

An Introduction to *OTUbase*

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```
> library("OTUbase")
```

The *OTUbase* package provides an organized structure for OTU (Operational Taxonomic Unit) data analysis. In addition, it provides a similar structure for general read-taxonomy classification type data. *OTUbase* provides some basic functions to analyze the data as well.

1 A simple workflow

This section walks through a simple workflow using a small example dataset. It demonstrates the main features of *OTUbase*. The data used for this example comes from a dataset described in "Microbial diversity in the deep sea and the underexplored 'rare biosphere'" by Sogin et al. (PNAS 2006). The complete dataset is available through PNAS. A random set of 1000 sequences was taken from this dataset.

1.1 Sample meta data

Sample metadata is collected along with the sample. This data may include any number of different pieces of information about the sample. In the example dataset, the meta data is provided in Table 1 of Sogin et al. This file is named 'sample_metadata.txt'. To be easily read by *OTUbase*, this file is in the form of an `AnnotatedDataFrame`.

1.2 Sequence preprocessing

This section describes the preprocessing steps necessary to generate the files used by *OTUbase*.

1.2.1 Sequence trimming and filtering

Many OTU data projects will begin with raw sequence reads from a next generation pyrosequencer. These reads may include primers, barcodes, and/or adapters that are not part of the actual read. The first step in the analysis pipeline is to trim the primers and barcodes from the read. A number of tools

are able to do this. Commands in Mothur are 'trim.seqs()' and 'filter.seqs()'. For this workflow we are assuming that these steps have already been done. When the barcodes are trimmed off the reads, a separate file is generally created that links the read with a sample identification. Mothur creates this file automatically and gives it a '.groups' extension. Each line of this file contains a read ID and the ID of the sample the read belongs to, separated by a tab. OTUbase requires this groups file.

1.3 Taxonomic classification and OTU generation

There are two main approaches used to analyse amplicon data. An OTU approach involves first clustering the sequences together by similarity into OTUs or Operational Taxonomic Units. These OTUs can then be used in richness calculations and in comparing two samples. An alternate approach attempts to classify each sequence into an existing taxonomy. The RDP classifier, for example, uses a Markov model to sort sequences into genus level classifications.

The data produced by these two approaches is slightly different. The OTU approach results in a list of sequences belonging to each OTU. The classification approach results in each sequence having a classification. OTUbase is able to use either of these types of data.

The data processing involved in OTU generation is described by Pat Schloss on the Mothur web site. Those interested can find the OTU generation steps for Sogin's data at .

The data processing involved in the RDP taxonomic classification is somewhat simpler and less computationally demanding. Details can be found on the RDP website .

1.3.1 Reducing the dataset to unique sequences

To decrease the computation time involved in both techniques, duplicate sequences in the dataset are removed first. These sequences can then be added back in after the processing is complete. Mothur removes the duplicate sequences with the command 'unique.seqs()' which also automatically generates a file that keeps track of which sequences have duplicates (called a name file).

In this workflow the duplicate sequences have been removed using Mothur. The name file is called 'sogin.names'

1.4 Importing files into an OTUbase object

OTUbase is able to automatically import a number of files generated during the data processing. These files include the sample file (the group file produced by Mothur), the OTU file (the list file produced by Mothur), and the meta data (in AnnotatedDataFrame format). In addition OTUbase inherits *ShortRead* which allows the user to include a fasta file and a quality file. OTUbase also recognizes the RDP taxonomic classification files that are in the 'fixed' format.

```
> dirPath <- system.file("extdata/Sogin_2006", package="OTUbase")
```

Usually `dirPath` will be the directory path containing the files that will be read by OTUbase.

Because there are two main approaches to data analysis (OTU and Taxonomic classification) we will look at both in parallel. To read in OTU related data the function `readOTUset()` is used. Likewise, to read in classification data the function `readTAXset()` is used.

