Package ‘statGraph’

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Type Package
Title Statistical Methods for Graphs
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Imports igraph, MASS, rARPACK, cluster, foreach, parallel, doParallel
Description Contains statistical methods to analyze graphs, such as graph
      parameter estimation, model selection based on the GIC (Graph Information
      Criterion), statistical tests to discriminate two or more populations of
      graphs (ANOGVA - Analysis of Graph Variability), correlation between
      graphs, and clustering of graphs.
      References: Takahashi et al. (2012) <doi:10.1371/journal.pone.0049949>,
      Futija et al. (2017) <doi:10.3389/fnins.2017.00066>,
      Fujita et al. (2017) <doi:10.1016/j.csda.2016.11.016>,
      Tang et al. (2017) <doi:10.3150/15-BEJ789>,
      Ghoshdastidar et al. (2017) <arXiv:1705.06168>,
      Ghoshdastidar et al. (2017) <arXiv:1707.00833>,
      Cerqueira et al. (2017) <doi:10.1109/TNSE.2017.2674026>,
      Fraiman and Fraiman (2018) <doi:10.1038/s41598-018-23152-5>,
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Description


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Details

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See Also

Useful links:

- https://www.ime.usp.br/~fujita/software.html

anogva

ANOGVA Analysis Of Graph Variability

Description

anogva statistically tests whether two or more sets of graphs are generated by the same random graph model. It is a generalization of the 'takahashi.test' function.

Usage

anogva(G, labels, maxBoot = 1000, bandwidth = "Silverman")

Arguments

G a list of undirected graphs (igraph type) or their adjacency matrices. The adjacency matrix of an unweighted graph contains only 0s and 1s, while the weighted graph may have nonnegative real values that correspond to the weights of the edges.

labels an array of integers indicating the labels of each graph.
maxBoot: integer indicating the number of bootstrap resamplings.
bandwidth: string showing which criterion is used to choose the bandwidth during the spectral density estimation. Choose between the following criteria: "Silverman" (default), "Sturges", "bcv", "ucv" and "SJ". "bcv" is an abbreviation of biased cross-validation, while "ucv" means unbiased cross-validation. "SJ" implements the methods of Sheather & Jones (1991) to select the bandwidth using pilot estimation of derivatives.

Value

A list containing:

- statistic: the statistic of the test.
- p.value: the p-value of the test.

References


Examples

```r
set.seed(1)
g1 <- g2 <- g3 <- list()
for (i in 1:20) {
  g1[[i]] <- igraph::sample_gnp(50, 0.50)
g2[[i]] <- igraph::sample_gnp(50, 0.50)
g3[[i]] <- igraph::sample_gnp(50, 0.52)
}
G <- c(g1, g2, g3)
label <- c(rep(1,20),rep(2,20),rep(3,20))
result <- anogva(G, label, maxBoot=50)
result
```
cerqueira.test  Andressa Cerqueira, Daniel Fraiman, Claudia D. Vargas and Florencia Leonardi non-parametric test of hypotheses to verify if two samples of random graphs were originated from the same probability distribution.

Description

Given two identically independently distributed (idd) samples of graphs G1 and G2, the test verifies if they have the same distribution by calculating the mean distance D from G1 to G2. The test rejects the null hypothesis if D is greater than the (1-alpha)-quantile of the distribution of the test under the null hypothesis.

Usage

cerqueira.test(G1, G2, maxBoot = 300)

Arguments

G1 the first iid sample of graphs to be compared. Must be a list of igraph objects.
G2 the second iid sample of graphs to be compared. Must be a list of igraph objects.
maxBoot integer indicating the number of bootstrap resamples (default is 300).

Value

A list containing:

W the value of the test.
p.value the p-value of the test.

References


Examples

```r
## Not run:
set.seed(42)

## test under H0
G1 <- G2 <- list()
for(i in 1:10){
  G1[[i]] <- igraph::sample_gnp(50,0.5)
  G2[[i]] <- igraph::sample_gnp(50,0.5)
}
k1 <- cerqueira.test(G1, G2)
k1
```
fast.eigenvalue.probability

Degree-based eigenvalue probability

Description

fast.eigenvalue.probability returns the probability of an eigenvalue given the degree and excess degree probability.

Usage

fast.eigenvalue.probability(deg_prob, q_prob, all_k, z, n_iter = 5000)

Arguments

deg_prob The degree probability of the graph.
q_prob The excess degree probability of the graph.
all_k List of sorted unique degrees greater than 1 of the graph.
z Complex number whose real part is the eigenvalue whose probability we want to obtain, the imaginary part is a small value (e.g., 1e-3).
n_iter The maximum number of iterations to perform.

Value

A complex number whose imaginary part absolute value corresponds to the probability of the given eigenvalue.

References

Examples

```r
set.seed(42)
G <- igraph::sample_smallworld(dim = 1, size = 10, nei = 2, p = 0.2)

# Obtain the degree distribution
deg_prob <- c(igraph::degree_distribution(graph = G, mode = "all"), 0.0)
k_deg <- seq(1, length(deg_prob)) - 1

c <- sum(k_deg * deg_prob)
q_prob <- c()
for(k in 0:(length(deg_prob) - 1)){
aux_q <- (k + 1) * deg_prob[k + 1]/c
q_prob <- c(q_prob, aux_q)
}

# Obtain the sorted unique degrees greater than 1
all_k <- c(1: length(q_prob))
valid_idx <- q_prob != 0
q_prob <- q_prob[valid_idx]
all_k <- all_k[valid_idx]

# Obtain the probability of the eigenvalue 0
z <- 0 + 0.01*1i
eigenval_prob <- -Im(fast.eigenvalue.probability(deg_prob, q_prob, all_k, z))
eigenval_prob
```

fast.graph.param.estimator

Degree-based graph parameter estimator

Description

fast.graph.param.estimator estimates the parameter of the complex network model using the degree-based spectral density and ternary search.

