Package ‘HMPTrees’

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Type Package

Title Statistical Object Oriented Data Analysis of RDP-based Taxonomic trees from Human Microbiome Data: Modeling, Visualization, and Two-Group Comparison

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Description In this package, we introduce Object Oriented Data Analysis (OODA) methods to analyze Human Microbiome taxonomic trees directly. We provide tools to model, compare, and visualize populations of taxonomic tree objects.

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HMPTrees-package .................................................. 2
checkTreeValidity ................................................... 3
compareTwoDataSets ............................................... 4
createAndPlot ....................................................... 5
createTrees ........................................................ 7
displayLegend ...................................................... 8
formatData ........................................................ 8
In this package, we provide tools to model, compare, and visualize populations of taxonomic tree objects. In particular, using a unimodal probability measure in the set of trees, we provide procedures to compute the MLE of the central tree and the LRT statistics for comparing the distributions of two metagenomic populations.

Details

HMP metagenomic sequences in a sample can be represented as a rooted taxonomic tree. Using supervised taxonomic methods a sequence is matched to a hierarchical taxa or taxonomy bins defined in a bacterial-taxonomy library such as, for example, the Ribosomal Database Project (RDP) (Cole, 2005). The supervised taxonomic analysis allows us to represent each sample (set of sequences) by a rooted taxonomic tree where the root corresponds to taxon at the Kingdom level, i.e., bacteria, and the leaves correspond to the taxa at the Genus level, and the width of the edges (paths) between taxonomic levels correspond to the 'abundances' of the descending taxon.

In particular, we combine RDP matches by adding RDP values of common taxon, which allows us to provide a measure of taxa abundance weighting on the confidence of each taxa assignment. The resulting taxonomic trees satisfy the following conditions: i) branches closer to the root have higher 'abundance' values than branches closer to leaves, and ii) the sum of the 'abundances' of all descending taxa under a common parent taxon cannot be larger than the 'abundance' of the corresponding parent taxon.

It is important to note that due to how the ape package works the following naming conventions apply to taxa names:

1. Colons cannot be used in the taxa names at all.
2. Each taxa name must be unique - you cannot have two separate branches both have a child named ‘unclassified’ for example. (We took the parent name and added a ‘U’ to the end to signify an unclassified in our data sets)

3. There can only be one top level node. (Bacteria and Archaea cannot both exist unless there is an additional single level above them for example)

Author(s)

Patricio S. La Rosa, Elena Deych, Berkley Shands, William D. Shannon

References


checkTreeValidity  
Check Validity of an RDP-Based Taxonomic Tree

Description

This function goes through every node in the tree and for each node it checks that the sum of that node’s children are less than or equal to the value of that node.

Usage

checkTreeValidity(data, samples = 1, epsilon = 0.0001, split = ".")

Arguments

data  
A data frame in which each column contains the rdp read counts for every taxa given in the row names.

samples  
A numeric vector that is used to select a single (or several) tree(s) out of a data set. A value of 0 will select all trees.

epsilon  
This value allows for rounding problems or other such small errors in the data such that the (parent + epsilon > sum(children)).

split  
This is the character that separates the taxa levels in the row names.
Value

A boolean vector that indicates the validity of every tree tested.

Author(s)

Berkley Shands, Patricio S. La Rosa, Elena Deych, William D. Shannon

Examples

data(saliva)
validTree <- checkTreeValidity(saliva, 1)
validTree

Description

This functions compares the distribution of two sets of RDP-based taxonomic trees using Likelihood-Ratio-Test statistics and a p-value is computed using bootstrapping.

Usage

compareTwoDataSets(data1, data2, numBootStraps = 1000, enableMC = FALSE, cores = 3)

Arguments

data1, data2  Data frames in which each column contains the rdp read counts for every taxa given in the row names.
numBootStraps The number of times to run the bootstrapping, the default is 1000.
enableMC When this is 'TRUE' it allows for parallel calculation of the bootstraps. (See Note 1 in Details).
cores The number of parallel processes to run if enableMC is 'TRUE'.

