2025 NDMC Metagenome workshop

Yincheng Chen

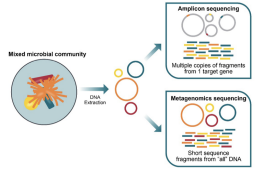
Created on March 28, 2025

Contents

Introduction to Microbiota Data	2
Practical guide to microbiota data	2
Pipeline for 16S rRNA amplicon	4
DADA2 pipeline tutorial	4
Diversity Analysis	8
Alpha diversity	9
Beta diversity	15
Community composition	20
Environmental Analysis	21
Mantel test	22
BioENV	22
CCA	23
DCA	26
RDA	29
Differential Taxa	30
ALDEx2	30
ANCOM	32
$\operatorname{edgeR} \ \ldots \ $	34
LEfSe	34
PreLect	37
Basal feature visualization	43
Functional Prediction	47
PICRUSt2	47
RPM	52

Co-Occurrence Network	5 6
SPIEC-EASI	. 56
SparCC	. 57
Cluster identification	. 58
Guide to Shotgun Metagenomic Data	5 9
Read bases pipeline	. 59
Assembly bases pipeline	. 60

Introduction to Microbiota Data

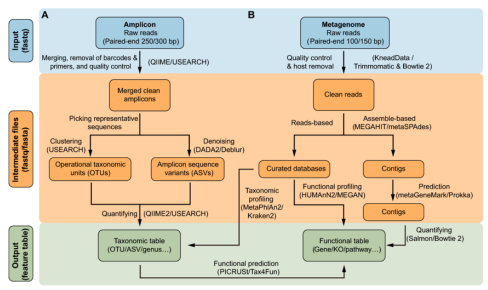


https://blog.crownbio.com/understanding-the-microbiome-using-genomic-sequencing-and-analysis

Pros and Cons

- · Quantify statistically
- Low resolution (genus level)
- · Low computing cost
- Biological functions.
- · High resolution (species/strain level)
- · High computing cost.

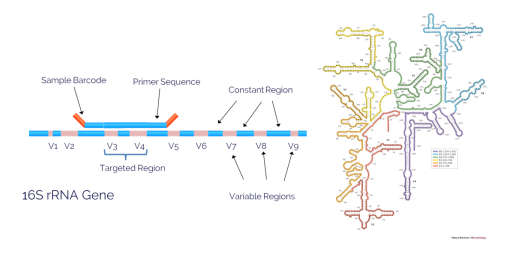
Practical guide to microbiota data



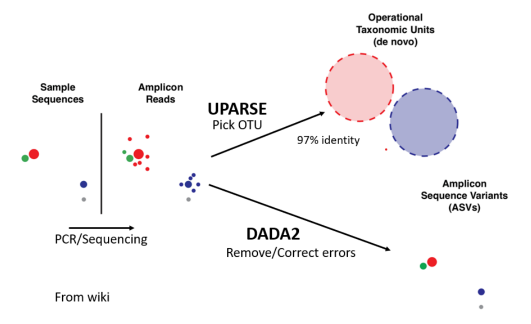
Liu, Y. X., Qin, Y., Chen, T., Lu, M., Qian, X., Guo, X., & Bai, Y. (2021). A practical guide to amplicon and metagenomic analysis of microbiome data. *Protein & cell*, *12*(5), 315-330.

16s rRNA amplicon

Highly conserved in bacteria and archaea, but contains nine hypervariable region V1-V9



Sequence process for amplicon data



Reference database:

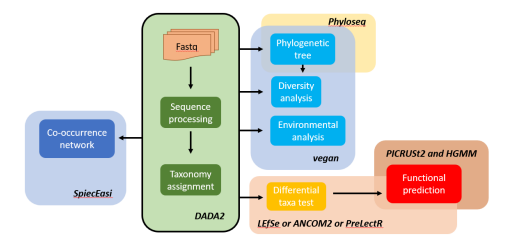
For bacteria

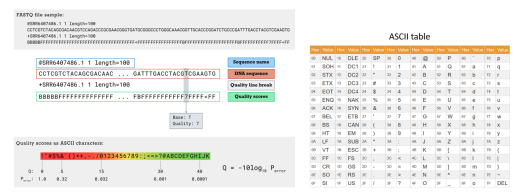
- 1. **SILVA** v138.2, (2024, Nov.)
- 2. **RDP** v19, (2023, Aug.)
- 3. GreenGenes v2, (2024, Sep.)

For fungi

• **UNITE v10**, (2025, Feb.)

Pipeline for 16S rRNA amplicon





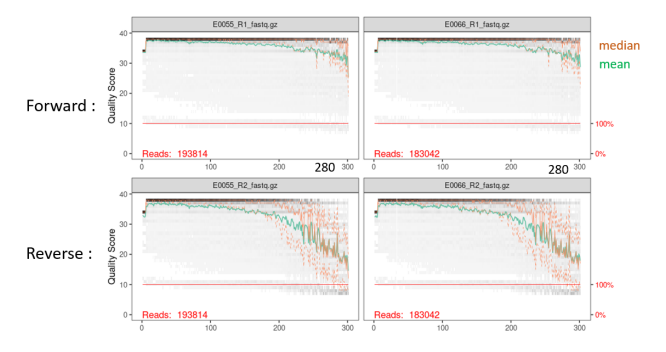
FASTQ format

Q=20, error rate = 0.01 (1%) Q=30, error rate = 0.001 (0.1%)

DADA2 pipeline tutorial

Please refer directly to the original page

Step 1. Inspect read quality profiles



Step 2. Filter and trim

The key parameters for this step are truncLen and trimLeft.

To prevent adapter sequence contamination, use trimLeft to remove bases from the 5' end of the reads.

To ensure sequence quality, use truncLen to truncate reads after assignment.

There are two important criteria to consider:

- After filtering, at least 80% of the reads should be retained.
- The V3-V4 region typically ranges from 360 to 420 nt in length, so the combined length of paired-end reads after truncation should be at least 400 nt to ensure sufficient overlap.

```
out <- filterAndTrim(fnFs, filtFs, fnRs, filtRs,</pre>
                              truncLen=c(280,200), trimLeft=10,
                              maxN=0, maxEE=c(2,2), truncQ=2, rm.phix=TRUE,
                              compress=TRUE, multithread=TRUE)
boxplot(out[,2]/out[,1])
                                                                                       c(200,200)
                                                                 c(280,200)
                                                              0.865
 > head(out)
 E0055_R1_fastq.gz
                            161537
 E0066_R1_fastq.gz
E0067_R1_fastq.gz
E0072_R1_fastq.gz
E0084_R1_fastq.gz
                   183042
                   241337
                            205745
                   156209
215914
                                                              0.845
Criterion1: out[,2]/out[,1] > 0.9 \#0.8
                                                              0.835
Criterion2: (280-10) + (200-10) > 400 #320
```

Step 3. Sequence processing

Follow the standard procedures and verify the length of the merged sequences.

```
# Learn the Error Rates
errF <- learnErrors(filtFs, multithread=TRUE)
...

# Sample Inference
dadaFs <- dada(filtFs, err=errF, multithread=TRUE)
...

# Merge paired reads
mergers <- mergePairs(dadaFs, filtFs, dadaRs, filtRs, verbose=TRUE)
seqtab <- makeSequenceTable(mergers)
dim(seqtab) # [n_smaple, n_ASVs]

# Inspect distribution of sequence lengths
hist(nchar(getSequences(seqtab)))

# Remove chimeras
seqtab.nochim <- removeBimeraDenovo(seqtab, method="consensus", multithread=TRUE, verbose=TRUE)
...

# Assign taxonomy
taxa <- assignTaxonomy(seqtab.nochim, "~/tax/silva_nr_v132_train_set.fa.gz", multithread=TRUE)
taxa <- addSpecies(taxa, "~/tax/silva_species_assignment_v132.fa.gz")</pre>
```

Step 4. Sample QC

```
getN <- function(x) sum(getUniques(x))</pre>
track <- cbind(out, sapply(dadaFs, getN), sapply(dadaRs, getN), sapply(mergers, getN), rowSums(seqtab.n</pre>
colnames(track) <- c("input", "filtered", "denoisedF", "denoisedR", "merged", "nonchim")</pre>
rownames(track) <- sample.names</pre>
mitochondria_id <- rownames(taxa)[taxa[,5] %in% "Mitochondria"]</pre>
unclassification_id <-rownames(taxa)[taxa[,1] != "Bacteria" | is.na(taxa[,2])]
keep_id <- rownames(taxa)[!rownames(taxa) %in% mitochondria_id)]
get_seqN <- function(table,list){</pre>
 if(length(list) == 0) { return(rep(0,nrow(table)))
 } else if(length(list) > 1) { return(rowSums(table[,list]))
  } else { return(sapply(table[,list],sum))}
}
track <- cbind(track, get_seqN(seqtab, mitochondria_id), get_seqN(seqtab,keep_id))</pre>
colnames(track) <- c("input", "filtered", "denoisedF", "denoisedR", "merged", "nonchim", "mitochondria"</pre>
track <- track[keep_sample,]</pre>
keep_sample <- rownames(track)[track[,8]/track[,6] > 0.5] # remove the sample with host contamination
seqtab <- seqtab[keep_sample,keep_id]</pre>
nozeroASV <- colnames(seqtab)[colSums(seqtab) > 0]
seqtab <- seqtab[,nozeroASV]</pre>
taxa <- taxa[nozeroASV,]
```

Step 5. Serial number and output

```
source("source/utils.R")
output_path <- '~/dada2_output'
DADA2Adapter(seqtab.nochim, taxa, output_path) # DADA2Adapter function is sourced from utils.R
dir(output_path)
data <- read.csv(paste0(output_path,'/ASV_table.txt'), sep = '\t')
taxa <- read.csv(paste0(output_path,'/ASV_taxa_table.txt'), sep = '\t')</pre>
```

Step 6. Phylogenetic tree building (1)

The phylogenetic tree can be directly constructed using DECIPHER. However, fitting a GTR model is time-consuming (5hr-10hr), and installing DECIPHER can be challenging.

```
library(dada2)
library(DECIPHER)
library(phangorn)
seqs <- getSequences(seqtab.nochim)</pre>
magnitude <- ceiling(log10(ncol(seqtab.nochim)))</pre>
ASVID <- sprintf(paste0("ASV%0", magnitude, "d"), 1:n_ASVs)
names(seqs) <- ASVID</pre>
alignment <- AlignSeqs(DNAStringSet(seqs), anchor=NA, verbose=TRUE)</pre>
phangAlign <- phyDat(as(alignment, "matrix"), type="DNA")</pre>
dm <- dist.ml(phangAlign)</pre>
treeNJ <- NJ(dm)</pre>
fit = pml(treeNJ, data=phangAlign)
fitGTR <- update(fit, k=4, inv=0.2)</pre>
fitGTR <- optim.pml(fitGTR, model="GTR", optInv=TRUE, optGamma=TRUE,</pre>
                     rearrangement = "stochastic", control = pml.control(trace = 0))
fitGTR$tree
write.tree(fitGTR$tree, file = paste0(output_path,'/ASV.tree'), append = FALSE,
           digits = 10, tree.names = FALSE)
```

Step 6. Phylogenetic tree building (2)

So we provider an alternative approach. Since 16S ribosomal sequences contained the non-coding sequence and secondary structure, the alignment method was considered. We suggest to use ssu-align to do alignment for phylogenetic inference. The detailed process is as follows.

```
# ssu-align installation
$ wget eddylab.org/software/ssu-align/ssu-align-0.1.1.tar.gz;
$ tar xfz ssu-align-0.1.1.tar.gz
$ cd ssu-align-0.1.1
$ ./configure
$ make
```

```
# install the man pages and programs in system-wide directories:
$ make install

$ export PATH="$PATH:/usr/local/bin"
$ export MANPATH="$MANPATH:/usr/local/share/man"
$ export SSUALIGNDIR="/usr/local/share/ssu-align-0.1.1"

# conduct SSU-alignment
$ cd ~/Course/NDMC_2025/tmp
$ ssu-align ASV.fasta ssuout
$ cd ssuout
```

The alignment result is generated in ssuout/ssuout.bacteria.stk.

We can convert the .stk file to FASTA format using alignment_transform.R, located in the source folder.

```
$ Rscript --vanilla ~/Course/NDMC_2025/source/alignment_transform.R ~/Course/NDMC_2025/tmp/ssuout/ssuou
```

In order to shorten the processing time, we use the FastTree for maximum-likelihood phylogenetic tree completion. If time is available, RAxML is suggested to get the better quality tree.

```
$ cd ~/Course/NDMC_2025/tmp
$ ~/bin/FastTree -gtr -nt ./ssuout/ssu_align.fasta > ./ASV.tree
```

Diversity Analysis

The fecal microbiota data which from PRJEB6070 (Zeller et al., 2016) is used in this demonstration. We subsetted this dataset, including 20 patients each from the normal and cancer groups, and used the DADA2 pipeline to complete the analysis up to the taxonomic assignment step.

Please ensure that the packages listed in prep_help.R are installed.

```
library(dada2)
library(dplyr)
library(DECIPHER)
library(phangorn)
library(matrixStats)
library(ggpubr)
library(ggplot2)
library(picante)
library(phyloseq)
library(patchwork)
library(rstatix)
library(tidyverse)
library(vegan)
library(scales)
```

```
dataset_path <- "~/Course/NDMC_2025/data/Zeller_CRC.RData"
load(dataset_path)</pre>
```

```
print(ls())
## [1] "assignModul"
                             "DADA2Adapter"
                                                 "dataset_path"
                             "ggrare"
                                                 "GMMviz"
## [4] "get_composition"
## [7] "LEfSe_preparation" "make_edgetable"
                                                 "make net"
## [10] "make_nodetable"
                             "meta"
                                                 "phyloseq_to_edgeR"
## [13] "seqtab"
                             "seqtab.nochim"
                                                 "taxa"
## [16] "track"
                            "track_ab"
                                                 "track_ta"
# Check patients condition
print(table(meta$Class))
##
## Cancer Normal
       20
print(table(meta$diagnosis))
## Adenoma Cancer Normal
         6
                20
                        14
# The clinical factor for time-to-event testing
meta$event <- ifelse(meta$Class == 'Cancer', 1, 0) # as events</pre>
meta$duration <- meta$age</pre>
                                                     # as duration
```

We designed the DADA2Adapter function to bridge the data into the PreLect pipeline.

Alpha diversity

Alpha diversity describes the species diversity within a community.