Usage

```r
fast.graph.param.estimator(
  G,
  model,
  lo = NULL,
  hi = NULL,
  eps = 0.001,
  from = NULL,
  to = NULL,
  npoints = 2000,
  numCores = 1
)
```
Arguments

G
The undirected unweighted graph (igraph type).

model
Either a string or a function:
A string that indicates one of the following models: "ER" (Erdos-Renyi random graph model), "GRG" (geometric random graph model), "WS" (Watts-Strogatz model), and "BA" (Barabasi-Albert model).
A function that returns a Graph generated by a graph model. It must contain two arguments: the first one corresponds to the graph size and the second to the parameter of the model.

lo
Smallest parameter value that the graph model can take.
If “model” is an string, then the default value of 0 is used for the predefined models ("ER", "GRG", "WS", and "BA").

hi
Largest parameter value that the graph model can take.
If “model” is an string, then the default values are used for the predefined models 1 for "ER", sqrt(2) for "GRG", 1 for "WS", and 5 for "BA").

eps
Desired precision of the parameter estimate.

from
Lower end of the interval that contain the eigenvalues to generate the degree-based spectral densities. The smallest eigenvalue of the adjacency matrix corresponding to “graph” is used if the value is not given.

to
Upper end of the interval that contain the eigenvalues to generate the degree-based spectral densities. The largest eigenvalue of the adjacency matrix corresponding to “graph” is used if the value is not given.

npoints
Number of points to discretize the interval <from,to>.

numCores
Number of cores to use for parallelization.

Value

Returns a list containing:

param
The degree-based parameter estimate. For the "ER", "GRG", "WS", and "BA" models, the parameter corresponds to the probability to connect a pair of vertices, the radius used to construct the geometric graph in a unit square, the probability to reconnect a vertex, and the scaling exponent respectively.

L1_dist
The L1 distance between the observed graph and the graph model with the estimated value.

Examples

set.seed(42)

### Example giving only the name of the model to use
G <- igraph::sample_smallworld(dim = 1, size = 15, nei = 2, p = 0.2)

# Obtain the parameter of the WS model
estimated.parameter1 <- fast.graph.param.estimator(G, "WS", lo = 0.1, hi = 0.5,
eps = 1e-1, npoints = 10,
estimated.parameter1

### Example giving a function instead of a model

```r
# Defining the model to use
G <- igraph::sample_smallworld(dim = 1, size = 5000, nei = 2, p = 0.2)
K <- as.integer(igraph::ecount(G)/igraph::vcount(G))
fun_WS <- function(n, param, nei = K){
  return (igraph::sample_smallworld(dim = 1, size = n, nei = nei, p = param))
}

# Obtain the parameter of the WS model
estimated.parameter2 <- fast.graph.param.estimator(G, fun_WS, lo = 0.0, hi = 1.0,
                                                  npoints = 100, numCores = 2)
```

## Not run:

### Example giving a function instead of a model

```r
# Defining the model to use
G <- igraph::sample_smallworld(dim = 1, size = 5000, nei = 2, p = 0.2)
K <- as.integer(igraph::ecount(G)/igraph::vcount(G))
fun_WS <- function(n, param, nei = K){
  return (igraph::sample_smallworld(dim = 1, size = n, nei = nei, p = param))
}

# Obtain the parameter of the WS model
estimated.parameter2 <- fast.graph.param.estimator(G, fun_WS, lo = 0.0, hi = 1.0,
                                                  npoints = 100, numCores = 2)
```

---

**Description**

`fast.spectral.density` returns the degree-based spectral density in the interval `<from`, `to>` by using `npoints` discretization points.

**Usage**

`fast.spectral.density(G, from = NULL, to = NULL, npoints = 2000, numCores = 1)`

**Arguments**

- **G**: The undirected unweighted graph (igraph type) whose spectral density we want to obtain.
- **from**: Lower end of the interval that contain the eigenvalues or smallest eigenvalue of the adjacency matrix of the graph. The smallest eigenvalue is used if the value is not given.
- **to**: Upper end of the interval that contain the eigenvalues or largest eigenvalue of the adjacency matrix of the graph. The largest eigenvalue is used if the value is not given.
- **npoints**: Number of discretization points of the interval `<from, to>`.
- **numCores**: Number of cores to use for parallelization.

**Value**

Returns the degree-based spectral density of the graph in the
References


Examples

```
set.seed(42)
G <- igraph::sample_smallworld(dim = 1, size = 100, nei = 2, p = 0.2)

# Obtain the degree-based spectral density
density <- fast.spectral.density(G = G, npoints = 80, numCores = 1)
density
```

fraiman.test Daniel Fraiman and Ricardo Fraiman test for network differences between groups with an analysis of variance test (ANOVA).

Description

Given a list of graphs, the test verifies if all the subpopulations have the same mean network, under the alternative that at least one subpopulation has a different mean network.

