# Pipeline

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## About.

This document contains a pipeline to go from raw Illumina MiSeq reads to a phyloseq object (along with some exploratory analysis) and is based on the workflow from the paper Characterising the bacterial gut microbiome of probiotic-supplmented very-preterm infants, which was based largely around this DADA2 workflow developed by *Callahan*, et al..

## Load required packages.

## Read quality.

Organise forward and reverse fastq filenames into own lists (check file format).

• First define the file path to the directory containing the fastq files (we will use this several times).

```
path <-"Data/new_data"

fnFs <- sort(list.files(path, pattern = "_R1_001.fastq.gz", full.names = TRUE))

fnRs <- sort(list.files(path, pattern = "_R2_001.fastq.gz", full.names = TRUE))</pre>
```

#### Extract sample names.

```
sample.names <- sapply(strsplit(basename(fnFs), "_"), `[`, 1)</pre>
```

Check quality of Forward and Reverse Reads (used to define truncLen in filtering).

## plotQualityProfile(fnFs[1:2])

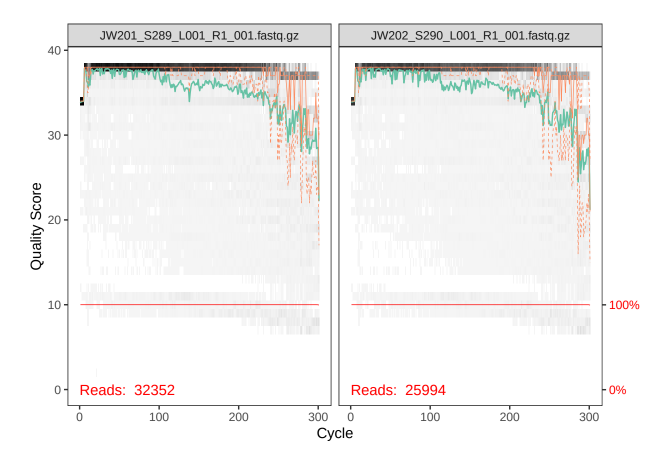


Figure 1: Quality of forward reads.

```
plotQualityProfile(fnRs[1:2])
```

## Assign names for filtered reads.

```
filtFs <- file.path(path, "filtered", paste0(sample.names, "_F_filt.fastq.gz"))
filtRs <- file.path(path, "filtered", paste0(sample.names, "_R_filt.fastq.gz"))</pre>
```

#### Filter and trim the reads.

- Paremeters based on data and quality plots.
- truncLean defined by when quality plots begin to drop off, but ensuring it is large enough to maintain read overlap (=>20bp) downstream.
- trimLeft is not needed as primers/barcodes already removed.

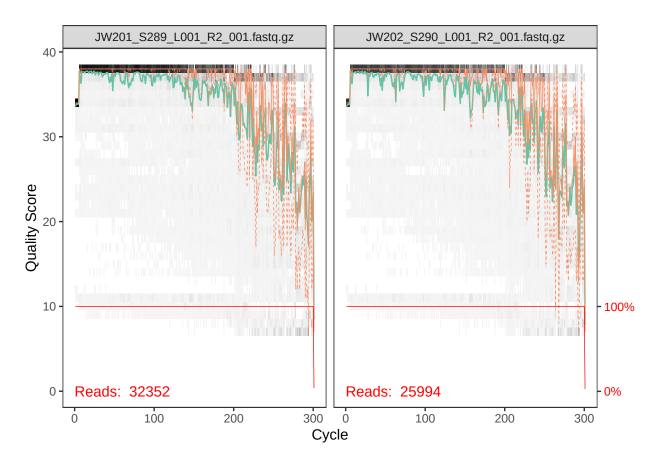


Figure 2: Quality of reverse reads.

- maxEE = c(2,2) is for filtering, where the higher the value the more relaxed filtering, allowing more reads to get through.
- Good quality data should allow for more stringent parameters (2 is stringent).
- The number of reads filtered is checked. If reads are too low, can alter parameters.

## Infer sequence variants.

#### Calculate Error Rates.

• Error rates are used for sample ineference downstream.

```
errF <- learnErrors(filtFs, multithread = TRUE)
errR <- learnErrors(filtRs, multithread = TRUE)</pre>
```

#### Plot error rates.

• Estimated error rates (black line) should be a good fit to observed rates (points) and error should decrease.

```
plotErrors(errF, nominalQ = TRUE)
plotErrors(errR, nominalQ = TRUE)
```

#### Dereplication.

- Combine indentical sequences into unique sequence bins.
- Name the derep-class objects by the sample name.

```
derepFs <- derepFastq(filtFs, verbose = TRUE)

derepRs <- derepFastq(filtRs, verbose = TRUE)

names(derepFs) <- sample.names

names(derepRs) <- sample.names</pre>
```

