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## MetAmp: a tool for Meta-Amplicon analysis User Manual

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# 1 Description

MetAmp tool was developed to analyze microbial amplicon data by combining several marker regions from 16S rRNA gene. Such marker regions serve as unique identifications for species. There are nine variable regions in total in bacterial 16S rRNA gene and even more marker regions, because marker region can contain itself several variable regions.

MetAmp was developed by Ilya Y. Zhbannikov (ilyaz@uidaho.edu, i.zhbannikov@mail.ru), James A. Foster (foster@uidaho.edu) and Janet Williams (janetw@uidaho.edu).

# 2 Installation

MetAmp is easy to install. However, you must have R ([www.r-project.org](http://www.r-project.org)), Python ([www.python.org](http://www.python.org)) and GCC (<https://gcc.gnu.org/>).

1. Download or clone MetAmp from our GitHub repository (<http://github.com/izhbannikov/MetAmp>), for example:

```
git clone http://github.com/izhbannikov/MetAmp
```

2. In Terminal: `cd` to the MetAmp directory and execute Makefile: `$make`. In some cases you have to be a system administrator, so you need to compile MetAmp with the following command: `$sudo make`. First, the installation script will check for required tools: R, Python and gcc. Then it builds and installs required packages and data for evaluation. You can also run `make install` to avoid checking stage.

3. Installation summary:

```
$git clone http://github.com/izhbannikov/MetAmp
$cd MetAmp/
$make
```

### 3 Quick start

You have to provide reference libraries, your amplicon data and output directory where all analysis results will be stored:

One marker region:

```
python metamp.py -r data/gold21/gold21.fasta \  
                  -r1 data/gold21/gold21_V13V31.fasta \  
                  -l1 data/even/SRR072220_V13V31_relabeled.fastq \  
                  -o test
```

Here:

- `metamp.py` - program name.
- `-r data/gold21/gold21.fasta` - passing the reference file, that contains whole 16S sequences, in forward and reverse complement.
- `-r1 data/gold/gold21_V13V31.fasta` - passing the reference file that contains marker (region V1-3) sequences extracted from `gold21.fasta` (whole 16S), in forward and reverse complement.
- `-l1 data/even/SRR072220_V13V31_relabeled.fastq` - amplicon empirical reads. "relabeled" means that a barcode sequences were attached to each read label.
- `-o test` - output directory with all analysis results.

Three variable regions:

```
python metamp.py -r data/gold21/gold21.fasta \  
                  -r1 data/gold21/gold21_V13V31.fasta \  
                  -l1 data/even/SRR072220_V13V31_relabeled.fastq \  
                  -o test
```

```

-l1 data/even/SRR072220_V13V31_relabeled.fastq \
-r2 data/gold21/gold21_V35V53.fasta \
-l2 data/even/SRR072220_V35V53_relabeled.fastq \
-r3 data/gold21/gold21_V69V96.fasta \
-l3 data/even/SRR072239_V69V96_relabeled.fastq \
-o test

```

In this case:

- `-r2 data/gold/gold21_V35V53.fasta` - a reference file that contains marker (region V3-5) sequences extracted from `gold21.fasta` (whole 16S), in forward and reverse complement.
- `-l2 data/even/SRR072220_V35V53_relabeled.fastq` - amplicon empirical reads.
- `-r3 data/gold/gold21_V69V96.fasta` - a reference file that contains marker (region V3-5) sequences extracted from `gold21.fasta` (whole 16S), in forward and reverse complement.
- `-l3 data/even/SRR072239_V69V96_relabeled.fastq` - amplicon empirical reads.

To get help from the program, you can use `python metamp.py -h`:

```

$python metamp.py -h
usage: metamp.py -o [--output] -r [--ref] <reference 16S seqs> \
        -r1 [--ref1] <reference marker seqs> \
        -l1 [--lib1] <your amplicon library> \

```

[options]

optional arguments:

-h, --help                    show this help message and exit

-o OUTPUT, --output OUTPUT  
                              Output directory.

-r REF16S, --ref REF16S  
                              Output file with keywords.