Usage

```
fraiman.test(G, maxBoot = 300)
```

Arguments

- `G` the undirected graphs to be compared. Must be a list of lists of igraph objects or a list of lists of adjacency matrices.
- `maxBoot` integer indicating the number of bootstrap resamples (default is 300).

Value

A list containing:

- `T` the value of the test.
- `p.value` the p-value of the test.

References

Examples

```r
## Not run:
set.seed(42)

## test under H0
a <- b <- G <- list()
for(i in 1:10){
a[[i]] <- igraph::sample_gnp(50, 0.5)
b[[i]] <- igraph::sample_gnp(50, 0.5)
}
G <- list(a, b)
k1 <- fraiman.test(G)
k1

## test under H1
a <- b <- G <- list()
for(i in 1:10){
a[[i]] <- igraph::sample_gnp(50, 0.5)
b[[i]] <- igraph::sample_gnp(50, 0.6)
}
G <- list(a, b)
k2 <- fraiman.test(G)
k2

## End(Not run)
```

ghoshdastidar.test

**Ghoshdastidar hypothesis testing for large random graphs.**

Description

Given two lists of graphs generated by the inhomogeneous random graph model, *ghoshdastidar.test* tests if they were generated by the same parameters.

Usage

```r
ghoshdastidar.test(G1, G2, maxBoot = 300, two.sample = FALSE)
```

Arguments

- **G1**
  - the first list of undirected graphs to be compared. Must be a list of matrices or igraph objects.
- **G2**
  - the second list of undirected graphs to be compared. Must be a list of matrices or igraph objects.
- **maxBoot**
  - integer indicating the number of bootstrap resamples (default is 300).
- **two.sample**
  - logical. If TRUE the sets contain only one graph each. If FALSE the sets contain more than one graph each (default is FALSE).
ghoshdastidar.test

Value

A list containing:

- `T`      the value of the test.
- `p.value` the p-value of the test (only returned when the parameter `two.sample` is FALSE).

References


Examples

```r
## Not run:
set.seed(42)

## test for sets with more than one graph each under H0
G1 <- G2 <- list()
for(i in 1:10){
  G1[[i]] <- as.matrix(igraph::get.adjacency(igraph::sample_gnp(50, 0.6)))
  G2[[i]] <- as.matrix(igraph::get.adjacency(igraph::sample_gnp(50, 0.6)))
}
D1 <- ghoshdastidar.test(G1, G2)
D1

## test for sets with more than one graph each under H1
G1 <- G2 <- list()
for(i in 1:10){
  G1[[i]] <- as.matrix(igraph::get.adjacency(igraph::sample_gnp(50, 0.6)))
  G2[[i]] <- as.matrix(igraph::get.adjacency(igraph::sample_gnp(50, 0.7)))
}
D2 <- ghoshdastidar.test(G1, G2)
D2

## test for sets with only one graph each under H0
G1 <- G2 <- list()
G1[[1]] <- igraph::sample_gnp(300, 0.6)
G2[[1]] <- igraph::sample_gnp(300, 0.6)
D3 <- ghoshdastidar.test(G1, G2, two.sample= TRUE)
D3

## test for sets with only one graph each under H1
G1 <- G2 <- list()
G1[[1]] <- igraph::sample_gnp(300, 0.6)
G2[[1]] <- igraph::sample_gnp(300, 0.7)
D4 <- ghoshdastidar.test(G1, G2, two.sample= TRUE)
D4

## End(Not run)
```
**GIC**

*Graph Information Criterion (GIC)*

**Description**

GIC returns the Kullback-Leibler divergence or L2 distance between an undirected graph and a given graph model.

**Usage**

```r
GIC(G, model, p = NULL, bandwidth = "Silverman", eigenvalues = NULL, dist = "KL")
```

**Arguments**

- **G**: the undirected graph (igraph type) or its adjacency matrix. The adjacency matrix of an unweighted graph contains only 0s and 1s, while the weighted graph may have nonnegative real values that correspond to the weights of the edges.

- **model**: either a list, a string, a function or a matrix describing a graph model:
  - A list that represents the spectral density of a model. It contains the components "x" and "y", which are numeric vectors of the same size. The "x" component contains the points at which the density was computed and the "y" component contains the observed density.
  - A string that indicates one of the following models: "ER" (Erdos-Renyi random graph), "GRG" (geometric random graph), "KR" (k regular random graph), "WS" (Watts-Strogatz model), and "BA" (Barabasi-Albert model). When the argument 'model' is a string, the user must also provides the model parameter by using the argument 'p'.
  - A function that returns a graph (represented by its adjacency matrix) generated by a graph model. It must contain two arguments: the first one corresponds to the graph size and the second to the parameter of the model. The model parameter will be provided by the argument 'p' of the 'GIC' function.
  - A matrix containing the spectrum of the model. Each column contains the eigenvalues of a graph generated by the model. To estimate the spectral density of the model, the method will estimate the density of the values of each column, and then will take the average density.
the model parameter. The user must provide a valid parameter if the argument 'model' is a string or a function. For the predefined models ("ER", "GRG", "KR", "WS", and "BA"), the parameter the probability to connect a pair of vertices, for the "ER" model (Erdos-Renyi random graph);
the radius used to construct the geometric graph in a unit square, for the "GRG" model (geometric random graph);
the degree 'k' of a regular graph, for the "KR" model (k regular random graph);
the probability to reconnect a vertex, for the "WS" model (Watts-Strogatz model);
and the scaling exponent, for the "BA" model (Barabasi-Albert model).

bandwidth string showing which criterion is used to choose the bandwidth during the spectral density estimation. Choose between the following criteria: "Silverman" (default), "Sturges", "bcv", "ucv" and "SJ". "bcv" is an abbreviation of biased cross-validation, while "ucv" means unbiased cross-validation. "SJ" implements the methods of Sheather & Jones (1991) to select the bandwidth using pilot estimation of derivatives.

eigenvalues optional parameter. It contains the eigenvalues of matrix G. Then, it can be used when the eigenvalues of G were previously computed. If no value is passed, then the eigenvalues of G will be computed by 'GIC'.

dist string indicating if you want to use the "KL" (default) or "L2" distances. "KL" means Kullback-Leibler divergence.

