# Package 'samovaR'

May 26, 2025

Type Package

```
Title R package for generating model metagenomes with specified properties
Description There is a fundamental problem in modern metagenomics: there are huge differences be-
      tween methodological approaches that strongly influence the results, while remaining out-
      side the attention of researchers. We propose an approach that utilizes de novo genera-
      tion of the artificial metagenomes - SamovaR.
URL https://github.com/ctlab/samovar/
License MIT + file LICENSE
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      xml2,
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      scclust,
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      htmlwidgets,
      progress,
      methods,
      dplyr,
      ggplot2,
      stringr,
      tibble
Suggests ggnewscale,
      shiny,
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      rmarkdown,
      testthat (>= 3.0.0),
      withr
```

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# VignetteBuilder knitr

# Config/testthat/edition 3

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annotation2samovar

Process annotation data.frame to SamovaR

# Description

Process annotation data.frame to SamovaR

# Usage

annotation2samovar(data)

# Arguments

data

Processed abundance table. Row names: sequence IDs, column names: annotators; true for true annotation

# Value

list of samovar data objects

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## **Examples**

```
library(samovaR)

data <- read_annotation_dir("data/test_annotations/")
viz_annotation(data)</pre>
```

build\_samovar

Build samovar object

## **Description**

Samovar network is a 2D-oriented graph with metadata and abundances of species per sample. Oriented graph could be used for network prediction, or for better generation some network could be used as initial (to be implemented) For better understanding of building database and using it in generation, visit github source

## Usage

```
build_samovar(
   samovar_data,
   dist_function = function(x) dist(x),
   network = F,
   k_means = F,
   min_cluster_size = F,
   plot_log = T
)
```

## **Arguments**

```
samovar_data samovar data after preprocessing stages

network FALSE or graph that can be used for generation. To be implemented

plot_log Logical or path for log plots output

distance_function
    function used for measuring distances between species based on samples

min_min_cluster_size
    FALSE or minimum number of species per cluster

max_min_cluster_size
    FALSE or minimum number of species per cluster
```

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concotion\_pour

Build samovar object

## **Description**

Samovar network is a 2D-oriented graph with metadata and abundances of species per sample. Oriented graph could be used for network prediction, or for better generation some network could be used as initial (to be implemented) For better understanding of building database and using it in generation, visit github source

#### Usage

```
concotion_pour(
  samovar_data,
  inner_method = "glm",
  inter_method = "glm",
  inner_model = "gaussian",
  inter_model = "gaussian",
  minimal_cluster = 2,
  probability_calculation = "oriented",
  cluster_connection = "mean"
)
```

#### **Arguments**

samovar\_data samovar data after preprocessing stages Character, glm, other to be implemented (bootstrap, bsPCA) inner\_method inter\_method Character, glm, other to be implemented (bootstrap, bsPCA) inner\_model Character, model processed by glm(). For glm mode only. quasipoisson by default Character, model processed by glm(). For glm mode only. quasipoisson by inter\_model default cluster\_connection Character (mean, median), or function. The way of cluster connection. If function, way of summarize all samples of species cluster FALSE or graph that can be used for generation. To be implemented network Character, co-occurrence calculation. cooccurrence If "simple", calculated as: P(A|B) = sum(A&B)/sum(A|B). If "oriented", calculated as P(A|B) = P(A&B|B)If "compositional", calculated as P(A|B) = P(A&B|B), and than sampled one of represented conditions of occurence

```
library(samovaR)
library(tidyverse)

# download data
teatree <- GMrepo_type2data(number_to_process = 2000)</pre>
```

GMrepo\_run 5

```
# filter
tealeaves <- teatree %>%
  teatree_trim(treshhold_species = 3, treshhold_samples = 3, treshhold_amount = 10^(-3))
# normalizing
## if you build teatree by your own, rescaling stage when building via teatree$rescale()
## or assigning teatree$min_value and teatree$max_value is required
## good approximation to normal distribution is required for glm generating methods
teabag <- tealeaves %>%
  tealeaves_pack(normalization_function = function(x) log10(x+1))
# clustering
concotion <- teabag %>%
  teabag_brew(min_cluster_size = 4, max_cluster_size = 6)
# remember: if you want to refilter, it is better to re-do welding stage to avoid crashes in future!
# building samovar
samovar <- concotion %>%
  concotion_pour()
```

GMrepo\_run

GMrepo run data class

# Description

GMrepo run data class

## **Slots**

metadata metadata DataFrame run character

GMrepo\_run2data

Get data from GMrepo\_run object

## **Description**

Get data from GMrepo\_run object

```
GMrepo_run2data(
  run,
  number_to_out = F,
  at_level = "species",
  QC_filter = "QCStatus"
)
```