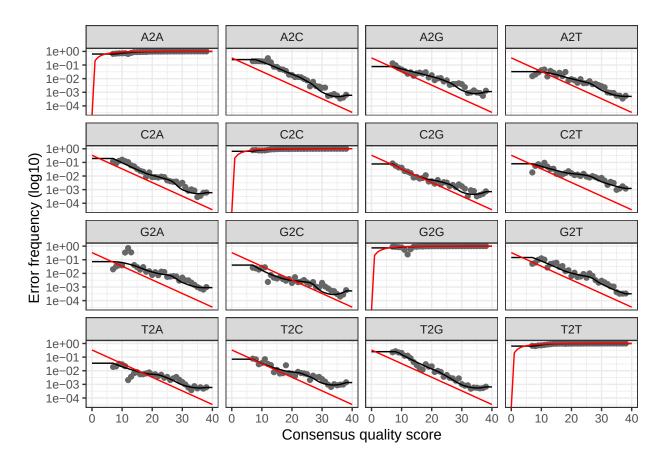


Figure 3: Error rates for forward reads

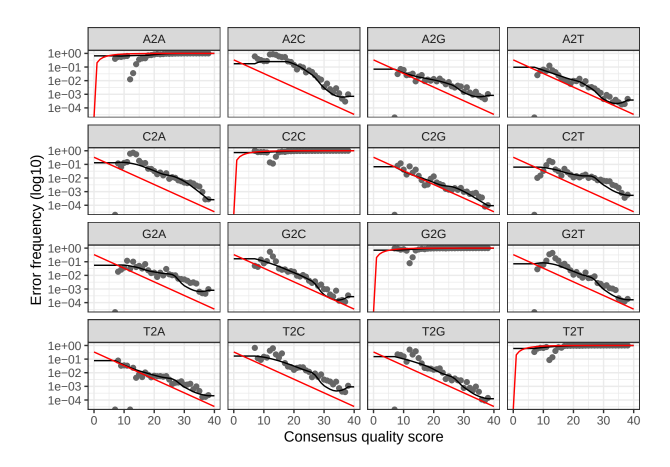


Figure 4: Error rates for reverse reads.

Sequence Inference.

```
dadaFs <- dada(derepFs, err = errF, multithread = F)
dadaRs <- dada(derepRs, err = errR, multithread = F)</pre>
```

Inspect denoised data.

```
dadaFs[[1]]
dadaRs[[1]]
```

Merge Paired Reads and inspect merged data.

- Removes paired reads that do not perfectly overlap.
- Arguments represent infered samples AND denoised reads.

```
mergers <- mergePairs(dadaFs, derepFs, dadaRs, derepRs, verbose = TRUE)</pre>
```

Construct amplicon sequence variance (ASV) table and remove chimeras.

Construct ASV table.

• Check dimentions and inspect distribution of sequence lengths.

```
seqtab <- makeSequenceTable(mergers)

dim(seqtab)

seqtab %>%
  getSequences() %>%
  nchar() %>%
  table()
```

Remove chimeras.

Track reads through pipeline.

```
getN <- function(x) sum(getUniques(x))
track <- cbind(out, sapply(dadaFs, getN), sapply(dadaRs, getN),</pre>
```

```
sapply(mergers, getN), rowSums(seqtab.nochim))
colnames(track) <- c("input", "filtered", "denoisedF", "denoisedR", "merged", "nonchim")
rownames(track) <- sample.names
head(track)</pre>
```

```
##
          input filtered denoisedF denoisedR merged nonchim
## JW201 32352
                  27583
                            27342
                                      27406 26036
                                                      14276
## JW202 25994
                   21958
                             21868
                                      21908 21762
                                                      20325
## JW203
            31
                      6
                                1
                                          3
                                                 0
                                                         0
## JW203b 30571
                  25513
                            24856
                                      25197 22249
                                                      7368
## JW204 25523
                  21198
                            20751
                                      21100 18947
                                                      10893
## JW205 31796
                  26653
                            25969
                                      26433
                                             23889
                                                      11348
```

## Contamination removal with MicroDecon.

```
library(microDecon)
```

## Read in metadata (needed for MicroDecon)

```
Metadata <- readxl::read_excel("Data/New_metadata.xlsx")</pre>
```

## Reformat data for MicroDecon.

- Function is data specific.
- Transpose sequencing table (post chimera removal) and convert to a dataframe.
- Reorder sequencing table by a prior grouping (days).
- Move blank sample columns to the start of the sequencing table.
- Turn row names into their own column as *MicroDecon* requires that the OTUs have a unique ID in column 1.

```
wrangle_microdecon <- function(seqtab.nochim){

# transpose data
microdecon.df <- t(seqtab.nochim) %>%
    as.data.frame()

microdecon.df <- microdecon.df %>%
    relocate("JW219B", "JW220") %>%
    tibble::rownames_to_column(var = "ID") # turn the rownames into the first column
}

microdecon.df <- wrangle_microdecon(seqtab.nochim)</pre>
```

#### Decontaminate data using decon().

- numb.ind is the number of columns for each priori grouping.
- taxa = F as there is no taxonomy in the dataframe.

```
decontaminated$decon.table
decontaminated$reads.removed
decontaminated$0TUs.removed
decontaminated$mean.per.group
decontaminated$sum.per.group
```

#### Check MicroDecon Outputs.

#### Reformat decon.table.

- Convert column 1 to row names.
- Remove blank average column (1).
- Save rownames as seperate vector to be added back, as row names are removed during apply().
- Convert numeric values to integers (for downstream analysis).
- Transpose data.

```
seqtab.microdecon <- decontaminated$decon.table %>%
  remove_rownames() %>%
  column_to_rownames(var = "ID") %>%
  select(-1) %>% # remove mean blank
  as.matrix() %>%
  t()
```

