OTU Picking

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All of the code in this page is meant to be run on the command line.	

Getting usage info from QIIME scripts

First let's move to the Guerrero Negro data directory. This directory contains sequences from different depths in the Guerrero Negro microbial in Mexico (Harris JK et al. ISME J. 2013 Jan;7(1):50-60).

```
# To be run on the command line
# you will need to put your correct path here
cd /your/path/to/8992repo/data/guerreronegro
```

We are going to run closed reference OTU picking, but first let's get usage information and list of options (be sure QIIME is loaded first)

```
pick_closed_reference_otus.py -h
```

```
## Usage: pick_closed_reference_otus.py [options] {-i/--input_fp INPUT_FP -o/--output_dir OUTPUT_DIR}
##
## [] indicates optional input (order unimportant)
## {} indicates required input (order unimportant)
##
##
##
## This script picks OTUs using a closed reference and constructs an OTU table.
## Taxonomy is assigned using a pre-defined taxonomy map of reference sequence OTU
## to taxonomy. If full-length genomes are provided as the reference sequences,
## this script applies the Shotgun UniFrac method.
##
## **Note:** If most or all of your sequences are failing to hit the reference,
```

```
## your sequences may be in the reverse orientation with respect to your reference
## database. To address this, you should add the following line to your parameters
## file (creating one, if necessary) and pass this file as -p:
## pick_otus:enable_rev_strand_match True
##
## Be aware that this doubles the amount of memory used.
##
##
##
## Example usage:
## Print help message and exit
## pick_closed_reference_otus.py -h
##
## Pick OTUs, assign taxonomy, and create an OTU table against a reference set of OTUs. ALWAYS SPECIFY
  pick_closed_reference_otus.py -i $PWD/seqs.fna -r $PWD/refseqs.fna -o $PWD/otus_w_tax/ -t $PWD/taxa
##
## Pick OTUs and create an OTU table against a reference set of OTUs without adding taxonomy assignment
   pick_closed_reference_otus.py -i $PWD/seqs.fna -r $PWD/refseqs.fna -o $PWD/otus/
##
## Pick OTUs, assign taxonomy, and create an OTU table against a reference set of OTUs using usearch_re
   pick_closed_reference_otus.py -i $PWD/seqs.fna -r $PWD/refseqs.fna -o $PWD/otus_usearch/ -p $PWD/us
##
## Pick OTUs using usearch_ref, assign taxonomy, and create an OTU table against a reference set of OTU
   pick_closed_reference_otus.py -i $PWD/seqs.fna -r $PWD/refseqs.fna -o $PWD/otus_usearch_ref/ -p $PW
## Pick OTUs, assign taxonomy, and create an OTU table against a reference set of OTUs using sortmerna.
   pick_closed_reference_otus.py -i $PWD/seqs.fna -r $PWD/refseqs.fna -o $PWD/otus_sortmerna/ -p $PWD/
##
##
## Options:
##
     --version
                           show program's version number and exit
##
     -h, --help
                           show this help message and exit
##
     -v, --verbose
                           Print information during execution -- useful for
##
                           debugging [default: False]
##
     -r REFERENCE_FP, --reference_fp=REFERENCE_FP
##
                           The reference sequences [default:
##
                           /macqiime/anaconda/lib/python2.7/site-packages/qiime_d
##
                           efault_reference/gg_13_8_otus/rep_set/97_otus.fasta].
                           NOTE: If you do not pass -r to this script, you will
##
                           be using QIIME's default reference sequences. In this
##
                           case, QIIME will copy the corresponding reference tree
##
##
                           to the output directory. This is the tree that should
##
                           be used to perform phylogenetic diversity analyses
##
                           (e.g., with core_diversity_analyses.py).
##
     -p PARAMETER_FP, --parameter_fp=PARAMETER_FP
                           path to the parameter file, which specifies changes to
##
##
                           the default behavior. See
                           http://www.qiime.org/documentation/file_formats.html
##
##
                           #qiime-parameters . [if omitted, default values will
##
                           be used]
     -t TAXONOMY_FP, --taxonomy_fp=TAXONOMY_FP
##
##
                           the taxonomy map [default:
##
                           /macqiime/anaconda/lib/python2.7/site-packages/qiime_d
##
                           efault_reference/gg_13_8_otus/taxonomy/97_otu_taxonomy
```

```
##
                            .txt]
##
     -s, --assign_taxonomy
                           Assign taxonomy to each sequence using
##
                           assign_taxonomy.py (this will override --taxonomy_fp,
##
##
                           if provided) [default: False]
##
     -f, --force
                           Force overwrite of existing output directory (note:
                           existing files in output_dir will not be removed)
##
##
                            [default: none]
##
                           Print the commands but don't call them -- useful for
     -w, --print_only
##
                           debugging [default: False]
##
     -a, --parallel
                           Run in parallel where available [default: False]
     -O JOBS_TO_START, --jobs_to_start=JOBS_TO_START
##
##
                           Number of jobs to start. NOTE: you must also pass -a
##
                           to run in parallel, this defines the number of jobs to
##
                           be started if and only if -a is passed [default: 1]
##
     --suppress_taxonomy_assignment
##
                           skip the taxonomy assignment step, resulting in an OTU
##
                           table without taxonomy (this will override
##
                            --taxonomy_fp and --assign_taxonomy, if provided)
##
                            [default: False]
##
     REQUIRED options:
##
       The following options must be provided under all circumstances.
##
##
##
       -i INPUT_FP, --input_fp=INPUT_FP
##
                           the input sequences [REQUIRED]
##
       -o OUTPUT_DIR, --output_dir=OUTPUT_DIR
##
                           the output directory [REQUIRED]
```

