_tippoint(
       mapping = aes(fill = logFC, size = -log10(FDR)),
       shape = 21,
       color = "grey"
     ) +
     scale_fill_viridis_c(
       option="C",
```

```
guide = guide_colorbar(
          title.theme = element_text(size = 7),
          label.theme = element_text(size = 5),
          barheight = unit(1.5, "cm"),
          barwidth = unit(.3, "cm")
       )
     ) +
     scale_size_continuous(
       range = c(.5, 6),
       guide = guide legend(
          key.width = .3,
          key.height = .3,
          label.theme = element_text(size = 5),
          title.theme = element_text(size = 7)
       )
     ) +
     geom text repel(
       data = td_filter(FDR <= .05 & abs(logFC) >= 2),
       mapping = aes(x = x, y = y, label = label),
       size = 2,
       min.segment.length = 0.1,
       segment.size = .25,
       segment.colour = 'grey18',
       colour = "black",
       bg.colour = 'white'
       \#max.overlaps = 60,
design <- "
  12
  13
  13
plot_list(pp, sample.tree, p, design = design, tag_levels = "A")
```

We compared the different result between the edgeR (Robinson, McCarthy, and Smyth 2010) and MicrobiotaProcess. We found the number of the different OTUs based on edgeR is more than the MicrobiotaProcess. We think this is because we didn't remove the low-abundance OTUs. This operation is generally needed in standard whole-transcriptome workflows. However, if it is preformed in the microbiome analysis, many low-abundance OTUs will be removed. More different OTUs were identified by the operation using edgeR (Robinson, McCarthy, and Smyth 2010).

Then we extract the same different OTUs, we found the abundance of same OTUs belonged to *Bifidobacterium*, *Faecalibacterium*, *Roseburia* and *Coprobacillus* were significantly decreased in CD group compared to the Control group, the abundance of several OTUs belonged to *Escherichia*, *Klebsiella* and *Haemophilus*, which belonged to Gammaproteobacteria, were significantly enriched in CD group.

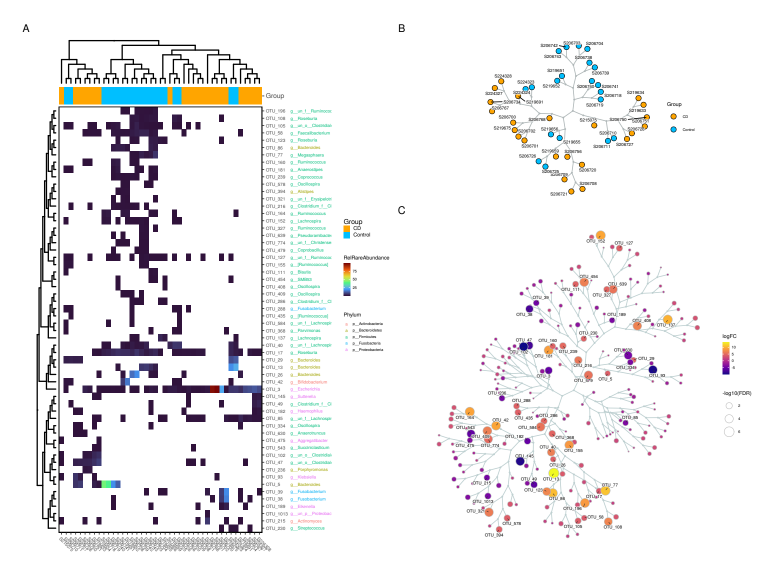


Fig. S15: The results of different OTUs based on the edgeR\_quasi\_likelihood with tidybulk A. The relative abundance heatmap of the different OTUs. B. The hierarchical cluster of samples based on the relative abundance of the different OTUs. C. The hierarchical cluster of OTUs based on the relative abundance of total OTUs, the different OTUs were labeled with their names. We found the cluster of different OTUs in the heatmap is consistent with the different OTUs in the background of total OTUs (C).

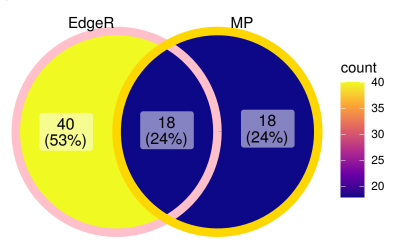


Fig. S16: The comparison of the different analysis result between the edgeR and MicrobiotaProcess

```
mpse2 %>%
    mp_extract_feature(addtaxa=T) %>%
```

```
dplyr::filter(OTU %in% do.call(intersect, unname(DE.method)))
```

```
## # A tibble: 18 x 22
     OTU
            ggupsetOfGroup logFC logCPM
                                          F PValue
                                                        FDR Kingdom Phylum Class
##
##
     <chr> <chr>> <chr>>
                          <dbl> <dbl> <dbl>
                                              <dbl>
                                                      <dbl> <chr> <chr> <chr>
  1 OTU_2~ <chr [2]>
                          -4.44
                                  9.41 9.00 3.50e-3 1.79e-2 k_Bac~ p_Ac~ c_A~
   2 OTU_42 <chr [1]>
                           8.97 12.1 29.0 5.79e-7 1.48e-5 k_Bac~ p_Ac~ c_A~
   3 OTU_86 <chr [1]>
                                 11.1 36.2 8.10e-8 4.66e-6 k_Bac~ p_Ba~ c_B~
                          10.4
##
  4 OTU_13 <chr [1]>
                          13.9
                                 14.5 48.9 1.48e-9 3.41e-7 k_Bac~ p_Ba~ c_B~
## 5 OTU 1~ <chr [1]>
                           9.78 10.5 33.9 1.74e-7 6.42e-6 k Bac~ p Fi~ c C~
## 6 OTU_1~ <chr [2]>
                           6.70 10.6 19.8 2.50e-5 2.21e-4 k_Bac~ p_Fi~ c_C~
## 7 OTU_17 <chr [2]>
                                      8.95 3.59e-3 1.79e-2 k__Bac~ p__Fi~ c__C~
                           3.48 13.6
## 8 OTU_1~ <chr [2]>
                           6.85 10.4 20.2 2.05e-5 1.96e-4 k_Bac~ p_Fi~ c_C~
## 9 OTU_40 <chr [2]>
                           6.29 12.7 17.8 5.74e-5 4.89e-4 k_Bac~ p_Fi~ c_C~
## 10 OTU_85 <chr [2]>
                          -3.88 11.0
                                       7.85 6.23e-3 2.71e-2 k_Bac~ p_Fi~ c_C~
## 11 OTU 58 <chr [2]>
                           3.61 11.9
                                      7.84 6.25e-3 2.71e-2 k__Bac~ p__Fi~ c__C~
## 12 OTU 1~ <chr [1]>
                           8.90
                                 9.65 33.6 1.95e-7 6.42e-6 k_Bac~ p_Fi~ c_C~
## 13 OTU_47 <chr [1]>
                          -7.37 11.8 20.7 1.69e-5 1.69e-4 k_Bac~ p_Fi~ c_C~
                          -9.73 10.6 28.2 1.20e-6 2.13e-5 k_Bac~ p_Fi~ c_C~
## 14 OTU_1~ <chr [1]>
                                      9.83 2.32e-3 1.27e-2 k__Bac~ p__Fi~ c__C~
## 15 OTU_1~ <chr [2]>
                          3.84 10.6
## 16 OTU_77 <chr [1]>
                                 11.5 37.4 5.50e-8 4.21e-6 k_Bac~ p_Fi~ c_C~
                          10.8
## 17 OTU_3 <chr [2]>
                          -5.35 18.2 15.8 1.40e-4 1.07e-3 k__Bac~ p__Pr~ c__G~
                          -9.39 10.3 27.5 1.56e-6 2.57e-5 k_Bac~ p_Pr~ c_G~
## 18 OTU_93 <chr [1]>
## # ... with 12 more variables: Order <chr>, Family <chr>, Genus <chr>,
      Species <chr>, RareAbundanceBySample <list>, RareAbundanceByGroup <list>,
## #
      LDAupper <dbl>, LDAmean <dbl>, LDAlower <dbl>, Sign_Group <chr>,
## #
      pvalue <dbl>, fdr <dbl>
```

## 2 the analysis of the other published pediatric CD stool samples

In the previous chapter, we described how to use *MicrobiotaProcess* to do the analysis of the 16s rDNA data. However, it also can be applied to metagenome or metatranscriptome species community data and the function data analysis. In this chapter, we used the example datasets about the other published pediatric CD stool microbial study (Douglas et al. 2018) to show how to used *MicrobiotaProcess* to do the related analysis. The datasets were obtained from the github<sup>2</sup>. However, to avoid duplication, we only show how to import the 16s dataset, we focused on the analysis of metagenomics and KEGG gene datasets.

## 2.1 Loading the 16s data and construction of MPSE class

cols <- c("orange", "deepskyblue")</pre>

cols2 <- c("deepskyblue", "yellow", "#FF9933")</pre>

The chapter is similar with the 1, so some operations can refer to the previous chapter 1.