Value
A real number corresponding to the Kullback-Leibler divergence or L2 distance between the observed graph and the graph model.

References


Examples
set.seed(1)
G <- as.matrix(igraph::get.adjacency(igraph::sample_gnp(n=50, p=0.5)))

# Using a string to indicate the graph model
result1 <- GIC(G, "ER", 0.5)
result1

# Using a function to describe the graph model
# Erdos-Renyi graph
The function `graph.acf` computes estimates of the autocorrelation function for graphs.

```r
model <- function(n, p) {
  return(as.matrix(igraph::get.adjacency(igraph::sample_gnp(n, p))))
}
result2 <- GIC(G, model, 0.5)
result2
```

**Description**

The function `graph.acf` computes estimates of the autocorrelation function for graphs.

**Usage**

```r
graph.acf(G, plot = TRUE)
```

**Arguments**

- **G**: a list of undirected graphs (igraph type) or their adjacency matrices. The adjacency matrix of an unweighted graph contains only 0s and 1s, while the weighted graph may have nonnegative real values that correspond to the weights of the edges.
- **plot**: logical. If TRUE (default) the `graph.acf` is plotted.

**Value**

An object of class `acf`.

**References**


**Examples**

```r
set.seed(1)
G <- list()
p <- array(0, 100)
p[1:3] <- rnorm(3)
for (t in 4:100) {
  p[t] <- 0.5*p[t-3] + rnorm(1)
}
ma <- max(p)
mi <- min(p)
p <- (p - mi)/(ma-mi)
for (t in 1:100) {
  G[[t]] <- igraph::sample_gnp(100, p[t])
}
```
Clustering Expectation-Maximization for Graphs (graph.cem)

Description

graph.cem clusters graphs following an expectation-maximization algorithm based on the Kullback-Leibler divergence between the spectral densities of the graph and of the random graph model.

Usage

```r
graph.cem(g, model, k, max_iter = 10, ncores = 1, bandwidth = "Sturges")
```

Arguments

g an list containing the graphs or their adjacency matrices to be clustered.

model a string that indicates one of the following random graph models: "ER" (Erdos-Renyi random graph), "GRG" (geometric random graph), "KR" (k regular graph), "WS" (Watts-Strogatz model), and "BA" (Barabasi-Albert model).

k an integer specifying the number of clusters.

max_iter the maximum number of expectation-maximization steps to execute.

ncores the number of cores to be used for the parallel processing. The default value is 1.

bandwidth string showing which criterion is used to choose the bandwidth during the spectral density estimation. Choose between the following criteria: "Silverman" (default), "Sturges", "bcv", "ucv" and "SJ". "bcv" is an abbreviation of biased cross-validation, while "ucv" means unbiased cross-validation. "SJ" implements the methods of Sheather & Jones (1991) to select the bandwidth using pilot estimation of derivatives.

Value

a list containing three fields: labels a vector of the same length of g containing the clusterization labels; a vector containing the estimated parameters for the groups. It has the length equals to k;

References


Tests for Association / Correlation Between Paired Samples of Graphs

**Examples**

```r
set.seed(42)
g <- list()
for(i in 1:2){
g[[i]] <- igraph::sample_gnp(n=10, p=0.5)
}
for(i in 3:4){
g[[i]] <- igraph::sample_gnp(n=10, p=1)
}
res <- graph.cem(g, model="ER", k=2, max_iter=1, ncores=1)
res
```

**Description**

`graph.cor.test` tests for association between paired samples of graphs, using Spearman’s rho correlation coefficient.

**Usage**

```r
graph.cor.test(G1, G2)
```

**Arguments**

- `G1`: a list of undirected graphs (igraph type) or their adjacency matrices. The adjacency matrix of an unweighted graph contains only 0s and 1s, while the weighted graph may have nonnegative real values that correspond to the weights of the edges.
- `G2`: a list of undirected graphs (igraph type) or their adjacency matrices. The adjacency matrix of an unweighted graph contains only 0s and 1s, while the weighted graph may have nonnegative real values that correspond to the weights of the edges.

**Value**

- `statistic`: the value of the test statistic.
- `p.value`: the p-value of the test.
- `estimate`: the estimated measure of association 'rho'.

**References**

Examples

```r
set.seed(1)
G1 <- G2 <- list()
p <- MASS::mvrnorm(50, mu=c(0,0), Sigma=matrix(c(1, 0.5, 0.5, 1), 2, 2))

ma <- max(p)
mi <- min(p)
p[,1] <- (p[,1] - mi)/(ma - mi)
p[,2] <- (p[,2] - mi)/(ma - mi)

for (i in 1:50) {
  G1[[i]] <- igraph::sample_gnp(50, p[i,1])
  G2[[i]] <- igraph::sample_gnp(50, p[i,2])
}
graph.cor.test(G1, G2)
```

---

**graph.entropy**

**Graph spectral entropy**

### Description

`graph.entropy` returns the spectral entropy of an undirected graph.

