dunfield_lab_qiime2_pipeline_e_pdf

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1 qiime2 v.2019.4 dunfield lab tutorial

- Relative paths are used
- All commands imply terminal is in the current working directory
- Knowledge of linux terminal is highly incouraged

1.1 Activate qiime2 environment

```
[]: conda activate qiime2-2019.4
```

1.2 Optionally enable qiime-specific autocompletion

```
source tab-qiime
```

1.3 Navigate to your working directory

- qiime2_lab_tutorial already folder contains raw data
- lab_pipeline folder contains trained NaiveBayes Classifier
- mapping file

Next step expects that the folder exists (example)

```
[]: cd ~/Desktop/qiime2_lab_tutorial/lab_pipeline/
```

1.4 Let's view the directory content

```
total 216584
-rwxrwxrwx 1 root root 50718 May 27 23:39
dunfield_lab_qiime2_pipeline_e.ipynb
-rwxrwxrwx 1 root root 70275 May 27 23:35 dunfield_lab_qiime2_pipeline.ipynb
-rwxrwxrwx 1 root root 2580 Mar 25 12:39 mappingfile_upd4.csv
-rwxrwxrwx 1 root root 221649684 May 18 09:59
v3v4_silva132_classifier_wps2_2groups.qza
```

1.5 Trim primers

- Various primer trimming tools exist
 - cutadapt
 - bbmap
 - qiime2 native tools
 - trimmomatic
 - manual removal...

It is critical to remove non-biological sequences from the data. We will remove our 16S V3-V4 region (Bacteria-specific primer set) primers sequences using cutadapt * f-primer CC-TACGGGNGGCWGCAG * r-primer GACTACHVGGGTATCTAATCC

1.5.1 Making directories

```
[58]: mkdir primer_trimmed_fastqs; mkdir cutadapt_logs
```

1.5.2 Primer trimming w cutadapt with a help of a little script:)

! Expects to contain our data in the raw_data folder in a parent directory

• Run the following script in a command line:

```
for file1 in ../raw_data/*_R1_*.fastq.gz; do
    file2="${file1%_R1_001.fastq.gz}_R2_001.fastq.gz"

fname1=`basename $file1`
    fname2=`basename $file2`
    `cutadapt --pair-filter any -j 4 --no-indels --discard-untrimmed \
    -g CCTACGGGNGGCWGCAG -G GACTACHVGGGTATCTAATCC \
    -o primer_trimmed_fastqs/$fname1 -p primer_trimmed_fastqs/$fname2 \
    $file1 $file2 \
    > cutadapt_logs/${fname1}_cutadapt_log.txt`
```

1.6 Import trimmed FASTQs as a QIIME2 artifact

To keep the directory clean you can put the artifact files in a new directory

```
[]: mkdir paired_reads_qza
```

1.6.1 Casava 1.8 single-end demultiplexed fastq

Format description

In the Casava 1.8 demultiplexed (single-end) format, there is one fastq.gz file for each sample in the study which contains the single-end reads for that sample. The file name includes the sample identifier and should look like L2S357_15_L001_R1_001.fastq.gz. The underscore-separated fields in this file name are:

```
the sample identifier,
the barcode sequence or a barcode identifier,
the lane number,
the direction of the read (i.e. only R1, because these are single-end reads), and
the set number.
```

Obtaining example data

1.6.2 Importing...

Imported primer_trimmed_fastqs as CasavaOneEightSingleLanePerSampleDirFmt to paired_reads_qza/reads_trimmed.qza

• Our reads are now ready to be used by qiime2

1.7 Quality control w/ deblur:

Currently deblur doesn't support paired-end reads ### Using VSEARCH for joining:

```
[62]: qiime vsearch join-pairs \
--i-demultiplexed-seqs paired_reads_qza/reads_trimmed.qza \
--o-joined-sequences paired_reads_qza/reads_trimmed_joined.qza
```

```
Saved SampleData[JoinedSequencesWithQuality] to: paired_reads_qza/reads_trimmed_joined.qza
```

