## Amplicon Bioinformatic Analysis: DADA2

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## Outline

Bioinformatic Goals

Get Data (pre-DADA2)
Validate Data (pre-DADA2)

Assemble Metadata Table (pre-DADA2)

Demultiplex (pre-DADA2)

Adapter Trimming (pre-DADA2)
Filter and Trim

Learn Error Rates

Dereplication Dereplication

Sample Inference
Merge Paired Reads

Merge Paired Reads
Construct Sequence Table

Remove Chimeras

Assign Taxonomy
Generate Phyloseq Object

Save Phyloseg as RDS

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#### Bioinformatic Goals

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#### Bioinformatic Analysis

#### Input: Raw FASTQ File(s)

M00698:36:000000000-AFBEL:1:1101:14738:1412 1:N:0:0

ABBBABBAFFFGGGGGGGGGGGGGGGCGCGF3FFGHHHHHHGGFGHEHHGGGEHHHHAGGHHGHHFFDHFHHHGEGGGG@F@H?GHH @MO0698:36:000000000-AFBEL:1:1101:16483:1412 1:N:0:0

#### Output: Count Table

	Sample 1	Sample 2	 Sample N
Bacteria 1			
Bacteria 2			
Bacteria N			

#### Naive Approach: Assumptions

- ► Library Prep is Perfect
- Sequencing is Perfect

1. Make an empty count table

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- 2. For each read in the FASTQ:

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- 2. For each read in the FASTQ:
  - 2.1 If read sequence is already in count table, add 1 to that row

- 1. Make an empty count table
- 2. For each read in the FASTQ:
  - 2.1 If read sequence is already in count table, add 1 to that row
  - 2.2 Otherwise add a new row for the sequence and set its count to 1

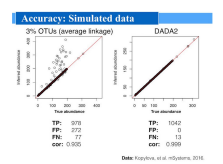
Sequence	Count	1.	CAGCT
		2.	TATAA
		3.	TATAA
		4.	TGCGC
		5.	CGGGC
		6.	TGCGC
		7.	TGCGC
		8.	CAGCT
		9.	CGGGC
		10.	TGCGC
		-	

# Naive Assumptions

- Library Prep is Perfect
- Sequencing is Perfect

## Tools for Bioinformatic Analysis

- ▶ "Clustering"
  - Mothur
  - UCLUST
  - ▶ UPARSE
- ▶ "Denoising"
  - ► DADA2
  - **▶** UNOISE3
  - Deblur



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<sup>a</sup>DADA2 Website

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#### Get Data: Sources

- Sequence Read Archive (SRA)
- MG-RAST (Metagenomic Rapid Annotations using Subsystems Technology)
- Sequencing Facility

#### Get Data: Tools

- curl
- ► wget
- ncftp
- rsync
- ▶ sftp
- ▶ SRA Toolkit

#### Get Data: Result

- FASTQ(s) (gzip'ed)
  - Undetermined\_S0\_L001\_I1\_001.fastq.gz
  - Undetermined\_S0\_L001\_R1\_001.fastq.gz
  - Undetermined\_S0\_L001\_R2\_001.fastq.gz
- Map File\*
  - mydata\_map.txt
- ► Checksum\*
  - md5sum.txt

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#### Validate Data: Input

- FASTQ(s) (gzip'ed)
  - Undetermined S0 L001 I1 001.fastq.gz
  - Undetermined S0 L001 R1 001.fastq.gz
  - Undetermined\_S0\_L001\_R2\_001.fastq.gz
- ► Checksum\*
  - md5sum.txt
- Map File\*
  - mydata\_map.txt

#### Validate Data: Output

```
$ md5sum -c md5sum.txt
mydata_map.txt: OK
Undetermined_SO_L001_I1_001.fastq.gz: OK
Undetermined_SO_L001_R1_001.fastq.gz: OK
Undetermined_SO_L001_R2_001.fastq.gz: OK
```

#### Validate Data: Tools

▶ md5sum

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## Assemble Metadata Table: Why?

