

# Package ‘mina’

January 26, 2026

**Title** Microbial community dIversity and Network Analysis

**Version** 1.19.0

**Description** An increasing number of microbiome datasets have been generated and analyzed with the help of rapidly developing sequencing technologies. At present, analysis of taxonomic profiling data is mainly conducted using composition-based methods, which ignores interactions between community members. Besides this, a lack of efficient ways to compare microbial interaction networks limited the study of community dynamics. To better understand how community diversity is affected by complex interactions between its members, we developed a framework (Microbial community dIversity and Network Analysis, mina), a comprehensive framework for microbial community diversity analysis and network comparison. By defining and integrating network-derived community features, we greatly reduce noise-to-signal ratio for diversity analyses. A bootstrap and permutation-based method was implemented to assess community network dissimilarities and extract discriminative features in a statistically principled way.

**Depends** R (>= 4.0.0)

**LinkingTo** Rcpp, RcppParallel, RcppArmadillo

**License** GPL

**Encoding** UTF-8

**Imports** methods, stats, Rcpp, MCL, RSpectra, apcluster, bigmemory, foreach, ggplot2, parallel, parallelDist, reshape2, plyr, biganalytics, stringr, Hmisc, utils

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**Enhances** doMC

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---

|        |                                                                                                                                                 |
|--------|-------------------------------------------------------------------------------------------------------------------------------------------------|
| .adj<- | <i>Setter for the slot ‘adj’ and ‘adj_sig’, the adjacency matrix of ‘norm’ and corresponding significant value matrix with ‘sig’ is ‘TRUE’.</i> |
|--------|-------------------------------------------------------------------------------------------------------------------------------------------------|

---

**Description**

Setter for the slot ‘adj’ and ‘adj\_sig’, the adjacency matrix of ‘norm’ and corresponding significant value matrix with ‘sig’ is ‘TRUE’.

Get the slot ‘adj’.

Get the slot ‘adj\_sig’.

**Usage**

```
.adj(x) <- value

## S4 replacement method for signature 'mina'
.adj(x) <- value

.adj(x)

adj_sig(x) <- value

## S4 replacement method for signature 'mina'
adj_sig(x) <- value

adj_sig(x)
```

**Arguments**

- x                    The ‘mina’ object.
- value                The value to set for the slot of the ‘mina’ object ‘x’.

**Value**

The 'adj' slot of the 'mina' object.

The slot 'adj\_sig' of the object.

---

`.dmr<-`*Setter and getter for the slot 'dmr'.*

---

**Description**

Setter and getter for the slot 'dmr'.

**Usage**

```
.dmr(x) <- value  
  
.dmr(x)  
  
## S4 replacement method for signature 'mina'  
.dmr(x) <- value  
  
## S4 method for signature 'mina'  
.dmr(x)
```

**Arguments**

|       |                                                         |
|-------|---------------------------------------------------------|
| x     | The 'mina' object.                                      |
| value | The value to set for the slot of the 'mina' object 'x'. |

**Value**

The 'dmr' slot of the 'mina' object.

**Examples**

```
maize <- new("mina", tab = maize_asv2, des = maize_des2)  
maize <- norm_tab(maize, method = "raref")  
maize <- fit_tabs(maize)  
maize <- com_dis(maize, method = "bray")  
maize <- dmr(maize)  
asv_dmr <- .dmr(maize)
```

---

|     |                                                    |
|-----|----------------------------------------------------|
| adj | <i>Calculate the correlation adjacency matrix.</i> |
|-----|----------------------------------------------------|

---

### Description

Calculate the correlation adjacency matrix.

### Usage

```
adj(x, method, ...)
```

### Arguments

|        |                                                                     |
|--------|---------------------------------------------------------------------|
| x      | An object of the class mina with 'norm' defined or a 'norm' matrix. |
| method | The correlation coefficient used for adjacency matrix.              |
| ...    | Additional parameters.                                              |

### Value

Adjacency matrix.

### Examples

```
maize <- new("mina", tab = maize_asv2, des = maize_des2)
maize <- norm_tab(maize, method = "raref", depth = 1000)
maize <- fit_tabs(maize)
maize <- adj(maize, method = "spearman", sig = FALSE)
```

---

|                       |                                                                                      |
|-----------------------|--------------------------------------------------------------------------------------|
| adj,matrix,ANY-method | <i>Calculate the adjacency matrix of 'norm' by correlation with matrix as input.</i> |
|-----------------------|--------------------------------------------------------------------------------------|

---

### Description

Calculate the adjacency matrix of 'norm' by correlation with matrix as input.

### Usage

```
## S4 method for signature 'matrix,ANY'
adj(x, method, sig = FALSE, threads = 80, nblocks = 400, ...)

## S4 method for signature 'matrix,character'
adj(x, method, sig = FALSE, threads = 80, nblocks = 400, ...)
```

**Arguments**

|         |                                                                                                                                      |
|---------|--------------------------------------------------------------------------------------------------------------------------------------|
| x       | An object of the class mina with 'norm' defined or a 'norm' matrix.                                                                  |
| method  | The correlation coefficient used for adjacency matrix.                                                                               |
| sig     | (optional) The asymptotic P-values, only applicable for Pearson and Spearman methods with 'mina' object as input, always FALSE here. |
| threads | The number of threads used for parallel running, 80 by default.                                                                      |
| nblocks | The number of row/column for splitting sub-matrix, 400 by default.                                                                   |
| ...     | Additional parameters.                                                                                                               |

**Value**

y The adjacency matrix.

**Examples**

```
asv_norm <- norm_tab(maize_asv2, method = "raref", depth = 1000)
asv_adj <- adj(asv_norm, method = "pearson")
```

---

|                     |                                                                                                   |
|---------------------|---------------------------------------------------------------------------------------------------|
| adj,mina,ANY-method | <i>Calculate the adjacency matrix of 'norm' by correlation with 'mina' class object as input.</i> |
|---------------------|---------------------------------------------------------------------------------------------------|

---

**Description**

Calculate the adjacency matrix of 'norm' by correlation with 'mina' class object as input.

**Usage**

```
## S4 method for signature 'mina,ANY'
adj(x, method, sig = FALSE, threads = 80, nblocks = 400, ...)

## S4 method for signature 'mina,character'
adj(x, method, sig = FALSE, threads = 80, nblocks = 400, ...)
```

**Arguments**

|         |                                                                                              |
|---------|----------------------------------------------------------------------------------------------|
| x       | An object of the class mina with 'norm' defined or a 'norm' matrix.                          |
| method  | The correlation coefficient used for adjacency matrix.                                       |
| sig     | The asymptotic P-values, only applicable for Pearson and Spearman methods, FALSE by default. |
| threads | The number of threads used for parallel running, 80 by default.                              |
| nblocks | The number of row/column for splitting sub-matrix, 400 by default.                           |
| ...     | Additional parameters.                                                                       |

Value

x The same ‘mina’ object with ‘adj’ added.

Examples

```
maize <- new("mina", tab = maize_asv2, des = maize_des2)
maize <- norm_tab(maize, method = "raref")
maize <- fit_tabs(maize)
maize <- adj(maize, method = "spearman", sig = FALSE)
```

---

|                 |                                                                                                   |
|-----------------|---------------------------------------------------------------------------------------------------|
| adj_method_list | <i>List of adjacency matrix calculation methods/ orrelations supported in <a href="#">adj</a></i> |
|-----------------|---------------------------------------------------------------------------------------------------|

---

Description

Correlation methods should be specified by exact string match.

Usage

```
adj_method_list
```

Format

- A list of character vectors.
- pearson** Pearson correlation.
  - spearman** Spearman correlation.
  - sparcc** SparCC correlation by spearman.

See Also

```
adj
```

Examples

```
? adj_method_list
```

---

|       |                                                                                                  |
|-------|--------------------------------------------------------------------------------------------------|
| bs_pm | <i>Inferring the network of different group of samples and test significance by permutation.</i> |
|-------|--------------------------------------------------------------------------------------------------|

---

## Description

Inferring the network of different group of samples and test significance by permutation.

## Usage

```
bs_pm(x, group, ...)
```

## Arguments

|       |                                                           |
|-------|-----------------------------------------------------------|
| x     | An object of class 'mina' with 'norm' and 'des' defined.  |
| group | The column name of descriptive file 'des' for comparison. |
| ...   | Additional parameters.                                    |

## Value

The network bootstrap and permutation result.

## Examples

```
maize <- new("mina", tab = maize_asv2, des = maize_des2)
maize <- norm_tab(maize, method = "raref")
maize <- fit_tabs(maize)
maize <- get_rep(maize, top = 5)
maize <- bs_pm(maize, group = "Compartment", per = 0.5)
```

---

|                       |                                                                                                  |
|-----------------------|--------------------------------------------------------------------------------------------------|
| bs_pm,mina,ANY-method | <i>Inferring the network of different group of samples and test significance by permutation.</i> |
|-----------------------|--------------------------------------------------------------------------------------------------|

---

## Description

Inferring the network of different group of samples and test significance by permutation.

**Usage**

```
## S4 method for signature 'mina,ANY'
bs_pm(
  x,
  group,
  g_size = 88,
  s_size = 30,
  rm = TRUE,
  per = 0.1,
  sig = TRUE,
  bs = 6,
  pm = 6,
  individual = FALSE,
  out_dir = "./",
  ...
)

## S4 method for signature 'mina,character'
bs_pm(
  x,
  group,
  g_size = 88,
  s_size = 30,
  rm = TRUE,
  per = 0.1,
  sig = TRUE,
  bs = 6,
  pm = 6,
  individual = FALSE,
  out_dir = "./",
  ...
)
```

**Arguments**

|                     |                                                                                                                                                                                             |
|---------------------|---------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| <code>x</code>      | An object of class ‘mina’ with @norm and @des defined.                                                                                                                                      |
| <code>group</code>  | The column name of descriptive file @des for comparison.                                                                                                                                    |
| <code>g_size</code> | The cutoff of group size used for filtering, default is 88.                                                                                                                                 |
| <code>s_size</code> | The number of samples used for network inference during bootstrap and permutation (when ‘sig’ is TRUE), it should be smaller than $g\_size / 2$ to make sure the randomness; default is 30. |
| <code>rm</code>     | Filtering the components present in less than ‘per’ of the samples from compared groups, default TRUE.                                                                                      |
| <code>per</code>    | The percentage of present samples for filtering, default is 0.1.                                                                                                                            |
| <code>sig</code>    | Whether to test the significance, skip the permutation when set as FALSE, default is TRUE.                                                                                                  |

|            |                                                                                                            |
|------------|------------------------------------------------------------------------------------------------------------|
| bs         | The times for bootstrap network inference, default is 6.                                                   |
| pm         | The times for permutated samples network inference, default is 6.                                          |
| individual | Whether to output the bootstrap and permutation results of each comparison individually, default is FALSE. |
| out_dir    | The output directory if 'individual' is TRUE, default is the current working directory                     |
| ...        | Additional parameters.                                                                                     |

**Value**

x The same object with @multi and @perm defined.

