

Package: rdiversity (via r-universe)

September 13, 2024

Type Package

Title Measurement and Partitioning of Similarity-Sensitive Biodiversity

Version 2.2.0

Date 2022-04-29

Maintainer Richard Reeve <richard.reeve@glasgow.ac.uk>

URL <https://github.com/boydorr/rdiversity>

BugReports <https://github.com/boydorr/rdiversity/issues>

Description Provides a framework for the measurement and partitioning of the (similarity-sensitive) biodiversity of a metacommunity and its constituent subcommunities. Richard Reeve, et al. (2016) <[arXiv:1404.6520v3](https://arxiv.org/abs/1404.6520v3)>.

License GPL-3

Depends R (>= 2.10)

Imports methods, reshape2, stats, stringdist, utils

Suggests ape, testthat, knitr, markdown, rmarkdown, covr

LazyData true

RoxygenNote 7.1.2

Encoding UTF-8

VignetteBuilder knitr

Collate 'ancestral_nodes.R' 'binary.R' 'chainsaw.R'
'check_partition.R' 'check_phypartition.R' 'check_similarity.R'
'class-distance.R' 'class-metacommunity.R' 'class-powermean.R'
'class-relativeentropy.R' 'class-similarity.R'
'descendant_tips.R' 'similarity.R' 'dist2sim.R' 'distance.R'
'metadiv.R' 'subdiv.R' 'metacommunity.R'
'diversity-components.R' 'diversity-measures.R' 'gen2dist.r'
'geneid.R' 'genevec.R' 'hs_parameters.R' 'inndiv.R'
'phy2branch.R' 'phy2dist.R' 'phy_abundance.R' 'phy_struct.R'
'power_mean.R' 'powermean.R' 'rdiversity-package.R'

'relativeentropy.R' 'repartition.R' 'smatrix.R' 'summarise.R'
 'tax2dist.R' 'taxfac.R' 'taxid.R' 'taxmask.R' 'taxvec.R'
 'tbar.R' 'zmatrix.R'

Repository <https://boydorr.r-universe.dev>

RemoteUrl <https://github.com/boydorr/rdiversity>

RemoteRef HEAD

RemoteSha 77e4212827691ca9d2c402214fd2eb61f5f9de40

Contents

rdiversity-package	3
as.binary	4
binAdd	5
binary	6
binSeq	7
byte	7
bytesNeeded	8
chainsaw	8
dist2sim	9
distance	10
distance-class	10
fillUpToBit	11
fillUpToByte	12
gen2dist	12
inndiv	13
is.binary	14
loadAttributes	15
metacommunity	15
metacommunity-class	17
metadiv	18
meta_gamma	19
negate	20
norm_alpha	21
norm_beta	22
norm_meta_alpha	23
norm_meta_beta	24
norm_meta_rho	25
norm_rho	26
norm_sub_alpha	27
norm_sub_beta	28
norm_sub_rho	29
Ops.binary	30
phy2branch	30
phy2dist	31
phy_abundance	31
phy_struct	32

powermean-class	32
power_mean	33
print.binary	34
raw_alpha	34
raw_beta	35
raw_gamma	36
raw_meta_alpha	37
raw_meta_beta	38
raw_meta_rho	39
raw_rho	40
raw_sub_alpha	41
raw_sub_beta	42
raw_sub_rho	42
relativeentropy-class	43
repartition	44
saveAttributes	45
similarity	45
similarity-class	46
subdiv	46
sub_gamma	48
summary.binary	49
switchEndianness	49
tax2dist	50
Index	52

rdiversity-package *rdiversity: diversity measurement in R*

Description

rdiversity is an R package based around a framework for measuring and partitioning biodiversity using similarity-sensitive diversity measures. It provides functionality for measuring alpha, beta and gamma diversity of metacommunities (*e.g.* ecosystems) and their constituent subcommunities, where similarity may be defined as taxonomic, phenotypic, genetic, phylogenetic, functional, and so on. It uses the diversity measures described in the arXiv paper, ‘*How to partition diversity*’.

Details

- For more information go to our GitHub page; <https://github.com/boydorr/rdiversity>
- Please raise an issue if you find any problems; <https://github.com/boydorr/rdiversity/issues>
- This package is cross-validated against our Julia package; <https://github.com/EcoJulia/Diversity.jl>

Author(s)

Sonia Mitchell
 Richard Reeve <richard.reeve@glasgow.ac.uk> (maintainer)

References

Reeve, R., T. Leinster, C. Cobbold, J. Thompson, N. Brummitt, S. Mitchell, and L. Matthews. 2016. How to partition diversity. (<https://arxiv.org/abs/1404.6520>)

 as.binary

as binary digit.

Description

Converts an integer (Base10) to a binary (Base2) number. It also converts a logical vector to a binary (Base2) number (see examples).

Usage

```
as.binary(x, signed=FALSE, littleEndian=FALSE, size=2, n=0, logic=FALSE)
```

Arguments

x	integer or logical vector.
signed	TRUE or FALSE. Unsigned by default. (two's complement)
littleEndian	if TRUE. Big Endian if FALSE.
size	in Byte. Needed if signed is set. (by default 2 Byte)
n	in Bit. Can be set if unsigned is set to TRUE. (by default 0 Bit = auto)
logic	If set to TRUE, x is expected as logical vector.

Details

The binary number is represented by a logical vector. The bit order usually follows the same endianness as the byte order. No floating-point support. If logic is set to TRUE an integer vector is interpreted as a logical vector (>0 becomes TRUE and 0 becomes FALSE)

- Little Endian (LSB) —> (MSB)
- Big Endian (MSB) <— (LSB)

Auto switch to signed if num < 0.

Value

a vector of class binary.

See Also

[is.binary](#) and [binary](#)

Examples

```
as.binary(0xAF)
as.binary(42)
as.binary(42, littleEndian=TRUE)
as.binary(c(0xAF, 0xBF, 0xFF))
as.binary(c(2,4,8,16,32), signed=TRUE, size=1)
as.binary(-1, signed=TRUE, size=1)
as.binary(1:7, n=3)
as.binary(sample(2^8,3),n=8)
as.binary(c(1,1,0), signed=TRUE, logic=TRUE)
as.binary(c(TRUE,TRUE,FALSE), logic=TRUE)
```

binAdd

Binary Addition (+)

Description

Adds two binary numbers. (x + y)

Usage

```
binAdd(x, y)
```

Arguments

x	summand 1 (binary vector)
y	summand 2 (binary vector)

Details

Little-Endian and unsigned is not supported at the moment. No floating point supported. if x or y is signed the return value will also be signed.

Value

The sum of x and y. Returns a binary vector.

See Also

`base::as.logical` , `base::is.logical`, `base::raw`

Examples

```
five <- as.binary(5); ten <- as.binary(10);
as.numeric(rdiversity:::binAdd(ten, five))
rdiversity:::binAdd(as.binary(c(0,1), logic=TRUE), as.binary(c(1,0), logic=TRUE))
```

 binary

Binary digit.

Description

Create objects of type binary.

Usage

```
binary(n, signed=FALSE, littleEndian=FALSE)
```

Arguments

n	length of vector. Number of bits
signed	TRUE or FALSE. Unsigned by default. (two's complement)
littleEndian	if TRUE. Big Endian if FALSE.

Details

The binary number is represented by a *logical* vector. The bit order usually follows the same endianness as the byte order. How to read:

- Little Endian (LSB) \rightarrow (MSB)
- Big Endian (MSB) \leftarrow (LSB)

The Big Endian endianness stores its MSB at the lowest adress. The Little Endian endianness stores its MSB at the highest adress.

e.g. `b <- binary(8)`.