Details

Notes:

1. Enabling parallel calculation requires the package doParallel.
2. Both data sets should be standardized to the same number of reads.
We are interested in assessing whether the distributions from two metagenomic populations are the same or different, which is equivalent to evaluating whether their respective parameters are the same or different. The corresponding hypothesis is given as follows:

\[ H_0 : (g^*_1, \tau_1) = (g^*_2, \tau_2) = (g^*_0, \tau_0) \text{ vs } H_A : (g^*_1, \tau_1) \neq (g^*_2, \tau_2), \]

where \((g^*_0, \tau_0)\) is the unknown common parameter vector. To evaluate this hypothesis we use the likelihood-ratio test (LRT) which is given by,

\[
\lambda = -2 \log \left( \frac{L(g^*_0, \tau_0; S_{1n}, S_{2m})}{L(g^*_1, \tau_1; S_{1n}) + L(g^*_2, \tau_2; S_{2m})} \right),
\]

where \(S_{1n}\) and \(S_{2m}\) are the sets containing \(n\) and \(m\) random samples of trees from each metagenomic population, respectively. We assume that the model parameters are unknown under both the null and alternative hypothesis, therefore, we estimate these using the MLE procedure proposed in La Rosa et al (see reference 2), and compute the corresponding p-value using non-parametric bootstrap.

**Value**

A p-value for the similarity of the two data sets based on the bootstrapping.

**Author(s)**

Patricio S. La Rosa, Elena Deych, Berkley Shands, William D. Shannon

**Examples**

```r
data(saliva)
data(stool)

### We use 1 for the number of bootstraps for computation time
### This value should be at least 1000 for an accurate result
numBootStraps <- 1
pval <- compareTwoDataSets(saliva, stool, numBootStraps)
pval
```

**createAndPlot**

Create and Plot a Tree from a Data Set

**Description**

This function combines the createTrees and plotTree functions to create and plot a set of trees.

**Usage**

```r
createAndPlot(data, samples = 1, level = "genus", colors, divisions, main, sub, showTipLabel = TRUE, showNodeLabel = FALSE, displayLegend = TRUE, onePerPage = FALSE)
```
Arguments

data A data frame in which each column contains the rdp read counts for every taxa
given in the row names.
samples A numeric vector that is used to select a single (or several) tree(s) out of a data
set. A value of 0 will select all trees.
level The depth the tree creation will go down to (kingdom, phylum, class, order,
family, genus, species, subspecies). 'genus' is the default.
colors A vector of colors to be applied to the branches in the plot.
divisions A vector of numbers to be used as break points to assign different colors.
main A custom title(s) for the plot(s). The default value is the column names.
sub A custom subtitle for the plot.
showTipLabel Hides the tip labels if 'FALSE' otherwise it shows all non-zero tip labels.
showNodeLabel Hides the interior node labels if 'FALSE' otherwise it shows all non-zero node
labels.
displayLegend Enables the display of a legend of the branch colors and divisions when 'TRUE'.
onePerPage If 'TRUE' one tree will be plotted per page, if 'FALSE' four will be displayed
per page.

Details

Notes:

1. For 'level' k, p, c, o, f, g, s and ss can be used in place of kingdom, phylum, class, order,
family, genus, species and subspecies respectively.
2. The values for division should directly relate to the values of your data, i.e. if your data ranges
from 0 to 50000 reads you should adjust the divisions to fit your data.

Value

A plot of the tree(s).

Author(s)

Berkley Shands, Patricio S. La Rosa, Elena Deych, William D. Shannon

Examples

data(saliva)

### Plots the trees in column 2 and 3 in 'Saliva'
createAndPlot(saliva, c(2:3))
Description

This function creates a list tree objects of type 'phylo' for use in plotting the trees.

Usage

createTrees(data, samples = 1, level = "genus")

Arguments

data A data frame in which each column contains the rdp read counts for every taxa given in the row names.
samples A numeric vector that is used to select a single (or several) tree(s) out of a data set. A value of 0 will select all trees.
level The depth the tree creation will go down to (kingdom, phylum, class, order, family, genus, species, subspecies). 'genus' is the default.

Details

For 'level' k, p, c, o, f, g, s and ss can be used in place of kingdom, phylum, class, order, family, genus, species and subspecies respectively.