**Examples**

```
maize <- new("mina", tab = maize_asv2, des = maize_des2)
maize <- norm_tab(maize, method = "raref")
maize <- fit_tabs(maize)
maize <- bs_pm(maize, group = "Compartment", per = 0.5)
```

---

|            |                                                                                                                                                                     |
|------------|---------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| check_mina | <i>Check the tab and des file. Return TRUE if they are NULL or both quantitative and descriptive files of same samples are included (i.e. the object is valid).</i> |
|------------|---------------------------------------------------------------------------------------------------------------------------------------------------------------------|

---

**Description**

Check the tab and des file. Return TRUE if they are NULL or both quantitative and descriptive files of same samples are included (i.e. the object is valid).

**Usage**

```
check_mina(x)
```

**Arguments**

x An object of class mina.

**Value**

TRUE if the object is valid.

**Examples**

```
data(maize)
check_mina(maize)
```

---

|               |                                                                                                                                   |
|---------------|-----------------------------------------------------------------------------------------------------------------------------------|
| check_mina_de | <i>Check the object and return TRUE if the object includes descriptive table contains the same samples as quantitative table.</i> |
|---------------|-----------------------------------------------------------------------------------------------------------------------------------|

---

**Description**

Check the object and return TRUE if the object includes descriptive table contains the same samples as quantitative table.

**Usage**

check\_mina\_de(x)

**Arguments**

x                      An object of class mina with ‘tab’ and ‘des’ defined.

**Value**

TRUE if the object contains non-empty descriptive table and has the same samples as quantitative table.

**Examples**

```
data(maize)
check_mina_de(maize)
```

---

|               |                                                                                    |
|---------------|------------------------------------------------------------------------------------|
| check_mina_qu | <i>Check the object and return TRUE if the object includes quantitative table.</i> |
|---------------|------------------------------------------------------------------------------------|

---

**Description**

Check the object and return TRUE if the object includes quantitative table.

**Usage**

check\_mina\_qu(x)

**Arguments**

x                      An object of class mina with ‘tab’ defined.

**Value**

TRUE if the object contains quantitative table and is not empty.

Examples

```
data(maize)
check_mina_qu(maize)
```

---

|     |                            |
|-----|----------------------------|
| cls | <i>Get the slot 'cls'.</i> |
|-----|----------------------------|

---

Description

Get the slot 'cls'.

Usage

```
cls(x)
```

Arguments

x                      The 'mina' object.

Value

The 'cls' slot of the object.

---

|       |                                   |
|-------|-----------------------------------|
| cls<- | <i>Setter for the slot 'cls'.</i> |
|-------|-----------------------------------|

---

Description

Setter for the slot 'cls'.

Usage

```
cls(x) <- value

## S4 replacement method for signature 'mina'
cls(x) <- value
```

Arguments

x                      The 'mina' object.  
value                  The value to set for the slot of the 'mina' object 'x'.

Value

The 'cls' slot of the 'mina' object.

---

|         |                                |
|---------|--------------------------------|
| cls_tab | <i>Get the slot 'cls_tab'.</i> |
|---------|--------------------------------|

---

**Description**

Get the slot 'cls\_tab'.

**Usage**

```
cls_tab(x)
```

**Arguments**

|   |                    |
|---|--------------------|
| x | The 'mina' object. |
|---|--------------------|

**Value**

The 'cls\_tab' slot of the object.

**Examples**

```
cls_tab(maize)
```

---

|           |                                       |
|-----------|---------------------------------------|
| cls_tab<- | <i>Setter for the slot 'cls_tab'.</i> |
|-----------|---------------------------------------|

---

**Description**

Setter for the slot 'cls\_tab'.

**Usage**

```
cls_tab(x) <- value

## S4 replacement method for signature 'mina'
cls_tab(x) <- value
```

**Arguments**

|       |                                                         |
|-------|---------------------------------------------------------|
| x     | The 'mina' object.                                      |
| value | The value to set for the slot of the 'mina' object 'x'. |

**Value**

The 'cls\_tab' slot of the 'mina' object.

---

|         |                                                                 |
|---------|-----------------------------------------------------------------|
| com_dis | <i>Calculate the community dissimilarity / distance matrix.</i> |
|---------|-----------------------------------------------------------------|

---

**Description**

Calculate the community dissimilarity / distance matrix.

**Usage**

```
com_dis(x, method = "bray", ...)
```

**Arguments**

|        |                                                                              |
|--------|------------------------------------------------------------------------------|
| x      | An object of the class minia with 'norm' defined or any quantitative matrix. |
| method | The dissimilarity / distance method used, default 'bray'.                    |
| ...    | Additional parameters.                                                       |

**Value**

The distance / dissimilarity matrix.

**Examples**

```
asv_norm <- norm_tab(maize_asv2, method = "raref", depth = 1000)
asv_dis <- com_dis(asv_norm, method = "bray")
```

---

|                           |                                                                                     |
|---------------------------|-------------------------------------------------------------------------------------|
| com_dis,matrix,ANY-method | <i>Calculate the community dissimilarity / distance matrix of the input matrix.</i> |
|---------------------------|-------------------------------------------------------------------------------------|

---

**Description**

Calculate the community dissimilarity / distance matrix of the input matrix.

**Usage**

```
## S4 method for signature 'matrix,ANY'
com_dis(x, method = "bray", threads = 80, nblocks = 400, ...)

## S4 method for signature 'matrix,character'
com_dis(x, method = "bray", threads = 80, nblocks = 400, ...)
```

**Arguments**

|         |                                                                                                   |
|---------|---------------------------------------------------------------------------------------------------|
| x       | A matrix of the quantitative table.                                                               |
| method  | The dissimilarity / distance method used, default 'bray'.                                         |
| threads | (optional, only needed when method == "tina") The number of threads used for parallel running.    |
| nblocks | (optional, only needed when method == "tina") The number of row / column for splitted sub-matrix. |
| ...     | Additional parameters.                                                                            |

**Value**

y The dissimilarity / distance matrix.

**Examples**

```
asv_norm <- norm_tab(maize_asv2, method = "raref", depth = 1000)
asv_dis <- com_dis(asv_norm, method = "bray")
```

---

com\_dis,mina,ANY-method

*Calculate the community dissimilarity / distance matrix of 'norm' with 'mina' class object as input.*

---

**Description**

Calculate the community dissimilarity / distance matrix of 'norm' with 'mina' class object as input.

**Usage**

```
## S4 method for signature 'mina,ANY'
com_dis(x, method = "bray", threads = 80, nblocks = 400, ...)

## S4 method for signature 'mina,character'
com_dis(x, method = "bray", threads = 80, nblocks = 400, ...)
```

**Arguments**

|         |                                                                                                   |
|---------|---------------------------------------------------------------------------------------------------|
| x       | An object of the class 'mina' with 'norm' defined.                                                |
| method  | The dissimilarity / distance method used, default 'bray'.                                         |
| threads | (optional, only needed when method == "tina") The number of threads used for parallel running.    |
| nblocks | (optional, only needed when method == "tina") The number of row / column for splitted sub-matrix. |
| ...     | Additional parameters.                                                                            |

**Value**

x The same 'mina' object with @dis added.

**Examples**

```
maize <- new("mina", tab = maize_asv2, des = maize_des2)
maize <- norm_tab(maize, method = "total")
maize <- fit_tabs(maize)
maize <- com_dis(maize, method = "bray")
```

---

|              |                                                                                                                                                   |
|--------------|---------------------------------------------------------------------------------------------------------------------------------------------------|
| com_dis_list | <i>List of dissimilarity / distance supported in <a href="#">com_dis</a>. Dissimilarity / distance should be specified by exact string match.</i> |
|--------------|---------------------------------------------------------------------------------------------------------------------------------------------------|

---

**Description**

List of dissimilarity / distance supported in [com\\_dis](#). Dissimilarity / distance should be specified by exact string match.

**Usage**

```
com_dis_list
```

**Format**

A list of character vectors indicate the dissimilarity / distance method used.

tina TINA from Schmidt\_et\_al\_2016

Jaccard Jaccard defined by [vegdist](#)

**weighted** Dissimilarity / distance method for weighted matrix:

bhjattacharyya from [parDist](#)

canberra from [parDist](#)

bray from [parDist](#)

chord from [parDist](#)

divergence from [parDist](#)

euclidean from [parDist](#)

fJaccard from [parDist](#)

geodesic from [parDist](#)

hellinger from [parDist](#)

kullback from [parDist](#)

manhattan from [parDist](#)

maximum from [parDist](#)

minkowski from [parDist](#)

podani from [parDist](#)  
soergel from [parDist](#)  
wave from [parDist](#)  
whittaker from [parDist](#)  
**unweighted** Dissimilarity / Distance for unweighted matrix:  
binary from [parDist](#)  
braun-blanquet from [parDist](#)  
consine from [parDist](#)  
dice from [parDist](#)  
fager from [parDist](#)  
faith from [parDist](#)  
hamman from [parDist](#)  
hamming from [parDist](#)  
kulczynski1 from [parDist](#)  
kulczynski2 from [parDist](#)  
michael from [parDist](#)  
mountford from [parDist](#)  
mozley from [parDist](#)  
ochiai from [parDist](#)  
phi from [parDist](#)  
russel from [parDist](#)  
simple matching from [parDist](#)  
simpson from [parDist](#)  
stiles from [parDist](#)  
tanimoto from [parDist](#)  
yule from [parDist](#)  
yule2 from [parDist](#)

## Examples

? com\_dis\_list

---

|          |                                                                            |
|----------|----------------------------------------------------------------------------|
| com_plot | <i>Visulization of components distance / dissimilarity in k dimension.</i> |
|----------|----------------------------------------------------------------------------|

---

## Description

Visulization of components distance / dissimilarity in k dimension.

## Usage

```
com_plot(x, match, ...)
```

## Arguments

|       |                                                                                           |
|-------|-------------------------------------------------------------------------------------------|
| x     | An object of class 'mina' with 'dmr' and 'des' defined.                                   |
| match | The column name of the components IDs in 'des' which exactly the same indicated in 'dmr'. |
| ...   | Additional parameters.                                                                    |

## Value

The PCoA plot.

## Examples

```
maize <- new("mina", tab = maize_asv2, des = maize_des2)
maize <- norm_tab(maize, method = "raref", depth = 5000)
maize <- fit_tabs(maize)
maize <- com_dis(maize, method = "bray")
maize <- dmr(maize)
p1a <- com_plot(maize, match = "Sample_ID", color = "Compartment")
p1b <- com_plot(maize, match = "Sample_ID", d1 = 3, d2 = 4,
color = "Compartment")
p2a <- com_plot(maize, match = "Sample_ID", color = "Host_genotype")
p2b <- com_plot(maize, match = "Sample_ID", d1 = 1, d2 = 3, color =
"Host_genotype")
p3a <- com_plot(maize, match = "Sample_ID", color = "Compartment", shape =
"Soil")
p3b <- com_plot(maize, match = "Sample_ID", d1 = 1, d2 = 4, color =
"Compartment", shape = "Soil")
```

---

 com\_plot,mina,ANY-method

*Visulization of components distance / dissimilarity in k dimension.*


---

## Description

Visulization of components distance / dissimilarity in k dimension.