1.7.1 Filter out low-quality reads.

This command will filter out low-quality reads based on the default options. (this step may take a while)

```
[63]: qiime quality-filter q-score-joined \
--i-demux paired_reads_qza/reads_trimmed_joined.qza \
--o-filter-stats filt_stats.qza \
--o-filtered-sequences paired_reads_qza/reads_trimmed_joined_filt.qza
```

```
Saved SampleData[JoinedSequencesWithQuality] to:
paired_reads_qza/reads_trimmed_joined_filt.qza
Saved QualityFilterStats to: filt_stats.qza
```

1.7.2 Deblur Workflow

This workflow is 16S sequences, for other amplicon regions, you can use the denoise-other option in the command and specify a reference database.

Note that you will need to trim all sequences to the same length with the –p-trim-length option. In order to determine the correct length to trim down to, run the following QC:

1.7.3 To find appropriate deblur parameters we need to summarize our joined reads

```
[64]: qiime demux summarize \
--i-data paired_reads_qza/reads_trimmed_joined_filt.qza \
--o-visualization reads_trimmed_joined_filt_summary.qzv
```

Saved Visualization to: reads_trimmed_joined_filt_summary.qzv

1.7.4 View the obtained visualization

```
[65]: qiime tools view reads_trimmed_joined_filt_summary.qzv
```

Press the 'q' key, Control-C, or Control-D to quit. This view may no longer be accessible or work correctly after quitting.[20069:20069:0527/234619.288113:ERROR:sandbox_linux.cc(364)] InitializeSandbox() called with multiple threads in process gpu-process. [20029:20050:0527/234619.290330:ERROR:browser_process_sub_thread.cc(209)] Waited 4 ms for network service Opening in existing browser session.

Press the 'q' key, Control-C, or Control-D to quit. This view may no longer be

1.7.5 Qiime help on importing/exporting/viewing artefacts

```
[66]: qiime tools --help
```

Usage: qiime tools [OPTIONS] COMMAND [ARGS]...

Tools for working with QIIME 2 files.

Options:

--help Show this message and exit.

Commands:

citations Print citations for a QIIME 2 result.

export Export data from a QIIME 2 Artifact or a

Visualization

extract a QIIME 2 Artifact or Visualization

archive.

Visualization.

validate Validate data in a QIIME 2 Artifact.

view View a QIIME 2 Visualization.

1.7.6 Explore provenance w/https://view.qiime2.org

Showing on denoise-16S

```
[67]: qiime deblur denoise-16S --help
```

Usage: qiime deblur denoise-16S [OPTIONS]

Perform sequence quality control for Illumina data using the Deblur workflow with a 16S reference as a positive filter. Only forward reads are supported at this time. The specific reference used is the 88% OTUs from Greengenes 13_8. This mode of operation should only be used when data were generated from a 16S amplicon protocol on an Illumina platform. The reference is only used to assess whether each sequence is likely to be 16S by a local alignment using SortMeRNA with a permissive e-value; the reference is not used to characterize the sequences.