6 GMrepo\_type2data

## **Arguments**

number\_to\_out False by default, maximum number of obtained data

at\_level "species" by default. level to obtain classification from GMrepo

QC\_filter QCStatus by default. Perform auto QC filtering based on metadata column, or

False for no checking.

runs GMrepo\_run object got by GMrepo\_type2run or created by user with new('GMrepo\_run',

metadata = data.frame(), run = run\_list)

## **Examples**

```
library(samovaR)
library(tidyverse)
# get data from GMrepo
data_GMrepo <- GMrepo_type2data(mesh_ids = "D006262", number_to_process = 1000)</pre>
# equal to:
run_GMrepo <- GMrepo_type2run(mesh_ids = "D006262", number_to_process = 1000)</pre>
data_GMrepo <- GMrepo_run2data(run_GMrepo)</pre>
# filter runs before obtaining data (OOP updating data!)
run_GMrepo$filter("checking", 1)
# view
data_GMrepo
# access to metadata
data_GMrepo$run
# access to data
data_GMrepo$data
# access to runs
data_GMrepo$run
# access to taxa
data_GMrepo$species
```

 ${\tt GMrepo\_type2} data$ 

Get data from GMrepo

## **Description**

Wrapper around GMrepo\_type2run and GMrepo\_run2data functions

```
GMrepo_type2data(
  mesh_ids = c("D006262"),
  number_to_process = F,
  number_to_out = F,
  at_level = "species",
  QC_filter = "QCStatus"
)
```

GMrepo\_type2run 7

## **Arguments**

mesh\_ids Character. All types of meshID to use. List of relations between meshID and

phenotype could be obtained using GMrepo\_meshID(). Health meshID by de-

faul

number\_to\_process

False by default, or maximum number of runs per meshID

number\_to\_out False by default, maximum number of obtained data

at\_level "species" by default. level to obtain classification from GMrepo

QC\_filter QCStatus by default. Perform auto QC filtering based on metadata column, or

False for no checking.

#### **Examples**

```
library(samovaR)
library(tidyverse)
# get data from GMrepo
data_GMrepo <- GMrepo_type2data(mesh_ids = "D006262", number_to_process = 1000)</pre>
# equal to:
run_GMrepo <- GMrepo_type2run(mesh_ids = "D006262", number_to_process = 1000)</pre>
data_GMrepo <- GMrepo_run2data(run_GMrepo)</pre>
# filter runs before obtaining data (OOP updating data!)
run_GMrepo$filter("checking", 1)
# view
data_GMrepo
# access to metadata
data_GMrepo$run
# access to data
data_GMrepo$data
# access to runs
data_GMrepo$run
# access to taxa
data_GMrepo$species
```

 ${\it GMrepo\_type2run}$ 

Get runs from GMrepo by meshID

## Description

Get runs from GMrepo by meshID

```
GMrepo_type2run(mesh_ids = c("D006262"), number_to_process = F)
```

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## **Arguments**

False by default, or maximum number of runs per meshID

## **Examples**

```
library(samovaR)
library(tidyverse)
# get data from GMrepo
data_GMrepo <- GMrepo_type2data(mesh_ids = "D006262", number_to_process = 1000)</pre>
# equal to:
run_GMrepo <- GMrepo_type2run(mesh_ids = "D006262", number_to_process = 1000)</pre>
data_GMrepo <- GMrepo_run2data(run_GMrepo)</pre>
# filter runs before obtaining data (OOP updating data!)
run_GMrepo$filter("checking", 1)
# view
data_GMrepo
# access to metadata
data_GMrepo$run
# access to data
data_GMrepo$data
# access to runs
data_GMrepo$run
# access to taxa
data_GMrepo$species
```

log\_plot

Print a log plot

# Description

Print a log plot

```
log_plot(plot_log, postfix, gg, mode = "ggplot", write = F)
```

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minmaxscale

Misc functions

# Description

Misc functions

## Usage

```
minmaxscale(x)
```

progress\_function

Progress bar

# Description

Progress bar

# Usage

```
progress_function(iters)
```

 $read\_annotation\_dir$ 

Read annotations produced with samovar pipeline from the directory

## **Description**

Read annotations produced with samovar pipeline from the directory

# Usage

```
read_annotation_dir(data_dir, sample_name_position = 0, ...)
```

# **Arguments**

```
data_dir Path to abundance table. Row names: sequence IDs, column names: annotators; true for true annotation sample_name_position Position to split the file basename by "." to extract sample names. 0 by default ... Parameters processed by read.csv()
```

```
library(samovaR)

data <- read_annotation_dir("data/test_annotations/")
viz_annotation(data)</pre>
```

