

Package ‘microSTASIS’

April 16, 2024

Title Microbiota STability ASsessment via Iterative cluStering

Version 1.2.0

Description The toolkit 'μSTASIS', or microSTASIS, has been developed for the stability analysis of microbiota in a temporal framework by leveraging on iterative clustering. Concretely, the core function uses Hartigan-Wong k-means algorithm as many times as possible for stressing out paired samples from the same individuals to test if they remain together for multiple numbers of clusters over a whole data set of individuals. Moreover, the package includes multiple functions to subset samples from paired times, validate the results or visualize the output.

License GPL-3

Imports BiocParallel, ggplot2, ggside, grid, rlang, stats, stringr, TreeSummarizedExperiment

Suggests BiocStyle, gghighlight, knitr, rmarkdown, methods, RefManageR, sessioninfo, SingleCellExperiment, SummarizedExperiment, testthat (>= 3.0.0)

biocViews GeneticVariability, BiomedicalInformatics, Clustering, MultipleComparison, Microbiome

BugReports <https://github.com/BiotechPedro/microSTASIS>

URL <https://doi.org/10.1093/bib/bbac055>

Encoding UTF-8

LazyData FALSE

Roxygen list(markdown = TRUE)

RoxygenNote 7.2.1

VignetteBuilder knitr

Depends R (>= 4.2.0)

Config/testthat/edition 3

git_url <https://git.bioconductor.org/packages/microSTASIS>

git_branch RELEASE_3_18

git_last_commit a3057b2

git_last_commit_date 2023-10-24

Repository Bioconductor 3.18

Date/Publication 2024-04-15

Author Pedro Sánchez-Sánchez [aut, cre]

(<https://orcid.org/0000-0002-4846-1813>),

Alfonso Benítez-Páez [aut] (<https://orcid.org/0000-0001-5707-4340>)

Maintainer Pedro Sánchez-Sánchez <bio.pedro.technology@gmail.com>

R topics documented:

clr	2
iterativeClustering	3
iterativeClusteringCV	4
microSTASIS	5
mSerrorCV	5
mSinternalPairedTimes	6
mSmetadataGroups	7
mSpreviz	8
pairedTimes	8
plotmSdynamics	10
plotmSheatmap	11
plotmSlinesCV	12
plotmSscatter	13
Index	14

clr	<i>Detected ASV from multiple individuals at four different sampling times.</i>
-----	---

Description

A dataset containing the amplicon sequence variants of 131 samples from the gut microbiota of 43 individuals. The values are transformed from counts by applying centred log-transformation (CLR).

Usage

```
data(clr)
```

Format

A data.frame with 131 rows and 226 variables

References

- Gloria M. Agudelo-Ochoa, Beatriz E. Valdés-Duque, Nubia A. Giraldo-Giraldo, Ana M. Jaillier-Ramírez, Adriana Giraldo-Villa, Irene Acevedo-Castaño, Mónica A. Yepes-Molina, Janeth Barbosa-Barbosa, Alfonso Benítez-Paéz, Gut microbiota profiles in critically ill patients, potential biomarkers and risk variables for sepsis, *Gut Microbes*, Volume 12, Issue 1, January 2020, <https://doi.org/10.1080/19490976.2019.170>
- Pedro Sánchez-Sánchez, Francisco J Santonja, Alfonso Benítez-Páez, Assessment of human microbiota stability across longitudinal samples using iteratively growing-partitioned clustering, *Briefings in Bioinformatics*, Volume 23, Issue 2, March 2022, bbac055, <https://doi.org/10.1093/bib/bbac055>

iterativeClustering *Stability of individuals after iteratively performing Hartigan-Wong k-means clustering.*

Description

Perform Hartigan-Wong `stats::kmeans()` algorithm as many times as possible. The values of `k` are from 2 to the number of samples minus 1. Those individuals whose paired samples are clustered under the same label sum 1. If paired samples are in different clusters, then sum 0, except when the euclidean distance between them is smaller to the ones of each sample to its centroid. This is done for all possible values of `k` and, finally, divided the sum by `k`, so obtaining a value between 0 and 1.

Usage

```
iterativeClustering(
  pairedTimes,
  BPPARAM = BiocParallel::bpparam(),
  common = "_"
)
```

Arguments

- | | |
|--------------------------|--|
| <code>pairedTimes</code> | list of matrices with paired times, i.e. samples to be stressed to multiple iterations. Output of <code>pairedTimes()</code> . |
| <code>BPPARAM</code> | supply a <code>BiocParallel</code> parameters object, e.g. <code>BiocParallel::SerialParam()</code> in the specific case of Windows OS or <code>BiocParallel::bpparam()</code> . |
| <code>common</code> | pattern that separates the ID and the sampling time. |

Value

μ STASIS stability score (mS) for the individuals from the corresponding paired times.