Inputs:

```
--i-demultiplexed-seqs ARTIFACT
SampleData[SequencesWithQuality |
    PairedEndSequencesWithQuality | JoinedSequencesWithQuality]
                         The demultiplexed sequences to be denoised.
[required]
Parameters:
  --p-trim-length INTEGER
                         Sequence trim length, specify -1 to disable
                         trimming.
[required]
  --p-left-trim-len INTEGER
    Range(0, None)
                         Sequence trimming from the 5' end. A value of
0 will
                         disable this trim.
                                                                   [default:
  --p-sample-stats / --p-no-sample-stats
                         If true, gather stats per sample.
                                                               [default:
  --p-mean-error NUMBER The mean per nucleotide error, used for
original
                         sequence estimate.
                                                               [default:
0.005]
  --p-indel-prob NUMBER Insertion/deletion (indel) probability
(same for N
                                                                [default:
                         indels).
0.017
  --p-indel-max INTEGER Maximum number of insertion/deletions.
[default: 3]
  --p-min-reads INTEGER Retain only features appearing at least
min-reads
```

```
times across all samples in the resulting feature
                         table.
                                                                  [default:
107
                         In each sample, discard all features with
  --p-min-size INTEGER
an
                         abundance less than min-size.
[default: 2]
  --p-jobs-to-start INTEGER
                         Number of jobs to start (if to run in parallel).
                                                                    [default:
1]
  --p-hashed-feature-ids / --p-no-hashed-feature-ids
                         If true, hash the feature IDs.
                                                             [default:
True]
Outputs:
  --o-table ARTIFACT FeatureTable[Frequency]
                         The resulting denoised feature table.
[required]
  --o-representative-sequences ARTIFACT
FeatureData[Sequence]
                         The resulting feature sequences.
[required]
  --o-stats ARTIFACT
                         Per-sample stats if requested.
    DeblurStats
[required]
Miscellaneous:
  --output-dir PATH
                         Output unspecified results to a directory
  --verbose / --quiet
                         Display verbose output to
stdout and/or stderr
                         during execution of this action. Or silence output if
                         execution is successful (silence is golden).
  --citations
                         Show citations and exit.
                         Show this message and exit.
  --help
```

1.7.7 Denoising w/ deblur

- Here I'm using the default behaviour of –p-min-reads = 10
- Reads are trimmed to 402nt which retains is at least 98% of the reads (this step may take a while depending on the size of your data ...)

```
[106]: qiime deblur denoise-16S \
--i-demultiplexed-seqs paired_reads_qza/reads_trimmed_joined_filt.qza \
--p-trim-length 402 \
--p-sample-stats \
--p-jobs-to-start 8 \
--p-min-reads 10 \
--output-dir deblur_output
```

```
Saved FeatureTable[Frequency] to: deblur_output/table.qza
Saved FeatureData[Sequence] to:
deblur_output/representative_sequences.qza
Saved DeblurStats to: deblur_output/stats.qza
```

1.7.8 Output is saved in the deblur_output folder

let's summarise our deblur output

```
[107]: qiime deblur visualize-stats \
    --i-deblur-stats deblur_output/stats.qza \
    --o-visualization deblur_output/deblur-stats.qzv
```

Saved Visualization to: deblur_output/deblur-stats.qzv

```
[108]: qiime tools view deblur_output/deblur-stats.qzv
```

Press the 'q' key, Control-C, or Control-D to quit. This view may no longer be accessible or work correctly after quitting.[29817:29838:0527/235316.922441:ERRO R:browser_process_sub_thread.cc(209)] Waited 5 ms for network service Opening in existing browser session.

Press the 'q' key, Control-C, or Control-D to quit. This view may no longer be

```
[109]: qiime feature-table summarize \
--i-table deblur_output/table.qza \
--o-visualization deblur_output/deblur_table_summary.qzv
```

Saved Visualization to: deblur_output/deblur_table_summary.qzv

```
[110]: qiime tools view deblur_output/deblur_table_summary.qzv
```

Press the 'q' key, Control-C, or Control-D to quit. This view may no longer be accessible or work correctly after quitting.[30210:30231:0527/235353.890697:ERRO R:browser_process_sub_thread.cc(209)] Waited 5 ms for network service Opening in existing browser session.

Press the 'q' key, Control-C, or Control-D to quit. This view may no longer be

1.7.9 Tabulate representative sequences

```
[111]: qiime feature-table tabulate-seqs \
    --i-data deblur_output/representative_sequences.qza \
    --o-visualization representative_sequences.qzv
```

Saved Visualization to: representative_sequences.qzv

```
[112]: qiime tools view representative_sequences.qzv
```

Press the 'q' key, Control-C, or Control-D to quit. This view may no longer be accessible or work correctly after quitting.[30607:30607:0527/235411.374129:ERROR:sandbox_linux.cc(364)] InitializeSandbox() called with multiple threads in process gpu-process. [30568:30589:0527/235411.374893:ERROR:browser_process_sub_thread.cc(209)] Waited 3 ms for network service Opening in existing browser session.

