

Package ‘MLVSBM’

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Type Package

Title A Stochastic Block Model for Multilevel Networks

Version 0.2.4

Description Simulation, inference and clustering of multilevel networks using a Stochastic Block Model framework as described in Chabert-Liddell, Barbillon, Donnet and Lazega (2021) <[doi:10.1016/j.csda.2021.107179](https://doi.org/10.1016/j.csda.2021.107179)>. A multilevel network is defined as the junction of two interaction networks, the upper level or inter-organizational level and the lower level or inter-individual level. The inter-level represents an affiliation relationship.

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ARI

*Compare two clustering with the Adjusted Rand Index***Description**

Compare two clustering with the Adjusted Rand Index

Usage

```
ARI(x, y)
```

Arguments

x A vector of integers, the clusters labels
y A vector of integers of the same length as x, the clusters labels

Value

A number between 0 (random clustering) and 1 (identical clustering)

Examples

```
ARI(x = c(1, 2, 1), y = c(2, 2, 1))
```

build_fold_matrix	<i>Title</i>
-------------------	--------------

Description

Title

Usage

```
build_fold_matrix(X, K)
```

Arguments

X	An adjacency matrix
K	An integer, the number of folds

Value

A matrix of the same size than X with class integer as coefficient

coef.FitMLVSBM	<i>Extract model coefficients</i>
----------------	-----------------------------------

Description

Extracts model coefficients from objects with class [FitMLVSBM](#)

Usage

```
## S3 method for class 'FitMLVSBM'
coef(object, ...)
```

Arguments

object	an R6 object of class FitMLVSBM
...	additional parameters for S3 compatibility. Not used

Value

List of parameters.

 FitMLVSBM

An R6 Class object, a fitted multilevel network once \$dovem() is done

Description

An R6 Class object, a fitted multilevel network once \$dovem() is done

An R6 Class object, a fitted multilevel network once \$dovem() is done

Public fields

vbound The vector of variational bound for monitoring convergence

Active bindings

affiliation_matrix Get the affiliation matrix

adjacency_matrix Get the list of adjacency matrices

nb_nodes Get the list of the number of nodes

nb_clusters Get the list of the number of blocks

parameters Get the list of the model parameters

membership Get the list of the variational parameters

independent Are the levels independent?

distribution Emission distribution of each level

directed Are the levels directed?

entropy Get the entropy of the model

bound Get the variational bound of the model

df_mixture Get the degrees of freedom of the mixture parameters

df_connect Get the degrees of freedom of the connection parameters

connect Get the number of possible observed connections

ICL Get the ICL model selection criterion of the model

full_penalty Get the penalty used to compute the ICL

Z Get the list of block memberships (vector form)

X_hat Get the list of the matrices of probability connection predictions

map Get the list of block memberships (matrix form)

penalty Get the ICL penalty

likelihood Compute the likelihood of both levels

complete_likelihood Get the complete likelihood of the model

Methods**Public methods:**

- `FitMLVSBM$new()`
- `FitMLVSBM$update_alpha()`
- `FitMLVSBM$update_pi()`
- `FitMLVSBM$update_gamma()`
- `FitMLVSBM$init_clustering()`
- `FitMLVSBM$clear()`
- `FitMLVSBM$m_step()`
- `FitMLVSBM$ve_step()`
- `FitMLVSBM$do_vem()`
- `FitMLVSBM$permute_empty_class()`
- `FitMLVSBM$plot()`
- `FitMLVSBM$show()`
- `FitMLVSBM$print()`
- `FitMLVSBM$clone()`

Method `new()`: Constructor for the FitMLVSBM class

Usage:

```
FitMLVSBM$new(
  Q = list(I = 1, O = 1),
  A = NA,
  X = NA,
  M = list(I = NA, O = NA),
  directed = NA,
  distribution = list("bernoulli", "bernoulli"),
  independent = FALSE
)
```

Arguments:

Q List of number of blocks
 A Affiliation matrix
 X List of adjacency matrices
 M List of Mask matrices
 directed List of boolean
 distribution List of string
 independent Boolean

