Analyzing dual-barcoded sequences from targeted amplicons

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1 Overview

This document provides instructions for the installation and execution of the *dbcAmplicons* package for the analysis of targeted genomic regions. Amplicon sequencing can be customized to very specific regions via PCR primers and dual-barcoded for a high degree of multiplexing within a single sequencing run. However, the amount of data generated as well as the complexity of the information can be an obstacle when it comes to analysis and interpretation. This package is designed to take the researcher from sequence data to figure generation in a simple pipeline.

2 Installation

Minimum requirements for the *dbcAmplicons* package: Python 2.X is required to run any and all of the elements in the package. Some applications within the package rely on FLASH or the RDP classifier to complete, so the user must determine of those external tools are necessary if the entire pipeline will not be

run.

2.1 Scripts

All executable scripts are located in the github repository dbcAmplicons.

2.2 Programs

External programs used in the pipeline:

FLASH

RDP classifier

Windows Systems

dbcAmplicons has not been tested on Windows systems, although Windows Powershell (active in XP systems and later) may be sufficient to run the analysis. It is highly recommended, however, that the user install a Linux-based virtual machine to run the pipeline.

Mac OSX

Mac users should first download and install Xcode and then download the app via a terminal by:

```
git clone https://github.com/msettles/dbcAmplicons.git
cd dbcAmplicons
python setup.py install
dbcAmplicons
```

If installation was successful, this will bring up the usage arguments.

Linux

For Linux users, some python dependencies may be required before the installation is successful. To install, open a terminal and type

```
sudo apt-get install python-dev
```

And type the sudo password at the prompt. Then, download the app via a terminal by:

```
git clone https://github.com/msettles/dbcAmplicons.git
cd dbcAmplicons
sudo python setup.py install
dbcAmplicons
```

If installation was successful, this will bring up the usage arguments.

3 Running dbcAmplicons

There are 4 levels of analysis within the *dbcAmplicons* package. The package was developed for command line usage to analyze high volumes of microbial 16S rRNA sequences. All components of the package are designed to flow directly into the next with no intermediate steps required.

```
dbcAmplicons [-h] [--version] {preprocess, join, classify, abundance}
```

$3.1 \quad dbc Amplicons \ preprocess$

dbcAmplicons preprocess takes four fastq files (forward read, index 1, reverse read, and index 2), the assigned barcode indices for each sample, and the sequences of the template-specific primers. dbcAmplicons preprocess will then assign each read in the fastq file to a specific project according to both barcode and primer sequences, and return fastq files with all of the identifying information in the read header. The syntax for usage is as follows:

dbcAmplicons preprocess [-h] -B barcodesFile [-d BARCODEDIFF]

```
[-P primerFile] [-D PRIMERDIFF] [-e PRIMEREND]
           [-S sampleFile] [-q MINQ] [-1 MINL]
           [-b BATCHSIZE] [-O PREFIX] [-U] [-u] [-v]
           [-1 read1 [read1 ...]] [-2 read2 [read2 ...]]
           [-3 read3 [read3 ...]] [-4 read4 [read4 ...]] [--test]
           [--keepPrimers] [--debug]
Options:
-h, --help Shows this help message and exits.
-B barcodesFile, --barcodes_file Barcodes file with barcodes used in the run
-d BARCODEDIFF, --integer The maximum number of mismatches allowed per barcode (default:1)
-P primerFile, --primer_file File with primer sequences
-D PRIMERDIFF, --integer The maximum number of mismatched allowed per primer (default:4)
-e PRIMEREND, --integer The required number of matching bases at end of primer (default:4)
-S sampleFile, --samples_metadata File denoting samples with primer and barcode asignments
-q MINQ, --minQ Trim 3' end of sequences to minQ (default:None)
-1 MINL, --minL If minQ is not None, only keep reads that are at least minL length (default:0)
-b BATCHSIZE, --integer Batch size in which to process reads (default:100,000)
-0 PREFIX, --output_prefix Output file basename (default:fastq_prefix)
-U, --output_unidentified Keep and output unidentified reads (default:False)
-u, --uncompressed Leave output files uncompressed (default:False)
-v, --silent Verbose output (default:False)
-1 read1 [read1 ...] Path to read 1 of an amplicon fastq four file set
-2 read2 [read2 ...] Path to read 2 of an amplicon fastq four file set
-3 read3 [read3 ...] Path to read 3 of an amplicon fastq four file set
-4 read4 [read4 ...] Path to read 4 of an amplicon fastq four file set
--test Exit after the first batch in order to test the inputs
--keepPrimers No primer trimming, leave as a part of the read (default:0)
--debug Show traceback on error
```