Press the 'q' key, Control-C, or Control-D to quit. This view may no longer be

1.8 Building phylogeny with FastTree

1.8.1 Making multiple-sequence alignment

We'll need to make a multiple-sequence alignment of the ASVs before running FastTree.

```
[113]: mkdir fast_tree_out
```

mkdir: cannot create directory fast_tree_out: File exists

```
[114]: qiime alignment mafft \
--i-sequences deblur_output/representative_sequences.qza \
--p-n-threads 8 \
--o-alignment fast_tree_out/rep_seqs_mafft.qza
```

Saved FeatureData[AlignedSequence] to: fast_tree_out/rep_seqs_mafft.qza

1.8.2 Filtering multiple-sequence alignment

Variable positions in the alignment need to be masked before FastTree is run, which can be done with this command:

```
[115]: qiime alignment mask --i-alignment fast_tree_out/rep_seqs_mafft.qza \
--o-masked-alignment fast_tree_out/rep_seqs_mafft_masked.qza
```

```
Saved FeatureData[AlignedSequence] to: fast_tree_out/rep_seqs_mafft_masked.qza
```

1.8.3 Running FastTree

Finally FastTree can be run on this masked multiple-sequence alignment:

```
qiime phylogeny fasttree \
--i-alignment fast_tree_out/rep_seqs_mafft_masked.qza \
--p-n-threads 4 \
--o-tree fast_tree_out/rep_seqs_aligned_masked_tree
```

```
Saved Phylogeny[Unrooted] to:
fast_tree_out/rep_seqs_aligned_masked_tree.qza
```

1.8.4 Make a rooted tree

Use midpoint root

```
[117]: qiime phylogeny midpoint-root \
--i-tree fast_tree_out/rep_seqs_aligned_masked_tree.qza \
--o-rooted-tree fast_tree_out/rep_seqs_mafft_masked_tree_rooted.qza
```

```
Saved Phylogeny[Rooted] to:
fast_tree_out/rep_seqs_mafft_masked_tree_rooted.qza
```

1.8.5 Generate rarefaction curves

- Useful QC step
- Determine maximum depth for the rarefaction using following (I'm using 8000):

```
[118]: qiime tools view deblur_output/deblur_table_summary.qzv
```

```
Press the 'q' key, Control-C, or Control-D to quit. This view may no longer be accessible or work correctly after quitting. [31267:31267:0527/235529.491841:ERROR:sandbox_linux.cc(364)] InitializeSandbox() called with multiple threads in process gpu-process. [31226:31247:0527/235529.502609:ERROR:browser_process_sub_thread.cc(209)] Waited 5 ms for network service Opening in existing browser session.
```

Press the 'q' key, Control-C, or Control-D to quit. This view may no longer be

[119]: qiime diversity alpha-rarefaction --help

```
Usage: qiime diversity alpha-rarefaction [OPTIONS]
```

Generate interactive alpha rarefaction curves by computing rarefactions between `min_depth` and `max_depth`. The number of intermediate depths to compute is controlled by the `steps` parameter, with n `iterations` being computed at each rarefaction depth. If sample metadata is provided, samples may be grouped based on distinct values within a metadata column.