- "Little Endian" : MSB at `b[1]` and LSB at `b[8]`.
- "Big Endian" : LSB at `b[1]` and MSB at `b[8]`.

No floating-point support.

Value

a vector of class `binary` of length `n`. By default filled with `zeros(0)`.

See Also

[as.binary](#) and [is.binary](#).

Examples

```
b <- rdiversity:::binary(8)
summary(b)
b <- rdiversity:::binary(16, signed=TRUE)
summary(b)
b <- rdiversity:::binary(32, littleEndian=TRUE)
summary(b)
```

binSeq	<i>Binary sequence</i>
--------	------------------------

Description

Binary sequence.

Usage

```
binSeq(x, ...)
```

Arguments

x	a sequence.
...	used for dec2bin().

Value

a sequence list of binary digits.

See Also

[binary](#)

Examples

```
rdiversity:::binSeq(0:4)
```

byte	<i>A simple helper function to return the size of one byte</i>
------	--

Description

Used to increase readability

Usage

```
byte()
```

Value

The size of one byte (8)

See Also

[fillUpToByte](#)

bytesNeeded	<i>Minimum number of "byte" needed to hold n "bit"</i>
-------------	--

Description

A simple helper function that returns the minimum number of byte needed to hold the amount of n bit.

Usage

```
bytesNeeded(n)
```

Arguments

n The number of bit.

Value

The number of minimum byte needed to hold n bit.

See Also

[fillUpToByte](#) or [byte](#)

Examples

```
ten <- as.binary(10)
rdiversity:::bytesNeeded(length(ten))
```

chainsaw	<i>Function to cut the phylogeny to a specified depth from the tip with the greatest distance from the root.</i>
----------	--

Description

Function to cut the phylogeny to a specified depth from the tip with the greatest distance from the root.

Usage

```
chainsaw(partition, ps, depth)
```


Arguments

partition	two-dimensional matrix of mode numeric with rows as types (species), columns as subcommunities, and each element containing the relative abundance of types in each subcommunity relative to the metacommunity as a whole. In the phylogenetic case, this corresponds to the proportional abundance of historical species, which is calculated from the proportional abundance of terminal taxa
ps	phy_struct() output
depth	proportion of total tree height to be conserved (taken as a proportion from the highest tip). Describes how far back we go in the tree, with 0 marking the date of the most recent tip, and 1 marking the most recent common ancestor. Numbers greater than 1 extend the root of the tree

Value

chainsaw() returns an object of class metacommunity

dist2sim	<i>Distance to similarity</i>
----------	-------------------------------

Description

Converts distance objects into similarity objects.

Usage

```
dist2sim(dist, transform, k = 1, normalise = TRUE, max_d)
```

Arguments

dist	object of class distance
transform	object of class character, can be either "linear" or "exponential"
k	scaling parameter
normalise	object of class logical, which when TRUE will normalise distances to one
max_d	object of class numeric

Details

Distances can be transformed either **linearly** or **exponentially**. That is $1 - k * dist$ for non-negative values, or $\exp(-k * dist)$, respectively. If normalise is true, then $dist = dist / max_d$.

Value

dist2sim(x) returns an object of class similarity.

distance	<i>Generate distance object</i>
----------	---------------------------------

Description

Container for class distance.

Usage

```
distance(distance, dat_id)

## S4 method for signature 'matrix,character'
distance(distance, dat_id)

## S4 method for signature 'matrix,missing'
distance(distance, dat_id)
```

Arguments

distance	distance matrix
dat_id	object of class character denoting the type of diversity being calculated. This can be "naive", "genetic", "taxonomic", and so on

Value

distance() returns an object of class distance.

distance-class	<i>distance-class</i>
----------------	-----------------------

Description

Container for class distance.

Usage

```
## S4 method for signature 'distance'
show(object)
```

Arguments

object	object of class distance
--------	--------------------------

Fields

- `distance` two-dimensional matrix of mode numeric with rows as types, columns as types, and elements containing the pairwise distance of types
- `dat_id` object of class character describing the class of distance / similarity being used, e.g. "naive", "taxonomic", and so on
- `components` list containing the components necessary to calculate similarity. This list is empty when `precompute_dist = TRUE` when calculating distance. When a pairwise distance matrix is too large and `precompute_dist = FALSE`, this list contains all the information required to calculate pairwise distance between types

fillUpToBit	<i>Fill up to bit (000..)</i>
-------------	-------------------------------

Description

Fills up the binary number with zeros(0) or ones(1), to the size n in bit.

Usage

```
fillUpToBit(x, n, value=FALSE)
```

Arguments

- `x` The binary number to fill up with zeros. (Any binary vector).
- `n` size in bit.
- `value` to fill up with FALSE(0) or fill up with TRUE(1).

Details

No floating point supported.

Value

binary number. A binary vector with the desired size.

See Also

[fillUpToByte](#).

Examples

```
rdiversity:::fillUpToBit(as.binary(c(1,1), logic=TRUE), n=4)
rdiversity:::fillUpToBit(as.binary(c(1,0,1), logic=TRUE), n=4, value=FALSE)
```

fillUpToByte	<i>Fill up to Byte (00000000..)</i>
--------------	-------------------------------------

Description

Fills up the binary number with zeros(0) or ones(1), to the size in Byte.

Usage

```
fillUpToByte(x, size=0, value=FALSE)
```

Arguments

x	The binary number to fill up with zeros. (Any binary vector).
size	in Byte. 0 = auto (smallest possible Byte).
value	to fill up with FALSE(0) or fill up with TRUE(1).

Details

No floating point supported.

Value

binary number. A binary vector with the desired size.

See Also

[fillUpToBit](#).

Examples

```
rdiversity:::fillUpToByte(as.binary(c(1,1), logic=TRUE), size=2)
rdiversity:::fillUpToByte(as.binary(c(1,0,1), logic=TRUE), size=2, value=FALSE)
```

gen2dist	<i>Genetic distance matrix</i>
----------	--------------------------------

Description

Converts a vcfR object to a matrix of pairwise genetic distances.

Usage

```
gen2dist(vcf, biallelic = FALSE)
```

Arguments

vcf object of class data.frame.
biallelic logical describing whether the data is biallelic or not (default).

Value

gen2dist(x) returns an object of class distance containing a matrix of pairwise genetic distances.

inddiv	<i>Calculate individual-level diversity</i>
--------	---

Description

Generic function for calculating individual-level diversity.

Usage

```
inddiv(data, qs)

## S4 method for signature 'powermean'
inddiv(data, qs)

## S4 method for signature 'relativeentropy'
inddiv(data, qs)

## S4 method for signature 'metacommunity'
inddiv(data, qs)
```

Arguments

data matrix of mode numeric; containing diversity components
qs vector of mode numeric containing q values

Details

data may be input as three different classes:

- power_mean: calculates raw and normalised subcommunity alpha, rho or gamma diversity by taking the powermean of diversity components
- relativeentropy: calculates raw or normalised subcommunity beta diversity by taking the relative entropy of diversity components
- metacommunity: calculates all subcommunity measures of diversity

Value

inddiv() returns a standard output of class rdiv

References

Reeve, R., T. Leinster, C. Cobbold, J. Thompson, N. Brummitt, S. Mitchell, and L. Matthews. 2016. How to partition diversity. arXiv 1404.6520v3:1–9.

See Also

[subdiv](#) for subcommunity-level diversity and [metadiv](#) for metacommunity-level diversity.