```
sample.da <- read.table("./data/CD_RF_microbiome/biscuit_metadata.txt", header=TRUE, check.names=FALSE, sep="\t
sample.da %<>% dplyr::select(1:5)
biom <- biomformat::read_biom("./data/CD_RF_microbiome/otu_table_w_tax_BISCUIT.biom")</pre>
mpse16s <- biom %>% as.MPSE
mpse16s
## # A MPSE-tibble (MPSE object) abstraction: 37,392 x 10
## # OTU=984 | Samples=38 | Assays=Abundance | Taxanomy=Kingdom, Phylum, Class, Order, Family, Genus, Speies
##
      OTU
              Sample Abundance Kingdom
                                           Phylum
                                                   Class Order Family Genus Speies
                         <dbl> <chr>
                                           <chr>
                                                   <chr> <chr> <chr> <chr> <chr> <chr>
##
      <chr>>
              <chr>
##
    1 358030
              S15
                             5 k_Bacteria p_Fir~ c_Cl~ o_C~ f_Ru~ g_u~ s_un~
    2 196271
              S15
                             O k_Bacteria p_Fir~ c_Cl~ o_C~ f_La~ g_u~ s_un~
   3 196270
                             2 k_Bacteria p_Fir~ c_Cl~ o_C~ f_un~ g_u~ s_un~
##
             S15
                             0 k_Bacteria p_Fir~ c_Cl~ o_C~ f_La~ g_u~ s_un~
    4 297149
              S15
   5 3604981 S15
                             O k_Bacteria p_Fir~ c_Cl~ o_C~ f_La~ g_B~ s_un~
##
   6 240755 S15
                             O k Bacteria p Pro~ c Ga~ o P~ f Pa~ g H~ s in~
   7 326482 S15
                             O k_Bacteria p_Bac~ c_Ba~ o_B~ f_Pr~ g_P~ s_co~
##
    8 4393540 S15
                             0 k__Bacteria p__Bac~ c__Ba~ o__B~ f__[B~ g__u~ s__un~
##
##
   9 4339144 S15
                             0 k_Bacteria p_Bac~ c_Ba~ o_B~ f_[0~ g_B~ s_un~
## 10 4369050 S15
                             O k_Bacteria p_Fus~ c_Fu~ o_F~ f_Fu~ g_F~ s_un~
## # ... with 37,382 more rows
mpse16s %<>% dplyr::left_join(sample.da, by=c("Sample"="sample_id"))
mpse16s
## # A MPSE-tibble (MPSE object) abstraction: 37,392 x 14
## # OTU=984 | Samples=38 | Assays=Abundance | Taxanomy=Kingdom, Phylum, Class, Order, Family, Genus, Speies
##
      OTU
              Sample Abundance disease response sex
                                                        age Kingdom Phylum Class
                                       <chr>
##
      <chr>>
              <chr>
                         <dbl> <chr>
                                                <chr> <dbl> <chr>
                                                                     <chr>
                                                                             <chr>
   1 358030
             S15
                             5 CN
                                       CN
                                                Male
                                                       15.4 k_Bact~ p_Fir~ c_Cl~
                             O CN
                                                       15.4 k__Bact~ p__Fir~ c__Cl~
    2 196271
                                       CN
##
             S15
                                                Male
    3 196270
                             2 CN
                                       CN
                                                       15.4 k__Bact~ p__Fir~ c__Cl~
##
             S15
                                                Male
##
    4 297149
             S15
                             O CN
                                       CN
                                                       15.4 k__Bact~ p__Fir~ c__Cl~
                                                Male
    5 3604981 S15
                             O CN
                                       CN
                                                Male
                                                       15.4 k__Bact~ p__Fir~ c__Cl~
    6 240755
                             O CN
                                       CN
                                                       15.4 k__Bact~ p__Pro~ c__Ga~
##
             S15
                                                Male
                                       CN
##
   7 326482
             S15
                             O CN
                                                Male
                                                       15.4 k__Bact~ p__Bac~ c__Ba~
                                       CN
   8 4393540 S15
                             O CN
                                                Male
                                                       15.4 k__Bact~ p__Bac~ c__Ba~
   9 4339144 S15
                                                       15.4 k__Bact~ p__Bac~ c__Ba~
                             O CN
                                       CN
                                                Male
## 10 4369050 S15
                             O CN
                                       CN
                                                       15.4 k__Bact~ p__Fus~ c__Fu~
                                                Male
## # ... with 37,382 more rows, and 4 more variables: Order <chr>, Family <chr>,
       Genus <chr>, Speies <chr>
```

<sup>&</sup>lt;sup>2</sup>https://github.com/LangilleLab/CD\_RF\_microbiome

## 2.2 Loading the KEGG data

The KEGG gene abundances were annotated based on the MGS data. It can also be imported as MPSE, and further analyzed using *MicrobiotaProcess*. Here, we only show how to identify the different gene using the  $mp\_diff\_analysis$  of *MicrobiotaProcess* (refer to chapter 2.3). Other operations are similar with the analysis of 16s rDNA data (refer to chapter 1).

```
KO.da <- read.table("./data/CD_RF_microbiome/biscuit_mgs_KOs.tsv", header=TRUE, sep = "\t", row.names=1, check.
# Building the MPSE object.
mpseKO <- MPSE(assays=list(Abundance = KO.da))
# merge the sample metadata information.
mpseKO %<>% left_join(sample.da, by=c("Sample"="sample_id"))
```

#### 2.2.1 Different analysis for KEGG genes abundance

The metrics of the KEGG genes is the relative abundance, here we used  $mp\_diff\_analysis$  to identify the difference KEGG genes with 'force = TRUE and relative = FALSE', meaning the relative abundance will be used directly.

```
mpseKO %<>%
  mp_diff_analysis(
    .abundance = Abundance,
  force = TRUE,
  relative = FALSE,
    .group = disease,
  filter.p = "pvalue"
)
```

Then we can perform the KEGG pathway enrichment analysis using clusterProfiler (Wu et al. 2021) and MicrobiomeProfiler (Chen and Yu 2021) developed by our team.

```
# perform KEGG pathway analysis with clusterProfiler and MicrobiomeProfiler
com.xx <- mpseKO %>%
    mp_extract_feature() %>% # Extracting the feature metadata information
    dplyr::filter(pvalue <= 0.05) %>% # Extracting the features pvalue <= 0.05
    compareCluster(OTU~Sign_disease, data=., fun=enrichKO)
# visualizing the enriched pathway with dotplot
p.dot <- dotplot(com.xx) +</pre>
         scale_color_gradientn(
           colours = c("#b3eebe", "#46bac2", "#371ea3"),
           guide = guide_colorbar(reverse=TRUE, order=1)
         ) +
         labs(x = NULL) +
         guides(size = guide_legend(override.aes=list(shape=1))) +
           panel.grid.major.y = element_line(linetype='dotted', color='#808080'),
           panel.grid.major.x = element_blank()
# with network plot
set.seed(1024)
p.net <- cnetplot(</pre>
           com.xx,
           layout = "fr",
           cex_label_category = 1.8
         scale_fill_manual(
           values = cols
p <- aplot::plot_list(p.net, p.dot, widths = c(3, 1), tag_levels="A")</pre>
```

A B

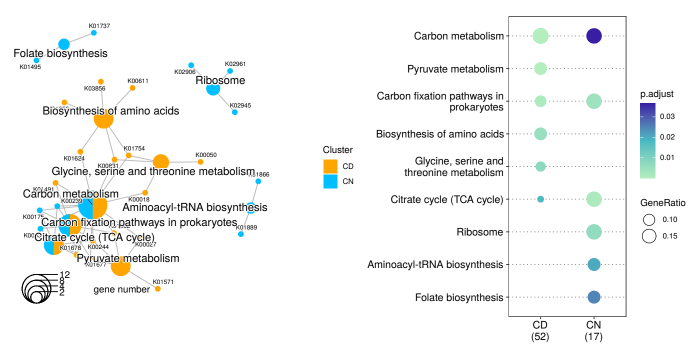


Fig. S17: The result of KEGG pathway enrichment analysis

### 2.3 Loading the MGS data

The taxa abundance data also can be analyzed by MicrobiotaProcess, Here we used the example data from output of MetaPhlAn (Segata et al. 2012) to show how to perform the related analysis using MicrobiotaProcess. The output of other taxa abundance can also be imported and converted to the MPSE object, and be further analyzed using MicrobiotaProcess, which can refer to chapter 2.2 and chapter 3.