Inputs:

```
--i-table ARTIFACT FeatureTable[Frequency]

Feature table to compute rarefaction curves from.

[required]

--i-phylogeny ARTIFACT Optional phylogeny for phylogenetic metrics.

Phylogeny[Rooted]

[optional]

Parameters:
```

--p-max-depth INTEGER The maximum rarefaction depth. Must be

```
greater than
       Range(1, None)
                             min-depth.
   [required]
     --p-metrics TEXT... Choices('margalef', 'simpson',
       'michaelis_menten_fit', 'simpson_e', 'gini_index', 'faith_pd',
       'mcintosh_d', 'mcintosh_e', 'enspie', 'doubles', 'brillouin_d',
       'goods_coverage', 'lladser_pe', 'ace', 'fisher_alpha', 'shannon',
       'dominance', 'heip_e', 'pielou_e', 'chao1', 'robbins', 'menhinick',
       'singles', 'berger_parker_d', 'observed_otus')
                             The metrics to be measured. By default computes
                             observed_otus, shannon, and if phylogeny is
                             provided, faith_pd.
   [optional]
     --m-metadata-file METADATA...
       (multiple arguments
                             The sample metadata.
       will be merged)
   [optional]
     --p-min-depth INTEGER
                             The minimum rarefaction depth.
      Range(1, None)
   [default: 1]
     --p-steps INTEGER
                             The number of rarefaction depths to
  include between
       Range(2, None)
                             min-depth and max-depth.
   [default: 10]
     --p-iterations INTEGER The number of rarefied feature tables to
  compute at
      Range(1, None)
                             each step.
   [default: 10]
  Outputs:
     --o-visualization VISUALIZATION
   [required]
  Miscellaneous:
     --output-dir PATH
                             Output unspecified results to a
  directory
     --verbose / --quiet
                             Display verbose output to
  stdout and/or stderr
                             during execution of this action. Or silence output
                             if execution is successful (silence is golden).
     --citations
                             Show citations and exit.
     --help
                             Show this message and exit.
[]:
[]:
[]:
```

```
[120]: qiime diversity alpha-rarefaction \
--i-table deblur_output/table.qza \
--p-max-depth 8000 \
--p-metrics simpson \
--p-metrics faith_pd \
--p-metrics dominance \
--p-metrics chao1 \
--p-metrics observed_otus \
--p-metrics shannon \
--p-steps 20 \
--i-phylogeny fast_tree_out/rep_seqs_mafft_masked_tree_rooted.qza \
--o-visualization rarefaction_curves.qzv
```

Saved Visualization to: rarefaction_curves.qzv

```
[121]: qiime tools view rarefaction_curves.qzv
```

Press the 'q' key, Control-C, or Control-D to quit. This view may no longer be accessible or work correctly after quitting.[31652:31673:0527/235612.774071:ERRO R:browser_process_sub_thread.cc(209)] Waited 3 ms for network service Opening in existing browser session.

Press the 'q' key, Control-C, or Control-D to quit. This view may no longer be

1.8.6 Using metadata w rarefaction

```
[122]: qiime diversity alpha-rarefaction \
--i-table deblur_output/table.qza \
--p-max-depth 8000 \
--p-steps 20 \
--i-phylogeny fast_tree_out/rep_seqs_mafft_masked_tree_rooted.qza \
--m-metadata-file mappingfile_upd4.csv \
--o-visualization rarefaction_metadata_curves.qzv
```

Plugin error from diversity:

The following IDs are not present in the metadata: 'nm1-9a', 'o1', 'o29' Debug info has been saved to /tmp/qiime2-q2cli-err-6vb4gl4x.log

1.8.7 Ups! We seem to have an error...

- We need to remove those samples from FeatureTable
- Also, we have 2 samples that contain no data, let's remove them as well

```
[123]: echo SampleID > samples-to-exclude.tsv
[]: echo nm1-9a >> samples-to-exclude.tsv
```

```
[]: echo o1 >> samples-to-exclude.tsv
[]: echo o29 >> samples-to-exclude.tsv
[]: echo o20 >> samples-to-exclude.tsv
[]: echo o7 >> samples-to-exclude.tsv
```