## Remove non-amplified samples and rename all.

```
seqtab.microdecon <- seqtab.microdecon %>%
  as.data.frame() %>%
  rownames_to_column("ID") %>%
  filter(ID != c("JW203", "JW211")) %>%
  mutate(ID = str_remove(ID, "b")) %>%
  mutate(ID = str_remove(ID, "JW")) %>%
  column_to_rownames("ID") %>%
  as.matrix()
```

#### Merging multiple sequence runs.

```
# Reading in pilot data for rmd.
seqtab.combined <- read_csv("Data/seqtab_original.csv") %>%
  column_to_rownames("ID") %>%
  as.matrix() %>%
  mergeSequenceTables(seqtab.microdecon)
```

## Assign taxonomy.

• With optional species addition (there is an agglomeration step downstream, so you can add species now for curiosities sake, and remove later for analysis).

```
taxa <- assignTaxonomy(seqtab.microdecon, "Data/silva_nr_v132_train_set.fa.gz")
taxa.print <- taxa # Removes sequence rownames for display only
rownames(taxa.print) <- NULL</pre>
```

#### Calculate percentage of NA taxa

```
sum(is.na(taxa))/prod(dim(taxa)) * 100

## [1] 19.73255

apply(taxa, 2, function(col)sum(is.na(col))/length(col)) * 100

## Kingdom Phylum Class Order Family Genus Species
## 0.000000 1.369863 2.054795 4.566210 9.589041 32.876712 87.671233
```

## Preprocessing: Creating a Phyloseq Object.

#### About.

Creating a phyloseq object to be used for analysis, and create different objects to be used for different types of analysis downstream.

## Load required packages.

## Import metadata.

```
Metadata <- readxl::read_excel("Data/New_metadata.xlsx") %>%
  select(-c(3, 18, 21)) %>%
  add_column("Primary_Group" = "SCN", "Type" = "Discharge") %>%
  rbind(
    readxl::read_excel("Data/Old_metadata.xlsx") %>%
```

```
separate(DOB, into = c("DOB", "Time"), sep = "\\s") %>%
    mutate(DOB = as.Date(DOB)) %>%
    select(1, 3:4, 9, 17:18, 20:35)) %>%
add_row(URN = "ZymoDNA1", ID = "ZymoDNA1", Type = "Control") %>%
add_row(URN = "ZymoDNA2", ID = "ZymoDNA2", Type = "Control") %>%
add_row(URN = "ZymoDNA4", ID = "ZymoDNA4", Type = "Control") %>%
mutate(Sample_ID = ID) %>%
column_to_rownames("Sample_ID") %>%
mutate(Mode_of_Delivery = str_replace(Mode_of_Delivery, "Ceaserean", "Cesarean")) %>%
mutate(Batch = 1:nrow(.)) %>%
mutate(Batch = if_else(Batch <= 20, "Run2", "Run1"))</pre>
```

## Construct the Phyloseq object.

• Includes: metadata, ASV table, taxonomy table and phylogenetic tree.

## Wrangling the metadata.

- And do some additional wrangling.
- Convert chraracters to factors.

```
sample_data(ps) <- sample_data(ps) %>%
  unclass() %>%
  as.data.frame() %>%
  mutate_if(is.character, as.factor) %>%
  mutate("Sample" = ID) %>% # needed to save it back into the original ps object
  column_to_rownames("Sample")
```

## Getting read counts

```
sample_data(ps) %>%
unclass() %>%
as.data.frame() %>%
mutate(TotalReads = sample_sums(ps)) %>%
ggplot(aes(TotalReads)) +
   geom_histogram() +
   ggtitle("Sequencing Depth")
```

## Filtering and normalisation.

#### Taxonomy filtering.

• Can check the number of phyla before and after transformation with table(tax\_table(ps)[, "Phylum"], exclude = NULL).

• Remove features with ambiguous and NA phylum annotation.

```
ps1 <- subset_taxa(ps, !is.na(Phylum) & !Phylum %in% c("", "uncharacterized"))
```

#### Remove problematic samples

• samples that produce NA values during transformations downstream in the otu\_table (for some reason it won't allow all within the same function, hence the pipe)

```
ps1 <- subset_samples(ps1, ID != "219") %>%
  subset_samples(ID != "141") %>%
  subset_samples(ID != "118")
```

```
# Total
sum(is.na(tax_table(ps1)))/prod(dim(tax_table(ps1))) * 100
```

#### Check percentages of NA values left.

```
## [1] 18.81614
```

```
# Per taxonomic rank
apply(tax_table(ps1), 2, function(col)sum(is.na(col))/length(col)) * 100

## Kingdom Phylum Class Order Family Genus Species
## 0.0000000 0.0000000 0.6944444 3.2407407 8.3333333 31.9444444 87.5000000
```

#### Prevelance filtering.

- Using an unsupervised method (relying on the data in this experiment) explore the prevelance of features in the dataset.
- Calculate the prevalence of each feature and store as a dataframe.
- Add taxonomy and total read counts.

