hundo Documentation

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Installation

This protocol leverages the work of Bioconda and depends on conda. For complete setup of these, please see:

https://bioconda.github.io/#using-bioconda

Really, you just need to make sure conda is executable and you've set up your channels (steps 1 and 2). Then:

```
conda install python>=3.6 click \
    pyyaml snakemake>=5.1.4 biopython
pip install hundo
```

To update to the newest version of Hundo, run

pip install --upgrade hundo

Alternatively, if you do not want any new executables in your environment you can install into a new conda environment, e.g. hundo_env:

```
conda create --name hundo_env python=3 click pyyaml snakemake biopython
conda activate hundo_env
pip install hundo
```

To leave the environment:

conda deactivate

Usage

Running samples through annotation requires that input FASTQs be paired-end, named in a semi-conventional style starting sample ID, contain "_R1" (or "_r1") and "_R2" (or "_r2") index identifiers, and have an extension ".fastq" or ".fq". The files may be gzipped and end with ".gz". By default, both R1 and R2 need to be larger than 10K in size, which corresponds to around 100 reads in a compressed fastq file. This cutoff is arbitrary and can be set using --prefilter-file-size.

Using the example data of the mothur SOP located in our tests directory, we can annotate across SILVA using:

```
cd example
hundo annotate \
    --filter-adapters qc_references/adapters.fa.gz \
    --filter-contaminants qc_references/phix174.fa.gz \
    --out-dir mothur_sop_silva \
    --database-dir annotation_references \
    --reference-database silva \
    mothur_sop_data
```

The data directory can optionally be a pattern containing a wildcard, such as:

```
hundo annotate \
    --filter-adapters qc_references/adapters.fa.gz \
    --filter-contaminants qc_references/phix174.fa.gz \
    --out-dir mothur_sop_silva \
    --database-dir annotation_references \
    --reference-database silva \
    'mothur_sop_data/F3D14*S20*.fastq.gz'
```

The string must be contained between single quotes so it isn't expanded into a space delimited list.

Or when data is spread across multiple directories, you can use a combination of paths and patterns in a comma separated list, like:

```
hundo annotate \
    --filter-adapters qc_references/adapters.fa.gz \
```

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```
--filter-contaminants qc_references/phix174.fa.gz \
--out-dir mothur_sop_silva \
--database-dir annotation_references \
--reference-database silva \
'collection1/LM_*.fastq.gz,collection2/rawdata'
```

Or if you have a case where you have lots of data directories, you can specify --input-dir multiple times:

```
hundo annotate \
    --filter-adapters qc_references/adapters.fa.gz \
    --filter-contaminants qc_references/phix174.fa.gz \
    --out-dir mothur_sop_silva \
    --database-dir annotation_references \
    --reference-database silva \
    --input-dir collection2/rawdata \
    --input-dir collection3/rawdata \
    'collection1/LM_*.fastq.gz'
```

Dependencies are installed by default in the results directory defined on the command line as --out-dir. If you want to re-use dependencies across many analyses and not have to re-install each time you update the output directory, use Snakemake's --conda-prefix:

```
hundo annotate \
    --out-dir mothur_sop_silva \
    --database-dir annotation_references \
    --reference-database silva \
    mothur_sop_data \
    --conda-prefix /Users/brow015/devel/hundo/example/conda
```

Tip: In instances where compute nodes do not have access to the internet, download the reference databases and conda packages in advance.

To download the references for SILVA, run:

```
hundo download --database-dir annotation_references \
    --jobs 5 --reference-database silva
```

To download the conda environment:

