Moderate-to-Severe diarrhea 16S dataset

April 25, 2017

This data package contains the information used to run the analyses found in "Diarrhea in young children from low-income countries leads to large-scale alterations in intestinal microbiota composition". Measurements are the number of reads annotated for a particular cluster within a given sample followed by filtering. Sequencing was performed on the 454 Flex platform. Data is stored as an MRexperiment-class object. The count matrix was generated using DNAclust (http://dnaclust.sourceforge.net/). For more details please refer to the paper.

The help file ?msd16s describes the example dataset.

1 The Data

We start by loading the library and the data.

```
> suppressMessages(library(metagenomeSeq))
> library(msd16s)
> data(msd16s)
```

This will load the msd16s object of class MRexperiment. As described in the metagenomeSeq vignette, print (or show) will display summary information.

> msd16s

```
MRexperiment (storageMode: environment)
assayData: 26044 features, 992 samples
element names: counts
protocolData: none
phenoData
sampleNames: 100259 100262 ... 602385 (992 total)
varLabels: Type Country ... Dysentery (5 total)
varMetadata: labelDescription
featureData
featureNames: 54 94 ... 276421 (26044 total)
```

```
fvarLabels: superkingdom phylum ... clusterCenter (10 total)
fvarMetadata: labelDescription
experimentData: use 'experimentData(object)'
Annotation:
```

The data in msd16s is the substrate for the analysis described in "Diarrhea in young children from low-income countries leads to large-scale alterations in intestinal microbiota composition". Included in the MRexperiment object are the counts, phenotype and feature information.

The phenotype information can be accessed with the phenoData and pData methods:

> phenoData(msd16s)

```
An object of class 'AnnotatedDataFrame' sampleNames: 100259 100262 ... 602385 (992 total) varLabels: Type Country ... Dysentery (5 total) varMetadata: labelDescription
```

> head(pData(msd16s))

	Туре	Country	Age	AgeFactor	Dysentery
100259	Case	Gambia	14	[12,18)	1
100262	${\tt Control}$	Gambia	24	[24,60)	0
100267	Case	Gambia	17	[12,18)	0
100274	Case	Gambia	36	[24,60)	0
100275	Case	Gambia	29	[24,60)	0
100277	Case	Gambia	29	[24,60)	0

The feature information including cluster representative sequence can be accessed with the featureData and fData methods:

> featureData(msd16s)

```
An object of class 'AnnotatedDataFrame' featureNames: 54 94 ... 276421 (26044 total) varLabels: superkingdom phylum ... clusterCenter (10 total) varMetadata: labelDescription
```

> head(fData(msd16s))

	superkingdom	phylum	class	order	family
54	Bacteria	Firmicutes	Bacilli	${\tt Lactobacillales}$	${\tt Lactobacillaceae}$
94	Bacteria	Firmicutes	Bacilli	${\tt Lactobacillales}$	${\tt Lactobacillaceae}$
113	Bacteria	Firmicutes	Bacilli	Lactobacillales	Lactobacillaceae

```
117
      Bacteria
                Firmicutes
                            Bacilli Lactobacillales Lactobacillaceae
145
      Bacteria Bacteroidetes Bacteroidia
                                    Bacteroidales
                                                 Prevotellaceae
202
      Bacteria Bacteroidetes Bacteroidia
                                    Bacteroidales
                                                 Bacteroidaceae
                               species OTU
54 Lactobacillus Lactobacillus sp. TSK G32-2
94 Lactobacillus Lactobacillus sp. TSK G32-2 94
113 Lactobacillus Lactobacillus sp. TSK G32-2 113
117 Lactobacillus Lactobacillus sp. TSK G32-2 117
                 Prevotella sp. DJF_RP53 145
145
     Prevotella
202
    Bacteroides
                    Bacteroides fragilis 202
54
                       ;cellular organisms;Bacteria;Firmicutes;Bacilli;Lactobacil
94
                       ;cellular organisms;Bacteria;Firmicutes;Bacilli;Lactobacil
113
                       ;cellular organisms;Bacteria;Firmicutes;Bacilli;Lactobacil
117
                       ;cellular organisms;Bacteria;Firmicutes;Bacilli;Lactobacil
145 ;cellular organisms;Bacteria;Bacteroidetes/Chlorobi group;Bacteroidetes;Bacteroidia
202
     cellular organisms;Bacteria;Bacteroidetes/Chlorobi group;Bacteroidetes;Bacteroid;
54
   94
                   113
                       CATGCTGCCTCCGTAGGAGTTTGGGCCGTGTCTCAGTCCCAATGTGGCCGATCAACC
117
                        145
                             CATGCTGCCTCCCGTAGGAGTTTGGACCGTGTCTCAGT
202
```

The raw or normalized counts matrix can be accessed with the MRcounts function:

> head(MRcounts(msd16s[,1:10]))

	100259	100262	100267	100274	100275	100277	100291	100292	100293	100294
54	0	0	0	0	0	0	0	0	0	0
94	0	0	0	0	0	0	0	0	0	0
113	0	0	0	0	0	0	0	0	0	0
117	0	0	0	0	0	0	0	0	0	0
145	0	0	0	0	0	0	0	0	0	0
202	0	0	0	0	0	0	0	0	0	0

Using this class, the object can be easily subsetted, for example:

```
> msd16s_bangladesh = msd16s[,pData(msd16s)$Country == "Bangladesh"]
```

MRexperiment (storageMode: environment) assayData: 26044 features, 206 samples

> msd16s_bangladesh

element names: counts

protocolData: none

phenoData

sampleNames: 600002 600005 ... 602385 (206 total)
varLabels: Type Country ... Dysentery (5 total)

varMetadata: labelDescription

featureData

featureNames: 54 94 ... 276421 (26044 total)

fvarLabels: superkingdom phylum ... clusterCenter (10 total)

fvarMetadata: labelDescription

experimentData: use 'experimentData(object)'

Annotation: