Package ‘robber’

September 20, 2021

Title Using Block Model to Estimate the Robustness of Ecological Network

Version 0.2.2

Description Implementation of a variety of methods to compute the robustness of ecological interaction networks with binary interactions as described in <arXiv:1910.10512>. In particular, using the Stochastic Block Model and its bipartite counterpart, the Latent Block Model to put a parametric model on the network, allows the comparison of the robustness of networks differing in species richness and number of interactions. It also deals with networks that are partially sampled and/or with missing values.

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URL https://github.com/Chabert-Liddell/robber

BugReports https://github.com/Chabert-Liddell/robber/issues

Depends R (>= 3.5)

Imports blockmodels (>= 1.1.2), ggplot2, GREMLINS, igraph, magrittr, pammtools, partitions, purrr, stats

Suggests dplyr, forcats, knitr, patchwork, pbmcapply, RColorBrewer, markdown, spelling, testthat, tibble, tidyverse

VignetteBuilder knitr

Encoding UTF-8

Language en-US

LazyData true

RoxygenNote 7.1.2

NeedsCompilation no

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Repository CRAN

Date/Publication 2021-09-20 14:50:09 UTC
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| auc_robustness_lbm | Fast computation of the LBM robustness statistic for uniform extinctions |

Description

This function provides much faster computation than robustness_lbm for the case where ext_seq = "uniform", when just the AUC statistic is needed and the robustness function is not needed. This is particularly useful if nr gets large.

Usage

auc_robustness_lbm(con, pi, rho, nr, nc)

Arguments

- con: A matrix, the connectivity parameter
- pi: A vector of length nrow(con), the proportion of row blocks
- rho: A vector of length ncol(con), the proportion of column blocks
- nr: An integer, the number of row (primary) species
- nc: An integer, the number of column (secondary) species

Value

The AUC of the LBM Robustness function for uniform primary extinctions sequences.
compare_robustness

Examples

```r
con <- matrix(c(.5,.3,.3,.1), 2, 2)
pi <- c(.25,.75)
rho <- c(1/3, 2/3)
nr <- 50
nc <- 30
auc_robustness_lbm(con, pi, rho, nr, nc)
```

**Description**

This function is designed to be used with a list of parameters as the one given by the function `get_lbm_param()`. The default parameters are designed such that the return value for a uniform extinction sequence is bounded above by 0.5.

**Usage**

```r
compare_robustness(
  list_param,
  dens = 0.0156,
  new_nr = 100,
  new_nc = 100,
  ext_seq = "uniform"
)
```

**Arguments**

- `list_param`: A list of list of LBM parameters, each list must contain at list `con`, `pi` and `rho` entries, such as the one returned by the function `get_lbm_param()`. The size of the network is optional.
- `dens`: The density (connectance) used to compare the network. The `con` parameters of `list_param` will be normalized so that the average probability of interaction in the network is equal to `dens`. Default to 0.0156.
- `new_nr`: The number of rows of the normalized networks. Default to 100.
- `new_nc`: The number of columns of the normalized networks. Default to 100.
- `ext_seq`: The distribution of the primary extinction sequence used to compare the networks. One of `c("uniform", "decreasing", "increasing")`. Default to "uniform".

**Value**

A list of the same length as `list_param`, the robustness (AUC) after normalization for all set of parameters.
get_lbm_param

Estimate the parameters of a LBM

Description

Estimate the parameters of a LBM

Usage

get_lbm_param(A, model_size = NULL, ...)

Arguments

A
model_size
...  
A matrix, the incident matrix of the network
A vector of size 2, the size of the model parameters. If NULL, the default, model will be chosen as the one with the highest ICL criterion among all fitted models during the inference.
Additional arguments pass to the inference function of package GREMLIN if there are missing value and blockmodels if none.