## Usage

```
## S4 method for signature 'mina,ANY'
com_plot(x, match, d1 = 1, d2 = 2, color, shape = NULL, ...)

## S4 method for signature 'mina,character'
com_plot(x, match, d1 = 1, d2 = 2, color, shape = NULL, ...)

## S4 method for signature 'mina,character'
com_plot(x, match, d1 = 1, d2 = 2, color, shape = NULL, ...)
```

## Arguments

|       |                                                                                        |
|-------|----------------------------------------------------------------------------------------|
| x     | An object of 'mina' with list 'dmr' defined.                                           |
| match | The column name of the components IDs in 'des' with exactly the same as rownames in x. |
| d1    | The dimension be visualized in x-axis, default '1'.                                    |
| d2    | The dimension be visualized in y-axis, default '2'.                                    |
| color | The column name in 'des' to be used for different color groups.                        |
| shape | The column name in 'des' to be used for different shape groups, default 'NULL'.        |
| ...   | Additional parameters.                                                                 |

## Value

p The plotted figure.  
The PCoA plot.

## Examples

```
maize <- new("mina", tab = maize_asv, des = maize_des)
maize <- norm_tab(maize, method = "raref")
maize <- fit_tabs(maize)
maize <- com_dis(maize, method = "bray")
maize <- dmr(maize)
p1a <- com_plot(maize, match = "Sample_ID", color = "Compartment")
p1b <- com_plot(maize, match = "Sample_ID", d1 = 3, d2 = 4,
  color = "Compartment")
p2a <- com_plot(maize, match = "Sample_ID", color = "Host_genotype")
```

```
p2b <- com_plot(maize, match = "Sample_ID", d1 = 1, d2 = 3, color =
"Host_genotype")
p3a <- com_plot(maize, match = "Sample_ID", color = "Compartment", shape =
"Soil")
p3b <- com_plot(maize, match = "Sample_ID", d1 = 1, d2 = 4, color =
"Compartment", shape = "Soil")
```

---

|        |                                                                                                                                                                                                         |
|--------|---------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| com_r2 | <i>Calculate the unexplained variance ratio using formula indicated in:<br/>Anderson, M.J. 2001. A new method for non-parametric multivariate<br/>analysis of variance. Austral Ecology, 26: 32–46.</i> |
|--------|---------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|

---

## Description

Calculate the unexplained variance ratio using formula indicated in: Anderson, M.J. 2001. A new method for non-parametric multivariate analysis of variance. Austral Ecology, 26: 32–46.

## Usage

```
com_r2(x, group)
```

## Arguments

|       |                                                                  |
|-------|------------------------------------------------------------------|
| x     | An object of class ‘mina’ with ‘dis’ and ‘des’ defined.          |
| group | The name(s) of column(s) defined as experimental setup group(s). |

## Value

Unexplained variance ratio.

## Examples

```
data(maize)
maize <- norm_tab(maize, method = "raref", depth = 5000)
maize <- fit_tabs(maize)
maize <- com_dis(maize, method = "bray")
com_r2(maize, group = c("Compartment", "Soil", "Host_genotype"))
```

---

```
com_r2,mina,ANY-method
```

*Function for unexplained variance ratio calculation indicated in Anderson, M.J. 2001. A new method for non-parametric multivariate analysis of variance. Austral Ecology, 26: 32–46.*

---

### Description

Function for unexplained variance ratio calculation indicated in Anderson, M.J. 2001. A new method for non-parametric multivariate analysis of variance. Austral Ecology, 26: 32–46.

### Usage

```
## S4 method for signature 'mina,ANY'
com_r2(x, group)

## S4 method for signature 'mina,character'
com_r2(x, group)
```

### Arguments

|       |                                                                  |
|-------|------------------------------------------------------------------|
| x     | An mina object with ‘dis’ and ‘des’ defined.                     |
| group | The name(s) of column(s) defined as experimental setup group(s). |

### Value

r2 The variance ratio cannot be explained by given groups.

### Examples

```
maize <- new("mina", tab = maize_asv2, des = maize_des2)
maize <- norm_tab(maize, method = "raref")
maize <- fit_tabs(maize)
maize <- com_dis(maize, method = "bray")
com_r2(maize, group = c("Compartment", "Soil", "Host_genotype"))
```

---

```
cp_cor
```

*Function for correlation coefficient calculation.*

---

### Description

Function for correlation coefficient calculation.

### Usage

```
cp_cor(mat)
```

**Arguments**

mat                    The input matrix for correlation calculation.

**Value**

The output correlation matrix.

---

|          |                                                                                                                                  |
|----------|----------------------------------------------------------------------------------------------------------------------------------|
| data-hmp | <i>Internal testing data of HMP project, including quantitative table (hmp_otu) and descriptive table (hmp_des) for testing.</i> |
|----------|----------------------------------------------------------------------------------------------------------------------------------|

---

**Description**

Internal testing data of HMP project, including quantitative table (hmp\_otu) and descriptive table (hmp\_des) for testing.

**Examples**

data(hmp)

---

|            |                                                                                                                                                                               |
|------------|-------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| data-maize | <i>Internal testing data of maize project, vegetative stage samples only, including quantitative table (maize_asv.rds) and descriptive table (maize_des.txt) for testing.</i> |
|------------|-------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|

---

**Description**

Internal testing data of maize project, vegetative stage samples only, including quantitative table (maize\_asv.rds) and descriptive table (maize\_des.txt) for testing.

**Examples**

data(maize)

---

|       |                                                                                                       |
|-------|-------------------------------------------------------------------------------------------------------|
| des<- | <i>Setter and getter for the slot 'des', which is the description and meta data of rows in 'tab'.</i> |
|-------|-------------------------------------------------------------------------------------------------------|

---

## Description

Setter and getter for the slot 'des', which is the description and meta data of rows in 'tab'.

## Usage

```
des(x) <- value

## S4 replacement method for signature 'mina'
des(x) <- value

des(x)

## S4 method for signature 'mina'
des(x)
```

## Arguments

|       |                                                         |
|-------|---------------------------------------------------------|
| x     | The 'mina' object.                                      |
| value | The value to set for the slot of the 'mina' object 'x'. |

## Value

The 'des' slot of the 'mina' object.

## Examples

```
des(maize) <- maize_des2
head(des(maize))
```

---

|       |                                              |
|-------|----------------------------------------------|
| dis<- | <i>Setter and getter for the slot 'dis'.</i> |
|-------|----------------------------------------------|

---

## Description

Setter and getter for the slot 'dis'.

Get the slot 'dis'

**Usage**

```

dis(x) <- value

dis(x)

## S4 replacement method for signature 'mina'
dis(x) <- value

## S4 method for signature 'mina'
dis(x)

```

**Arguments**

x                      The ‘mina’ object.

value                  The value to set for the slot of the ‘mina’ object ‘x’.

**Value**

The ‘dis’ slot of the ‘mina’ object.

**Examples**

```

maize_norm <- norm_tab(maize_asv2, method = "total")
dis(maize) <- com_dis(maize_norm, method = "bray")
dis(maize)[1:5, 1:5]

```

---

|        |                                                                |
|--------|----------------------------------------------------------------|
| dis_bs | <i>Getter for the slots ‘dis_bs’, ‘dis_pm’ and ‘dis_stat’.</i> |
|--------|----------------------------------------------------------------|

---

**Description**

Getter for the slots ‘dis\_bs’, ‘dis\_pm’ and ‘dis\_stat’.

**Usage**

```

dis_bs(x)

dis_pm(x)

dis_stat(x)

```

**Arguments**

x                      The ‘mina’ object.

**Value**

The ‘dis\_bs’, ‘dis\_pm’ and ‘dis\_stat’ slots of the ‘mina’ object.

**Examples**

```

maize <- new("mina", tab = maize_asv2, des = maize_des2)
maize <- norm_tab(maize, method = "raref")
maize <- fit_tabs(maize)
maize <- bs_pm(maize, group = "Compartment", per = 0.5)
maize <- net_dis(maize, method = "Jaccard")
dis_bs(maize)
dis_pm(maize)
dis_stat(maize)

```

---

dis\_bs<-                      *Setter for the slots 'dis\_bs', 'dis\_pm' and 'dis\_stat'.*

---

**Description**

Setter for the slots 'dis\_bs', 'dis\_pm' and 'dis\_stat'.

**Usage**

```

dis_bs(x) <- value

dis_pm(x) <- value

dis_stat(x) <- value

## S4 replacement method for signature 'mina'
dis_bs(x) <- value

## S4 replacement method for signature 'mina'
dis_pm(x) <- value

## S4 replacement method for signature 'mina'
dis_stat(x) <- value

```

**Arguments**

|       |                                                         |
|-------|---------------------------------------------------------|
| x     | The 'mina' object.                                      |
| value | The value to set for the slot of the 'mina' object 'x'. |

**Value**

The 'dis\_bs', 'dis\_pm' and 'dis\_stat' slots of the 'mina' object.

---

|     |                                                                                          |
|-----|------------------------------------------------------------------------------------------|
| dmr | <i>Dimensionality reduction of community dissimilarity / distance for visualization.</i> |
|-----|------------------------------------------------------------------------------------------|

---

**Description**

Dimensionality reduction of community dissimilarity / distance for visualization.

**Usage**

```
dmr(x, k = 2)
```

**Arguments**

|   |                                                                    |
|---|--------------------------------------------------------------------|
| x | An object of class 'mina' with 'dis' defined or a distance matrix. |
| k | The dimension number after reduction.                              |

**Value**

The dimensionality reduction results.

**Examples**

```
maize <- new("mina", tab = maize_asv2, des = maize_des2)
maize <- norm_tab(maize, method = "raref")
maize <- fit_tabs(maize)
maize <- com_dis(maize, method = "bray")
maize <- dmr(maize)
```

---

|                   |                                                         |
|-------------------|---------------------------------------------------------|
| dmr,matrix-method | <i>Dimensionality reduction of the distance matrix.</i> |
|-------------------|---------------------------------------------------------|

---

**Description**

Dimensionality reduction of the distance matrix.

**Usage**

```
## S4 method for signature 'matrix'
dmr(x, k = 4)
```

**Arguments**

|   |                                                             |
|---|-------------------------------------------------------------|
| x | A distance matrix.                                          |
| k | The number of dimensionality after reduction, 4 by default. |

**Value**

y The coordinates of components indicated in distance matrix in k dimension.

**Examples**

```
maize <- new("mina", tab = maize_asv2, des = maize_des2)
maize <- norm_tab(maize, method = "raref")
maize <- fit_tabs(maize)
maize <- com_dis(maize, method = "bray")
asv_dis <- dis(maize)
asv_dis_dmr <- dmr(asv_dis, k = 4)
```

---

|                 |                                                                |
|-----------------|----------------------------------------------------------------|
| dmr,mina-method | <i>Dimensionality reduction of the ‘dis’ included in mina.</i> |
|-----------------|----------------------------------------------------------------|

---

**Description**

Dimensionality reduction of the ‘dis’ included in mina.

**Usage**

```
## S4 method for signature 'mina'
dmr(x, k = 4)
```

**Arguments**

|   |                                                             |
|---|-------------------------------------------------------------|
| x | An object of the class ‘mina’ with ‘dis’ defined.           |
| k | The number of dimensionality after reduction, 4 by default. |

**Value**

x The same object with ‘dmr’ added.