pick_closed_reference_otus.py is a "workflow" script that runs other scripts.

Ask pick_closed_reference_otus.py to print the other commands it would run. The -f tells it to force overwriting the directory closedref, in case that directory is already there. The -r tells it where the reference sequences are. The -t tells it where the reference taxonomy map is.

```
pick_closed_reference_otus.py -i seqs.fna -o closedref -f -r ../ref/greengenes/97_otus.fasta -t ../ref/
```

```
## pick_otus.py -i seqs.fna -o closedref/uclust_ref_picked_otus -r ../ref/greengenes/97_otus.fasta -m u
## make_otu_table.py -i closedref/uclust_ref_picked_otus/seqs_otus.txt -t ../ref/greengenes/97_otu_taxo
```

Picking closed reference OTUs with QIIME

Now actually run the script. The -v tells it to be "verbose", printing updates as it runs. Use time to report how long it takes.

```
time pick_closed_reference_otus.py -i seqs.fna -o closedref -r ../ref/greengenes/97_otus.fasta -t ../re
## Pick OTUs
```

```
## pick_otus.py -i seqs.fna -o closedref/uclust_ref_picked_otus -r ../ref/greengenes/97_otus.fasta -m u
## Make OTU table
## make_otu_table.py -i closedref/uclust_ref_picked_otus/seqs_otus.txt -t ../ref/greengenes/97_otu_taxo
```

```
## real 0m37.278s
## user 0m30.371s
## sys 0m3.007s
```

Picking closed-reference OTUs with NINJA-OPS

Compare to NINJA-OPS. Installation instructions are at https://github.com/GabeAl/NINJA-OPS. Note NINJA-OPS is much faster.

```
time python /path/to/your/ninja/directory/bin/ninja.py -i seqs.fna -o ninja
```

```
## Ninja 1.3.1
## Ninja database directory is is/Users/danknights/Copy/research/ninja/src/databases/greengenes97
## Running Ninja filter...
## "/Users/danknights/Copy/research/ninja/src/bin/ninja_filter_mac" "seqs.fna" "/Users/danknights/Dropb
## Not using paired-end reads
## Performing NINJA replicon-denoising at 1 compacted reads.
## Number of sequences: 27389
## Total reads considered: 27389
##
## DONE SORTING.
## 18 Samples found.
## Ninja filter time: 0.135617017746
## Running Bowtie...
## bowtie2-align-s --no-head -x /Users/danknights/Copy/research/ninja/src/databases/greengenes97/greeng
## Bowtie time: 5.48249912262
## Running Ninja parse...
## "/Users/danknights/Copy/research/ninja/src/bin/ninja_parse_filtered_mac" "/Users/danknights/Dropbox/
## Opened /Users/danknights/Dropbox/docs/teaching/mice8992-2016-spring/repo/data/guerreronegro/ninja/ni
## Total OTUs available: 99322
## Number of unique samples: 18, max reads: 14793
## Total reads expanded: 23138
## Run complete.
## Ninja parse time: 0.142162799835
## real 0m5.862s
## user 0m11.703s
## sys 0m0.595s
```