• Plot the relationship between prevelance and total read count for each feature. This provides information on outliers and ranges of features.

```
prevdf %>%
  subset(Phylum %in% get_taxa_unique(ps1, "Phylum")) %>%
  ggplot(aes(TotalAbundance, Prevalence / nsamples(ps1),color=Phylum)) +
  geom_hline(yintercept = 0.05, alpha = 0.5, linetype = 1) +
  geom_point(size = 2, alpha = 0.7) +
  scale_x_log10() +
  xlab("Total Abundance") + ylab("Prevalence [Frac. Samples]") +
  facet_wrap(~Phylum) + theme(legend.position="none")
```

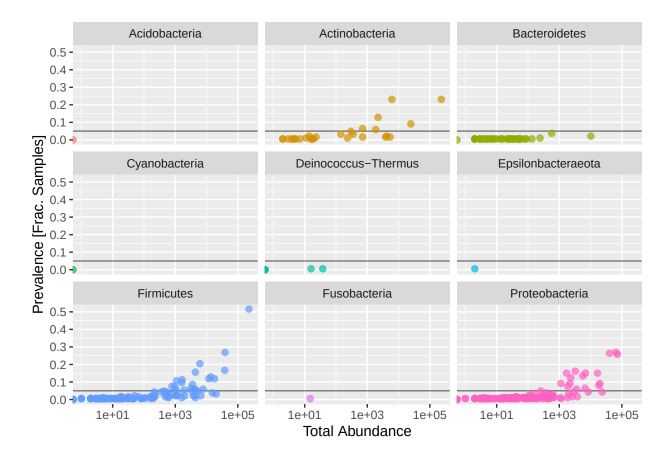


Figure 5: Scatterplot exploring the relationship between prevelance and abundance of phyla.

• Define prevalence threshold based on the plot (~1% is standard) and apply to ps object (if prevelance is too low don't designate a threshold).

```
prevalenceThreshold = 0.01 * nsamples(ps1)
keepTaxa = rownames(prevdf)[(prevdf$Prevalence >= prevalenceThreshold)]
ps2 = prune_taxa(keepTaxa, ps1)
```

Subset phyloseq object for data to be analyzed.

```
ps2 <- subset_samples(ps2, Type == "Discharge")
```

• Explore the relationship on the filtered data set.

```
prevdf %>%
  subset(Phylum %in% get_taxa_unique(ps2, "Phylum")) %>%
  ggplot(aes(TotalAbundance, Prevalence / nsamples(ps2),color=Phylum)) +
  geom_hline(yintercept = 0.05, alpha = 0.5, linetype = 1) +
  geom_point(size = 2, alpha = 0.7) +
  scale_x_log10() +
  xlab("Total Abundance") + ylab("Prevalence [Frac. Samples]") +
  facet_wrap(~Phylum) + theme(legend.position="none")
```

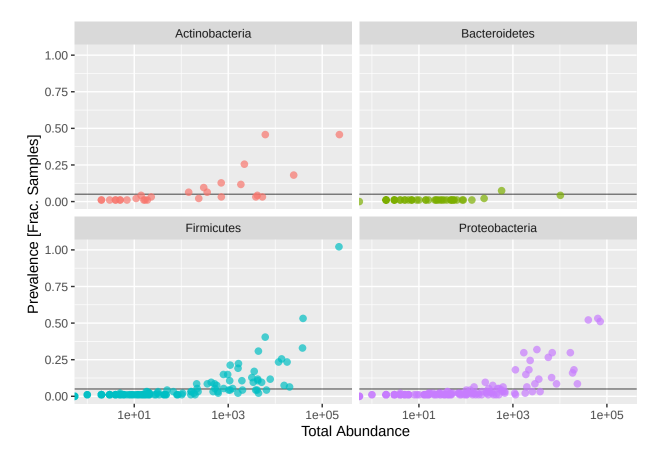


Figure 6: Scatterplot exploring the relationship between prevelance and abundance of phyla on data passed through a prevalence threshold.

#### Aggolmerate taxa.

• Combine features that descend from the same genus as most species have not been identified due to the poor taxonomic depth in 16S, a result of the length of the fragment amplified from the 16SrRNA

gene.

• Can check how many genera would be present after filtering by running length(get\_taxa\_unique(ps2, taxonomic.rank = "Genus")) and/or ntaxa(ps3) will give the number of post agglomeration taxa.

```
ps3 = tax_glom(ps2, "Genus", NArm = TRUE)
```

#### Normalisation.

- Plot a refraction curve to see if total sum scaling will surfice.
- Define colours and lines.
- Step = step size for sample sizes in rarefaction curve.