```
> soginOTU <- readOTUset(dirPath=dirPath, level="0.03", samplefile="sogin.groups", fastafilename=fastafilename)
> soginOTU
```

```
Class: OTUsetF
Number of Sequences: 1000 reads
Sequence Width: 56..100 cycles
Number of OTUs: 399
Number of Samples: 8
sampleData: T      ncol: 6
assignmentData: F
```

The level is the OTU classification level desired (many clustering levels may be present in one otufilename). The default is '0.03'. The samplefile connects the read ID to the sample it belongs to. The fastafilename and the associated quality file are optional. Their inclusion may make the reading of the data significantly slower. The otufilename must be in Mothur format. The sampleADF is the sample meta data file.

```
> soginTAX <- readTAXset(dirPath=dirPath, fastafilename='sogin.fasta', sampleADF='sample_metadata.txt')
> soginTAX
```

```
Class: TAXsetF
Number of Sequences: 1000 reads
Sequence Width: 56..100 cycles
Number of Samples: 8
sampleData: T      ncol: 6
assignmentData: F
```

The `readTAXset` function only differs from the `readOTUset` function slightly. Notably different is the absence of an otufilename and the presence of a taxfile (in this case the RDP fixed output). Also included in the `readTAXset` function is the namefile. This file is the Mothur names file and should be included when the dataset has been reduced to unique sequences.

1.5 Accessing data in OTUbase objects

OTUbase provides a number of accessor functions that allow the user to easily access the data contained in the OTUbase object. `sread`, `quality`, and `id` are inherited from the *ShortRead* package and allow access to the sequence, the quality, and the sequence id. In addition, `sampleID`, `sData`, and `aData` provide access to the sample ID, the sample meta data, and the assignment meta data respectively (when available).

```

> head(id(soginOTU))

BStringSet object of length 6:
      width seq
[1]    14 D4WT9DQ06DVGFR
[2]    14 D4WT9DQ05C6YNI
[3]    14 D4WT9DQ12HNYQ2
[4]    14 D4WT9DQ01APOUQ
[5]    14 D4WT9DQ09FLPTJ
[6]    14 D27LU0R02A82DK

> head(sread(soginOTU))

DNAStringSet object of length 6:
      width seq
[1]    60 TGCCTTTGACATCCTCGGAACGGT...GGTGCCTTCGGGAACCGAGAGAC
[2]    71 TGGACTTGACATGTTAGTGTA AAC...AGCTTGCTCAAAGACACTATCAC
[3]    58 CGGGCTTGAAGTGCAAGCGACAAC...GATTTCCGCAAGGACGCTTGTAG
[4]    64 TGGTCTTGACATCCCGGGAATCTC...CCTCATTAGAGGAGCCTGGTGAC
[5]    60 AGGACTTGACATCCAGAGAACTCG...GGTGCCTTCGGGAACCTCTGTGAC
[6]    59 ATCCCTTGACATCCTGCGAACTTT...TGGTGCCTTCGGGAACGCAGTGAC

> head(sampleID(soginOTU))

[1] "53R"  "53R"  "115R" "FS312" "112R" "FS312"

> head(sData(soginOTU))

      Site Lat_N Long_W Depth Temperature
53R  Labrador seawater  58.3 -29.133 1,400      3.5
55R   Oxygen minimum  58.3 -29.133   500      7.1
112R Lower deep water  50.4 -25.000 4,121      2.3
115R   Oxygen minimum  50.4 -25.000   550      7.0
137  Labrador seawater  60.9 -38.516 1,710      3.0
138  Labrador seawater  60.9 -38.516   710      3.5

      Cells
53R  6.4 × 104
55R  1.8 × 105
112R 3.9 × 104
115R 1.5 × 105
137  3.3 × 104
138  5.2 × 104

```

There are a couple accessors specific to OTUset or TAXset. To access the OTU IDs stored in OTUset objects, `otuID` is used. Likewise, to access the taxonomic classifications stored in TAXset objects, `tax` is used.