Rarefaction normalization

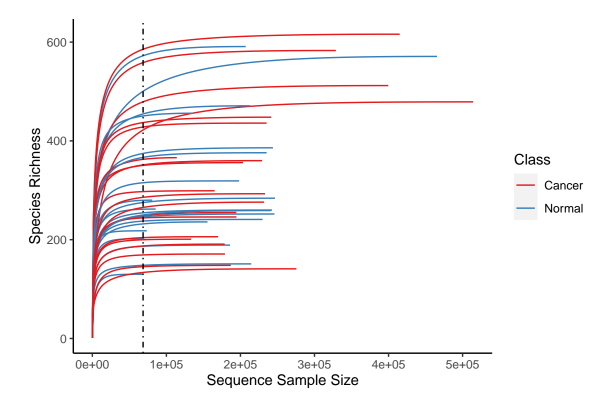
Since various range of sequencing depth in data, we usually conduct the rarefaction normalization to 'fairly' compare the diversity metrics.

```
# Rarefaction curve
ASV = otu_table(data, taxa_are_rows = TRUE)
TAX = tax_table(as.matrix(taxa))
```

```
physeq = phyloseq(ASV, TAX)
sampledata = sample_data(meta)
physeq = merge_phyloseq(physeq, sampledata)
min_depth <- min(colSums(data))
print(min_depth)</pre>
```

[1] 68528

```
# ggrare function is sourced from utils.R
ggrare(physeq, step = 1500, colour = "Class", se = FALSE) +
    scale_colour_brewer(palette="Set1") +
    geom_vline(xintercept = min_depth, linetype = 'dotdash') +
    theme(axis.line = element_line(linetype = 1,colour = 'black'),
    panel.background = element_rect(I(0)),
    panel.grid.major = element_line(colour = NA),
    panel.grid.minor = element_line(colour = NA))
```



The upper picture show the library size vs. number of observed species, in rarefaction process, we limited the all samples in minimum depth (black dotdash), and then randomly discarding reads from larger samples until the number of remaining samples is equal to this threshold

```
# Rarefaction
print(colSums(data)[1:16])

## ERR475473 ERR475476 ERR475478 ERR475480 ERR475483 ERR475484 ERR475485 ERR475493

## 73475 155494 207049 68528 70074 85513 80933 203534

## ERR475500 ERR475504 ERR475513 ERR475518 ERR475521 ERR475527 ERR475528 ERR475529

## 514255 465384 415085 242616 399663 328836 136463 133480
```

```
min <- min(colSums(data))</pre>
data_rarefied <- t(rrarefy(t(data), min))</pre>
## Warning in rrarefy(t(data), min): function should be used for observed counts,
## but smallest count is 2
data_rarefied <- data_rarefied[rowSums(data_rarefied) > 0,]
print(colSums(data_rarefied)[1:16])
## ERR475473 ERR475476 ERR475478 ERR475480 ERR475483 ERR475484 ERR475485 ERR475493
       68528
                 68528
                            68528
                                      68528
                                                 68528
                                                           68528
                                                                      68528
## ERR475500 ERR475504 ERR475513 ERR475518 ERR475521 ERR475527 ERR475528 ERR475529
       68528
                 68528
                            68528
                                      68528
                                                 68528
                                                           68528
                                                                      68528
```

After rarefaction, All the sample size are equality, we can calculate the various alpha index to each sample.

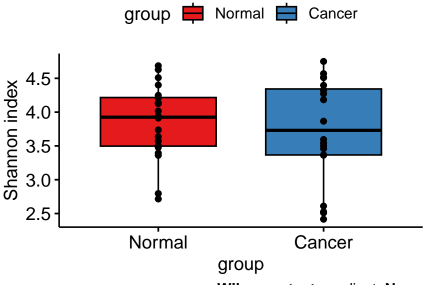
```
dir(output_path)
                            "ASV_taxa_table.txt" "ASV.fasta"
## [1] "ASV_table.txt"
                            "for_PICRUSt2.tsv"
## [4] "ASV.tree"
                                                 "LEfSe_tmp"
## [7] "PICRUSt2"
                            "PreLect_tmp"
                                                 "ssuout"
# load the data we need
tree <- read.tree(paste0(output_path, '/ASV.tree'))</pre>
                          #check the root by farthest phylogenetic distance ASV
a <- cophenetic(tree)
rowSums(a)[order(rowSums(a), decreasing = T)][1:8]
## ASV2154 ASV1810 ASV0186 ASV3210 ASV2490 ASV2300 ASV1408 ASV1814
## 3417.648 3349.283 3266.763 3223.351 3205.014 3199.102 3170.137 3145.933
tree <- root(tree, "ASV2154", resolve.root = T)</pre>
AlphaIndex <- data.frame(Shannon = diversity(t(data_rarefied) ,index = "shannon"),
                         Chao1 = estimateR(t(data_rarefied))[2,],
                         Simpson = diversity(t(data_rarefied) ,index = "simpson"),
                         invSimpson = diversity(t(data_rarefied) ,index = "invsimpson"),
                         PD = pd(t(data_rarefied), tree)[,1],
                         group = meta$Class)
```

Shannon diversity

$$H_{sw} = -\sum_{i=1}^s (\frac{n_i}{N}) \ln(\frac{n_i}{N})$$

Shannon and Wiener (1963) is base on information theory, N is total number of individual, and n_i is the number of individual belong in species i. The more the number of species and the more evenly distributed the individuals are, the higher the index it get. Therefore, H_{sw} can be regarded as equitability.

```
pwc <- wilcox_test(Shannon ~ group, paired = F, p.adjust.method = "None", data = AlphaIndex)
pwc <- pwc %>% add_xy_position(x = "group")
ggboxplot(AlphaIndex, x = "group", y = "Shannon", add = "point", fill = "group") +
    scale_fill_brewer(palette = "Set1") + ylab("Shannon index") +
    stat_pvalue_manual(pwc, hide.ns = TRUE) +
    labs(caption = get_pwc_label(pwc))
```

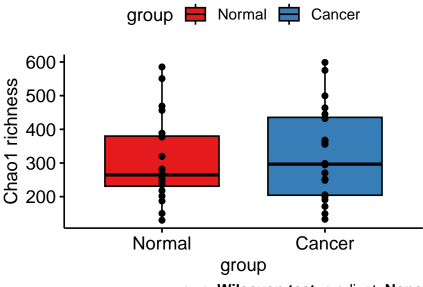


Chao1 richness

$$S = S_{obs} + \frac{F_1^2}{2F_2}$$

In Chao1 estimator (Chao, A. 1984). S_{obs} is indicated the number of observed species, F_1 is the number of species which only once. and F_2 is the number of species which twice in community. Higher F_1 indicates that the number of non-observed species is likely to be higher. It is expected that the community will have a relatively high abundance of species. but F_2 show the species have occurred at least twice, then the chance of new species occurring in the community is low.

```
pwc <- wilcox_test(Chao1 ~ group, paired = F, p.adjust.method = "None", data = AlphaIndex)
pwc <- pwc %>% add_xy_position(x = "group")
ggboxplot(AlphaIndex, x = "group", y = "Chao1", add = "point", fill = "group") +
    scale_fill_brewer(palette = "Set1") + ylab("Chao1 richness") +
    stat_pvalue_manual(pwc, hide.ns = TRUE) +
    labs(caption = get_pwc_label(pwc))
```

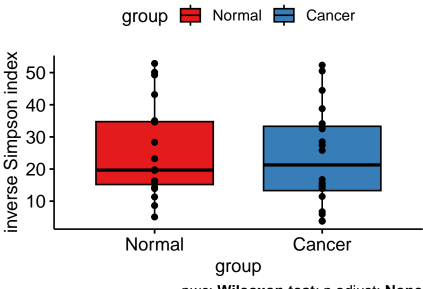


Simpson index

$$D_s = \sum_{i=1}^s (\frac{n_i}{N})^2 \ ; \quad D_{s'} = \frac{1}{D_s}$$

Simpson (1949) is measure the dominance in single community. if some species is dominant in community, the Simpson will be higher. so it can be regarded as concentration index. In other words, the inverse Simpson index is show the evenness in community.

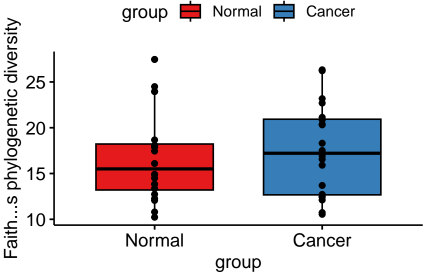
```
pwc <- wilcox_test(Simpson ~ group, paired = F, p.adjust.method = "None", data = AlphaIndex)
pwc <- pwc %>% add_xy_position(x = "group")
ggboxplot(AlphaIndex, x = "group", y = "invSimpson", add = "point", fill = "group") +
    scale_fill_brewer(palette = "Set1") + ylab("inverse Simpson index") +
    stat_pvalue_manual(pwc, hide.ns = TRUE) +
    labs(caption = get_pwc_label(pwc))
```



Phylogenetic diversity

Faith's Phylogenetic Diversity (Faith D., 1992) which is defined as the sum of the branch lengths of a phylogenetic tree connecting all species, this means that PD indicates Feature diversity.

```
pwc <- wilcox_test(PD ~ group, paired = F, p.adjust.method = "None", data = AlphaIndex)
pwc <- pwc %>% add_xy_position(x = "group")
ggboxplot(AlphaIndex, x = "group", y = "PD", add = "point", fill = "group") +
    scale_fill_brewer(palette = "Set1") + ylab("Faith's phylogenetic diversity") +
    stat_pvalue_manual(pwc, hide.ns = TRUE) +
    labs(caption = get_pwc_label(pwc))
```



Beta diversity

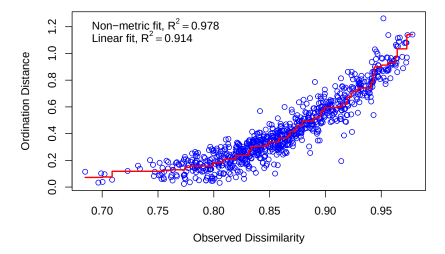
Bray-Curtis distance

$$D_{BC} = \frac{\sum_{i=1}^{S} |M_{i1} - M_{i2}|}{\sum_{i=1}^{S} M_{i1} + M_{i2}}$$

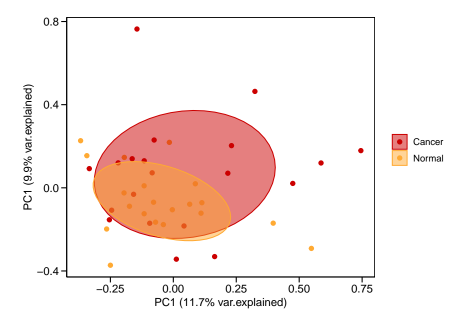
In Bray-Curtis distance, S is indicates the total number of species in two communities, M_{i1} is means the number of species i in community 1, and so on. This method is similar to Sørensen index. and usually utilizes non-metric multidimensional scaling nMDS for dimension reduction.

```
NMDS=metaMDS(t(data_rarefied), distance = "bray")
## Square root transformation
## Wisconsin double standardization
## Run 0 stress 0.149106
## Run 1 stress 0.1503017
## Run 2 stress 0.1491056
## ... New best solution
## ... Procrustes: rmse 0.0005981917 max resid 0.002768234
## ... Similar to previous best
## Run 3 stress 0.1491057
## ... Procrustes: rmse 1.548925e-05 max resid 7.090235e-05
## ... Similar to previous best
## Run 4 stress 0.1503017
## Run 5 stress 0.1491056
## ... New best solution
## ... Procrustes: rmse 4.216836e-05 max resid 0.0001434376
## ... Similar to previous best
## Run 6 stress 0.1491057
## ... Procrustes: rmse 0.0001441396 max resid 0.0005599521
## ... Similar to previous best
```

```
## Run 7 stress 0.1912499
## Run 8 stress 0.1491055
## ... New best solution
## ... Procrustes: rmse 0.0002371133 max resid 0.001049122
## ... Similar to previous best
## Run 9 stress 0.1503017
## Run 10 stress 0.1503017
## Run 11 stress 0.1503084
## Run 12 stress 0.1491058
## ... Procrustes: rmse 0.0001865112 max resid 0.0008890889
## ... Similar to previous best
## Run 13 stress 0.1503084
## Run 14 stress 0.1491059
## ... Procrustes: rmse 0.0002518848 max resid 0.001206103
## ... Similar to previous best
## Run 15 stress 0.1503017
## Run 16 stress 0.1491055
## ... Procrustes: rmse 0.0001621694 max resid 0.0007371812
## ... Similar to previous best
## Run 17 stress 0.2091891
## Run 18 stress 0.1503017
## Run 19 stress 0.1491058
## ... Procrustes: rmse 0.0002246621 max resid 0.00102273
## ... Similar to previous best
## Run 20 stress 0.1503017
## *** Best solution repeated 5 times
NMDSplot <- as.data.frame(NMDS$points)</pre>
NMDSplot$group <- meta$Class</pre>
prop <- cmdscale(vegdist(t(data_rarefied), method = "bray"), k = 2, eig = T, add = T )</pre>
prop <- round(prop$eig*100/sum(prop$eig),1)</pre>
print(prop[1:8]) # chick the proportion of variance explained
## [1] 11.7 9.9 7.6 5.7 5.4 4.5 3.8 3.7
stressplot(NMDS) # chick the fitness in nMDS
```



```
ggscatter(NMDSplot, x = "MDS1", y = "MDS2",combine = T, color = 'group',
    ellipse.type = "norm", ellipse = T,ellipse.level = 0.5, ellipse.alpha = 0.5, repel = TRUE) +
    scale_color_manual(values = c("#CC0000","#FFAA33"))+
    scale_fill_manual(values = c("#CC0000","#FFAA33")) +
    xlab(paste0(c('PC1 (', prop[1],'% var.explained)'), collapse = "")) +
    ylab(paste0(c('PC1 (', prop[2],'% var.explained)'), collapse = "")) +
    theme(panel.background = element_rect(fill = 'transparent'),
        panel.grid = element_blank(),
        axis.ticks.length = unit(0.4,"lines"),
        axis.ticks = element_line(color='black'),
        axis.line = element_line(color = "black"),
        legend.title=element_blank(),
        legend.position = 'right')
```



Ellipse type can choose the convex, confidence, t, euclid.

Unifrac distance

• unWeighted

$$U_{uw} = \frac{\sum_{i=1}^{N} l_i |A_i - B_i|}{\sum_{i=1}^{N} max(A_i + B_i)}$$

• Weighted

$$U_{w} = \frac{\sum_{i=1}^{n} b_{i} |\frac{A_{i}}{A_{T}} - \frac{B_{i}}{B_{T}}|}{\sum_{j=1}^{S} L_{j}}$$

```
ASV = otu_table(data_rarefied, taxa_are_rows = TRUE)

TAX = tax_table(as.matrix(taxa))

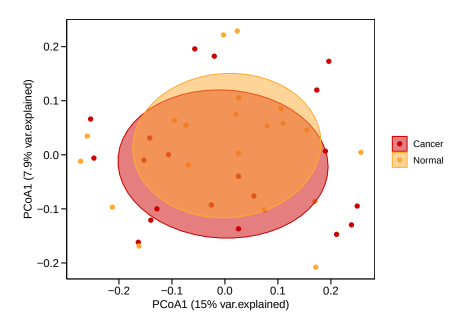
physeq = phyloseq(ASV, TAX, tree)

Unif = UniFrac(physeq, weighted = F, normalized = F, parallel = F)  # if weighted = TRUE; then weighted

Unif_d <- pcoa(Unif)

Unifplot <- data.frame(axis1 = as.numeric(Unif_d$vectors[,1]),
```

```
axis2 = as.numeric(Unif_d$vectors[,2]))
Unifplot$group <- meta$Class</pre>
prop <- cmdscale(Unif, k = 2, eig = T, add = T)</pre>
prop <- round(prop$eig*100/sum(prop$eig),1)</pre>
print(prop[1:8]) # chick the proportion of variance explained
## [1] 15.0 7.9 6.3 4.0 3.7 3.4 3.3 3.2
ggscatter(Unifplot, x = "axis1", y = "axis2",combine = T, color = 'group',
          ellipse.type = "norm", ellipse = T,ellipse.level = 0.5, ellipse.alpha = 0.5, repel = TRUE) +
          scale_color_manual(values = c("#CC0000","#FFAA33"))+
          scale_fill_manual(values = c("#CC0000","#FFAA33")) +
          xlab(paste0(c('PCoA1 (', prop[1],'% var.explained)'), collapse = "")) +
          ylab(pasteO(c('PCoA1 (', prop[2],'% var.explained)'), collapse = "")) +
          theme(panel.background = element_rect(fill = 'transparent'),
                panel.grid = element_blank(),
                axis.ticks.length = unit(0.4, "lines"),
                axis.ticks = element_line(color='black'),
                axis.line = element_line(colour = "black"),
                legend.title=element_blank(),
                legend.position = 'right')
```



ANOSIM

Analysis of similarities (ANOSIM) is a non-parametric statistical test widely used in the field of ecology. As an ANOVA-like test, where instead of operating on raw data, operates on a ranked dissimilarity matrix.