**Examples**

```
maize <- new("mina", tab = maize_asv2, des = maize_des2)
maize <- norm_tab(maize, method = "raref")
maize <- fit_tabs(maize)
maize <- com_dis(maize, method = "bray")
maize <- dmr(maize)
```

---

|            |                                                                                                                                                               |
|------------|---------------------------------------------------------------------------------------------------------------------------------------------------------------|
| filter_mat | <i>Function for filtering of matrix, rows present in less than 'p' columns will be removed. After row filtering, columns with a sum of 0 will be removed.</i> |
|------------|---------------------------------------------------------------------------------------------------------------------------------------------------------------|

---

**Description**

Function for filtering of matrix, rows present in less than 'p' columns will be removed. After row filtering, columns with a sum of 0 will be removed.

**Usage**

```
filter_mat(x, p)
```

**Arguments**

|   |                                        |
|---|----------------------------------------|
| x | The input matrix to be filtered.       |
| p | The cutoff for non-zero column number. |

**Value**

x The same matrix after filtering.

---

|          |                                                                                                                                  |
|----------|----------------------------------------------------------------------------------------------------------------------------------|
| fit_tabs | <i>Filter the quantitative and descriptive table to make them have the same samples, the intersect samples will be remained.</i> |
|----------|----------------------------------------------------------------------------------------------------------------------------------|

---

**Description**

Filter the quantitative and descriptive table to make them have the same samples, the intersect samples will be remained.

**Usage**

```
fit_tabs(x)
```

**Arguments**

|   |                                                                                                                      |
|---|----------------------------------------------------------------------------------------------------------------------|
| x | An object of the class mina with 'tab' and 'des' defined or a quantitative matrix (need parameter des in this case). |
|---|----------------------------------------------------------------------------------------------------------------------|

**Value**

Same 'mina' object but fitted 'tab' and 'des' (as well as 'norm' if defined)

**Examples**

```
data(maize)
maize <- fit_tabs(maize)
maize <- norm_tab(maize, method = "raref")
maize <- fit_tabs(maize)
```

---

|                      |                                                                                                                                                                                                                                                                    |
|----------------------|--------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| fit_tabs,mina-method | <i>Filter the quantitative and descriptive table to make them have the same samples, samples present in both tables are remained. If 'norm' table exist in the 'mina' object, descriptive table will be filtered again to only keep samples present in 'norm'.</i> |
|----------------------|--------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|

---

**Description**

Filter the quantitative and descriptive table to make them have the same samples, samples present in both tables are remained. If 'norm' table exist in the 'mina' object, descriptive table will be filtered again to only keep samples present in 'norm'.

**Usage**

```
## S4 method for signature 'mina'
fit_tabs(x)
```

**Arguments**

x                      An object of class mina.

**Value**

x The same object as input with fitted 'tab', 'des' and 'norm' (if defined).

**Examples**

```
{
data(maize)
maize <- fit_tabs(maize)
maize <- norm_tab(maize, method = "total")
maize <- fit_tabs(maize)
}
```

---

|              |                                                                    |
|--------------|--------------------------------------------------------------------|
| get_contrast | <i>Get the contrast between two networks / adjacency matrices.</i> |
|--------------|--------------------------------------------------------------------|

---

**Description**

Get the contrast between two networks / adjacency matrices.

**Usage**

```
get_contrast(x, y)
```

**Arguments**

|   |                                                                |
|---|----------------------------------------------------------------|
| x | The network/adjacency matrix of one of the compared condition. |
| y | The other network/adjacency matrix.                            |

**Value**

The contrast between two matrices.

---

|                  |                                                                                     |
|------------------|-------------------------------------------------------------------------------------|
| get_contrast_grp | <i>Get the contrast of groups of nodes between two networks/adjacency matrices.</i> |
|------------------|-------------------------------------------------------------------------------------|

---

**Description**

Get the contrast of groups of nodes between two networks/adjacency matrices.

**Usage**

```
get_contrast_grp(x, y, grp)
```

**Arguments**

|     |                                                                |
|-----|----------------------------------------------------------------|
| x   | The network/adjacency matrix of one of the compared condition. |
| y   | The other network/adjacency matrix.                            |
| grp | The group information.                                         |

**Value**

The contrast between two matrices.

---

|            |                                                              |
|------------|--------------------------------------------------------------|
| get_dis_df | <i>Function for getting distance data frame from 'dist'.</i> |
|------------|--------------------------------------------------------------|

---

**Description**

Function for getting distance data frame from 'dist'.

**Usage**

```
get_dis_df(x)
```

**Arguments**

x                      The object of class 'dist'.

**Value**

The data frame of distance matrix.

---

|        |                                                                            |
|--------|----------------------------------------------------------------------------|
| get_ja | <i>Get the Jaccard distance between two networks / adjacency matrices.</i> |
|--------|----------------------------------------------------------------------------|

---

**Description**

Get the Jaccard distance between two networks / adjacency matrices.

**Usage**

```
get_ja(x, y)
```

**Arguments**

x                      The network/adjacency matrix of one of the compared condition.  
y                      The other network/adjacency matrix.

**Value**

The Jaccard distance between two matrices.

---

|         |                                                                             |
|---------|-----------------------------------------------------------------------------|
| get_ja0 | <i>Get the Jaccard0 distance between two networks / adjacency matrices.</i> |
|---------|-----------------------------------------------------------------------------|

---

**Description**

Get the Jaccard0 distance between two networks / adjacency matrices.

**Usage**

```
get_ja0(x, y)
```

**Arguments**

|   |                                                                |
|---|----------------------------------------------------------------|
| x | The network/adjacency matrix of one of the compared condition. |
| y | The other network/adjacency matrix.                            |

**Value**

The Jaccard0 distance between two matrices.

---

|             |                                                                             |
|-------------|-----------------------------------------------------------------------------|
| get_ja0_grp | <i>Get the Jaccard0 distance between two networks / adjacency matrices.</i> |
|-------------|-----------------------------------------------------------------------------|

---

**Description**

Get the Jaccard0 distance between two networks / adjacency matrices.

**Usage**

```
get_ja0_grp(x, y, grp)
```

**Arguments**

|     |                                                                |
|-----|----------------------------------------------------------------|
| x   | The network/adjacency matrix of one of the compared condition. |
| y   | The other network/adjacency matrix.                            |
| grp | The table with group information.                              |

**Value**

The Jaccard0 distance between two matrices.

---

|            |                                                                            |
|------------|----------------------------------------------------------------------------|
| get_ja_grp | <i>Get the Jaccard distance between two networks / adjacency matrices.</i> |
|------------|----------------------------------------------------------------------------|

---

**Description**

Get the Jaccard distance between two networks / adjacency matrices.

**Usage**

```
get_ja_grp(x, y, grp)
```

**Arguments**

- x                   The network/adjacency matrix of one of the compared condition.
- y                   The other network/adjacency matrix.
- grp                 The table with group information.

**Value**

The Jaccard distance between two matrices.

---

|                 |                                                                                                             |
|-----------------|-------------------------------------------------------------------------------------------------------------|
| get_net_cls_tab | <i>Get the cluster table 'cls_tab' from quantitative table 'norm' and network clustering results 'cls'.</i> |
|-----------------|-------------------------------------------------------------------------------------------------------------|

---

**Description**

Get the cluster table 'cls\_tab' from quantitative table 'norm' and network clustering results 'cls'.

**Usage**

```
get_net_cls_tab(x_norm, x_cls, uw = FALSE)
```

**Arguments**

- x\_norm             The normalized quantitative table used for netowrk inference and clustering.
- x\_cls              The network clustering table.
- uw                 By summing up the number of present components of each cluster instead of relative abundance, default is FALSE.

**Value**

x\_cls The quantitative table with clusters in rows.

**Examples**

```

maize <- new("mina", tab = maize_asv2, des = maize_des2)
maize <- norm_tab(maize, method = "raref", depth = 1000)
maize <- fit_tabs(maize)
maize_norm <- norm(maize)
maize_adj <- adj(maize_norm, method = "spearman")
maize_cls <- net_cls(maize_adj, method = "ap", cutoff = 0.5)
maize_cls_tab <- get_net_cls_tab(maize_norm, maize_cls)

```

---

```
get_net_cls_tab,matrix,data.frame-method
```

*Get the cluster table 'cls\_tab' from quantitative table 'norm' and network clustering results 'cls'.*

---

**Description**

Get the cluster table 'cls\_tab' from quantitative table 'norm' and network clustering results 'cls'.

**Usage**

```

## S4 method for signature 'matrix,data.frame'
get_net_cls_tab(x_norm, x_cls, uw = FALSE)

```

**Arguments**

|        |                                                                                                                 |
|--------|-----------------------------------------------------------------------------------------------------------------|
| x_norm | The normalized quantitative table used for network inference and clustering.                                    |
| x_cls  | The network clustering table.                                                                                   |
| uw     | By summing up the number of present components of each cluster instead of relative abundance, default is FALSE. |

**Value**

x\_cls The quantitative table with clusters in rows.

**Examples**

```

maize <- new("mina", tab = maize_asv2, des = maize_des2)
maize <- norm_tab(maize, method = "raref", depth = 1000)
maize <- fit_tabs(maize)
maize_norm <- norm(maize)
maize_adj <- adj(maize_norm, method = "spearman")
maize_cls <- net_cls(maize_adj, method = "ap", cutoff = 0.5)
maize_cls_tab <- get_net_cls_tab(maize_norm, maize_cls)

```

---

|        |                                                                                            |
|--------|--------------------------------------------------------------------------------------------|
| get_r2 | <i>Same function as ‘com_r2’ with matrix and corresponding descriptive table as input.</i> |
|--------|--------------------------------------------------------------------------------------------|

---

### Description

Same function as ‘com\_r2’ with matrix and corresponding descriptive table as input.

### Usage

```
get_r2(x, des, group)
```

### Arguments

|       |                                                                                   |
|-------|-----------------------------------------------------------------------------------|
| x     | Dissimilarity / distance matrix which indicate variances.                         |
| des   | The descriptive table of samples which define the groups.                         |
| group | The name(s) of column(s) used as experimental setup group(s) in descriptive file. |

### Value

r2 The variance ratio cannot be explained by given groups.

### Examples

```
maize <- new("mina", tab = maize_asv2, des = maize_des2)
maize <- norm_tab(maize, method = "raref")
maize <- fit_tabs(maize)
maize <- com_dis(maize, method = "bray")
get_r2(dis(maize), des(maize), group = c("Compartment", "Soil"))
```

---

get\_r2,matrix,ANY,ANY-method

*Function for unexplained variance ratio calculation indicated in Anderson, M.J. 2001. A new method for non-parametric multivariate analysis of variance. Austral Ecology, 26: 32–46.*

---

### Description

Function for unexplained variance ratio calculation indicated in Anderson, M.J. 2001. A new method for non-parametric multivariate analysis of variance. Austral Ecology, 26: 32–46.

**Usage**

```
## S4 method for signature 'matrix,ANY,ANY'
get_r2(x, des, group)

## S4 method for signature 'matrix,data.frame,ANY'
get_r2(x, des, group)

## S4 method for signature 'matrix,data.frame,character'
get_r2(x, des, group = c("Host_genotype", "Compartment", "Soil", "Management"))
```

**Arguments**

|       |                                                                                   |
|-------|-----------------------------------------------------------------------------------|
| x     | Dissimilarity / distance matrix which indicate variances.                         |
| des   | The descriptive table of samples which define the groups.                         |
| group | The name(s) of column(s) used as experimental setup group(s) in descriptive file. |

**Value**

r2 The variance ratio cannot be explained by given groups.