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read\_annotation\_table Build samovar data object from file or environment

# Description

Build samovar data object from file or environment

#### Usage

```
read_annotation_table(data, metadata = F, ...)
```

# Arguments

| 1 4  | Data frame or path to ab   | 1 (1          | D : :                 | . 1 1                     |
|------|----------------------------|---------------|-----------------------|---------------------------|
| data |                            |               |                       |                           |
| uata | Data.Hallic Ol Datil to at | junuance inc. | IXOW HAILICS IS USHIE | as succies list, coluilli |

names as sample list. Unique names required

samovar base class

metadata Data.frame or path to metadata file
... Parameters processed by read.csv()

# Description

samovar\_base

samovar base class

## **Slots**

samovar\_base samovar\_data object

method method to obtain samovar\_base

inner\_cluster\_graph\_method list of graphs in matrix form of inner cluster connections

inter\_cluster\_graph\_method list of graphs in matrix form of inter cluster connections

inner\_cluster\_graph\_prob list of co-occurrence probabilities in matrix form of inner cluster members

inter\_cluster\_graph\_prob list of co-occurrence probabilities in matrix form between clusters
properties concotion\_pour() properties

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samovar\_boil

Generate artificial data

#### **Description**

Use pre-built samovar\_data with its parameters

## Usage

```
samovar_boil(
  samovar_base,
  N = 1,
  init_sp = F,
  init_ab = F,
  avoid_zero_generations = T,
  seed = 42
)
```

#### **Arguments**

samovar data after preprocessing and building stages samovar\_base Ν number of artificial samples to generate init\_sp species vector for initializing data generation, or FALSE for usage most common taxa, auto for choosing random taxa init\_ab species amount vector (values from 0 to 1) for initializing data generation, or FALSE for mean initial taxa assignment, or auto for usage from known edf for each species from init\_sp avoid\_zero\_generations logical, avoid zero-based generations or not. FALSE might results in underdistributed communities, while TRUE in over-represented with species from different clusters possibly come from different samples groups seed initial seed for the seeds generation

```
library(samovaR)
library(tidyverse)

# download and prepare data
samovar <- GMrepo_type2data(number_to_process = 2000) %>%
  teatree_trim(treshhold_species = 3, treshhold_samples = 3, treshhold_amount = 10^(-3)) %>%
  tealeaves_pack(normalization_function = function(x) log10(x+1)) %>%
  teabag_brew(min_cluster_size = 4, max_cluster_size = 6) %>%
  concotion_pour()

# generate
new_data <- samovar %>%
  samovar_boil(N = 100)
```

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samovar\_data

samovar data class

# Description

samovar data class

#### **Slots**

```
description metadata DataFrame
data DataFrame with species abundances. No NA pass
run character, runs
species character, runs
normalization_function normalization function for samples
reverse_normalization_function reverse normalization function
min_value minimal value after scaling
max_value maximal value after scaling
cluster character vector, enumerated clusters for each species
cluster_size named numeric, cluster sizes per cluster
```

samovar\_run

Samovar run data class

# Description

Samovar run data class

#### **Slots**

```
metadata metadata DataFrame
data data
run character, samle IDs
```

## Methods

export(Class) Returns the result of coercing the object to Class. No effect on the object itself.

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teabag\_brew

Build samovar object

## **Description**

Samovar network is a 2D-oriented graph with metadata and abundances of species per sample. Oriented graph could be used for network prediction, or for better generation some network could be used as initial (to be implemented) For better understanding of building database and using it in generation, visit github source

# Usage

```
teabag_brew(
  samovar_data,
  dist_function = function(x) dist(x),
  network = F,
  min_cluster_size = 10,
  max_cluster_size = 100,
  plot_log = F
)
```

## **Arguments**

```
network FALSE or graph that can be used for generation. To be implemented min_cluster_size
FALSE or minimum number of species per cluster
max_cluster_size
FALSE or minimum number of species per cluster
plot_log Logical or path for log plots output
data samovar data after preprocessing stages
distance_function
```

function used for measuring distances between species based on samples

tealeaves\_pack

Scale species abundances

# Description

Scale species abundances

```
tealeaves_pack(
  samovar_data,
  normalization_function = function(x) log10(x + 1),
  plot_log = T
)
```

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#### **Arguments**

```
samovar_data Samovar data object to rescale
normalization_function
Function using for rescaling
plot_log Logical or path for log plots output
```