Associate barcode with Sample

- Label
- Animal
- ► Site
- Phenotype
- Treatment
- Date
- . . .

# Assemble Metadata Table: Input

- Existing Map
- Publication
- Notes

# Assemble Metadata Table: Output Metadata Table (Mapping File)

#Sample	ID Barc	odeSequence LinkerPrimerSequ	ence	Treatment
PC.354	AGCACGAGCCTA	YATGCTGCCTCCCGTAGGAGT	Control	20061218
PC.355	AACTCGTCGATG	YATGCTGCCTCCCGTAGGAGT	Control	20061218
PC.356	ACAGACCACTCA	YATGCTGCCTCCCGTAGGAGT	Control	20061126
PC.481	ACCAGCGACTAG	YATGCTGCCTCCCGTAGGAGT	Control	20070314
PC.593	AGCAGCACTTGT	YATGCTGCCTCCCGTAGGAGT	Control	20071210
PC.607	AACTGTGCGTAC	YATGCTGCCTCCCGTAGGAGT	Fast	20071112
PC.634	ACAGAGTCGGCT	YATGCTGCCTCCCGTAGGAGT	Fast	20080116
PC.635	ACCGCAGAGTCA	YATGCTGCCTCCCGTAGGAGT	Fast	20080116
PC.636	ACGGTGAGTGTC	YATGCTGCCTCCCGTAGGAGT	Fast	20080116

DOB Description
Control\_mouse\_\_I.D.\_354
Control\_mouse\_\_I.D.\_355
Control\_mouse\_\_I.D.\_356
Control\_mouse\_\_I.D.\_481
Control\_mouse\_\_I.D.\_693
Fasting\_mouse\_\_I.D.\_607
Fasting\_mouse\_\_I.D.\_635
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Fasting\_mouse\_\_I.D.\_636

#### Assemble Metadata Table: Tools

- Excel
- ► Text Editor
- Script

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Generate Phyloseq Object

Save Phyloseq as RDS

Demultiplex: Why?

Split FASTQ File(s) by sample <sup>1</sup>

<sup>&</sup>lt;sup>1</sup>Some data comes demultiplexed

## Demultiplex: Input

- Sequence FASTQ(s)
  - Undetermined\_S0\_L001\_I1\_001.fastq.gz
  - Undetermined S0 L001 R1 001.fastq.gz
- ▶ Barcode FASTQ or Trimmed Versions <sup>2</sup>
  - Undetermined S0 L001 R2 001.fastq.gz
- Map File
  - mydata\_map.txt

<sup>&</sup>lt;sup>2</sup>Some facilities incorporate barcodes in the sequence FASTQ, these will need to be extracted

#### Demultiplex: Output

#### Demultiplexed FASTQs

- sampleA\_R1.fastq.gz
- sampleB\_R1.fastq.gz
- sampleC\_R1.fastq.gz
- **.** . . .
- sampleA\_R2.fastq.gz
- sampleB\_R2.fastq.gz
- sampleC\_R2.fastq.gz
- **.** . . .

## Demultiplex: Tools

- split\_libraries\_fastq.py + split\_sequence\_file\_on\_sample\_ids.py
- fastq\_multx

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## Adapter Trimming: Why?

Remove adapter contamination

Necessary for amplicons with large variation in length (e.g. ITS)

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## Adapter Trimming: Input

#### Adapter Sequence

 $my\_adapter.fasta$ 

#### Demultiplexed FASTQs

- sampleA\_R1.fastq.gz
  - sampleB\_R1.fastq.gz
- ► sampleC\_R1.fastq.gz
- ▶ sampleA\_R2.fastq.gz
- sampleB\_R2.fastq.gz
- ▶ sampleC\_R2.fastq.gz
- **>** . . .