Given a matrix of rank dissimilarities between a set of samples, each solely belong to one treatment group, the ANOSIM tests whether we can reject the null hypothesis that the similarity between groups is greater than or equal to the similarity within the groups.

The test statistic R is calculated in the following way:

$$R = \frac{\bar{r_B} - \bar{r_W}}{M/2}$$

where $\bar{r_B}$ is the average of rank similarities of pairs of samples (or replicates) originating from different sites, $\bar{r_W}$ is the average of rank similarity of pairs among replicates within sites, and M = n(n-1)/2 where n is the number of samples.

```
anosim(vegdist(t(data_rarefied), method = "bray"), meta$Class)
## Call:
## anosim(x = vegdist(t(data_rarefied), method = "bray"), grouping = meta$Class)
## Dissimilarity: bray
## ANOSIM statistic R: 0.08716
##
         Significance: 0.004
##
## Permutation: free
## Number of permutations: 999
anosim(Unif, meta$Class)
##
## Call:
## anosim(x = Unif, grouping = meta$Class)
## Dissimilarity:
## ANOSIM statistic R: 0.01125
##
         Significance: 0.274
## Permutation: free
## Number of permutations: 999
```

adonis2

Permutational Multivariate Analysis of Variance (adonis),

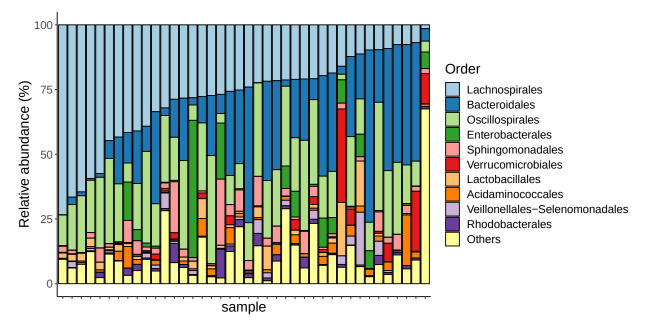
```
adonis2(t(data_rarefied) ~ Class, data = meta, method= "bray")
## Permutation test for adonis under reduced model
## Terms added sequentially (first to last)
## Permutation: free
## Number of permutations: 999
## adonis2(formula = t(data_rarefied) ~ Class, data = meta, method = "bray")
          Df SumOfSqs
                          R2
                               F Pr(>F)
           1 0.5381 0.04186 1.66 0.015 *
## Class
## Residual 38 12.3178 0.95814
## Total
         39 12.8559 1.00000
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
adonis2(Unif ~ Class, data = meta)
## Permutation test for adonis under reduced model
## Terms added sequentially (first to last)
## Permutation: free
## Number of permutations: 999
## adonis2(formula = Unif ~ Class, data = meta)
                         R2 F Pr(>F)
          Df SumOfSqs
## Class 1 0.1782 0.02849 1.1142 0.244
```

```
## Residual 38 6.0787 0.97151
## Total 39 6.2569 1.00000
```

Community composition

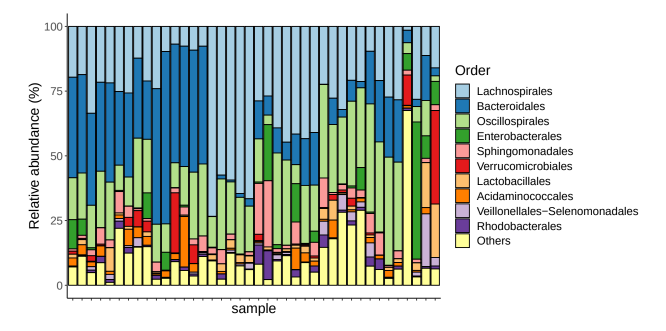
sort by major species decrease

```
# choose the taxonomic rank for visualization
print(colnames(taxa))
## [1] "Kingdom" "Phylum" "Class"
                                     "Order"
                                              "Family"
                                                                    "Species"
# qet_composition function is sourced from utils.R
compair <- get_composition(data, taxa, 'Order', meta, 'Class')</pre>
ggplot(compair, aes(x = sample, y = percentage, fill = taxa)) +
  geom_bar(stat="identity",colour = "black") + scale_fill_brewer(palette = "Paired") +
  labs(fill = "Order") + ylab("Relative abundance (%)") +
  #facet_grid(~group, space="free", scales="free") + # separate the group
  theme(axis.text.x = element_blank(),
        axis.line = element_line(linetype = 1,colour = 'black'),
        panel.background = element rect(I(0)),
        panel.grid.major = element_line(colour = NA),
        panel.grid.minor = element_line(colour = NA),
        text = element_text(size=16))
```



sort by hierarchical clustering

```
compair_2D <- reshape(compair[,1:3], idvar = "sample", timevar = "taxa", direction = "wide")
rownames(compair_2D) <- compair_2D[,1]
compair_2D <- compair_2D[,2:ncol(compair_2D)]
horder <- hclust(dist(compair_2D), method = 'ward.D')
horder <- horder$labels[horder$order]
compair$sample <- factor(compair$sample, levels = horder)</pre>
```



Environmental Analysis

To demonstrate how the environmental factor impacting in microbiota, we use wetland microbiota which characterised by ions concentration. the soil samples were collected from 4 wetland in southern Taiwan and named as AA, BB, CC, DD respectively. and each sample had be measured with LCMS.

```
wetland <- read.table("~/Course/NDMC_2025/data/wetland/ASV_table.txt", sep = "\t", stringsAsFactors = F
ion <- read.table("~/Course/NDMC_2025/data/wetland/ion.txt", sep = "\t", stringsAsFactors = F)
wetland <- wetland[,rownames(ion)]</pre>
head(wetland[,1:5])
                  AA1 AA2 AA3 AA4
                                    AA5
## ASV V3V400009
                    0 196
                             0
                               13
## ASV V3V400024
                     0
                             0
                         0
## ASV_V3V400036 2846
                        26 552
                                 0 2030
## ASV_V3V400040
                    0
                         0
                             0
                                 0
## ASV_V3V400042 821 793 947 633
                                    427
## ASV V3V400043
                                      0
head(ion)
                            Mg
                                    Ca
                                                         S04
                                                                  EC
                                                                           Salinity
```

```
## AA1
        538.0 45.76
                       93.40
                              78.33
                                       775.3
                                              0.0 321.9
                                                           3600 7.143 1.610759
                                              0.0 164.2
## AA2
        691.1 47.13
                       87.07
                               49.96
                                       703.3
                                                           4020 7.675 1.799610
## AA3
        716.1 56.79 124.20 151.00
                                       920.1
                                              0.0 418.2
                                                           4330 6.862 2.040961
        651.2 80.48
## AA4
                       97.89
                              89.33
                                       727.9
                                              0.0 244.4
                                                           3640 6.330 1.755768
## AA5
        576.9
                0.00
                       85.44
                               83.32
                                      748.3
                                              0.0 326.7
                                                           3510 6.670 1.650724
## BB1 23830.0 922.90 2989.00 1494.00 37540.0 266.6 5298.0 137700 6.701 74.292982
```

Mantel test

The Mantel test computes the Pearson or Spearman correlation between two distance matrices and evaluates its significance using a permutation test.

```
min <- min(colSums(wetland))</pre>
wetland_rarefied <- t(rrarefy(t(wetland), min))</pre>
wetland_rarefied <- wetland_rarefied[rowSums(wetland_rarefied) > 0,]
manteldf <- data.frame(Factor = 0, rho = 0, p_value = 0)
count <- 1
for(i in 1:ncol(ion)){
 Mantel_res <- mantel(dist(ion[,i]),vegdist(t(wetland_rarefied)), method = "spearman")</pre>
  manteldf[count,1] <- colnames(ion)[i]</pre>
 manteldf[count,2] <- round(Mantel_res$statistic,4)</pre>
  manteldf[count,3] <- round(Mantel_res$signif,4)</pre>
  count = count + 1
}
manteldf
##
        Factor
                   rho p_value
## 1
            Na 0.7629 0.001
## 2
            K 0.1049
                         0.121
## 3
            Mg 0.6716
                         0.001
## 4
            Ca 0.7391
                         0.001
## 5
            Cl 0.7424
                         0.001
            Br 0.1665
## 6
                         0.033
## 7
           SO4 0.6853
                         0.001
## 8
            EC 0.7235
                         0.001
## 9
            pH 0.2262
                         0.020
## 10 Salinity 0.7553
                         0.001
```

BioENV

This finds the best subset of environmental variables, so that the Euclidean distances of scaled environmental variables have the maximum (rank) correlation with community dissimilarities.

```
bio <- bioenv(as.formula(paste("t(wetland_rarefied)~ ", paste(colnames(ion), collapse = " + "))), ion)
## 1023 possible subsets (this may take time...)
bio <- summary(bio)
df <- data.frame(Factor = bio$variables, rank = bio$size, rho = bio$correlation)
df
## Factor rank rho
## 1 Na 1 0.7629216
## 2 Ca Salinity 2 0.7699462</pre>
```

```
## 3
                          Na Ca Salinity
                                           3 0.7783320
## 4
                       Na Ca EC Salinity
                                           4 0.7773259
## 5
                    Na Ca Cl EC Salinity
                                           5 0.7741496
## 6
                 Na Ca Cl EC pH Salinity
                                           6 0.7634334
## 7
              Na Mg Ca Cl EC pH Salinity
                                           7 0.7617276
## 8
          Na Mg Ca Cl SO4 EC pH Salinity
                                           8 0.7566520
## 9
       Na Mg Ca Cl Br SO4 EC pH Salinity
                                           9 0.7420710
## 10 Na K Mg Ca Cl Br SO4 EC pH Salinity 10 0.6766375
```

CCA

Canonical Correspondence Analysis (CCA) is an ordination technique that directly relates species composition to environmental variables. CCA is based on Chi-square distances, which measure differences in species composition between samples, and uses a constrained ordination approach to maximize the correlation between species distributions and environmental gradients.

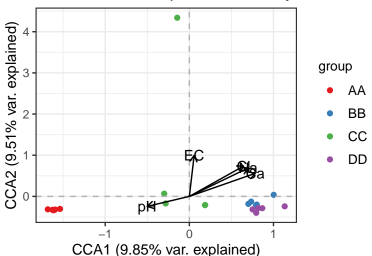
Given a species abundance matrix and an environmental variable matrix, CCA seeks to find axes that maximize the weighted correlation between species composition and environmental variables. The species scores are derived based on weighted averaging, ensuring that species are positioned optimally along the environmental gradients. This makes CCA particularly suitable for ecological datasets where species exhibit unimodal responses to environmental gradients.

```
\#CCA \leftarrow cca(as.formula(paste("t(wetland_rarefied)~", paste(colnames(ion), collapse = " + "))), ion)
CCA <- cca(t(wetland rarefied)~ Na + Ca + Cl+ EC + Salinity + pH, ion)
#cca_temp <- anova.cca(CCA)</pre>
cca_term <- anova.cca(CCA, by="terms")</pre>
cca term
## Permutation test for cca under reduced model
## Terms added sequentially (first to last)
## Permutation: free
## Number of permutations: 999
## Model: cca(formula = t(wetland_rarefied) ~ Na + Ca + Cl + EC + Salinity + pH, data = ion)
##
            Df ChiSquare
                              F Pr(>F)
## Na
            1
                 0.8836 2.0777 0.001 ***
## Ca
             1
                  0.5973 1.4045 0.029 *
## Cl
             1
                  0.5803 1.3645
                                 0.022 *
## EC
                  0.6755 1.5884 0.002 **
             1
## pH
             1
                  0.5788 1.3611 0.020 *
## Residual 14
                  5.9536
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
CCAscores <- scores(CCA, display = "sites") %>%
               as.data.frame() %>% rownames_to_column("site")
CCAscores$group <- substr(rownames(ion),1,2)</pre>
cc <- data.frame(CCA$CCA$biplot)</pre>
ggplot() + geom_point(data = CCAscores, aes(x = CCA1, y = CCA2, color=group), alpha=1) +
    geom_vline(xintercept = c(0), color = "grey70", linetype = 2) +
    geom_hline(yintercept = c(0), color = "grey70", linetype = 2) +
```

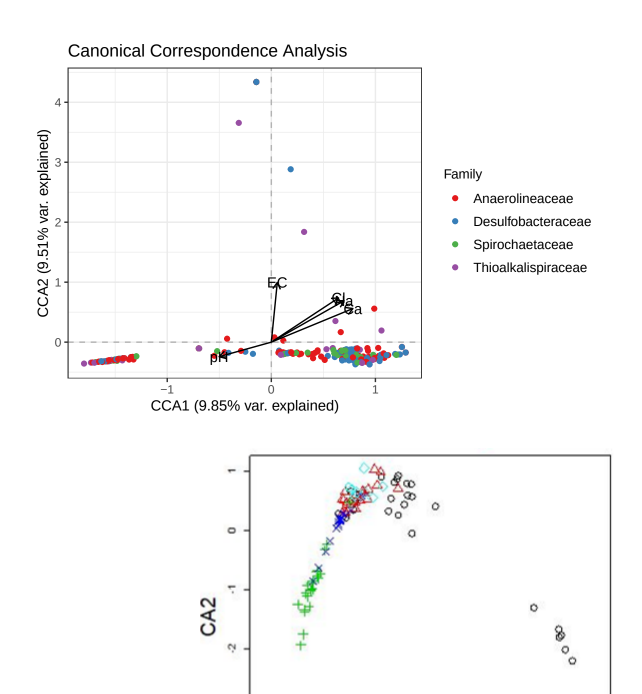
geom_segment(data = cc, aes(x = 0, y = 0, xend = CCA1, yend = CCA2), arrow = arrow(length = unit(0.

```
scale_color_brewer(palette = 'Set1') +
geom_text(data = cc, aes(x = CCA1, y = CCA2, label = rownames(cc))) + theme_bw() +
labs(x = paste0("CCA1 (",round(CCA$CCA$eig[1] / CCA$tot.chi*100,2), "% var. explained)"),
    y = paste0("CCA2 (",round(CCA$CCA$eig[2] / CCA$tot.chi*100,2), "% var. explained)"),
    title = "Canonical Correspondence Analysis") +
theme(legend.text.align = 0, legend.title.align = 0,
    legend.title=element_text(size=10),legend.text=element_text(size=10))
```