```
# This is the output of MetaPhlAn2, which might need to specific the 'linenum'
# base on the first several rows whether to contain the metadata information
mpseMGS <- mp import metaphlan("./data/CD RF microbiome/metaphlan2 out merged species.tsv", linenum=1)
# rename the column names of MPSE.
colnames(mpseMGS) <- mpseMGS %>% mp_extract_sample %>% pull(2)
mpseMGS %<>% left_join(sample.da, by=c("Sample"="sample_id"))
mpseMGS
## # A MPSE-tibble (MPSE object) abstraction: 4,370 x 14
   # OTU=115 | Samples=38 | Assays=Abundance | Taxanomy=Kingdom, Phylum, Class, Order, Family, Genus
##
      OTU
              Sample Abundance unknown1 disease response sex
                                                                    age Kingdom Phylum
##
      <chr>
              <chr>
                          <dbl> <chr>
                                                           <chr> <dbl> <chr>
                                          <chr>
                                                  <chr>
                                                                                 <chr>
##
    1 s_un_~ S12
                           0
                                S12
                                          CN
                                                  CN
                                                           Fema~
                                                                    8.6 k__Arc~ p__Eu~
                                                                    8.6 k__Bac~ p__Ac~
    2 s Bif~ S12
                           0
                                S12
                                          CN
                                                  CN
##
                                                           Fema~
                                                                    8.6 k__Bac~ p__Ac~
##
    3 s Bif~ S12
                           0
                                S12
                                          CN
                                                  CN
                                                           Fema~
##
    4 s__Bif~ S12
                           Ω
                                S12
                                          CN
                                                  CN
                                                           Fema~
                                                                    8.6 k__Bac~ p__Ac~
    5 s Col~ S12
                           0
                                S12
                                          CN
                                                  CN
                                                                    8.6 k Bac~ p Ac~
##
                                                           Fema~
    6 s__Col~ S12
                           0
                                S12
                                                  CN
                                                                    8.6 k__Bac~ p__Ac~
##
                                          CN
                                                           Fema~
    7 s__un_~ S12
##
                           0
                                S12
                                          CN
                                                  CN
                                                           Fema~
                                                                    8.6 k__Bac~ p__Ac~
                                                  CN
##
    8 s_un_~ S12
                           0
                                S12
                                          CN
                                                           Fema~
                                                                    8.6 k__Bac~ p__Ac~
##
    9 s__Bac~ S12
                           6.34 S12
                                          CN
                                                  CN
                                                           Fema~
                                                                    8.6 k__Bac~ p__Ba~
   10 s__Bac~ S12
                           0
                                S12
                                          CN
                                                  CN
                                                                    8.6 k__Bac~ p__Ba~
                                                           Fema~
     ... with 4,360 more rows,
                                and 4 more variables: Class <chr>, Order <chr>,
       Family <chr>, Genus <chr>
```

#### 2.3.1 Alpha diversity analysis in MGS level

The metric of Metagenomics data usually is relative abundance. But the some functions of MicrobiotaProcess need to require the abundance is count (in default). To process the relative abundance (not integer), We can specific 'force = TRUE', which meaning the corresponding functions will be calculated directly without rarefied.

```
mpseMGS %<>%
    mp_cal_alpha(
        .abundance = Abundance,
        force = TRUE
    )

p <- mpseMGS %>%
        mp_plot_alpha(
        .group = disease,
        .alpha = c(Observe, Shannon, Pielou)
    ) +
    scale_color_manual(values = cols) +
    scale_fill_manual(values = cols) +
    theme(legend.position = "none")

p
```

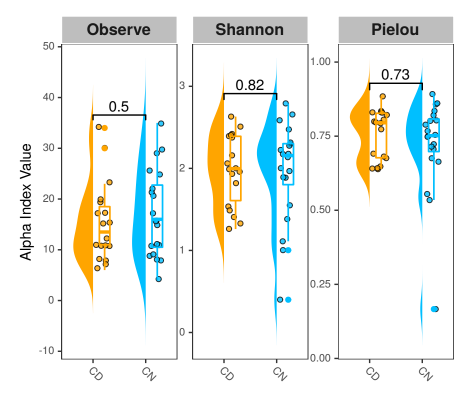


Fig. S18: The alpha diversity boxplot based on MGS data

#### 2.3.2 Beta diversity analysis in MGS level

We used  $mp\_cal\_dist$  to calculated the distance between the samples, then used  $mp\_plot\_dist$  to display the distance with heatmap (Fig.S19.A) and boxplot (Fig.S19.B), then the distance was used to perform the PCoA analysis (Fig.S19.C).

```
mpseMGS %<>%
   mp_decostand(
      .abundance = Abundance,
      method = "hellinger"
    )
mpseMGS %<>%
    mp_cal_dist(
      .abundance = hellinger,
      distmethod = "bray"
    )
mpseMGS %<>%
    mp_cal_pcoa(
      .abundance = hellinger,
      distmethod = "bray"
    )
p1 <- mpseMGS %>%
      mp_plot_dist(
        .distmethod = bray,
        .group=c(disease, response)
      ) %>%
      set_scale_theme(
        x = scale_fill_manual(
              values = cols,
              guide = guide_legend(
                           keywidth = 0.5,
                           keyheight = 0.5,
                           label.theme=element_text(size=6)
                 )
            ),
        aes_var = disease
      ) %>%
      set_scale_theme(
        x = scale_fill_manual(
              values=cols2,
              guide = guide_legend(
                           keywidth = 0.5,
                           keyheight = 0.5,
                           label.theme=element_text(size=6)
                 )
            ),
        aes_var = response
      ) %>%
      set_scale_theme(
        x = scale_size_continuous(
              range = c(1, 3),
              guide = guide_legend(
                           keywidth = 0.5,
                           keyheight = 0.5,
                           label.theme=element_text(size=6)
                 )
            ),
        aes_var = bray
```

```
p2 <- mpseMGS %>%
      mp_plot_dist(
        .distmethod = bray,
        .group = disease,
       group.test = TRUE
      ) +
      scale_color_manual(
        values = c("orange", "#00A08A", "deepskyblue")
      scale_fill_manual(
        values = c("orange", "#00A08A", "deepskyblue")
p3 <- mpseMGS %>%
      mp_plot_ord(
       .ord = pcoa,
       .group = disease,
        .size = Observe,
        .starshape = response,
        show.side = FALSE
      ) +
      scale_starshape_manual(values = c(1, 13, 15)) +
      scale fill manual(
       values=cols,
        guide=guide_legend(
          keywidth = 0.3,
          keyheight = 0.3,
          label.element = element_text(size = 6),
          override.aes = list(size = 2, starshape = 15)
        )
      ) +
      scale_size_continuous(
       range = c(1, 3),
        guide = guide_legend(
          keywidth = 0.3,
          keyheight = 0.3,
          label.element = element_text(size = 6),
          override.aes = list(starshape = 15)
        )
      )
design <- "\n111\n111\n111\n233\n233\n"
aplot::plot_list(p1, p2, p3, design = design, tag_levels = "A")
```

Then we used  $mp\_adonis$  to perform the Permutational Multivariate Analysis of Variance based on the distance.

```
mpseMGS %<>%
    mp_adonis(
        .abundance = Abundance,
        .formula = ~ disease + response,
        distmethod = "bray",
        permutation = 9999,
        action = "add"
    )
# the result can be extracted with mp_extract_internal_attr
# mpseMGS %>% mp_extract_internal_attr(name = adonis)
```

#### 2.3.3 Different analysis in MGS level

Here, we also used  $mp\_diff\_analysis$  to detect the difference taxa, we also specified the 'force = TRUE' and 'relative = FALSE', meaning the metric of abundance (abundance) was used to perform the analysis directly without rarefied and calculated the relative abundance (Fig.S20).