-r1 REF1, --ref1 REF1  
                              Reference marker sequences for marker type #1

-r2 REF2, --ref2 REF2  
                              Reference marker sequences for marker type #2

-r3 REF3, --ref3 REF3  
                              Reference marker sequences for marker type #3

-r4 REF4, --ref4 REF4  
                              Reference marker sequences for marker type #4

-r5 REF5, --ref5 REF5  
                              Reference marker sequences for marker type #5

-r6 REF6, --ref6 REF6  
                              Reference marker sequences for marker type #6

-r7 REF7, --ref7 REF7  
                              Reference marker sequences for marker type #7

-r8 REF8, --ref8 REF8  
                              Reference marker sequences for marker type #8

-r9 REF9, --ref9 REF9

```

Reference marker sequences for marker type #9
-11 LIB1, --lib1 LIB1
    Amplicon sequences for marker #1
-12 LIB2, --lib2 LIB2
    Amplicon sequences for marker #2
-13 LIB3, --lib3 LIB3
    Amplicon sequences for marker #3
-14 LIB4, --lib4 LIB4
    Amplicon sequences for marker #4
-15 LIB5, --lib5 LIB5
    Amplicon sequences for marker #5
-16 LIB6, --lib6 LIB6
    Amplicon sequences for marker #6
-17 LIB7, --lib7 LIB7
    Amplicon sequences for marker #7
-18 LIB8, --lib8 LIB8
    Amplicon sequences for marker #8
-19 LIB9, --lib9 LIB9
    Amplicon sequences for marker #9
-qual QUAL, --qual QUAL
    Quality score threshold (default=15)
-minlen MINLEN, --minlen MINLEN
    Minimum read length (default=250)

```

### 3.1 Test data

The following test data sets were used:

- data/even/
- data/staggered/

The even and staggered are the Human mock community pyrosequence data (SRX021555), even and staggered community, 21 microbial species and one eucariotic species.

### 3.2 Results

#### 3.2.1 Clustering results

By default it is `clusters.clstr` but the name can be changed (by changing `clust_filename` parameter in `config.R` file). This file has the uc format and described at <http://drive5.com/usearch/manual/ucout.html>. Below we provide an example and copy the format description from [www.drive5.com](http://www.drive5.com):

H	0	250	99.6	+	0	0	250M	read11;barcodelabel=TCAG;	OTU_1
H	1	250	100.0	+	0	0	250M	read6;barcodelabel=TCAG;	OTU_2
H	1	250	100.0	+	0	0	250M	read12;barcodelabel=TCAG;	OTU_2
H	5	250	100.0	+	0	0	250M	read7;barcodelabel=TCAG;	OTU_4
H	22	250	98.8	+	0	0	250M	read22;barcodelabel=TCAG;	OTU_20

Description of uc format is directly taked from <http://drive5.com/usearch/manual/ucout.html>

Field	Description
1	Record type S, H, C or N (see table below).
2	Cluster number (0-based).
3	Sequence length (S, N and H) or cluster size (C).
4	For H records, percent identity with target.
5	For H records, the strand: + or - for nucleotides, . for proteins.
6	Not used
7	Not used
8	Compressed alignment or the symbol '=' (equals sign).
9	Label of query sequence (always present).
10	Label of target sequence (H records only).

Record	Description
H	Hit. Represents a query-target alignment
S	Centroid (clustering only)
C	Cluster record (clustering only)
N	No hit

### 3.2.2 OTU table

An OTU table, where rows are OTUs and columns are barcodes (by default - `otu_table.txt`).

Example:

```
OTUId    TCAG
OTU_8    123
OTU_9    158
OTU_2    341
...      ....
```

### 3.2.3 Coordinates

A text file that contain coordinates for each point, including reference points (by default - `coordinates.crd`)

### 3.2.4 Log

A text file containing all messages during the analysis process (by default - `log.txt`)

## 4 MetAmp workflow

In the following sections we present key stages of our meta-amplicon analysis pipeline.

### 4.1 Input data

Input data can be any of the following NGS libraries: Roche 454, Ion Torrent, Illumina.



Roche 454 and Ion Torrent provide single-end sequences in special binary Sequence Flowgram Format (SFF). Before using SFF files you have to convert them into FASTQ format.

Illumina paired-end reads should be merged into longer single-end sequences.

**Important!** - Before analysis, you may need to perform re-labeling of your read headers: a read header should contain `barcodeLabel` (see more at [www.drive5.com](http://www.drive5.com)), for example:

```
@read1;barcodeLabel=TCAG;  
GATGAACGCTGGCGGCGTGCCTAATACATGCAAGT...AT  
+  
IIIAAAIIIIIIIIIIIIIIHHHIAAAAAAAAAA...II
```

If you use Illumina sequence data, you also have to merge overlapping paired-end reads. Later I will provide the scripts that can do these things above, but for now you can simply use scripts provided at [www.drive5.com](http://www.drive5.com).