## Adapter Trimming: Output

#### Trimmed FASTQs

- sampleA\_R1.trim.fastq.gz
- ► sampleB R1.trim.fastq.gz
- sampleC\_R1.trim.fastq.gz
- **.** . . .
- sampleA\_R2.trim.fastq.gz
- sampleB\_R2.trim.fastq.gz
- sampleC\_R2.trim.fastq.gz
- •

#### Synchronized Trimming

Depending on settings, some reads may be thrown out during trimming. It is essential that if a read is thrown out, its paired read is thrown out too. Most trimming software will do this for you if you input R1 and R2 files when you run.

# Adapter Trimming: Tools

- fastq\_mcf
- ► Trimmomatic
- cutadapt
- seqtk
- ► etc

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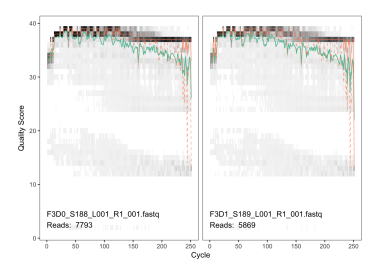
Generate Phylosea Obiec

Save Phyloseq as RDS

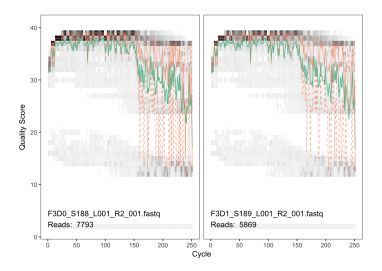
## Filter and Trim: Why?

- Remove low quality parts of reads
- Remove reads that are low quality overall

### R1 Read Quality



## R2 Read Quality



# Filter and Trim: Input

Trimmed FASTQs (or Demultiplexed)

- sampleA\_R1.trim.fastq.gz
- sampleB\_R1.trim.fastq.gz
- sampleC\_R1.trim.fastq.gz
- **•** . . .
- sampleA\_R2.trim.fastq.gz
- sampleB\_R2.trim.fastq.gz
- sampleC\_R2.trim.fastq.gz
- **.** . . .

# Filter and Trim: Output

Trimmed and filtered FASTQs

### Filter and Trim: Tools

dada2::filterAndTrim()

▶ truncQ: Truncate reads at the first instance of a quality score less than or equal to truncQ.

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- truncLen: Truncate reads after truncLen bases. Don't use for ITS

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- minQ: After truncation, reads contain a quality score less than minQ will be discarded.

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- truncLen: Truncate reads after truncLen bases. Don't use for ITS
- trimLeft: The number of nucleotides to remove from the start of each read.
- minQ: After truncation, reads contain a quality score less than minQ will be discarded.
- maxEE: After truncation, reads with higher than maxEE "expected errors" will be discarded.

```
EE = sum(10^{(-Q/10)})
```

- truncQ: Truncate reads at the first instance of a quality score less than or equal to truncQ.
- truncLen: Truncate reads after truncLen bases. Don't use for ITS
- trimLeft: The number of nucleotides to remove from the start of each read.
- minQ: After truncation, reads contain a quality score less than minQ will be discarded.
- ► maxEE: After truncation, reads with higher than maxEE "expected errors" will be discarded.
  FE = gym(10°( 0/10))
  - $EE = sum(10^(-Q/10))$
- rm.phix: Discard reads that match against the phiX genome

#### Filter and Trim: Notes

Paired-End Reads need to be run simultaneously to keep them in sync

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#### Learn Error Rates

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# Learn Error Rates: Why?

Build an error model from data

Phred	A:A	A:T	A:C	A:G	C:A		G:G
1							
2							
3							
40							

#### Learn Error Rates: Input

Filtered and Trimmed FASTQs

# Learn Error Rates: Output

error model

Phred	A:A	A:T	A:C	A:G	C:A		G:G
1							
2							
3							
40							

#### Learn Error Rates: Tools

dada2::learnErrors()

#### Learn Error Rates: Notes

Separate error models need to be built for R1 and R2

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## Dereplication: Why?