```

> head(otuID(soginOTU))

```

```
[1] "otu221" "otu250" "otu116" "otu385" "otu59" "otu95"
```

```
> head(tax(soginTAX))
```

```

  root root_score  domain domain_score  phylum
1 Root          1.0 Bacteria          0.91 Proteobacteria
2 Root          1.0 Bacteria          0.91 Actinobacteria
3 Root          1.0 Bacteria          0.96 Actinobacteria
4 Root          1.0 Bacteria          1.00 Proteobacteria
5 Root          1.0 Bacteria          1.00 Proteobacteria
6 Root          1.0 Bacteria          1.00 Proteobacteria
  phylum_score      class class_score
1          0.62 Gammaproteobacteria    0.46
2          0.10      Actinobacteria    0.10
3          0.16      Actinobacteria    0.16
4          1.00 Deltaproteobacteria    1.00
5          0.98 Gammaproteobacteria    0.97
6          1.00 Gammaproteobacteria    1.00
  order order_score      family
1 Oceanospirillales    0.07  Halomonadaceae
2 Bifidobacteriales    0.04 Bifidobacteriaceae
3 Bifidobacteriales    0.06 Bifidobacteriaceae
4 Desulfobacterales    1.00  Desulfobulbaceae
5 Oceanospirillales    0.54 Oceanospirillaceae
6 Thiotrichales        1.00  Francisellaceae
  family_score      genus genus_score
1          0.06 Modicisalibacter    0.03
2          0.04  Metascardovia      0.04
3          0.06  Parascardovia      0.02
4          1.00  Desulfocapsa      1.00
5          0.51 Oceanospirillum    0.44
6          0.63   Francisella      0.63

```

1.6 First data analysis steps

Now that the data is in the OTUbase object, we can now generate tables and figures that help analyze it. One of the first steps in many analyses is the generation of an abundance table. There is an OTUbase method that does this.

```
> abundOTU <- abundance(soginOTU, weighted=F, collab='Site')
> head(abundOTU)
```

```

      s
o      Lower deep water Oxygen minimum Labrador seawater
otu1      0          0          2
otu10     2          0          0
otu100    1          0          0

```

```

otu101          0          0          0
otu102          0          0          0
otu103          0          0          0
s
o      Labrador seawater Labrador seawater Oxygen minimum
otu1          1          1          0
otu10         0          0          0
otu100        0          0          0
otu101        0          0          0
otu102        1          0          0
otu103        0          0          0
s
o      Bag City Marker 52
otu1          0          0
otu10         0          0
otu100        1          0
otu101        2          0
otu102        0          0
otu103        1          2

> abundTAX <- abundance(soginTAX, weighted=F, taxCol='genus', collab='Site')
> head(abundTAX)

s
o      Lower deep water Oxygen minimum
Abiotrophia          0          0
Acetivibrio          0          0
Acinetobacter        0          0
Actibacter           0          0
Aestuariicola        0          0
Agromonas            0          0
s
o      Labrador seawater Labrador seawater
Abiotrophia          0          1
Acetivibrio          0          0
Acinetobacter        0          1
Actibacter           0          0
Aestuariicola        0          0
Agromonas            0          0
s
o      Labrador seawater Oxygen minimum Bag City
Abiotrophia          0          0          0
Acetivibrio          0          0          1
Acinetobacter        0          0          0
Actibacter           0          0          1
Aestuariicola        0          0          1

```

Agromonas	1	0	0
s			
o	Marker 52		
Abiotrophia	0		
Acetivibrio	0		
Acinetobacter	0		
Actibacter	3		
Aestuariicola	0		
Agromonas	4		

It should be noted that the abundance method for TAXset objects requires one extra piece of information, the column of the classification desired. The abundance can be generated from any of them (genus, family, etc). Other options are also available in the abundance methods. For example, the abundance can be generated based on any column in the assignment data. For more on the abundance method please see the help documentation.

One of the strengths of OTUbase is that by being in the R environment it can take advantage of a number of available data analysis packages. One of these packages is *vegan*. *vegan* is an R package that provides many tools to analyze ecological type data. It includes diversity estimation and cluster analysis.

Using the functions provided by *vegan* and the abundance table previously generated:

