Package ‘graph4lg’

May 4, 2020

Type Package

Title Build Graphs for Landscape Genetics Analysis

Version 0.5.0

Author Paul Savary

Maintainer Paul Savary <savarypaul660@gmail.com>


Depends R(>= 3.1.0)

License GPL-2

Encoding UTF-8

LazyData true

Imports stringr, adegenet, stats, spatstat, Matrix, vegan, utils, methods, pegas, MASS, igraph, ggplot2, tidyr, sp, rgdal, sf, diveRsity, raster, foreign, ecodist

RdMacros Rdpack

RoxygenNote 7.0.2

Suggests knitr, rmarkdown, Rdpack

VignetteBuilder knitr

NeedsCompilation no
Repository CRAN
Date/Publication 2020-05-04 08:30:08 UTC

R topics documented:

add_nodes_attr ............................................ 3
compar_r_fisher ........................................... 4
convert_cd .................................................. 5
data_ex_genind ........................................... 7
data_ex_gstud ............................................. 8
data_ex_loci ............................................... 8
data_simul_genind ........................................ 9
data_tuto .................................................... 10
df_to_pw_mat ............................................... 10
dist_max_corr .............................................. 11
genepop_to_genind ........................................ 13
genind_to_genepop ....................................... 15
gen_graph_indep .......................................... 16
gen_graph_thr ............................................ 19
gen_graph_topo .......................................... 20
graphab_to_igraph ........................................ 21
graph_modul_compar ...................................... 23
graph_node_compar ....................................... 26
graph_plan ................................................. 28
graph_plot_compar ....................................... 29
graph_topo_compar ....................................... 30
graph_to_df ............................................... 32
graph_to_shp .............................................. 33
gstud_to_genind ......................................... 34
g_percol ..................................................... 35
kernel_param ............................................... 36
loci_to_genind ........................................... 37
mat_gen_dist ............................................... 38
mat_geo_dist .............................................. 40
mat_pw_dps ............................................... 41
mat_pw_d_j ............................................... 42
mat_pw_fst ............................................... 43
mat_pw_gst ............................................... 44
plot_graph_lg ............................................. 46
plot_graph_modul ....................................... 47
plot_w_hist ............................................... 49
pop_gen_index ............................................. 50
pts_pop_ex ............................................... 51
pts_pop_simul .......................................... 52
pw_mat_to_df ............................................. 52
reorder_mat .............................................. 53
scatter_dist .............................................. 54
add_nodes_attr

Add attributes to the nodes of a graph

Description
The function adds attributes to the nodes of a graph from either an object of class data.frame or from a shapefile layer. The node IDs in the input objects must be the same as in the graph object.

Usage

```r
add_nodes_attr(
  graph,
  input = "df",
  data,
  dir_path = NULL,
  layer = NULL,
  index = "Id",
  include = "all"
)
```

Arguments

- **graph**: A graph object of class igraph.
- **input**: A character string indicating the nature of the input data from which come the attributes to add to the nodes.
  - If `input = "shp"`, then attributes come from the attribute table of a shapefile layer of type point.
  - If `input = "df"`, then attributes come from an object of class data.frame In both cases, input attribute table or dataframe must have a column with the exact same values as the node IDs.
- **data** (only if `input = "df"`) The name of the object of class data.frame with the attributes to add to the nodes.
- **dir_path** (only if `input = "shp"`) The path (character string) to the directory containing the shapefile layer of type point whose attribute table contains the attributes to add to the nodes.
- **layer** (only if `input = "shp"`) The name (character string) of the shapefile layer of type point (without extension, ex.: "nodes" refers to "nodes.shp" layer) whose attribute table contains the attributes to add to the nodes.
- **index**: The name (character string) of the column with the nodes names in the input data (column of the attribute table or of the dataframe).
include A character string (vector) indicating which columns of the input data will be added as nodes’ attributes. By default, ‘include = "all"’, i.e. every column of the input data is added. Alternatively, ‘include’ can be a vector with the names of the columns to add (ex.: "c('x', 'y', 'pop_name')").

Details

The graph can be created with the function graphab_to_igraph from shapefile layers created with GRAPHAB. Values of the metrics computed at the node level with GRAPHAB can then be added to such a graph with this function.

Value

A graph object of class igraph

Author(s)

P. Savary

Examples

data("data_tuto")
graph <- data_tuto[[3]]
df_nodes <- data.frame(Id = igraph::V(graph)$name,  
    Area = runif(50, min = 10, max = 60))
graph <- add_nodes_attr(graph,  
    data = df_nodes,  
    input = "df",  
    index = "Id",  
    include = "Area")

compar_r_fisher Compare two correlation coefficients obtained from different sample sizes

Description

The function compares two correlation coefficients obtained from different sample sizes using Z-Fisher transformation.

Usage

compar_r_fisher(data)
**Argument**

**data**

An object of class `data.frame` with at least 4 columns of data used to perform the test. 4 columns must be called "n1", "n2", "r1" and "r2".

- n1 and n2 are the sizes of the samples from which r1 and r2 were computed respectively.
- r1 and r2 are Pearson’s correlation coefficients

**Details**

The Z-Fisher method consists in computing z scores from the correlation coefficients and to compare these z scores. z scores are computed as follows : Let n1 and r1 be the sample size and the correlation coefficient, z1 = (1/2)*log((1+r1) / (1-r1)) Then, a test’s statistic is computed from z1 and z2 : Z = (z1-z2) / sqrt((1/(n1-3)) + (1/(n2-3))) If Z is above the limit given by the alpha value, then the difference between r1 and r2 is significant

**Value**

An object of class `data.frame` with the same columns as 'data' and 4 columns more : z1, z2 (respective z-scores), Z (test’s statistic) and p (p-value) of the test.

**Author(s)**

P. Savary

**Examples**

```r
df <- data.frame(n1 = rpois(n = 40, lambda = 85),
n2 = rpois(n = 40, lambda = 60),
r1 = runif(n = 40, min = 0.6, max = 0.85),
r2 = runif(n = 40, min = 0.55, max = 0.75))
data <- compar_r_fisher(df)
```

---

**convert_cd**

*Fit a model to convert cost-distances into Euclidean distances*

**Description**

The function fits a model to convert cost-distances into Euclidean distances as implemented in GRAPHAB software.