• Perform total sum scaling on agglomerated dataset.

```
ps4 <- transform_sample_counts(ps3, function(x) x / sum(x))
```

- Explore normalisation with violin plots.
- Compares differences in scale and distribution of the abundance values before and after transformation.
- Using arbitrary subset, based on Phylum = Firmicutes, for plotting (ie. can explore any taxa to observe transformation).

```
plot_abundance = function(physeq, Title = "Abundance",
                          Facet = "Order", Color = "Phylum", variable = "Type"){
    subset_taxa(physeq, Phylum %in% c("Firmicutes")) %>%
   psmelt() %>%
    subset(Abundance > 0) %>%
    ggplot(mapping = aes_string(x = variable, y = "Abundance", color = Color, fill = Color)) +
      geom_violin(fill = NA) +
      geom_point(size = 1, alpha = 0.3, position = position_jitter(width = 0.3)) +
     facet_wrap(facets = Facet) +
      scale_y_log10()+
      theme(legend.position="none") +
      labs(title = Title)
}
grid.arrange(nrow = 2, (plot_abundance(ps3, Title = "Abundance",
                          Color = "Type", variable = "Type")),
                        plot_abundance(ps4, Title = "Relative Abundance",
                          Color = "Type", variable = "Type"))
```

## Data exploration and univariate analysis.

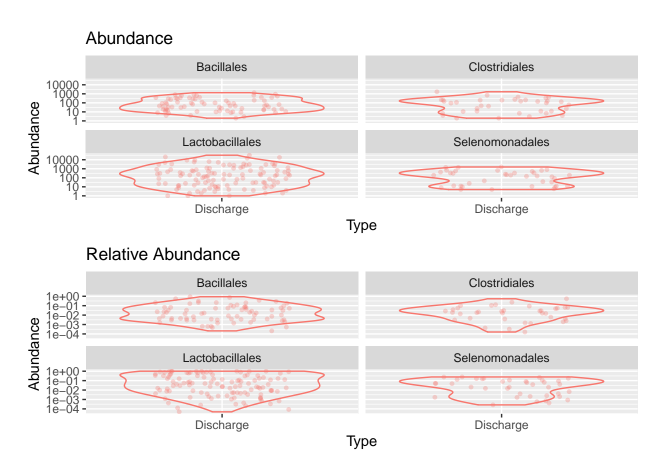


Figure 7: Violin plots exploring of distribution of abundance in Firmicutes before and after normalisation of data.

## About.

This section again uses the phyloseq package (along with several others) to explore the data using bar, violin and ordination plot. This then leads into a collection of univariate analyses, including; alpha and beta diversity, and also taxonomic differential abundance.

## Load required packages.

## Taxanomic distribution.

#### Bar charts

- Use plot\_bar\_auto() function wrapped around phyloseq's plot\_bar() to explore the distribution of taxa at the genus and phylum levels.
- Subset transformed data (relative abundance) to only the top20 taxa.

```
top20 <- names(sort(taxa_sums(ps4), decreasing=TRUE))[1:20]
ps.top20 <- prune_taxa(top20, ps4)

plot_bar_auto <- function(ps, taxonomy){
  plot_bar(ps, fill = taxonomy) +
     facet_wrap(~Primary_Group, scales = "free_x") +
     labs(title = paste0("Level:", taxonomy), y = "Abundance") +
     theme(legend.position = "bottom", legend.title = element_blank(),
     axis.title.x = element_blank(), axis.text.x = element_blank(),
     axis.ticks = element_blank())
}

plot_bar_auto(ps.top20, "Genus")</pre>
```

## Investigate specific taxa

```
ps4 %>%
  subset_taxa(Genus == "Bifidobacterium") %>%
  plot_bar_auto(., "Genus")
```

Calculate the number samples containing a given taxa by creating a samples\_with\_taxa() function.

• Define a function takes the phyloseq object, taxonomy level and taxanomic name (with the later two as strings).

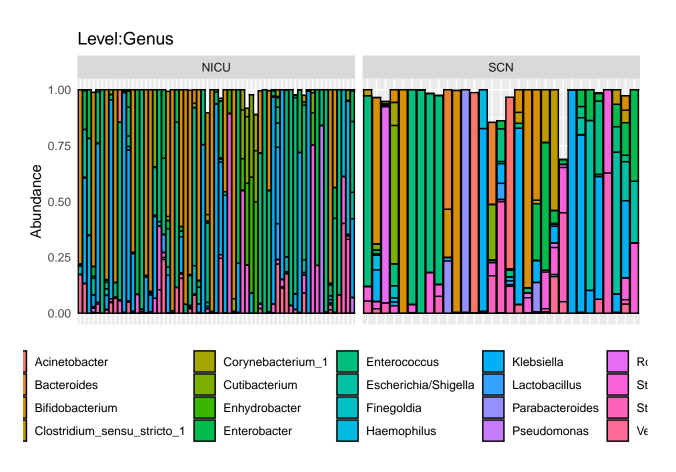


Figure 8: Bar plots of the taxonomic distribution (relative abundance) at the genus levels.

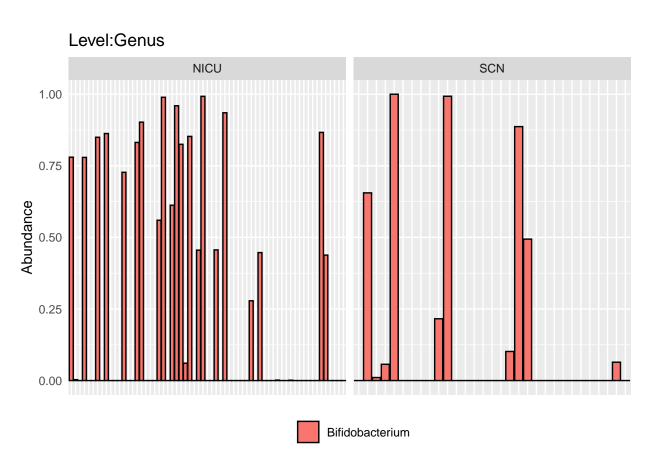


Figure 9: Bar plot of the taxonomic distribution of Bifidobacterium.