Returns: A FitMLVSBM object

Method `update_alpha()`: Update the connection parameters for the M step

Usage:

```
FitMLVSBM$update_alpha(safeguard = 2 * .Machine$double.eps)
```

Arguments:

safeguard Parameter live in a compact [safeguard, 1-safeguard]

Method `update_pi()`: Update the upper level mixture parameter for the M step

Usage:

```
FitMLVSBM$update_pi(safeguard = 0.001)
```

Arguments:

`safeguard` Parameter live in a compact [`safeguard`, 1-`safeguard`]

Method `update_gamma()`: Update the lower level mixture parameter for the M step

Usage:

```
FitMLVSBM$update_gamma(safeguard = 1e-06)
```

Arguments:

`safeguard` Parameter live in a compact [`safeguard`, 1-`safeguard`]

Method `init_clustering()`: `init_clustering` Initial clustering for VEM algorithm

Usage:

```
FitMLVSBM$init_clustering(
  safeguard = 2 * .Machine$double.eps,
  method = "hierarchical",
  Z = NULL
)
```

Arguments:

`safeguard` Parameter live in a compact [`safeguard`, 1-`safeguard`]

`method` Algorithm used to initiate the clustering, either "spectral", "hierarchical" or "merge_split" (if Z is provided)

`Z` Initial clustering if provided

Method `clear()`: Reset all parameters

Usage:

```
FitMLVSBM$clear()
```

Method `m_step()`: `m_step` Compute the M step of the VEM algorithm

Usage:

```
FitMLVSBM$m_step(safeguard = 1e-06)
```

Arguments:

`safeguard` Parameter live in a compact [`safeguard`, 1-`safeguard`]

Method `ve_step()`: Compute the VE step of the VEM algorithm

Usage:

```
FitMLVSBM$ve_step(threshold = 1e-06, fixPointIter = 10, safeguard = 1e-06)
```

Arguments:

`threshold` The convergence threshold

`fixPointIter` The maximum number of fixed point iterations

`safeguard` Parameter live in a compact [`safeguard`, 1-`safeguard`]

Method `do_vem()`: Launch a Variational EM algorithm

Usage:

```
FitMLVSBM$do_vem(
  init = "hierarchical",
  threshold = 1e-06,
  maxIter = 1000,
  fixPointIter = 100,
  safeguard = 1e-06,
  Z = NULL
)
```

Arguments:

`init` The method for `self$init_clustering`
`threshold` The convergence threshold
`maxIter` The max number of VEM iterations
`fixPointIter` The max number of fixed point iterations for VE step
`safeguard` Parameter live in a compact [`safeguard`, `1-safeguard`]
`Z` Initial clustering if provided

Method `permute_empty_class()`: `permute_empty_class` Put empty blocks numbers at the end

Usage:

```
FitMLVSBM$permute_empty_class()
```

Method `plot()`: Plot of FitMLVSBM objects

Usage:

```
FitMLVSBM$plot(type = c("matrix"), ...)
```

Arguments:

`type` A string for the type of plot, just "matrix" for now

Returns: a `ggplot2` object

Method `show()`: print method

Usage:

```
FitMLVSBM$show(type = "Multilevel Stochastic Block Model")
```

Arguments:

`type` character to tune the displayed name

Method `print()`: print method

Usage:

```
FitMLVSBM$print()
```

Method `clone()`: The objects of this class are cloneable with this method.

Usage:

```
FitMLVSBM$clone(deep = FALSE)
```

Arguments:

`deep` Whether to make a deep clone.

Description

a fitted level of a unilevel network once `$do_vem()` is done

Public fields

`vbound` vector of variational bound for convergence monitoring

Active bindings

`adjacency` Get the adjacency matrix

`mask` Get the mask matrix for dealing with NA

`nb_nodes` Get the number of nodes of the level

`nb_clusters` Get the number of blocks

`distribution` Get the distribution used for the connections

`directed` Get if the level is directed or not

`mixture_parameter` Access the block proportions

`connectivity_parameter` Access the connectivity matrix

`membership` Access the variational parameters

`entropy` Get the entropy of the model

`bound` Get the variational bound of the model

`df_mixture` Get the degree of freedom of the block proportion

`df_connect` Get the degree of freedom of the connection parameters

`connect` Get the number of observed dyads

`ICL` Get the ICL model selection criterion

`penalty` Get the penalty used for computing the ICL

`Z` Access the vector of block membership (clustering)