**Usage**

```r
convert_cd(
  mat_euc,
  mat_ld,
  to_convert,
  method = "log-log",
)```
Arguments

mat_euc A symmetric matrix with pairwise geographical Euclidean distances between populations or sample sites. It will be the explanatory variable, and only values from the off diagonal lower triangle will be used.

mat_ld A symmetric matrix with pairwise landscape distances between populations or sample sites. These distances can be cost-distances or resistance distances, among others. It will be the explained variable, and only values from the off diagonal lower triangle will be used.

to_convert A numeric value or numeric vector with Euclidean distances to convert into cost-distances.

method A character string indicating the method used to fit the model.

- If `method = "log-log"` (default), then the model takes the following form: 
  \[
  \log(\text{ld}) \sim A + B \times \log(\text{euc})
  \]
- If `method = "lm"`, then the model takes the following form: 
  \[
  \text{ld} \sim A + B \times \text{euc}
  \]

fig Logical (default = TRUE) indicating whether a figure is plotted

line_col (if `fig = TRUE`) Character string indicating the color used to plot the line (default: "blue"). It must be a hexadecimal color code or a color used by default in R.

pts_col (if `fig = TRUE`) Character string indicating the color used to plot the points (default: "#999999"). It must be a hexadecimal color code or a color used by default in R.

Details

IDs in `mat_euc` and `mat ld` must be the same and refer to the same sampling site or populations, and both matrices must be ordered in the same way. Matrix of Euclidean distance `mat_euc` can be computed using the function `mat_geo_dist`. Matrix of landscape distance `mat ld` can be computed using GRAPHAB software. Before the log calculation, 0 distance values are converted into 1, so that they are 0 after this calculation.

Value

A list of output (converted values, estimated parameters, R2) and optionally a ggplot2 object to plot

Author(s)

P. Savary
References


Examples

data("data_tuto")
mat_ld <- data_tuto[[2]][1:10, 1:10] * 1000
mat_euc <- data_tuto[[1]][1:10, 1:10] * 50000
to_convert <- c(30000, 40000)
res <- convert_cd(mat_euc = mat_euc,
                  mat_ld = mat_ld,
                  to_convert = to_convert, fig = FALSE)

Description

Genetic dataset from genetic simulation on CDPOP 200 individuals, 10 populations 20 microsatellite loci (3 digits coding) 100 generations simulated

Usage

data_ex_genind

Format

An object of type ’genind’

Details

The simulation was made with CDPOP during 100 generations. Dispersal was possible between the 10 populations. Its probability depended on the cost distance between populations, calculated on a simulated resistance surface (raster). Mutations were not possible. There were initially 600 alleles in total (many disappeared because of drift). Population stayed constant with a sex-ratio of 1. Generations did not overlap. This simulation includes a part of stochasticity and these data result from only 1 simulation run.

References


Examples

data("data_ex_genind")
length(unique(data_ex_genind@pop))
**data_ex_gstud**

**data_ex_gstud genetic dataset**

**Description**
Genetic dataset from genetic simulation on CDPOP 200 individuals, 10 populations 20 microsatellite loci (3 digits coding) 100 generations simulated

**Usage**
```
data_ex_gstud
```

**Format**
A `data.frame` with columns:

- **ID** Individual ID
- **POP** Population name
- **LOCI-1 to LOCI-20** 20 loci columns with microsatellite data with 3 digits coding, alleles separated by ":", and blank missing data (class 'locus' from `gstudio`)

**Examples**
```
data("data_ex_gstud")
str(data_ex_gstud)
length(unique(data_ex_gstud$POP))
```

---

**data_ex_loci**

**data_ex_loci genetic dataset**

**Description**
Genetic dataset from genetic simulation on CDPOP 200 individuals, 10 populations 20 microsatellite loci (3 digits coding) 100 generations simulated

**Usage**
```
data_ex_loci
```

**Format**
An object of class 'loci' and 'data.frame' with the columns:

- **population** Population name
- **Other columns** 20 loci columns with microsatellite data with 3 digits coding, alleles separated by "/", and missing data noted "NA/NA"

Row names correspond to individuals’ ID
data_simul_genind

Examples

data("data_ex_loci")
length(unique(data_ex_loci$population))

data_simul_genind data_simul_genind genetic dataset

Description

Genetic dataset from genetic simulation on CDPOP 1500 individuals, 50 populations 20 microsatellite loci (3 digits coding) 50 generations simulated

Usage

data_simul_genind

Format

An object of type 'genind'

Details

The simulation was made with CDPOP during 50 generations. Dispersal was possible between the 50 populations. Its probability depended on the cost distance between populations, calculated on a simulated resistance surface (raster). Mutations were not possible. There were initially 600 alleles in total (many disappeared because of drift). Population stayed constant with a sex-ratio of 1. Generations did not overlap. This simulation includes a part of stochasticity and these data result from only 1 simulation run.

References


Examples

data("data_simul_genind")
length(unique(data_simul_genind@pop))
data_tuto

Description
Data used to generate the vignette
Data used to generate the vignette

Usage
data_tuto
data_tuto

Format
Several outputs or inputs to show how the package works in a list

- **dmc**  Output of the function 'dist_max_corr'
- **graph_ci**  Genetic independence graph example
- **mat_dps**  Genetic distance matrix example
- **mat_pg**  Second genetic distance matrix example

Examples

data("data_tuto")
mat_dps <- data_tuto[[1]]
str(mat_dps)
data("data_tuto")
mat_dps <- data_tuto[[1]]
str(mat_dps)

---

df_to_pw_mat

Convert an edge-list data.frame into a pairwise matrix

Description
The function converts an edge-list data.frame into a symmetric pairwise matrix

Usage
df_to_pw_mat(data, from, to, value)
Arguments

<table>
<thead>
<tr>
<th>Argument</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>data</td>
<td>An object of class data.frame</td>
</tr>
<tr>
<td>from</td>
<td>A character string indicating the name of the column with the ID of the origins</td>
</tr>
<tr>
<td>to</td>
<td>A character string indicating the name of the column with the ID of the arrivals</td>
</tr>
<tr>
<td>value</td>
<td>A character string indicating the name of the column with the values corresponding to each pair</td>
</tr>
</tbody>
</table>

Details

The matrix is a symmetric matrix. Be careful, you shall not provide a data.frame with different values corresponding to the pair 1-2 and 2-1 as an example. Ideally, for a complete matrix, data should have n(n-1)/2 rows if values are computed between n objects.

Value

A pairwise matrix

Author(s)

P. Savary

Examples

```r
data(pts_pop_simul)
suppressWarnings(mat_geo <- mat_geo_dist(pts_pop_simul,
                                           ID = "ID",
                                           x = "x",
                                           y = "y")
g <- gen_graph_topo(mat_w = mat_geo,
                     mat_topo = mat_geo,
                     topo = "comp")
df <- data.frame(igraph::as_edgelist(g))
df$w <- igraph::E(g)$weight
df_to_pw_mat(df, from = "X1", to = "X2", value = "w")
```

Description

The function enables to compute the distance at which the correlation between genetic distance and landscape distance is maximal, using a method similar to that employed by van Strien et al. (2015). Iteratively, distance threshold values are tested. For each value, all the population pairs separated by a landscape distance larger than the threshold are removed before the Mantel correlation coefficient between genetic distance and landscape distance is computed. The distance threshold at which the correlation is the strongest is then identified. A figure showing the evolution of the correlation coefficients when landscape distance threshold increases is plotted.
Usage

dist_max_corr(
  mat_gd,
  mat_ld,
  interv,
  from = NULL,
  to = NULL,
  fig = TRUE,
  thr_gd = NULL,
  line_col = "black",
  pts_col = "#999999"
)

Arguments

mat_gd A symmetric matrix with pairwise genetic distances between populations or sample sites.

mat_ld A symmetric matrix with pairwise landscape distances between populations or sample sites. These distances can be Euclidean distances, cost-distances or resistance distances, among others.

interv A numeric value indicating the interval between the different distance thresholds for which the correlation coefficients are computed.

from (optional) The minimum distance threshold value at which the correlation coefficient is computed.

to (optional) The maximum distance threshold value at which the correlation coefficient is computed.

fig Logical (default = TRUE) indicating whether a figure is plotted.

thr_gd (optional) A numeric value used to remove genetic distance values from the data before the calculation. All genetic distances values above 'thr_gd' are removed from the data. This parameter can be used especially when there are outliers.

line_col (optional, if fig = TRUE) A character string indicating the color used to plot the line (default: "blue"). It must be a hexadecimal color code or a color used by default in R.

pts_col (optional, if fig = TRUE) A character string indicating the color used to plot the points (default: "#999999"). It must be a hexadecimal color code or a color used by default in R.

Details

IDs in 'mat_gd' and 'mat_ld' must be the same and refer to the same sampling sites or populations, and both matrices must be ordered in the same way. The correlation coefficient between genetic distance and landscape distance computed is a Mantel correlation coefficient. If there are less than 50 pairwise values, the correlation is not computed, as in van Strien et al. (2015). Such a method can be subject to criticism from a strict statistical point of view given correlation coefficients computed from samples of different size are compared. The matrix of genetic distance 'mat_gd' can be computed using mat_gen_dist. The matrix of landscape distance 'mat_ld' can be computed...
using \texttt{mat\_geo\_dist} when the landscape distance needed is a Euclidean geographical distance. Mantel correlation coefficients are computed using the function \texttt{mantel}.

\textbf{Value}

A list of objects:

- The distance at which the correlation is the highest.
- The vector of correlation coefficients at the different distance thresholds
- The vector of the different distance thresholds
- A ggplot2 object to plot

\textbf{Author(s)}

P. Savary

\textbf{References}


\textbf{Examples}

data("data\_tuto")
mat\_gen <- data\_tuto[[1]]
mat\_dist <- data\_tuto[[2]]*1000
res\_dmc <- dist\_max\_corr(mat\_gd = mat\_gen, 
mat\_ld = mat\_dist, 
from = 32000, to = 42000, 
interv = 5000, 
fig = FALSE)

\textbf{Description}

The function converts a text file in the format used by GENEPOP software into a genind object

\textbf{Usage}

genepop\_to\_genind(path, n\_loci, pop\_names = NULL)
Arguments

- **path**: A character string with the path leading to the GENEPOP file in format .txt, or alternatively the name of this file in the working directory.
- **n.loci**: The number of loci in the GENEPOP file (integer).
- **pop_names**: (optional) Populations’ names in the same order as in the GENEPOP file. Vector object (class character) of the same length as the number of populations. Without this parameter, populations are numbered from 1 to the number of populations.

Details

This function uses functions from **pegas** package. GENEPOP file should only include microsatellites loci with allele names of length 3. The loci line(s) must not start with a spacing.

Value

An object of type `genind`.

Author(s)

P. Savary

References


See Also

For more details about GENEPOP file formatting: [http://genepop.curtin.edu.au/help_input.html#Input](http://genepop.curtin.edu.au/help_input.html#Input) For the opposite conversion, see `genind_to_genepop`. The output file can be used to compute pairwise FST matrix with `mat_pw_fst`

Examples