Examples

```
# Define metacommunity
pop <- cbind.data.frame(A = c(1,1), B = c(2,0), C = c(3,1))
row.names(pop) <- paste0("sp", 1:2)
pop <- pop/sum(pop)
meta <- metacommunity(pop)

# Calculate subcommunity gamma diversity (takes the power mean)
g <- raw_gamma(meta)
inndiv(g, 0:2)

# Calculate subcommunity beta diversity (takes the relative entropy)
b <- raw_beta(meta)
inndiv(b, 0:2)

# Calculate all measures of individual diversity
inndiv(meta, 0:2)
```

is.binary

is Binary Vector

Description

test for object "binary".

Usage

```
is.binary(x)
```

Arguments

x object to test.

Value

TRUE or FALSE.

See Also

[as.binary](#) and [binary](#)

loadAttributes	<i>loadAttributes</i>
----------------	-----------------------

Description

Helper function load Attributes

Usage

```
loadAttributes(x, l)
```

Arguments

x	x
l	l

metacommunity	<i>Metacommunity</i>
---------------	----------------------

Description

Functions to generate a metacommunity object.

Usage

```
metacommunity(partition, similarity)

## S4 method for signature 'data.frame,missing'
metacommunity(partition)

## S4 method for signature 'numeric,missing'
metacommunity(partition)

## S4 method for signature 'matrix,missing'
metacommunity(partition)

## S4 method for signature 'missing,similarity'
metacommunity(partition, similarity)

## S4 method for signature 'numeric,similarity'
metacommunity(partition, similarity)

## S4 method for signature 'data.frame,similarity'
metacommunity(partition, similarity)

## S4 method for signature 'matrix,similarity'
metacommunity(partition, similarity)
```

Arguments

partition	two-dimensional matrix of mode numeric with rows as types, columns as sub-communities, and elements containing the relative abundances of types in sub-communities. For phylogenetic diversity, see <i>Details</i>
similarity	(optional) object of class similarity

Value

metacommunity() returns an object of class metacommunity (see *Fields*).

Fields

type_abundance	two-dimensional matrix of mode numeric with rows as types (species), columns as subcommunities, and each element containing the relative abundance of types in each sub-community relative to the metacommunity as a whole. In the phylogenetic case, this corresponds to the proportional abundance of historical species, which is calculated from the proportional abundance of terminal taxa
similarity	two-dimensional matrix of mode numeric with rows as types, columns as types, and elements containing pairwise similarities between types
similarity_components	list containing the components necessary to calculate similarity. This list is empty when precompute_dist = TRUE when calculating distance. When a pairwise distance matrix is too large and precompute_dist = FALSE, this list contains all the information required to calculate pairwise distance between types
similarity_parameters	list containing parameters associated with converting pairwise distances to similarities (the dist2sim() arguments)
ordinariness	two-dimensional matrix of mode numeric with rows as types, columns as sub-communities, and elements containing the ordinariness of types within subcommunities
subcommunity_weights	vector of mode numeric containing subcommunity weights
type_weights	two-dimensional matrix of mode numeric, with rows as types, columns as sub-communities, and elements containing weights of types within a subcommunity
dat_ID	object of class character denoting the type of diversity being calculated. This can be "naive", "genetic", "taxonomic", and so on
raw_abundance	[Phylogenetic] two-dimensional matrix of mode numeric with rows as types, columns as subcommunities, and elements containing the relative abundance of present day species
raw_structure	[Phylogenetic] two-dimensional matrix of mode numeric with rows as historical species, columns as present day species, and elements containing historical species lengths within lineages
parameters	[Phylogenetic] data.frame containing parameters associated with each historic species in the phylogeny

See Also

[metacommunity-class](#)

Examples

```
# Naive-type
partition <- cbind(a = c(1,1,1,0,0), b = c(0,1,0,1,1))
row.names(partition) <- paste0("sp", 1:5)
partition <- partition / sum(partition)
meta <- metacommunity(partition)
```

metacommunity-class *metacommunity-class*

Description

Container for class metacommunity.

Usage

```
## S4 method for signature 'metacommunity'
show(object)
```

Arguments

object object of class metacommunity

Fields

type_abundance two-dimensional matrix of mode numeric with rows as types (species), columns as subcommunities, and each element containing the relative abundance of types in each subcommunity relative to the metacommunity as a whole. In the phylogenetic case, this corresponds to the proportional abundance of historical species, which is calculated from the proportional abundance of terminal taxa

similarity two-dimensional matrix of mode numeric with rows as types, columns as types, and elements containing the pairwise similarity of types

similarity_components list containing the components necessary to calculate similarity. This list is empty when `precompute_dist = TRUE` when calculating distance. When a pairwise distance matrix is too large and `precompute_dist = FALSE`, this list contains all the information required to calculate pairwise distance between types

similarity_parameters list containing parameters associated with converting pairwise distances to similarities (the `dist2sim()` arguments)

ordinariness two-dimensional matrix of mode numeric with rows as types, columns as subcommunities, and elements containing the ordinariness of types within subcommunities

subcommunity_weights vector of mode numeric containing subcommunity weights

type_weights two-dimensional matrix of mode numeric, with rows as types, columns as subcommunities, and elements containing weights of types within a subcommunity

`dat_id` object of class character describing the class of distance / similarity being used, e.g. "naive", "taxonomic", and so on

`raw_abundance` [Phylogenetic] two-dimensional matrix of mode numeric with rows as types, columns as subcommunities, and elements containing the relative abundance of present day species

`raw_structure` [Phylogenetic] two-dimensional matrix of mode numeric with rows as historical species, columns as present day species, and elements containing historical species lengths within lineages

`parameters` [Phylogenetic] data.frame containing parameters associated with each historic species in the phylogeny

metadiv	<i>Metacommunity-level diversity</i>
---------	--------------------------------------

Description

Generic function for calculating metacommunity-level diversity.

Usage

```
metadiv(data, qs)

## S4 method for signature 'powermean'
metadiv(data, qs)

## S4 method for signature 'relativeentropy'
metadiv(data, qs)

## S4 method for signature 'metacommunity'
metadiv(data, qs)
```

Arguments

<code>data</code>	matrix of mode numeric; containing diversity components
<code>qs</code>	vector of mode numeric containing q values

Details

`data` may be input as one of three different classes:

- `powermean`: raw or normalised metacommunity alpha, rho or gamma diversity components; will calculate metacommunity-level raw or normalised metacommunity alpha, rho or gamma diversity
- `relativeentropy`: raw or normalised metacommunity beta diversity components; will calculate metacommunity-level raw or normalised metacommunity beta diversity
- `metacommunity`: will calculate all metacommunity measures of diversity

Value

metadiv() returns a standard output of class rdiv

References

Reeve, R., T. Leinster, C. Cobbold, J. Thompson, N. Brummitt, S. Mitchell, and L. Matthews. 2016. How to partition diversity. arXiv 1404.6520v3:1–9.

See Also

[inndiv](#) for type-level diversity and [subdiv](#) for subcommunity-level diversity.

Examples

```
# Define metacommunity
pop <- data.frame(a = c(1,3), b = c(1,1))
pop <- pop / sum(pop)
meta <- metacommunity(pop)

# Calculate metacommunity gamma diversity (takes the power mean)
g <- raw_gamma(meta)
metadiv(g, 0:2)

# Calculate metacommunity beta diversity (takes the relative entropy)
b <- raw_beta(meta)
metadiv(b, 0:2)

# Calculate all measures of metacommunity diversity
metadiv(meta, 0:2)
```

meta_gamma	<i>Metacommunity gamma diversity</i>
------------	--------------------------------------

Description

Calculates similarity-sensitive metacommunity gamma diversity (the metacommunity similarity-sensitive diversity). This measure may be calculated for a series of orders, represented as a vector of qs .