```
mpseMGS %<>%
    mp_diff_analysis(
       .abundance = Abundance,
       force = TRUE,
       relative = FALSE,
       .group = disease,
       filter.p = "pvalue"
    )
library(forcats)
trda <- mpseMGS %>% mp_extract_tree()
p <- ggtree(trda, layout = 'radial') +</pre>
     geom_tiplab(size = 1.8, offset = 11) +
     geom_hilight(
         mapping = aes(
           subset = nodeClass == "Phylum",
           node = node,
           fill = label
     )
p2 <- p +
      ggnewscale::new_scale_fill() +
      geom_fruit(
         data = td_unnest(AbundanceBySample, names_repair=tidyr::tidyr_legacy),
         geom = geom_star,
         mapping = aes(
            x = fct_reorder(Sample, disease, .fun=min),
            size = Abundance,
            fill = disease,
            subset = Abundance > 0
         ),
         starshape = 13,
         offset = 0.02,
         pwidth = 1,
         grid.params = list(linetype=2)
      ) +
      scale size continuous(name="Relative Abundance (%)",range = c(1, 3)) +
      scale_fill_manual(values = cols)
p3 <- p2 +
      ggnewscale::new_scale("fill") +
      geom_fruit(
         geom = geom_col,
         mapping = aes(
                       x = LDAmean,
                       fill = Sign_disease,
                       subset = !is.na(LDAmean)
                       ),
         orientation = "y",
         offset = .05,
         pwidth = 0.5,
         width = 0.5, # the parameter of geom_col
         axis.params = list(axis = "x",
                             title = "Log10(LDA)",
```

```
title.height = 0.001,
                            title.size = 2,
                            text.size = 1.8,
                            vjust = 1),
         grid.params = list(linetype = 1)
      ) +
      ggnewscale::new_scale("size") +
      geom_point(
         data=td_filter(!is.na(pvalue)),
         mapping = aes(size = -log10(pvalue),
                       fill = Sign_disease
                   ),
         shape = 21
      ) +
      scale_size_continuous(range=c(0.5, 3)) +
      scale_fill_manual(values=cols) +
      theme(
           legend.key.height = unit(0.3, "cm"),
           legend.key.width = unit(0.3, "cm"),
           legend.spacing.y = unit(0.02, "cm"),
           legend.text = element_text(size = 7),
           legend.title = element_text(size = 9),
      )
рЗ
```

Next, we extract the abundance of the different species, then using ggplot2 (Wickham 2011) to visualize them (Fig.S21).

```
deT <- mpseMGS %>% mp_extract_tree() %>% dplyr::filter(!is.na(fdr) & isTip, keep.td=F) %>% dplyr::pull(label)
mpseMGS %>%
    mp_extract_abundance(taxa.class="OTU") %>%
    dplyr::filter(label %in% deT) %>%
    tidyr::unnest(AbundanceBySample) %>%
    ggplot(mapping=aes(x=disease, y=Abundance, fill=disease)) +
    geom_boxplot() +
    facet_wrap(facets = vars(label), nrow = 1, scales = "free", strip.position = "right") +
    ggsignif::geom_signif(comparisons=list(c("CD", "CN"))) +
    scale_fill_manual(values=cols, guide="none") +
    labs(x=NULL, y="relative abundance (%)")
```

#### The analysis of the mosquito ecology data using MicrobiotaProcess 3

MicrobiotaProcess also can be used to perform the other related ecology data analysis, besides the microbial community data. Here, we used an example data about a Mosquito ecology study (REISKIND et al. 2017) to show how to use MicrobiotaProcess to perform the analysis of the related ecology study. The data was obtained from the github<sup>3</sup>.

## Loading data and Construction of MPSE object

The 1 to 14 columns are the sample metadata including the study site, and habitat, etc. and the others columns represent the abundance of mosquito species the in each sample.

```
data <- read.csv("./data/Mosquito_ecology/data.csv", row.names=1)</pre>
abun.d <- data[, 14:36]
sample.d <- data[, 1:13]</pre>
# We implements `MPSE` function to build the `MPSE` object, which requires the abundance table (matrix-like).
mpse <- MPSE(assays=list(Abundance=t(abun.d)), colData=sample.d)</pre>
mpse
## # A MPSE-tibble (MPSE object) abstraction: 1,035 x 16
## # OTU=23 | Samples=45 | Assays=Abundance | Taxanomy=NULL
##
      OTU
               Sample Abundance Region Transect Habitat DeciduousForest
##
      <chr>>
               <chr>>
                           <int> <chr> <chr>
                                                  <chr>>
                                                                     <dbl>
               DU1.1
                              19 Durham DU1
                                                  Field
##
   1 Cx.sal
                                                                      125.
##
    2 Ae.albo DU1.1
                               0 Durham DU1
                                                  Field
                                                                      125.
##
    3 Ae.cin
               DU1.1
                               1 Durham DU1
                                                  Field
                                                                      125.
##
    4 Ae.vex
               DU1.1
                              16 Durham DU1
                                                  Field
                                                                      125.
   5 Ps.fer
               DU1.1
                               1 Durham DU1
                                                  Field
                                                                      125.
                             372 Durham DU1
##
   6 Cx.err
               DU1.1
                                                  Field
                                                                      125.
    7 Ps.col
               DU1.1
                             104 Durham DU1
                                                  Field
                                                                      125.
##
##
   8 Ae.tris DU1.1
                               0 Durham DU1
                                                                      125.
                                                  Field
   9 Cx.pip.q DU1.1
                               2 Durham DU1
                                                  Field
                                                                      125.
## 10 Ae.can
               DU1.1
                               0 Durham DU1
                                                  Field
                                                                      125.
## # ... with 1,025 more rows, and 9 more variables: EvergreenForest <dbl>,
       Grassland <dbl>, MixedForest <dbl>, ShrubScrub <dbl>, BarrenLand <dbl>,
## #
       Building <dbl>, Pavement <dbl>, CultivatedCrops <dbl>, TrapNights <int>
## #
```

#### Alpha diversity analysis of the Mosquito ecology study 3.2

0 Durham DU1

The MicrobiotaProcess provides some verbs of dplyr, which allows user to explore the MPSE class effectively and develop reproducible and human-readable pipelines

```
cols = c("lightgoldenrod1", "orange", "chartreuse2", "chartreuse4", "darkgreen")
# Adjusting the order of Habitat
mpse %<>%
   dplyr::mutate(
     Habitat = factor(
       Habitat,
       levels = c("Field", "NearField", "Edge", "NearForest", "Forest")
    )
   )
mpse
## # A MPSE-tibble (MPSE object) abstraction: 1,035 x 16
## # OTU=23 | Samples=45 | Assays=Abundance | Taxanomy=NULL
##
      OTU
               Sample Abundance Region Transect Habitat DeciduousForest
##
      <chr>
               <chr>>
                           <int> <chr> <chr>
                                                 <fct>
                                                                    <dbl>
    1 Cx.sal
               DU1.1
                              19 Durham DU1
```

2 Ae.albo DU1.1

##

Field

Field

125.

125.

<sup>&</sup>lt;sup>3</sup>https://github.com/rgriff23/Mosquito ecology

```
3 Ae.cin
               DU1.1
                              1 Durham DU1
                                                Field
                                                                    125.
##
                                                                    125.
##
   4 Ae.vex
               DU1.1
                             16 Durham DU1
                                                Field
                              1 Durham DU1
##
   5 Ps.fer
               DU1.1
                                                Field
                                                                    125.
##
               DU1.1
                            372 Durham DU1
                                                Field
                                                                    125.
   6 Cx.err
                            104 Durham DU1
##
   7 Ps.col
              DU1.1
                                                Field
                                                                    125.
   8 Ae.tris DU1.1
                              0 Durham DU1
                                                Field
                                                                    125.
##
## 9 Cx.pip.q DU1.1
                              2 Durham DU1
                                                Field
                                                                    125.
## 10 Ae.can DU1.1
                              0 Durham DU1
                                                Field
                                                                    125.
## # ... with 1,025 more rows, and 9 more variables: EvergreenForest <dbl>,
       Grassland <dbl>, MixedForest <dbl>, ShrubScrub <dbl>, BarrenLand <dbl>,
## #
       Building <dbl>, Pavement <dbl>, CultivatedCrops <dbl>, TrapNights <int>
# force=TRUE meaning the Abundance will be used to calculate the alpha index without rarefaction
mpse %<>% mp_cal_alpha(.abundance=Abundance, force=TRUE)
# test the relationship between the Observe Species and Habitat or Shannon and Habitat.
tb1 <- mpse %>% mp_extract_sample() %>% lm(formula=Observe ~ Habitat, data=.) %>% anova() %>% broom::tidy()
tb2 <- mpse %>% mp_extract_sample() %>% lm(formula=Shannon ~ Habitat, data=.) %>% anova() %>% broom::tidy()
```

The result of ANOVA test revealed that the richness of the mosquito species was significantly associated with the **habitat**. Then the result was visualized by  $mp\_plot\_alpha$  (Fig.S22).