Value

A list of 'phylo' objects that can be passed to plotTree to plot them.

Author(s)

Berkley Shands, Patricio S. La Rosa, Elena Deych, William D. Shannon

Examples

data(saliva)

mySalivaTree <- createTrees(saliva, 4)
**displayLegend**  
*Displays Tree Plot Legend*

**Description**
This function displays a legend that shows the tree branch sizes/colors divisions.

**Usage**
```
displayLegend(colors, divisions, title = "Confidence Value")
```

**Arguments**
- **colors**
  A vector of colors to be used in the plot from lowest ranking to highest ranking.
- **divisions**
  A vector of numbers from lowest to highest to separate the tree branches into the color ranking.
- **title**
  The title for the legend. The default is 'Confidence Value'.

**Details**
The values for division should directly relate to the values of your data, i.e. if your data ranges from 0 to 50000 reads you should adjust the divisions to fit your data.

**Value**
A blank plot that contains a legend.

**Author(s)**
Berkley Shands, Patricio S. La Rosa, Elena Deych, William D. Shannon

**Examples**
```
displayLegend(c("red", "orange", "blue"), c(.1, 100, 10000))
```

---

**formatData**  
*Formats a Data Set*

**Description**
This function will take a data set and format it by removing low count trees, and/or normalizing counts.

**Usage**
```
formatData(data, countThreshold = 1000, normalizeThreshold = 10000)
```

**Description**
This function will take a data set and format it by removing low count trees, and/or normalizing counts.

**Usage**
```
formatData(data, countThreshold = 1000, normalizeThreshold = 10000)
```
generateTree

Arguments

data A data frame in which each column contains the rdp read counts for every taxa
given in the row names.
countThreshold A cut off threshold for reads - all trees with fewer than this number of reads will
be removed.
normalizeThreshold
  All the trees that are not removed will be normalized to this many reads.

Details

When removing trees with too few reads, the cuts off is based on the value of the top level node,
not the sum of all the reads in a sample.

Value

A new data set that is trimmed and standardized based on the specified parameters. The new data is
also reordered alphabetically according to row labels.

Author(s)

Patricio S. La Rosa, Elena Deych, Berkley Shands, William D. Shannon

Examples

data(throat)

throat <- formatData(throat, 1000, 10000)

generateTree

Generate Test Trees

Description

This function will take several initial trees and will randomly populate new trees based on the
originals.

Usage

generateTree(data, nreads = 10000, nsamps = 50, theta = 0, level = "genus", split = ".")

Arguments

data A data frame in which each column contains the rdp read counts for every taxa
given in the row names.
nreads The maximum number of reads the created a tree will contain.
nsamps The number of trees to generate.
 theta  When theta is between 0 and 1 the base tree is generated by using the \texttt{dirmult} function and modified by the theta value, otherwise the base tree is generated by the \texttt{dmNmom} function.

 level  The depth the tree will go down to (kingdom, phylum, class, order, family, genus, species, subspecies). Genus is the default.

 split  This is the character that separates the taxa levels in the row names.

\section*{Details}
This function requires that the package \texttt{HMP} be installed.

\section*{Value}
A data frame containing the generated tree(s).

\section*{Author(s)}
Patricio S. La Rosa, Elena Deych, Berkley Shands, William D. Shannon

\section*{Examples}
\begin{verbatim}
data(saliva)

gndata <- generateTree(saliva, 7000, 2)
\end{verbatim}

---

\begin{verbatim}
getMLEandLoglike \hspace{1cm} Get MLE and Log Likelihood of a Data Set
\end{verbatim}

\section*{Description}
This function takes a data set and computes the MLE and its Log-Likelihood value.