```r
path_in <- system.file('extdata', 'gpop_simul_10_g100_04_20.txt',
                        package = 'graph4lg')
file_n <- file.path(tempdir(), "gpop_simul_10_g100_04_20.txt")
file.copy(path_in, file_n, overwrite = TRUE)
genepop_to_genind(path = file_n, n.loci = 20,
                   pop_names = as.character(order(as.character(1:10))))
file.remove(file_n)
```
**Description**

The function converts an object of class `genind` into a GENEPOP file. It then allows to use the functionalities of the GENEPOP software and its derived package `GENEPOP` on R, as well as some functions from other packages (differentiation test, F-stats calculations, HWE test...). It is designed to be used with diploid microsatellite data and alleles coded with 3 digits.

**Usage**

```r
genind_to_genepop(x, output = "data.frame")
```

**Arguments**

- `x`: An object of class `genind` from package `adegenet`.
- `output`: A character string indicating the option used to select what the function will return:
  - If `output = "data.frame"` (default), then the function will return an object 'x' of class `data.frame` ready to be saved as a text file with the following command: `write.table(x, file = "file_name.txt", quote=FALSE, row.names=FALSE, col.names=FALSE)`.
  - If `output = "path_to_file/file_name.txt"`, then the function will write a text file named 'file_name.txt' in the directory corresponding to 'path_to_file'. Without 'path_to_file', the text file is written in the current working directory. The text file has the format required by GENEPOP software.

**Value**

An object of type `data.frame` if output = "data.frame". If output is the path and/or the file name of a text file, then nothing is returned in R environment but a text file is created with the specified file name, either in the current working directory or in the specified folder.

**Warning**

- **Confusion**: Do not confound this function with `genind2genpop` from `adegenet`. The latter converts an object of class `genind` into an object of class `genpop`, whereas `genind_to_genepop` converts an object of class `genind` into a text file compatible with GENEPOP software (Rousset, 2008).

- **Allele coding**: This function can handle genetic data with different alleles coding lengths (for example alleles '99' and '101' at one locus). BUT, it is highly advisable not to use it with this type of data because applications using GENEPOP file won't be able to use the output file.

- **Individuals order**: When individuals in input data are not ordered by populations, individuals from the same population can be separated by individuals from other populations. It can be problematic when calculating then pairwise distance matrices. Therefore, in such a case, individuals are ordered by populations and populations ordered in alphabetic order.
Author(s)

P. Savary

References


See Also

For more details about GENEPOP file formatting: [http://genepop.curtin.edu.au/help_input.html#Input](http://genepop.curtin.edu.au/help_input.html#Input). For the opposite conversion, see `genepop_to_genind`. The output file can be used to compute pairwise FST matrix with `mat_pw_fst`.

Examples

```r
data(data_ex_genind)
x <- data_ex_genind
df_genepop <- suppressWarnings(genind_to_genepop(x,
output = "data.frame"))
```

---

gen_graph_indep Create an independence graph of genetic differentiation from genetic data of class genind

Description

The function allows to create genetic graphs from genetic data by applying the conditional independence principle. Populations whose allelic frequencies covary significantly once the covariance with the other populations has been taken into account are linked on the graphs.

Usage

```r
gen_graph_indep(
  x,
  dist = "basic",
  cov = "sq",
  pcor = "magwene",
  alpha = 0.05,
  test = "EED",
  adj = "none",
  output = "igraph"
)
```
Arguments

- **x**: An object of class genind that contains the multilocus genotype (format 'locus') of the individuals as well as their population and their geographical coordinates.

- **dist**: A character string indicating the method used to compute the multilocus genetic distance between populations.
  - If `dist = 'basic'` (default), then the multilocus genetic distance is computed using a Euclidean genetic distance formula (Excoffier et al., 1992).
  - If `dist = 'weight'`, then the multilocus genetic distance is computed as in Fortuna et al. (2009). It is a Euclidean genetic distance giving more weight to rare alleles.
  - If `dist = 'PG'`, then the multilocus genetic distance is computed as in popgraph::popgraph function, following several steps of PCA and SVD (Dyer et Nason, 2004).
  - If `dist = 'PCA'`, then the genetic distance is computed following a PCA of the matrix of allelic frequencies by population. It is a Euclidean genetic distance between populations in the multidimensional space defined by all the independent principal components.

- **cov**: A character string indicating the formula used to compute the covariance matrix from the distance matrix.
  - If `cov = 'sq'` (default), then the covariance matrix is calculated from the matrix of squared distances as in Everitt et Hothorn (2011).
  - If `cov = 'dist'`, then the covariance matrix is calculated from the matrix of distances as in Dyer et Nason (2004) and popgraph function.

- **pcor**: A character string indicating the way the partial correlation matrix is computed from the covariance matrix.
  - If `pcor = 'magwene'`, the steps followed are the same as in Magwene (2001) and in popgraph::popgraph function. It is the recommended option as it meets mathematical requirements.
  - If `pcor = 'other'`, the steps followed are the same as used by Fortuna et al. (2009). They are not consistent with the approach of Magwene (2001).

- **alpha**: A numeric value corresponding to the statistical tolerance threshold used to test the difference from 0 of the partial correlation coefficients. By default, `alpha=0.05`.

- **test**: A character string indicating the method used to test the significance of the partial correlation coefficients.
  - If `test = 'EED'` (default), then the Edge Exclusion Deviance criterion is used (Whittaker, 2009). Although other methods exist, this is the most common and thus the only one implemented here.

- **adj**: A character string indicating the way of adjusting p-values to assess the significance of the p-values.
  - If `adj = 'none'` (default), there is no p-value adjustment correction.
  - If `adj = 'holm'`, p-values are adjusted using the sequential Bonferroni correction (Holm, 1979).
If `adj = 'bonferroni'`, p-values are adjusted using the classic Bonferroni correction
If `adj = 'BH'`, p-values are adjusted using Benjamini et Hochberg (1995) correction controlling false discovery rate

output
A character string indicating the matrices included in the output list.
- If `output = 'all'` (default), then D (distance matrix), C (covariance matrix), Rho (partial correlation matrix), M (graph incidence matrix) and S (strength matrix) are included
- If `output = 'dist_graph'`, then the distance matrix D is returned only with the values corresponding to the graph edges
- If `output = 'str_graph'`, then the strength values matrix S is returned only with the values corresponding to the graph edges
- If `output = 'inc'`, then the binary adjacency matrix M is returned
- If `output = 'igraph'`, then a graph of class igraph is returned

Details
The function allows to vary many parameters such as the genetic distance used, the formula used to compute the covariance, the statistical tolerance threshold, the p-values adjustment, among others.

Value
A list of objects of class matrix, an object of class matrix or a graph object of class igraph

Author(s)
P. Savary

References
Examples