1.8.8 Filtering out samples

```
[124]: qiime feature-table filter-samples \
    --p-exclude-ids \
    --i-table deblur_output/table.qza \
    --m-metadata-file samples-to-exclude.tsv \
    --o-filtered-table id-filtered-deblur-table.qza
```

Saved FeatureTable[Frequency] to: id-filtered-deblur-table.qza

1.8.9 Let's run it again

• Pay attention that we are supplying updated Feature Table as its –i-table argument

```
[125]: qiime diversity alpha-rarefaction \
--i-table id-filtered-deblur-table.qza \
--p-max-depth 8000 \
--p-steps 20 \
--i-phylogeny fast_tree_out/rep_seqs_mafft_masked_tree_rooted.qza \
--m-metadata-file mappingfile_upd4.csv \
--o-visualization rarefaction_metadata_curves.qzv
```

Saved Visualization to: rarefaction_metadata_curves.qzv

```
[126]: qiime tools view rarefaction_metadata_curves.qzv
```

Press the 'q' key, Control-C, or Control-D to quit. This view may no longer be accessible or work correctly after quitting. [32156:32156:0527/235720.460100:ERROR:sandbox_linux.cc(364)] InitializeSandbox() called with multiple threads in process gpu-process. [32116:32137:0527/235720.466302:ERROR:browser_process_sub_thread.cc(209)] Waited 5 ms for network service Opening in existing browser session.

Press the 'q' key, Control-C, or Control-D to quit. This view may no longer be

1.8.10 Optional step to re-summarize our filtered Feature Table

```
[127]: qiime feature-table summarize \
    --i-table id-filtered-deblur-table.qza \
    --o-visualization id-filtered-deblur-table.qzv \
    --m-sample-metadata-file mappingfile_upd4.csv
```

Saved Visualization to: id-filtered-deblur-table.qzv

```
[128]: qiime tools view id-filtered-deblur-table.qzv
```

Press the 'q' key, Control-C, or Control-D to quit. This view may no longer be accessible or work correctly after quitting. [32516:32516:0527/235731.148830:ERROR:sandbox_linux.cc(364)] InitializeSandbox() called with multiple threads in process gpu-process. [32474:32495:0527/235731.158822:ERROR:browser_process_sub_thread.cc(209)] Waited 3 ms for network service Opening in existing browser session.

Press the 'q' key, Control-C, or Control-D to quit. This view may no longer be

1.9 Assign taxonomy

- Could be assigned to ASVs using a Naive-Bayes classifier
- This classifier was trained using SILVA 132 database and is specific for v3v4 region
- Contains edits for WPS-2 (Rubrimentifilales and AS-11)
- Could be trained de novo, but RAM intensive
- Qiime version sensitive

(this step may take a long time to complete ...)

```
[129]: qiime feature-classifier classify-sklearn \
--i-reads deblur_output/representative_sequences.qza \
--i-classifier v3v4_silva132_classifier_wps2_2groups.qza \
--output-dir taxonomy
```

Saved FeatureData[Taxonomy] to: taxonomy/classification.qza

1.9.1 Our taxonomy folder now contains classification.qza file

let's explore the results..

Following command export the classification as a tsv-file

```
[130]: qiime tools export \
--input-path taxonomy/classification.qza \
--output-path taxonomy
```

Exported taxonomy/classification.qza as TSVTaxonomyDirectoryFormat to directory taxonomy

1.9.2 At last..., Our Beloved Bar-Chart:)

```
[131]: qiime taxa barplot \
--i-table id-filtered-deblur-table.qza \
--i-taxonomy taxonomy/classification.qza \
--m-metadata-file mappingfile_upd4.csv \
--o-visualization taxonomy/taxa_barplot.qzv
```

Saved Visualization to: taxonomy/taxa_barplot.qzv

```
[132]: qiime tools view taxonomy/taxa_barplot.qzv
```

Press the 'q' key, Control-C, or Control-D to quit. This view may no longer be accessible or work correctly after quitting. [6326:6375:0528/003825.571767:ERROR:browser_process_sub_thread.cc(209)] Waited 3 ms for network service Opening in existing browser session.