Usage

```
meta_gamma(meta, qs)
```

Arguments

meta	object of class metacommunity
qs	vector of mode numeric containing q values

Value

meta_gamma returns a standard output of class rdiv

References

R. Reeve, T. Leinster, C. Cobbold, J. Thompson, N. Brummitt, S. Mitchell, and L. Matthews. 2016. How to partition diversity. arXiv 1404.6520v3:1–9.

Examples

```
pop <- data.frame(a = c(1,3), b = c(1,1))
row.names(pop) <- paste0("sp", 1:2)
pop <- pop/sum(pop)
meta <- metacommunity(pop)

# Calculate metacommunity gamma diversity
meta_gamma(meta, 0:2)
```

negate

Binary Negation (!)

Description

Negates the binary number x. Negation $x \rightarrow -x$ or $-x \rightarrow x$

Usage

```
negate(x)
```

Arguments

x The number to be negated. A binary vector is expected.

Details

An »unsigned« number will be returned as »signed« regardless of whether the value is negative. No floating point supported.

Value

The negated number of x. Returns a binary vector with signed=TRUE

See Also

[switchEndianess](#) or [fillUpToByte](#).

Examples

```
summary(rdiversity::negate(as.binary(5, signed=TRUE)))
summary(rdiversity::negate(as.binary(-5, signed=TRUE)))
summary(rdiversity::negate(as.binary(5, signed=FALSE)))
```

norm_alpha	<i>Normalised alpha (low level diversity component)</i>
------------	---

Description

Calculates the low-level diversity component necessary for calculating normalised alpha diversity.

Usage

```
norm_alpha(meta)
```

Arguments

meta object of class metacommunity

Details

Values generated from norm_alpha() may be input into subdiv() and metadiv() to calculate normalised subcommunity and metacommunity alpha diversity.

Value

norm_alpha returns an object of class powermean

References

R. Reeve, T. Leinster, C. Cobbold, J. Thompson, N. Brummitt, S. Mitchell, and L. Matthews. 2016. How to partition diversity. arXiv 1404.6520v3:1–9.

Examples

```
pop <- data.frame(a = c(1,3), b = c(1,1))
row.names(pop) <- paste0("sp", 1:2)
pop <- pop/sum(pop)
meta <- metacommunity(pop)

# Calculate normalised alpha component
a <- norm_alpha(meta)
subdiv(a, 1)
metadiv(a, 1)
```

norm_beta	<i>Normalised beta (low level diversity component)</i>
-----------	--

Description

Calculates the low-level diversity component necessary for calculating normalised beta diversity.

Usage

```
norm_beta(meta)
```

Arguments

meta object of class metacommunity

Details

Values generated from `norm_beta()` may be input into `subdiv()` and `metadiv()` to calculate normalised subcommunity and metacommunity beta diversity.

Value

`norm_beta` returns an object of class `relativeentropy`

References

R. Reeve, T. Leinster, C. Cobbold, J. Thompson, N. Brummitt, S. Mitchell, and L. Matthews. 2016. How to partition diversity. arXiv 1404.6520v3:1–9.

Examples

```
pop <- data.frame(a = c(1,3), b = c(1,1))
row.names(pop) <- paste0("sp", 1:2)
pop <- pop/sum(pop)
meta <- metacommunity(pop)

# Calculate normalised beta component
b <- norm_beta(meta)
subdiv(b, 1)
metadiv(b, 1)
```

norm_meta_alpha	<i>Normalised metacommunity alpha diversity</i>
-----------------	---

Description

Calculates similarity-sensitive normalised metacommunity alpha diversity (the average similarity-sensitive diversity of subcommunities). This measure may be calculated for a series of orders, represented as a vector of qs .

Usage

```
norm_meta_alpha(meta, qs)
```

Arguments

meta	object of class metacommunity
qs	vector of mode numeric containing q values

Value

norm_meta_alpha returns a standard output of class rdiv

References

R. Reeve, T. Leinster, C. Cobbold, J. Thompson, N. Brummitt, S. Mitchell, and L. Matthews. 2016. How to partition diversity. arXiv 1404.6520v3:1–9.

Examples

```
pop <- data.frame(a = c(1,3), b = c(1,1))
row.names(pop) <- paste0("sp", 1:2)
pop <- pop/sum(pop)
meta <- metacommunity(pop)

# Calculate normalised metacommunity alpha diversity
norm_meta_alpha(meta, 0:2)
```

norm_meta_beta	<i>Normalised metacommunity beta diversity</i>
----------------	--

Description

Calculates similarity-sensitive normalised metacommunity beta diversity (the effective number of distinct subcommunities). This measure may be calculated for a series of orders, represented as a vector of qs .

Usage

```
norm_meta_beta(meta, qs)
```

Arguments

meta	object of class metacommunity
qs	vector of mode numeric containing q values

Value

norm_meta_beta returns a standard output of class rdiv

References

R. Reeve, T. Leinster, C. Cobbold, J. Thompson, N. Brummitt, S. Mitchell, and L. Matthews. 2016. How to partition diversity. arXiv 1404.6520v3:1–9.

Examples

```
pop <- data.frame(a = c(1,3), b = c(1,1))
row.names(pop) <- paste0("sp", 1:2)
pop <- pop/sum(pop)
meta <- metacommunity(pop)

# Calculate normalised metacommunity beta diversity
norm_meta_beta(meta, 0:2)
```

norm_meta_rho	<i>Normalised metacommunity rho diversity</i>
---------------	---

Description

Calculates similarity-sensitive normalised metacommunity rho diversity (the average representativeness of subcommunities). This measure may be calculated for a series of orders, represented as a vector of qs .

Usage

```
norm_meta_rho(meta, qs)
```

Arguments

meta	object of class metacommunity
qs	vector of mode numeric containing q values

Value

norm_meta_rho returns a standard output of class rdiv

References

R. Reeve, T. Leinster, C. Cobbold, J. Thompson, N. Brummitt, S. Mitchell, and L. Matthews. 2016. How to partition diversity. arXiv 1404.6520v3:1–9.

Examples

```
pop <- data.frame(a = c(1,3), b = c(1,1))
row.names(pop) <- paste0("sp", 1:2)
pop <- pop/sum(pop)
meta <- metacommunity(pop)

# Calculate normalised metacommunity rho diversity
norm_meta_rho(meta, 0:2)
```

norm_rho	<i>Normalised rho (low level diversity component)</i>
----------	---

Description

Calculates the low-level diversity component necessary for calculating normalised rho diversity.

Usage

```
norm_rho(meta)
```

Arguments

meta object of class metacommunity

Details

Values generated from `norm_rho()` may be input into `subdiv()` and `metadiv()` to calculate normalised subcommunity and metacommunity rho diversity.

Value

`norm_rho` returns an object of class `powermean`

References

R. Reeve, T. Leinster, C. Cobbold, J. Thompson, N. Brummitt, S. Mitchell, and L. Matthews. 2016. How to partition diversity. arXiv 1404.6520v3:1–9.

Examples

```
pop <- data.frame(a = c(1,3), b = c(1,1))
row.names(pop) <- paste0("sp", 1:2)
pop <- pop/sum(pop)
meta <- metacommunity(pop)

# Calculate normalised rho component
r <- norm_rho(meta)
subdiv(r, 1)
metadiv(r, 1)
```

norm_sub_alpha	<i>Normalised subcommunity alpha diversity</i>
----------------	--

Description

Calculates similarity-sensitive normalised subcommunity alpha diversity (the diversity of subcommunity j in isolation). This measure may be calculated for a series of orders, represented as a vector of qs .