- It then gets the ASV name from the *phyloseq* tax\_table() by filtering with *dply* and *lazyeval*. ( *lazyeval* is needed because of two concepts;non-standard evaluation and lazy evaluation.
- paste() is then used to concatenate the ASVs and collapse to insert the 'or' symbol.
- The function then matches the ASV names to the otu\_table() of the *phyloseq* object to select the desired column(s) that represent the taxa of interest, and then counts the number of rows that have any of the selected taxa with counts greater than 0 to get the number of samples with that taxa present.

```
samples_with_taxa <- function(ps_object, taxonomy_level, taxa){
    ASV <- tax_table(ps_object) %>%
        unclass() %>%
        as.data.frame() %>%
        filter_(interp(~y == x, .values=list(y = as.name(taxonomy_level), x = taxa))) %>%
        row.names() %>%
        paste(collapse = " | ")

    otu_table(ps_object) %>%
        as.data.frame() %>%
        select(matches(ASV)) %>%
        filter_all(any_vars( . > 0)) %>%
        nrow()
}

ps4 %>%
    subset_samples(Primary_Group == "NICU") %>%
samples_with_taxa(., "Genus", "Bifidobacterium")
```

## Beta diversity

- Use distance and ordination methods to explore the relationship between metadata.
- We calculate the distances using pruned, transformed (TSS) and non-agglomerated data.

```
ps2.TSS <- ps2 %>%
    transform_sample_counts(function(x) x / sum(x))
```

- We can then create distance matrices and plots for this data subset using several methods:
- e.g. bray-curtis or weighted unifrac distances with PCoA, NMDS, etc.
- Define a function that ordindates the previously transformed data, extarcts the eigenvalues, and creates a dissimilarity plot.
- Extract eigenvalues from ordination.

```
ordination_plots <- function(filtered_ps, variable, vis_method, dist_method){
# ordinate
ps_ordination <- ordinate(filtered_ps, method = vis_method, distance = dist_method)
# get eignenvalues
evals <- ps_ordination$values$Eigenvalues
# generate plot
plot_ordination(filtered_ps, ps_ordination, color = variable,
    title = "PCoA (Bray-Curtis)") +
    labs(col = variable) +
    coord_fixed(sqrt(evals[2] / evals[1])) +
    geom_point(size = 5) +
    stat_ellipse(typre = "norm", linetype = 2) +</pre>
```

```
scale_color_hue(labels = c("Probiotic-treated", "Non-treated"))
}
ordination_plots(ps2.TSS, "Primary_Group", "PCoA", "bray")
```

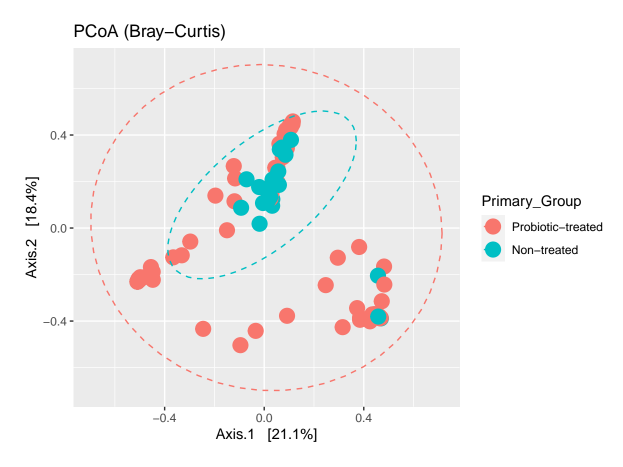


Figure 10: PCoA plot of Bray-Curtis distances coloured by date.

#### Statistical test: PERMANOVA.

- ps2 transformed.
- Preforming permutational anova for group-level differences based on dissimilarity.
- Extract of otu table and metadata from phyloseq object.
- Use adonis() from the *vegan* package to perform the PERMANOVA.
- Homogeneity Condition
- Significant PERMANOVA means one of three things:
- there is a difference in the location of the samples (i.e. the average community composition).
- there is a difference in the dispersion of the samples (i.e. the variability in the community composition).
- there is a difference in both the location and the dispersion.
- If you get a significant PERMANOVA you'll want to distinguish between the three options by checking the homogeneity condition using permdisp(). If you get a non-significant result the first option above is correct.
- betadisper() gives a measure of the dispersion within groups. Thus, if the PERMANOVA test is significant and the permdisp is not, the significant result in your communities is due to a mean shift in community composition and not from increased variance within groups.

```
permanova_func <- function(ps2.TSS){</pre>
# permanova
ps_otu <- data.frame(otu_table(ps2.TSS))</pre>
ps_metadata <- data.frame(sample_data(ps2.TSS))</pre>
permanova <- adonis(ps_otu ~Primary_Group, data = ps_metadata, method = "bray")</pre>
permanova <- tableGrob(as.data.frame(permanova$aov.tab)) %>%
  annotate_figure(fig.lab = "PERMANOVA", fig.lab.face = "bold", fig.lab.size = 15)
# homogeneity condition
dist <- vegdist(ps_otu)</pre>
homogeneity <- as.data.frame(anova(betadisper(dist, ps_metadata$Primary_Group))) %%
  tableGrob() %>%
  annotate_figure(fig.lab = "Homogeneity Condition", fig.lab.face = "bold", fig.lab.size = 15)
# combine ouputs in a grid
grid.arrange(permanova, homogeneity, ncol = 1)
}
permanova_func(ps2.TSS)
```