```
> estrichOTU <- apply(abundOTU, 2, estimateR)
> estrichOTU
```

s			
	Lower deep water	Oxygen minimum	
S.obs	57.000000	44.000000	
S.chao1	192.125000	176.000000	
se.chao1	60.975679	74.630319	
S.ACE	283.067602	154.791782	
se.ACE	7.841178	8.026768	
s			
	Labrador seawater	Labrador seawater	
S.obs	49.000000	48.000000	
S.chao1	254.000000	159.42857	
se.chao1	110.794024	53.88188	
S.ACE	230.006548	320.32000	
se.ACE	5.532269	11.21429	
s			
	Labrador seawater	Oxygen minimum	Bag City
S.obs	37.000000	29.000000	110.00000
S.chao1	145.750000	191.500000	384.61538
se.chao1	62.860021	104.644533	96.01630
S.ACE	147.625000	182.685606	579.22305
se.ACE	3.619804	2.553564	16.82521

```

s
  Marker 52
S.obs      130.00000
S.chao1    308.00000
se.chao1   54.70480
S.ACE      392.65701
se.ACE     12.75554

> estrichTAX <- apply(abundTAX, 2, estimateR)
> estrichTAX

```

```

s
  Lower deep water Oxygen minimum
S.obs           48.00000    34.000000
S.chao1         118.00000    64.000000
se.chao1        33.36542    17.882622
S.ACE           202.70270    77.238156
se.ACE          10.26136     5.658764

```

```

s
  Labrador seawater Labrador seawater
S.obs           41.00000    35.000000
S.chao1         173.00000   122.750000
se.chao1        74.62941    52.070518
S.ACE           167.54231   132.763430
se.ACE           4.56640     5.203785

```

```

s
  Labrador seawater Oxygen minimum Bag City
S.obs           29.000000    22.000000  91.000000
S.chao1         67.000000   107.500000 253.750000
se.chao1        23.913688    59.310721  62.402381
S.ACE           98.141354   194.379259 290.053022
se.ACE          7.521336     9.549327   9.670711

```

```

s
  Marker 52
S.obs           88.000000
S.chao1        157.789474
se.chao1       26.353821
S.ACE          190.277732
se.ACE          8.276204

```

The vegan function `vegdist` and `hclust` have been combined into one OTUbase wrapper for convenience. This allows the user to quickly cluster the samples. This clustering can be done using a number of different distance and clustering methods.

```
> clusterSamples(soginOTU, distmethod='jaccard', clustermethod='complete', collab='Site')
```

```
Call:
hclust(d = d, method = clustermethod)
```

```
Cluster method : complete
Distance       : jaccard
Number of objects: 8
```

```
> clusterSamples(soginTAX, taxCol='genus', distmethod='jaccard', clustermethod='complete', c
```

```
Call:
hclust(d = d, method = clustermethod)
```

```
Cluster method : complete
Distance       : jaccard
Number of objects: 8
```

The user is encouraged to explore many functions available through *vegan* and other R packages. Commonly useful ones can then be brought into OTUbase to make their use more efficient.

2 Advanced features

A number of other functions are available. While the implementation is incomplete, `subOTUset()` is a function that allows the user to extract any OTUs or samples from the dataset to be analyzed separately. This makes it possible to remove one or more OTUs or samples from the analysis. Eventually this will be implemented using the more traditional '[' notation.

```
> soginReduced <- subOTUset(soginOTU, samples=c("137", "138", "53R", "55R"))
> soginReduced
```