**Examples**

```
maize <- new("mina", tab = maize_asv2, des = maize_des2)
maize <- norm_tab(maize, method = "raref")
maize <- fit_tabs(maize)
maize <- com_dis(maize, method = "bray")
x <- dis(maize)
des <- des(maize)
get_r2(x, des, group = c("Compartment", "Soil"))
```

get\_rep

---

*Get the representative community members by extracting the most abundant and prevalent compositions.*

---

**Description**

Get the representative community members by extracting the most abundant and prevalent compositions.

Get the representative community members.

**Usage**

```
get_rep(x, ...)

## S4 method for signature 'matrix'
get_rep(x, top = 5)
```

**Arguments**

|     |                                                                         |
|-----|-------------------------------------------------------------------------|
| x   | A quantitative matrix with samples in columns and compositions in rows. |
| ... | Additional parameters.                                                  |
| top | The percent of the most abundant and prevalent members.                 |

**Value**

The matrix with samples in columns and representative compositions in rows.

The matrix with samples in columns and representative compositions in rows.

**Examples**

```
data(maize_asv)
maize_asv_rep <- get_rep(maize_asv)
data(maize_asv)
maize_asv_rep <- get_rep(maize_asv, top = 5)
```

---

|                     |                                                  |
|---------------------|--------------------------------------------------|
| get_rep,mina-method | <i>Get the representative community members.</i> |
|---------------------|--------------------------------------------------|

---

**Description**

Get the representative community members.

**Usage**

```
## S4 method for signature 'mina'
get_rep(x, top = 5)
```

**Arguments**

|     |                                                         |
|-----|---------------------------------------------------------|
| x   | An object of the class ‘mina’ with @norm define.        |
| top | The percent of the most abundant and prevalent members. |

**Value**

The same object with @norm replaced by the representative members.

**Examples**

```
maize <- new("mina", tab = maize_asv, des = maize_des)
maize <- norm_tab(maize, method = "raref")
maize <- get_rep(maize, top = 5)
```

---

`get_spectra`*Function for calculation of eigenvalue of given matrix.*

---

**Description**

Function for calculation of eigenvalue of given matrix.

**Usage**

```
get_spectra(x, k = 100)
```

**Arguments**

|   |                              |
|---|------------------------------|
| x | The input matrix.            |
| k | Get the first k eigenvalues. |

**Value**

y The vector of the first k eigenvalues.

---

`get_stat`*Function for distance statistic and significance test.*

---

**Description**

Function for distance statistic and significance test.

**Usage**

```
get_stat(x, p = NULL)
```

**Arguments**

|   |                                      |
|---|--------------------------------------|
| x | The bootstrap distance data frame.   |
| p | The permutation distance data frame. |

**Value**

The statistics of network comparison.

---

|         |                                                                      |
|---------|----------------------------------------------------------------------|
| hmp_des | <i>Design file for HMP project, including 2711 samples in total.</i> |
|---------|----------------------------------------------------------------------|

---

**Description**

Design file for HMP project, including 2711 samples in total.

**Format**

A data frame with columns:

**Sample\_ID** The unique ID of the microbial profiling sample.

**Sex** The gender of the host human.

**Run\_center** The lab processing the sample sequencing.

**Subsite** The subsite of body where samples were collected.

**Site** The site of body where samples were collected.

**Description** The further details about the samples.

**Source**

HMP project.

**Examples**

```
data(hmp_des)
```

---

|         |                                                                                                                                           |
|---------|-------------------------------------------------------------------------------------------------------------------------------------------|
| hmp_otu | <i>OTU table of HMP project, data downloaded from <a href="https://www.hmpdacc.org/hmp/HMQCP/">https://www.hmpdacc.org/hmp/HMQCP/</a></i> |
|---------|-------------------------------------------------------------------------------------------------------------------------------------------|

---

**Description**

OTU table of HMP project, data downloaded from <https://www.hmpdacc.org/hmp/HMQCP/>

**Format**

A matrix with samples in columns and OTUs in rows.

**Source**

HMP project.

**Examples**

```
data(hmp_otu)
```

---

|           |                                                                   |
|-----------|-------------------------------------------------------------------|
| maize_asv | <i>ASV table of maize project, vegetative stage samples only.</i> |
|-----------|-------------------------------------------------------------------|

---

**Description**

ASV table of maize project, vegetative stage samples only.

**Format**

A matrix with samples in columns and ASVs in rows. Unnormalized table including 12765 ASVs from 420 samples.

**Source**

RECONSTRUCT project, maize microbiome part.

**Examples**

```
data(maize_asv)
```

---

|            |                                                                                                                     |
|------------|---------------------------------------------------------------------------------------------------------------------|
| maize_asv2 | <i>Subset of ASV table of maize project, ASVs appear in less than 100 samples were filtered for later analysis.</i> |
|------------|---------------------------------------------------------------------------------------------------------------------|

---

**Description**

Subset of ASV table of maize project, ASVs appear in less than 100 samples were filtered for later analysis.

**Format**

A matrix with samples in columns and ASVs in rows. Unnormalized table including 1219 ASVs from 313 samples.

**Source**

RECONSTRUCT project, maize microbiome part.

**Examples**

```
data(maize_asv2)
```

---

|           |                                                                                                     |
|-----------|-----------------------------------------------------------------------------------------------------|
| maize_des | <i>Design file of maize project, vegetative stage samples only, including 528 samples in total.</i> |
|-----------|-----------------------------------------------------------------------------------------------------|

---

### Description

Design file of maize project, vegetative stage samples only, including 528 samples in total.

### Format

A data frame with columns:

**Sample\_ID** The unique ID of the microbial profiling sample.

**Host\_genotype** The genotype of the plant host maize.

**Compartment** The compartment of the microbial sample comes from.

**Soil** The soil of the sampled microbiome.

**Management** The management of the soil where microbial sample from.

### Source

RECONSTRUCT project, maize microbiome part.

### Examples

```
data(maize_des)
```

---

|            |                                                                          |
|------------|--------------------------------------------------------------------------|
| maize_des2 | <i>Subset of design file of maize project, 313 samples are included.</i> |
|------------|--------------------------------------------------------------------------|

---

### Description

Subset of design file of maize project, 313 samples are included.

### Format

A data frame with columns:

**Sample\_ID** The unique ID of the microbial profiling sample.

**Host\_genotype** The genotype of the plant host maize.

**Compartment** The compartment of the microbial sample comes from.

**Soil** The soil of the sampled microbiome.

**Management** The management of the soil where microbial sample from.

**Source**

RECONSTRUCT project, maize microbiome part.

**Examples**

```
data(maize_des2)
```

---

|                   |                                                                                                                                                                                                                                                                                                                                                                                                                |
|-------------------|----------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| mat_or_NULL-class | <i>Modified from <a href="https://github.com/joey711/phyloseq/blob/master/R/allClasses.R">https://github.com/joey711/phyloseq/blob/master/R/allClasses.R</a> Use setClassUnion to define the unholy NULL-data union as a virtual class. This is a way of dealing with the expected scenarios in which one or more of the component data classes is not available, in which case NULL will be used instead.</i> |
|-------------------|----------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|

---

**Description**

Modified from <https://github.com/joey711/phyloseq/blob/master/R/allClasses.R> Use setClassUnion to define the unholy NULL-data union as a virtual class. This is a way of dealing with the expected scenarios in which one or more of the component data classes is not available, in which case NULL will be used instead.

---

|            |                                                                            |
|------------|----------------------------------------------------------------------------|
| mina-class | <i>Class "mina" includes the quantitative table and descriptive table.</i> |
|------------|----------------------------------------------------------------------------|

---

**Description**

Class "mina" includes the quantitative table and descriptive table.

**Slots**

**tab** The quantitative table of the dataset.

**des** The descriptive table of the samples listed in @tab.

**norm** The normalized quantitative table of @tab.

**dis** The distance / dissimilarity matrix between samples in @tab.

**dmr** The list of dimensionality reduction result, includes points and variance.

**adj** The adjacency matrix between pairwise compositions (e.g. OTUs/ASVs)

**adj\_sig** The P-value matrix of adjacency matrix, only applicable for Pearson and Spearman correlation adjacency matrices.

**cls** The cluster information for each composition.

**cls\_tab** The cluster quantitative table.

**multi** The list of subsampled adjacency matrices for each environment.

perm The list of permuted adjacency matrices for each pairwise environmental comparison.  
 dis\_bs The distance between networks of different environmental communities.  
 dis\_pm The distance between networks of permuted groups.  
 dis\_stat The average distance between subsampled environmental community networks, permuted networks and corresponding significance.

### Author(s)

Rui Guan <https://github.com/Guan06>

### Examples

```
maize <- new("mina", tab = maize_asv, des = maize_des)
```

---

|         |                                                           |
|---------|-----------------------------------------------------------|
| multi<- | <i>Setter and getter for the slot 'multi' and 'perm'.</i> |
|---------|-----------------------------------------------------------|

---

### Description

Setter and getter for the slot 'multi' and 'perm'.

### Usage

```
multi(x) <- value

multi(x)

## S4 replacement method for signature 'mina'
multi(x) <- value

## S4 method for signature 'mina'
multi(x)

perm(x) <- value

perm(x)

## S4 replacement method for signature 'mina'
perm(x) <- value

## S4 method for signature 'mina'
perm(x)
```

### Arguments

|       |                                                         |
|-------|---------------------------------------------------------|
| x     | The 'mina' object.                                      |
| value | The value to set for the slot of the 'mina' object 'x'. |

**Value**

The ‘multi’ or/and ‘perm’ slot of the ‘mina’ object.

---

|         |                                                          |
|---------|----------------------------------------------------------|
| net_cls | <i>Network clustering of sparsed adjacacency matrix.</i> |
|---------|----------------------------------------------------------|

---

**Description**

Network clustering of sparsed adjacacency matrix.

**Usage**

```
net_cls(x, method, ...)
```

**Arguments**

|        |                                               |
|--------|-----------------------------------------------|
| x      | An object of class ‘mina’ with ‘adj’ defined. |
| method | The clustering method used.                   |
| ...    | Additional parameters.                        |

**Value**

The network clustering results.

**Examples**

```
maize <- new("mina", tab = maize_asv2, des = maize_des2)
maize <- norm_tab(maize, method = "raref", depth = 1000)
maize <- fit_tabs(maize)
maize <- adj(maize, method = "spearman")
maize <- net_cls(maize, method = "mcl", cutoff = 0.4, neg = FALSE)
```

---

|                           |                                                                    |
|---------------------------|--------------------------------------------------------------------|
| net_cls,matrix,ANY-method | <i>Network clustering based on the sparsed adjacacency matrix.</i> |
|---------------------------|--------------------------------------------------------------------|

---

**Description**

Network clustering based on the sparsed adjacacency matrix.