Find out how many sequences were assigned ("Total count") by uclust_ref (QIIME default) and how many with NINJA-OPS using the biom summarize-table command.

```
biom summarize-table -i closedref/otu_table.biom > closedref/stats.txt
head closedref/stats.txt

## Num samples: 18
## Num observations: 1750
## Total count: 22879
## Table density (fraction of non-zero values): 0.192
```

```
##
## Counts/sample summary:
## Min: 950.0
## Max: 1758.0
## Median: 1262.000
## Mean: 1271.056
biom summarize-table -i ninja/ninja_otutable.biom > ninja/stats.txt
head ninja/stats.txt
## Num samples: 18
## Num observations: 2131
## Total count: 23138
## Table density (fraction of non-zero values): 0.193
## Counts/sample summary:
## Min: 962.0
## Max: 1773.0
## Median: 1276.000
## Mean: 1285.444
Try NINJA-OPS on the Global Gut data set. It can process 1 million sequences in the Global Gut data set
in under 20 seconds on a Macbook.
cd ../globalgut
time python /path/to/your/ninja/directory/bin/ninja.py -i seqs.fna -o ninja
## Ninja 1.3.1
## Ninja database directory is is/Users/danknights/Copy/research/ninja/src/databases/greengenes97
## Running Ninja filter...
## "/Users/danknights/Copy/research/ninja/src/bin/ninja_filter_mac" "seqs.fna" "/Users/danknights/Dropb
## Not using paired-end reads
## Performing NINJA replicon-denoising at 1 compacted reads.
## Number of sequences: 1000000
## Total reads considered: 1000000
## DONE SORTING.
## 489 Samples found.
## Ninja filter time: 3.66370201111
## Running Bowtie...
## bowtie2-align-s --no-head -x /Users/danknights/Copy/research/ninja/src/databases/greengenes97/greeng
##
## Bowtie time: 67.1731250286
## Running Ninja parse...
## "/Users/danknights/Copy/research/ninja/src/bin/ninja_parse_filtered_mac" "/Users/danknights/Dropbox/
## Opened /Users/danknights/Dropbox/docs/teaching/mice8992-2016-spring/repo/data/globalgut/ninja/ninja_
## Total OTUs available: 99322
## Number of unique samples: 489, max reads: 117369
## Total reads expanded: 942750
## Run complete.
## Ninja parse time: 0.650639057159
```

##

```
## real 1m11.605s
## user 0m47.767s
## sys 0m1.438s
```

Picking de novo OTUs with QIIME

Run the workflow script, pick_de_novo_otus.py with -w to print the basic commands it would run.

```
cd ../guerreronegro
pick_de_novo_otus.py -i seqs.fna -o openref -f -w

## pick_otus.py -i seqs.fna -o openref/uclust_picked_otus
## pick_rep_set.py -i openref/uclust_picked_otus/seqs_otus.txt -f seqs.fna -l openref/rep_set//seqs_rep
## assign_taxonomy.py -o openref/uclust_assigned_taxonomy -i openref/rep_set//seqs_rep_set.fasta
## make_otu_table.py -i openref/uclust_picked_otus/seqs_otus.txt -t openref/uclust_assigned_taxonomy/se
## align_seqs.py -i openref/rep_set//seqs_rep_set.fasta -o openref/pynast_aligned_seqs
## filter_alignment.py -o openref/pynast_aligned_seqs -i openref/pynast_aligned_seqs/seqs_rep_set_align
## make_phylogeny.py -i openref/pynast_aligned_seqs/seqs_rep_set_aligned_pfiltered.fasta -o openref/rep
```