\section*{Usage}
\begin{verbatim}
getMLEandLoglike(data, maxSteps = 50, weight = NULL)
\end{verbatim}

\section*{Arguments}
\begin{verbatim}
data \hspace{1cm} A data frame in which each column contains the rdp read counts for every taxa given in the row names.

maxSteps \hspace{1cm} The maximum number of times to bootstrap for the MLE.

weight \hspace{1cm} A vector to weight the MLE tree.
\end{verbatim}
Details

A unimodal probability model for graph-valued random objects has been derived and applied previously to several types of graphs (cluster trees, digraphs, and classification and regression trees) (For example, Banks and Constantine, 1998; Shannon and Banks, 1999). Here we apply this model to HMP trees constructed from RDP matches. Let $G$ be the finite set of taxonomic trees with elements $g$, and $d : G \times G \to R^+$ an arbitrary metric of distance on $G$. We have the probability measure $H(g^*, \tau)$ defined by

$$P(g; g^*, \tau) = c(g^*, \tau) \exp(-\tau d(g^*, g)), \text{forall} g \in G,$$

where $g^*$ is the modal or central tree, $\tau$ is a concentration parameter, and $c(g^*, \tau)$ is the normalization constant. The distance measure between two trees is the Euclidean norm of the difference between their corresponding adjacency-vectors. To estimate the parameters $(g^*, \tau)$, we use the maximum likelihood estimate (MLE) procedure described in La Rosa et al. (see reference 2)

Value

A list containing the MLE, log-likelihood, and tau.

Author(s)

Patricio S. La Rosa, Elena Deych, Berkley Shands, William D. Shannon

Examples

data(saliva)

### We use 1 for the maximum number of steps for computation time
### This value should be much higher to ensure an accurate result
numSteps <- 1
mle <- getMLEandLoglike(saliva, numSteps)$mleTree

mergeDataSets

Merge Multiple Data Sets

Description

This function can take any number of data sets, calculate their individual and combined MLEs and then merge them.

Usage

mergeDataSets(data, calcMLE = TRUE, uniqueNames = FALSE)
Arguments

- **data**: A list of data frames in which each column contains the rdp read counts for every taxa given in the row names.
- **calcMLE**: If 'FALSE' the MLEs for the data sets will not be calculated, otherwise they are added to the end.
- **uniqueNames**: If 'TRUE' the column names in the combined data set will be appended to insure uniqueness, otherwise the column names will follow the naming process from the `merge` function.

Details

Although not required, all data sets should be standardized to the same number of reads before merging.

Value

A single data set containing all the data from the input data sets, in addition to their individual MLEs and a combined MLE if requested.

Author(s)

Berkley Shands, Patricio S. La Rosa, Elena Deych, William D. Shannon

Examples

```r
data(saliva)
data(stool)

dataComb <- mergeDataSets(list(saliva, stool), FALSE)
```

---

**plotTree**

Plots a Tree Object

Description

This function takes one or more 'phylo' objects and plots them.

Usage

```r
plotTree(trees, colors, divisions, main, sub, showTipLabel = TRUE, showNodeLabel = FALSE, displayLegend = TRUE)
```
**plotTree**

**Arguments**

- **trees**: A list that contains at least one tree of type ‘phylo’.
- **colors**: A vector of colors to be applied to the branches in the plot.
- **divisions**: A vector of numbers to be used as break points to assign different colors.
- **main**: A custom title(s) for the plot(s). The default value is the column names.
- **sub**: A custom subtitle for the plot.
- **showTipLabel**: Hides the tip labels if ‘FALSE’ otherwise it shows all non-zero tip labels.
- **showNodeLabel**: Hides the interior node labels if ‘FALSE’ otherwise it shows all non-zero node labels.
- **displayLegend**: Enables the display of a legend of the branch colors and divisions when ‘TRUE’.

**Details**

Notes:

1. The `phylo` type is a product of the `ape` package and the `createTrees` function in this package produces a list of `phylo` type objects for use with this function.

2. The values for division should directly relate to the values of your data, i.e. if your data ranges from 0 to 50000 reads you should adjust the divisions to fit your data.

**Value**

A plot of the tree(s).

**Author(s)**

Berkley Shands, Patricio S. La Rosa, Elena Deych, William D. Shannon

**Examples**

```r
data(saliva)

### Creates a object of type 'phylo' for the 4th tree in 'Saliva'
### Then plots it
salivaTree <- createTrees(saliva, 4)
plotTree(salivaTree, displayLegend=FALSE)
```
plotTreeDataMDS  
Plot an MDS Plot of a Group of Trees

Description
This function can take any number of data sets and plots them on an MDS plot to show relative
closeness to one another.