**Usage**

```
## S4 method for signature 'matrix,ANY'
net_cls(x, method, cutoff = 0.4, neg = FALSE, ...)

## S4 method for signature 'matrix,character'
net_cls(x, method, cutoff = 0.4, neg = FALSE, ...)
```

**Arguments**

|        |                                                                                                    |
|--------|----------------------------------------------------------------------------------------------------|
| x      | Adjacency matrix used for clustering.                                                              |
| method | The clustering method used.                                                                        |
| cutoff | The cutoff for the sparsed adjacency matrix, default 0.4.                                          |
| neg    | Whether to keep the negative edges, cannot be TRUE when using 'mcl' for clustering. Default FALSE. |
| ...    | Additional parameters.                                                                             |

**Value**

y The cluster table.

**Examples**

```
asv_norm <- norm_tab(maize_asv2, method = "raref", depth = 1000)
asv_adj <- adj(asv_norm, method = "spearman")
asv_cls <- net_cls(asv_adj, method = "mcl")
```

---

net\_cls,mina,ANY-method

*Network clustering based on the sparsed adjacency matrix.*

---

**Description**

Network clustering based on the sparsed adjacency matrix.

**Usage**

```
## S4 method for signature 'mina,ANY'
net_cls(x, method, cutoff = 0.4, neg = FALSE, ...)

## S4 method for signature 'mina,character'
net_cls(x, method, cutoff = 0.4, neg = FALSE, ...)
```

**Arguments**

|        |                                                                                                    |
|--------|----------------------------------------------------------------------------------------------------|
| x      | An object of class 'mina' with 'adj' defined.                                                      |
| method | The clustering method used.                                                                        |
| cutoff | The cutoff for the sparsed adjacency matrix, default 0.4.                                          |
| neg    | Whether to keep the negative edges, cannot be TRUE when using 'mcl' for clustering. Default FALSE. |
| ...    | Additional parameters.                                                                             |

**Value**

x The same 'mina' class with @cls added.

**Examples**

```
maize <- new("mina", tab = maize_asv2, des = maize_des2)
maize <- norm_tab(maize, method = "raref", depth = 1000)
maize <- fit_tabs(maize)
maize <- adj(maize, method = "spearman")
maize <- net_cls(maize, method = "mcl", cutoff = 0.4, neg = FALSE)
maize <- net_cls(maize, method = "ap", cutoff = 0.4, neg = FALSE)
```

---

|             |                                                               |
|-------------|---------------------------------------------------------------|
| net_cls_tab | <i>Get the cluster table 'cls_tab' from 'norm' and 'cls'.</i> |
|-------------|---------------------------------------------------------------|

---

**Description**

Get the cluster table 'cls\_tab' from 'norm' and 'cls'.

**Usage**

```
net_cls_tab(x, uw = FALSE)
```

**Arguments**

|    |                                                                                                                  |
|----|------------------------------------------------------------------------------------------------------------------|
| x  | An object of class 'mina' with 'norm' and 'cls' defined.                                                         |
| uw | By summing up the number of present components of each cluster instead of relative abundances, default is FALSE. |

**Value**

The network cluster relative abundance table.

**Examples**

```
maize <- new("mina", tab = maize_asv2, des = maize_des2)
maize <- norm_tab(maize, method = "raref", depth = 1000)
maize <- fit_tabs(maize)
maize <- adj(maize, method = "spearman")
maize <- net_cls(maize, method = "ap", cutoff = 0.5)
maize <- net_cls_tab(maize)
```

---

```
net_cls_tab,mina-method
```

*Get the cluster table 'cls\_tab' from quantitative table 'norm' and network clustering results 'cls'.*

---

### Description

Get the cluster table 'cls\_tab' from quantitative table 'norm' and network clustering results 'cls'.

### Usage

```
## S4 method for signature 'mina'
net_cls_tab(x, uw = FALSE)
```

### Arguments

|    |                                                                                                                 |
|----|-----------------------------------------------------------------------------------------------------------------|
| x  | An object of class 'mina' with 'norm' and 'cls' defined.                                                        |
| uw | By summing up the number of present components of each cluster instead of relative abundance, default is FALSE. |

### Value

x The same 'mina' object with 'cls\_tab' added.

### Examples

```
maize <- new("mina", tab = maize_asv2, des = maize_des2)
maize <- norm_tab(maize, method = "raref", depth = 1000)
maize <- fit_tabs(maize)
maize <- adj(maize, method = "spearman")
maize <- net_cls(maize, method = "mcl", cutoff = 0.5)
maize <- net_cls_tab(maize)
```

---

```
net_dis
```

*Calculate the network distance of 'multi' and test the significance when 'perm' is defined.*

---

### Description

Calculate the network distance of 'multi' and test the significance when 'perm' is defined.

### Usage

```
net_dis(x, method, ...)
```

**Arguments**

`x`                    An object of class 'mina' with 'multi' (and 'perm' if 'sig' is TRUE) defined.  
`method`                The distance to be calculated, "spectra" and "Jaccard" are available.  
`...`                   Additional parameters.

**Value**

The network comparison result.

**Examples**

```
maize <- new("mina", tab = maize_asv2, des = maize_des2)
maize <- norm_tab(maize, method = "raref")
maize <- fit_tabs(maize)
maize <- get_rep(maize, top = 5)
maize <- bs_pm(maize, group = "Compartment")
maize <- net_dis(maize, method = "spectra", evk = 30)
```

---

net\_dis,mina,ANY-method

*Calculate the network distance of 'multi' and test the significance when 'perm' is defined.*

---

**Description**

Calculate the network distance of 'multi' and test the significance when 'perm' is defined.

**Usage**

```
## S4 method for signature 'mina,ANY'
net_dis(
  x,
  method,
  evk = 100,
  egv = TRUE,
  dir = "./",
  sig = TRUE,
  skip = TRUE,
  ...
)

## S4 method for signature 'mina,character'
net_dis(
  x,
  method,
  evk = 100,
  egv = TRUE,
```

```

    dir = "./",
    sig = TRUE,
    skip = TRUE,
    ...
  )

```

### Arguments

|        |                                                                                                                          |
|--------|--------------------------------------------------------------------------------------------------------------------------|
| x      | An object of class ‘mina’ with ‘multi’ (and ‘perm’ if sig is TRUE) defined.                                              |
| method | The distance to be calculated, "spectra" and "Jaccard" are available.                                                    |
| evk    | The first ‘evk’ eigenvalues will be used for ‘spectra’ distance, the default is 100.                                     |
| egv    | Whether to output the eigenvectors for Spectral distance, the default is TRUE, only validate when ‘method == "spectra"’. |
| dir    | The folder to output the eigenvectors, only validate when ‘egv == TRUE’.                                                 |
| sig    | Whether to test the significance, if TRUE (by default), ‘perm’ is needed.                                                |
| skip   | Whether to skip the comparison when the dimension of adjacency matrix is smaller than setted ‘evk’.                      |
| ...    | Additional parameters.                                                                                                   |

### Value

x The same ‘mina’ object with ‘net\_dis’ defined.

### Examples

```

maize <- new("mina", tab = maize_asv2, des = maize_des2)
maize <- norm_tab(maize, method = "raref")
maize <- fit_tabs(maize)
maize <- bs_pm(maize, group = "Compartment")
maize <- net_dis(maize, method = "Jaccard")

```

---

|              |                                                                                     |
|--------------|-------------------------------------------------------------------------------------|
| net_dis_indi | <i>Calculate the network distance of bootstrap and permutation when applicable.</i> |
|--------------|-------------------------------------------------------------------------------------|

---

### Description

Calculate the network distance of bootstrap and permutation when applicable.

Calculate the network distance of bootstrap and permutation when applicable.

**Usage**

```
net_dis_indi(x, method, ...)

## S4 method for signature 'character,ANY'
net_dis_indi(
  x,
  method,
  evk = 100,
  sig = TRUE,
  skip = TRUE,
  egv = TRUE,
  dir = "./",
  ...
)

## S4 method for signature 'character,character'
net_dis_indi(
  x,
  method,
  evk = 100,
  sig = TRUE,
  skip = TRUE,
  egv = TRUE,
  dir = "./",
  ...
)
```

**Arguments**

|        |                                                                                                                          |
|--------|--------------------------------------------------------------------------------------------------------------------------|
| x      | The folder store the network inference results. defined.                                                                 |
| method | The distance to be calculated, "spectra" and "Jaccard" are available.                                                    |
| ...    | Additional parameters.                                                                                                   |
| evk    | The first 'evk' eigenvalues will be used for 'spectra' distance, the default is 100.                                     |
| sig    | Whether to test the significance, if TRUE (by default), permutation results should be included in the folder 'x'.        |
| skip   | Whether to skip the comparison when the dimension of adjacency matrix is smaller than setted 'evk', default TRUE.        |
| egv    | Whether to output the eigenvectors for Spectral distance, the default is TRUE, only validate when 'method == "spectra"'. |
| dir    | The folder to output the eigenvectors, only validate when 'egv == TRUE'.                                                 |

**Value**

y The 'mina' object with 'dis\_bs', 'dis\_pm' and 'dis\_stat'.

y The 'mina' object with 'dis\_bs', 'dis\_pm' and 'dis\_stat'.

## Examples

```
## Not run:
data(maize)
maize <- norm_tab(maize, method = "raref")
maize <- fit_tabs(maize)
maize <- get_rep(maize, top = 5)
maize <- bs_pm(maize, group = "Compartment", individual = TRUE, out_dir =
"./individual_bs_pm/")
maize_stat1 <- net_dis_indi(x = "./individual_bs_pm/", method = "spectra")
maize_stat2 <- net_dis_indi(x = "./individual_bs_pm/", method = "Jaccard")
maize_stat3 <- net_dis_indi(x = "./individual_bs_pm/", method = "spectra",
evk = 100, skip = TRUE)

## End(Not run)
## Not run:
data(maize)
norm(maize) <- maize_asv2
maize <- fit_tabs(maize)
maize <- get_rep(maize, top= 5)
maize <- bs_pm(maize, group = "Compartment", individual = TRUE, out_dir =
"./individual_bs_pm/")
maize_stat1 <- net_dis_indi("./individual_bs_pm/", method = "spectra")
maize_stat2 <- net_dis_indi("./individual_bs_pm/", method = "Jaccard")
maize_stat3 <- net_dis_indi("./individual_bs_pm/", method = "spectra",
evk = 100, skip = TRUE)

## End(Not run)
```

---

net\_dis\_pcoa

*Visulization of spectra network distance as PCoA.*


---

## Description

Visulization of spectra network distance as PCoA.

Visulization of spectra network distance as PCoA.

## Usage

```
net_dis_pcoa(x)

## S4 method for signature 'character'
net_dis_pcoa(x)
```

## Arguments

x                      The folder with all egv files generated by net\_dis\_indi().

**Value**

p The plotted figure.

p The plotted figure.