Canonical Correspondence Analysis



```
wetland_taxa <- read.table("~/Course/NDMC_2025/data/wetland/ASV_taxa.csv", sep = ",", stringsAsFactors</pre>
finalmodel<- ordistep(CCA, scope=formula(CCA))</pre>
## Start: t(wetland_rarefied) ~ Na + Ca + Cl + EC + Salinity + pH
CCAscores <- data.frame(scores(finalmodel, display = "sp"))</pre>
CCAscores$sample <- rownames(wetland_rarefied)</pre>
CCAscores$Family <- wetland_taxa$Family</pre>
CCAscores <- CCAscores [CCAscores Family %in% c('Desulfobacteraceae', 'Anaerolineaceae', 'Spirochaetaceae'
cc <- data.frame(finalmodel$CCA$biplot)</pre>
ggplot() +
  geom_point(data = CCAscores, aes(x = CCA1, y = CCA2, color = Family)) +
  geom vline(xintercept = c(0), color = "grey70", linetype = 2) +
  geom_hline(yintercept = c(0), color = "grey70", linetype = 2) +
  geom_segment(data = cc, aes(x = 0, y = 0, xend = CCA1, yend = CCA2), arrow = arrow(length = unit(0.2,
  geom_text(data = cc, aes(x = CCA1, y = CCA2, label = rownames(cc))) +
  theme_bw() + scale_color_brewer(palette = 'Set1') +
  labs(x = paste0("CCA1 (",round(CCA$CCA$eig[1] / CCA$tot.chi*100,2), "% var. explained)"),
       y = paste0("CCA2 (",round(CCA$CCA$eig[2] / CCA$tot.chi*100,2), "% var. explained)"),
       title = "Canonical Correspondence Analysis") +
  theme(legend.text.align = 0, legend.title.align = 0,
        legend.title=element_text(size=10),legend.text=element_text(size=10))
```



The arch effect in CCA

A common artifact in CCA is the arch effect, a curvature in the ordination space that arises when species distributions follow strong unimodal patterns. This occurs because Chi-square distances emphasize differences in species presence and absence, leading to distortions in the ordination space. As a result, sites along

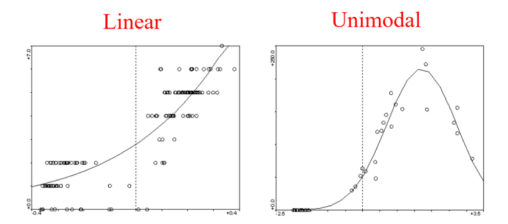
0

CA1

2

3

a continuous environmental gradient may appear curved instead of following a linear progression.



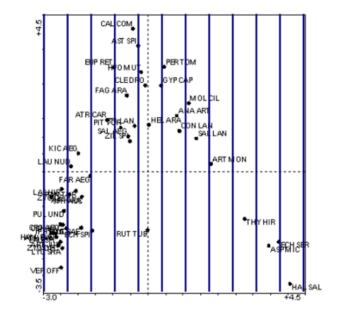
x-axis is environmental gradient and y-axis is abundance of species.

reference :

- David Zelený Lab at NTU
- slideplayer

DCA

Detrending by Segments



To correct the arch effect, Detrended Correspondence Analysis (DCA) was introduced. DCA is a modification of Correspondence Analysis (CA) that removes the arch effect through segment-wise detrending and local standardization.

Segment-wise Detrending DCA first computes an initial ordination using Correspondence Analysis (CA). However, instead of allowing the first axis to curve, DCA splits the axis into equal segments and removes the local mean within each segment. This eliminates the arch effect and ensures that the ordination axis follows a straight gradient.

Local Standardization In addition to detrending, DCA adjusts species scores to equalize species variance across the axis. This prevents overemphasis on dominant species and ensures that species with different dispersions are treated equally.

DCA First Axis and Gradient Length The first DCA axis (DCA1) represents the primary gradient in species composition. The length of the axis, measured in standard deviation (SD) units, indicates the degree of species turnover:

• If first axis length < 2 SD:

species distributions follow a linear pattern, making Redundancy Analysis (RDA) more appropriate.

• If first axis length > 4 SD:

species distributions follow a unimodal response, meaning CCA is the better choice.

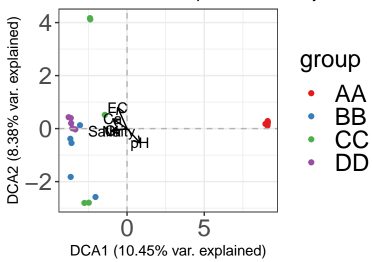
```
DCA <- decorana(t(wetland_rarefied))</pre>
DCA
##
## Call:
## decorana(veg = t(wetland rarefied))
## Detrended correspondence analysis with 26 segments.
## Rescaling of axes with 4 iterations.
## Total inertia (scaled Chi-square): 9.2689
##
##
                            DCA1
                                   DCA2
                                          DCA3
                                                  DCA4
## Eigenvalues
                          0.9682 0.7771 0.6458 0.4624
                         0.9682 0.7762 0.6419 0.4392
## Additive Eigenvalues
## Decorana values
                          0.9693 0.8123 0.6090 0.3957
                         12.8999 6.9736 5.7240 3.5045
## Axis lengths
```

```
#dca_factor <- envfit(as.formula(paste("DCA~ ", paste(colnames(ion), collapse = " + "))), data = ion)
dca_factor <- envfit(DCA~ Na + Ca + Cl+ EC + Salinity + pH, data = ion)
dca_factor
##
## ***VECTORS
##
##
                DCA1
                         DCA2
                                  r2 Pr(>r)
## Na
            -0.99542 -0.09556 0.4575 0.008 **
            -0.93041 0.36653 0.5559 0.002 **
## Ca
## Cl
            -0.99632 -0.08571 0.3943
                                      0.016 *
## EC
            -0.60555 0.79581 0.0241
                                     0.773
## Salinity -0.99586 -0.09094 0.4272 0.011 *
            0.84129 -0.54058 0.3720 0.017 *
## pH
## ---
```

```
## Permutation: free
## Number of permutations: 999
dcv <- data.frame(dca factor$vectors$arrows)</pre>
DCA_sum <- as.data.frame(summary(DCA)$site.scores)</pre>
##
## Call:
## decorana(veg = t(wetland_rarefied))
## Detrended correspondence analysis with 26 segments.
## Rescaling of axes with 4 iterations.
## Total inertia (scaled Chi-square): 9.2689
##
                           DCA1
                                  DCA2 DCA3 DCA4
## Eigenvalues
                         0.9682 0.7771 0.6458 0.4624
## Additive Eigenvalues 0.9682 0.7762 0.6419 0.4392
## Decorana values
                         0.9693 0.8123 0.6090 0.3957
## Axis lengths
                        12.8999 6.9736 5.7240 3.5045
DCAscores <- DCA_sum %>% rownames_to_column("site")
DCAscores$group <- substr(rownames(ion),1,2)</pre>
ggplot() + geom_point(data = DCAscores, aes(x = DCA1, y = DCA2, color = group), alpha=1) +
    geom_vline(xintercept = c(0), color = "grey70", linetype = 2) +
    geom_hline(yintercept = c(0), color = "grey70", linetype = 2) +
   geom_segment(data = dcv, aes(x = 0, y = 0, xend = DCA1, yend = DCA2), arrow = arrow(length = unit(0
   geom_text(data = dcv, aes(x = DCA1, y = DCA2, label = rownames(dcv))) + theme_bw() +
   scale_color_brewer(palette = 'Set1') +
   labs(x = paste0("DCA1 (",round(DCA$evals[1] / DCA$totchi*100,2), "% var. explained)"),
         y = paste0("DCA2 (",round(DCA$evals[2] / DCA$totchi*100,2), "% var. explained)"),
         title = "Detrended Correspondence Analysis")+
    theme(legend.text.align = 0, legend.title.align = 0,
      legend.title=element_text(size=18),
      legend.text=element_text(size=18),
      axis.text = element_text(size=18))
```

Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1

Detrended Correspondence Analysis



RDA

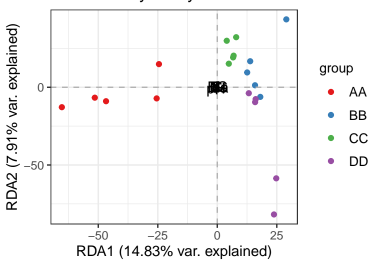
Redundancy Analysis (RDA) is a linear constrained ordination method similar to Principal Component Analysis (PCA), but with environmental constraints. Instead of using Chi-square distances, RDA applies Euclidean distances, making it suitable when species responses are approximately linear with respect to environmental gradients.

Given a species data matrix and an environmental matrix , RDA performs a multivariate regression of on , followed by PCA on the fitted values. This results in ordination axes that explain the maximum variation in species data constrained by the environmental variables.

```
\#RDA \leftarrow rda(as.formula(paste("t(wetland_rarefied) \sim ", paste(colnames(ion), collapse = " + "))), ion)
RDA <- rda(t(wetland_rarefied)~ Na + Ca + Cl+ EC + Salinity + pH, ion)
rda_temp <- anova.cca(RDA)</pre>
rda_term <- anova.cca(RDA, by="terms")
rda_term
## Permutation test for rda under reduced model
## Terms added sequentially (first to last)
## Permutation: free
## Number of permutations: 999
##
## Model: rda(formula = t(wetland_rarefied) ~ Na + Ca + Cl + EC + Salinity + pH, data = ion)
##
            Df Variance
                              F Pr(>F)
## Na
             1
                  783986 2.3209 0.001 ***
## Ca
                 541133 1.6020
                                 0.051
             1
                 392550 1.1621
## Cl
             1
                                 0.269
## EC
                 362663 1.0736
                                 0.371
             1
                 457295 1.3538 0.134
## pH
             1
## Residual 14 4729101
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
RDA_sum <- summary(RDA)</pre>
```

RDAscores <- as.data.frame(RDA sum\$sites[,1:2]) %>% rownames to column("site")

Redundancy Analysis



Differential Taxa

ALDEx2

ALDEx2 which generated Monte Carlo samples of Dirichlet distributions for each sample, using a uniform prior, performed CLR transformation of each realization, and then performed Wilcoxon tests on the transformed realizations.

```
library(ALDEx2)
## Loading required package: zCompositions
## Loading required package: MASS
##
## Attaching package: 'MASS'
## The following object is masked from 'package:rstatix':
##
## select
## The following object is masked from 'package:patchwork':
##
```

```
## The following object is masked from 'package:dplyr':
##
       select
## Loading required package: NADA
## Loading required package: survival
##
## Attaching package: 'NADA'
## The following object is masked from 'package: IRanges':
##
       cor
## The following object is masked from 'package:S4Vectors':
##
## The following object is masked from 'package:stats':
##
       cor
## Loading required package: truncnorm
ALDEx2_result <- aldex(reads=data, conditions = meta$Class,</pre>
                      mc.samples = 128, test="t", effect=TRUE,
                       include.sample.summary = FALSE, verbose=T, denom="all")
## aldex.clr: generating Monte-Carlo instances and clr values
## operating in serial mode
## removed rows with sums equal to zero
## computing center with all features
## data format is OK
## dirichlet samples complete
## transformation complete
## aldex.ttest: doing t-test
## running tests for each MC instance:
## |-----(25%)-----(50%)-----(75%)------|
## aldex.effect: calculating effect sizes
## operating in serial mode
## sanity check complete
## rab.all complete
## rab.win complete
## rab of samples complete
## within sample difference calculated
## between group difference calculated
## group summaries calculated
## effect size calculated
## summarizing output
selected result <- ALDEx2 result[ALDEx2 result$wi.ep < 0.05,]</pre>
selected_result <- cbind(rownames(selected_result),selected_result)</pre>
head(selected_result)
          rownames(selected_result) rab.all rab.win.Cancer rab.win.Normal
## ASV0001
                            ASV0001 14.373547 14.630241 13.8742929
                            ASV0007 9.648390
ASV0119 8.466592
                                                 12.282246
## ASV0007
                                                                  7.7414882
## ASV0119
                                                  3.945452 10.0351473
                            ASV0130 3.991660
## ASV0130
                                                   1.643844
                                                                 9.4433972
## ASV0146
                            ASV0146 8.222159
                                                  9.565661
                                                                  7.4461137
## ASV0178
                                                  5.716942
                            ASV0178 1.850530
                                                                   0.4256441
```

```
diff.btw diff.win
                                  effect
                                           overlap
                                                         we.ep
## ASV0001 -1.377995 2.364327 -0.5127501 0.3161593 0.009924686 0.9111672
## ASV0007 -4.198266 5.508847 -0.6886866 0.2265625 0.001738385 0.8659766
## ASV0119 4.845204 7.793662 0.6220667 0.2638564 0.008007031 0.8930373
## ASV0130 4.215072 9.212734 0.4781253 0.3099141 0.056589218 0.9238886
## ASV0146 -1.937207 5.932652 -0.3029744 0.3093750 0.149241061 0.9396661
## ASV0178 -5.289013 6.269317 -0.7701952 0.2107729 0.003434822 0.7223806
##
                 wi.ep
                          wi.eBH
## ASV0001 0.041240625 0.9270136
## ASV0007 0.003136204 0.8139919
## ASV0119 0.009039480 0.8642710
## ASV0130 0.049115638 0.9198037
## ASV0146 0.046041349 0.9261615
## ASV0178 0.003249730 0.6543478
#write.table(selected_result, "~/Course/NDMC_2025/tmp/ALDEx2.txt", quote=FALSE, sep="\t", col.names = F
```