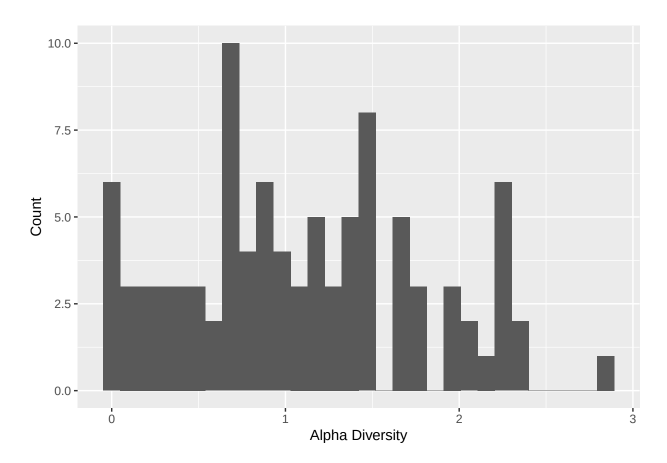
• Explore the major contributors to the differences.

```
major_contributors <- function(ps2.TSS, variable){</pre>
# perform permanova
ps_otu <- data.frame(otu_table(ps2.TSS))</pre>
ps_metadata <- data.frame(sample_data(ps2.TSS))</pre>
permanova <- adonis(ps_otu ~Primary_Group, data = ps_metadata, method = "bray")</pre>
# coefficients
coef <- coefficients(permanova)[paste0(variable, "1"),]</pre>
top.coef <- coef[rev(order(abs(coef)))[1:20]]</pre>
genus_contributors <- tax_table(ps2.TSS) %>%
    unclass() %>%
    as.data.frame() %>%
    select("Genus") %>%
    rownames_to_column(var = "ASV") %>%
    right_join((as.data.frame(top.coef) %>%
    rownames_to_column(var = "ASV"))) %>%
    select(!"ASV")
return(genus_contributors)
major_contributors(ps2.TSS, "Primary_Group")
```

## Alpha diversity.

• Define a function that calculates Shannon Index, Obsverved (richness) & Chao1 diversity, and binds it to our original metadata dataframe, which can then be used for analysis.

• Create histogram to examine distribution.



• Test for normality.

#### shapiro.test(ps\_metadata\$Shannon)

```
##
## Shapiro-Wilk normality test
##
## data: ps_metadata$Shannon
## W = 0.97086, p-value = 0.03382
```

#### Statistical test: compare mean/median between groups.

• define a function that performs a Wilcoxin test on the three diversity metrics and binds them (Shannon Index, Richness & Chao1).

```
## # A tibble: 3 x 8
##
     'Diversity Measure' group1 group2
                                           p p.adj p.format p.signif method
##
     <chr>
                         <chr> <chr> <dbl> <dbl> <chr>
                                                             <chr>>
                                                                      <chr>>
## 1 Shannon
                                NICU
                                       0.278 0.28 0.28
                         SCN
                                                                      Wilcoxon
                                                             ns
## 2 Observed
                         SCN
                                NICU
                                       0.171 0.17 0.17
                                                                      Wilcoxon
                                                             ns
## 3 Chao1
                                       0.171 0.17 0.17
                         SCN
                                NICU
                                                             ns
                                                                      Wilcoxon
```

#### Plot alpha diversity.

- Use plot\_richness() from *phyloseq*, which estimates alpha diversity metrics using *vegan* and plots them, taking standard *ggplot2 geoms*\_ for the plot design.
- use ps2 non-transformed data for alpha.