Usage
plotTreeDataMDS(data, main = "Tree MDS Comparisons", calcMLE = TRUE, mleTitles,
dotColors = c("red", "orange", "blue", "green", "yellow", "purple"),
dotSizes = c(1, 2), showNames = FALSE, returnCoords = FALSE)

Arguments
- data: A single data frame or a list of a data frames in which each column contains the
  rdp read counts for every taxa given in the row names.
- main: A title for the MDS plot.
- calcMLE: If ‘FALSE’ the MLEs for the data sets will not be calculated and plotted.
- mleTitles: A vector of labels for each MLE data point on the MDS plot.
- dotColors: The colors to be used when plotting the points and/or MLE points on the MDS
  plot. (See Notes in Details)
- dotSizes: A vector in which the first value is the data points CEX and the second value is
  the MLEs CEX.
- showNames: When ‘TRUE’ the column name will be plotted above each corresponding point.
- returnCoords: When ‘TRUE’ this function will return the x and y coordinates for every plotted
  point.

Details
Notes:
1. If the length of dot colors is at least twice the length of data, the colors will correspond in
   the following manner: (data 1’s points, data 1’s MLE, data 2’s points, data 2’s MLE, etc)
2. If the length of dot colors is less than twice the length of data, the colors will correspond in
   the following manner: (data 1’s points & data 1’s MLE, data 2’s points & data 2’s MLE, etc)
3. If the length of dot colors is less than the length of data, dotColors will be set by the rainbow
   function and the colors will correspond as above

Value
A MDS plot of the data.
Author(s)

Berkley Shands, Patricio S. La Rosa, Elena Deych, William D. Shannon

Examples

data(saliva)
data(stool)

plotTreeDataMDS(list(saliva, stool), mleTitles=c("Saliva", "Stool"))

saliva  

Saliva Data Set

Description

A data set containing all taxa from 24 subjects.

Usage

data(saliva)

Format

The format is a data frame of 454 rows by 24 columns, with each column being a separate subject and each row being a different taxa denoted by the row names. The taxonomical levels are separated by a "." in their names (Bacteria.Phylum.Class....). The values in each column are the sum of values that each taxa had in an RDP file. It should also be noted that the samples are normalized to 7000 reads and any level that ends with a U was unclassified in the RDP file.

stool  

Stool Data Set

Description

A data set containing all taxa from 24 subjects.

Usage

data(stool)

Format

The format is a data frame of 371 rows by 24 columns, with each column being a separate subject and each row being a different taxa denoted by the row names. The taxonomical levels are separated by a "." in their names (Bacteria.Phylum.Class....). The values in each column are the sum of values that each taxa had in an RDP file. It should also be noted that the samples are normalized to 7000 reads and any level that ends with a U was unclassified in the RDP file.
### throat  

**Throat Data Set**

**Description**

A data set containing all taxa from 22 subjects.

**Usage**

```r
data(throat)
```

**Format**

The format is a data frame of 529 rows by 22 columns, with each column being a separate subject and each row being a different taxa denoted by the row names. The taxanomical levels are separated by a '.' in their names (Bacteria.Phylum.Class....). The values in each column are the sum of values that each taxa had in an RDP file. It should also be noted that the samples have not been normalized and should be used with 'formatData'. Also any level that ends with a U was unclassified in the RDP file.

---

### transformHMPtoHMPTree  

**Transform data from HMP to HMPTrees**

**Description**

This function will take data in the format needed in the package 'HMP' to the format used by 'HMPTrees'.

**Usage**

```r
transformHMPtoHMPTree(data)
```

**Arguments**

- `data` A data frame of values in which each column is a separate taxa level and each row is a separate sample.

**Details**

This function does the opposite of `transformHMPTreetoHMP`.