```
p.alpha <- mpse %>%
     mp_plot_alpha(.group = Habitat, .alpha = c(Observe, Shannon), test = NULL) +
     scale_fill_manual(values = cols) +
     scale_color_manual(values = cols) +
     theme(legend.position = "none")
library(ggpp)
# building the table layer
tb1 %<>% dplyr::slice(1) %>% select(statistic, p.value) %>% round(3)
tb2 %<>% dplyr::slice(1) %>% select(statistic, p.value) %>% round(3)
df <- tibble(npcx=c(0.9, 0.9), npcy=c(0.05, 0.05), tb=list(tb1, tb2), Measure=c("Observe", "Shannon"))
p.alpha <- p.alpha +
           geom_table_npc(
             data = df,
             mapping = aes(
               npcx = npcx,
               npcy = npcy,
               label = tb
             ),
             table.theme = ttheme_gtminimal
p.alpha
```

### 3.3 Beta Diversity Analysis of the Mosquito ecology study

Here, we use the cca (constrained correspondence analysis) to test which environment factor is related to the Mosquito species in the habitat (Fig.S23).

```
Pavement+
             CultivatedCrops
    )
mpse
## # A MPSE-tibble (MPSE object) abstraction: 1,035 x 26
## # OTU=23 | Samples=45 | Assays=Abundance, NormAbun | Taxanomy=NULL
               Sample Abundance NormAbun Region Transect Habitat DeciduousForest
##
      OTU
                                    <dbl> <chr> <chr>
##
      <chr>>
                           <int>
                                                            <fct>
               <chr>
                                                                               <dbl>
   1 Cx.sal
##
               DU1.1
                              19
                                    0.436 Durham DU1
                                                           Field
                                                                               125.
    2 Ae.albo DU1.1
                               0
                                    0
                                           Durham DU1
##
                                                           Field
                                                                               125.
##
    3 Ae.cin
               DU1.1
                               1
                                    0.1
                                           Durham DU1
                                                           Field
                                                                               125.
##
   4 Ae.vex
               DU1.1
                              16
                                    0.4
                                           Durham DU1
                                                           Field
                                                                               125.
##
   5 Ps.fer
               DU1.1
                               1
                                    0.1
                                           Durham DU1
                                                           Field
                                                                               125.
                             372
                                    1.93 Durham DU1
##
   6 Cx.err
               DU1.1
                                                           Field
                                                                               125.
##
   7 Ps.col
               DU1.1
                             104
                                    1.02 Durham DU1
                                                           Field
                                                                               125.
##
   8 Ae.tris DU1.1
                               0
                                    0
                                           Durham DU1
                                                           Field
                                                                               125.
##
   9 Cx.pip.q DU1.1
                               2
                                    0.141 Durham DU1
                                                           Field
                                                                               125.
## 10 Ae.can
               DU1.1
                               0
                                    0
                                           Durham DU1
                                                            Field
                                                                                125.
## # ... with 1,025 more rows, and 18 more variables: EvergreenForest <dbl>,
       Grassland <dbl>, MixedForest <dbl>, ShrubScrub <dbl>, BarrenLand <dbl>,
## #
       Building <dbl>, Pavement <dbl>, CultivatedCrops <dbl>, TrapNights <int>,
## #
       Observe <dbl>, Chao1 <dbl>, ACE <dbl>, Shannon <dbl>, Simpson <dbl>,
## #
       Pielou <dbl>, CCA1 (25.28%) <dbl>, CCA2 (7.34%) <dbl>, CCA3 (3.39%) <dbl>
The raw result of pCCA was added the internal_attr, which can be extract by mp_extract_internal_attr with specific
name=cca. Then it can be performed the significance test using the functions of vegan (Oksanen et al. 2020), such as
anova.cca, permutest.
# Extract the raw result of cca analysis
# And significance test with anova
mpse %>%
    mp_extract_internal_attr(name=cca) %>%
    anova()
## Permutation test for cca under reduced model
## Permutation: free
## Number of permutations: 999
##
## Model: cca(formula = x ~ DeciduousForest + EvergreenForest + Grassland + MixedForest + ShrubScrub + Conditio
                               F Pr(>F)
##
            Df ChiSquare
## Model
             5
                 0.38999 4.4365 0.001 ***
## Residual 35
                 0.61534
## Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
Further we used mp\_envfit to identity the environment variables that were significantly associated with the mosquito commu-
# fits environmental vectors onto cca
mpse %<>%
    mp_envfit(
       .ord = cca,
       .env = c(
          DeciduousForest,
          EvergreenForest,
          Grassland,
          MixedForest,
          ShrubScrub
```

),

action = "add",

```
permutation = 9999
    )
# Extract the raw result of envfit analysis
mpse %>% mp_extract_internal_attr(name=cca_envfit)
##
## ***VECTORS
##
##
                        CCA1
                                 CCA2
                                          CCA3
                                                   r2 Pr(>r)
## DeciduousForest 0.42979 0.90272 -0.01945 0.3804 0.0020 **
## EvergreenForest 0.91539 -0.34612 -0.20559 0.5557 0.0001 ***
## Grassland
                   -0.97679 -0.21356  0.01639  0.7216  0.0001 ***
## MixedForest
                    0.77120 -0.25826  0.58186  0.1936  0.0929 .
## ShrubScrub
                   -0.73942 0.22976 -0.63283 0.2595 0.0537 .
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
## Permutation: free
## Number of permutations: 9999
Then we used mp\_plot\_ord to visualize the result of pCCA (Fig.S23).
# visualization only pCCA
f <- mpse %>%
     mp_plot_ord(
       .ord = cca,
       .group = Habitat,
       .size = Observe,
       .starshape = Region,
       show.side = FALSE,
       show.envfit = FALSE,
       colour = 'black',
       bg.colour = 'white'
     ) +
     scale_starshape_manual(values=c(1, 13, 15)) +
     scale_fill_manual(
        values = cols,
        guide = guide_legend(
          override.aes = list(starshape=15)
     ) +
     scale_size_continuous(
       range = c(1, 3),
       guide = guide_legend(override.aes = list(starshape=15))
     ) +
        legend.key.height = unit(0.3, "cm"),
        legend.key.width = unit(0.3, "cm"),
        legend.spacing.y = unit(0.02, "cm"),
        legend.text = element_text(size = 7),
        legend.title = element_text(size = 9),
     )
# visualization with envfit result
p <- mpse %>%
     mp_plot_ord(
       .ord = cca,
       .group = Habitat,
       .size = Observe,
       .starshape = Region,
       show.side = FALSE,
```

```
show.envfit = TRUE,
       colour = "black",
       bg.colour = "white"
    ) +
     scale_starshape_manual(values=c(1, 13, 15)) +
     scale_fill_manual(
       values = cols,
       guide = guide_legend(
          override.aes = list(starshape=15)
       )
    ) +
     scale_size_continuous(
      range = c(1, 3),
      guide = guide_legend(override.aes = list(starshape=15))
     ) +
     theme (
        legend.key.height = unit(0.3, "cm"),
        legend.key.width = unit(0.3, "cm"),
        legend.spacing.y = unit(0.02, "cm"),
       legend.text = element_text(size = 7),
       legend.title = element_text(size = 9),
     )
aplot::plot_list(f, p, tag_levels="A")
```

# 3.4 The distribution of Mosquito species in the study.

We used  $mp\_cal\_abundance$  and  $mp\_plot\_abundance$  to calculated and visualized the abundance of the Mosquito species in the study (Fig.S24).

```
cols2 <- c("deepskyblue", "yellow", "#FF9933")</pre>
# The theme and scale of fill of heatmap
Abund.char <- list(
           scale_fill_viridis_c(option = "H"),
           theme (
             axis.text.x = element_text(size = 6),
             axis.text.y = element_text(size = 8),
             legend.title = element_text(size = 7),
             legend.text = element_text(size = 5),
             legend.key.width = unit(0.3, "cm"),
             legend.key.height = unit(0.3, "cm")
           )
      )
# The theme and legend of annotate bar of 'Habitat' variable
Habitat.char <- list(</pre>
           scale fill manual(values = cols),
           theme(
             legend.key.height = unit(0.3, "cm"),
             legend.key.width = unit(0.3, "cm"),
             legend.spacing.y = unit(0.02, "cm"),
             legend.text = element_text(size = 7),
             legend.title = element_text(size = 9)
           )
# The theme and legend of annotate bar of 'Region' variable
Region.char <- list(
```