**Examples**

```
## Not run:
data(maize)
norm(maize) <- maize_asv2
maize <- fit_tabs(maize)
maize <- bs_pm(maize, group = "Compartment", individual = TRUE, out_dir =
"./individual_bs_pm/")
maize <- net_dis_indi("./individual_bs_pm/", method = "spectra", egv = TRUE,
dir = "./egv_folder/")
p <- net_dis_pcoa("./egv_folder/")

## End(Not run)
## Not run:
data(maize)
maize <- norm_tab(maize)
maize <- fit_tabs(maize)
maize <- get_rep(maize, top = 5)
maize <- bs_pm(maize, group = "Compartment", individual = TRUE, out_dir =
"./individual_bs_pm/")
maize <- net_dis_indi("./individual_bs_pm/", method = "spectra", egv = TRUE,
dir = "./egv_folder/")
p <- net_dis_pcoa("./egv_folder/")

## End(Not run)
```

---

net\_dis\_plot

*Visulization of network distance, average distances are used for tile plot.*

---

**Description**

Visulization of network distance, average distances are used for tile plot.

Visulization of network distance, average distances are used for tile plot.

**Usage**

```
net_dis_plot(x, d = "BS", ...)

## S4 method for signature 'mina'
net_dis_plot(x, d = "BS", sig = TRUE)
```

**Arguments**

|     |                                                             |
|-----|-------------------------------------------------------------|
| x   | An object of 'mina' with slot 'dis_stat' defined.           |
| d   | The distance to be plotted, could be "BS" or "PM".          |
| ... | Additional parameters.                                      |
| sig | If 'TRUE', indicating significant distance with gold guild. |

**Value**

p The plotted figure.  
p The plotted figure.

**Examples**

```
maize <- new("mina", tab = maize_asv2, des = maize_des2)
maize <- norm_tab(maize, method = "raref")
maize <- fit_tabs(maize)
maize <- bs_pm(maize, group = "Compartment")
maize <- net_dis(maize, method = "Jaccard")
p <- net_dis_plot(maize)
maize <- new("mina", tab = maize_asv2, des = maize_des2)
maize <- norm_tab(maize, method = "raref")
maize <- fit_tabs(maize)
maize <- bs_pm(maize, group = "Compartment")
maize <- net_dis(maize, method = "Jaccard")
p <- net_dis_plot(maize, d = "BS")
```

---

net\_grp\_cmp

---

*Compare the group features between networks.*


---

**Description**

Compare the group features between networks.  
Compare the group features between networks.

**Usage**

```
net_grp_cmp(x, cmp = "contrast", dir = "./", grp)

## S4 method for signature 'character'
net_grp_cmp(x, cmp = "contrast", dir = "./", grp)
```

**Arguments**

|     |                                                                    |
|-----|--------------------------------------------------------------------|
| x   | The folder with all network inference results generated by bs_pm() |
| cmp | The compared feature of grp, default 'contrast'.                   |
| dir | The directory to store the alculated node features.                |
| grp | The table with group information.                                  |

**Examples**

```
## Not run:
net_node_cmp("./individual_bs_pm/", f = "contrast", dir = "./", grp =
cls_tab(maize))

## End(Not run)
## Not run:
net_node_cmp("./individual_bs_pm/", f = "contrast", dir = "./", grp =
cls_tab(maize))

## End(Not run)
```

---

|              |                                                    |
|--------------|----------------------------------------------------|
| net_node_cmp | <i>Compare the node features between networks.</i> |
|--------------|----------------------------------------------------|

---

**Description**

Compare the node features between networks.

Compare the node features between networks.

**Usage**

```
net_node_cmp(x, cmp = "contrast", dir = "./")

## S4 method for signature 'character,character'
net_node_cmp(x, cmp = "contrast", dir = "./")
```

**Arguments**

|     |                                                                    |
|-----|--------------------------------------------------------------------|
| x   | The folder with all network inference results generated by bs_pm() |
| cmp | The compared feature of node, default 'contrast'.                  |
| dir | The directory to store the alculated node features.                |

**Examples**

```
## Not run:
net_node_cmp("./individual_bs_pm/", f = "contrast", dir = "./")

## End(Not run)
## Not run:
net_node_cmp("./individual_bs_pm/", f = "contrast", dir = "./")

## End(Not run)
```

---

|        |                                                                         |
|--------|-------------------------------------------------------------------------|
| norm<- | <i>Setter and getters for the slot 'norm', normalized 'tab' matrix.</i> |
|--------|-------------------------------------------------------------------------|

---

### Description

Setter and getters for the slot 'norm', normalized 'tab' matrix.

### Usage

```
norm(x) <- value

## S4 replacement method for signature 'mina'
norm(x) <- value

norm(x)

## S4 method for signature 'mina'
norm(x)
```

### Arguments

|       |                                                         |
|-------|---------------------------------------------------------|
| x     | The 'mina' object.                                      |
| value | The value to set for the slot of the 'mina' object 'x'. |

### Value

The 'norm' slot of the 'mina' object.

### Examples

```
norm(maize) <- norm_tab(maize_asv2, method = "total")
norm(maize)[1:5, 1:5]
```

---

|               |                                                                                                                                                                                                     |
|---------------|-----------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| norm_by_raref | <i>Function for normalization by rarefying the samples into the same depth, modified from <b>phyloseq</b>, find it <a href="https://rdrr.io/bioc/phyloseq/man/rarefy_even_depth.html">here</a>.</i> |
|---------------|-----------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|

---

### Description

Function for normalization by rarefying the samples into the same depth, modified from **phyloseq**, find it [here](https://rdrr.io/bioc/phyloseq/man/rarefy_even_depth.html).

### Usage

```
norm_by_raref(x, depth = 1000, replace = TRUE)
```

**Arguments**

|         |                                                                                                                      |
|---------|----------------------------------------------------------------------------------------------------------------------|
| x       | A quantitative table with sample in columns and compositions in rows.                                                |
| depth   | The depth for rarefying, 1000 by default.                                                                            |
| replace | Whether to sample with replacement (TRUE) or without replacement (FALSE). Default TRUE for computational efficiency. |

**Value**

A normalized quantitative table.

---

|               |                                                                                 |
|---------------|---------------------------------------------------------------------------------|
| norm_by_total | <i>Function for normalization, by total number of the reads in each sample.</i> |
|---------------|---------------------------------------------------------------------------------|

---

**Description**

Function for normalization, by total number of the reads in each sample.

**Usage**

```
norm_by_total(x)
```

**Arguments**

|   |                                                                        |
|---|------------------------------------------------------------------------|
| x | A quantitative table with samples in columns and compositions in rows. |
|---|------------------------------------------------------------------------|

**Value**

A normalized quantitative table.

---

|          |                                                     |
|----------|-----------------------------------------------------|
| norm_tab | <i>Normalize the slot 'tab' for later analysis.</i> |
|----------|-----------------------------------------------------|

---

**Description**

Normalize the slot 'tab' for later analysis.

**Usage**

```
norm_tab(x, method, ...)
```

**Arguments**

|        |                                                                                 |
|--------|---------------------------------------------------------------------------------|
| x      | The input mina object with quantitative tab / a matrix needed to be normalized. |
| method | The method used for the normalization of quantitative table.                    |
| ...    | Additional parameters.                                                          |

**Value**

Normalized quantitative table.

**Examples**

```
maize <- new("mina", tab = maize_asv2, des = maize_des2)
maize <- norm_tab(maize, method = "total")
```

---

norm\_tab,matrix,character-method

*Normalize the quantitative matrix.*

---

**Description**

Normalize the quantitative matrix.

**Usage**

```
## S4 method for signature 'matrix,character'
norm_tab(x, method, depth = 1000, replace = TRUE, multi = 1, ...)
```

**Arguments**

|                      |                                                                                                                                          |
|----------------------|------------------------------------------------------------------------------------------------------------------------------------------|
| <code>x</code>       | A quantitative matrix with samples in columns and compositions in rows.                                                                  |
| <code>method</code>  | The method used for normalization.                                                                                                       |
| <code>depth</code>   | The depth for rarefying, 1000 by default.                                                                                                |
| <code>replace</code> | Whether to sample with replacement (TRUE by default) or without replacement (FALSE) when using method 'raref'.                           |
| <code>multi</code>   | Rarefy the table for multiple times, 1 by default, indicate the times of rarefaction want to be repeated, only validate for rarefaction. |
| <code>...</code>     | Additional parameters.                                                                                                                   |

**Value**

The normalized quantitative matrix.

`x_norm` Normalized matrix of the quantitative table.

**Examples**

```
data(maize_asv2)
maize_asv_norm <- norm_tab(maize_asv2, method = "total")
maize_asv_norm <- norm_tab(maize_asv2, method = "raref", depth = 1000,
replace = TRUE, multi = 3)
```

---

norm\_tab,mina,ANY-method

*Normalize the quantitative table with mina input.*


---

## Description

Normalize the quantitative table with mina input.

## Usage

```
## S4 method for signature 'mina,ANY'
norm_tab(x, method, depth = 1000, replace = TRUE, multi = 1, ...)

## S4 method for signature 'mina,character'
norm_tab(x, method, depth = 1000, replace = TRUE, multi = 1, ...)
```

## Arguments

|         |                                                                                                                                              |
|---------|----------------------------------------------------------------------------------------------------------------------------------------------|
| x       | An object of the class mina with @tab defined.                                                                                               |
| method  | The method used for normalization.                                                                                                           |
| depth   | The depth for subsampling by rarefying, 1000 by default.                                                                                     |
| replace | Whether to sample with replacement (TRUE by default) or without replacement (FALSE) when using method 'reref'.                               |
| multi   | Rarefy the table for multiple times, FALSE by default, indicate the times of rarefaction want to be repeated, only validate for rarefaction. |
| ...     | Additional parameters.                                                                                                                       |

## Value

x An object of the class mina with @norm added.

## Examples

```
maize <- new("mina", tab = maize_asv2, des = maize_des2)
maize <- norm_tab(maize, method = "reref", depth = 1000, replace = TRUE,
multi = 3)
```

---

`norm_tab_method_list`    *List of normalization methods supported in [norm\\_tab](#)*

---

### Description

Normalization methods should be specified by exact string match.

### Usage

```
norm_tab_method_list
```

### Format

A list of character vectors.

`raref` By downsampling all samples to specific depth.

`total` Devided by the total read of each sample.

### See Also

[norm\\_tab](#)

### Examples

```
? norm_tab_method_list
```

---

`pcoa_plot`    *Visulization of components distance / dissimilarity in k dimension.*

---

### Description

Visulization of components distance / dissimilarity in k dimension.

Visulization of components distance / dissimilarity in k dimension.

### Usage

```
pcoa_plot(x, des, match, ...)
```

```
## S4 method for signature 'list,data.frame,character'
```

```
pcoa_plot(x, des, match, d1 = 1, d2 = 2, color, shape = NULL, ...)
```

**Arguments**

|       |                                                                                        |
|-------|----------------------------------------------------------------------------------------|
| x     | A list generated by 'dmr'.                                                             |
| des   | The corresponding descriptive table.                                                   |
| match | The column name of the components IDs in 'des' with exactly the same as rownames in x. |
| ...   | Additional parameters.                                                                 |
| d1    | The dimension be visualized in x-axis, default '1'.                                    |
| d2    | The dimension be visualized in y-axis, default '2'.                                    |
| color | The column name in 'des' to be used for different color groups.                        |
| shape | The column name in 'des' to be used for different shape groups, default 'NULL'.        |

**Value**

- p The plotted figure.
- p The plotted PCoA.