```r
data(data_ex_genind)
dist_graph_test <- gen_graph_indep(x = data_ex_genind, dist = "basic",
    cov = "sq", pcor = "magwene",
    alpha = 0.05, test = "EED",
    adj = "none", output = "igraph")
```

---

### Description

The function allows to construct a genetic graph whose links’ weights are larger or lower than a specific threshold.

### Usage

```r
gen_graph_thr(mat_w, mat_thr = NULL, thr, mode = "larger")
```

### Arguments

- `mat_w`: A symmetric (pairwise) matrix whose elements will be the links’ weights.
- `mat_thr`: (optional) A symmetric (pairwise) distance matrix whose values will be used for the pruning based on the threshold value.
- `thr`: The threshold value (logically between min(mat_thr) and max(mat_thr)).
- `mode`: 
  - If `mode = "larger"` (default), all the links whose weight is larger than `thr` are removed.
  - If `mode = "lower"`, all the links whose weight is lower than `thr` are removed.

### Details

If `mat_thr` is not defined, `mat_w` is used for the pruning. Matrices `mat_w` and `mat_thr` must have the same dimensions and the same rows’ and columns’ names. Values in `mat_thr` matrix must be positive. Negative values from `mat_w` are transformed into zeros. The function works only for undirected graphs.

### Value

A graph object of class igraph

### Author(s)

P. Savary
**Examples**

```r
code_snippet
```

**Description**

The function constructs a genetic graph with a specific topology from genetic and/or geographical distance matrices.

**Usage**

```r
gen_graph_topo(mat_w, mat_topo = NULL, topo = "gabriel", k = NULL)
```

**Arguments**

- `mat_w`: A symmetric (pairwise) matrix whose elements will be the links’ weights.
- `mat_topo`: (optional) A symmetric (pairwise) distance matrix whose values will be used for the pruning method.
- `topo`: Which topology does the created graph have?
  - If `topo = 'gabriel'` (default), the resulting graph will be a Gabriel graph (Gabriel et al., 1969). It means that there is a link between nodes x and y if and only if $d_{xy}^2 \leq \min(\sqrt{d_{xz}^2 + d_{yz}^2})$, with z any other node of the graph.
  - If `topo = 'mst'`, the resulting graph will have the topology of a minimum spanning tree. It means that the graph will not include any cycle (tree) and it will be the subgraph with a tree topology with the minimum total links’ weight (based on `mat_topo` values).
  - If `topo = 'percol'`, if the link of the resulting graph with the minimum weight is removed, then the graph breaks into two components.
  - If `topo = 'comp'`, a complete graph whose links are weighted with values from `mat_w` is created.
  - If `topo = 'knn'`, a k-nearest neighbor graph whose links are weighted with values from `mat_w` is created. If the distance between node i and node j is among the k-th smallest distances between node i and the other nodes according to distances in matrix `mat_topo`, then there is a link between i and j in the resulting graph. Therefore, a node can be connected to more than two nodes because the nearest node to node j is not necessarily among the k nearest neighbors to node i. Let $d_1$ be the smallest distance from node
i to other nodes, if there are k nodes or more at this distance from node i, they are all connected to i. If there are less than k nodes connected to i at a distance \( d_1 \), then we consider nodes at a distance \( d_2 \) from i. In the latter case, all the nodes at a distance \( d_2 \) from i are connected to i.

\( k \) (if ‘topo = ‘knn”) An integer which indicates the number of nearest neighbors considered to create the K-nearest neighbor graph. \( k \) must be lower than the total number of nodes minus 1.

Details

If ‘mat_topo’ is not defined, ‘mat_w’ is used for the pruning. Matrices ‘mat_w’ and ‘mat_topo’ must have the same dimensions and the same rows’ and columns’ names. Values in ‘mat_topo’ matrix must be positive. Negative values from ‘mat_w’ are transformed into zeros. The function works only for undirected graphs. Note that the topology ‘knn’ works best when ‘mat_topo’ contains distance values from a continuous value range, thereby avoiding equal distances between a node and the others. are more than \( k \) nodes located at distances in the \( k \)-th smallest distances

Value

A graph object of class igraph

Author(s)

P. Savary

References


Examples

```r
mat_w <- mat_gen_dist(x = data_ex_genind, dist = 'DPS')
suppressWarnings(mat_topo <- mat_geo_dist(pts_pop_ex, ID = "ID",
    x = "x",
    y = "y"))
mat_topo <- mat_topo[row.names(mat_w), colnames(mat_w)]
graph <- gen_graph_topo(mat_w, mat_topo, topo = "mst")
```

**Description**

The function imports a landscape graph created with GRAPHAH software and converts it into a graph object of class igraph. The graph has weighted links and is undirected. Nodes have spatial coordinates. Other nodes attributes can be included. It takes shapefiles layers created with GRAPHAH as input.
Usage

graphab_to_igraph(
  dir_path,
  nodes = "patches",
  links,
  weight = "cost",
  fig = FALSE,
  crds = FALSE
)

Arguments

dir_path       A character string indicating the path of the GRAPHAB project directory. This directory normally contains several spatial layer files in format .shp:
                      • the spatial layer of the habitat patches corresponding to the nodes of the graph (usually named 'patches.shp').
                      • (alternatively) an exported spatial layer of the nodes (faster option).
                      • the link spatial layer file used to import the graph.

nodes          A character string indicating the names of the node spatial layer in format .shp (without extension, ex.: "nodes" refers to "nodes.shp" layer). This layer has been created with GRAPHAB and has therefore coordinates in a projected co-ordinates reference system. Default: nodes = "patches", referring to the spatial polygon layer of the habitat patches.

links          A character string indicating the name of the link spatial layer in format .shp (without extension, ex.: "link_graph" refers to "link_graph.shp" layer). This layer has been created with GRAPHAB and has therefore coordinates in a projected coordinates reference system. It includes in the attribute tables between patches Euclidean as well as cost-distance. These distances are used to weight the link.

weight         A character string ("euc" or "cost") indicating whether to weight the links with Euclidean distance or cost-distance (default) values.

fig            Logical (default = FALSE) indicating whether to plot a figure of the resulting spatial graph. The figure is plotted using function plot_graph_lg. The plotting can be long if the graph has many nodes and links.

crds           Logical (default = FALSE) indicating whether to create an object of class data.frame with the nodes spatial coordinates. Such a data.frame has 3 columns: 'ID', 'x', 'y'.

Details

Nodes attributes can be added to the graph using the function add_nodes_attr.

Value

A graph object of class igraph (if crds = FALSE) or a list of objects: a graph object of class igraph and a data.frame with the nodes spatial coordinates (if crds = TRUE).
Author(s)

P. Savary

References


Examples

```r
path <- system.file('extdata', package='graph4lg')
links <- "liens_simul_10"
graph <- graphab_to_igraph(dir_path = path,
                          nodes = "patches_10",
                          links = links,
                          fig = FALSE)
```

