Gleesso

December 13, 2017

batch_converter

batch community converter

Description

convert community (give the same indice) of a batch of graphs to the community of a reference graph

Usage

```
batch_converter(graph_batch, graph_ref)
```

Arguments

graph_batch : a list of graph to convert

graph_ref : the reference used for conversion

 $community_contrast_dashboard$

community_contrast_dashboard

Description

create a graphic with a facet by community and a boxplot by modality of the contrast vector

Usage

```
community_contrast_dashboard(community_table, contrast_name, contrast_vector,
  nrow = 4, levels_class = c("low", "high"), color_vect = NULL,
  offset = 10^-8)
```

Arguments

community_table

: a table of community composition generated by the Compute_community_abondance

function

contrast_name : how the contrast column should be named

contrast_vector

: the contrast vector

community_converter community_converter

Description

Community converter function take to two nodes table as argument and give a translation of each community to the other graph based on jacquart distance If I may, it is automated community translation

Usage

```
community_converter(nodes_graph1, nodes_graph2, join_type = "inner")
```

Arguments

nodes_graph1 is the table of nodes with the walktrap column properly filled nodes_graph2 is the table of nodes of the second graph

community_taxa_abundance

community_taxa_abundance

Description

This function generate latex table to help the interpretation of community: * Table of the list of species in each community * Compute the proportion of taxa for each community and create: * create a table listing taxa and their proportion * create a barplot of taxa proportion

Usage

```
community_taxa_abundance(Nodes, MGS_by_taxo_species, file_output,
  community_kind = "walktrap_community", width_graph = 14,
  height_graph = 5, community_levels = NULL, prevalence_level = 0.01)
```

Arguments

Nodes graph table

MGS_by_taxo_species

: Nodes abundance

file_output : file prefix for relative abundance figure

Compute_community_abondance

compute community abundances

Description

Compute the sum of abundance of species in each community given the abundance matrix by species and the graph community Also compute the p-value of difference of abundance given a contrast vector with to class (2 classes)

Usage

```
Compute_community_abondance(Nodes, abundance, species_taxo, contrast = NULL,
   community_kind = "walktrap_community")
```

Arguments

Nodes: The network node table with the a community attribution column

abundance: The Metagenomic species abundance table

species_taxo: The Metagenomic species taxonomy table

contrast: a boolean vector to form two group of samples. for each community the rank

test difference of abundance p-value is calculated between the two groups.

community kind: the algorithm of used to compute the community: "spinglass_community",

"walktrap_community"

Value

a table of community abundance and composition

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Compute_graph

Compute Glasso

Description

Compute the Glasso model from the MGS abundances on individuals with a true value in the selection vector.

Usage

```
Compute_graph(MGS_abundance, selection_vector, fout,
  abundance_treshold = 10^-7, occurrence_treshold = 0.05, nlambda = 20,
  lambda.min.ratio = 0.1, lambda = NULL, rep.num = 20)
```

Arguments

selection_vector

: a boolean vector to select a subset of the cohort (the model will be infered on

samples with a TRUE value)

MGS_abundance: The Metagenomic species abundance table

fout: where to save the model object

community: Should the community structure be calculated?

nlambda: Number of regularisation parameter that will be tested (see huge::huge() docu-

mentation)

lambda.min.ratio:

the smallest value of lambda as a fraction of its maximum (see huge::huge()

documentation)

occurence_treshold:

minimum fraction of samples where a species must be present to be taken into

account in the analysis

abundance_threshold:

minimum mean abundances for a species to be included in the analysis

rep.num: Number of subsampling to compute the edge stability with "StarS" (see huge::huge.select

documentation)

lambda: A sequence of regularisation parameter. If not null, it will override the automatic

computation of the lambda sequence (with nlambda and lambda.min.ratio)

concordance_table 5

concordance_table concordance_table

Description

Enable one to assess if community found in diverse cohort are the same Used to generate the alluvial plot

Usage

```
concordance_table(nlist, Graph_tags, join_type = "outer")
```

Arguments

nlist : list of graphs nodes table with the walktrap_community information Available

Graph_tags : list of graph labels

join_type : how to join graph row (outer joins or inner join). Outer join means that all

species present in at least one graph will be taken into account. Inner join means

that only species present in all graphs will treated.