Usage

```
norm_sub_alpha(meta, qs)
```

Arguments

meta	object of class metacommunity
qs	vector of mode numeric containing q values

Value

norm_sub_alpha returns a standard output of class rdiv

References

R. Reeve, T. Leinster, C. Cobbold, J. Thompson, N. Brummitt, S. Mitchell, and L. Matthews. 2016. How to partition diversity. arXiv 1404.6520v3:1–9.

Examples

```
pop <- data.frame(a = c(1,3), b = c(1,1))
row.names(pop) <- paste0("sp", 1:2)
pop <- pop/sum(pop)
meta <- metacommunity(pop)

# Calculate normalised subcommunity alpha diversity
norm_sub_alpha(meta, 0:2)
```

norm_sub_beta	<i>Normalised subcommunity beta diversity</i>
---------------	---

Description

Calculates similarity-sensitive normalised subcommunity beta diversity (an estimate of the effective number of distinct subcommunities). This measure may be calculated for a series of orders, represented as a vector of qs .

Usage

```
norm_sub_beta(meta, qs)
```

Arguments

meta	object of class metacommunity
qs	vector of mode numeric containing q values

Value

norm_sub_beta returns a standard output of class rdiv

References

R. Reeve, T. Leinster, C. Cobbold, J. Thompson, N. Brummitt, S. Mitchell, and L. Matthews. 2016. How to partition diversity. arXiv 1404.6520v3:1–9.

Examples

```
pop <- data.frame(a = c(1,3), b = c(1,1))
row.names(pop) <- paste0("sp", 1:2)
pop <- pop/sum(pop)
meta <- metacommunity(pop)

# Calculate normalised subcommunity beta diversity
norm_sub_beta(meta, 0:2)
```

norm_sub_rho	<i>Normalised subcommunity rho diversity</i>
--------------	--

Description

Calculates similarity-sensitive normalised subcommunity rho diversity (the representativeness of subcommunity j). This measure may be calculated for a series of orders, represented as a vector of qs .

Usage

```
norm_sub_rho(meta, qs)
```

Arguments

meta	object of class metacommunity
qs	vector of mode numeric containing q values

Value

norm_sub_rho returns a standard output of class rdiv

References

R. Reeve, T. Leinster, C. Cobbold, J. Thompson, N. Brummitt, S. Mitchell, and L. Matthews. 2016. How to partition diversity. arXiv 1404.6520v3:1–9.

Examples

```
pop <- data.frame(a = c(1,3), b = c(1,1))
row.names(pop) <- paste0("sp", 1:2)
pop <- pop/sum(pop)
meta <- metacommunity(pop)

# Calculate normalised subcommunity rho diversity
norm_sub_rho(meta, 0:2)
```

 Ops.binary

Group Generic Ops

Description

Group generic Ops operators

Usage

```
## S3 method for class 'binary'
Ops(e1, e2)
```

Arguments

e1	e1
e2	e2

 phy2branch

Phylogenetic similarity

Description

Packages all inputs into an object of class similarity.

Usage

```
phy2branch(tree, partition, depth = 1)
```

Arguments

tree	object of class phylo.
partition	two-dimensional matrix of mode numeric with rows as types (terminal taxa), columns as subcommunities, and each element containing the relative abundance of types in each subcommunity relative to the metacommunity as a whole.
depth	proportion of total tree height to be conserved (taken as a proportion from the highest tip). Describes how much evolutionary history should be retained, with 0 marking the date of the most recent tip, and 1 (the default) marking the most recent common ancestor. Numbers greater than 1 extend the root of the tree.

Value

phy2branch() returns an object of class similarity.

phy2dist	<i>Phylogenetic pairwise tip distance matrix</i>
----------	--

Description

Converts any phylo object to a matrix of pairwise tip-to-tip distances.

Usage

```
phy2dist(tree, precompute_dist = TRUE)
```

Arguments

tree	object of class phylo.
precompute_dist	object of class logical or numeric. When TRUE (by default) a distance matrix is generated and stored in slot distance, when FALSE no distance matrix is generated, and when numeric a distance matrix is generated until the number of species exceeds the defined value.

Value

phy2sim(x) returns an object of class distance containing a matrix of pairwise tip-to-tip distances.

phy_abundance	<i>Relative abundance of historical species</i>
---------------	---

Description

Calculates the relative abundance of historical species.

Usage

```
phy_abundance(partition, structure_matrix)
```

Arguments

partition	two-dimensional matrix of mode numeric with rows as types (species), columns as subcommunities, and each element containing the relative abundance of types in each subcommunity relative to the metacommunity as a whole. In the phylogenetic case, this corresponds to the proportional abundance of historical species, which is calculated from the proportional abundance of terminal taxa
structure_matrix	output\$structure of phy_struct().

phy_struct	<i>Calculate phylogenetic structure matrix</i>
------------	--

Description

Converts an object into class phylo into class phy_struct.

Usage

```
phy_struct(tree, partition)
```

Arguments

tree	object of class phylo
partition	two-dimensional matrix of mode numeric with rows as types (species), columns as subcommunities, and each element containing the relative abundance of types in each subcommunity relative to the metacommunity as a whole. In the phylogenetic case, this corresponds to the proportional abundance of historical species, which is calculated from the proportional abundance of terminal taxa

Value

phy_struct() returns a list containing:

\$structure	- each row denotes historical species, columns denote terminal taxa
\$tbar - the average distance from root to tip for all terminal taxa	
\$parameters	- information associated with each historical species
\$tree	- object of class phylo

powermean-class	<i>powermean-class</i>
-----------------	------------------------

Description

Container for class powermean.

Fields

results	data.frame containing rdiversity output
measure	object of class character naming the diversity measure being calculated

- `type_abundance` two-dimensional matrix of mode numeric with rows as types (species), columns as subcommunities, and each element containing the relative abundance of types in each subcommunity relative to the metacommunity as a whole. In the phylogenetic case, this corresponds to the proportional abundance of historical species, which is calculated from the proportional abundance of terminal taxa
- `ordinariness` two-dimensional matrix of mode numeric with rows as types, columns as subcommunities, and elements containing the ordinariness of types within subcommunities
- `subcommunity_weights` vector of mode numeric containing subcommunity weights
- `type_weights` two-dimensional matrix of mode numeric, with rows as types, columns as subcommunities, and elements containing weights of types within a subcommunity
- `dat_id` object of class character describing the class of distance / similarity being used, e.g. "naive", "taxonomic", and so on
- `similarity_components` list containing the components necessary to calculate similarity. This list is empty when `precompute_dist = TRUE` when calculating distance. When a pairwise distance matrix is too large and `precompute_dist = FALSE`, this list contains all the information required to calculate pairwise distance between types
- `similarity_parameters` list containing parameters associated with converting pairwise distances to similarities (the `dist2sim()` arguments)

power_mean

Power mean of vector elements

Description

`power_mean()` calculates the power mean of a set of values.

Usage

```
power_mean(values, order = 1, weights = rep(1, length(values)))
```

Arguments

<code>values</code>	Values for which to calculate mean.
<code>order</code>	Order of power mean.
<code>weights</code>	Weights of elements, normalised to 1 inside function.

Details

Calculates the order-th power mean of a single set of non-negative values, weighted by weights; by default, weights are equal and order is 1, so this is just the arithmetic mean. Equal weights and a order of 0 gives the geometric mean, and an order of -1 gives the harmonic mean.

Value

Weighted power mean

Examples

```
values <- sample(1:50, 5)
power_mean(values)
```

print.binary	<i>Print method for binary number.</i>
--------------	--

Description

This method prints the binary number.

Usage

```
## S3 method for class 'binary'
print(x, ...)
```

Arguments

x	any binary number.
...	further arguments.

Value

Output in ones and zeros (binary vector).

