# Package 'msd16s'

December 19, 2024

Maintainer Joseph N. Paulson <pre><jpaulson@umiacs.umd.edu></jpaulson@umiacs.umd.edu></pre>
Author Joseph N. Paulson, Hector Corrada Bravo, Mihai Pop
<b>Version</b> 1.27.0
License Artistic-2.0
Title Healthy and moderate to severe diarrhea 16S expression data
<b>Description</b> Gut 16S sequencing expression data from 992 healthy and moderate-to-severe diarrhetic samples used in 'Diarrhea in young children from low-income countries leads to large-scale alterations in intestinal microbiota composition'.
LazyData yes
<b>Depends</b> R (>= 2.10), Biobase, metagenomeSeq,
<pre>URL http://www.cbcb.umd.edu/research/projects/GEMS-pathogen-discovery</pre>
biocViews ExperimentData, SequencingData, MicrobiomeData
git_url https://git.bioconductor.org/packages/msd16s
git_branch devel
git_last_commit 07e57c4
git_last_commit_date 2024-10-29
Repository Bioconductor 3.21
Date/Publication 2024-12-19
Contents
msd16s-package
Index

2 msd16s

msd16s-package	Curated dataset of many healthy and moderate-to-severe diarretic gut 16s samples on the 454 FLEX platform.

## Description

Data used in 'Diarrhea in young children from low-income countries leads to large-scale alterations in intestinal microbiota composition'. Measurements are number of reads clustered into OTUs (operational taxanomic units) by DNAClust.

#### Author(s)

Joseph N. Paulson

#### References

Diarrhea in young children from low-income countries leads to large-scale alterations in intestinal microbiota composition.

msd16s	Curated dataset of many healthy and moderate-to-severe diarretic gut 16s samples on the 454 FLEX platform.

## **Description**

Data used in 'Diarrhea in young children from low-income countries leads to large-scale alterations in intestinal microbiota composition'. Measurements are number of reads annotated for a particular cluster within a given sample followed by filtering. Sequencing was performed on the 454 Flex platform.

## format

Data is stored as an MRexperiment-class object. Using MRcounts one can obtain the 16S count matrix produced using using DNAclust (http://dnaclust.sourceforge.net/). The pData function accesses a data frame with the following columns:

Type: Status of samples: Case, Control

Country: Country of origin

Age: Month

AgeFactor: Month group

Dysentery: Dysentteric (1) non-dysenterric (0) indicator

The fData function accesses a data frame with the following columns:

OTU: OTU cluster id

msd16s

```
Taxonomy: Full taxonomic profile
superkingdom: superkingdom
phylum: phylum
class: class
order: order
family: family
genus: genus
species: species
clusterCenter: The OTU cluster's representative sequence
```

## Author(s)

Joseph N. Paulson

#### References

'Diarrhea in young children from low-income countries leads to large-scale alterations in intestinal microbiota composition'

## See Also

MRexperiment-class for the class definition, cumNorm to normalize the counts.

## Examples

```
data(msd16s)
head(pData(msd16s))
head(fData(msd16s))
```

## **Index**

```
* datasets
msd16s, 2
cumNorm, 3
msd16s, 2
msd16s-package, 2
```