Summarize reads into unique observed reads, with quality summary and count

- 1. CAGCT
- 2. TATAA
- 3. TATAA
- 4. TGCGC
- 5. CGGGC
- 6. TGCcC
- 7. TGCGC
- 8. CAGCT
- CGGGa
- 10. TGCGC

Sequence	Count	Quality	
CAGCT	2	99989	
TATAA	2	99998	
TGCGC	3	99988	
CGGGC	1	99999	
TGCcC	1	99948	
CGGGa	1	99993	

### Dereplication: Input

Filtered and Trimmed FASTQs

# Dereplication: Output

Unique reads with summarized quality and counts

### Dereplication: Tools

dada2::derepFastq()

#### Dereplication: Notes

Dereplication is done separately for R1 and R2

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#### Sample Inference

Merge Paired Read

Construct Sequence Table

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# Sample Inference: Why?

Attempt to determine the true sequences from which reads were derived

Sequence	Count	Quality
CAGCT	2	99989
TATAA	2	99998
TGCGC	3	99988
CGGGC	1	99999
TGCcC	1	99948
CGGGa	1	99993

Sequence	Count
CAGCT	2
TATAA	2
TGCGC	4
CGGGC	2

# Sample Inference: Input

- ► Dereplicated Reads
- ► Error Model

## Sample Inference: Output

Inferred read sequences with counts

# Sample Inference: Tools

dada2::dada()

# Sample Inference: Notes

Sample Inference is done separately for R1 and R2

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Sample Inference

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Construct Sequence Table

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#### Merge Paired Reads: Why?

Collapse read pairs into a single sequence for each inferred amplicon

R1: ATACCCTAGTGC

R2: CCCTAGTGCCGT

Merged: ATACCCTAGTGCCGT

# Merge Paired Reads: Input

- ▶ R1
  - Inferred Sequences
  - Dereplicated Sequences
- ► R2
  - Inferred Sequences
  - Dereplicated Sequences

# Merge Paired Reads: Output

Inferred amplicon sequences

#### Merge Paired Reads: Tools

dada2::mergePairs()

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Remove Chimeras

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Generate Phyloseq Object

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#### Construct Sequence Table: Why?

#### Generate count table

	Sample 1	Sample 2	 Sample N
Bacteria 1			
Bacteria 2			
Bacteria N			

#### Construct Sequence Table: Input

Merged sequences

# Construct Sequence Table: Output

Count table

#### Construct Sequence Table: Tools

dada2::makeSequenceTable()

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# Remove Chimeras: Why?

Library preparation is imperfect, so it generates chimeric amplicons

# Remove Chimeras: Input

Count Table

# Remove Chimeras: Output

Count table without chimeras

#### Remove Chimeras: Tools

dada2::removeBimeraDenovo()

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#### Assign Taxonomy

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## Assign Taxonomy: Why?

Relate sequences in our count table to specific bacteria

## Assign Taxonomy: Input

Chimera-free merged sequences

#### Assign Taxonomy: Output

Mapping from sequences to specific bacteria

#### Assign Taxonomy: Tools

dada2::assignTaxonomy()

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#### Generate Phyloseq Object: Why?

Phyloseq objects organize multiple aspects of our results and ease downstream analysis and visualization

# Generate Phyloseq Object: Input

- Count Table
- Metadata Table
- ► Taxonomic Assignment
- Phylogenetic Tree (optional)

## Generate Phyloseq Object: Output

Phyloseq Object

## Generate Phyloseq Object: Tools

phyloseq::phyloseq()

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## Save Phyloseq as RDS: Why?

- Generating the final phyloseq object from raw FASTQs is time consuming, we would prefer to not repeat it everytime we want to play with the results
- ► The Phyloseq object is a very space efficient representation of the processed data

# Save Phyloseq as RDS: Input

- Phyloseq object
- ▶ Name for RDS file

# Save Phyloseq as RDS: Output RDS file

# Save Phyloseq as RDS: Tools

readr::write\_rds()