Examples

```
data(clr)
times <- pairedTimes(data = clr, sequential = TRUE, common = "_0_")
mS <- iterativeClustering(pairedTimes = times, common = "_")
```

 microSTASIS

Microbiota Stability Assessment via Iterative cluStering

Description

The toolkit 'μSTASIS' has been developed for the stability analysis of microbiota in a temporal framework by leveraging on iterative clustering. Concretely, the core function uses Hartigan-Wong k-means algorithm as many times as possible for stressing out paired samples from the same individuals to test if they remain together for multiple numbers of clusters over a whole data set of individuals. Moreover, the package includes multiple functions to subset samples from paired times, validate the results or visualize the output.

 mSerrorCV

Compute the mean absolute error (MAE) in percentage after [iterativeClusteringCV\(\)](#).

Description

Compute the mean absolute error after the cross validation or plot lines connecting the stability values for each subset of the original matrix of paired times.

Usage

```
mSerrorCV(pairedTime, CVklist, k = 1L)
```

Arguments

`pairedTime` input matrix with paired times whose stability has being assessed. One of the lists output of [pairedTimes\(\)](#).

`CVklist` list resulting from [iterativeClusteringCV\(\)](#).

`k` integer; number of individuals to subset from the data. The same as used in [iterativeClusteringCV\(\)](#).

Value

A vector with MAE values for each individual's mS score.

Examples

```
data(clr)
times <- pairedTimes(data = clr[, 1:20], sequential = TRUE, common = "_0_")
mS <- iterativeClustering(pairedTimes = times, common = "_")
cv_klist_t1_t25_k2 <- iterativeClusteringCV(pairedTimes = times,
                                           results = mS, name = "t1_t25",
                                           common = "_0_", k = 2L)
MAE_t1_t25 <- mSerrorCV(pairedTime = times$t1_t25,
```

```
CVklist = cv_klist_t1_t25_k2, k = 2L)
MAE <- mSpreviz(results = list(MAE_t1_t25),
               times = list(t1_t25 = times$t1_t25))
plotmSheatmap(results = MAE, times = c("t1_t25", "t25_t26"), label = TRUE,
              high = 'red2', low = 'forestgreen', midpoint = 5)
```

mSinternalPairedTimes *Internal function for [pairedTimes\(\)](#).*

Description

Internal function for [pairedTimes\(\)](#).

Usage

```
mSinternalPairedTimes(data, specifiedTimePoints, common = "_")
```

Arguments

data matrix with rownames including ID, common pattern and sampling time.

specifiedTimePoints character vector to specify the selection of concrete paired times.

common pattern separating the ID and the sampling time in rownames.

Value

A list of matrices with the same number of columns as input and with samples from paired sampling times as rows.

Examples

```
data(clr)
t1_t2 <- mSinternalPairedTimes(data = clr,
                              specifiedTimePoints = c("1", "25"),
                              common = "_0_")
```

mSmetadataGroups	<i>Easily extract groups of individuals from sample metadata.</i>
------------------	---

Description

Easily extract groups of individuals from sample metadata.

Usage

```
mSmetadataGroups(  
  metadata,  
  samples,  
  individuals,  
  variable,  
  common,  
  ID,  
  timePoints  
)
```

Arguments

metadata	input data.frame with data corresponding to samples. It can be the <code>SummarizedExperiment::colData()</code> from the <code>TreeSummarizedExperiment</code> .
samples	vector from metadata corresponding to the samples ID, if applicable; should be NULL if ID and timePoints are provided from a <code>TreeSummarizedExperiment</code> , for example.
individuals	vector of individuals; first column of the <code>mSpreviz()</code> output.
variable	column name with the variable used for grouping individuals.
common	pattern that separates the ID and the sampling time in rownames, if applicable.
ID	If applicable, one of the <code>colData()</code> colnames from the <code>TreeSummarizedExperiment</code> should be given as individuals.
timePoints	If applicable, one of the <code>colData()</code> colnames from the <code>TreeSummarizedExperiment</code> should be given as sampling times.

Value

A vector with the same length as the number of rows in the `mSpreviz()` output.

Examples

```
data(clr)  
times <- pairedTimes(data = clr, sequential = TRUE, common = "_0_")  
mS <- iterativeClustering(pairedTimes = times, common = "_")  
results <- mSpreviz(results = mS, times = times)  
metadata <- data.frame(Sample = rownames(clr), age = c(rep("youth", 65),  
  rep("old", 131-65)))
```

```
group <- mSmetadataGroups(metadata = metadata, samples = metadata$Sample,
                          common = "_0_", individuals = results$individual,
                          variable = "age")
```

mSpreviz	<i>Process the <code>iterativeClustering()</code> output to a new format ready for the implemented visualization functions.</i>
----------	---

Description

Process the `iterativeClustering()` output to a new format ready for the implemented visualization functions.