Description

The Function also compute the community structure of the graph with various algorithms (betweeness community, walktrap community...) community specified by the user

Usage

```
create_graph(file_input, file_output, MGS_by_taxo_species, species_taxo,
nspins = 20, mod_rep = 10, community = FALSE, additional_info = NULL,
spinglass_opt = FALSE, variability_treshold = NULL)
```

Arguments

spinglass_opt : Should the number of spin of the spinglass community be optimized?

file_input: emplacement of the GLASSO model object file_output: where to save the network representation file

MGS_by_taxo_species:

species_taxo:

The Metagenomic species abundance table
The Metagenomic species taxonomy table

nspins: number of spin for the spinglass community detection algorithm

```
community: Should the community structure be calculated? additional_info:
```

a vector or data.frame containing information to add to the nodes table of the

variability_treshold:

The maximum mean variability for graph edge presence. If null, the optimal covariance matrix will correspond to a variability of 0.05

Description

create a gephi file for nodes with a robust community attribution

Usage

```
create_graph_robust_community_tags(model_folder, fout, abund_by_species,
  taxo_by_species, model_tag, Robust_table_community,
  Nodes_table_on_all_samples, variability_treshold = NULL)
```

Arguments

fout : where to save the csv tables with nodes with the robust community column

abund_by_species

: abundance mean group by species or

taxo_by_species

: taxo grouped at the species level

Robust_table_community

: Community attribution

Nodes tables computed on all samples

```
draw_community_total_abundance
```

draw community total abundance treshold

Description

A function to draw a barplot of the mean abundance of community accross samples

Usage

```
draw_community_total_abundance(abund)
```

Arguments

abund : table of community abundance accross samples

Description

draw a barplot of the number of species by community

Usage

```
draw_community_total_species_count(community_table)
```

Arguments

```
community_table
```

: Community table with the species composition line

Description

function to handily extract only the abondance of community by samples tables (and remove the extra information from the table like community composition)

Usage

```
extract_community_abundance_table(Community_table)
```

Arguments

```
Community_table
```

: a table of community composition generated by the Compute_community_abondance function $\,$

8 Gleesso_bootstrap

```
generate_graph_from_tables
```

generate_graph_from_tables create a gephi graph from a Nodes table and Glasso model object

Description

generate_graph_from_tables create a gephi graph from a Nodes table and Glasso model object

Usage

```
generate_graph_from_tables(fout, nodes_viz_att, fgraph_model,
  variability_treshold = NULL)
```

Gleesso_bootstrap

bootstrap Gleesso_pipeline

Description

Apply the bootstrap pipeline to a fraction of the cohort. A factor vector can be supplied to stratify the different samples

Usage

```
Gleesso_bootstrap(N_bootstrap, fraction, tag_model, variability_treshold,
  community_table_folder, model_folder, graph_folder, MGS_file, taxo_file,
  stratifying_vector = NULL, ...)
```

Arguments

N_bootstrap : number of different bootstrap samples that should be drawn

fraction : fraction of the initial dataset that should be drawn to form each bootstrap

samples

tag_model :
stratifying_vector

: a factor vector that represent a class that should be evenly scattered between

bootstrap samples

... : parameters to pass to the

Graphs_folder : folder to output all graphs and all bootstrap samples (to keep track of which

individual was used in each iteration)

Gleesso_pipeline 9

Description

Launching the complete glasso analysis (data prep, graph inference, community detection, community abundances computation) Warning: folder shouldn't be indicated with an / at the end

Usage

```
Gleesso_pipeline(data_folder, MGS_file, taxo_file, model_folder, graph_folder,
    selection_vector, tag_model, tag_graph, community = TRUE, nlambda = 20,
    lambda.min.ratio = 0.1, occurence_treshold = 0.05,
    abundance_treshold = 10^-7, variability_treshold = NULL,
    analysis_step = NULL, species_mode = TRUE)
```

Arguments

data_folder : where the community abundance table will be written

selection_vector

: a boolean vector to select a subset of the cohort (the model will be infered on

samples with a TRUE value)

tag_model : a tag that will be inserted in output file to recognize the model parameter

tag_graph : a tag that will be inserted in output file to recognize the graph

species_mode : should the graph inference be done on MGS (FALSE) or with MGS of the

same specied merged togethere (TRUE)