**Value**

A data frame in which each column contains the rdp read counts for every taxa given in the row names.
transformHMPTreetoHMP

**Author(s)**
Berkley Shands, Patricio S. La Rosa, Elena Deych, William D. Shannon

**Examples**

data(saliva)

```r
### Trims saliva to only contain the class level
salivaClass <- trimToTaxaLevel(saliva, "class", TRUE)

### This transforms the saliva data set but retains
### any zero rows that may exist.
transSaliva <- transformHMPTreetoHMP(salivaClass, FALSE, 0)

### saliva2 should be the same as salivaClass
saliva2 <- transformHMPtoHMPTree(transSaliva)
```

---

**transformHMPTreetoHMP  Transform data from HMPTrees to HMP**

**Description**
This function will take data in the format needed in the package 'HMPTrees' to the format used by 'HMP'.

**Usage**

```r
transformHMPTreetoHMP(data, elimZero = FALSE, zeroValue = 1e-05)
```

**Arguments**

data  A data frame in which each column contains the rdp read counts for every taxa
given in the row names.

elimZero  If 'TRUE' this will eliminate taxa levels that contain only zero values and 'zeroValue' is ignored.

zeroValue  If 'elimZero' is 'FALSE' this parameter will fill in one zero with the number provided, at each all zero taxa level. The default is 0.00001. (See Details)

**Details**
This function does the opposite of transformHMPtoHMPTree.

Some functions will not work properly in the HMP package if a taxa level exists with all zeros. 'elimZero' and 'zeroValue' allow for an easy way to remove such levels or modify them with a small value in order to use them. Generally the HMP package requires data sets that contain only 1 taxa level and therefore 'trimToTaxaLevel' should also be used with this function.
Value

A data set of values in which each column is a separate taxa level and each row is a separate sample.

Author(s)

Berkley Shands, Patricio S. La Rosa, Elena Deych, William D. Shannon

Examples

data(saliva)

### Trims saliva to only contain the class level
salivaClass <- trimToTaxaLevel(saliva, "class", TRUE)

### This transforms the saliva data set but retains
### any zero rows that may exist.
transSaliva <- transformHMPtreeToHMP(salivaClass, FALSE, 0)

---

trimToTaxaLevel  Trim a Tree to a Given Level

Description

This function will take a tree and either remove all nodes lower than the given level or will remove all nodes not of the given level.

Usage

trimToTaxaLevel(data, level = "genus", eliminateParentNodes = FALSE, trimBelow = NULL, split = ".")

Arguments

data  A data frame in which each column contains the rdp read counts for every taxa given in the row names.

level  The depth the tree will go down to (kingdom, phylum, class, order, family, genus, species, subspecies). Genus is the default.

eliminateParentNodes

If 'TRUE' the data set returned will only contain rows at the level specified by 'myTaxaLevel'. If 'FALSE' the data set returned will contain all the nodes up to the level specified by 'myTaxaLevel'.

trimBelow

If 'NULL' the function will pull out only the data at the level specified by 'myTaxaLevel'. If 'TRUE' the function will remove all the levels below the specified level. If 'FALSE' the function will remove all the levels above the specified level.

split  This is the character that separates the taxa levels in the row names.
trimToTaxaLevel

Details

Notes:

1. For 'level' k, p, c, o, f, g, s and ss can be used in place of kingdom, phylum, class, order, family, genus, species and subspecies respectively.
2. Numbers can also be used for 'level', with no maximum limit.
3. The option to 'eliminateParentNodes' only works when 'trimBelow' is NULL.

Value

A new data set that has been trimmed to the level selected.

Author(s)

Berkley Shands, Patricio S. La Rosa, Elena Deych, William D. Shannon

Examples

data(saliva)

### Trims saliva to only contain the class level
salivaClass <- trimToTaxaLevel(saliva, "class", TRUE)
Index

*Topic datasets
  saliva, 15
  stool, 15
  throat, 16
*Topic package
  HMPTrees-package, 2

tcheckTreeValidity, 3
ccompareTwoDataSets, 4
ccreateAndPlot, 5
ccreateTrees, 7
ddisplayLegend, 8
dformatData, 8
ggenerateTree, 9
ggetMLEandLoglike, 10

HMPTrees (HMPTrees-package), 2
HMPTrees-package, 2
mergeDataSets, 11

plotTree, 12
plotTreeDataMDS, 14

saliva, 15
stool, 15

throat, 16
ttransformHMPtoHMPTree, 16
ttransformHMPTreesToHMP, 17
ttrimToTaxaLevel, 18