```
scale_fill_manual(values = cols2),
             legend.key.height = unit(0.3, "cm"),
             legend.key.width = unit(0.3, "cm"),
             legend.spacing.y = unit(0.02, "cm"),
             legend.text = element_text(size = 7),
             legend.title = element_text(size = 9)
      )
# visualization of the count abundance.
p.count <- mpse %>%
    mp_cal_abundance(
      .abundance = Abundance,
      force = T,
      relative = F
    ) %>%
    mp_plot_abundance(
      .abundance = Abundance,
      force = T,
      relative = F,
      geom = "heatmap",
      topn = "all",
      .group = c(Habitat, Region)
    ) %>%
    set_scale_theme(
      x = Abund.char,
      aes_var = Abundance
    ) %>%
    set_scale_theme(
     x = Habitat.char,
     aes var = Habitat
    ) %>%
    set_scale_theme(
      x = Region.char,
      aes_var = Region
# visualization of the relative abundance
p.rel <- mpse %>%
    mp_cal_abundance(
      .abundance = Abundance,
      force = T,
      relative = T
    ) %>%
    mp_plot_abundance(
      .abundance = Abundance,
      force = T,
      relative = T,
      geom = "heatmap",
      topn = "all",
      .group = c(Habitat, Region)
    ) %>%
    set_scale_theme(
      x = Abund.char,
      aes_var = RelAbundance
    ) %>%
    set_scale_theme(
      x = Habitat.char,
      aes_var = Habitat
```

```
) %>%
set_scale_theme(
    x = Region.char,
    aes_var = Region
)
aplot::plot_list(p.count, p.rel, tag_levels="A")
```

Then We can use  $mp\_diff\_analysis$  to identify the species that have significant difference abundance between the **field** and **forest**. We found the Cx.sal ( $Culex\ salinarius$ ) and Ps.col ( $Psorophora\ columbiae$ ) were significantly enriched in **field**, However, the Ae.albo ( $Aedes\ albopicta$ ), Ae.cin ( $Aedes\ cinereus$ ), Ps.fer ( $Psorophora\ ferox$ ), Ae.tris ( $Aedes\ triseriatus$ ), Ae.can ( $Aedes\ canadensis$ ), Ae.hen ( $Aedes\ hendersoni$ ), Ae.atl ( $Aedes\ atlanticus$ ) and Ae.dup ( $Aedes\ dupreei$ ) were significantly enriched in the **forest** 

```
dplyr::filter(Habitat %in% c("Field", "Forest")) %>%
    dplyr::mutate(Habitat = as.vector(Habitat)) %>%
    mp diff analysis(.abundance=Abundance, force=T, relative=T, .group=Habitat) %>%
    mp_extract_feature() %>%
    dplyr::filter(fdr<=0.05) %>%
    print(width=200)
## # A tibble: 10 x 8
##
      OTU
              AbundanceBySample LDAupper LDAmean LDAlower Sign_Habitat
                                                                             pvalue
##
      <chr>
              st>
                                     <dbl>
                                             <dbl>
                                                       <dbl> <chr>
                                                                              <dbl>
##
    1 Cx.sal
              <tibble [18 x 6]>
                                      4.96
                                              4.92
                                                        4.87 Field
                                                                           0.00705
##
   2 Ae.albo <tibble [18 x 6]>
                                      4.83
                                              4.79
                                                        4.75 Forest
                                                                           0.000229
                                      4.36
   3 Ae.cin <tibble [18 x 6]>
                                              4.31
                                                        4.25 Forest
                                                                           0.0159
   4 Ps.fer <tibble [18 x 6]>
                                      4.94
                                              4.90
                                                        4.87 Forest
##
                                                                           0.00122
   5 Ps.col <tibble [18 x 6]>
##
                                      5.26
                                              5.24
                                                        5.22 Field
                                                                           0.000327
##
   6 Ae.tris <tibble [18 x 6]>
                                      4.49
                                              4.46
                                                        4.43 Forest
                                                                           0.000530
   7 Ae.can \langle \text{tibble } [18 \times 6] \rangle
                                      4.28
                                              4.24
                                                        4.19 Forest
                                                                           0.0119
##
   8 Ae.hen <tibble [18 x 6]>
                                      4.28
                                              4.23
                                                        4.18 Forest
                                                                           0.000483
##
   9 Ae.atl <tibble [18 x 6]>
                                      4.59
                                              4.56
                                                        4.52 Forest
                                                                           0.00311
                                      4.03
##
  10 Ae.dup <tibble [18 x 6]>
                                              3.96
                                                        3.88 Forest
                                                                           0.0119
##
          fdr
##
        <dbl>
##
   1 0.0211
   2 0.00278
   3 0.0334
##
   4 0.00513
##
##
   5 0.00278
   6 0.00278
##
   7 0.0278
##
   8 0.00278
##
   9 0.0109
## 10 0.0278
```

# 4 Session information

mpse %>%

Here is the output of sessionInfo() on the system on which this document was compiled:

en\_US.UTF-8 ## ctype ## Asia/Shanghai tz 2021-12-28 ## date ## - Packages ----date lib source \* version package ## 1.56.1 2021-10-29 [1] Bioconductor ## AnnotationDbi ## ape 5.6 2021-12-22 [1] local \* 0.1.1 ## aplot 2021-09-22 [1] CRAN (R 4.1.1) 0.2.1 0.3.1 1.3.0 2019-03-21 [1] CRAN (R 4.1.1) ## assertthat attempt ## 2020-05-03 [1] CRAN (R 4.1.1) 2021-10-27 [1] CRAN (R 4.1.1) ## backports \* 2.54.0 ## Biobase 2021-10-26 [1] Bioconductor \* 0.40.0 2021-10-26 [1] Bioconductor ## BiocGenerics ## BiocManager 1.30.16 2021-06-15 [1] CRAN (R 4.1.1) ## BiocParallel 1.28.0 2021-10-26 [1] Bioconductor 1.22.0 2021-10-26 [1] Bioconductor ## biomformat ## Biostrings 2.62.0 2021-10-26 [1] Bioconductor ## 4.0.4 2020-08-04 [1] CRAN (R 4.1.1) bit bit64 4.0.5 2020-08-30 [1] CRAN (R 4.1.1) ## 1.0-7 1.2.2 0.24 ## bitops 2021-04-24 [1] CRAN (R 4.1.1) blob 2021-07-23 [1] CRAN (R 4.1.1) ## 0.24 2021-09-02 [1] CRAN (R 4.1.1) ## bookdown ## broom 0.7.10 2021-10-31 [1] CRAN (R 4.1.1) 2021-10-06 [1] CRAN (R 4.1.1) bslib 0.3.1 ## 1.0.6 2021-08-19 [1] CRAN (R 4.1.1) ## cachem3.7.0 2021-04-20 [1] CRAN (R 4.1.1) ## callr 7.3-19 ## class 2021-05-03 [1] CRAN (R 4.1.1) ## classInt 0.4 - 32020-04-07 [1] CRAN (R 4.1.1) ## cli 3.1.0 2021-10-27 [1] CRAN (R 4.1.1) ## 2.1.2 2021-04-17 [1] CRAN (R 4.1.1) cluster clusterProfiler \* 4.2.0 2021-10-26 [1] Bioconductor 0.2-18 ## codetools 2020-11-04 [1] CRAN (R 4.1.1) \* 1.4-2 2.0-2 0.3.1 \* 1.0.4 0.4.3 1.4.2 ## 2021-10-08 [1] CRAN (R 4.1.1) coin colorspace 2021-06-24 [1] CRAN (R 4.1.1) ## 2020-12-17 [1] CRAN (R 4.1.1) ## config conflicted 2019-06-21 [1] CRAN (R 4.1.1) ## 2020-11-24 [1] CRAN (R 4.1.1) ## corrr 2021-10-29 [1] CRAN (R 4.1.1) crayon 1.14.2 2021-09-27 [1] CRAN (R 4.1.1) ## data.table DBI 1.1.1 2021-01-15 [1] CRAN (R 4.1.1) ## 0.20.0 2021-10-26 [1] Bioconductor ## DelayedArray 1.4.0 2021-09-28 [1] CRAN (R 4.1.1) ## desc 2021-09-23 [1] CRAN (R 4.1.1) ## digest 0.6.28 ## DO.db 2.9 2021-12-13 [1] Bioconductor ## 2021-09-03 [1] CRAN (R 4.1.1) dockerfiler 0.1.4 DOSE 3.20.1 2021-11-18 [1] Bioconductor 2015-07-09 [1] CRAN (R 4.1.1) ## downloader 0.4 ## 1.0.7 2021-06-18 [1] CRAN (R 4.1.1) dplyr ## DT 0.19 2021-09-02 [1] CRAN (R 4.1.1) ## 1.1.0 2021-02-20 [1] CRAN (R 4.1.1) dtplyr ## e1071 1.7-9 2021-09-16 [1] CRAN (R 4.1.1) 2021-10-26 [1] Bioconductor ## edgeR \* 3.36.0 0.3.2 2021-04-29 [1] CRAN (R 4.1.1) ellipsis ## enrichplot \* 1.14.1 2021-10-31 [1] Bioconductor ## evaluate 0.14 2019-05-28 [1] CRAN (R 4.1.1) 0.5.0 2.1.0 1.1.0 ## fansi 2021-05-25 [1] CRAN (R 4.1.1) farver 2021-02-28 [1] CRAN (R 4.1.1) 2021-01-25 [1] CRAN (R 4.1.1) ## fastmap 1.1-3 2021-07-23 [1] CRAN (R 4.1.1) fastmatch