3.1.1 Inputs

At a minimum, dbcAmplicons preprocess requires a properly formatted reference barcode sequence file to demultiplex raw sequences. For maximum information, the following three inputs should be used, all in a tab-separated text format:

- 1. Reference barcode sequences file.
- 2. Reference primer sequences file.
- 3. Sample metadata file with sample IDs, primer IDs, and barcode IDs.

1. Reference barcode sequences file

dbcAmplicons preprocess requires a tab-separated text file with as few extraneous characters as possible. The barcode sequence reference file should have three columns: the first column is the user-defined barcode name; the second column is the Read 2 sequence, also referred to as Index 1 on the P7 end of the sequence; and the third column is the Read 3 sequence, also known as Index 2 on the P5 end of the sequence. Example format is below:

#BarcodeID	Read2	Read3
Barcode1	TAAGGCGA	TAGATCGC
Barcode2	CGTACTAG	CTCTCTAT
Barcode3	TAAGGCGA	TATCCTCT
Barcode4	CGTACTAG	AGAGTAGA
Barcode5	TAAGGCGA	GTAAGGAG

dbcAmplicons preprocess searches through the Read 2 fastq file from CASAVA for Index 1. As noted below (Figure 1), Index 2 is sequenced in reverse complement orientation relative to the P7 adaptor end of the sequence, so use of dbcAmplicons requires that the user know the orientation of the barcodes that were attached to the P7 end. Nextera kit users will often need to reverse complement their Index 1 sequences. TruSeq kit users will often not need to reverse complement their Index 1 sequences.

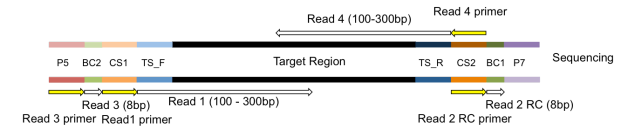


Figure 1: The orientation of sequencing reads on the Illumina Miseq. Note that Read 2 (Barcode 1) is sequenced in reverse complement orientation relative to Read 4.

2. Reference primer sequences file

The reference primer sequences file, similar to the barcode reference file, should also be a tab-separated text file. An example primer sheet is below:

# Read	Pair_ID	Primer_ID	Sequence
P5	PrimerPair1	Primer1Forward	GTAGAGTTTGATCCTGGCTCAG
P5	PrimerPair2	Primer2Forward	CGTAGAGTTTGATCATGGCTCAG
P5	PrimerPair3	Primer3Forward	ACGTAGAGTTTGATTCTGGCTCAG
P5	DegeneratePair1	Degenerate1Forward	GTGARTCATCGAATCTTTG
P5	DegeneratePair2	Degenerate2Forward	CGTGARTCATCGAATCTTTG
P7	PrimerPair1	Primer1Reverse	GTCCTCCGCTTATTGATATGC
P7	PrimerPair2	Primer2Reverse	TGTCCTCCGCTTATTGATATGC
P7	PrimerPair3	Primer3Reverse	ATGTCCTCCGCTTATTGATATGC
P7	DegeneratePair1	Degenerate1Reverse	GGGACTACHVGGGTWTCTAAT
P7	DegeneratePair2	Degenerate2Reverse	TGGGACTACHVGGGTWTCTAAT

There are 4 necessary columns in the primer reference sheet. dbcAmplicons will ignore any columns that are outside of the 4 named columns above, so the user may include additional information to each sample if necessary. Column 1 (Read) indicates the orientation of the primer. The user can denote this by the Illumina-specific adapter (P7 or P5), or the user may also use R1, R2, Read1, Read2, F, R, Forward, and Reverse. The "Pair_ID" column is the user-specified name for the primer pair that was used. The "Primer_ID" column denotes the name of each individual primer, and should correspond with the "Read" column. The "Sequence" column contains the full primer sequence without the CS-tag sequences. Note that the "Sequence" column accepts IUPAC nucleotide codes and treats each variation as a separate primer for searching.