Usage

```
mSpreviz(results, times)
```

Arguments

results	list; output of <code>iterativeClustering()</code> .
times	list; output of <code>pairedTimes()</code> .

Value

A data frame ready for its use under the implemented visualization functions and others.

Examples

```
data(clr)
times <- pairedTimes(data = clr, sequential = TRUE, common = "_0_")
mS <- iterativeClustering(pairedTimes = times, common = "_")
results <- mSpreviz(results = mS, times = times)
```

pairedTimes	<i>Generate one or multiple matrices with paired times.</i>
-------------	---

Description

Generate one or multiple matrices with paired times.

Usage

```
pairedTimes(data, ...)

## S4 method for signature 'matrix'
pairedTimes(data, sequential, common, specifiedTimePoints)

## S4 method for signature 'TreeSummarizedExperiment'
pairedTimes(
  data,
  sequential,
  assay,
  alternativeExp,
  ID,
  timePoints,
  specifiedTimePoints
)
```

Arguments

<code>data</code>	input object: either a matrix with rownames including ID, common pattern and sampling time, or a <code>TreeSummarizedExperiment</code> object.
<code>...</code>	Additional argument list that might not ever be used.
<code>sequential</code>	TRUE if paired times to analyse are sequential and present the desired alphanumerical order.
<code>common</code>	If <code>is.matrix(data)</code> , pattern that separates the ID and the sampling time in rownames.
<code>specifiedTimePoints</code>	character vector to specify the selection of concrete paired times.
<code>assay</code>	If <code>class(data) == "TreeSummarizedExperiment"</code> , name of the assay to use.
<code>alternativeExp</code>	If <code>class(data) == "TreeSummarizedExperiment"</code> , name of the alternative experiment to use (if applicable).
<code>ID</code>	If <code>class(data) == "TreeSummarizedExperiment"</code> , one of the <code>colData(data)</code> colnames should be given as individuals.
<code>timePoints</code>	If <code>class(data) == "TreeSummarizedExperiment"</code> , one of the <code>colData(data)</code> colnames should be given as sampling times.

Value

A list of matrices with the same number of columns as input and with samples from paired sampling times as rows.

Examples

```
data(clr)
times_b <- pairedTimes(data = clr, sequential = TRUE, common = "_0_")
times_b <- pairedTimes(data = clr, sequential = FALSE, common = "_0_",
  specifiedTimePoints = c("1", "26"))
```

plotmSdynamics	<i>Generate boxplots of the stability dynamics throughout sampling times by groups.</i>
----------------	---

Description

Generate boxplots of the stability dynamics throughout sampling times by groups.

Usage

```
plotmSdynamics(results, groups, points = TRUE, linetype = 2)
```

Arguments

results	input data.frame resulting from <code>mSpreviz()</code> .
groups	vector with the same length as individuals, i.e. the number of rows in the <code>mSpreviz()</code> output.
points	logical; FALSE to only visualize boxplots or TRUE to also add individual points.
linetype	numeric; type of line to connect the median value of paired times; 0 to avoid the line.

Value

A plot with as many boxes as paired times by group in the form of a `ggplot2::ggplot()` object.

Examples

```
data(c1r)
times <- pairedTimes(data = c1r, sequential = TRUE, common = "_0_")
mS <- iterativeClustering(pairedTimes = times, common = "_")
results <- mSpreviz(results = mS, times = times)
metadata <- data.frame(Sample = rownames(c1r), age = c(rep("youth", 65),
  rep("old", 131-65)))
group <- mSmetadataGroups(metadata = metadata, samples = metadata$Sample,
  common = "_0_", individuals = results$individual,
  variable = "age")
plotmSdynamics(results, groups = group, points = TRUE, linetype = 0)
```

plotmSheatmap	<i>Plot a heatmap of the stability results.</i>
---------------	---

Description

Plot a heatmap of the stability results.

Usage

```
plotmSheatmap(  
  results,  
  order = NULL,  
  times,  
  label = FALSE,  
  low = "red2",  
  mid = "yellow",  
  high = "forestgreen",  
  midpoint = 0.5  
)
```

Arguments

results	input data.frame resulting from <code>mSpreviz()</code> .
order	NULL object or character: none, mean or median; if the individuals should be sorted by any of those statistics of the stability values.
times	character; names of the paired times to plot, i.e. colnames of results.
label	logical; TRUE to print the mS score or FALSE to not.
low	color for the lowest value.
mid	color for the middle value.
high	color for the highest values.
midpoint	value to situate the middle.