```
##
                            1.20.0
                                       2021-10-26 [1] Bioconductor
    fgsea
                                       2021-01-27 [1] CRAN (R 4.1.1)
##
    forcats
                          * 0.5.1
                            1.5.1
                                       2020-10-15 [1] CRAN (R 4.1.1)
##
    foreach
##
    fs
                            1.5.0
                                       2020-07-31 [1] CRAN (R 4.1.1)
                                       2021-10-25 [1] CRAN (R 4.1.1)
##
    generics
                            0.1.1
    {\tt GenomeInfoDb}
                          * 1.30.0
                                       2021-10-26 [1] Bioconductor
##
    {\tt GenomeInfoDbData}
                                       2021-10-29 [1] Bioconductor
##
                            1.2.7
##
    GenomicRanges
                          * 1.46.0
                                       2021-10-26 [1] Bioconductor
##
    ggalluvial
                            0.12.3
                                       2020-12-05 [1] CRAN (R 4.1.1)
                            0.3.3
                                       2021-03-05 [1] CRAN (R 4.1.1)
##
    ggforce
##
    ggfun
                            0.0.4
                                       2021-09-17 [1] CRAN (R 4.1.1)
                            0.2.0
##
    ggh4x
                                       2021-08-21 [1] CRAN (R 4.1.1)
##
    gghalves
                            0.1.1
                                       2020-11-08 [1] CRAN (R 4.1.1)
##
    ggnewscale
                          * 0.4.5
                                       2021-01-11 [1] CRAN (R 4.1.1)
##
                          * 3.3.5
                                       2021-06-25 [1] CRAN (R 4.1.1)
    ggplot2
##
    ggplotify
                            0.1.0
                                       2021-09-02 [1] CRAN (R 4.1.1)
                          * 0.4.2
                                       2021-07-31 [1] CRAN (R 4.1.1)
##
    ggpp
##
    ggraph
                            2.0.5
                                       2021-02-23 [1] CRAN (R 4.1.1)
                          * 0.9.1
                                       2021-01-15 [1] CRAN (R 4.1.1)
##
    ggrepel
                            0.2.0
                                       2021-12-11 [1] CRAN (R 4.1.1)
##
    ggside
                            0.6.3
                                       2021-09-09 [1] CRAN (R 4.1.1)
##
    ggsignif
                          * 1.0.3
                                       2021-12-03 [1] Github (xiangpin/ggstar@5e1f018)
##
    ggstar
                          * 3.3.0.901 2021-12-10 [1] Bioconductor
##
    ggtree
##
    ggtreeExtra
                          * 1.5.1
                                       2021-11-24 [1] Bioconductor
                                       2020-05-05 [1] CRAN (R 4.1.1)
                            0.3.0
##
    ggupset
                                       2021-07-07 [1] CRAN (R 4.1.1)
##
    ggVennDiagram
                          * 1.1.4
                                       2021-11-07 [1] CRAN (R 4.1.1)
##
    glue
                            1.5.0
##
    GO.db
                            3.14.0
                                       2021-11-30 [1] Bioconductor
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    {\tt GOSemSim}
                            2.20.0
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                            0.7.1
##
    graphlayouts
                                       2020-10-26 [1] CRAN (R 4.1.1)
                            2.3
                                       2017-09-09 [1] CRAN (R 4.1.1)
##
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##
    gridGraphics
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    gtable
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##
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                                       2021-08-25 [1] CRAN (R 4.1.1)
                            0.5.2
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    httr
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##
##
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                                       2021-10-15 [1] CRAN (R 4.1.1)
                          * 2.28.0
                                       2021-10-26 [1] Bioconductor
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                            1.0.13
##
    iterators
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##
    jquerylib
                            0.1.4
                                       2021-04-26 [1] CRAN (R 4.1.1)
##
    jsonlite
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                                       2020-12-09 [1] CRAN (R 4.1.1)
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                            1.34.0
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##
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##
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                                       2021-09-22 [1] CRAN (R 4.1.1)
##
    lazyeval
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                                       2019-03-15 [1] CRAN (R 4.1.1)
##
    libcoin
                            1.0-9
                                       2021-09-27 [1] CRAN (R 4.1.1)
    lifecycle
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                                       2021-09-24 [1] CRAN (R 4.1.1)
##
##
    limma
                          * 3.50.0
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##
    locfit
                            1.5 - 9.4
                                       2020-03-25 [1] CRAN (R 4.1.1)
##
                          * 2.0.1
                                       2020-11-17 [1] CRAN (R 4.1.1)
    magrittr
##
   MASS
                            7.3-54
                                       2021-05-03 [1] CRAN (R 4.1.1)
                                       2021-06-01 [1] CRAN (R 4.1.1)
##
   Matrix
                            1.3 - 4
                          * 1.6.0
                                       2021-10-26 [1] Bioconductor
   MatrixGenerics
```

```
matrixStats
                          * 0.61.0
                                      2021-09-17 [1] CRAN (R 4.1.1)
##
                                      2021-01-26 [1] CRAN (R 4.1.1)
##
    memoise
                            2.0.0
                            1.8-38
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##
    mgcv
##
    MicrobiomeProfiler
                          * 1.0.0
                                      2021-10-26 [1] Bioconductor
    MicrobiotaProcess
                          * 1.7.4.991 2021-12-28 [1] Bioconductor
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##
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                                      2021-09-28 [1] CRAN (R 4.1.1)
    mime
                            0.2-23
                                      2020-03-05 [1] CRAN (R 4.1.1)
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                                      2021-04-29 [1] CRAN (R 4.1.1)
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##
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                            0.5.0
    mvtnorm
                                      2021-10-08 [1] CRAN (R 4.1.1)
##
                            1.1-3
##
    nlme
                            3.1-153
                                      2021-09-07 [1] CRAN (R 4.1.1)
##
    patchwork
                          * 1.1.1
                                      2020-12-17 [1] CRAN (R 4.1.1)
##
    permute
                            0.9 - 5
                                      2019-03-12 [1] CRAN (R 4.1.1)
                                      2021-10-18 [1] CRAN (R 4.1.1)
##
    pillar
                            1.6.4
    pkgbuild
                            1.2.0
                                      2020-12-15 [1] CRAN (R 4.1.1)
##
    pkgconfig
                            2.0.3
                                      2019-09-22 [1] CRAN (R 4.1.1)
                            1.2.2
                                      2021-09-11 [1] CRAN (R 4.1.1)
##
    pkgload
##
    plyr
                            1.8.6
                                      2020-03-03 [1] CRAN (R 4.1.1)
                            0.1 - 7
                                      2013-12-03 [1] CRAN (R 4.1.1)
##
    png
                            1.10-0
                                      2019-03-14 [1] CRAN (R 4.1.1)
##
    polyclip
    preprocessCore
                            1.56.0
                                      2021-10-26 [1] Bioconductor
##
                            1.1.1
                                      2020-01-24 [1] CRAN (R 4.1.1)
##
    prettyunits
                            3.5.2
                                      2021-04-30 [1] CRAN (R 4.1.1)
##
    processx
##
    promises
                            1.2.0.1
                                      2021-02-11 [1] CRAN (R 4.1.1)
                                      2021-06-07 [1] CRAN (R 4.1.1)
                            0.4 - 26
##
    proxy
                                      2021-02-28 [1] CRAN (R 4.1.1)
##
                            1.6.0
    ps
                                      2020-04-17 [1] CRAN (R 4.1.1)
##
    purrr
                            0.3.4
    qvalue
                            2.26.0
                                      2021-10-26 [1] Bioconductor
##
##
    R6
                            2.5.1
                                      2021-08-19 [1] CRAN (R 4.1.1)
##
    RColorBrewer
                            1.1-2
                                      2014-12-07 [1] CRAN (R 4.1.1)
                            1.0.7
                                      2021-07-07 [1] CRAN (R 4.1.1)
##
    Rcpp
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                                      2021-09-17 [1] CRAN (R 4.1.1)
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## [1] /mnt/d/UbuntuApps/R/4.1.1/lib/R/library