3. Sample metadata file

The sample metadata file contains information to help split each sample into different projects along with the sample, primer, and barcode identifications. This feature used when the user wants to split out a single run's worth of data for multiple projects and can be done at several points along the *dbc Amplicons* analysis pipeline, in *dbcAmplicons classify* and also in *dbcAmplicons abundance*. The correct format for the sample metadata file is below:

SampleID	PrimerPairID	BarcodeID	ProjectID
Amp1	PrimerPair1	Barcode1	Idaho/amplicon
Amp2	PrimerPair2	Barcode2	Idaho/amplicon
Amp3	PrimerPair3	Barcode3	Idaho/amplicon
Car1	DegeneratePair1	Barcode4	Idaho/car
Car2	DegeneratePair2	Barcode5	Idaho/car

As in the primer reference sequences sheet, there are 4 columns in the sample metadata file. The "SampleID" column is the user's designation for each sample that was sequenced. The "PrimerPairID" column and the "BarcodeID" column should assign each of the primers and barcodes used to individual samples. Lastly, the "ProjectID" column denotes the output path for each sample. In the above table, sequences identified from all 5 samples will be placed in a folder named "Idaho" and from there will be divided into folders called "amplicon" and "car." If the user wants all primer pairs to be searched for all samples, dbcAmplicons preprocess will accept the regular expression "*" as a command to search all reference primers.

3.1.2 Outputs

Once dbcAmplicons preprocess has completed, a summary IdentifiedBarcodes.txt file is automatically generated within the executing directory that contains the number of reads that were assigned to each barcode and primer combination. Additionally, two fastq.gz files will be created for each user-specified output folder. Each of the fastq files will contain all of the information that was input at the beginning of dbcAmplicons preprocess for each identified read within the header of a sequence. Any primers found in the sequences are trimmed from the final output files (Figure 2).

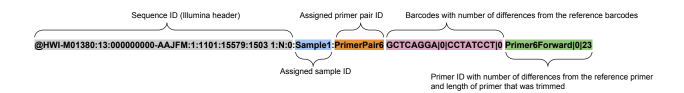


Figure 2: Example output header from $dbcAmplicons\ preprocess$ with annotations for sample ID, barcode ID and primer ID.

3.2 dbcAmplicons join

dbcAmplicons join takes the two fastq files generated from dbcAmplicons preprocess and joins them into a single long read using FLASH. The syntax is as follows:

Options:

- -h, --help Show this help message and exit
- -O PREFIX, --output_path Path for output files (default:joined)
- -u, --uncompressed Leave output files uncompressed (default:False)
- -x NUM, --max-mismatch-density Maximum allowed ratio between the number of mismatched base

pairs and the overlap length. Two reads will not be combined with a given overlap if that overlap results in a mismatched base density higher than this value. Note: Any occurrence of an "N" in either read is ignored and not counted towards the mismatches or overlap length. (default:0.25)

- -t NTHREADS, --integer Set the number of worker threads. (default:1)
- -p path, --flash_path Path to the application FLASH (default: FLASH)
- -v, --verbose Verbose output (default:False)
- -1 read1 Path to read1 of an amplicon fastq (or fastq.gz) two file set
- -2 read2 Path to read2 of an amplicon fastq (or fastq.gz) two file set

3.2.1 Outputs

There are 5 output files from dbcAmplicons join. The successfully combined sequences will be in a fastq file with an .extendedFrags.fastq.gz extension. Any sequences that were not combined will be in the fastq files with the .notCombined_1/2 extension. Two text files, joined.hist and joined.histogram list the number of sequences and total lengths of the combined reads; joined.hist by integer, and joined.histogram graphically. For more information about FLASH output, please refer to the developer site.