`X_hat` Get the connection probability matrix

`X_likelihood` adjacency part of the log likelihood

`Z_likelihood` block part of the log likelihood

`likelihood` complete log likelihood

Methods**Public methods:**

- `FitSBM$new()`
- `FitSBM$update_alpha()`
- `FitSBM$update_pi()`
- `FitSBM$init_clustering()`
- `FitSBM$m_step()`
- `FitSBM$ve_step()`
- `FitSBM$do_vem()`
- `FitSBM$permute_empty_class()`
- `FitSBM$clear()`
- `FitSBM$clone()`

Method `new()`: Constructor for FitSBM R6 class

Usage:

```
FitSBM$new(
  Q = 1,
  X = NULL,
  M = NULL,
  directed = FALSE,
  distribution = "bernoulli"
)
```

Arguments:

Q Number of blocks
 X Adjacency matrix
 M Mask matrix
 directed boolean
 distribution string (only "bernoulli")

Returns: A new FitSBM object

Method `update_alpha()`: Update the connection parameter for the M step

Usage:

```
FitSBM$update_alpha(safeguard = 1e-06)
```

Arguments:

safeguard Parameter live in a compact [safeguard, 1-safeguard]

Method `update_pi()`: Update the upper level mixture parameter for the M step

Usage:

```
FitSBM$update_pi(safeguard = 1e-06)
```

Arguments:

safeguard Parameter live in a compact [safeguard, 1-safeguard]

Method `init_clustering()`: `init_clustering` Initial clustering for VEM algorithm

Usage:

```
FitSBM$init_clustering(safeguard = 1e-06, method = "hierarchical", Z = NULL)
```

Arguments:

safeguard Parameter live in a compact [safeguard, 1-safeguard]

method Algorithm used to initiate the clustering, either "spectral", "hierarchical" or "merge_split"
(if Z is provided)

Z Initial clustering if provided

Method `m_step()`: `m_step` Compute the M step of the VEM algorithm

Usage:

```
FitSBM$m_step(safeguard = 1e-06)
```

Arguments:

safeguard Parameter live in a compact [safeguard, 1-safeguard]

Method `ve_step()`: Compute the VE step of the VEM algorithm

Usage:

```
FitSBM$ve_step(threshold = 1e-06, fixPointIter = 100, safeguard = 1e-06)
```

Arguments:

threshold The convergence threshold

fixPointIter The maximum number of fixed point iterations

safeguard Parameter live in a compact [safeguard, 1-safeguard]

Method `do_vem()`: Launch a Variational EM algorithm

Usage:

```
FitSBM$do_vem(
  init = "hierarchical",
  threshold = 1e-06,
  maxIter = 1000,
  fixPointIter = 100,
  safeguard = 1e-06,
  Z = NULL
)
```

Arguments:

init The method for `self$init_clustering`

threshold The convergence threshold

maxIter The max number of VEM iterations

fixPointIter The max number of fixed point iterations for VE step

safeguard Parameter live in a compact [safeguard, 1-safeguard]

Z Initial clustering if provided

Method `permute_empty_class()`: `permute_empty_class` Put empty blocks numbers at the end

Usage:

```
FitSBM$permute_empty_class()
```

Method `clear()`: Reset all parameters

Usage:

`FitSBM$clear()`

Method `clone()`: The objects of this class are cloneable with this method.

Usage:

`FitSBM$clone(deep = FALSE)`

Arguments:

`deep` Whether to make a deep clone.

hierarClust *Perform a Hierarchical Clustering*

Description

Perform a Hierarchical Clustering

Usage

`hierarClust(X, K)`

Arguments

`X` An Adjacency Matrix
`K` the number of wanted clusters

Value

A vector : The clusters labels

merge_clust *Merge a list of clusters*

Description

Merge a list of clusters

Usage

`merge_clust(Z, Q)`

Arguments

`Z` a vector of cluster memberships
`Q` the number of original clusters

Value

A list of $Q(Q-1)/2$ clustering of $Q-1$ clusters

MLVSBM

R6Class for multilevel object

Description

Store all simulation parameters and list of fittedmodels. Methods for global inference and model selection are included.