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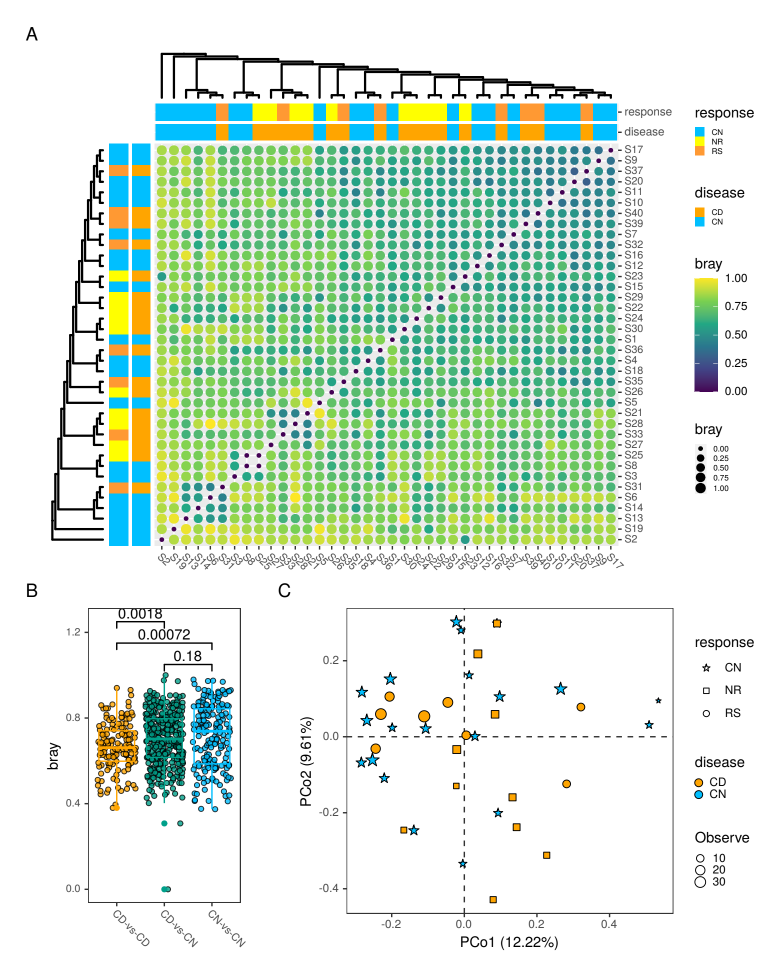


Fig. S19: The distance heatmap and boxplot and the PCoA plot based on MGS data

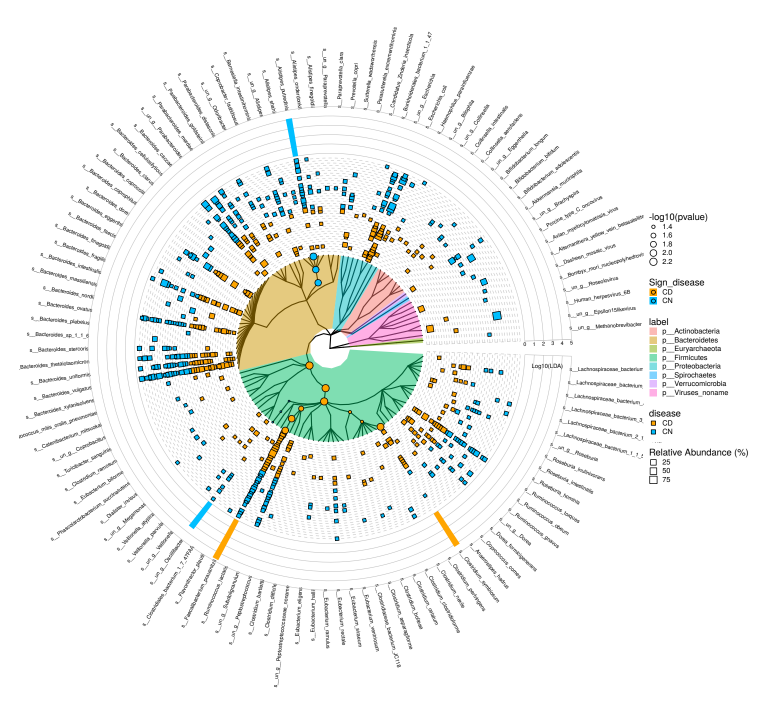


Fig. S20: The result of different analysis based on MGS data

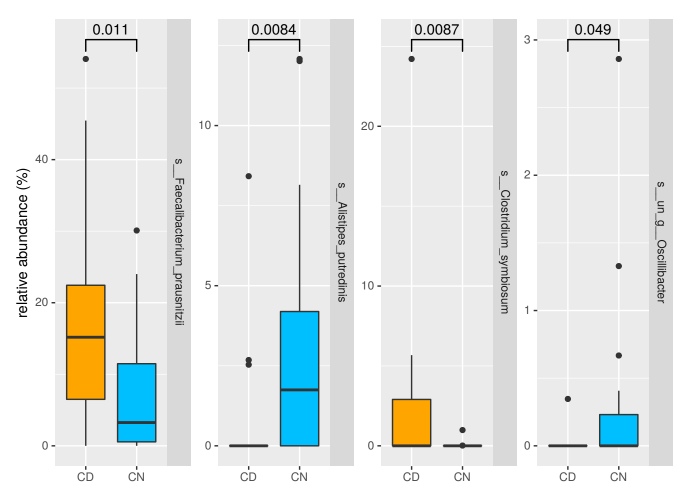


Fig. S21: The abundance boxplot of the different species between the CD and control group

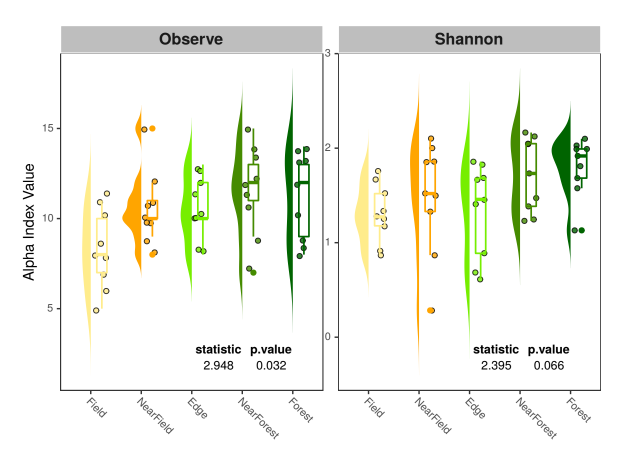


Fig. S22: The raincloud plot of the alpha diversity of the Mosquito ecology community. The result of the alpha diversity analysis about the Mosquito ecology study showed that the Mosquito species richness gradually increases from field to forest (field --> near field --> edge --> near field --> forest).

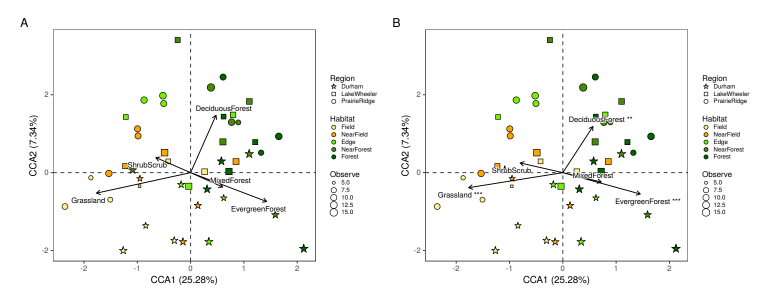


Fig. S23: The CCA plot of the Mosquito ecology study. Each point represents one sample, the size of the points represents the observe species of the corresponding sample, the color of the points represents the habitat of the corresponding sample, the shape of points represents the Region of the corresponding sample. And the arrows represent the environment factors, the marked ones by star represent significant related to the Mosquito communities in the study (\* 0.05, \*\* 0.01, \*\*\* 0.001).

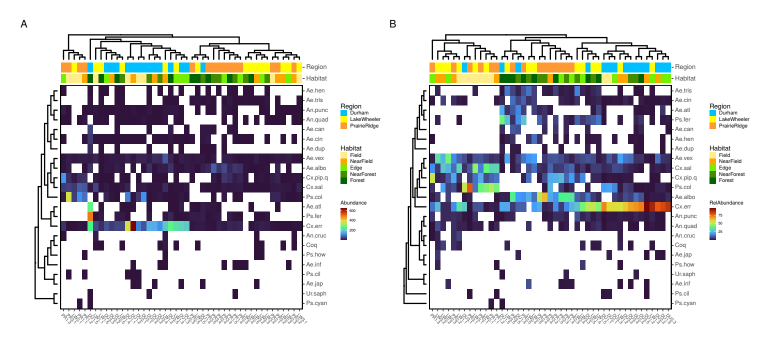


Fig. S24: The heatmap of the abundance and relative abundance of the Mosquito species.