ANCOM

ANCOM first examined the abundance table to identify outlier zeros and structural zeros, Outlier zeros, identified by finding outliers in the distribution of taxon counts within each sample grouping, were ignored during differential abundance analysis, and replaced with NA. Structural zeros, taxa that were absent in one grouping but present in the other, were ignored during data analysis and automatically called as differentially abundant. Using the main function ANCOM, all additive log-ratios for each taxon were then tested for significance using Wilcoxon rank-sum tests, and p-values were FDR-corrected using the BH method. ANCOM-II then applied a detection threshold as described in the original paper, whereby a taxon was called as DA if the number of corrected p-values reaching nominal significance for that taxon was greater than 60% of the maximum possible number of significant comparisons.

```
library(compositions)
## Welcome to compositions, a package for compositional data analysis.
## Find an intro with "? compositions"
##
## Attaching package: 'compositions'
## The following object is masked from 'package:NADA':
##
##
##
  The following object is masked from 'package:ape':
##
##
       balance
## The following objects are masked from 'package: IRanges':
##
##
       cor, cov, var
## The following objects are masked from 'package:S4Vectors':
##
##
       cor, cov, var
## The following objects are masked from 'package:BiocGenerics':
##
##
       normalize, var
## The following objects are masked from 'package:stats':
##
##
       anova, cor, cov, dist, var
## The following object is masked from 'package:graphics':
```

```
##
##
       segments
## The following objects are masked from 'package:base':
       %*%, norm, scale, scale.default
library(exactRankTests)
## Package 'exactRankTests' is no longer under development.
## Please consider using package 'coin' instead.
library(nlme)
source('~/Course/NDMC 2025/source/ancom v2.1.R')
meta$Sample <- rownames(meta)</pre>
prepro <- feature_table_pre_process(feature_table = data, meta_data = meta,</pre>
                                     sample_var = 'Sample', group_var = 'Class',
                                     out_cut = 0.05, zero_cut = 0.90,
                                     lib_cut = 1000, neg_lb=FALSE)
feature_table <- prepro$feature_table</pre>
metadata <- prepro$meta_data</pre>
struc_zero <- prepro$structure_zeros</pre>
main_var <- 'Class'</pre>
p_adj_method = "BH"
alpha=0.05
adj_formula=NULL
rand_formula=NULL
ANCOM_result <- ANCOM(feature_table = feature_table, meta_data = metadata,
             struc_zero = struc_zero, main_var = main_var, p_adj_method = p_adj_method,
             alpha=alpha, adj formula = adj formula, rand formula = rand formula)
ANCOM_result <- ANCOM_result$out
ANCOM_result <- ANCOM_result[ANCOM_result$W != 0, ]</pre>
head(ANCOM_result)
       taxa_id
                 W detected_0.9 detected_0.8 detected_0.7 detected_0.6
## 6
       ASV0007 457
                         FALSE
                                        FALSE
                                                      TRUE
                                                                    TRUE
## 28 ASV0030
                          FALSE
                                        FALSE
                                                     FALSE
                                                                   FALSE
## 40 ASV0042
                          FALSE
                                        FALSE
                                                     FALSE
                                                                   FALSE
                1
## 43 ASV0045
                                        FALSE
                                                                   FALSE
               3
                          FALSE
                                                     FALSE
## 103 ASV0119 467
                                                                    TRUE
                          FALSE
                                        FALSE
                                                      TRUE
## 113 ASV0130 2
                          FALSE
                                        FALSE
                                                     FALSE
                                                                   FALSE
#write.table(out, "~/Course/NDMC_2025/tmp/ANCOM.txt", quote=FALSE, sep="\t", col.names = F, row.names =
ANCOM_selected <- ANCOM_result[ANCOM_result$detected_0.6, ]
head(ANCOM selected)
##
                 W detected_0.9 detected_0.8 detected_0.7 detected_0.6
       taxa id
## 6 ASV0007 457
                          FALSE
                                        FALSE
                                                      TRUE
                                                                    TRUE
## 103 ASV0119 467
                          FALSE
                                        FALSE
                                                      TRUE
                                                                    TRUE
## 155 ASV0178 495
                          FALSE
                                        FALSE
                                                      TRUE
                                                                    TRUE
## 182 ASV0222 395
                          FALSE
                                        FALSE
                                                     FALSE
                                                                    TRUE
## 255 ASV0323 Inf
                           TRUE
                                         TRUE
                                                      TRUE
                                                                    TRUE
## 468 ASV0842 Inf
                           TRUE
                                         TRUE
                                                      TRUE
                                                                    TRUE
#write.table(out, "~/Course/NDMC_2025/tmp/ANCOM_thr.txt", quote=FALSE, sep="\t", col.names = F, row.nam
```

edgeR

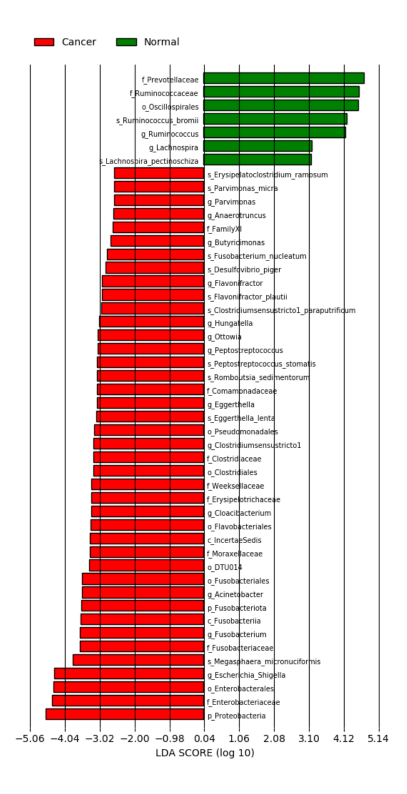
We added a pseudocount of 1 to the data and used the function calcNormFactors from the edgeR to compute relative log expression normalization factors. Negative binomial dispersion parameters were then estimated using the functions estimateCommonDisp followed by estimateTagwiseDisp to shrink featurewise dispersion estimates through an empirical Bayes approach. We then used the exactTest for negative binomial data to identify features that differ between the specified groups. The resulting p-values were then corrected for multiple testing with the BH method with the function topTags.

```
library(edgeR)
## Loading required package: limma
## Attaching package: 'limma'
## The following object is masked from 'package:BiocGenerics':
##
##
       plotMA
ASV <- phyloseq::otu_table(data, taxa_are_rows = T)
sampledata <- phyloseq::sample data(meta, errorIfNULL = T)</pre>
phylo <- phyloseq::merge_phyloseq(ASV, sampledata)</pre>
test <- phyloseq_to_edgeR(physeq = phylo, group = "Class") # phyloseq_to_edgeR function is sourced from
et = exactTest(test)
out = topTags(et, n=nrow(test$table), adjust.method="BH", sort.by="PValue")
edgeR_result <- out@.Data[[1]]</pre>
edgeR_selected <- edgeR_result[edgeR_result$FDR < 0.05,]</pre>
edgeR_selected <- cbind(rownames(edgeR_selected), edgeR_selected)</pre>
head(edgeR_selected)
           rownames(edgeR_selected)
                                          logFC
                                                    logCPM
                                                                 PValue
## ASV0075
                             ASV0075 -10.005478 11.509109 4.983343e-12 1.089973e-08
## ASV0029
                             ASV0029 -11.378337 12.878259 6.000401e-12 1.089973e-08
## ASV0178
                             ASV0178 -7.512891 9.946950 1.006243e-11 1.218560e-08
## ASV0073
                             ASV0073 -10.009413 11.513042 3.960265e-11 2.895228e-08
## ASV0083
                             ASV0083 -9.866453 11.370704 4.697678e-11 2.895228e-08
## ASV0198
                             ASV0198 -8.189402 9.708196 5.451478e-11 2.895228e-08
# write.table(subout, "~/Course/NDMC 2025/tmp/edgeR.txt", quote=FALSE, sep="\t", col.names = F, row.nam
```

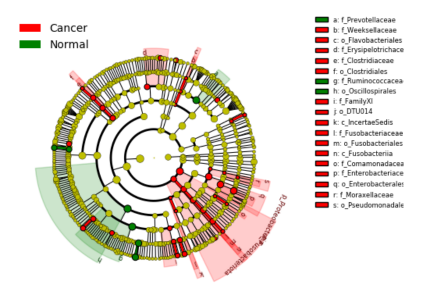
LEfSe

LEfSe performed a Kruskal-Wallis (which in our two-group case reduces to the Wilcoxon rank-sum) hypothesis test to identify potential differentially abundant features, followed by linear discriminant analysis (LDA) of class labels on abundances to estimate the effect sizes for significant features. From these, only those features with scaled LDA analysis scores above the threshold score of 2.0 (default) were called as differentially abundant.

```
## k_Bacteria|p_Proteobacteria "13.0343654304185" "6.13142629297594"
## k_Bacteria|p_Firmicutes
                               "58.7764545763865" "40.1578195943252"
##
                               [,3]
                                                 [.4]
## Class
                               "Normal"
                                                  "Normal"
                               "ERR475478"
## sampleID
                                                 "ERR475480"
## k_Bacteria
                               "89.5623741239996" "98.1584169974317"
## k_Archaea
                               "3.39050176528261" "0"
## k_Bacteria|p_Proteobacteria "6.6023018705717" "16.199217837964"
                               "71.9902052171225" "69.8721690403923"
## k_Bacteria|p_Firmicutes
                               Γ.51
## Class
                               "Normal"
                               "ERR475483"
## sampleID
## k_Bacteria
                               "98.2261609156035"
## k_Archaea
## k_Bacteria|p_Proteobacteria "7.60909895253589"
## k_Bacteria|p_Firmicutes
                              "88.7233496018495"
dir.create("~/Course/NDMC_2025/tmp/LEfSe_tmp")
write.table(ret_tab, "~/Course/NDMC_2025/tmp/LEfSe_tmp/tmp_in.txt", quote=FALSE, sep="\t", col.names = 1
# LEfSe execution by docker
$ docker run -u $UID:$(id -g) -it --rm -v /home/yincheng23/Course/NDMC_2025/tmp/LEfSe_tmp:/tmp yincheng
$ format_input.py /tmp/tmp_in.txt /tmp/data_in -s -1 -u 2 -o 1000000
$ run_lefse.py /tmp/data_in /tmp/LEfSe_res.txt
$ plot_res.py /tmp/LEfSe_res.txt /tmp/LEfSe_res.png --dpi 100
$ plot_cladogram.py /tmp/LEfSe_res.txt /tmp/cladogram.png --format png --dpi 100
$ cat /tmp/LEfSe_res.txt | awk '{if($3>2){print $0}}' > /tmp/LEfSe_res_selected.txt # filtering by /LD
result <- read.csv("~/Course/NDMC_2025/tmp/LEfSe_tmp/LEfSe_res.txt", sep = '\t', header = F)</pre>
colnames(result) <- c('taxa','log10LDA','tendency','U','p')</pre>
head(result)
```



Cladogram



PreLect

PreLectR is an R package implementing the PreLect algorithm, which integrates L1 regularization with an inverted prevalence penalty to select universal and informative features in sparse data.

It supports four tasks: binary classification, multi-class classification, regression, and time-to-event analysis.

Binary classification :
$$J(\mathbf{w}) = BCE(\mathbf{y}, \hat{\mathbf{y}}) + \lambda \sum_{j} \frac{|\mathbf{w}_{j}|}{p_{j}}$$

Regression :
$$J(\mathbf{w}) = \text{MSE}(\mathbf{y}, \hat{\mathbf{y}}) + \lambda \sum_{j} \frac{|\mathbf{w}_{j}|}{p_{j}}$$

Multi-class classification :
$$J(\mathbf{w}) = \frac{1}{c} \sum_{l=1}^{c} \left(\text{BCE}(\mathbf{y}_{l}, \hat{\mathbf{y}}_{l}) + \lambda \sum_{j=1}^{d} \frac{|\mathbf{w}_{j,l}|}{p_{j,l}} \right)$$

Time-to-event :
$$J(\mathbf{w}) = h_0(t) \cdot e^{\sum x_i \cdot w} + \lambda \sum_j \frac{|\mathbf{w}_j|}{p_j}$$

Based on previous studies, we recommend using variance-stabilizing transformation (VST) for data normalization.

```
vst_table <- assay(vst)</pre>
# feature-wise z-standardization
data scaled <- t(scale(t(as.matrix(vst table))))</pre>
```

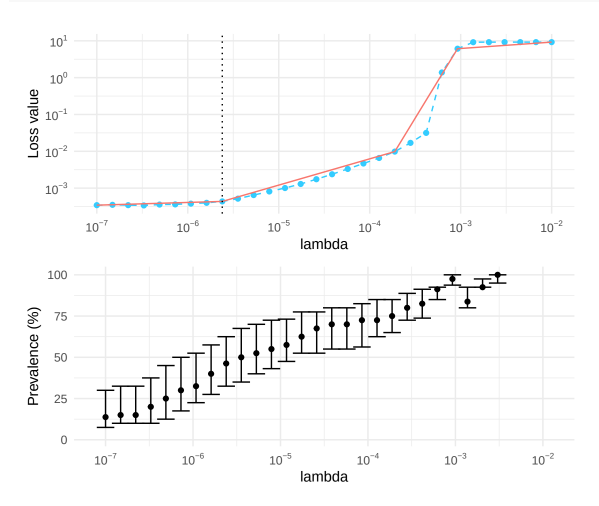
We will only use z-standardized table data_scaled and raw count table data for the subsequent analyses.

```
Automatically perform lambda scanning to identify 30 lambda values for examination.
meta$Class <- factor(meta$Class, levels = c("Normal", "Cancer")) # assign "Normal" as control sample
lrange <- AutoScanning(data_scaled, data, meta$Class, step =30)</pre>
length(lrange)
## [1] 30
exp(lrange)
## [1] 1.000000e-07 1.487352e-07 2.212216e-07 3.290345e-07 4.893901e-07
## [6] 7.278954e-07 1.082637e-06 1.610262e-06 2.395027e-06 3.562248e-06
## [11] 5.298317e-06 7.880463e-06 1.172102e-05 1.743329e-05 2.592944e-05
## [16] 3.856620e-05 5.736153e-05 8.531679e-05 1.268961e-04 1.887392e-04
## [21] 2.807216e-04 4.175319e-04 6.210169e-04 9.236709e-04 1.373824e-03
## [26] 2.043360e-03 3.039195e-03 4.520354e-03 6.723358e-03 1.000000e-02
# Examining the testing lambda.
dir.create("~/Course/NDMC_2025/tmp/PreLect_tmp")
output_path <-"~/Course/NDMC_2025/tmp/PreLect_tmp"</pre>
tuning_res <- LambdaTuningParallel(data_scaled, data, meta$Class, lrange, n_cores=10, outpath=output_pa
head(tuning res$TuningResult)
     Feature_number Percentage Prevalence AUC loss_history error_history
## 1
              1051 0.2892926
                                    0.15 0.75 0.0003444818 9.998251e-05
## 2
               988 0.2719516
                                    0.15 0.75 0.0003522416 9.997643e-05
## 3
               943 0.2595651
                                    0.15 0.75 0.0003449498 9.998842e-05
                                     0.20 0.75 0.0003388232 9.999399e-05
## 4
                805 0.2215800
                                     0.25 0.75 0.0003578406 9.999683e-05
## 5
                653 0.1797413
## 6
                556 0.1530416
                                     0.30 0.75 0.0003616958 9.990256e-05
##
      loglmbd
## 1 -16.11810
## 2 -15.72110
## 3 -15.32410
## 4 -14.92710
## 5 -14.53011
## 6 -14.13311
head(tuning_res$PvlDistSummary)
         llmbd max
                                   q1 min
                      q3
                             q2
                                        0
                                        0
```

```
##
0
0
## 6 -14.13311 1 0.500 0.3000 0.175
       0
```

Determine the optimal lambda by partitioning tree.

lmbd_picking <- LambdaDecision(tuning_res\$TuningResult, tuning_res\$PvlDistSummary, maxdepth=5, minbucke
lmbd_picking\$selected_lmbd_plot/lmbd_picking\$pvl_plot</pre>



```
print(lmbd_picking$opt_lmbd)
## [1] 2.395027e-06
```