Value

A heatmap of the stability values in the form of a `ggplot2::ggplot()` object.

Examples

```
data(clr)  
times <- pairedTimes(data = clr, sequential = TRUE, common = "_0_")  
mS <- iterativeClustering(pairedTimes = times, common = "_")  
results <- mSpreviz(results = mS, times = times)  
plotmSheatmap(results = results, order = "mean", times = c("t1_t25", "t25_t26"),  
  label = TRUE)
```

plotmSlinesCV *Plot the stability values after `iterativeClusteringCV()`.*

Description

Plot lines connecting the mS score for each subset of the original matrix of paired times.

Usage

```
plotmSlinesCV(pairedTime, CVklist, k = 1L, points = TRUE, sizeLine = 0.5)
```

Arguments

pairedTime	input matrix with paired times whose stability has being assessed. One of the lists output of <code>pairedTimes()</code> .
CVklist	list resulting from <code>iterativeClusteringCV()</code> .
k	integer; number of individuals to subset from the data. The same as used in <code>iterativeClusteringCV()</code> .
points	logical; if plotting, FALSE to only plot lines and TRUE to add points on the mS score, i.e. result from <code>iterativeClusteringCV()</code> .
sizeLine	numeric; if plotting, size of the multiple lines.

Value

A line plot in the form of a `ggplot2::ggplot()` object with the values of stability for the multiple subsets and the original matrix of paired samples (points).

Examples

```
data(clr)
times <- pairedTimes(data = clr[, 1:20], sequential = TRUE, common = "_0_")
mS <- iterativeClustering(pairedTimes = times, common = "_")
cv_klist_t1_t25_k2 <- iterativeClusteringCV(pairedTimes = times,
                                           results = mS, name = "t1_t25",
                                           common = "_0_", k = 2L)
plotmSlinesCV(pairedTime = times$t1_t25, CVklist = cv_klist_t1_t25_k2, k = 2L)
```

plotmScatter	<i>Plot a scatter and side boxplot of the stability results.</i>
--------------	--

Description

Plot a scatter and side boxplot of the stability results.

Usage

```
plotmScatter(results, order = NULL, times, gridLines = FALSE, sideScale = 0.3)
```

Arguments

results	input data.frame resulting from <code>mSpreviz()</code> .
order	NULL object or character: mean or median; if the individuals should be sorted by any of those statistics of the stability values.
times	a vector with the names of each paired time, e.g. "t1_t2".
gridLines	logical; FALSE to print a blank background or TRUE to include a gray grid.
sideScale	numeric; scale of the side boxplot.

Value

A scatter plot and a side boxplot of the stability values in the form of a `ggplot2::ggplot()` object.

Examples

```
data(c1r)
times <- pairedTimes(data = c1r, sequential = TRUE, common = "_0_")
mS <- iterativeClustering(pairedTimes = times, common = "_")
results <- mSpreviz(results = mS, times = times)
plotmScatter(results = results, order = "median", times = c("t1_t25",
                                                         "t25_t26"),
             gridLines = TRUE, sideScale = 0.2)
```

Index

* datasets

clr, [2](#)

BiocParallel::bpparam(), [3](#), [4](#)

BiocParallel::SerialParam(), [3](#), [4](#)

clr, [2](#)

ggplot2::ggplot(), [10–13](#)

iterativeClustering, [3](#)

iterativeClustering(), [4](#), [8](#)

iterativeClusteringCV, [4](#)

iterativeClusteringCV(), [5](#), [12](#)

microSTASIS, [5](#)

mSerrorCV, [5](#)

mSinternalPairedTimes, [6](#)

mSmetadataGroups, [7](#)

mSpreviz, [8](#)

mSpreviz(), [7](#), [10](#), [11](#), [13](#)

pairedTimes, [8](#)

pairedTimes(), [3–6](#), [8](#), [12](#)

pairedTimes, matrix, matrix-method
(pairedTimes), [8](#)

pairedTimes, matrix-method
(pairedTimes), [8](#)

pairedTimes, TreeSummarizedExperiment, TreeSummarizedExperiment-method
(pairedTimes), [8](#)

pairedTimes, TreeSummarizedExperiment-method
(pairedTimes), [8](#)

plotmSdynamics, [10](#)

plotmSheatmap, [11](#)

plotmSlinesCV, [12](#)

plotmSscatter, [13](#)

stats::kmeans(), [3](#)

SummarizedExperiment::colData(), [7](#)