### Usage

```r
graph.entropy(G = NULL, bandwidth = "Silverman", eigenvalues = NULL)
```

### Arguments

- **G**
  - the undirected graph (igraph type) or its adjacency matrix. The adjacency matrix of an unweighted graph contains only 0s and 1s, while the weighted graph may have nonnegative real values that correspond to the weights of the edges.

- **bandwidth**
  - string showing which criterion is used to choose the bandwidth during the spectral density estimation. Choose between the following criteria: "Silverman" (default), "Sturges", "bcv", "ucv" and "SJ". "bcv" is an abbreviation of biased cross-validation, while "ucv" means unbiased cross-validation. "SJ" implements the methods of Sheather & Jones (1991) to select the bandwidth using pilot estimation of derivatives.

- **eigenvalues**
  - optional parameter. It contains the eigenvalues of matrix G. Then, if the eigenvalues of matrix G have already been computed, this parameter can be used instead of G. If no value is passed, then the eigenvalues of G will be computed by `graph.entropy`.

### Value

- a real number corresponding to the graph spectral entropy.
**References**


**Examples**

```r
G <- igraph::sample_gnp(n=100, p=0.5)
entropy <- graph.entropy(G)
entropy
```

---

**graph.hclust**

_Hierarchical cluster analysis on a list of graphs._

**Description**

Given a list of graphs, `graph.hclust` builds a hierarchy of clusters according to the Jensen-Shannon divergence between graphs.

**Usage**

```r
graph.hclust(G, k, method = "complete", bandwidth = "Silverman")
```

**Arguments**

- `G` a list of undirected graphs (igraph type) or their adjacency matrices. The adjacency matrix of an unweighted graph contains only 0s and 1s, while the weighted graph may have nonnegative real values that correspond to the weights of the edges.
- `k` the number of clusters.
- `method` the agglomeration method to be used. This should be (an unambiguous abbreviation of) one of "ward.D", "ward.D2", "single", "complete", "average" (= UPGMA), "mcquitty" (= WPGMA), "median" (= WPGMC) or "centroid" (= UPGMC).
- `bandwidth` string showing which criterion is used to choose the bandwidth during the spectral density estimation. Choose between the following criteria: "Silverman" (default), "Sturges", "bcv", "ucv" and "SJ". "bcv" is an abbreviation of biased cross-validation, while "ucv" means unbiased cross-validation. "SJ" implements the methods of Sheather & Jones (1991) to select the bandwidth using pilot estimation of derivatives.
**Value**

A list containing:

- `hclust`: an object of class *hclust* which describes the tree produced by the clustering process.
- `cluster`: the clustering labels for each graph.

**References**


**Examples**

```r
set.seed(1)
G <- list()
for (i in 1:5) {
  G[[i]] <- igraph::sample_gnp(50, 0.5)
}
for (i in 6:10) {
  G[[i]] <- igraph::sample_smallworld(1, 50, 8, 0.2)
}
for (i in 11:15) {
  G[[i]] <- igraph::sample_pa(50, power = 1, directed = FALSE)
}
graph.hclust(G, 3)
```

---

**graph.kmeans**

*K-means for Graphs*

**Description**

`graph.kmeans` clusters graphs following a k-means algorithm based on the Jensen-Shannon divergence between the spectral densities of the graphs.

**Usage**

```r
graph.kmeans(x, k, nstart = 2)
```
**Arguments**

- **x**: a list containing the graphs or their adjacency matrices to be clustered.
- **k**: an integer specifying the number of clusters.
- **nstart**: the number of trials of k-means clusterizations. The algorithm returns the clusterization with the best silhouette.

**Value**

a vector of the same length of x containing the clusterization labels.

**References**


**Examples**

```r
set.seed(42)
g <- list()for(i in 1:5){
  g[[i]] <- igraph::sample_gnp(30, p=0.2)
}for(i in 6:10){
  g[[i]] <- igraph::sample_gnp(30, p=0.5)
}res <- graph.kmeans(g, k=2, nstart=2)
res
```

---

**graph.model.selection**  
*Graph model selection*

**Description**

`graph.model.selection` selects the graph model that best approximates the observed graph according to the Graph Information Criterion (GIC).

**Usage**

```r
graph.model.selection(
  G,
  models = NULL,
  parameters = NULL,
  eps = 0.01,
```
Graph model selection

```r
bandwidth = "Silverman",
eigenvalues = NULL
```

**Arguments**

- **G**: the undirected graph (igraph type) or its adjacency matrix. The adjacency matrix of an unweighted graph contains only 0s and 1s, while the weighted graph may have nonnegative real values that correspond to the weights of the edges.

- **models**: either a vector of strings, a list of functions or a list of arrays describing graph models:
  - A vector of strings containing some of the following models: "ER" (Erdos-Renyi random graph), "GRG" (geometric random graph), "KR" (k regular random graph), "WS" (Watts-Strogatz model), and "BA" (Barabasi-Albert model).
  - A list of functions. Each function returns a graph (represented by its adjacency matrix) generated by a graph model and has two arguments: the graph size and the model parameter, in this order.
  - A list of arrays. Each element of the list is a three-dimensional array containing the precomputed spectrum of each model. Let M be a graph model. For each parameter p considered for M, the array of model M contains the eigenvalues of graphs randomly generated by M with parameter p. The position (i,j,k) of the array contains the j-th eigenvalue of the k-th graph that generated by M with the i-th parameter. The attribute 'rownames' of the array corresponds to the parameters converted to string.
  - If the argument "models" is NULL, then the "ER", "WS", and "BA" models will be considered for the model selection.

- **parameters**: list of numeric vectors. Each vector contains the values that will be considered for the parameter estimation of the corresponding model. If the user does not provide the argument 'parameters', then default values are used for the pre-defined models ("ER", "GRG", "KR", "WS", and "BA"). The default vector corresponds to a sequence from
  - 0 to 1 with step 'eps' for the "ER" model (Erdos-Renyi random graph), in which the parameter corresponds to the probability to connect a pair of vertices;
  - 0 to sqrt(2) with step 'eps' for the "GRG" model (geometric random graph), in which the parameter corresponds to the radius used to construct the geometric graph in a unit square;
  - 0 to 'n' with step 'n*eps' for the "KR" model (k regular random graph), in which the parameter corresponds to the degree 'k' of a regular graph;
  - 0 to 1 with step 'eps' for the "WS" model (Watts-Strogatz model), in which the parameter corresponds to the probability to reconnect a vertex;
  - and 0 to 3 with step 'eps' for the "BA" model (Barabasi-Albert model), in which the parameter corresponds to the scaling exponent.

- **eps**: precision of the grid (default is 0.01).

- **bandwidth**: string showing which criterion is used to choose the bandwidth during the spectral density estimation. Choose between the following criteria: "Silverman" (default), "Sturges", "bcv", "ucv" and "SJ". "bcv" is an abbreviation of biased
cross-validation, while "ucv" means unbiased cross-validation. "SJ" implements the methods of Sheather & Jones (1991) to select the bandwidth using pilot estimation of derivatives.

eigenvalues optional parameter. It contains the eigenvalues of matrix G. Then, it can be used when the eigenvalues of G were previously computed. If no value is passed, then the eigenvalues of G will be computed by 'graph.model.selection'.

Value

A list containing:

model the indice(s) or name(s) of the selected model(s), i.e. the model(s) that minimize(s) the Graph Information Criterion (GIC).

estimates a matrix in which each row corresponds to a model, the column "param" corresponds to the parameter estimate, and the column "GIC" corresponds to the Graph Information Criterion (GIC), i.e. the Kullback-Leibler divergence between the observed graph and the model.

References


Examples

```r
## Example using an igraph object as input data
set.seed(1)
G <- igraph::sample_gnp(n=30, p=0.5)

# Using strings to indicate the graph models
result1 <- graph.model.selection(G, models=c("ER", "WS"), eps=0.5)
result1

## Using functions to describe the graph models
# Erdos-Renyi graph
model1 <- function(n, p) {
  return(igraph::sample_gnp(n, p))
}
# Watts-Strogatz small-world graph
model2 <- function(n, pr, K=8) {
  return(igraph::sample_smallworld(1, n, K, pr))
}
parameters <- list(seq(0.01, 0.99, 0.49), seq(0.01, 0.99, 0.49))
```
result2 <- graph.model.selection(G, list(model1, model2), parameters)
result2

graph.mult.scaling

Multidimensional scaling of graphs

Description

graph.mult.scaling performs multidimensional scaling of graphs. It takes the Jensen-Shannon divergence between graphs (JS) and uses the 'cmdscale' function from the 'stats' package to obtain a set of points such that the distances between the points are similar to JS.

Usage

graph.mult.scaling(
    G,
    plot = TRUE,
    bandwidth = "Silverman",
    type = "n",
    main = "",
    ...
)

Arguments

G a list of undirected graphs (igraph type) or their adjacency matrices. The adjacency matrix of an unweighted graph contains only 0s and 1s, while the weighted graph may have nonnegative real values that correspond to the weights of the edges.

plot logical. If TRUE (default) the points chosen to represent the Jensen-Shannon divergence between graphs are plotted.

bandwidth string showing which criterion is used to choose the bandwidth during the spectral density estimation. Choose between the following criteria: "Silverman" (default), "Sturges", "bcv", "ucv" and "SJ". "bcv" is an abbreviation of biased cross-validation, while "ucv" means unbiased cross-validation. "SJ" implements the methods of Sheather & Jones (1991) to select the bandwidth using pilot estimation of derivatives.

type what type of plot should be drawn. The default value is "n", which indicates that the points will not be plotted (i.e. only the labels of the graphs will be plotted).

main title of the plot (default value is "").

... additional plotting parameters. See 'plot' function from the 'graphics' package for the complete list.
Value

A matrix in which each column corresponds to a coordinate and each row corresponds to a graph (point). Then, each row gives the coordinates of the points chosen to represent the Jensen-Shannon divergence between graphs.

References


Examples

```r
set.seed(1)
G <- list()
for (i in 1:5) {
  G[[i]] <- igraph::sample_gnp(50, 0.5)
}
for (i in 6:10) {
  G[[i]] <- igraph::sample_smallworld(1, 50, 8, 0.2)
}
for (i in 11:15) {
  G[[i]] <- igraph::sample_pa(50, power = 1, directed = FALSE)
}
graph.mult.scaling(G)
```

Description

`graph.param.estimator` estimates the parameter that best approximates the model to the observed graph according to the Graph Information Criterion (GIC).

Usage

```r
graph.param.estimator(
  G,
  model,
  parameters = NULL,
  eps = 0.01,
  bandwidth = "Silverman",
)```
graph.param.estimator

eigenvalues = NULL,
spectra = NULL,
classic = FALSE
)

Arguments

G
the undirected graph (igraph type) or its adjacency matrix. The adjacency matrix of an unweighted graph contains only 0s and 1s, while the weighted graph may have nonnegative real values that correspond to the weights of the edges.

model
either a string or a function:
A string that indicates one of the following models: "ER" (Erdos-Renyi random graph), "GRG" (geometric random graph), "KR" (k regular random graph), "WS" (Watts-Strogatz model), and "BA" (Barabasi-Albert model).
A function that returns a graph (represented by its adjacency matrix) generated by a graph model. It must contain two arguments: the first one corresponds to the graph size and the second to the parameter of the model.

parameters
numeric vector containing the values that will be considered for the parameter estimation. The 'graph.param.estimator' will return the element of 'parameter' that minimizes the Kullback-Leiber divergence. If the user does not provide the argument 'parameters', and 'model' is an array, then the values considered for the parameter estimation are the rownames converted to a numeric vector. If 'model' is a string, then default values are used for the predefined models ("ER", "GRG", "KR", "WS", and "BA"). The default 'parameter' argument corresponds to a sequence from
0 to 1 with step 'eps' for the "ER" model (Erdos-Renyi random graph), in which the parameter corresponds to the probability to connect a pair of vertices;
0 to \(\sqrt{2}\) with step 'eps' for the "GRG" model (geometric random graph), in which the parameter corresponds to the radius used to construct the geometric graph in a unit square;
0 to 'n' with step 'n*eps' for the "KR" model (k regular random graph), in which the parameter of the model corresponds to the degree 'k' of a regular graph;
0 to 1 with step 'eps' for the "WS" model (Watts-Strogatz model), in which the parameter corresponds to the probability to reconnect a vertex;
and 0 to 3 with step 'eps' for the "BA" model (Barabasi-Albert model), in which the parameter corresponds to the scaling exponent.

eps
precision of the grid (default is 0.01) when 'classic' is TRUE.

bandwidth
string showing which criterion is used to choose the bandwidth during the spectral density estimation. Choose between the following criteria: "Silverman" (default), "Sturges", "bcv", "ucv" and "SJ". "bcv" is an abbreviation of biased cross-validation, while "ucv" means unbiased cross-validation. "SJ" implements the methods of Sheather & Jones (1991) to select the bandwidth using pilot estimation of derivatives.

eigenvalues
optional parameter. It contains the eigenvalues of matrix G. Then, it can be used when the eigenvalues of G were previously computed. If no value is passed, then the eigenvalues of G will be computed by 'graph.param.estimator'.
spectra

optional parameter containing the precomputed spectrum of the model. It is a three-dimensional array in which the first dimension corresponds to all parameters that will be explored in the grid, the second dimension has the same size of the given graph, and the third one corresponds to graphs randomly generated by the model. Thus, the position (i,j,k) contains the j-th eigenvalue of the k-th graph generated with the i-th parameter. The attribute 'rownames' of the array corresponds to the parameters converted to string. If spectra is NULL (default), then 'model' is used to generate random graphs and their spectra are computed automatically.

classic

logical. If FALSE (default) parameter is estimated using ternary search. If TRUE parameter is estimated using grid search.

Value

A list containing:

param

the parameter estimate. For the "ER", "GRG", "KR", "WS", and "BA" models, the parameter corresponds to the probability to connect a pair of vertices, the radius used to construct the geometric graph in a unit square, the degree k of a regular graph, the probability to reconnect a vertex, and the scaling exponent, respectively.

KLD

the Kullback-Leibler divergence between the observed graph and the graph model with the estimated parameter.

References


Examples

```r
set.seed(1)
G <- igraph::sample_gnp(n=50, p=0.5)

# Using a string to indicate the graph model
result1 <- graph.param.estimator(G, "ER", eps=0.25)
result1

## Not run:
# Using a function to describe the graph model
# Erdos-Renyi graph
set.seed(1)
model <- function(n, p) {
  return(igraph::sample_gnp(n, p))
```

sp.anogva <- graph.param.estimator(G, model, seq(0.2, 0.8, 0.1))
result2

## End(Not run)

**sp.anogva**

Semi-Parametric Analysis Of Graph Variability (ANOGVA)

**Description**

sp.anogva statistically tests whether two or more graphs are generated by the same model and set of parameters.

**Usage**

```r
sp.anogva(
  G,
  model,
  maxBoot = 500,
  spectra = NULL,
  eps = 0.01,
  classic = FALSE,
  bandwidth = "Silverman"
)
```

**Arguments**

- **G**
  
a list of undirected graphs (igraph type) or their adjacency matrices. The adjacency matrix of an unweighted graph contains only 0s and 1s, while the weighted graph may have nonnegative real values that correspond to the weights of the edges.

- **model**
  
  A string that indicates one of the following models: "ER" (Erdos-Renyi random graph model), "GRG" (geometric random graph model), "WS" (Watts-Strogatz random graph model), and "BA" (Barabasi-Albert random graph model).

- **maxBoot**
  
  integer indicating the number of bootstrap resamples (default is 500).

- **spectra**
  
  optional parameter containing the precomputed spectrum of the model. It is a three-dimensional array in which the first dimension corresponds to all parameters that will be explored in the parameter estimation, the second dimension has the same size of the given graph, and the third one corresponds to graphs randomly generated by the model. Thus, the position (i,j,k) contains the j-th eigenvalue of the k-th graph generated with the i-th parameter. The attribute 'rownames' of the array corresponds to the parameters converted to string. If spectra is NULL (default), then model is used to generate random graphs and their spectra are computed automatically.
sp.anogva

eps (default is 0.01) precision of the grid when 'classic' = TRUE.

classic logical. If FALSE (default) parameter is estimated using ternary search, if TRUE parameter is estimated using grid search.

bandwidth string showing which criterion is used to choose the bandwidth during the spectral density estimation. Choose between the following criteria: "Silverman" (default), "Sturges", "bcv", "ucv" and "SJ". "bcv" is an abbreviation of biased cross-validation, while "ucv" means unbiased cross-validation. "SJ" implements the methods of Sheather & Jones (1991) to select the bandwidth using pilot estimation of derivatives.

Value

A list containing:

- parameters a vector containing the estimated parameters for each graph.
- F.value the F statistic of the test.
- p.value the p-value of the test.

References


Examples

```r
## Not run:
set.seed(42)
model <- "ER"
G <- list()

# Under H0
G[[1]] <- igraph::sample_gnp(50, 0.5)
G[[2]] <- igraph::sample_gnp(50, 0.5)
G[[3]] <- igraph::sample_gnp(50, 0.5)
result1 <- sp.anogva(G, model, maxBoot = 300)
result1

# Under H1
G[[1]] <- igraph::sample_gnp(50, 0.5)
G[[2]] <- igraph::sample_gnp(50, 0.55)
G[[3]] <- igraph::sample_gnp(50, 0.5)
result2 <- sp.anogva(G, model, maxBoot = 300)
result2

## End(Not run)
```
takahashi.test

Test for the Jensen-Shannon divergence between graphs

Description

takahashi.test tests whether two sets of graphs were generated by the same random graph model. This bootstrap test is based on the Jensen-Shannon (JS) divergence between graphs.

Usage

takahashi.test(G1, G2, maxBoot = 1000, bandwidth = "Silverman")

Arguments

G1 a list of undirected graphs (igraph type) or their adjacency matrices. The adjacency matrix of an unweighted graph contains only 0s and 1s, while the weighted graph may have nonnegative real values that correspond to the weights of the edges.

G2 a list of undirected graphs (igraph type) or their adjacency matrices. The adjacency matrix of an unweighted graph contains only 0s and 1s, while the weighted graph may have nonnegative real values that correspond to the weights of the edges.

maxBoot integer indicating the number of bootstrap resamplings.

bandwidth string showing which criterion is used to choose the bandwidth during the spectral density estimation. Choose between the following criteria: "Silverman" (default), "Sturges", "bcv", "ucv" and "SJ". "bcv" is an abbreviation of biased cross-validation, while "ucv" means unbiased cross-validation. "SJ" implements the methods of Sheather & Jones (1991) to select the bandwidth using pilot estimation of derivatives.

Details

Given two lists of graphs, 'G1' and 'G2', 'takahashi.test' tests H0: "JS divergence between 'G1' and 'G2' is 0" against H1: "JS divergence between 'G1' and 'G2' is larger than 0".

Value

A list containing:

JSD the Jensen-Shannon divergence between 'G1' and 'G2'.

p.value the p-value of the test.
References


Examples

```r
set.seed(1)
G1 <- G2 <- list()
for (i in 1:20) {
  G1[[i]] <- igraph::sample_gnp(n=50, p=0.5)
}
for (i in 1:20) {
  G2[[i]] <- igraph::sample_gnp(n=50, p=0.51)
}
result <- takahashi.test(G1, G2, maxBoot=100)
result
```

---

tang.test  

_Tang hypothesis testing for random graphs._

Description

Given two independent finite-dimensional random dot product graphs, tang.test tests if they have generating latent positions that are drawn from the same distribution.

Usage

tang.test(G1, G2, dim, sigma = NULL, maxBoot = 200)

Arguments

- **G1**: the first undirected graph to be compared. Must be an igraph object.
- **G2**: the second undirected graph to be compared. Must be an igraph object.
- **dim**: dimension of the adjacency spectral embedding.
- **sigma**: a real value indicating the kernel bandwidth. If NULL (default) the bandwidth is calculated by the method.
- **maxBoot**: integer indicating the number of bootstrap resamples (default is 200).
Value

A list containing:

- `T` the value of the test.
- `p.value` the p-value of the test.

References


Examples

```r
set.seed(42)

## test under H0
lpvs <- matrix(rnorm(200), 20, 10)
lpv <- apply(lpv, 2, function(x) { return (abs(x)/sqrt(sum(x^2))) })
G1 <- igraph::sample_dot_product(lpv)
G2 <- igraph::sample_dot_product(lpv)
D1 <- tang.test(G1, G2, 5)
D1

## test under H1
lpvs2 <- matrix(rnorm(200), 20, 10)
lpv <- apply(lpv, 2, function(x) { return (abs(x)/sqrt(sum(x^2))) })
G2 <- suppressWarnings(igraph::sample_dot_product(lpv2))
D2 <- tang.test(G1, G2, 5)
D2
```
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