Value

A list of the LBM parameters. The list is set

Examples

# When the size of the model is unknown:
data("seeddispersal")
param <- get_lbm_param(seeddispersal, ncores = 1L)
do.call(auc_robustness_lbm, param)

# For a fixed number of blocks (useful for comparison)
param <- get_lbm_param(seeddispersal, model_size = c(1, 3), ncores = 1L)
do.call(auc_robustness_lbm, param)

# For data with missing observations
A <- seeddispersal
hostparasite

A binarized host-parasite network

Description

Locality: Tomsk-Tumen

Usage

hostparasite

Format

A matrix of binary entries with 27 rows and 26 columns

Source

http://www.web-of-life.es/

plot.robber

Plot function pour robber class

Description

Plot function pour robber class

Usage

## S3 method for class 'robber'
plot(x, type = "function", add = FALSE, ...)

Arguments

x An object of class robber
type Type of the plot, just function
add Whether it should be added to a previous ggplot
... lty, col, ... to be passed to geom_step() in ggplot2
Value

A ggplot2 object

Examples

data("hostparasite", package = "robber")
rob <- robustness_emp(hostparasite)
plot(rob)

---

pollination  A plant-pollinator network

Description


Usage

pollination

Format

A matrix of binary entries with 17 rows and 51 columns

Source

http://www.web-of-life.es/

---

robustness_emp  Empirical Robustness of a Network

Description

Compute the robustness of an ecological network by averaging over a great number of randomly generated primary extinctions sequences

Usage

robustness_emp(  
  A,
  ext_seq = "uniform",
  nb_iter = 300,
  net_type = "bipartite",
  method = "ordered",
  power = 1,
  block = NULL
)
robustness_emp

Arguments

A  A binary incident matrix

ext_seq  A string, the rule for the primary extinctions sequences, one of "uniform", the
default for uniform extinctions sequences, "decreasing" and "increasing" for
primary extinctions sequences by increasing and decreasing degree order
sequence on the row species

nb_iter  An integer, the number of generated sequences. Default to 300.

net_type  A string, the type of network. For now, only "bipartite" is available.

method  A string used when ext_seq = "decreasing" or ext_seq = "increasing":

  • "linear" (default) set an extinction probability for each row species that is
    linear in its degree. It is a shortcut for method = "power" and power = 1.
  • "ordered" strictly follow the degree order, i.e. the most connected species
    will always goes last for ext_seq = "increasing" or first for ext_seq =
    "decreasing".

power  An integer (default to 1). Only used when method = "power", the power to
which the degree are elevated to compute the primary extinctions sequences for
"increasing" and "decreasing" ext_seq. For 1, it is equivalent to method =
"linear", for 0, it is the same as ext_seq = "uniform". The largest the number
the closest it will be to method = "ordered".

block  A vector of row species memberships for method = "block"

Value

A list (robber object) with the following fields:

  • $model, $ext_seq, $method, power
  • $fun is the robustness function, a vector of size (nrow(A) +1)
  • $sd Standard deviation of the secondary extinctions
  • $auc the area under the curve of the robustness function

Examples

A <- matrix(c(1, 0, 0,
              0, 1, 0,
              0, 0, 1,
              1, 1, 1), nrow = 4, ncol = 3, byrow = TRUE)
my_rob <- robustness_emp(A, ext_seq = "increasing")
my_rob$fun
my_rob$auc
robustness_lbm  Robustness for LBM

Description

Compute the robustness for a given set of Latent Block Model parameters.

Usage

```r
robustness_lbm(
  con = NULL,
  pi = NULL,
  rho = NULL,
  nr = NULL,
  nc = NULL,
  ext_seq = "uniform",
  method = "exact",
  approx_threshold = 10000,
  net = NULL,
  ...
)
```

Arguments

- **con**: A matrix, the connectivity parameter
- **pi**: A vector of length `nrow(con)`, the proportion of row blocks
- **rho**: A vector of length `ncol(con)`, the proportion of column blocks
- **nr**: An integer, the number of row (primary) species
- **nc**: An integer, the number of column (secondary) species
- **ext_seq**: A string, the rule for the primary extinctions sequences, one of:
  - "uniform", the default for uniform extinctions sequences,
  - "decreasing" and "increasing" for primary extinctions sequences by increasing and decreasing row blocks connection probability,
  - "natural" primary extinctions sequences with the block ordering given in the function parameter.
  - "blocks" primary extinctions sequences for all blocks permutation.
- **method**: A string, the method used to compute the robustness by block. One of:
  - "exact" (the default), average on all possible networks
  - "no_empty_block" restricts the networks space to the ones with no empty block of primary species
  - "expected" restricts the networks space to the ones that have approximately the same number of species per block than the expected one. Do not do anything for `ext_seq = "uniform"`. 
approx_threshold
A numeric, the maximum size of the possible block partition allowed for exact robustness by block calculation. Higher threshold gives more precise results at the cost of computation times and possibly memory problem. Do not do anything for ext_seq = "uniform". When the threshold is exceeded the robustness is computed by a Monte Carlo approximation with approx_threshold number of samples.

net
A network, if given, the function will fit a LBM to obtain the parameters of the network and then compute the robustness.

