# Metagenome Analysis of Premature Birth

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## Overview

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- Methods
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Introduction

#### Microbiome

- Microbiota: the microorganisms which live inside & on humans (Turnbaugh et al., 2007)
- Microbiome:  $10^{13}$  to  $10^{14}$  microorganisms whose which collective genome (Gill et al., 2006)



Figure: Concept of a core human microbiome (Turnbaugh et al., 2007)

#### rRNA

- Ribosomal RNA
- Well-known as a key to phylogeny (Olsen & Woese, 1993)

# Premature Birth (Preterm Birth; PTB)

#### PTB:

- PTB < 37 GW (Gestational week)</p>
- Normal ≥ 37 GW

#### Detailed PTB:

- Extremely PTB < 28 GW
- 2 28 GW  $\leq$  Very PTB < 32 GW
- $32 \text{ GW} \leq \text{Late PTB} < 37 \text{ GW}$
- Normal > 37 GW
- (J. Tucker & McGuire, 2004; Voronkov, Solonovych, Liashenko, & Revenko, 2018)

## Materials

# 16S rRNA Sequencing

**16S rRNA sequencing** is the *reference method* for bacterial taxonomy & identification (Mignard & Flandrois, 2006)
Three main reasons (Janda & Abbott, 2007):

- 16S rRNA exists in almost all bacteria
- Functions of the 16S rRNA has not changed over time
- 16S rRNA is large enough for bioinformatics

# Train/Test Data vs. Validate Data

- JBNU/Helixco data
  - First data
  - Second data
  - Stool data

Table: Sample Information

	Data	Participants	Samples	Remarks
_	First	24	107	-
	Second	35	288	-
	Stool	63	126	Stool

## Methods

## Methods

Qiime 2 Workflow

# Qiime 2 Workflow

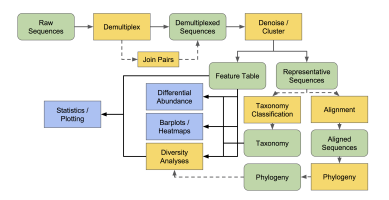


Figure: QIIME 2 workflow (Bolyen et al., 2019; Mandal, Van Treuren, White, Eggesbø, et al., 2015; McDonald et al., 2012)

# Filitering with Quality Score

#### Drawback between:

- Longer sequence read
- Higher quality value
- $\therefore$  Select the maximum length n where:

$$\forall n_i \in \{n_k | \text{MedianQualityScore} \ge 30\}$$
  
$$\exists! n \in \{n_i\} : n \ge n_i$$
 (1)

# **Denoising Techniques**

- DADA2: Amplicon Sequence Variants (ASVs) (Callahan et al., 2016)
- Deblur: Operational Taxonomic Units (OTUs) (Amir et al., 2017)



Figure: Denoising Algorithms

# Taxonomy Classification

- Greengenes (GG) (DeSantis et al., 2006)
- SILVA (Pruesse et al., 2007; Quast et al., 2012)



Figure: Taxonomy Classifications

"A **higher** performance at taxonomic levels above *genus level*; but performance appears to **drop** at *species level*" (Gihawi et al., 2019)

# Merging Denoising/Taxonomy

Merging multiple IDs (ASVs or OTUs) into one, which have

- Different IDs
- Identified as same taxonomy

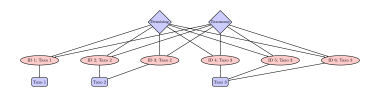


Figure: Example Diagram for Merging Denoising/Taxonomy

## Methods

Abundance Test

## **ANCOM**

- Analysis of composition of microbiome (Mandal, Van Treuren, White, Eggesbø, et al., 2015)
- ANCOM detects significantly abundant taxa, while maintain high statistical power
- Find taxa that can divide each classes

## Methods

**Diversity Indices** 

# **Diversity Indices**

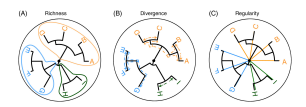


Figure: Three dimensions of phylogenic information (C. M. Tucker et al., 2017)

- A quantitative measure that shows richness, divergence, and regularity (C. M. Tucker et al., 2017)
- Alpha diversity indices: the richness of taxa at a single community
- Beta diversity indices: the taxonomic differentiation between communities

# Alpha Diversity Indices

- Evenness index
- Faith's Phylogenetic Diversity (Faith PD) index
- Oberseved Features index
- Shannon's Diversity index

# Beta Diversity Indices

- Bray-Curtis distance index
- Jaccard distnace index
- Unweighted UniFrac distance index
- Weighted UniFrac distance index

## Methods

Miscellaneous

# t-distributed Stochastic Neighbor Embedding (t-SNE)

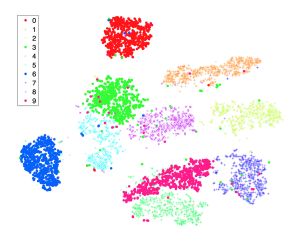


Figure: t-SNE with handwritten data (Maaten & Hinton, 2008)