Active bindings

nb_nodes List of the umber of nodes for each levels
simulation_parameters List of parameters of the MLVSBM
affiliation_matrix Access the affiliation matrix
adjacency_matrix Access the list of adjacency_matrix
memberships Access the list of the clusterings
fittedmodels Get the list of selected fitted FitMLVSBM objects
ICL A summary table of selected fitted models and ICL model selection criterion
ICL_sbm Summary table of ICL by levels
tmp_fittedmodels A list of all fitted FitMLVSBM objects
fittedmodels_sbm A list of selected fitted FitSBM objects of each levels
max_clusters Access the list of maximum model size
min_clusters Access the list of minimum model size
directed Access the list of boolean for levels direction
directed Access the list of the distribution used for each levels

Methods**Public methods:**

- `MLVSBM$estimate_level()`
- `MLVSBM$estimate_sbm_neighbours()`
- `MLVSBM$estimate_sbm_from_neighbours()`
- `MLVSBM$estimate_sbm()`
- `MLVSBM$mceestimate()`
- `MLVSBM$estimate_from_neighbours()`
- `MLVSBM$estimate_neighbours()`
- `MLVSBM$merge_split_membership()`
- `MLVSBM$mc_ms_estimate()`
- `MLVSBM$estimate_one()`

- `MLVSBM$estimate_all_bm()`
- `MLVSBM$new()`
- `MLVSBM$findmodel()`
- `MLVSBM$clearmodels()`
- `MLVSBM$addmodel()`
- `MLVSBM$simulate()`
- `MLVSBM$clone()`

Method `estimate_level()`:*Usage:*

```
MLVSBM$estimate_level(  
  level = "lower",  
  Q_min = 1,  
  Q_max = 10,  
  Z = NULL,  
  init = "hierarchical",  
  depth = 1,  
  nb_cores = NULL  
)
```

Method `estimate_sbm_neighbours()`:*Usage:*

```
MLVSBM$estimate_sbm_neighbours(  
  level = "lower",  
  Q = NULL,  
  Q_min = 1,  
  Q_max = 10,  
  fit = NULL,  
  nb_cores = NULL,  
  init = NULL  
)
```

Method `estimate_sbm_from_neighbours()`:*Usage:*

```
MLVSBM$estimate_sbm_from_neighbours(  
  level = "lower",  
  Q = NULL,  
  fits = NULL,  
  nb_cores = NULL  
)
```

Method `estimate_sbm()`:*Usage:*

```
MLVSBM$estimate_sbm(level = "lower", Q = Q, Z = NULL, init = "hierarchical")
```

Method `mceestimate()`:*Usage:*

```
MLVSBM$mceestimate(Q, Z = NULL, init = "hierarchical", independent = FALSE)
```

Method estimate_from_neighbours():

Usage:

```
MLVSBM$estimate_from_neighbours(  
  Q,  
  models = NULL,  
  independent = FALSE,  
  nb_cores = nb_cores  
)
```

Method estimate_neighbours():

Usage:

```
MLVSBM$estimate_neighbours(  
  Q,  
  fit = NULL,  
  independent = independent,  
  nb_cores = NULL  
)
```

Method merge_split_membership():

Usage:

```
MLVSBM$merge_split_membership(  
  fitted = private$fitted[[length(private$fitted)]]  
)
```

Method mc_ms_estimate():

Usage:

```
MLVSBM$mc_ms_estimate(Z = NA, independent = FALSE, nb_cores = NULL)
```

Method estimate_one():

Usage:

```
MLVSBM$estimate_one(  
  Q,  
  Z = NULL,  
  independent = FALSE,  
  init = "hierarchical",  
  nb_cores = NULL  
)
```

Method estimate_all_bm():

Usage:

```
MLVSBM$estimate_all_bm(  
  Q = NULL,  
  Z = NULL,  
  independent = FALSE,  
  clear = TRUE,  
  nb_cores = NULL  
)
```