```r

graph_modul_compar(x, y, mode = "graph", nb_modul = NULL, algo = "fast_greedy", weight = "inv", data = NULL)
```

Description

The function computes the Adjusted Rand Index (ARI) to compare two graphs’ partitions into modules or clusters more generally. Both graphs must have the same number of nodes, but not necessarily the same number of links. They must also have the same nodes’ names and in the same order.

Arguments

- `x` The first graph object
  - If `mode = 'graph'` (default), `x` is a graph object of class `igraph`. Then, its nodes must have the same names as in graph `y`.  

Graph Module Comparison

- If `mode = 'data.frame'`, `x` refers to a column of the `data.frame` named `data`. Then `x` must be a character string indicating the name of the column of `data` with the modules' labels of the nodes in the first graph. In that case, the column can be of class numeric, character or factor but will be converted into a numeric vector in any case.
- If `mode = 'vector'`, `x` is a vector of class character, factor or numeric. In that case, it must have the same length as `vector y` and will be converted into a numeric vector.

**y**
The second graph object. Same classes possible as for `x`. Must be of the same format as `x`.

**mode**
A character string indicating whether `x` and `y` are igraph objects, vectors or columns from a data.frame. `mode` can be 'graph', 'data.frame' or 'vector'.

**nb_modul**
(if `x` and `y` are igraph objects) A numeric value or numeric vector with 2 elements indicating the number of modules to create in both graphs.

- If `nb_modul` is a numeric value, then the same number of modules are created in both graphs.
- If `nb_modul` is a numeric vector of length 2, then the numbers of modules created in graphs `x` and `y` are the first and second elements of `nb_modul`, respectively.

**algo**
(if `x` and `y` are igraph objects) A character string indicating the algorithm used to create the modules with `igraph`.

- If `algo = 'fast_greedy'` (default), function `cluster_fast_greedy` from `igraph` is used (Clauset et al., 2004).
- If `algo = 'walktrap'` (default), function `cluster_walktrap` from `igraph` is used (Pons et Latapy, 2006) with 4 steps (default options).
- If `algo = 'louvain'`, function `cluster_louvain` from `igraph` is used (Blondel et al., 2008). In that case, the number of modules created in each graph is imposed.
- If `algo = 'optimal'`, function `cluster_optimal` from `igraph` is used (Brandes et al., 2008) (can be very long). In that case, the number of modules created in each graph is imposed.

**weight**
(optional, if `x` and `y` are igraph objects) A character string or character vector indicating how to weight graphs' links during the calculation of the modularity.

- If `weight = 'inv'` (default), then links are weighted with the inverse values of their initial weights.
- If `weight = 'w'`, then links are weighted with their initial weights values.
- If `weight = 'none'`, then links are not weighted during the calculation.

Two different weightings can be used to create the modules of the two graphs.

- If `weight` is a character string, then the same algorithm is used for both graphs.
- If `weight` is a character vector of length 2, then the link weighting used by the algorithm to create the modules of graphs `x` and `y` is determined by the first and second elements of `weight`, respectively.
If the graphs’ links are not weighted, then this argument is ignored. Links with large weights are considered as stronger connections in the modularity calculation.

**Data** (if x and y are columns from a data.frame) An object of class data.frame with at least two columns and as many rows as there are nodes in the graphs compared. The columns indicate the modules of each node in 2 different classifications.

**Details**

This index takes values between -1 and 1. It measures how often pairs of nodes pertaining to the same module in one graph also pertain to the same module in the other graph. Therefore, large values indicate that both partitions are similar. The Rand Index can be defined as the frequency of agreement between two classifications into discrete classes. It is the number of times a pair of elements are classified into the same class or in two different classes in both compared classifications, divided by the total number of possible pairs of elements. The Rand Index is between 0 and 1 but its maximum value depends on the number of elements. Thus, another 'adjusted' index was created, the Adjusted Rand Index. According to the Hubert et Arabie's formula, the ARI is computed as follows: \( ARI = \frac{Index - Expected\ Index}{Maximum\ Index - Expected\ Index} \) where the values of Index, Expected index and Maximum index are computed from a contingency table. This function uses `adjustedRandIndex` from package `mclust` which applies the Hubert and Arabie’s formula for the ARI. This function works for undirected graphs only.

**Value**

The value of the ARI

**Author(s)**

P. Savary

**References**


**Examples**

```r
data(data_ex_genind)
data(pts_pop_ex)
mat_dist <- suppressWarnings(graph4lg::mat_geo_dist(data=pts_pop_ex,
  ID = "ID",
  x = "x",
  y = "y"))
```
graph_node_compar

Compare the local properties of the nodes from two graphs

Description
The function computes a correlation coefficient between the graph-theoretic metric values computed
at the node-level in two graphs sharing the same nodes. It allows to assess whether the connectivity
properties of the nodes in one graph are similar to that of the same nodes in the other graph. Alternatively, the correlation is computed between a graph-theoretic metric values and the values of an
attribute associated to the nodes of a graph.

Usage
graph_node_compar(
  x,
  y,
  metrics = c("siw", "siw"),
  method = "spearman",
  weight = TRUE,
  test = TRUE
)

Arguments
x                   An object of class igraph. Its nodes must have the same names as in graph y.
y                   An object of class igraph. Its nodes must have the same names as in graph x.
metrics             Two-elements character vector specifying the graph-theoretic metrics computed
                     at the node-level in the graphs or the nodes’ attribute values to be correlated to
                     these metrics. Graph-theoretic metrics can be:
                     • Degree (metrics = c("deg",...))
                     • Closeness centrality index (metrics = c("close",...))
                     • Betweenness centrality index (metrics = c("btw",...))
                     • Strength (sum of the weights of the links connected to a node) (metrics = c("str",...))
                     • Sum of the inverse weights of the links connected to a node (metrics = c("siw",...), default)
• Mean of the inverse weights of the links connected to a node (metrics = c("miw",...))

Nodes’ attributes must have the same names as in the igraph object, and must refer to an attribute with numerical values. The vector metrics is composed of two character values. When a nodes’ attribute has the same name as a metric computable from the graph, nodes’ attributes are given priority.

**method**  
A character string indicating which correlation coefficient is to be computed ("pearson", "kendall" or "spearman" (default)).

**weight**  
Logical which indicates whether the links are weighted during the calculation of the centrality indices betweenness and closeness. (default: weight = TRUE). Links’ weights are interpreted as distances when computing the shortest paths. They should then be inversely proportional to the strength of the relationship between nodes (e.g. to fluxes).

**test**  
Logical. Should significance testing be performed? (default = TRUE)

**Details**

The correlation coefficients between the metrics can be computed in different ways, as initial assumptions (e.g. linear relationship) are rarely verified. Pearson’s r, Spearman’s rho and Kendall’s tau can be computed (from function `cor`). When x is similar to y, then the correlation is computed between two metrics characterizing the nodes of the same graph.

**Value**

A list summarizing the correlation analysis.

**Author(s)**

P. Savary

**Examples**

data(data_ex_genind)
data(pts_pop_ex)
mat_dist <- suppressWarnings(graph4lg::mat_geo_dist(data = pts_pop_ex,
   ID = "ID",
   x = "x",
   y = "y"))
mat_dist <- mat_dist[order(as.character(row.names(mat_dist))),
   order(as.character(colnames(mat_dist)))]
graph_obs <- gen_graph_thr(mat_w = mat_dist, mat_thr = mat_dist,
   thr = 9500, mode = "larger")
mat_gen <- mat_gen_dist(x = data_ex_genind, dist = "DPS")
graph_pred <- gen_graph_topo(mat_w = mat_gen, mat_topo = mat_dist,
   topo = "gabriel")
res_cor <- graph_node_compar(x = graph_obs, y = graph_pred,
   metrics = c("siw", "siw"), method = "spearman",
   test = TRUE, weight = TRUE)
graph_plan

Create a graph with a minimum planar graph topology

Description

The function constructs a graph with a minimum planar graph topology

Usage

graph_plan(crds, ID = NULL, x = NULL, y = NULL, weight = TRUE)

Arguments

crds  A data.frame with the spatial coordinates of the point set (the graph nodes). It must have three columns:
• ID: A character string indicating the name of the points (graph nodes).
• x: A numeric or integer indicating the longitude of the graph nodes.
• y: A numeric or integer indicating the latitude of the graph nodes.

ID  A character string indicating the name of the column of crds with the point IDs

x  A character string indicating the name of the column of crds with the point longitude

y  A character string indicating the name of the column of crds with the point latitude

weight  A character string indicating whether the links of the graph have weights (TRUE)(default) or not (FALSE)

Details

A delaunay triangulation is performed in order to get the planar graph.

Value

A planar graph of class igraph

Author(s)

P. Savary

Examples

data(pts_pop_ex)
g_plan <- graph_plan(crds = pts_pop_ex, 
  ID = "ID",
  x = "x",
  y = "y")
**graph_plot_compar**

Visualize the topological differences between two spatial graphs on a map

---

**Description**

The function enables to compare two spatial graphs by plotting them highlighting the topological similarities and differences between them. Both graphs should share the same nodes and cannot be directed graphs.

**Usage**