```
hundo annotate \
    --out-dir mothur_sop_silva \
    --database-dir annotation_references \
    --reference-database silva \
    mothur_sop_data \
    --conda-prefix /Users/brow015/devel/hundo/example/conda \
    --create-envs-only
```

chapter $\mathbf{3}$

Annotation Parameters

Argument list, definitions and default values for hundo annotate:

Argument	Type or Choice	Description	De- fault
prefilter-	fi lNaTE siz GER	e Any FASTQ file size smaller than this in bytes is omitted from being pro- cessed.	10000
jobs	INTE- GER	Use at most this many cores in parallel. The total running tasks at any given time will be jobs divided by threads.	auto
out-dir	TEXT	Results output directory.	cur-
			rent
			di-
			rec-
			tory
no-conda		Do not use conda environments. Requires that all dependencies are in- stalled and executable.	FALS
dryrun		Do not execute anything, just show the commands that will be executed by Snakemake.	FALS
author	TEXT	Will show in footer of summary HTML document.	110.00
		arthdal aligner; <i>blast</i> is more sensitive while <i>vsearch</i> is much faster	unan blast
aligner threads	INTE-	When a step is multi-threaded, use this many threads. This is all or a subset	8
	GER	ofjobs.	
database-d	1 rTEXT	Directory containing reference data or new directory into which to down-	'ref-
		load reference data.	er- ences
filter-ada	+TEXT	File path to adapters FASTA to use for trimming read ends.	None
		s File path to FASTA to use for filtering reads.	None
		m Kiner ensistatches allowed during adapter trim process.	1
	GER		
reference-	cmleNaTEmat GER	cheigth of kmer to search against contaminant sequences.	27
reduced-km	erlNiTEn GER	Look for shorter kmers at read tips down to this length; 0 disables.	8
minimum-pa	s sNAG- re	a Rassing stingle-end read length prior to merging.	100
	GER		
minimum-ba	selNoftEali GER	t $Regions$ with average quality below this will be trimmed.	10
minimum-me	r deNTE eng GER	t Minimum allowable read length after merging.	150
allow-merg		r Allow merging of staggered reads by VSEARCH.	FALS
max-diffs	INTE-	Maximum number of different bases allowable in overlap.	5
	GER	I I I I I I I I I I I I I I I I I I I	_
min-overla	INTE-	When merging, the minimum length of overlap between reads.	16
	GER		
maximum-exp	eFdtQA4T-e	r After merging, the allowable limit of erroneous bases.	1
reference-	h liEn& Fa-	f Definera file path or set to true to use BLAST reference database.	TRU
		b When and user ing, do not create any clusters with fewer than this many	2
	GER	representative sequences.	
percent-of	- FLQAT ab	1 Maximum difference between an OTU member sequence and the repre- sentative sequence of that OTU.	3
reference-	1 d t i bit a borrege	mesite at the sequence of that of 0.	'silva
TETETEIICE	1 CL DIG KANGED CA	Unite for ITS. References will be downloaded as needed during the exe-	51174
		cution of the workflow to the location set usingdatabase-dir.	
		cFilter out alignments below this bitscore threshold and do not use them in	100
	GER	the LCA calculation.	100
blast-top-			0.95
_		ing alignment.	
read-ident	t FL-OAT u	i Wheen mapping reads back to OTU seed sectrapper 9. 9Annification Para	amete
		quire this fraction of sequence identity between sequence and reference.	0.77
min-pid	FLOAT	Minimum percent ID required from VSEARCH hits in order to be retained for LCA calculation	0.85

Annotation Output

An interactive example is available at https://pnnl.github.io/hundo/.

OTU.biom

Biom table with raw counts per sample and their associated taxonomic assignment formatted to be compatible with downstream tools like phyloseq.

OTU.fasta

Representative DNA sequences of each OTU.

OTU.tree

Newick tree representation of aligned OTU sequences.

OTU.txt

Tab-delimited text table with columns OTU ID, a column for each sample, and taxonomy assignment in the final column as a comma delimited list.

OTU_aligned.fasta

OTU sequences after alignment using MAFFT.

all-sequences.fasta

Quality-controlled, dereplicated DNA sequences of all samples. The header of each record identifies the sample of origin and the count resulting from dereplication.

blast-hits.txt

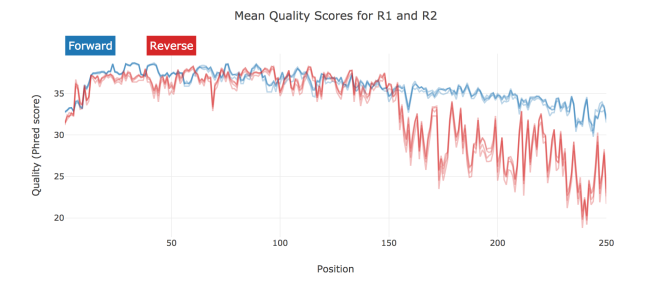
The BLAST assignments per OTU sequence.

summary.html

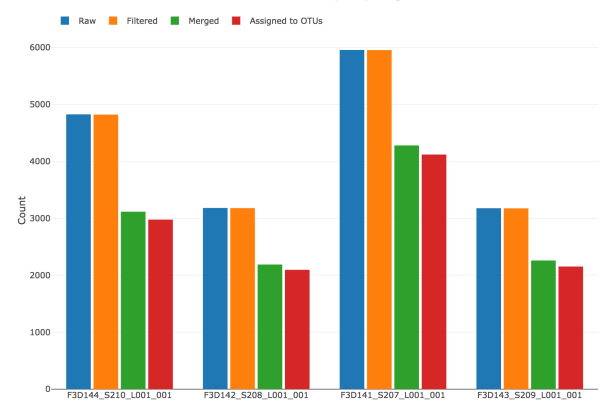
Captures and summarizes data of the experimental dataset. Things like sequence quality:

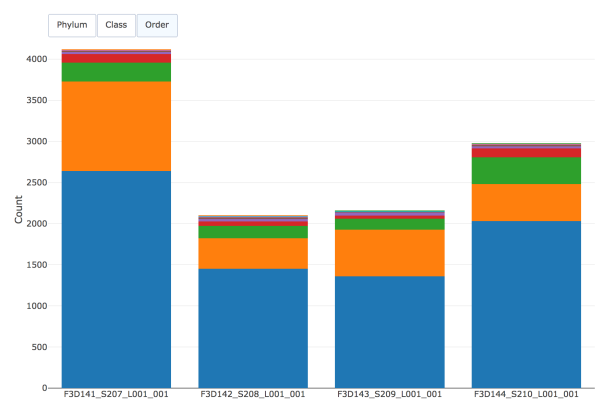
And counts per sample at varying stages of pre-processing:

Taxonomies are also summarized per sample across phylum, class, and order:



Counts Per Sample By Stage





Assigned Taxonomy Per Sample (Level: Order)

Results Example

https://pnnl.github.io/hundo/

Summary

Snakemake-based amplicon processing protocol for 16S and ITS sequences:

- Performs quality control based on quality, can trim adapters, and remove sequences matching a contaminant database;
- Handles paired-end read merging;
- Integrates de novo and reference-based chimera filtering;
- Clusters sequences and annotates using databases that are downloaded as needed;
- Generates standard outputs for these data like a newick tree, a tabular OTU table with taxonomy, and .biom.

This workflow is built using Snakemake and makes use of Bioconda to install its dependencies.