Method `new()`: Constructor for R6 class MLVSBM

Usage:

```
MLVSBM$new(
  n = NULL,
  X = NULL,
  A = NULL,
  Z = NULL,
  directed = NULL,
  sim_param = NULL,
  distribution = list("bernoulli", "bernoulli")
)
```

Arguments:

`n` A list of size 2, the number of nodes

`X` A list of 2 adjacency matrices

`A` The affiliation matrix

`Z` A list of 2 vectors, the blocks membership

`directed` A list of 2 booleans

`sim_param` A list of MLVSBM parameters for simulating networks

`distribution` The distributions of the interactions ("bernoulli")

Returns: A MLVSBM object

Method `findmodel()`: Find a fitted model of a given size

Usage:

```
MLVSBM$findmodel(nb_clusters = NA, fit = NA)
```

Arguments:

`nb_clusters` A list of the size of the model

`fit` if `fit = "best"` return the best model according to the ICL

Returns: A FitMLVSBM object

Method `clearmodels()`: delete all fitted models

Usage:

```
MLVSBM$clearmodels()
```

Method `addmodel()`: Added a FitMLVSBM object to the list of fitted model

Usage:

```
MLVSBM$addmodel(fit)
```

Arguments:

`fit` The FitMLVSBM object to be added

Method `simulate()`:

Usage:

```
MLVSBM$simulate()
```

Method `clone()`: The objects of this class are cloneable with this method.

Usage:

```
MLVSBM$clone(deep = FALSE)
```

Arguments:

deep Whether to make a deep clone.

mlvsbm_create_network *Create a MLVSBM object from observed data*

Description

Create a MLVSBM object from observed data

Usage

```
mlvsbm_create_network(  
  X,  
  A,  
  directed = NULL,  
  distribution = list("bernoulli", "bernoulli")  
)
```

Arguments

X	A list of 2 squares binary matrices, the first one being the individual or lower level the second one being the organizational or upper level
A	A matrix the affiliation matrix with individuals in rows and organizations in columns
directed	A list of 2 boolean are the upper and lower level directed or not. Default will check if the matrix are symmetric or not.
distribution	A list for the distribution of X, only "bernoulli" is implemented

Value

An unfitted MLVSBM object corresponding to the multilevel network

Examples

```
ind_adj <- matrix(stats::rbinom(n = 10**2, size = 1, prob = .2),  
                 nrow = 10, ncol = 10)  
org_adj <- matrix(stats::rbinom(n = 10**2, size = 1, prob = .3),  
                 nrow = 10, ncol = 10)  
affiliation <- diag(1, 10)  
my_mlvsbm <- mlvsbm_create_network(X = list(I = ind_adj, O = org_adj),  
                                  directed = list(I = FALSE, O = FALSE),  
                                  A = affiliation)
```

mlvsbm_estimate_network

Infer a multilevel network (MLVSBM object), the original object is modified

Description

The inference use a greedy algorithm to navigate between model size. For a given model size, the inference is done via a variational EM algorithm. The returned model is the one with the highest ICL criterion among all visited models.

By default the algorithm fits a single level SBM for each level, before inferring the multilevel network. This step can be skipped by specifying an initial clustering with the `init_clustering`. Also, a given model size can be force by setting the parameters `nb_clusters` to a given value.

Usage

```
mlvsbm_estimate_network(  
  mlv,  
  nb_clusters = NULL,  
  init_clustering = NULL,  
  nb_cores = NULL,  
  init_method = "hierarchical"  
)
```

Arguments

<code>mlv</code>	A MLVSBM object, the network to be inferred.
<code>nb_clusters</code>	A list of 2 integers, the model size. If left to NULL, the algorithm will navigate freely. Otherwise it will navigate between the specified model size and its neighbors.
<code>init_clustering</code>	A list of 2 vectors of integers of the same length as the number of node of each level. If specified, the algorithm will start from this clustering, then navigate freely.
<code>nb_cores</code>	An integer, the number of cores to use. Default to 1 for Windows and <code>detectCores()/2</code> for Linux and MacOS
<code>init_method</code>	One of "hierarchical" (the default) or "spectral", "spectral" might be more efficient but can lead to some numeric errors. Not used when <code>int_clustering</code> is given.