```r
graph_plot_compar(x, y, crds)
```

**Arguments**

- `x`: A graph object of class `igraph`. Its nodes must have the same names as in graph `y`.
- `y`: A graph object of class `igraph`. Its nodes must have the same names as in graph `x`.
- `crds`: A `data.frame` with the spatial coordinates of the graph nodes (both `x` and `y`). It must have three columns:
  - `ID`: Name of the graph nodes (character string). The names must be the same as the node names of the graphs of class `igraph` (`igraph::V(graph)$name`)
  - `x`: Longitude of the graph nodes (numeric or integer).
  - `y`: Latitude of the graph nodes (numeric or integer).

**Details**

The graphs `x` and `y` of class `igraph` must have node names (not necessarily in the same order as IDs in `crds`, given a merging is done).

**Value**

A ggplot2 object to plot

**Author(s)**

P. Savary

**Examples**

```r
data(pts_pop_ex)
data(data_ex_genind)
mat_w <- mat_gen_dist(data_ex_genind, dist = "DPS")
mat_dist <- mat_geo_dist(data = pts_pop_ex,
                         ID = "ID",
```

---
```r
x = "x",
y = "y"
mat_dist <- mat_dist[order(as.character(row.names(mat_dist))),
                   order(as.character(colnames(mat_dist)))]
g1 <- gen_graph_topo(mat_w = mat_w, topo = "mst")
g2 <- gen_graph_topo(mat_w = mat_w, mat_topo = mat_dist, topo = "gabriel")
g <- graph_plot_compar(x = g1, y = g2,
                       crds = pts_pop_ex)
```

---

**graph_topo_compar**

*Compute an index comparing graph topologies*

**Description**

The function computes several indices in order to compare two graph topologies. One of the graph has the "true" topology the other is supposed to reproduce. The indices are then a way to assess the reliability of the latter graph. Both graphs must have the same number of nodes, but not necessarily the same number of links. They must also have the same node names and in the same order.

**Usage**

```r
graph_topo_compar(obs_graph, pred_graph, mode = "mcc", directed = FALSE)
```

**Arguments**

- **obs_graph**: A graph object of class `igraph` with `n` nodes. It is the observed graph that `pred_graph` is supposed to approach.
- **pred_graph**: A graph object of class `igraph` with `n` nodes. It is the predicted graph that is supposed to be akin to `obs_graph`.
- **mode**: A character string specifying which index to compute in order to compare the topologies of the graphs.
  - If `mode = 'mcc'` (default), the Matthews Correlation Coefficient (MCC) is computed.
  - If `mode = 'kappa'`, the Kappa index is computed.
  - If `mode = 'fdr'`, the False Discovery Rate (FDR) is computed.
  - If `mode = 'acc'`, the Accuracy is computed.
  - If `mode = 'sens'`, the Sensitivity is computed.
  - If `mode = 'spec'`, the Specificity is computed.
  - If `mode = 'prec'`, the Precision is computed.
- **directed**: Logical (TRUE or FALSE) specifying whether both graphs are directed or not.
Details

The indices are calculated from a confusion matrix counting the number of links that are in the "observed" graph ("true") and also in the "predicted" graph (true positives : TP), that are not in the "observed" graph but in the "predicted" graph (false negatives : FN), that are not in the "observed" graph but in the "predicted" graph (false positives : FP) and that are not in the "observed" graph and not in the "predicted" graph neither (true negatives: TN). K is the total number of links in the graphs. K is equal to $n \times (n - 1)$ if the graphs are directed and to $\frac{n \times (n - 1)}{2}$ if they are not directed, with $n$ the number of nodes. OP = TP + FN, ON = TN + FP, PP = TP + FP and PN = FN + TN.

The Matthews Correlation Coefficient (MCC) is computed as follows:

$$MCC = \frac{TP \times TN - FP \times FN}{\sqrt{(TP + FP)(TP + FN)(TN + FP)(TN + FN)}}$$

The Kappa index is computed as follows:

$$Kappa = \frac{K \times (TP + TN) - (ON \times PN) - (OP \times PP)}{K^2 - (ON \times PN) - (OP \times PP)}$$

The False Discovery Rate (FDR) is calculated as follows:

$$FDR = \frac{FP}{TP + FP}$$

The Accuracy is calculated as follows:

$$Acc = \frac{TP + TN}{K}$$

The Sensitivity is calculated as follows:

$$Sens = \frac{TP}{TP + FN}$$

The Specificity is calculated as follows:

$$Spec = \frac{TN}{TN + FP}$$

The Precision is calculated as follows:

$$Prec = \frac{TP}{TP + FP}$$

Self loops are not taken into account.

Value

The value of the index computed

Author(s)

P. Savary

References


Examples

```r
data(data_ex_genind)
data(pts_pop_ex)
mat_dist <- suppressWarnings(graph4lg::mat_geo_dist(data=pts_pop_ex,
  ID = "ID",
  x = "x",
  y = "y"))
mat_dist <- mat_dist[order(as.character(row.names(mat_dist))),
  order(as.character(colnames(mat_dist)))]
graph_obs <- gen_graph_thr(mat_w = mat_dist, mat_thr = mat_dist,
```
thr = 15000, mode = "larger")
mat_gen <- mat_gen_dist(x = data_ex_genind, dist = "DPS")
graph_pred <- gen_graph_topo(mat_w = mat_gen, mat_topo = mat_dist,
topo = "gabriel")

graph_topo_compar(obs_graph = graph_obs,
pred_graph = graph_pred,
mode = "mcc",
directed = FALSE)

---

**graph_to_df**

*Convert a graph into a edge list data.frame*

**Description**

The function converts a graph into a edge list data.frame

**Usage**

`graph_to_df(graph, weight = TRUE)`

**Arguments**

- `graph` A graph object of class `igraph`
- `weight` Logical. If TRUE (default), then the column 'link' of the output data.frame contains the weights of the links. If FALSE, it contains only 0 and 1.

**Details**

The 'graph' nodes must have names. Links must have weights if 'weight = TRUE'.

**Value**

An object of class `data.frame` with a link ID, the origin nodes ('from') and arrival nodes ('to') and the link value ('link')(weighted or binary)

**Author(s)**

P. Savary

**Examples**

data(pts_pop_ex)
suppressWarnings(mat_geo <- mat_geo_dist(pts_pop_ex,
ID = "ID",
x = "x",
y = "y"))
g1 <- gen_graph_thr(mat_w = mat_geo,
mat_thr = mat_geo,
thr = 20000)

---


`g1_df <- graph_to_df(g1, weight = TRUE)`

---

**Description**

The function enables to export a spatial graph to shapefile layers.

**Usage**