PreLect execution and get the property for each feature

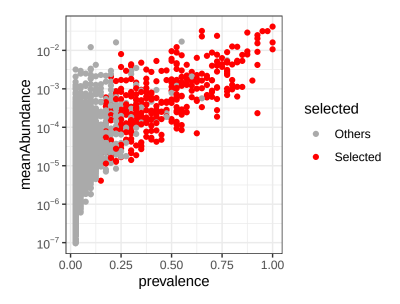
```
# We strongly suggest using 100000 for max_iter to achieve more accurate results.
s=Sys.time()
prevalence <- GetPrevalence(data)
PreLect_out <- PreLect(data_scaled, prevalence, meta$Class, lambda=lmbd_picking$opt_lmbd, max_iter = 10
print(Sys.time()-s)
## Time difference of 2.99019 secs

featpropt <- FeatureProperty(data, meta$Class, PreLect_out, task="classification")</pre>
```

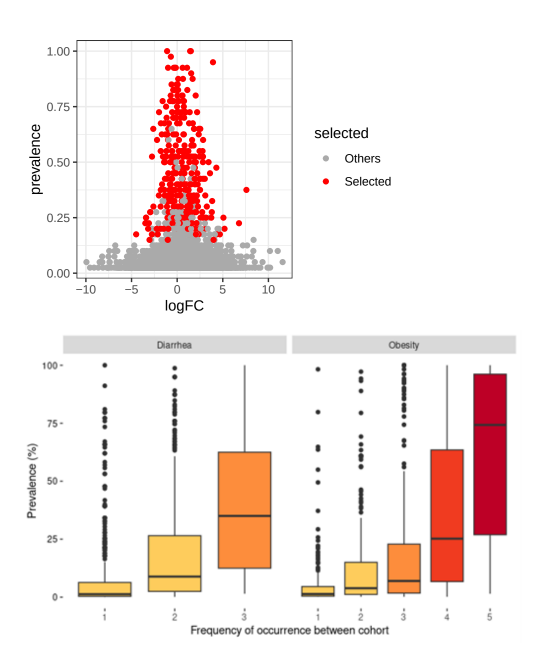
print(paste(nrow(featpropt[featpropt\$selected == 'Selected',]), 'features were selected'))

```
## [1] "336 features were selected"
print(paste('median of prevalence :', median(featpropt$prevalence[featpropt$selected == 'Selected'])))
## [1] "median of prevalence : 0.425"
write.table(featpropt, "~/Course/NDMC_2025/tmp/PreLect_tmp/feature_propt.txt", quote=FALSE, sep="\t")
head(featpropt)
                           coef tendency selected meanAbundance
           FeatName
                                                                   variance
## ASV0001 ASV0001 0.54988067
                                  Cancer Selected
                                                     0.04150302
                                                                 117734960
## ASV0002
           ASV0002 -0.42573824
                                  Normal Selected
                                                     0.03167390
                                                                   64237504
## ASV0003 ASV0003 0.00000000
                                    <NA>
                                           Others
                                                     0.01213962 1555924984
## ASV0004 ASV0004 -0.33145665
                                  Normal Selected
                                                     0.03201389 288009324
## ASV0005 ASV0005 -0.32683697
                                                     0.02903994
                                  Normal Selected
                                                                   41768447
## ASV0006 ASV0006 -0.03458693
                                  Normal Selected
                                                     0.02485074
                                                                   72058376
##
           prevalence prevalence_case prevalence_control
                                                               logFC
## ASV0001
                1.000
                                 1.00
                                                    1.00 1.4998663
## ASV0002
                0.975
                                 0.95
                                                     1.00 -0.6884894
## ASV0003
                0.100
                                 0.05
                                                    0.15 11.0217666
## ASV0004
                0.650
                                 0.60
                                                    0.70 -2.6154169
                                                    0.95 0.6372900
## ASV0005
                                 0.80
                0.875
## ASV0006
                0.925
                                 0.90
                                                     0.95 -0.1468191
```

Selection profile visualization and evaluation



```
ggplot(featpropt, aes(x = logFC, y = prevalence, color=selected)) + geom_point() +
    scale_color_manual(values = c('Selected'='red', 'Others'='#AAAAAA')) +
    theme_bw()+ theme(panel.background = element_rect(fill = "white", colour = "white"))
```



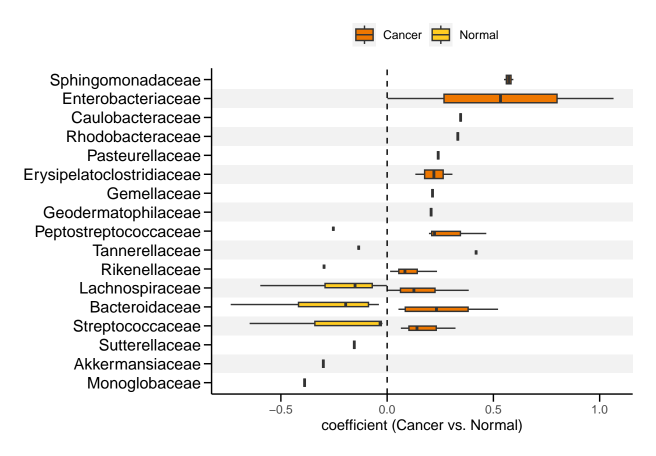
We observed that features present in multiple datasets tend to have higher prevalence within those datasets. This finding suggests that features with higher prevalence in one dataset are more likely to be universal across different datasets.

[1] 0.875

```
result <- TaxaProperty(featpropt, taxa, "Family", pvl_filter = 0.5)

mycolor <- c("Normal" = "#FFCC22", "Cancer" = "#EE7700")

result$effectSizePlot + scale_fill_manual(values = mycolor) +
    geom_hline(yintercept = 0, color='black', linetype='dashed')</pre>
```



```
head(result$selectedInfo)
##
           FeatName
                          coef tendency selected meanAbundance variance
## ASV0001 ASV0001 0.5498807
                                                    0.04150302 117734960
                                 Cancer Selected
## ASV0002 ASV0002 -0.4257382
                                 Normal Selected
                                                    0.03167390 64237504
## ASV0005
           ASV0005 -0.3268370
                                 Normal Selected
                                                    0.02903994 41768447
           ASV0007 1.0654049
## ASV0007
                                 Cancer Selected
                                                    0.03158703 178336011
## ASV0008
           ASV0008 -0.3002699
                                 Normal Selected
                                                    0.02299001 312317781
## ASV0009 ASV0009 -0.4994069
                                 Normal Selected
                                                    0.02042962 45214122
##
           prevalence prevalence_case prevalence_control
                                                              logFC
                                 1.00
## ASV0001
                1.000
                                                    1.00 1.4998663
## ASV0002
                0.975
                                 0.95
                                                    1.00 -0.6884894
## ASV0005
                0.875
                                 0.80
                                                    0.95
                                                         0.6372900
                0.950
                                 1.00
## ASV0007
                                                    0.90
                                                          3.9252053
## ASV0008
                0.650
                                 0.65
                                                    0.65
                                                          1.8688587
## ASV0009
                0.925
                                 0.85
                                                    1.00 0.5991814
##
                         taxa
## ASV0001 Sphingomonadaceae
```

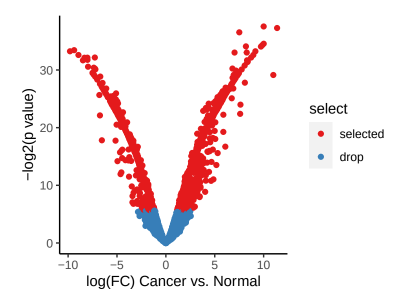
```
## ASV0002 Lachnospiraceae
## ASV0005 Bacteroidaceae
## ASV0007 Enterobacteriaceae
## ASV0008 Akkermansiaceae
## ASV0009 Lachnospiraceae
```

Basal feature visualization

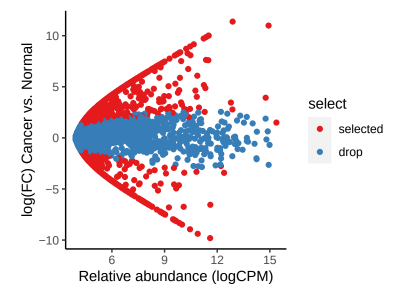
we use edgeR to demonstrate the several plots.

```
library(edgeR)
ASV <- phyloseq::otu_table(data, taxa_are_rows = T)
sampledata <- phyloseq::sample_data(meta, errorIfNULL = T)</pre>
phylo <- phyloseq::merge_phyloseq(ASV, sampledata)</pre>
test <- phyloseq_to_edgeR(physeq = phylo, group = "Class")</pre>
et = exactTest(test)
out = topTags(et, n=nrow(test$table), adjust.method="BH", sort.by="PValue")
edgeR_result <- out@.Data[[1]]</pre>
edgeR_selected <- edgeR_result[edgeR_result$FDR < 0.05,]</pre>
et = exactTest(test)
et = et$table
et$select <- "drop"
et$select[rownames(et) %in% rownames(edgeR_selected)] = "selected"
data_freq <- data
for(i in 1:ncol(data_freq)){
 data_freq[,i] <- data_freq[,i]/colSums(data_freq)[i]</pre>
}
get_pvl <- function(m){</pre>
 m_ <- m
 m_ <- as.matrix(m_)</pre>
 return(as.numeric(rowSums(m_)/ncol(m_)))
variance <- c()</pre>
for(i in 1:nrow(data_freq)){
  variance <- c(variance, log(sd(data_freq[i,])**2))</pre>
et$variation <- variance</pre>
et$prevalence <- get_pvl(data)</pre>
et$neglog2p <- -log(et$PValue,2)
et$select <- factor(et$select, levels = c('selected', 'drop'))</pre>
ggplot(et, aes(x = logFC, y = neglog2p, color = select)) + geom_point() +
 xlab('log(FC) Cancer vs. Normal') + ylab('-log2(p value)') +
  scale_color_brewer(palette = 'Set1') + theme(axis.line = element_line(linetype = 1,colour = 'black'),
```

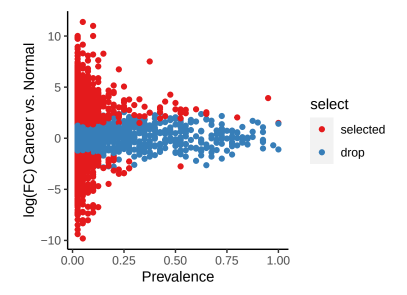
```
panel.background = element_rect(I(0)),
panel.grid.major = element_line(colour = NA),
panel.grid.minor = element_line(colour = NA))
```



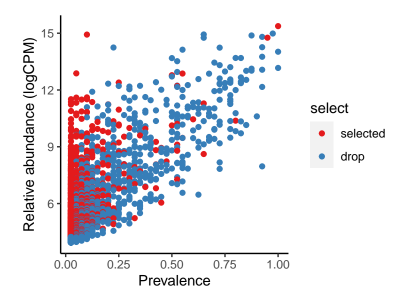
```
ggplot(et, aes(x = logCPM, y = logFC, color = select)) + geom_point() +
    xlab('Relative abundance (logCPM)') + ylab('log(FC) Cancer vs. Normal') +
    scale_color_brewer(palette = 'Set1') +
    theme(axis.line = element_line(linetype = 1,colour = 'black'),
        panel.background = element_rect(I(0)),
        panel.grid.major = element_line(colour = NA),
        panel.grid.minor = element_line(colour = NA))
```



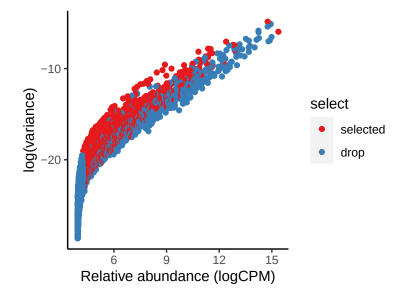
```
ggplot(et, aes(x = prevalence, y = logFC, color = select)) + geom_point() +
    xlab('Prevalence') + ylab('log(FC) Cancer vs. Normal') +
    scale_color_brewer(palette = 'Set1') +
    theme(axis.line = element_line(linetype = 1,colour = 'black'),
        panel.background = element_rect(I(0)),
        panel.grid.major = element_line(colour = NA),
        panel.grid.minor = element_line(colour = NA))
```



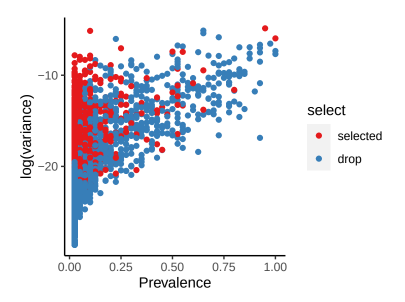
```
ggplot(et, aes(x = prevalence, y = logCPM, color = select)) + geom_point() +
    xlab('Prevalence') + ylab('Relative abundance (logCPM)') +
    scale_color_brewer(palette = 'Set1') +
    theme(axis.line = element_line(linetype = 1,colour = 'black'),
        panel.background = element_rect(I(0)),
        panel.grid.major = element_line(colour = NA),
        panel.grid.minor = element_line(colour = NA))
```



```
ggplot(et, aes(x = logCPM, y = variance, color = select)) + geom_point() +
xlab('Relative abundance (logCPM)') + ylab('log(variance)') +
scale_color_brewer(palette = 'Set1') +
theme(axis.line = element_line(linetype = 1,colour = 'black'),
    panel.background = element_rect(I(0)),
    panel.grid.major = element_line(colour = NA),
    panel.grid.minor = element_line(colour = NA))
```



```
ggplot(et, aes(x = prevalence, y = variance, color = select)) + geom_point() +
    xlab('Prevalence') + ylab('log(variance)') +
    scale_color_brewer(palette = 'Set1') +
    theme(axis.line = element_line(linetype = 1,colour = 'black'),
        panel.background = element_rect(I(0)),
        panel.grid.major = element_line(colour = NA),
        panel.grid.minor = element_line(colour = NA))
```

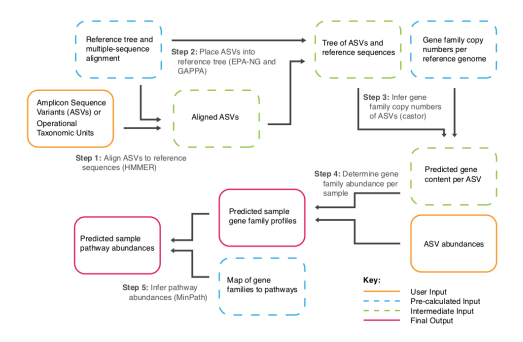


Functional Prediction

PICRUSt2

Prepare files for running PICRUSt2 by extracting the selected ASVs.

```
# please ensure the "ASV.fasta" is in "output_path" directory
output_path <- '~/Course/NDMC_2025/tmp'</pre>
dir(output_path)
## [1] "ASV_table.txt"
                             "ASV_taxa_table.txt" "ASV.fasta"
## [4] "ASV.tree"
                             "for_PICRUSt2.tsv"
                                                    "LEfSe_tmp"
                             "PreLect tmp"
## [7] "PICRUSt2"
                                                    "ssuout"
selected_id <- featpropt$FeatName[featpropt$selected == 'Selected'] # selected by PreLect</pre>
data_sub <- data[selected_id, ] # if you want to use whole ASVs, just data_sub <- data
data_sub <- rbind(colnames(data_sub), data_sub)</pre>
colnames(data_sub) <- NULL</pre>
rownames(data_sub)[1] <- "#OTU ID"</pre>
write.table(data_sub, paste0(output_path,"/for_PICRUSt2.tsv"), sep = "\t", quote=F, col.names
                                                                                                     =F)
```



Conduct the PICRUSt2 pipeline via docker

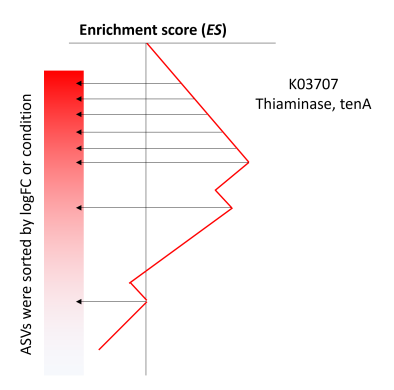
```
# Bash
# run the PICRUSt2 pipeline
# ensure the "ASV. fasta" and "for_PICRUSt2.tsv" files are in the same directory
# and set the working directory to that location before running this script.
$
 cd ~/Course/NDMC_2025/tmp
 docker run \
$
     -u $UID:$(id -g) \setminus
$
                                      # perform this work as the current user.
$
     -i -t \
                                     # interactive terminal mode, allowing you to interact with the cont
$
     --rm \
                                     # automatically remove the container after it exits to save disk sp
$
     -v $(pwd):/tmp \
                                     # bind mount the current directory ($(pwd)) to /tmp in the containe
$
                                     # specify the Docker image to use; it will be pulled from Docker Hu
     yincheng23/picrust2:0.2.0 \
$
     sh /bin/Run_picrust2.sh 10
                                     # use the shell to execute the built-in script (Run picrust2.sh) wi
# the results will store as ~/Course/NDMC_2025/tmp/PICRUSt2
```

output file : ./PICRUSt2/KO/pred_metagenome_contrib.tsv - Represents the relationship between functions and taxa.

output file : ./ $PICRUSt2/KO/pred_metagenome_unstrat_descrip.tsv$ - Contains KO (KEGG Orthology) abundances for each sample along with detailed gene descriptions.

output file: ./PICRUSt2/KO/pred_metagenome_unstrat.tsv.gz - Provides KO abundances for each sample.