Value

A FitMLVSBM object, the best inference of the network

Examples

```

my_mlvsbm <- MLVSBM::mlvsbm_simulate_network(
  n = list(I = 10, O = 20), # Number of nodes for the lower level and the upper level
  Q = list(I = 2, O = 2), # Number of blocks for the lower level and the upper level
  pi = c(.3, .7), # Block proportion for the upper level, must sum to one
  gamma = matrix(c(.9, .2, # Block proportion for the lower level,
    .1, .8), # each column must sum to one
    nrow = 2, ncol = 2, byrow = TRUE),
  alpha = list(I = matrix(c(.8, .2,
    .2, .1),
    nrow = 2, ncol = 2, byrow = TRUE), # Connection matrix
  O = matrix(c(.99, .3,
    .3, .1),
    nrow = 2, ncol = 2, byrow = TRUE)), # between blocks
  directed = list(I = FALSE, O = FALSE), # Are the upper and lower level directed or not ?
  affiliation = "preferential") # How the affiliation matrix is generated
fit <- MLVSBM::mlvsbm_estimate_network(mlv = my_mlvsbm, nb_cores = 1)

```

`mlvsbm_log_likelihood` *Compute the complete log likelihood of a multilevel network for a given clustering of the nodes.*

Description

This function is useful to compute the likelihood for clusters obtained by different methods.

Usage

```
mlvsbm_log_likelihood(mlv, clustering)
```

Arguments

<code>mlv</code>	A MLVSBM object, the network data
<code>clustering</code>	A list of 2 vectors of integers of the same length as the number of node of each level.

Value

A numeric, the log likelihood of the multilevel network for the given clustering.

Examples

```

my_mlvsbm <- MLVSBM::mlvsbm_simulate_network(
  n = list(I = 40, O = 20), # Number of nodes for the lower level and the upper level
  Q = list(I = 2, O = 2), # Number of blocks for the lower level and the upper level
  pi = c(.3, .7), # Block proportion for the upper level, must sum to one
  gamma = matrix(c(.9, .2, # Block proportion for the lower level,
    .1, .8), # each column must sum to one
    nrow = 2, ncol = 2, byrow = TRUE),

```

```

alpha = list(I = matrix(c(.8, .2,
                        .2, .1),
                      nrow = 2, ncol = 2, byrow = TRUE), # Connection matrix
            O = matrix(c(.99, .3,
                        .3, .1),
                      nrow = 2, ncol = 2, byrow = TRUE)), # between blocks
directed = list(I = FALSE, O = FALSE), # Are the upper and lower level directed or not ?
affiliation = "preferential" # How the affiliation matrix is generated
mlvsbm_log_likelihood(mlv = my_mlvsbm, clustering = my_mlvsbm$memberships)

```

mlvsbm_simulate_network

Create a simulated multilevel network (MLVSBM object)

Description

Create a simulated multilevel network (MLVSBM object)

Usage

```

mlvsbm_simulate_network(
  n,
  Q,
  pi,
  gamma,
  alpha,
  directed,
  affiliation = "uniform",
  distribution = list("bernoulli", "bernoulli"),
  no_empty_org = FALSE,
  no_isolated_node = FALSE
)

```

Arguments

n	A list of 2 positive integers, the number of individuals and organizations.
Q	A list of 2 positive integers, the number of clusters of individuals and organizations.
pi	A vector of probabilities of length Q_O , the mixture parameter for the organizations.
gamma	A $Q_I \times Q_O$ matrix with each column summing to one, the mixture parameters for the individuals
alpha	A list of 2 matrices, a $Q_I \times Q_I$ matrix giving the connectivity probabilities of the individuals and a $Q_O \times Q_O$ matrix giving the connectivity probabilities of the organizations.
directed	A list of 2 logical. Is the individual level a directed network ? Is the inter-organizational level a directed network?

affiliation	The distribution under which the affiliation matrix is simulated in <code>c("uniform", "preferential")</code> .
distribution	A list for the distribution of X, only "bernoulli" is implemented.
no_empty_org	A logical with FALSE as default, should every organizations have at least one affiliated individual? Needs to have $n_I \geq n_O$.
no_isolated_node	A logical, if TRUE then the network is simulated again until all nodes are connected.