See Also

[summary.binary](#) provides some additional information.

raw_alpha	<i>Raw alpha (low level diversity component)</i>
-----------	--

Description

Calculates the low-level diversity component necessary for calculating alpha diversity.

Usage

```
raw_alpha(meta)
```

Arguments

meta	object of class metacommunity
------	-------------------------------

Details

Values generated from `raw_alpha()` may be input into `subdiv()` and `metadiv()` to calculate raw subcommunity and metacommunity alpha diversity.

Value

`raw_alpha` returns an object of class `powermean`

References

R. Reeve, T. Leinster, C. Cobbold, J. Thompson, N. Brummitt, S. Mitchell, and L. Matthews. 2016. How to partition diversity. arXiv 1404.6520v3:1–9.

Examples

```
pop <- data.frame(a = c(1,3), b = c(1,1))
row.names(pop) <- paste0("sp", 1:2)
pop <- pop/sum(pop)
meta <- metacommunity(pop)

# Calculate raw alpha component
a <- raw_alpha(meta)
subdiv(a, 1)
metadiv(a, 1)
```

`raw_beta`*Raw beta (low level diversity component)*

Description

Calculates the low-level diversity component necessary for calculating raw beta diversity.

Usage

```
raw_beta(meta)
```

Arguments

`meta` object of class `metacommunity`

Details

Values generated from `raw_beta()` may be input into `subdiv()` and `metadiv()` to calculate raw subcommunity and metacommunity beta diversity.

Value

`raw_beta` returns an object of class `relativeentropy`

References

R. Reeve, T. Leinster, C. Cobbold, J. Thompson, N. Brummitt, S. Mitchell, and L. Matthews. 2016. How to partition diversity. arXiv 1404.6520v3:1–9.

Examples

```
pop <- data.frame(a = c(1,3), b = c(1,1))
row.names(pop) <- paste0("sp", 1:2)
pop <- pop/sum(pop)
meta <- metacommunity(pop)

# Calculate raw beta component
b <- raw_beta(meta)
subdiv(b, 1)
metadiv(b, 1)
```

raw_gamma

Gamma (low level diversity component)

Description

Calculates the low-level diversity component necessary for calculating gamma diversity.

Usage

```
raw_gamma(meta)
```

Arguments

meta object of class metacommunity

Details

Values generated from `raw_gamma()` may be input into `subdiv()` and `metadiv()` to calculate sub-community and metacommunity gamma diversity.

Value

`raw_gamma` returns an object of class `powermean`

References

R. Reeve, T. Leinster, C. Cobbold, J. Thompson, N. Brummitt, S. Mitchell, and L. Matthews. 2016. How to partition diversity. arXiv 1404.6520v3:1–9.

Examples

```
pop <- cbind.data.frame(A = c(1,1), B = c(2,0), C = c(3,1))
row.names(pop) <- paste0("sp", 1:2)
pop <- pop/sum(pop)
meta <- metacommunity(pop)

# Calculate gamma component
g <- raw_gamma(meta)
subdiv(g, 1)
metadiv(g, 1)
```

raw_meta_alpha	<i>Raw metacommunity alpha diversity</i>
----------------	--

Description

Calculates similarity-sensitive raw metacommunity alpha diversity (the naive-community metacommunity diversity). This measure may be calculated for a series of orders, represented as a vector of qs .

Usage

```
raw_meta_alpha(meta, qs)
```

Arguments

meta	object of class metacommunity
qs	vector of mode numeric containing q values

Value

raw_meta_alpha returns a standard output of class rdiv

References

R. Reeve, T. Leinster, C. Cobbold, J. Thompson, N. Brummitt, S. Mitchell, and L. Matthews. 2016. How to partition diversity. arXiv 1404.6520v3:1–9.

Examples

```
pop <- data.frame(a = c(1,3), b = c(1,1))
row.names(pop) <- paste0("sp", 1:2)
pop <- pop/sum(pop)
meta <- metacommunity(pop)

# Calculate raw metacommunity alpha diversity
raw_meta_alpha(meta, 0:2)
```

raw_meta_beta	<i>Raw metacommunity beta diversity</i>
---------------	---

Description

Calculates similarity-sensitive raw metacommunity beta diversity (the average distinctiveness of subcommunities). This measure may be calculated for a series of orders, represented as a vector of qs .

Usage

```
raw_meta_beta(meta, qs)
```

Arguments

meta	object of class metacommunity
qs	vector of mode numeric containing q values

Value

raw_meta_beta returns a standard output of class rdiv

References

R. Reeve, T. Leinster, C. Cobbold, J. Thompson, N. Brummitt, S. Mitchell, and L. Matthews. 2016. How to partition diversity. arXiv 1404.6520v3:1–9.

Examples

```
pop <- data.frame(a = c(1,3), b = c(1,1))
row.names(pop) <- paste0("sp", 1:2)
pop <- pop/sum(pop)
meta <- metacommunity(pop)

# Calculate raw metacommunity beta diversity
raw_meta_beta(meta, 0:2)
```

raw_meta_rho	<i>Raw metacommunity rho diversity</i>
--------------	--

Description

Calculates similarity-sensitive raw metacommunity rho diversity (the average redundancy of sub-communities). This measure may be calculated for a series of orders, represented as a vector of qs .

Usage

```
raw_meta_rho(meta, qs)
```

Arguments

meta	object of class metacommunity
qs	vector of mode numeric containing q values

Value

raw_meta_rho returns a standard output of class rdiv

References

R. Reeve, T. Leinster, C. Cobbold, J. Thompson, N. Brummitt, S. Mitchell, and L. Matthews. 2016. How to partition diversity. arXiv 1404.6520v3:1–9.

Examples

```
pop <- data.frame(a = c(1,3), b = c(1,1))
row.names(pop) <- paste0("sp", 1:2)
pop <- pop/sum(pop)
meta <- metacommunity(pop)

# Calculate metacommunity rho diversity
raw_meta_rho(meta, 0:2)
```

raw_rho	<i>Raw rho (low level diversity component)</i>
---------	--

Description

Calculates the low-level diversity component necessary for calculating raw rho diversity.

Usage

```
raw_rho(meta)
```

Arguments

meta object of class metacommunity

Details

Values generated from `raw_rho()` may be input into `subdiv()` and `metadiv()` to calculate raw subcommunity and metacommunity rho diversity.

Value

`raw_rho` returns an object of class `powermean`

References

R. Reeve, T. Leinster, C. Cobbold, J. Thompson, N. Brummitt, S. Mitchell, and L. Matthews. 2016. How to partition diversity. arXiv 1404.6520v3:1–9.

Examples

```
pop <- data.frame(a = c(1,3), b = c(1,1))
row.names(pop) <- paste0("sp", 1:2)
pop <- pop/sum(pop)
meta <- metacommunity(pop)

# Calculate raw rho component
r <- raw_rho(meta)
subdiv(r, 1)
metadiv(r, 1)
```

raw_sub_alpha	<i>Raw subcommunity alpha diversity</i>
---------------	---

Description

Calculates similarity sensitive raw subcommunity alpha diversity (an estimate of naive-community metacommunity diversity). This measure may be calculated for a series of orders, represented as a vector of qs .

Usage

```
raw_sub_alpha(meta, qs)
```

Arguments

meta	object of class metacommunity
qs	vector of mode numeric containing q values

Value

raw_sub_alpha returns a standard output of class rdiv

References

R. Reeve, T. Leinster, C. Cobbold, J. Thompson, N. Brummitt, S. Mitchell, and L. Matthews. 2016. How to partition diversity. arXiv 1404.6520v3:1–9.