Since PICRUSt2 predicts gene abundance based on taxa abundance, differential expression analysis using this approach may be less convincing. Therefore, we adopted a GSEA strategy, ranking taxa by fold change based on their abundance. Using insights provided by PICRUSt2, we examined which species carry specific genes to assess the KOs that are actively expressed or suppressed.



Conduct the GSEA with permutation test via GSEATestwithFC or GSEATest

```
# load the KO-taxa relationship file
# output_path is the directory you save the for_PICRUSt2.tsv file
KOindex <- read.table(paste0(output_path, "/PICRUSt2/KO/pred_metagenome_contrib.tsv"), sep = "\t", heade
# extract the selected ASV identifiers
selected_id <- featpropt$FeatName[featpropt$selected == 'Selected']</pre>
# Provide the raw count table for logFC calculation, subsetting with the selected ASVs.
# Additionally, specify the labels for each sample and the name of the case sample group.
GSEAresult <- GSEATestwithFC(KOindex, data[selected_id, ], meta$Class, "Cancer")
## Building the KO-to-taxa mapper...
## Done. In total, 5755 KOs need to be processed.
## Shuffling the labels for GSEA...
## Performing GSEA to identify activated KOs...
   |-----
## Shuffling the labels for GSEA...
## Performing GSEA to identify suppressed KOs...
## Done.
# If the grouping variable is continuous (e.g., body mass index), use the following line instead:
# GSEAresult <- GSEATest(KOindex, data[selected_id, ], meta$body.mass.index)</pre>
```

Selected the enriched KOs with z-score

```
Actived_result <- GSEAresult$Actived_KO
Actived_result <- Actived_result[!is.na(Actived_result$z),]
Actived_result <- Actived_result[Actived_result$z > 2,]
Actived_result <- Actived_result[Actived_result$p < 0.05,]
nrow(Actived_result)
## [1] 630
head(Actived_result)
## KO ES z p
## 1 K02415 0.3022222 2.092140 0.037
## 9 K07067 0.5423197 3.000060 0.006
## 11 K09154 0.8323353 2.352198 0.022
## 21 K00342 0.3318762 2.717256 0.013
## 36 K14415 0.4094937 2.272980 0.029
## 40 K01087 0.5552147 2.681651 0.013
```

Since PICRUSt2 does not provide detailed information for each KO, we preprocess the KO-pathway table using the KEGG API.

```
kodb_path <- system.file("exdata", "total_KO_pathway.rds", package = "PreLectR")</pre>
KOinfo <- readRDS(kodb_path)</pre>
head(KOinfo)
         KO
                   symbol
                                                         name
## 1 K00001 E1.1.1.1, adh alcohol dehydrogenase [EC:1.1.1.1] map00010
## 2 K00001 E1.1.1.1, adh alcohol dehydrogenase [EC:1.1.1.1] map00071
## 3 K00001 E1.1.1.1, adh alcohol dehydrogenase [EC:1.1.1.1] map00350
## 4 K00001 E1.1.1.1, adh alcohol dehydrogenase [EC:1.1.1.1] map00620
## 5 K00001 E1.1.1.1, adh alcohol dehydrogenase [EC:1.1.1.1] map00625
## 6 K00001 E1.1.1.1, adh alcohol dehydrogenase [EC:1.1.1.1] map00626
##
                                        pathway
## 1
                  Glycolysis / Gluconeogenesis
## 2
                        Fatty acid degradation
## 3
                           Tyrosine metabolism
                           Pyruvate metabolism
## 5 Chloroalkane and chloroalkene degradation
                       Naphthalene degradation
## 6
```

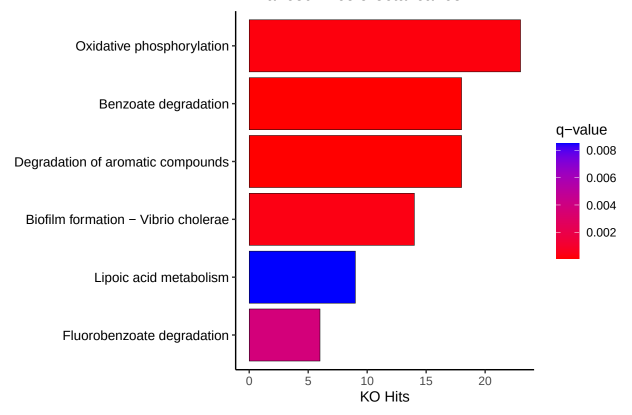
Examining the condition enriched pathway with Fisher's exact test via PathwayEnrichment

```
KOinfo <- KOinfo[KOinfo$KO %in% unique(KOindex$function.), ]</pre>
Actived_result <- GSEAresult$Actived_KO</pre>
Actived result <- Actived result[!is.na(Actived result$z), ]
Actived_result <- Actived_result[Actived_result$z > 2,]
Actived_result <- Actived_result[Actived_result$p < 0.05,]</pre>
selected_KOs <- Actived_result$KO</pre>
enrichPW <- PathwayEnrichment(selected_KOs, KOinfo)</pre>
enrichPW$q <- p.adjust(enrichPW$p, method = 'fdr')</pre>
enrichPW <- enrichPW[enrichPW$q < 0.05, ]</pre>
nrow(enrichPW)
## [1] 6
head(enrichPW)
##
                                      pathway
                                                      id count
                                                                    ratio
```

```
## 26
        Degradation of aromatic compounds map01220
                                                      18 0.3829787 9.848597e-07
## 31
                   Lipoic acid metabolism map00785
                                                       9 0.4090909 2.743792e-04
## 34
                Oxidative phosphorylation map00190
                                                      23 0.3026316 4.214420e-06
                     Benzoate degradation map00362
## 64
                                                      18 0.4000000 4.614879e-07
               Fluorobenzoate degradation map00364
## 112
                                                      6 0.6666667 1.007485e-04
## 133 Biofilm formation - Vibrio cholerae map05111
                                                      14 0.4000000 7.841760e-06
##
       odds_ratio
## 26
        5.161160 9.159195e-05
## 31
        5.756061 8.505756e-03
        3.605573 2.612941e-04
## 34
## 64
        5.543134 8.583675e-05
## 112 16.614397 3.747845e-03
## 133 5.572086 3.646418e-04
```

```
enrichPW$pathway <- factor(enrichPW$pathway, levels = enrichPW$pathway[order(enrichPW$count)])
ggplot(enrichPW, aes(x = pathway, y = count, fill = q)) + ggtitle('Enhanced in colorectal cancer') +
    geom_bar(stat="identity",colour = "black",size = 0.1) + coord_flip() + labs(y = 'KO Hits', fill = "q
    scale_fill_continuous(low = "red", high = "blue", limits = range(enrichPW$q)) +
    theme(axis.text.y = element_text(color='black', size='10'),
        axis.line = element_line(linetype = 1,colour = 'black'),
        axis.title.y = element_blank(),
        panel.background = element_rect(I(0)),
        panel.grid.major = element_line(colour = NA),
        panel.grid.minor = element_line(colour = NA)) + coord_flip()</pre>
```

Enhanced in colorectal cancer



RPM

Reference Pathways Mapper (RPM) utilizes a reference pathways database, which is a flat file listing pathway/module reactions in sequential order. Tab-separated reactions indicate alternative reactions (OR operation), while line breaks and comma-separated reactions represent reactions that are all required for pathway completion (AND operation).

Below is a snippet from the human gut metabolic modules (GMMs) database, as described by Vieira-Silva et al. 2016.

```
MF0001 arabinoxylan degradation
K01209 K15921 K01181 K01198 K15531 K18205
///
MF0003 pectin degradation I
K01051
K01184,K01213
               K18650
///
MF0103 mucin degradation
K01186
K05970
K01132 K01135 K01137 K01205
K01207 K12373 K14459
K01205 K01207 K12373 K01227 K13714
K01206
///
```

Installation According to original repository

```
$ wget https://github.com/omixer/omixer-rpmR/releases/download/0.3.3/omixerRpm_0.3.3.tar.gz
$ R CMD INSTALL omixerRpm_0.3.3.tar.gz
```

```
library(omixerRpm)
library(dplyr)
library(tidyr)

KOtable <- read.table(pasteO(output_path,"/PICRUSt2/KO/pred_metagenome_contrib.tsv"), sep = "\t", heade

colnames(KOtable)[2] <- 'fun'
sumtb <- KOtable %>% group_by(sample, fun) %>% summarise(sum_rel_function_abun=sum(taxon_rel_function_a'
tmp <- spread(sumtb, key = sample, value = sum_rel_function_abun)
colnames(tmp)[1] <- 'entry'
tmp[is.na(tmp)] <- 0

mods <- rpm(tmp, minimum.coverage=0.3, annotation = 1)
## [1] "Loaded GMMs.v1.07"</pre>
```

```
db <- loadDefaultDB()</pre>
## [1] "Loaded GMMs.v1.07"
getNames(db, mods@annotation[1,])
## [1] "ribose degradation"
coverage <- asDataFrame(mods, "coverage")</pre>
module <- mods@abundance</pre>
rownames(module) <- coverage$Description</pre>
head(module)
##
                                 ERR475527 ERR475549
                                                        ERR475504
                                                                   ERR475529
## ribose degradation
                                35.16937498 35.17516 3.098085e+01 55.4419850
## tyrosine degradation II
                                             0.00000 0.000000e+00
                                0.00736405
                                                                    0.1532172
## aspartate degradation I
                                56.20638043 68.65780 5.569382e+01 54.3297498
## tryptophan degradation
                                5.20525631
                                             1.12031 1.160618e+01
                                                                     3.2142130
## tyrosine degradation I
                                50.58997819 101.04308 5.424949e+01 104.9491610
## galacturonate degradation II 0.00000000
                                             0.00000 9.507594e-04
                                                                     0.0000000
##
                                ERR475528 ERR475545
                                                        ERR475500
                                                                     ERR475588
## ribose degradation
                                27.8606223 27.09122669 63.79307368 50.59532850
                                0.0159962 0.00000000 0.14151356 0.00000000
## tyrosine degradation II
## aspartate degradation I
                                79.0703436 87.57625174 87.98962719 63.41611505
                                10.6104558 21.77601754 2.87128420 33.18194936
## tryptophan degradation
## tyrosine degradation I
                                51.9888339 60.34117553 53.45470956 41.96002481
## galacturonate degradation II 0.0000000 0.01126817 0.01038632 0.03272184
##
                               ERR475547 ERR475541 ERR475485
                                                                 ERR475540
## ribose degradation
                                 31.55949 28.4148131 25.326445 23.52926496
## tyrosine degradation II
                                 0.00000 0.1387234 0.000000
                                                                0.08120360
## aspartate degradation I
                                 87.24203 59.8965261 49.600167 117.38540049
## tryptophan degradation
                                 15.62138 4.5938590 3.226059 16.50346978
## tyrosine degradation I
                                 52.56882 51.8888387 50.360550 58.77502133
## galacturonate degradation II
                                0.00000 0.0175545 0.000000
                                                                 0.02160463
##
                               ERR475562
                                            ERR475584
                                                        ERR475521
                                                                      ERR475565
## ribose degradation
                                 21.27139 22.234725150 21.73869043 18.653580678
## tyrosine degradation II
                                 0.00000 0.019633212 0.08697272 0.095860634
## aspartate degradation I
                                 61.78199 51.502887344 71.82182553 36.918989696
## tryptophan degradation
                                 21.02328 4.870319828 13.28846530 12.576196223
## tyrosine degradation I
                                 56.55310 51.969782663 55.30241396 52.756376669
## galacturonate degradation II
                                 0.00000 0.005960082 0.04027999 0.002396516
##
                                  ERR475586 ERR475580
                                                         ERR475484 ERR475561
## ribose degradation
                                46.99615938 34.6893540 27.80085184 36.651911
## tyrosine degradation II
                                0.43654951 0.0000000
                                                       0.03075402 0.000000
## aspartate degradation I
                                78.71544166 48.5301653 63.21129882 53.275009
## tryptophan degradation
                                14.27417268 8.0092019
                                                        6.06292426 1.126657
## tyrosine degradation I
                                50.86031496 50.9725785 101.77878685 50.407778
## galacturonate degradation II 0.01942096 0.0221183
                                                        0.00000000 0.000000
                                 ERR475483 ERR475560 ERR475590
##
                                                                   ERR475518
## ribose degradation
                                 29.1709203 45.736165 29.9555807 28.04525333
## tyrosine degradation II
                                 0.0000000 0.000000 0.0000000 0.01163850
## aspartate degradation I
                                 58.3060655 41.854488 72.7521866 49.14149961
## tryptophan degradation
                                 0.6992597 2.599723 19.2800891 3.64808639
                                102.9874650 49.899148 54.3941577 53.93726862
## tyrosine degradation I
## galacturonate degradation II 0.0000000 0.000000 0.1025416 0.02168992
##
                                  ERR475534
                                               ERR475478
                                                           ERR475555
## ribose degradation
                                31.02395184 3.450413e+01 3.441916e+01 48.548405549
## tyrosine degradation II
                                0.05004837 1.083518e-01 0.000000e+00 0.127484993
```

```
## aspartate degradation I 55.09630176 5.241802e+01 6.256350e+01 57.795267824
## tryptophan degradation
                             2.10620205 6.076617e+00 3.097498e-01 1.738492718
## tyrosine degradation I
                             49.95218205 1.067859e+02 1.007209e+02 51.958954529
## galacturonate degradation II 0.00000000 8.334751e-03 3.997287e-03 0.009393631
##
                              ERR475513
                                         ERR475579
                                                   ERR475554
                                                                ERR475576
## ribose degradation
                             31.59691328 22.92837864 18.73800856 37.82823763
## tyrosine degradation II
                            90.11875161 91.17260136 76.39798496 148.87959127
## aspartate degradation I
## tryptophan degradation
                             25.45861537 23.86014480 1.94103675 59.91196697
## tyrosine degradation I
                             57.19484500 45.59180272 50.44075281 74.74717910
## galacturonate degradation II 0.07131394 0.02515217 0.00000000 0.01037918
##
                               ERR475476 ERR475553
                                                     ERR475493
                                                                 ERR475591
## ribose degradation
                             23.014066166 40.9080176 2.808007e+01 27.068650441
## tyrosine degradation II
                            ## aspartate degradation I
                             67.459882221 85.0328572 4.551332e+01 62.929806811
## tryptophan degradation
                            12.719718609  0.5422964  5.907959e+00  8.902784854
## tyrosine degradation I
                            53.791136776 99.3264616 1.021285e+02 56.030165476
## galacturonate degradation II 0.009710515 0.0000000 7.034617e-03 0.048438766
##
                             ERR475473 ERR475550 ERR475594
                                                             ERR475480
## ribose degradation
                            19.1812121 32.08871268 62.5142559 31.15863997
## tyrosine degradation II
                            0.1721522 0.03903622 0.3954753
                                                            0.00000000
## aspartate degradation I
                             58.6562929 78.48988082 94.8305828 35.21222110
## tryptophan degradation
                            4.5486796 0.62264043 5.3451859
                                                             0.16811702
## tyrosine degradation I
                             58.1781156 49.20786881 61.6465806 106.09704137
## galacturonate degradation II 0.0000000 0.00000000 0.2272061 0.03641655
```

