Package 'iNEXT.OTU'

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Type Package			
Title Interpolation and Extrapolation for UniFrac based on dissimilarity measure			
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Description iNEXT.OTU provides the function to calculate UniFrac for multiple assemblages.			
Depends R (>= 4.0)			
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R topics documented:			
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gghierPD

ggplot2 extension for an hierPD object

Description

gghierPD: the ggplot extension for hierPD object to plot order q against to hierarchical phylogenetic diversity decomposition and dissimilarity measure.

Usage

```
gghierPD(output, method = "A")
```

Arguments

output the output from hierPD.

method (method = "A") diversity(alpha, gamma); (method = "B") beta diversity; (method

= "D") dissimilarity measure based on multiplicative decomposition.

Value

a figure for hierarchical phylogenetic diversity decomposition or dissimilarity measure.

Examples

```
data("antechinus")
data("antechinus_mat")
data("antechinus_tree")
hier_output <- hierPD(antechinus, mat = antechinus_mat, tree = antechinus_tree, q = seq(0, 2, 0.2))
gghierPD(hier_output, method = "A")</pre>
```

ggiNEXTOTU

ggplot2 extension for an iNEXT.OTU object

Description

ggiNEXTOTU: the ggplot extension for iNEXTOTU object to plot coverage- or sample-sized-based rarefaction/extrapolation curves for phylogenetic diversity decomposition and dissimilarity measure.

Usage

```
ggiNEXTOTU(output, type = "B", scale = "fixed", transp = 0.4)
```

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Arguments

output	the output from iNEXTOTU
type	(required only when base = "coverage"), selection of plot type: type = 'B' for plotting the gamma, alpha, and beta diversity; type = 'D' for plotting 4 turnover dissimilarities.
scale	Are scales shared across all facets (the default, "fixed"), or do they vary across rows ("free_x"), columns ("free_y"), or both rows and columns ("free")?
transp	a value between 0 and 1 for controlling transparency. transp = 0 is completely transparent, default is 0.4.

Value

a figure for phylogenetic diversity decomposition or dissimilarity measure.

Examples

```
data("tongue_cheek")
data("tongue_cheek_tree")
output <- iNEXTOTU(tongue_cheek, q=c(0,1,2), nboot = 0, PDtree = tongue_cheek_tree)
ggiNEXTOTU(output, scale = 'free', transp = 0.4)</pre>
```

hierPD

function to calculate hierarchical phylogenetic gamma, alpha, beta diversity and dissimilarity measure

Description

hierPD: function to calculate empirical estimates for hierarchical phylogenetic gamma, alpha, beta diversity and dissimilarity measure

Usage

```
hierPD(
   data,
   mat,
   tree,
   q = seq(0, 2, 0.2),
   weight = "size",
   nboot = 20,
   conf = 0.95,
   type = "mle",
   decomposition = "relative"
)
```

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Arguments

data	data should be input as a matrix/data.frame (species by assemblages).
mat	hierarchical structure of data should be input as a matrix.
tree	a phylogenetic tree in Newick format for all observed species in the pooled assemblage.
q	a numerical vector specifying the diversity orders. Default is $seq(0, 2, 0.2)$.
weight	weight for relative decomposition. Default is "size".
nboot	a positive integer specifying the number of bootstrap replications when assessing sampling uncertainty and constructing confidence intervals. Bootstrap replications are generally time consuming. Enter \emptyset to skip the bootstrap procedures. Default is $2\emptyset$.
conf	a positive number < 1 specifying the level of confidence interval. Default is 0.95.
type	estimate type: estimate (type = "est"), empirical estimate (type = "mle"). Default is "mle".
decomposition	relative decomposition: (decomposition = "relative"), Absolute decomposition: (decomposition = "absolute").

Value

a data frames with hierarchical phylogenetic diversity (gamma, alpha, and beta) and four types dissimilarity measure.

Examples

```
data("antechinus")
data("antechinus_mat")
data("antechinus_tree")
hier_output <- hierPD(antechinus, mat = antechinus_mat, tree = antechinus_tree, q = seq(0, 2, 0.2))</pre>
```

iNEXTOTU function to calculate phylogenetic gamma, alpha, beta diversity and dissimilarity measure

Description

iNEXTOTU: function to calculate interpolation and extrapolation for phylogenetic gamma, alpha, beta diversity and dissimilarity measure

Usage

```
iNEXTOTU(
   data,
   q = c(0, 1, 2),
   base = "coverage",
   level = NULL,
   nboot = 10,
   conf = 0.95,
```

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```
PDtree = NULL,
PDreftime = NULL
)
```

Arguments

data OTU data can be input as a matrix/data.frame (species by assemblages), or a

list of matrices/data.frames, each matrix represents species-by-assemblages

abundance matrix.

q a numerical vector specifying the diversity orders. Default is c(0, 1, 2).

base sample-sized-based rarefaction and extrapolation for gamma and alpha diversity

(base = "size") or coverage-based rarefaction and extrapolation for gamma, alpha and beta diversity (base = "coverage"). Default is base = "coverage".

level A numerical vector specifying the particular value of sample coverage (between

0 and 1 when base = "coverage") or sample size (base = "size"). level = 1 (base = "coverage") means complete coverage (the corresponding diversity

represents asymptotic diversity).

If base = "size" and level = NULL, then this function computes the gamma and

alpha diversity estimates up to double the reference sample size.

If base = "coverage" and level = NULL, then this function computes the gamma and alpha diversity estimates up to one (for q = 1, 2) or up to the coverage of double the reference sample size (for q = 0); the corresponding beta diversity is

computed up to the same maximum coverage as the alpha diversity.

nboot a positive integer specifying the number of bootstrap replications when assessing

sampling uncertainty and constructing confidence intervals. Bootstrap replications are generally time consuming. Enter 0 to skip the bootstrap procedures.

Default is 10.

conf a positive number < 1 specifying the level of confidence interval. Default is

0.95.

PDtree a phylogenetic tree in Newick format for all observed species in the pooled

assemblage.

PDreftime a numerical value specifying reference time for PD. Default is NULL (i.e., the age

of the root of PDtree).

Value

If base = "coverage", return a list of seven data frames with three coverage-based diversity (gamma, alpha, and beta) and four types dissimilarity measure. If base = "size", return a list of two data frames with two diversity (gamma and alpha).

Examples

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tongue_cheek

tongue and cheek OTU count data

Usage

```
data("tongue_cheek")
```

Format

A data frame with 2549 observations on the following 2 variables.

Cheek a numeric vector

Tongue a numeric vector

Examples

```
data(tongue_cheek)
```

tongue_cheek_tree

phylogenetic for tongue and cheek data

Usage

```
data("tongue_cheek_tree")
```

Format

The format is: List of 4 \$ edge : int [1:25400, 1:2] 12702 12703 12704 12704 12705 12706 12706 12707 12707 12705 ... \$ edge.length: num [1:25400] 2.20e-03 9.91e-01 6.61e-03 7.87e-05 6.38e-03 ... \$ Nnode : int 12700 \$ tip.label : chr [1:12701] "OTU_97.15099" "OTU_97.13686" "OTU_97.30326" "OTU_97.26112" ... - attr(*, "class")= chr "phylo" - attr(*, "order")= chr "cladewise"

Examples

```
data(tongue_cheek_tree)
```

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