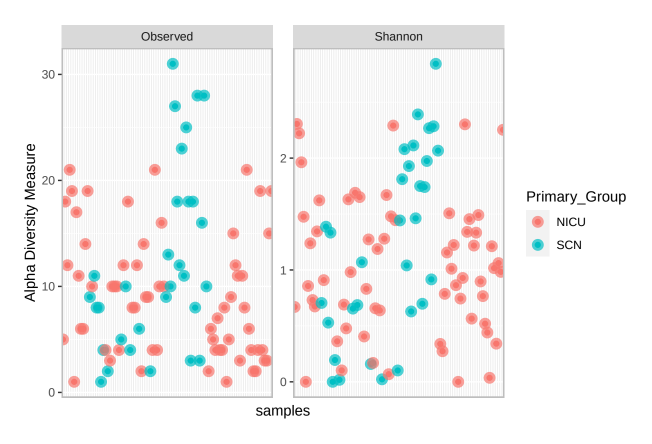
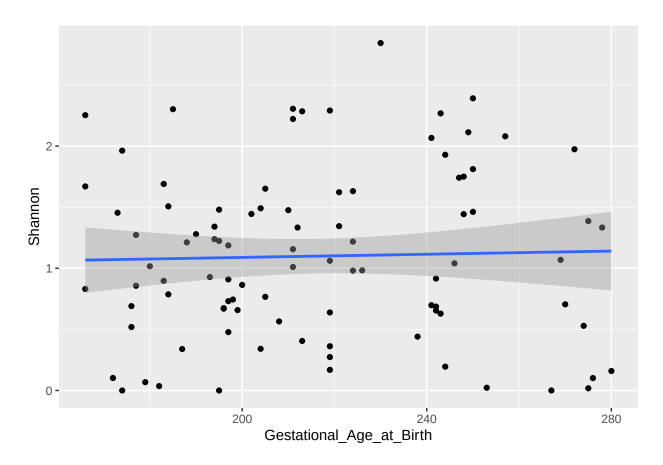


Figure 11: Scatterplot of richness and shannon diversity metrics coloured by ddPCR.

• shannon diversity for continuous variable.

```
ps_metadata %>%
ggplot(aes(x = Gestational_Age_at_Birth, y = Shannon)) +
  geom_point() +
  geom_smooth(method = "lm", se = T)
```

```
## 'geom_smooth()' using formula 'y ~ x'
```



## Differential abundance analysis with DESeq2.

- Define function for calculating geometric means and estimating size factors.
- Define function to filter out taxa with small counts and low occurance. *count* and *samples* arguments need to be applied as numerical values.

```
calc_geo_means <- function(deseq_object){
# geometric mean
gm_mean = function(x, na.rm = TRUE){
    exp(sum(log(x[x > 0]), na.rm = na.rm) / length(x))
}
geoMeans <- apply(counts(deseq_object), 1, gm_mean)
# size factors
estimateSizeFactors(deseq_object, geoMeans = geoMeans)</pre>
```

```
deseq_filter <- function(deseq_object, count, samples){
  nc <- counts(deseq_object, normalized = TRUE)
  filtered <- rowSums(nc >= count) >= samples # filter = abundance of 10 in 60 samples.
  deseq_object[filtered,]
}
```

- Define a function to extract the results.
- Extract the results, order by p value, selects significant (<0.05) results, binds this data to the *tax\_table* from the *phyloseq* object to get the taxonomic information, and then select and order the desired columns.

```
# function for liklihood ratio test
get_deseq_res_lrt <- function(deseq_object){</pre>
  res = results(deseq_object)
  res = res[order(res$padj, na.last = NA), ]
  sigtab = res[(res$padj < 0.1), ]</pre>
  sigtab = cbind(as(sigtab, "data.frame"),
          as(tax_table(ps3)[rownames(sigtab), ], "matrix"))
  sigtab %>%
  arrange(padj) %>%
  select("log2FoldChange", "lfcSE", "padj", "Genus")
}
# function for Walds test and continuous variables
get_deseq_res_cont <- function(deseq_object, contrast_variable){</pre>
  res = results(deseq_object, name = contrast_variable)
  res = res[order(res$padj, na.last = NA), ]
  sigtab = res[(res$padj < 0.1), ]
  sigtab = cbind(as(sigtab, "data.frame"),
          as(tax_table(ps3)[rownames(sigtab), ], "matrix"))
  sigtab %>%
  arrange(padj) %>%
  select("log2FoldChange", "lfcSE", "padj", "Genus")
}
# function for Walds test and categorical variables
get_deseq_res_cat <- function(desq_object, contrast_variable, level1, level2){</pre>
  res = results(desq_object, contrast = c(contrast_variable, level1, level2))
  res = res[order(res$padj, na.last = NA), ]
  sigtab = res[(res$padj < 0.1), ]</pre>
  sigtab = cbind(as(sigtab, "data.frame"),
    as(tax_table(ps3)[rownames(sigtab), ], "matrix"))
  sigtab %>%
  arrange(padj) %>%
  select("log2FoldChange", "lfcSE", "padj", "Genus") %>%
  add_column(Variable = paste0(contrast_variable, level1)) # label the base level
```

- Convert from *phyloseq* to *deseq* object.
- Calculate geometric means and filter to the most abundant and frequent taxa.
- Use Deseq() to perform the normalisation and analysis.

• Extract the results.

```
# LRT
phyloseq_to_deseq2(ps3, ~ Primary_Group) %>%
    calc_geo_means() %>%
    deseq_filter(10,30) %>%
    DESeq(fitType = "local", test = "LRT", reduced = ~ 1) %>%
    get_deseq_res_lrt() %>%
    remove_rownames()
```

```
# Wald
phyloseq_to_deseq2(ps3, ~ Primary_Group) %>%
    calc_geo_means() %>%
    deseq_filter(10,30) %>%
    DESeq(fitType = "local", test = "Wald") %>%
    get_deseq_res_cat("Primary_Group", "NICU", "SCN") %>%
    remove_rownames()
```

```
## log2FoldChange lfcSE padj Genus Variable ## 1 3.913908 1.501674 0.06405668 Bifidobacterium Primary_GroupNICU
```