**Examples**

```

maize <- new("mina", tab = maize_asv2, des = maize_des2)
maize <- norm_tab(maize, method = "raref")
maize <- fit_tabs(maize)
maize <- com_dis(maize, method = "bray")
maize <- dmr(maize)
asv_dmr <- .dmr(maize)
des <- des(maize)
p1a <- pcoa_plot(asv_dmr, des, match = "Sample_ID", color = "Compartment")
p1b <- pcoa_plot(asv_dmr, des, match = "Sample_ID", d1 = 3, d2 = 4, color =
"Compartment")
p2a <- pcoa_plot(asv_dmr, des, match = "Sample_ID", color = "Host_genotype")
p2b <- pcoa_plot(asv_dmr, des, match = "Sample_ID", d1 = 1, d2 = 3, color =
"Host_genotype")
p3a <- pcoa_plot(asv_dmr, des, match = "Sample_ID", color = "Compartment",
shape = "Soil")
p3b <- pcoa_plot(asv_dmr, des, match = "Sample_ID", d1 = 1, d2 = 4, color =
"Compartment", shape = "Soil")
maize <- new("mina", tab = maize_asv2, des = maize_des2)
maize <- norm_tab(maize, method = "raref")
maize <- fit_tabs(maize)
maize <- com_dis(maize, method = "bray")
maize <- dmr(maize)
asv_dmr <- .dmr(maize)
des <- des(maize)
p1a <- pcoa_plot(asv_dmr, des, match = "Sample_ID", color = "Compartment")
p1b <- pcoa_plot(asv_dmr, des, match = "Sample_ID", d1 = 3, d2 = 4, color =
"Compartment")
p2a <- pcoa_plot(asv_dmr, des, match = "Sample_ID", color = "Host_genotype")
p2b <- pcoa_plot(asv_dmr, des, match = "Sample_ID", d1 = 1, d2 = 3, color =
"Host_genotype")

```

```
p3a <- pcoa_plot(asv_dmr, des, match = "Sample_ID", color = "Compartment",
  shape = "Soil")
p3b <- pcoa_plot(asv_dmr, des, match = "Sample_ID", d1 = 1, d2 = 4, color =
  "Compartment", shape = "Soil")
```

---

rarefaction\_subsample *Rarefaction subsample function, one sample, modified from a internal function in **phyloseq**, find it [here](https://rdrr.io/bioc/phyloseq/man/rarefy_even_depth.html).*

---

### Description

Rarefaction subsample function, one sample, modified from a internal function in **phyloseq**, find it [here](#).

### Usage

```
rarefaction_subsample(x, depth = 1000, replace = TRUE)
```

### Arguments

|         |                                                                                              |
|---------|----------------------------------------------------------------------------------------------|
| x       | A column of quantitative table.                                                              |
| depth   | The depth for rarefying, 1000 by default.                                                    |
| replace | Whether to sample with or without replacement, TRUE by default for computational efficiency. |

### Value

The vector rarefied to defined depth.

---

|              |                                                          |
|--------------|----------------------------------------------------------|
| re_format_AP | <i>Convert APResult (apcluster output) to dataframe.</i> |
|--------------|----------------------------------------------------------|

---

### Description

Modified from <https://rdrr.io/github/jefferis/flycircuit/src/R/clustering.R> #sym-as.data.frame.APResult

### Usage

```
re_format_AP(x)
```

### Arguments

|   |                                            |
|---|--------------------------------------------|
| x | an APResult object from <b>apcluster</b> . |
|---|--------------------------------------------|

### Value

y A data frame with columns 'ID', 'Exemplar', 'Cluster' and 'Cluster\_size'.

---

|               |                                               |
|---------------|-----------------------------------------------|
| re_format_MCL | <i>Convert mcl (mcl output) to dataframe.</i> |
|---------------|-----------------------------------------------|

---

**Description**

Modified from <https://rdr.io/github/jefferis/flycircuit/src/R/clustering.R> #sym-as.data.frame.APResult

**Usage**

```
re_format_MCL(x, names)
```

**Arguments**

|       |                                            |
|-------|--------------------------------------------|
| x     | an 'mcl' object from <a href="#">mcl</a> . |
| names | The names of clustered components.         |

**Value**

y A data frame with columns 'ID', 'Cluster' and 'Cluster\_size'.

**See Also**

[mcl](#)

---

|         |                                                                                                                                                                                                                                                                                                                                    |
|---------|------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| sim_par | <i>Function for community similarity calculation used by 'tina', modified from <a href="https://github.com/defleury/Schmidt_et_al_2016_community_similarity/blob/master/functions.community_similarity.R">https://github.com/defleury/Schmidt_et_al_2016_community_similarity/blob/master/functions.community_similarity.R</a></i> |
|---------|------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|

---

**Description**

Function for community similarity calculation used by 'tina', modified from [https://github.com/defleury/Schmidt\\_et\\_al\\_2016\\_community\\_similarity/blob/master/functions.community\\_similarity.R](https://github.com/defleury/Schmidt_et_al_2016_community_similarity/blob/master/functions.community_similarity.R)

**Usage**

```
sim_par(x, y, sim_method = "w_ja", threads = 80, nblocks = 400)
```

**Arguments**

|            |                                                                                                                          |
|------------|--------------------------------------------------------------------------------------------------------------------------|
| x          | An quantitative matrix.                                                                                                  |
| y          | The Cij matrix, which is correlation matrix of adjusted sparcc matrix of x.                                              |
| sim_method | The method for similarity, "w_ja" and "uw_ja" are available for weighted and unweighted Jaccard similarity respectively. |
| threads    | The number of threads used for parallel running, 80 by default.                                                          |
| nblocks    | The number of row / column for splitted sub-matrix, 400 by default.                                                      |

**Value**

s The output similarity matrix.

**Examples**

```
## Not run:
data(maize_asv)
maize_tab <- maize_asv[1 : 1000, 1 : 200]
asv <- norm_tab(maize_tab, method = "raref", depth = 100)
asv[is.na(asv)] <- 0
asv_sparcc <- sparcc(asv, threads = 8, nblocks = 40)
tmp.S <- adj(asv_sparcc, method = "spearman")
y <- 0.5 * (tmp.S + 1)
s <- sim_par(asv_sparcc, y, sim_method = "w_ja", threads = 8, nblocks = 40)

## End(Not run)
```

---

|        |                                                                                                                                                                                                                                                                                        |
|--------|----------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| sparcc | <i>Function for ‘sparcc’ correlation calculation. Modified from Schmidt et al. 2016, find the scripts <a href="https://github.com/defleury/Schmidt_et_al_2016_community_similarity">here</a> and the SparCC paper <a href="https://doi.org/10.1371/journal.pcbi.1002687">here</a>.</i> |
|--------|----------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|

---

**Description**

Function for ‘sparcc’ correlation calculation. Modified from Schmidt et al. 2016, find the scripts [here](#) and the SparCC paper [here](#).

**Usage**

```
sparcc(x, threads = 80, nblocks = 400)
```

**Arguments**

|         |                                                                     |
|---------|---------------------------------------------------------------------|
| x       | An matrix for correlation calculation.                              |
| threads | The number of threads used for parallel running, 80 by default.     |
| nblocks | The number of row /column for splitting sub-matrix, 400 by default. |

**Value**

y The adjacency matrix.

**Examples**

```
## Not run:
asv_sparcc <- sparcc(maize_asv2, threads = 2, nblocks = 40)

## End(Not run)
```

---

|       |                                              |
|-------|----------------------------------------------|
| tab<- | <i>Setter and getter for the slot 'tab'.</i> |
|-------|----------------------------------------------|

---

## Description

Setter and getter for the slot 'tab'.

## Usage

```
tab(x) <- value

## S4 replacement method for signature 'mina'
tab(x) <- value

tab(x)

## S4 method for signature 'mina'
tab(x)
```

## Arguments

|       |                                                         |
|-------|---------------------------------------------------------|
| x     | The 'mina' object.                                      |
| value | The value to set for the slot of the 'mina' object 'x'. |

## Value

The 'tab' slot of the 'mina' object.  
The 'tab' slot of the 'mina' object.  
The 'tab' slot of the 'mina' object.  
The 'tab' slot of the 'mina' object.

## Examples

```
tab(maize) <- maize_asv2
tab(maize)[1:5, 1:5]
```

---

|      |                                                                                                                                                                  |
|------|------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| tina | <i>TINA community dissimilarity used in <a href="#">com_dis</a>. Function for ‘tina’ dissimilarity/distance calculation. Modified from Schmidt et al., 2016.</i> |
|------|------------------------------------------------------------------------------------------------------------------------------------------------------------------|

---

**Description**

TINA community dissimilarity used in [com\\_dis](#). Function for ‘tina’ dissimilarity/distance calculation. Modified from Schmidt et al., 2016.

**Usage**

```
tina(x, ...)
```

**Arguments**

|     |                                          |
|-----|------------------------------------------|
| x   | An matrix for dissimilarity calculation. |
| ... | Additional parameters.                   |

**Value**

The output ‘tina’ dissimilarity matrix.

**Examples**

```
## Not run:
asv_norm <- norm_tab(maize_asv2, method = "raref", depth = 1000)
asv_dis <- com_dis(asv_norm, method = "bray")
asv_dis <- com_dis(asv_norm, method = "tina", threads = 8, nblocks = 40)
asv_tina <- tina(asv_norm, cor_method = "spearman", sim_method = "w_ja",
threads = 8, nblocks = 40)

## End(Not run)
```

---

|                    |                                                                                                                                                                                                                           |
|--------------------|---------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| tina,matrix-method | <i>Function for ‘tina’ dissimilarity calculation. Modified from Schmidt et al., 2016. Person and Spearman could be used for correlation and weighted and unweighted Jaccard could be used for similarity calculation.</i> |
|--------------------|---------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|

---

**Description**

Function for ‘tina’ dissimilarity calculation. Modified from Schmidt et al., 2016. Person and Spearman could be used for correlation and weighted and unweighted Jaccard could be used for similarity calculation.

**Usage**

```
## S4 method for signature 'matrix'
tina(
  x,
  cor_method = "spearman",
  sim_method = "w_ja",
  threads = 80,
  nblocks = 400,
  ...
)
```

**Arguments**

|            |                                                                                                                          |
|------------|--------------------------------------------------------------------------------------------------------------------------|
| x          | A matrix for dissimilarity calculation.                                                                                  |
| cor_method | The method for correlation, "pearson" and "spearman" are available.                                                      |
| sim_method | The method for similarity, "w_ja" and "uw_ja" are available for weighted and unweighted Jaccard similarity respectively. |
| threads    | The number of threads used for parallel running, 80 by default.                                                          |
| nblocks    | The number of row and column for splitted sub-matrix, 400 by default.                                                    |
| ...        | Additional parameters.                                                                                                   |

**Value**

t The output 'tina' dissimilarity matrix.

**Examples**

```
## Not run:
asv_norm <- norm_tab(maize_asv2, method = "raref", depth = 1000)
asv_dis <- com_dis(asv_norm, method = "bray")
asv_dis <- com_dis(asv_norm, method = "tina", threads = 8, nblocks = 40)
asv_tina <- tina(asv_norm, cor_method = "spearman", sim_method = "w_ja",
  threads = 8, nblocks = 40)

## End(Not run)
```

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