Press the 'q' key, Control-C, or Control-D to quit. This view may no longer be

1.10 Finally, let's calculate core diversity metrics

- For this step we need to select a reasonable rarefaction value
- Let's have a look at our FeatureTable again

```
[133]: qiime tools view id-filtered-deblur-table.qzv
```

Press the 'q' key, Control-C, or Control-D to quit. This view may no longer be accessible or work correctly after quitting. [6800:6800:0528/003836.431192:ERROR:sandbox_linux.cc(364)] InitializeSandbox() called with multiple threads in process gpu-process. [6759:6780:0528/003836.436214:ERROR:browser_process_sub_thread.cc(209)] Waited 5 ms for network service Opening in existing browser session.

Press the 'q' key, Control-C, or Control-D to quit. This view may no longer be

400 seems to be a good number in this case, we are loosing only 2 samples

```
[134]: qiime diversity core-metrics-phylogenetic \
--i-phylogeny fast_tree_out/rep_seqs_mafft_masked_tree_rooted.qza \
--i-table id-filtered-deblur-table.qza \
--p-sampling-depth 400 \
--m-metadata-file mappingfile_upd4.csv \
--output-dir core-metrics
```

```
Saved FeatureTable[Frequency] to: core-metrics/rarefied table.qza
Saved SampleData[AlphaDiversity] % Properties('phylogenetic') to: core-
metrics/faith_pd_vector.qza
Saved SampleData[AlphaDiversity] to: core-
metrics/observed_otus_vector.qza
Saved SampleData[AlphaDiversity] to: core-metrics/shannon vector.qza
Saved SampleData[AlphaDiversity] to: core-metrics/evenness_vector.qza
Saved DistanceMatrix % Properties('phylogenetic') to: core-
metrics/unweighted_unifrac_distance_matrix.qza
Saved DistanceMatrix % Properties('phylogenetic') to: core-
metrics/weighted_unifrac_distance_matrix.qza
Saved DistanceMatrix to: core-metrics/jaccard_distance_matrix.qza
Saved DistanceMatrix to: core-metrics/bray_curtis_distance_matrix.qza
Saved PCoAResults to: core-metrics/unweighted unifrac pcoa results.qza
Saved PCoAResults to: core-metrics/weighted_unifrac_pcoa_results.qza
Saved PCoAResults to: core-metrics/jaccard pcoa results.gza
Saved PCoAResults to: core-metrics/bray_curtis_pcoa_results.qza
Saved Visualization to: core-metrics/unweighted_unifrac_emperor.qzv
Saved Visualization to: core-metrics/weighted unifrac emperor.qzv
Saved Visualization to: core-metrics/jaccard_emperor.qzv
Saved Visualization to: core-metrics/bray_curtis_emperor.qzv
```

let's view an ordination plot

```
[135]: qiime tools view core-metrics/weighted_unifrac_emperor.qzv
```

```
Press the 'q' key, Control-C, or Control-D to quit. This view may no longer be accessible or work correctly after quitting. [7169:7226:0528/003856.615757:ERROR:browser_process_sub_thread.cc(209)] Waited 3 ms for network service [7278:7278:0528/003856.616153:ERROR:sandbox_linux.cc(364)] InitializeSandbox() called with multiple threads in process gpu-process. Opening in existing browser session.
```

Press the 'q' key, Control-C, or Control-D to quit. This view may no longer be

1.10.1 Alpha diversity group significance test

• An example of just one metric

```
qiime diversity alpha-group-significance \
--i-alpha-diversity core-metrics/faith_pd_vector.qza \
--m-metadata-file mappingfile_upd4.csv \
--o-visualization core-metrics/faith-pd-group-significance.qzv
```

Saved Visualization to: core-metrics/faith-pd-group-significance.qzv

[137]: qiime tools view core-metrics/faith-pd-group-significance.qzv

Press the 'q' key, Control-C, or Control-D to quit. This view may no longer be accessible or work correctly after quitting. [7683:7683:0528/003911.628849:ERROR:sandbox_linux.cc(364)] InitializeSandbox() called with multiple threads in process gpu-process. [7643:7664:0528/003911.631947:ERROR:browser_process_sub_thread.cc(209)] Waited 5 ms for network service Opening in existing browser session.