Value
A list and a robber type object:

- $fun the robustness function, a vector of size nr +1
- $auc the area under the curve of the robustness function
- $block a vector of size length(pi), the block ordering for primary extinctions sequence by blocks. NULL if ext_seq = "uniform".
- $model, $method, $ext_seq, $param.

Examples
con <- matrix(c(.5,.3,.3,.1), 2, 2)
p <- c(.25,.75)
rho <- c(1/3, 2/3)
nr <- 50
nc <- 30
my_rob <- robustness_lbm(con, pi, rho, nr, nc, ext_seq = "natural")
my_rob$fun
my_rob$auc
# A easier alternative way, if you don't know the parameters of the network:
data(hostparasite)
(robustness_lbm(net = hostparasite, ncores = 1L))

seddispersal A seed-dispersal network

Description

Usage
seddispersal
simulate_lbm

Simulate a bipartite interaction network using block model

Description

Simulate a bipartite interaction network using block model

Usage

simulate_lbm(con, pi, rho, nr, nc, method = "gnp")

Arguments

con A matrix, the connectivity between blocks. If method = "gnp" then each entry is the probability of interactions given 2 blocks. If method = "gnm" then each entry is the number of interactions between 2 blocks.

pi A vector of the same length as nrow(con), the block mixture for the row species. If method = "gnp", then pi is a probability vector, if method = "gnm", then pi is the number of species per block, must sum to nr.

rho A vector of the same length as ncol(con), the block mixture for the column species. If method = "gnm", then rho is a probability vector, if method = "gnm", then rho is the number of species per block, must sum to nc.

nr The number of row Species

nc The number of column Species

method One of "gnp" (the default) where the blocks size and number of interactions is random and "gnm" where the blocks size and number of interactions are fixed.

Value

A list of 3 elements:

- A The incident matrix of size nr x nc
- Z A vector of length nr, the block assignment of the row species
- W A vector of length nc, the block assignment of the column species
Examples

# For a random number of interactions and blocks sizes
con <- matrix(c(0.5, 0.3, 0.3, 0.1), 2, 2)
pi <- c(0.25, 0.75)
rho <- c(1/3, 2/3)
nr <- 50
nc <- 30
simulate_lbm(con, pi, rho, nr, nc, method = "gnp")

# For a fixed number of interactions and blocks sizes
con <- matrix(c(50, 120, 60, 80), 2, 2)
pi <- c(10, 40)
rho <- c(10, 20)
nr <- 50
nc <- 30
simulate_lbm(con, pi, rho, nr, nc, method = "gnm")

var_auc_unif_lbm

Compute the Conditional variance of the AUC of the LBM Robustness

Description

Compute the Conditional variance of the AUC of the LBM Robustness

Usage

var_auc_unif_lbm(con, pi, rho, nr, nc)

Arguments

con A matrix, the connectivity parameter
pi A vector of length nrow(con), the proportion of row blocks
rho A vector of length ncol(con), the proportion of column blocks
nr An integer, the number of row (primary) species
nc An integer, the number of column (secondary) species

Value

A numeric, the variance

Examples

con <- matrix(c(0.5, 0.3, 0.3, 0.1), 2, 2)
pi <- c(0.25, 0.75)
rho <- c(1/3, 2/3)
nr <- 50
nc <- 30
var_auc_unif_lbm(con, pi, rho, nr, nc)
Compute the Conditional variance of the LBM Robustness term by term

### Description
Compute the Conditional variance of the LBM Robustness term by term

### Usage
```
var_fun_unif_lbm(con, pi, rho, nr, nc)
```

### Arguments
- **con**: A matrix, the connectivity parameter
- **pi**: A vector of length `nrow(con)`, the proportion of row blocks
- **rho**: A vector of length `ncol(con)`, the proportion of column blocks
- **nr**: An integer, the number of row (primary) species
- **nc**: An integer, the number of column (secondary) species

### Value
A vector, the variance after m extinctions

### Examples
```
con <- matrix(c(.5,.3,.3,.1), 2, 2)
p <- c(.25,.75)
rho <- c(1/3, 2/3)
nr <- 50
nc <- 30
var_fun_unif_lbm(con, pi, rho, nr, nc)
```
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