MGS_file: The Metagenomic species abundance file in the RDS format

model_folder: where to save the graphical Lasso model object (output of the spiec.easi func-

tion)

graph_folder: where the graphical representation of the model will be saved (in the gephi for-

mat)

community: Should the community structure be calculated?

nlambda: Number of regularisation parameter that will be tested (see huge::huge() docu-

mentation)

lambda.min.ratio:

the smallest value of lambda as a fraction of its maximum (see huge::huge()

documentation)

occurence_treshold:

minimum fraction of samples where a species must be present to be taken into

account in the analysis

abundance_threshold:

minimum mean abundances for a species to be included in the analysis

variability_treshold:

The maximum mean variability for graph edge presence. If null, the optimal covariance matrix will correspond to a variability of 0.05

analysis_step:

At which step the analysis should be started (0: from scratch, 1: model inferences, 2: save gephi network, 3: Community detection). If NULL (default), the step will be infered from the files present in the output folders. Use analysis_step=0 to force computation from scratch.

opt_spinglass_com

Computing spinglass communities and their modularity for a range of number of spin We then retrieve the optimal number of spin according to modularity

Description

Computing spinglass communities and their modularity for a range of number of spin We then retrieve the optimal number of spin according to modularity

Usage

```
opt_spinglass_com(con.grph, mod_rep, nspins)

parrallel_coord_community

parrallel_coord_community
```

Description

Alluvial plot of concordance of community Draw a parrallel coord graph of community belonging for different graph object. Enable one to compare and understand the stability or discrepancy between graph community

Usage

```
parrallel_coord_community(graph_node_list, Graph_tags, measure = "sum_ab",
    color_graph = 1, join_type = "inner")
```

Arguments

color_graph : index of the graph used to color the parallel coordiante plot

join_type : take the intersection or the union of species?

graph_node_list:

a sequence of nodes tables with community annotated

Graph_tags: a sequence of str which are the name of nodes tables of graph_node_list

measure: the weight attributed to each CAG either "sum" of abundance or "count" of

objects

+ 11

overload '+' operator to allow character strings concatenation

Description

overload '+' operator to allow character strings concatenation

Usage

```
"+"(e1, e2)
```

Robust_table_community

Community attribution stability table

Description

Compute the stability of species community attribution from bootstraped graphs and community table abondance for robust attribution.

Usage

```
Robust_table_community(graphs_folder, alluvial_diagnostic_file, taxo,
   N_alluvial = 10, join_type = "outer", stability_treshold = 0.6,
   silhouette_treshold = 0.1)
```

Arguments

graphs_folder : folder where all graphs are placed (with the / at the end please ^^)

N_alluvial : number of graph to represent on a graph

join_type : Should we work on the union of species or the intersection

alluvial_diagnostic

: file name for the alluvial graph

Value

```
a list object containing the following elements: staby: table of species stability with community assignation for all graphs stab_n_taxo: table of species stability and taxonomy Robust_community_stability_[...]
```

 $: abundance \ of \ communities \ for \ species \ with \ a \ stability \ above \ the \ specified \ treshold \ Robust_community_stability_[...]_silhout$

: idem but species also have a silhouette above a specified treshold

```
Silhouette\_to\_community \\ Silhouette\_to\_community
```

Description

Compute the silhouette cluster metric for all species

Usage

```
Silhouette_to_community(my_dist, Nodes_with_com)
```

Arguments

my_dist : distance matrix of species to all species

 $Nodes_with_com: Nodes table with a walktrap_community attribution$

Description

stability_index function look at the walktrap community of each species in a list of graph then compute the number of graph where the species as been attributed to the same community

Usage

```
stability_index_converter(graph_list, join_type = "outer")
```

Arguments

graph_batch : a list of graph with walktrap community converted to a ref

walktrap_distance 13

walktrap_distance wal

walktrap_distance

Description

reproduce the distance used in the walktrap community detection algorithm

Usage

```
walktrap_distance(pos.grph, n_steps)
```

Arguments

pos.graph: an igraph object that contain only positive edges n_steps: number of steps of the random walk on the graph

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