```r
graph_to_shp(
  graph,
  crds,
  mode = "both",
  crds_crs,
  layer_name,
  dir_path,
  metrics = FALSE
)
```

**Arguments**

- **graph**
  A graph object of class `igraph`

- **crds**
  (if `mode = 'spatial'`) A `data.frame` with the spatial coordinates of the graph nodes. It must have three columns:
  - **ID**: Name of the graph nodes (will be converted into character string). The names must the same as the node names of the graph object of class `igraph` (`igraph::V(graph)$name`)
  - **x**: Longitude (numeric or integer) of the graph nodes in the coordinates reference system indicated with the argument `crds_crs`.
  - **y**: Latitude (numeric or integer) of the graph nodes in the coordinates reference system indicated with the argument `crds_crs`.

- **mode**
  Indicates which shapefile layers will be created
  - If `mode = 'both'` (default), then two shapefile layers are created, one for the nodes and another for the links.
  - If `mode = 'node'`, a shapefile layer is created for the nodes only.
  - If `mode = 'link'`, a shapefile layer is created for the links only.

- **crds_crs**
  A character string indicating the Coordinates Reference System of the spatial coordinates of the nodes and of the shapefile layers created. The projection and datum are given in the PROJ.4 format.

- **layer_name**
  A character string indicating the suffix of the name of the layers to be created.
**gstud_to_genind**

Convert a file from `gstudio` or `popgraph` into a genind object

**Description**

The function converts a file formatted to use `gstudio` or `popgraph` package into a genind object (`adegenet` package)

**Usage**

```
gstud_to_genind(x, pop_col, ind_col = NULL)
```
Arguments

- **x**: An object of class `data.frame` with loci columns in format `locus` (defined in package `gstudio`) with as many rows as individuals and as many columns in format `locus` as there are loci and additional columns.
- **pop_col**: A character string indicating the name of the column with populations’ names in `x`.
- **ind_col**: (optional) A character string indicating the name of the column with individuals’ ID in `x`.

Details

This function uses functions from the `pegas` package. It can handle genetic data where alleles codings do not have same length, (99:101, for example). If the names of the loci include '.' characters, they will be replaced by '_'.

Value

An object of class `genind`.

Author(s)

P. Savary

Examples

```r
data("data_ex_gstud")
x <- data_ex_gstud
pop_col <- "POP"
ind_col <- "ID"
data_genind <- gstud_to_genind(x, pop_col, ind_col)
```

---

**g_percol**

*Prune a graph using the 'percolation threshold' method*

Description

The function allows to prune a graph by removing the links with the largest weights until the graph breaks into two components. The returned graph is the last graph with only one component.

Usage

```r
g_percol(x, val_step = 20)
```

Arguments

- **x**: A symmetric matrix of pairwise distances between nodes.
- **val_step**: The number of classes to create to search for the threshold value without testing all the possibilities. By default, `val_step = 20`.
Kernel param

Description

The function computes the constant parameters of a dispersal kernel with a negative exponential distribution.

Usage

```
kernel_param(p, d_disp, mode = "A")
```

Arguments

- **p**: A numeric value indicating the dispersal probability at a distance equal to 'd_disp' under a negative exponential distribution.
- **d_disp**: A numeric value indicating the distance to which dispersal probability is equal to 'p' under a negative exponential distribution.
- **mode**: A character string indicating the value to return:
  - If 'mode = 'A" (default), the returned value 'alpha' is such that \(\exp(-\alpha \cdot d_{disp}) = p\)
  - If 'mode = 'B", the returned value 'alpha' is such that \(10^{-\alpha \cdot d_{disp}} = p\)

Details

If the resulting parameter when mode = "A" is \(a\) and the resulting parameter when mode = "B" is \(b\), then we have: \(p = \exp(-a \cdot d_{disp}) = 10^{b \cdot d_{disp}}\) and \(a = b \cdot \ln(10)\)

Value

A numeric value
**loci_to_genind**

**Author(s)**

P. Savary

**Examples**

```r
p <- 0.5
d_disp <- 3000
alpha <- kernel_param(p, d_disp, mode = "A")
```

---

**Description**

This function is exactly the same as loci2genind from pegas package

**Usage**

```r
loci_to_genind(x, ploidy = 2, na.alleles = c("NA"))
```

**Arguments**

- `x` An object of class loci to convert
- `ploidy` An integer indicating the ploidy level (by default, 'ploidy = 2')
- `na.alleles` A character vector indicating the coding of the alleles to be treated as missing data (by default, 'na.alleles = c("NA")')

**Value**

An object of class genind

**Author(s)**

P. Savary

**Examples**

```r
data("data_ex_loci")
genind <- loci_to_genind(data_ex_loci, ploidy = 2, na.alleles = "NA")
```
mat_gen_dist

Compute a pairwise matrix of genetic distances between populations

Description

The function computes a pairwise matrix of genetic distances between populations and allows to implement several formula.

Usage

mat_gen_dist(x, dist = "basic", null_val = FALSE)

Arguments

x
An object of class genind that contains the multilocus genotypes (format 'locus') of the individuals as well as their populations.

dist
A character string indicating the method used to compute the multilocus genetic distance between populations

- If `dist = "basic"` (default), then the multilocus genetic distance is computed using a formula of Euclidean genetic distance (Excoffier et al., 1992)
- If `dist = "weighted"`, then the multilocus genetic distance is computed as in Fortuna et al. (2009). It is a Euclidean genetic distance giving more weight to rare alleles
- If `dist = "PG"`, then the multilocus genetic distance is computed as in popgraph::popgraph function, following several steps of PCA and SVD (Dyer et Nason, 2004).
- If `dist = "DPS"`, then the genetic distance used is equal to 1 - the proportion of shared alleles (Bowcock, 1994)
- If `dist = "FST"`, then the genetic distance used is the pairwise FST (Weir et Cockerham, 1984)
- If `dist = "FST_lin"`, then the genetic distance used is the linearised pairwise FST (Weir et Cockerham, 1984)(FST_lin = FST/(1-FST))
- If `dist = "PCA"`, then the genetic distance is computed following a PCA of the matrix of allelic frequencies by population. It is a Euclidean genetic distance between populations in the multidimensional space defined by all the independent principal components.
- If `dist = "GST"`, then the genetic distance used is the G' ST (Hedrick, 2005)
- If `dist = "D"`, then the genetic distance used is Jost’s D (Jost, 2008)

null_val
(optional) Logical. Should negative and null FST, FST_lin, GST or D values be replaced by half the minimum positive value? This option allows to compute Gabriel graphs from these "distances". Default is null_val = FALSE. This option only works if `dist = "FST"` or `FST_lin` or `GST` or `D"`
Details

Negative values are converted into 0. Euclidean genetic distance \( d_{ij} \) between population \( i \) and \( j \) is computed as follows:

\[
d^2_{ij} = \sum_{k=1}^{n} (x_{ki} - x_{kj})^2
\]

where \( x_{ki} \) is the allelic frequency of allele \( k \) in population \( i \) and \( n \) is the total number of alleles. Note that when 'dist = 'weight", the formula becomes

\[
d^2_{ij} = \sum_{k=1}^{n} \left( \frac{1}{(K \ast p_k)} \right) (x_{ki} - x_{kj})^2
\]

where \( K \) is the number of alleles at the locus of the allele \( k \) and \( p_k \) is the frequency of the allele \( k \) in all populations. Note that when 'dist = 'PCA", \( n \) is the number of conserved independent principal components and \( x_{ki} \) is the value taken by the principal component \( k \) in population \( i \).

Value

An object of class \texttt{matrix}

Author(s)

P. Savary

References


Examples

data(data_ex_genind)
x <- data_ex_genind
D <- mat_gen_dist(x = x, dist = "basic")
\textbf{mat\_geo\_dist} \quad \textit{Compute Euclidean geographic distances between points}

\section*{Description}

The function computes Euclidean geographic distance between points given their coordinates in a metric projected Coordinates Reference System.

\section*{Usage}

\begin{verbatim}
mat_geo_dist(data, ID = NULL, x = NULL, y = NULL)
\end{verbatim}

\section*{Arguments}

\begin{itemize}
  \item \texttt{data} \quad An object of class:
    \begin{itemize}
      \item \texttt{data.frame} with 3 columns: 2 columns with the point spatial coordinates and another column with point IDs
      \item \texttt{SpatialPointsDataFrame}
    \end{itemize}
  \item \texttt{ID} \quad (if \texttt{data} is of class \texttt{data.frame}) A character string indicating the name of the column of \texttt{data} with the point IDs
  \item \texttt{x} \quad (if \texttt{data} is of class \texttt{data.frame}) A character string indicating the name of the column of \texttt{data} with the point longitude
  \item \texttt{y} \quad (if \texttt{data} is of class \texttt{data.frame}) A character string indicating the name of the column of \texttt{data} with the point latitude
\end{itemize}

\section*{Details}

Calculate classical Euclidean geographic distance (in m) between two points using Pythagora's theorem

\section*{Value}

A pairwise matrix of geographic distances between points

\section*{Author(s)}

P. Savary

\section*{Examples}

\begin{verbatim}
data(pts_pop_simul)
mat_dist <- mat_geo_dist(data=pts_pop_simul,
                          ID = "ID",
                          x = "x",
                          y = "y")
\end{verbatim}
**mat_pw_dps**

Compute a pairwise genetic distance matrix between populations using Bowcock et al. (1994) formula

---

**Description**

The function computes the pairwise DPS, a genetic distance based on the proportion of shared alleles.

**Usage**