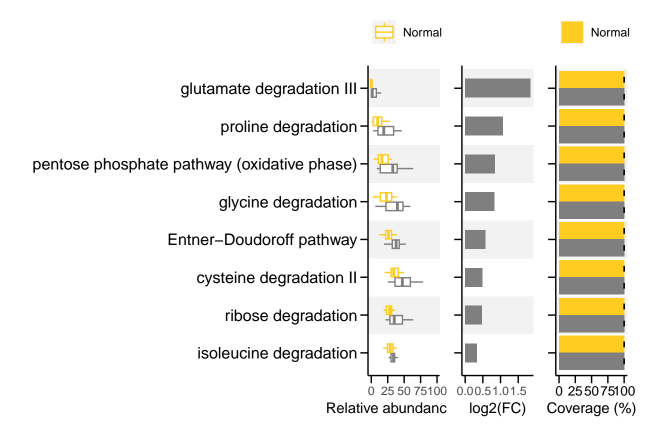
Pathway mapping

Module visualization Please ensure that the dataframe meta has a column named group, which records the grouping information.

Only binary grouping is supported in the following process.

```
meta$group <- meta$Class # pleas ensure this</pre>
# GMMviz function is sourced from utils.R
results <- GMMviz(module, meta, 'Normal') # please assign the control name as the third parameter.
sigdf <- results$significance_df</pre>
boxdf <- results$relative_abundance_df</pre>
covdf <- results$coverage_df</pre>
sigdf <- sigdf[sigdf$p < 0.01, ]</pre>
boxdf <- boxdf[boxdf$module %in% sigdf$module,]</pre>
covdf <- covdf[covdf$module %in% sigdf$module,]</pre>
module_order <- sigdf$module[order(sigdf$log2FC, decreasing = F)]</pre>
sigdf$module <- factor(sigdf$module, levels = module_order)</pre>
boxdf$module <- factor(boxdf$module, levels = module_order)</pre>
covdf$module <- factor(covdf$module, levels = module_order)</pre>
mycolor <- c("Normal" = "#FFCC22", "Cancer" = "#EE7700") # color assignment
p1 <- ggplot(boxdf, aes(x = module, y = value, color = group)) + coord_flip() +
      scale x discrete() +
```

```
ylab("Relative abundance (%)") + scale_color_manual(values = mycolor) +
      theme(panel.background = element_rect(fill = 'transparent'),
        panel.grid = element_blank(),
        axis.ticks.length = unit(0.4, "lines"),
        axis.ticks = element_line(color='black'),
        axis.line = element_line(colour = "black"),
        axis.text.y = element_text(colour='black',size=12),
        axis.title.y = element blank(),
        legend.title=element blank(),
        legend.position = 'top')
sing = 1
for (i in 1:(nrow(sigdf)-1)){
  sing = sing * -1
 p1 <- p1 + annotate('rect', xmin = i+0.5, xmax = i+1.5, ymin = -Inf, ymax = Inf,
                      fill = ifelse(sing > 0, 'white', 'gray95'))
p1 <- p1 + geom_boxplot(outlier.shape = NA, width = 0.5)
p2 <- ggplot(sigdf, aes(x = module, y = log2FC, fill = group)) + coord_flip() +
      geom_bar(stat = "identity", width=.5, position = "dodge") + scale_x_discrete() +
      scale_fill_manual(values = mycolor) + ylab("log2(FC)") +
      theme(panel.background = element_rect(fill = 'transparent'),
        panel.grid = element_blank(),
        axis.ticks.length = unit(0.4, "lines"),
        axis.ticks = element line(color='black'),
        axis.line = element_line(colour = "black"),
        #axis.text = element text(colour='black', size=10),
        axis.text.y = element_blank(),
        axis.title.y = element_blank(),
        legend.title=element_blank(),
       legend.position = 'top')
sing = 1
for (i in 1:(nrow(sigdf)-1)){
 sing = sing * -1
 p2 \leftarrow p2 + annotate('rect', xmin = i+0.5, xmax = i+1.5, ymin = -Inf, ymax = Inf,
                      fill = ifelse(sing > 0, 'white', 'gray95'))
p2 <- p2 + geom_bar(stat = "identity", width=.5, position = "dodge")
p3 <- ggplot(covdf, aes(x = module, y = mean, fill = group)) +
      scale_x_discrete() + coord_flip() + scale_fill_manual(values = mycolor) +
      theme(panel.background = element_rect(fill = 'transparent'),
        panel.grid = element blank(),
        axis.ticks.length = unit(0.4, "lines"),
       axis.ticks = element_line(color='black'),
       axis.line = element_line(colour = "black"),
        axis.text = element_text(colour='black',size=10),
        axis.text.y = element_blank(),
        axis.title.y = element_blank(),
       legend.title=element_blank(),
        legend.position = 'top')
sing = 1
for (i in 1:(nrow(sigdf)-1)){
```



Co-Occurrence Network

```
library(Rcpp)
library(igraph)
library(gtools)
library(Matrix)
library(SpiecEasi)
sourceCpp("source/Microbial_network.cpp")
```

SPIEC-EASI

SPIEC-EASI(**SP**arse **InversE** Covariance **E**stimation for **E**cological **A**ssociation **I**nference) infers direct microbial interactions from compositional data using a graphical model. It transforms counts into log-ratios,

estimates a sparse inverse covariance matrix with regularization (e.g., Graphical Lasso), and builds a network where non-zero elements indicate direct species relationships, overcoming spurious correlations.

```
pvl_filter <- 0.1</pre>
SpiecEasi_threshold <- 0.7
# filter low prevalence feature
pdata <- data
pdata[pdata > 0] <- 1</pre>
keep_id <- rownames(pdata)[rowSums(pdata) > ncol(pdata)*pvl_filter]
filerted_data <- data[keep_id,]</pre>
dim(filerted data)
## [1] 645 40
# run SpiecEas mb = Neighborhood Selection, glasso = graph LASSO
SPIEC_EASI <- spiec.easi(t(filerted_data), method='mb', lambda.min.ratio=1e-2, nlambda=15)
## Applying data transformations...
## Selecting model with pulsar using stars...
## Fitting final estimate with mb...
## done
adjm <- getOptMerge(SPIEC_EASI)</pre>
adjm <- as.matrix(adjm)</pre>
adjm[adjm < SpiecEasi_threshold] <- 0</pre>
edge_table <- make_edgetable(adjm, filerted_data)</pre>
node_table <- make_nodetable(edge_table, data, taxa)</pre>
```

SparCC

SparCC (Sparse Correlations for Compositional data) infers microbial interactions from compositional data by estimating correlations while accounting for the sum constraint. It assumes most species pairs are uncorrelated, uses log-ratio variance to approximate correlations, and applies a sparse model to focus on strong relationships. Iterative refinement reduces noise, producing a correlation network for ecological insights.

```
pvl filter <- 0.1</pre>
SparCC threshold <- 0.3</pre>
SparCC conf threshold <- 0.05
# filter low prevalence feature
pdata <- data
pdata[pdata > 0] <- 1</pre>
keep_id <- rownames(pdata)[rowSums(pdata) > ncol(pdata)*pvl_filter]
filerted_data <- data[keep_id,]</pre>
dim(filerted_data)
# run SparCC
sparcc_cor <- sparcc(t(filerted_data))</pre>
sparcc_boost <- sparccboot(t(filerted_data), R = 100, ncpus = 10)</pre>
sparcc_p <- pval.sparccboot(sparcc_boost)</pre>
cors <- sparcc_p$cors</pre>
pvals <- sparcc_p$pvals</pre>
# adjacency matrix constructing
```

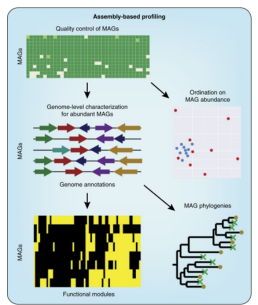
```
sparCCpcors <- diag(0.5, nrow = dim(sparcc_cor$Cor)[1], ncol = dim(sparcc_cor$Cor)[1])
sparCCpcors[upper.tri(sparCCpcors, diag=FALSE)] <- cors
sparCCpcors <- sparCCpcors + t(sparCCpcors)
sparCCpval <- diag(0.5, nrow = dim(sparcc_cor$Cor)[1], ncol = dim(sparcc_cor$Cor)[1])
sparCCpval[upper.tri(sparCCpval, diag=FALSE)] <- pvals
sparCCpval <- sparCCpval + t(sparCCpval)
rownames(sparCCpcors) <- rownames(filerted_data)
colnames(sparCCpcors) <- rownames(filerted_data)
rownames(sparCCpval) <- rownames(filerted_data)
sparCCpcors[abs(sparCCpcors) < SparCC_threshold | sparCCpval > SparCC_conf_threshold] = 0
edge_table <- make_edgetable(sparCCpcors, filerted_data)
node_table <- make_nodetable(edge_table, data, taxa)</pre>
```

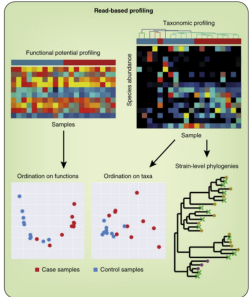
Cluster identification

```
net <- make_net(edge_table)</pre>
net_pack <- graph_from_incidence_matrix(net, mode = "total", weighted = TRUE)</pre>
#clustB <- cluster edge betweenness(net pack)</pre>
clustW <- cluster_walktrap(net_pack)</pre>
clustL <- cluster louvain(net pack)</pre>
clustG <- cluster_fast_greedy(net_pack)</pre>
clustLE <- cluster_leading_eigen(net_pack)</pre>
Modularity <- data.frame(Method = c("Random walks", "Louvain", "Greedy", "Non-negative eigenvector"),
                          Modularity = c(modularity(clustW), modularity(clustL),
                                           modularity(clustG),modularity(clustLE)))
Modularity
Modul_number_cutoff <- 3 # if the number of members in a cluster is lower than the cutoff, discard thi
edge_table <- assignModul(edge_table,clustW, "RandomWalks", Modul_number_cutoff)</pre>
edge_table <- assignModul(edge_table,clustL, "Louvain", Modul_number_cutoff)</pre>
edge_table <- assignModul(edge_table,clustG, "Greedy", Modul_number_cutoff)</pre>
edge_table <- assignModul(edge_table,clustLE, "Leading_eigen", Modul_number_cutoff)</pre>
write.csv(edge_table, "edge_table.csv", row.names = F,quote = F)
write.csv(node_table, "node_table.csv", row.names = F,quote = F)
```

Next, you can visualize with Gephi.

Guide to Shotgun Metagenomic Data





Read bases pipeline

Shotgun metagenomics involves sequencing all genetic material in a sample, generating vast amounts of short DNA fragments known as read bases. These raw reads undergo quality control, filtering, and taxonomic or functional annotation.

The Huttenhower Lab has developed a comprehensive pipeline called bioBakery Workflows, which includes:

• KneadData: Read quality control and host contamination removal

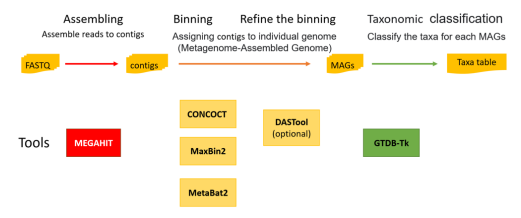
• **HUMAnN**: Functional profiling

• PhyloPhlAn: Phylogenetic analysis

Our lab also developed a user friendly bioBakery Workflows named BacNex

Assembly bases pipeline

Assembly bases, on the other hand, refer to contigs or scaffolds reconstructed from overlapping reads, providing longer genomic sequences for better functional and phylogenetic analysis.



			loU	Precision	Recall	number
MOCK HMP (human gut) Taxa number : 20	Reads-based Assembly-based	Concoct	0.208	0.556	0.250	9
		MaxBin2	0.520	0.722	0.650	18
		Metabat	0.520	0.722	0.650	18
		Concoct::DASTool	0.125	0.429	0.150	7
		MaxBin2::DASTool	0.480	0.706	0.600	17
		Metabat::DASTool	0.500	0.750	0.600	16
		humann3	0.692	0.750	0.900	24
		metaphlan4	0.571	0.667	0.800	24
	= 1					
	Rea		loU	Precision	Recall	number
	Rea	Concoct	IoU 0.294	Precision 0.606		number 33
	Rea	Concoct MaxBin2				
	Rea		0.294	0.606	0.364	33
моск	Rea	MaxBin2	0.294 0.294	0.606 0.619	0.364 0.473	33 42
Environment	Rea	MaxBin2 Metabat	0.294 0.294 0.357	0.606 0.619 0.625	0.364 0.473 0.455	33 42 40
	Rear	MaxBin2 Metabat Concoct::DASTool	0.294 0.294 0.357 0.308	0.606 0.619 0.625 0.667	0.364 0.473 0.455 0.364	33 42 40 30
Environment	Rea	MaxBin2 Metabat Concoct::DASTool MaxBin2::DASTool	0.294 0.294 0.357 0.308 0.328	0.606 0.619 0.625 0.667 0.700	0.364 0.473 0.455 0.364 0.382	33 42 40 30 30

While read-based analysis offers high-resolution species profiling, assembly-based approaches improve gene prediction and strain-level characterization. Choosing between them depends on study objectives, sample complexity, and computational resources.