Press the 'q' key, Control-C, or Control-D to quit. This view may no longer be

1.10.2 Beta diversity group significance test

• lets test weighted unifrac

```
[138]: qiime diversity beta-group-significance \
--i-distance-matrix core-metrics/weighted_unifrac_distance_matrix.qza \
--m-metadata-file mappingfile_upd4.csv \
--m-metadata-column Location \
--p-pairwise \
--o-visualization core-metrics/unweighted-unifrac-bodysite-significance.qzv
```

Saved Visualization to: core-metrics/unweighted-unifrac-bodysite-significance.qzv

[139]: qiime tools view core-metrics/unweighted-unifrac-bodysite-significance.qzv

Press the 'q' key, Control-C, or Control-D to quit. This view may no longer be accessible or work correctly after quitting. [8054:8054:0528/003925.363841:ERROR:sandbox_linux.cc(364)] InitializeSandbox() called with multiple threads in process gpu-process. [8012:8033:0528/003925.379743:ERROR:browser_process_sub_thread.cc(209)] Waited 5 ms for network service Opening in existing browser session.

Press the 'q' key, Control-C, or Control-D to quit. This view may no longer be

1.11 Bonus part: Exporting Feature Tables (biom files)

- qiime2 keeps taxonomy separately
- therefore exporting biom files with taxonomy needs some additional steps

Here we are slightly modifying the header of the taxonomy file

[140]: sed -i -e '1 s/Feature/#OTUID/' -e '1 s/Taxon/taxonomy/' taxonomy/taxonomy.tsv

1.11.1 Let's export the biom file into the folder and convert to tsv there

```
[141]: qiime tools export \
--input-path id-filtered-deblur-table.qza \
--output-path id-filtered-deblur-table-exported
```

Exported id-filtered-deblur-table.qza as BIOMV210DirFmt to directory id-filtered-deblur-table-exported

```
[142]: biom add-metadata \
-i id-filtered-deblur-table-exported/feature-table.biom \
-o id-filtered-deblur-table-exported/feature-table_w_tax.biom \
--observation-metadata-fp taxonomy/taxonomy.tsv \
--sc-separated taxonomy
```

1.11.2 And our familiar biom-convert:)

```
[143]: biom convert \
-i id-filtered-deblur-table-exported/feature-table_w_tax.biom \
-o id-filtered-deblur-table-exported/feature-table.tsv \
--to-tsv --header-key taxonomy
```

1.11.3 Finally, let's convert our biom table as % at a specific taxon level

```
[]: mkdir taxa-levels
```

collapsing to a level

```
[]: qiime taxa collapse \
--i-table id-filtered-deblur-table.qza \
--i-taxonomy taxonomy/classification.qza \
--p-level 2 \
--o-collapsed-table taxa-levels/table-12.qza
```

converting counts to fractions

```
[]: qiime feature-table relative-frequency \
--i-table taxa-levels/table-12.qza \
--o-relative-frequency-table taxa-levels/frac-table-12.qza
```

• exporting a biom file

```
[]: qiime tools export \
--input-path taxa-levels/frac-table-12.qza \
--output-path taxa-levels/frac-table-12
```

exporting biom to tsv

```
[]: biom convert \
    -i taxa-levels/frac-table-l2/feature-table.biom \
    -o taxa-levels/frac-table-l2/feature-table.tsv \
    --to-tsv \
    --header-key taxonomy
```

1.12 Good-bye!