$3.3 \quad dbc Amplicons \ classify$

The current iteration of *dbcAmplicons classify* uses the RDP classifier for fungal LSU, bacterial 16S, and archaeal 16S to identify sequences. Normally, the RDP classifier does not take single reads that Usage is as follows:

Options:

- -h, --help Show this help message and exit.
- -b BATCHSIZE, --integer Batch size for read processing (default:100,000)
- -q MINQ, --minQ Trim 3' end of sequences to minQ (paired-end reads only) (default:None)
- -1 MINL, --minL If minQ is not None, only keep reads that are at least minL length (paired-end reads only) (default:0)
- -p PROCS, --integer Number of processors to use (default:1)
- --rdpPath Path to the RDP classifier
- -g <arg>, --arg Choice of classification between 16S microbial rRNA or fungal LSU (default:16SrRNA)
- -S sampleFile, --sample_metadata Sample metadata file with primer and barcode designations
- -O outputPrefix, --output_path Path for output files (default:classify)
- -1 read1 Read 1 of an amplicon fastq two file set
- -2 read2 Read 2 of an amplicon fastq two file set
- -U single-end amplicon Output from dbcAmplicons join or other joined paired reads
- -v, --silent Verbose output (default:False)
- --debug Show traceback on error

3.3.1 Outputs

For more information on the RDP classifier and its classification algorithm, please refer to the RDP Wiki. The RDP classifier within *dbcAmplicons classify* outputs a single file with each sequence classified according to its closest identified taxonomic rank using bootstrap values for confidence. The read is classified within the most specific taxonomic rank that meets the boostrap threshold of 50%.

$3.4 \quad dbc Amplicons \ abundance$

dbcAmplicons abundance takes the output from dbcAmplicons classify and builds abundance tables from the information. Usage:

Options:

- -h, --help Show this help message and exit
- -r <arg>, --rank Taxonomic rank from which to build table, allowable values: domain, phylum, class, order, family, genus, and species (if performed), (default:genus)
- -t VALUE, --integer Threshold bootstrap value to use for assignment, first taxon level greater than threshold
- -S FILE, --sample_metadata File with primers
- -O FILE_PREFIX, --output_path Path for output files (default:table)
- -F FILE [FILE ...] Fixrank formatted classification file generated by dbcAmplicons classify
- -v, --silent Verbose output (default:False)
- --debug Show traceback on error

3.4.1 Outputs

dbcAmplicons abundance will generate three .txt files. The abundance.txt file lists the taxon abundance with the read counts. The proportions.txt lists the proportions of reads per sample, and the taxa_counts.txt lists a summary of the lowest level classification and the number of counts for that level.

4 Supplementary Scripts

4.1 reduce_amplicons

reduce_amplicons

Options:

- -h, --help Show this help message and exit
- -p PROGRAM Comma separated list of the functions of consensus, ambiguities, occurrence (default: consensus)
- -s MIN-SEQ, --integer Minimum number of reads (default:5)
- -f MIN-FREQ, --frequency Minimum frequecy of reads (default:0.05)
- --trim-1=TRIM-1 Number of bases to trim from the end of read 1 (default:0)
- --trim-2=TRIM-2 Number of bases to trim from the end of read 2 (default:0)
- -r, --reuse Reuse the reads within the folder (default:False)
- -o OUTPUTDIR, --outputDir Directory to place output files (default:[basename].reduced)
- -c CPUS, --integer Number of processors to use; choose 0 to use all available cores (default:0)

4.2 convert2ReadTo4Read

A script for back-coverting two Illumina sequence files that have been processed for their barcodes back to a four read set to be reprocessed using dbcAmplicons. Usage:

```
convert2ReadTo4Read.py [-h] [--version] [-b BATCHSIZE]
      [-0 PREFIX] [-u] [-v] [-1 read1 [read1 ...]]
      [-2 read2 [read2 ...]] [--debug]
```

Options:

- -h, -help Show this help message and exit
- -version Show program's version number and exit
- -b BATCHSIZE, -batchsize Batch size for read processing (default:100,000)
- -O PREFIX, -output_path Path for output files (default:None)
- -u, -uncompressed Leave output files uncompressed (default:False)
- -v, -silent Verbose output (default:False)
- -1 read1 [read1 ...] Read 1 of an amplicon fastq two file set
- -2 read2 [read2 ...] Read2 of an amplicon fastq two file set
- -debug Show traceback on error

5 References