# **Examples**

```
library(samovaR)
library(tidyverse)
# download data
teatree <- GMrepo_type2data(number_to_process = 2000)</pre>
# filter
tealeaves <- teatree %>%
  teatree\_trim(treshhold\_species = 3, treshhold\_samples = 3, treshhold\_amount = 10^(-3))
# normalizing
## if you build teatree by your own, rescaling stage when building via teatree$rescale()
## or assigning teatree$min_value and teatree$max_value is required
## good approximation to normal distribution is required for glm generating methods
teabag <- tealeaves %>%
  tealeaves\_pack(normalization\_function = function(x) log10(x+1))
# clustering
concotion <- teabag %>%
  teabag_brew(min_cluster_size = 4, max_cluster_size = 6)
# remember: if you want to refilter, it is better to re-do welding stage to avoid crashes in future!
# building samovar
samovar <- concotion %>%
  concotion_pour()
```

teatree\_trim

Filter species and samples from samovar\_data object

#### **Description**

Filter species and samples from samovar\_data object

```
teatree_trim(
  samovar_data,
  metadata_filter = F,
  treshhold_amount = 10^(-5),
  treshhold_samples = 1,
  treshhold_species = 1,
  drop_species = F,
  drop_unclassified = T
)
```

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## **Arguments**

```
samovar_data Samovar data object to filter

metadata_filter

False, character or data.frame with 2 columns: first contain metadata names for filtering, and second values per column

treshhold_amount

Minimum value to conclude as not the noise.

treshhold_samples

Minimum number of representing samples to keep species.

treshhold_species

Minimum number of representing species to keep samples.

drop_unclassified

Drop unknown and unclassified ranks. True by default
```

## **Examples**

```
library(samovaR)
library(tidyverse)
# download data
teatree <- GMrepo_type2data(number_to_process = 2000)</pre>
# filter
tealeaves <- teatree %>%
  teatree_trim(treshhold_species = 3, treshhold_samples = 3, treshhold_amount = 10^(-3))
# normalizing
## if you build teatree by your own, rescaling stage when building via teatree$rescale()
## or assigning teatree$min_value and teatree$max_value is required
## good approximation to normal distribution is required for glm generating methods
teabag <- tealeaves %>%
  tealeaves_pack(normalization_function = function(x) log10(x+1))
# clustering
concotion <- teabag %>%
  teabag_brew(min_cluster_size = 4, max_cluster_size = 6)
# remember: if you want to refilter, it is better to re-do welding stage to avoid crashes in future!
# building samovar
samovar <- concotion %>%
  concotion_pour()
```

viz\_annotation

Visualize annotation results

# Description

Visualize annotation results

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## Usage

```
viz_annotation(
  data,
  type = c("f1", "R2", "cv"),
  show_top = 10,
  output_dir = NULL,
  plot = T
)
```

## **Arguments**

data Processed abundance table. Row names: sequence IDs, Column names:

• annotators: (starting with taxID\_);

• true: for true annotation

• length: length of sequence

• sample

type character vector.

• if present column true\_annotation: could be one of "f1", "R2", or their

combination

• else: "cross-validation"

show\_top integer. Number of top annotations to show.

output\_dir character. Directory to save the plots. If NULL, plots are not saved.

plot logical. If TRUE, plots are printed.

# Value

list of ggplot objects

# **Examples**

```
library(samovaR)

data <- read_annotation_dir("data/test_annotations/")
viz_annotation(data)</pre>
```

 $viz\_composition$ 

Visualize composition

# Description

Visualize composition

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## Usage

```
viz_composition(
  data,
  reord_samples = "fpc",
  reord_species = "amount",
  type = "column",
  top = 15,
  interactive = F,
  ggplot_add = F,
  bottom_legend = F
```

# **Arguments**

data data.frame with dimensions of species \* samples, or samovar objects: samovar\_data,

samovar\_base, samovar\_run or GMrepo\_run (with data)

reord\_samples character, fpc, fpc\_scaled, hcl, amount, tsne, or none reorder of samples on plot

reord\_species character, same for reord\_samples

type character, column or tile for composition visualize, or donut (to implement) for

mean composition visualization

top integer, number of top-represented taxa to show, or FALSE to show all

interactive logical. ggplot or plotly object to return ggplot\_add functions to add to ggplot object, or FALSE.

bottom\_legend vector length of samples to show on plot as a color legend, or FALSE

```
library(samovaR)
library(tidyverse)

# Download data
teatree <- GMrepo_type2data(number_to_process = 1000)

#Composition
viz_composition(teatree)</pre>
```

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