Examples

```
pop <- data.frame(a = c(1,3), b = c(1,1))
row.names(pop) <- paste0("sp", 1:2)
pop <- pop/sum(pop)
meta <- metacommunity(pop)

# Calculate raw subcommunity alpha diversity
raw_sub_alpha(meta, 0:2)
```

raw_sub_beta	<i>Raw subcommunity beta diversity</i>
--------------	--

Description

Calculates similarity-sensitive raw subcommunity beta diversity (the distinctiveness of subcommunity j). This measure may be calculated for a series of orders, represented as a vector of qs .

Usage

```
raw_sub_beta(meta, qs)
```

Arguments

meta	object of class <code>metacommunity</code>
qs	vector of mode numeric containing q values

Value

`raw_sub_beta` returns a standard output of class `rdiv`

References

R. Reeve, T. Leinster, C. Cobbold, J. Thompson, N. Brummitt, S. Mitchell, and L. Matthews. 2016. How to partition diversity. arXiv 1404.6520v3:1–9.

Examples

```
pop <- data.frame(a = c(1,3), b = c(1,1))
row.names(pop) <- paste0("sp", 1:2)
pop <- pop/sum(pop)
meta <- metacommunity(pop)

# Calculate raw subcommunity beta diversity
raw_sub_beta(meta, 0:2)
```

raw_sub_rho	<i>Raw subcommunity rho diversity</i>
-------------	---------------------------------------

Description

Calculates similarity-sensitive raw subcommunity rho diversity (the redundancy of subcommunity j). This measure may be calculated for a series of orders, represented as a vector of qs .

Usage

```
raw_sub_rho(meta, qs)
```

Arguments

`meta` object of class `metacommunity`
`qs` vector of mode numeric containing q values

Value

`raw_sub_rho` returns a standard output of class `rdiv`

References

R. Reeve, T. Leinster, C. Cobbold, J. Thompson, N. Brummitt, S. Mitchell, and L. Matthews. 2016. How to partition diversity. arXiv 1404.6520v3:1–9.

Examples

```
pop <- data.frame(a = c(1,3), b = c(1,1))
row.names(pop) <- paste0("sp", 1:2)
pop <- pop/sum(pop)
meta <- metacommunity(pop)

# Calculate raw subcommunity rho diversity
raw_sub_rho(meta, 0:2)
```

relativeentropy-class *relativeentropy-class*

Description

Container for class `relativeentropy`.

Fields

`results` data.frame containing `rdiversity` output
`measure` object of class `character` naming the diversity measure being calculated
`type_abundance` two-dimensional matrix of mode numeric with rows as types (species), columns as subcommunities, and each element containing the relative abundance of types in each subcommunity relative to the metacommunity as a whole. In the phylogenetic case, this corresponds to the proportional abundance of historical species, which is calculated from the proportional abundance of terminal taxa
`ordinariness` two-dimensional matrix of mode numeric with rows as types, columns as subcommunities, and elements containing the ordinariness of types within subcommunities

`subcommunity_weights` vector of mode numeric containing subcommunity weights

`type_weights` two-dimensional matrix of mode numeric, with rows as types, columns as subcommunities, and elements containing weights of types within a subcommunity

`dat_id` object of class character describing the class of distance / similarity being used, e.g. "naive", "taxonomic", and so on

`similarity_components` list containing the components necessary to calculate similarity. This list is empty when `precompute_dist = TRUE` when calculating distance. When a pairwise distance matrix is too large and `precompute_dist = FALSE`, this list contains all the information required to calculate pairwise distance between types

`similarity_parameters` list containing parameters associated with converting pairwise distances to similarities (the `dist2sim()` arguments)

repartition

Repartition metacommunity

Description

Randomly reshuffles the relative abundance of types (*e.g.* species) in a metacommunity (whilst maintaining the relationship between the relative abundance of a particular species across subcommunities). In the case of a phylogenetic metacommunity, the relative abundance of terminal taxa are randomly reshuffled and the relative abundance of types (historical species) are calculated from the resulting partition.

Usage

```
repartition(meta, new_partition)
```

Arguments

`meta` object of class `metacommunity`.

`new_partition` two-dimensional matrix of mode numeric with rows as types (species), columns as subcommunities, and each element containing the relative abundance of types in each subcommunity relative to the metacommunity as a whole. In the phylogenetic case, this corresponds to the proportional abundance of terminal taxa. If this argument is missing, all species / tips will be shuffled

Value

`repartition()` returns an object of class `metacommunity`

saveAttributes	<i>saveAttributes</i>
----------------	-----------------------

Description

Helper function save Attributes

Usage

```
saveAttributes(x)
```

Arguments

x	x
---	---

similarity	<i>Generate similarity object</i>
------------	-----------------------------------

Description

Container for class similarity.

Usage

```
similarity(similarity, dat_id)

## S4 method for signature 'matrix,character'
similarity(similarity, dat_id)

## S4 method for signature 'matrix,missing'
similarity(similarity, dat_id)
```

Arguments

similarity	similarity matrix
dat_id	object of class character denoting the type of diversity being calculated. This can be "naive", "genetic", "taxonomic", and so on

Value

similarity() returns an object of class similarity.

similarity-class	<i>similarity-class</i>
------------------	-------------------------

Description

Container for class similarity.

Usage

```
## S4 method for signature 'similarity'
show(object)
```

Arguments

object object of class similarity

Fields

similarity two-dimensional matrix of mode numeric with rows as types, columns as types, and elements containing the pairwise similarity of types

dat_id object of class character describing the class of distance / similarity being used, e.g. "naive", "taxonomic", and so on

components list containing the components necessary to calculate similarity. This list is empty when precompute_dist = TRUE when calculating distance. When a pairwise distance matrix is too large and precompute_dist = FALSE, this list contains all the information required to calculate pairwise distance between types

parameters list containing parameters associated with converting pairwise distances to similarities (the dist2sim() arguments)

subdiv	<i>Calculate subcommunity-level diversity</i>
--------	---

Description

Generic function for calculating subcommunity-level diversity.

Usage

```
subdiv(data, qs)
```

```
## S4 method for signature 'powermean'
subdiv(data, qs)
```

```
## S4 method for signature 'relativeentropy'
subdiv(data, qs)
```

```
## S4 method for signature 'metacommunity'  
subdiv(data, qs)
```

Arguments

data	matrix of mode numeric; containing diversity components
qs	vector of mode numeric containing q values

Details

data may be input as one of three different classes:

- powermean: raw or normalised metacommunity alpha, rho or gamma diversity components; will calculate subcommunity-level raw or normalised metacommunity alpha, rho or gamma diversity
- relativeentropy: raw or normalised metacommunity beta diversity components; will calculate subcommunity-level raw or normalised metacommunity beta diversity
- metacommunity: will calculate all subcommunity measures of diversity

Value

subdiv() returns a standard output of class rdiv

References

Reeve, R., T. Leinster, C. Cobbold, J. Thompson, N. Brummitt, S. Mitchell, and L. Matthews. 2016. How to partition diversity. arXiv 1404.6520v3:1–9.

See Also

[inddiv](#) for type-level diversity and [metadiv](#) for metacommunity-level diversity.

Examples

```
# Define metacommunity  
pop <- data.frame(a = c(1,3), b = c(1,1))  
row.names(pop) <- paste0("sp", 1:2)  
pop <- pop/sum(pop)  
meta <- metacommunity(pop)  
  
# Calculate subcommunity gamma diversity (takes the power mean)  
g <- raw_gamma(meta)  
subdiv(g, 0:2)  
  
# Calculate subcommunity beta diversity (takes the relative entropy)  
b <- raw_beta(meta)  
subdiv(b, 0:2)  
  
# Calculate all measures of subcommunity diversity  
subdiv(meta, 0:2)
```

sub_gamma	<i>Subcommunity gamma diversity</i>
-----------	-------------------------------------

Description

Calculates similarity-sensitive subcommunity gamma diversity (the contribution per individual toward metacommunity diversity). This measure may be calculated for a series of orders, represented as a vector of *qs*.