## 4.2 Data denoising

Data denoising is necessary for downstream analysis because it removes the majority of 'foreign' nucleotides, such as barcodes and primers, and low-quality regions. We suggest you to use SeqyClean tool for this operation, since it is the most comprehensive tool for sequence denoising. You can download SeqyClean here: <https://bitbucket.org/izhbannikov/seqyclean>. Using SeqyClean is simple and straightforward. Nevertheless, we provided a special script that does all the data de-noising.

## 4.3 Analysis workflow

1. Building a reference topology of microbial populations, where pairwise distances are computed from applying pairwise alignment of complete reference 16S gene sequences from

RDP database, see Figure 1(a), top plane. We use the percent sequence identity in order to compute the distances between sequences.

2. Computing a guide "amplicon" topology for each reference 16S sequence. To do this, we extract marker regions (for example V1- V3 and V6-V9) from reference complete 16S sequences ("reference points") and place them on to the plane with the same methodology that was used for building the reference topology of complete 16S (see Figure 1(a), bottom plane, hollow green circles).
3. add empirical amplicon sequences (in fact, amplicon consensus sequences), obtained through amplicon sequencing of microbial data (Figure 1b, bottom, filled blue circles). This topology is the same empirical topology as if the empirical data would be clustered with existing methods
4. Normalized empirical topologies are formed foreach marker through mapping of each empirical point back to the reference 16S sequence topology. Mapping between the reference topology and guide "amplicon" topology is achieved with affine transformation. Guide sequences are so match with the corresponding reference 16S sequences, from where they were obtained from (see Figure 1a, top, solid green circles), and, in turn, carry the empirical amplicon sequences (or consensus amplicon sequences) back to the 16S plane.
5. Repeating stages 2-4 for each variable region.
6. Clustering and building OTUs.
7. Final statistics.

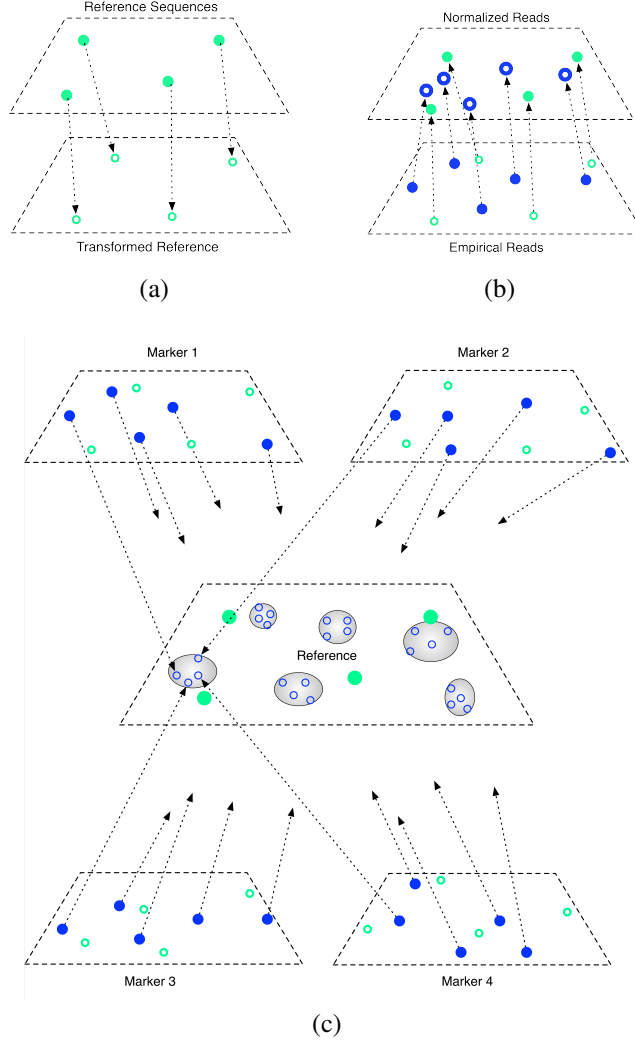


Figure 1: Illustration of “meta-amplicon” analysis algorithm. (a) Computing reference (top plane) and guide (bottom plane) topologies. Pairwise sequence dissimilarity is used to compute distances between reference sequences. The same approach is used for guide sequences, originated from corresponding whole 16S gene sequences. Reference and guide points are then placed on to the planes with multidimensional scaling. (b) The empirical amplicon sequences (hollow blue circles) are placed on to the bottom plane along with the guide circles (hollow green) circles and with the same metric. Such guide sequences are then mapped back to the reference 16S plane, carrying empirical sequences (solid blue circles). (c) Then this is repeated for each of marker (variable) region.