Now actually run the script. Note that the OTU picking step is quick on this data set. The other steps will take a long time (more than 10 minutes). So instead, kill the script by typing ctrl-c.

```
pick_de_novo_otus.py -i seqs.fna -o openref -f -v
```

Instead let's just run the pick_otus.py step, with several different OTU picking methods.

```
time pick_otus.py -i seqs.fna -o uclust -m uclust
make_otu_table.py -i uclust/seqs_otus.txt -o uclust/otu_table.biom
biom summarize-table -i uclust/otu_table.biom | head -n 5
```

```
##
## real 0m3.236s
## user 0m2.688s
## sys 0m0.388s
## Num samples: 18
## Num observations: 4074
## Total count: 27389
## Table density (fraction of non-zero values): 0.118

time pick_otus.py -i seqs.fna -o swarm -m swarm
make_otu_table.py -i swarm/seqs_otus.txt -o swarm/otu_table.biom
biom summarize-table -i swarm/otu_table.biom | head -n 5
```

```
##
## real 0m4.681s
## user 0m3.816s
## sys 0m0.506s
## Num samples: 18
## Num observations: 10025
## Total count: 27389
## Table density (fraction of non-zero values): 0.075
```

```
time pick_otus.py -i seqs.fna -o cdhit -m cdhit
make_otu_table.py -i cdhit/seqs_otus.txt -o cdhit/otu_table.biom
biom summarize-table -i cdhit/otu table.biom | head -n 5
##
## real 0m32.281s
## user 0m28.681s
## sys 0m1.139s
## Num samples: 18
## Num observations: 4520
## Total count: 27389
## Table density (fraction of non-zero values): 0.121
time pick_otus.py -i seqs.fna -o sumaclust -m sumaclust
make_otu_table.py -i sumaclust/seqs_otus.txt -o sumaclust/otu_table.biom
biom summarize-table -i sumaclust/otu_table.biom | head -n 5
##
## real 1m18.702s
## user 1m14.010s
## sys 0m1.423s
## Num samples: 18
## Num observations: 3552
## Total count: 27389
## Table density (fraction of non-zero values): 0.125
```

Customizing workflow scripts

We can customize workflow scripts using a parameter file. You will need some kind of text editor to edit the parameter file, such as Notepad or Notepad++ for Windows, TextWrangler or Sublime for Mac, or Emacs or vi for the Mac/Linux command line. For more information, please see the official QIIME explaining parameter files here: http://qiime.org/documentation/qiime_parameters_files.html.

For example, let's assume we want to customize the OTU picking method used by (pick_otus.py) as part of the pick_de_novo_otus.py workflow, changing it from the default uclust to swarm. To find the parameter name, we can either run pick_otus.py -h or read the description at qiime.org: http://qiime.org/scripts/pick_otus.html (found by Googling QIIME pick_otus.py). We see that the parameter we need is called otu_picking_method, and that the valid options are sortmerna, mothur, trie, uclust_ref, usearch, usearch_ref, blast, usearch61, usearch61_ref, sumaclust, swarm, prefix_suffix, cdhit, uclust.

Therefore we need to add pick_otus:otu_picking_method swarm to a text file. This can be done with emacs as follows, if you have emacs on your system:

```
emacs parameters-swarm.txt
```

Then add these lines to the top of the file:

```
pick_otus:otu_picking_method swarm
pick_otus:similarity .99
```

Then save the file with ctrl-x ctrl-s, then exit with ctrl-x ctrl-c.

Now we can run the workflow script with the custom parameters file:

```
pick_de_novo_otus.py -i seqs.fna -o swarm99 -p parameters-swarm.txt
```

Making OTU tables human-readable

You can convert OTU tables to tab-delimited output using biom convert:

```
biom convert -i closedref/otu_table.biom -o closedref/otu_table.txt --to-tsv
```

The output file can now be opened in Excel.

Follow-up exercises

- 1. How can you use filter_otus_from_otu_table.py to determine the number of singleton OTUs produced by each OTU picking method?
- 2. How many OTUs remain from each method after filtering singletons?
- 3. Run OTU picking on another data set from EBI or QIITA.
- 4. Try (loading OTU tables into R)[]