Value

An MLVSBM object, a simulated multilevel network with levels, affiliations and memberships.

Examples

```
my_mlvsbm <- MLVSBM::mlvsbm_simulate_network(
  n = list(I = 10, O = 20), # Number of nodes for the lower level and the upper level
  Q = list(I = 2, O = 2), # Number of blocks for the lower level and the upper level
  pi = c(.3, .7), # Block proportion for the upper level, must sum to one
  gamma = matrix(c(.9, .2, # Block proportion for the lower level,
                  .1, .8), # each column must sum to one
                nrow = 2, ncol = 2, byrow = TRUE),
  alpha = list(I = matrix(c(.8, .2,
                           .2, .1),
                         nrow = 2, ncol = 2, byrow = TRUE), # Connection matrix
              O = matrix(c(.99, .3,
                          .3, .1),
                        nrow = 2, ncol = 2, byrow = TRUE)), # between blocks
  directed = list(I = FALSE, O = FALSE)) # Are the upper and lower level directed
```

plot.FitMLVSBM

Multilevel SBM Plot

Description

basic matrix plot method for a FitMLVSBM object

Usage

```
## S3 method for class 'FitMLVSBM'
plot(x, type = c("matrix"), ...)
```

Arguments

x	an R6 object of class <code>FitMLVSBM</code>
type	A string for the type of plot, just "matrix" for now
...	additional parameters. block ordering with <code>order = c("affiliation", "degree", "natural")</code>

Details

Basic matrix plot method for a FitMLVSBM object

Value

a ggplot2 object

predict.FitMLVSBM *Model Predictions*

Description

Make predictions from an SBM.

Usage

```
## S3 method for class 'FitMLVSBM'
predict(object, ...)
```

Arguments

object an R6 object of class `FitMLVSBM`
... additional parameters for S3 compatibility. Not used

Value

A list with the following entries:

dyads A list of matrix with the probability of each dyads

nodes A list of vectors with the clustering of each nodes

simulate_adjacency *Simulation an adjacency matrix*

Description

Simulation an adjacency matrix

Usage

```
simulate_adjacency(  
  Z,  
  n,  
  alpha,  
  directed,  
  distribution = "bernoulli",  
  no_isolated_node = FALSE  
)
```

Arguments

<code>Z</code>	A vector of integer of size n , the label
<code>n</code>	An integer, the number of rows or columns of the matrix
<code>alpha</code>	A $\max(Z) \times \max(Z)$ matrix, the connectivity parameters
<code>directed</code>	A boolean, Is the network directed or not ?
<code>distribution</code>	The distribution of the indices: only "bernoulli"
<code>no_isolated_node</code>	A boolean, may row and column of adjacency matrices sum to 0

Value

A $n \times n$ adjacency matrix

`simulate_affiliation` *Simulate of matrix of affiliation*

Description

Simulate of matrix of affiliation

Usage

```
simulate_affiliation(n, m, affiliation = "uniform", no_empty_org = FALSE)
```

Arguments

<code>n</code>	An integer, the number of individuals
<code>m</code>	An integer, the number of organizations
<code>affiliation</code>	The type of affiliation between c("uniform", "preferential")
<code>no_empty_org</code>	A Boolean. Force all columns to have at least a 1. Must have $n > m$.

Value

A $n \times m$ affiliation matrix, with a unique 1 on each rows

spcClust	<i>Perform a spectral clustering</i>
----------	--------------------------------------

Description

Perform a spectral clustering

Usage

```
spcClust(X, K)
```

Arguments

X	an Adjacency matrix
K	the number of clusters

Value

A vector : The clusters labels

split_clust	<i>Merge a list of clusters</i>
-------------	---------------------------------

Description

Merge a list of clusters

Usage

```
split_clust(X, Z, Q)
```

Arguments

X	an adjacency matrix
Z	a vector of cluster memberships
Q	The number of maximal clusters

Value

A list of Q clustering of Q+1 clusters

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