```
Class: OTUsetF
Number of Sequences: 222 reads
Sequence Width: 56..100 cycles
Number of OTUs: 127
Number of Samples: 4
sampleData: T      ncol: 6
assignmentData: F
```

3 The structure of an OTUbase object

OTUbase objects include a number of possible slots. Inherited from *ShortRead* are `sread`, `id`, and `quality`. These slots, along with `otuID`, `tax`, and `sampleID` are all of identical length and order. For example, the first row in the `id` slot is connected to the first rows in the `sread`, `quality`, `otuID`, and `sampleID` slots. In other words, the first `id` represents the first sequence that has a quality described

by the first row of the quality slot; it is a member of the otu listed in the otuID slot and a member of the sample listed in the first row of the sample slot.

In addition there are two AnnotatedDataFrames. The sampleData data frame is linked to the sampleID slot through the sample IDs. The assignmentData data frame is linked to the otuID slot through the OTU IDs.

There are slight differences in the OTUset objects and the TAXset objects. In the TAXset objects, the assignmentData data frame is not explicitly linked to the tax slot.

4 Conclusions and directions for development

OTUbase provides an organization and structure for OTU data and taxonomic classification data produced during the analysis of amplicon sequences. This allows the user to quickly and easily analyze amplicon data.

While the structure and a few basic functions are available withing OTUbase, there are a large number of possible improvements and extensions that have yet to be developed. OTUbase provides a structure for the data but functions for downstream analysis are not yet included. Future development will include a better integration of OTUbase with other available R packages such as vegan and the inclusion of a wider variety of functions for data analysis.

5 References

Sogin, M., H. Morrison, J. Huber, D. Welch, S. Huse, P. Neal, J. Arrieta, and G. Herndl. 2006. Microbial diversity in the deep sea and the underexplored "rare biosphere." *Proc. Natl. Acad. Sci. U. S. A.* 103:12115-12120

Schloss PD, et al. (2009) Introducing mothur: Open-source, platform-independent, community-supported software for describing and comparing microbial communities. *Appl Environ Microbiol* 75:7537-7541

Wang, Q, G. M. Garrity, J. M. Tiedje, and J. R. Cole. 2007. Naïve Bayesian Classifier for Rapid Assignment of rRNA Sequences into the New Bacterial Taxonomy. *Appl Environ Microbiol.* 73(16):5261-7

```
> toLatex(sessionInfo())
```

- R version 4.1.0 (2021-05-18), x86_64-pc-linux-gnu
- Locale: LC_CTYPE=en_US.UTF-8, LC_NUMERIC=C, LC_TIME=en_GB, LC_COLLATE=C, LC_MONETARY=en_US.UTF-8, LC_MESSAGES=en_US.UTF-8, LC_PAPER=en_US.UTF-8, LC_NAME=C, LC_ADDRESS=C, LC_TELEPHONE=C, LC_MEASUREMENT=en_US.UTF-8, LC_IDENTIFICATION=C
- Running under: Ubuntu 20.04.2 LTS
- Matrix products: default
- BLAS: /home/biocbuild/bbs-3.13-bioc/R/lib/libRblas.so
- LAPACK: /home/biocbuild/bbs-3.13-bioc/R/lib/libRlapack.so
- Base packages: base, datasets, grDevices, graphics, methods, parallel, stats, stats4, utils
- Other packages: Biobase 2.52.0, BiocGenerics 0.38.0, BiocParallel 1.26.0, Biostrings 2.60.0, GenomeInfoDb 1.28.0, GenomicAlignments 1.28.0, GenomicRanges 1.44.0, IRanges 2.26.0, MatrixGenerics 1.4.0, OTUbase 1.42.0, Rsamtools 2.8.0, S4Vectors 0.30.0, ShortRead 1.50.0, SummarizedExperiment 1.22.0, XVector 0.32.0, lattice 0.20-44, matrixStats 0.58.0, permute 0.9-5, vegan 2.5-7
- Loaded via a namespace (and not attached): DelayedArray 0.18.0, GenomeInfoDbData 1.2.6, MASS 7.3-54, Matrix 1.3-3, RColorBrewer 1.1-2, RCurl 1.98-1.3, bitops 1.0-7, cluster 2.1.2, compiler 4.1.0, crayon 1.4.1, grid 4.1.0, hwriter 1.3.2, jpeg 0.1-8.1, latticeExtra 0.6-29, mgcv 1.8-35, nlme 3.1-152, png 0.1-7, rstudioapi 0.13, splines 4.1.0, tools 4.1.0, zlibbioc 1.38.0

Table 1: The output of `sessionInfo` on the build system after running this vignette.