# Python Packages

- Pandas (McKinney et al., 2011)
- Scikit-Learn (Pedregosa et al., 2011)
- SciPy (Virtanen et al., 2020)
- Matplotlib (Hunter, 2007)
- Seaborn (Waskom et al., 2020)
- Statannot

## Results

## Results

Filtering Results

# Quality Score from JBNU/Helixco Data



Figure: Quality Score Plot

- $\ell_{Forward} = 300$
- $\ell_{Reverse} = 245$

## Results

t-SNE with Clinical Information

# Workflow for t-SNE with Site/Premature Information

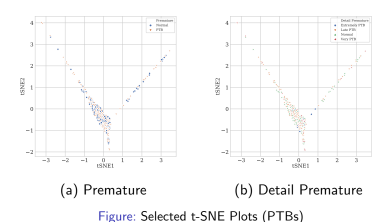


Figure: Workflow of t-SNE for Site/Premature Information

### **Used Clinical Information**

- Apgar score
- 2 C-section
- OPAP (Continuous positive airway pressure)
- Oetail premature & Premature
- Operation of the second of
- Gender
- Gestational diabetes for maternity
- Ostational week (Gestational age)
- Mospitalized day
- Hypertension for maternity
- LGA (Large for GA) & SGA (Small for GA)
- Maternal age
- Antibiotic usage for maternity & for neonate
- PROM (Premature rupture of membranes)
- Preterm labor
- Usage of respirator for neonate
- Sepsis against neonate
- Birth weight of neonate

## Selected t-SNE Plots I



## Selected t-SNE Plots II

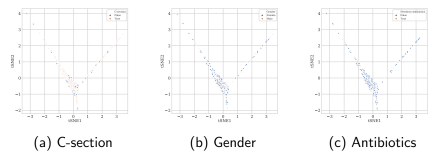


Figure: Selected t-SNE Plots (Not severe condition)

### Selected t-SNE Plots III

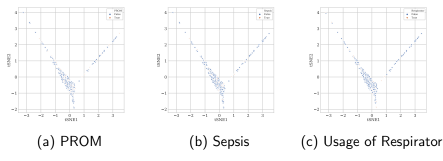


Figure: Selected t-SNE Plots (Severe conditions)

## Selected t-SNE Plots IV

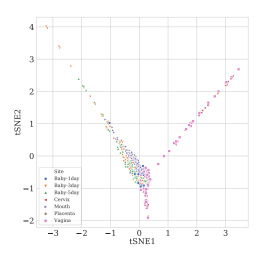


Figure: t-SNE Plot with Site

## Results

Bacterial Abundance Test with ANCOM

# ANCOM?

- Analysis of composition of microbiomes
- ANCOM can be used for analyzing the composition of microbiomes in multiple populations (Mandal, Van Treuren, White, Eggesbø, et al., 2015)
- Differential abundance testing
- clr: Centered log(Ratio)
- W: a count of the number of sub-hypothesis which have passed for given species

### ANCOM with ...

#### Site selection:

- 1 Neonatal mouth: 1-day, 3-day, and 5-day
- Cervix
- Maternal mouth
- Vagina

#### PTB:

- Premature
- Oetail premature

### ANCOM with Neonatal Mouth

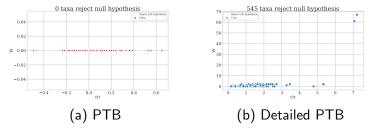


Figure: ANCOM with Neonatal Mouth

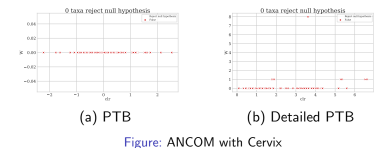
### ANCOM with Maternal Mouth



Figure: ANCOM with Maternal Mouth

 Bacteria Proteobacteria Alphaproteobacteria Rickettsiales mitochondria family

### **ANCOM** with Cervix



# ANCOM with Vagina

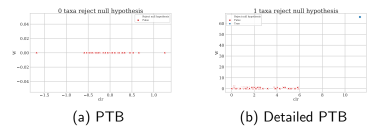


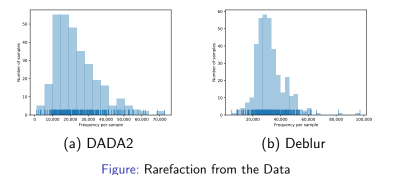
Figure: ANCOM with Vagina

 Bacteria Proteobacteria Epsilonproteobacteria Campylobacterales Campylobacteraceae Campylobacter genus

# Results

Diversity Index

### Rarefaction from First Data



•  $min(\ell_{DADA2})$ : 1046

• *min*(ℓ<sub>Deblur</sub>): 4864

# Alpha-Diversity

# Beta-Diversity

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