## 4.4 Output files

- Clustering results (`*.clstr`) - by default it is `clusters.clstr` but the name can be changed (by editing `clust_filename` parameter in `config.R` file)
- An OTU table, where rows are OTUs and columns are barcodes (by default - `otu_table.txt`).
- A text file that contain coordinates for each point, including reference points (by default - `coordinates.crd`)
- A text file containing all messages during the analysis process (by default - `log.txt`)

## 5 Provided data

We provided some data that can be used in your projects. For evaluation purposes we provided small data set. Description of provided data is given below:

- gold folder contains 'Gold Dataset' (`gold.fa`) containing the ChimeraSlayer reference database in the Broad Microbiome Utilities (<http://microbiomeutil.sourceforge.net/>), version `microbiomeutil-r20110519`. This contains more than 5000 sequences in both orientations (forward and reverse-complemented.).
- In addition, we supply a dataset that contains 1500 microbial 16S sequences extracted from the 'Gold Dataset'.
- Evaluation data set contains a small set of reference sequences (21 sequences) and empirical amplicon sequences (`data/even.zip` and `data/staggered.zip`). Amplicon sequences contain sequence data from Human Mock Community. More information provided here: <http://www.hmpdacc.org/HMMC/>

Evaluation:

Name	Description
even	Even community: SRR053818, SRR072220, SRR072239
staggered	Staggered community: SRR072221, SRR072223, SRR072237
gold21.fasta	contains 21 reference 16S sequences
gold21_V13V31.fasta	contains 21 reference marker sequences (1-3 regions)
gold21_V35V53.fasta	contains 21 reference marker sequences (3-5 regions)
gold21_V69V96.fasta	contains 21 reference marker sequences (6-9 regions)

The even and staggered are the Human mock community (HMP) pyrosequence data (SRX021555: <http://www.ncbi.nlm.nih.gov/sra?term=SRX021555>). Detailed description of these datasets (and sequencing protocols) is under the following link: <http://www.hmpdacc.org/HMMC/>

#### 5.0.1 Directory map and descriptions of folders

```
MetAmp
├── data
│   ├── gold21
│   ├── gold1500
│   └── gold5000
├── src
├── Packages
│   └── blastparser
├── bin
└── python_scripts
```

**data** - contains reference sequences, evaluation data and other data that can potentially be useful

**data/gold** - contains 'gold' reference sequences from Broad Institute. **data/gold1500** - contains 1500 'gold' reference sequences extracted from the previous 'gold' dataset. **data/gold21** - contains 21 'gold' reference sequences extracted from the previous 'gold' dataset. This dataset was used to evaluate MetAmp **src** - contains source files.

**Packages** - contains installation files.

**Packages/blastparser** - a parser for blastn files.

**bin** - binary files (alignment and clustering programs).

**python\_scripts** - Python scripts from [www.drive5.com](http://www.drive5.com).

## 6 Parameters

You can adjust parameters in file `config.R`. It contains parameters such as path to temporary directory, names of the output files, etc. Below we describe those parameters in details.

### 6.1 `config.R`

**usearch** and **usearch7** - paths to alignment tool (we use USEARCH, see [www.drive5.com](http://www.drive5.com))

**R\_LIBS** - Path to installed BLASTParser package (BLASTParser parses output files from USEARCH and creates distance matrices)

**clust\_filename** - Filename of file with final clusters

**otu\_table\_filename** - Filename of final OTU table

**coord\_filename** - Filename of output file with NMDS coordinates of each reference and empirical point (see 'Analysis workflow')

**chime\_ref** - Chimeric reference database

**tmp\_dir** - A directory that contains temporary files

**keep\_tmp\_files** - Keep or not temporary files

## 7 Acknowledgements

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