Usage

```
sub_gamma(meta, qs)
```

Arguments

<i>meta</i>	object of class <code>metacommunity</code>
<i>qs</i>	vector of mode <code>numeric</code> containing <i>q</i> values

Value

`sub_gamma` returns a standard output of class `rdiv`

References

R. Reeve, T. Leinster, C. Cobbold, J. Thompson, N. Brummitt, S. Mitchell, and L. Matthews. 2016. How to partition diversity. arXiv 1404.6520v3:1–9.

Examples

```
pop <- data.frame(a = c(1,3), b = c(1,1))
row.names(pop) <- paste0("sp", 1:2)
pop <- pop/sum(pop)
meta <- metacommunity(pop)

# Calculate subcommunity gamma diversity
sub_gamma(meta, 0:2)
```

summary.binary	<i>Summary method for binary number.</i>
----------------	--

Description

This method provides information about the attributes of the binary number.

Usage

```
## S3 method for class 'binary'  
summary(object, ...)
```

Arguments

object	binary number.
...	further arguments.

Value

Contains the following information:

- Signedness : unsigned or signed
- Endianess : Big-Endian or Little-Endian
- value<0 : negative or positive number
- Size[bit] : Size in bit
- Base10 : Decimal(Base10) number.

See Also

[print.binary](#)

switchEndianess	<i>Switch Endianess.</i>
-----------------	--------------------------

Description

Switch little-endian to big-endian and vice versa.

Usage

```
switchEndianess(x, stickyBits=FALSE)
```

Arguments

x	binary number. Any binary number.
stickyBits	Bits wont change if set TRUE. Only the attribute will be switched.

Value

switch little-endian to big-endian and vice versa.

See Also

[fillUpToByte](#).

Examples

```
x <- as.binary(c(1,1,0,0), logic=TRUE); print(x); summary(x);
y <- rdiversity:::switchEndianess(x); print(y); summary(y);
y <- rdiversity:::switchEndianess(x, stickyBits=TRUE); print(y); summary(y);
```

tax2dist

Generate taxonomic distance matrix

Description

Calculates taxonomic distances between species.

Usage

```
tax2dist(lookup, tax_distance, precompute_dist = TRUE)
```

Arguments

lookup	data.frame with colnames corresponding to nested taxonomic levels, e.g. c('Species', 'Genus', 'Family', 'Subclass')
tax_distance	vector with the distances attributed to taxonomic levels defined in lookup. The highest distance is the distance attributed to species that are not the same at any recorded taxonomic level. e.g. c(Species = 0, Genus = 1, Family = 2, Subclass = 3, Other = 4) from Shimatani.
precompute_dist	object of class logical or numeric. When TRUE (by default) a distance matrix is generated and stored in slot distance, when FALSE no distance matrix is generated, and when numeric a distance matrix is generated until the number of species exceeds the defined value.

Value

tax2dist() returns an object of class distance containing a matrix of pairwise taxonomic distances

References

Shimatani, K. 2001. On the measurement of species diversity incorporating species differences. *Oikos* 93:135–147.

Examples

```
# Create Lookup table
Species <- c("tenuifolium", "asterolepis", "simplex var.grandiflora", "simplex var.ochracea")
Genus <- c("Protium", "Quararibea", "Swartzia", "Swartzia")
Family <- c("Burseraceae", "Bombacaceae", "Fabaceae", "Fabaceae")
Subclass <- c("Sapindales", "Malvales", "Fabales", "Fabales")
lookup <- cbind.data.frame(Species, Genus, Family, Subclass)

# Assign values for each level (Shimatani's taxonomic distance)
tax_distance <- c(Species = 0, Genus = 1, Family = 2, Subclass = 3, Other = 4)

# Generate pairwise distances
distance <- tax2dist(lookup, tax_distance)
similarity <- dist2sim(distance, "linear")
```

Index

as.binary, [4](#), [6](#), [14](#)

binAdd, [5](#)

binary, [5](#), [6](#), [7](#), [14](#)

binSeq, [7](#)

byte, [7](#), [8](#)

bytesNeeded, [8](#)

chainsaw, [8](#)

dist2sim, [9](#)

distance, [10](#)

distance, matrix, character-method
(distance), [10](#)

distance, matrix, missing-method
(distance), [10](#)

distance-class, [10](#)

fillUpToBit, [11](#), [12](#)

fillUpToByte, [7](#), [8](#), [11](#), [12](#), [20](#), [50](#)

gen2dist, [12](#)

inndiv, [13](#), [19](#), [47](#)

inndiv, metacommunity-method (inndiv), [13](#)

inndiv, powermean-method (inndiv), [13](#)

inndiv, relativeentropy-method (inndiv),
[13](#)

is.binary, [5](#), [6](#), [14](#)

loadAttributes, [15](#)

meta_gamma, [19](#)

metacommunity, [15](#)

metacommunity, data.frame, missing-method
(metacommunity), [15](#)

metacommunity, data.frame, similarity-method
(metacommunity), [15](#)

metacommunity, data.frame-method
(metacommunity), [15](#)

metacommunity, matrix, missing-method
(metacommunity), [15](#)

metacommunity, matrix, similarity-method
(metacommunity), [15](#)

metacommunity, matrix-method
(metacommunity), [15](#)

metacommunity, missing, similarity-method
(metacommunity), [15](#)

metacommunity, numeric, missing-method
(metacommunity), [15](#)

metacommunity, numeric, similarity-method
(metacommunity), [15](#)

metacommunity, numeric-method
(metacommunity), [15](#)

metacommunity, similarity-method
(metacommunity), [15](#)

metacommunity-class, [17](#)

metadiv, [14](#), [18](#), [47](#)

metadiv, metacommunity-method (metadiv),
[18](#)

metadiv, powermean-method (metadiv), [18](#)

metadiv, relativeentropy-method
(metadiv), [18](#)

negate, [20](#)

norm_alpha, [21](#)

norm_beta, [22](#)

norm_meta_alpha, [23](#)

norm_meta_beta, [24](#)

norm_meta_rho, [25](#)

norm_rho, [26](#)

norm_sub_alpha, [27](#)

norm_sub_beta, [28](#)

norm_sub_rho, [29](#)

Ops.binary, [30](#)

phy2branch, [30](#)

phy2dist, [31](#)

phy_abundance, [31](#)

phy_struct, 32
power_mean, 33
powermean-class, 32
print.binary, 34, 49

raw_alpha, 34
raw_beta, 35
raw_gamma, 36
raw_meta_alpha, 37
raw_meta_beta, 38
raw_meta_rho, 39
raw_rho, 40
raw_sub_alpha, 41
raw_sub_beta, 42
raw_sub_rho, 42
rdiversity (rdiversity-package), 3
rdiversity-package, 3
relativeentropy-class, 43
repartition, 44

saveAttributes, 45
show, distance-method (distance-class),
10
show, metacommunity-method
(metacommunity-class), 17
show, similarity-method
(similarity-class), 46
similarity, 45
similarity, matrix, character-method
(similarity), 45
similarity, matrix, missing-method
(similarity), 45
similarity-class, 46
sub_gamma, 48
subdiv, 14, 19, 46
subdiv, metacommunity-method (subdiv), 46
subdiv, powermean-method (subdiv), 46
subdiv, relativeentropy-method (subdiv),
46
summary.binary, 34, 49
switchEndianess, 20, 49

tax2dist, 50