```r
mat_pw_dps(x)
```

**Arguments**

- `x`: An object of class `genind`

**Details**

The formula used is inspired from MSA software:

\[
D_{PS} = 1 - \frac{\sum D \sum K \min(f_{a_{kd}i}, f_{a_{kd}j})}{D}
\]

such as \(a_{kd}\) is the allele \(k\) at locus \(d\), \(D\) is the total number of loci, \(K\) is the allele number at each locus, \(\gamma_{a_{kd}ij} = 0\) if individuals \(i\) and \(j\) do not share allele \(a_{kd}\), \(\gamma_{a_{kd}ij} = 1\) if one of individuals \(i\) and \(j\) has a copy of \(a_{kd}\), \(\gamma_{a_{kd}ij} = 2\) if both individuals have 2 copies of \(a_{kd}\) (homozygotes) \(f_{a_{kd}i}\) is allele \(a_{kd}\) frequency in individual \(i\) (0, 0.5 or 1). More information in: Bowcock et al., 1994 and MSA manual. This function uses functions from adegenet package. Note that in the paper of Bowcock et al. (1994), the denominator is \(2D\). But, in MSA software manual, the denominator is \(D\).

**Value**

A pairwise matrix of genetic distances between populations

**Author(s)**

P. Savary

**References**


**Examples**

```r
data("data_ex_genind")
dist_bowcock <- mat_pw_dps(data_ex_genind)
```
mat_pw_d_j

Compute a pairwise Jost's D matrix between populations

Description

The function computes the pairwise Jost's D matrix between populations from an object of class genind or directly from a GENEPOP file.

Usage

mat_pw_d_j(x, pop_names = NULL)

Arguments

x

An object of class genind, or the character string indicating the path of the GENEPOP file.

pop_names

(optional) A vector of class character of the same length as the number of populations (row and column number in the returned matrix). It contains the name of the populations.

Details

The formula used is that of Jost (2008) This functions uses directly the function diffCalc from diversity. See http://genepop.curtin.edu.au/help_input.html for details on the GENEPOP file format and see Raymond (1995) for detail about GENEPOP software.

Value

A pairwise matrix of G'ST with as many rows and columns as there are populations in the input data.

Warnings

The order of populations matters :

- If x is an object of class genind, individuals are re-ordered by populations and populations are ordered in alphabetic order.
- If x is the path to a GENEPOP file, population order in pop_names must be the same as in the GENEPOP file.

Negative values are converted into 0

Author(s)

P. Savary
References


Examples

data("data_ex_genind")
mat_d_j <- mat_pw_d_j(data_ex_genind)

path_in <- system.file("extdata", "gpop_simul_10_g100_04_20.txt", package = "graph4lg")
file_n <- file.path(tempdir(), "gpop_simul_10_g100_04_20.txt")
file.copy(path_in, file_n, overwrite = TRUE)
mat_pw_d_j(x = file_n, pop_names = as.character(order(as.character(1:10))))
file.remove(file_n)

---

mat_pw_fst

*Compute a pairwise FST matrix between populations*

Description

The function computes the pairwise FST matrix between populations from an object of class genind or directly from a GENEPOP file.

Usage

```r
mat_pw_fst(x, pop_names = NULL)
```

Arguments

- `x` An object of class genind, or the character string indicating the path of the GENEPOP file.
- `pop_names` (optional) A vector of class character of the same length as the number of populations (row and column number in the returned matrix). It contains the name of the populations.

Details

The formula used is that of Weir et Cockerham (1984). This function uses directly the function diffCalc from diveRsity. See [http://genepop.curtin.edu.au/help_input.html](http://genepop.curtin.edu.au/help_input.html) for details on the GENEPOP file format and see Raymond (1995) for detail about GENEPOP software.

Value

A pairwise matrix of FST with as many rows and columns as there are populations in the input data.
Warnings

The order of populations matters:

• If x is an object of class genind, individuals are re-ordered by populations and populations are ordered in alphabetic order.
• If x is the path to a GENEPOP file, population order in pop_names must be the same as in the GENEPOP file.

Negative values are converted into 0

Author(s)

P. Savary

References


Examples

data("data_ex_genind")
mat_d_j <- mat_pw_d_j(data_ex_genind)
path_in <- system.file('extdata', 'gpop_simul_10_g100_04_20.txt', package = 'graph4lg')
file_n <- file.path(tempdir(), "gpop_simul_10_g100_04_20.txt")
file.copy(path_in, file_n, overwrite = TRUE)
mat_pw_fst(x = file_n, pop_names = as.character(order(as.character(1:10))))
file.remove(file_n)

mat_pw_gst

*Compute a pairwise G’ST matrix between populations*

Description

The function computes the pairwise G’ST matrix between populations from an object of class genind or directly from a GENEPOP file.

Usage

mat_pw_gst(x, pop_names = NULL)

Arguments

x An object of class genind, or the character string indicating the path of the GENEPOP file.

pop_names (optional) A vector of class character of the same length as the number of populations (row and column number in the returned matrix). It contains the name of the populations.
Details

The formula used is that of Hedrick (2005) This functions uses directly the function diffCalc from diveRsity. See [http://genepop.curtin.edu.au/help_input.html](http://genepop.curtin.edu.au/help_input.html) for details on the GENEPOP file format and see Raymond (1995) for detail about GENEPOP software.

Value

A pairwise matrix of G'ST with as many rows and columns as there are populations in the input data.

Warnings

The order of populations matters:

- If x is an object of class genind, individuals are re-ordered by populations and populations are ordered in alphabetic order.
- If x is the path to a GENEPOP file, population order in pop_names must be the same as in the GENEPOP file.

Negative values are converted into 0

Author(s)

P. Savary

References


Examples

data("data_ex_genind")
mat_gst <- mat_pw_gst(data_ex_genind)
path_in <- system.file("extdata", "gpop_simul_10_g100_04_20.txt", 
package = "graph4lg")
file_n <- file.path(tempdir(), "gpop_simul_10_g100_04_20.txt")
file.copy(path_in, file_n, overwrite = TRUE)
mat_pw_gst(x = file_n, pop_names = as.character(order(as.character(1:10))))
file.remove(file_n)
The function enables to plot graphs, whether spatial or not.

Usage

```r
plot_graph_lg(
  graph,
  crds,
  mode = "spatial",
  weight = TRUE,
  width = "w",
  pts_col = "#F2B950"
)
```

Arguments

- **graph**: A graph object of class `igraph`
- **crds**: (if `mode = 'spatial'`) A `data.frame` with the spatial coordinates of the graph nodes. It must have three columns:
  - ID: A character string indicating the name of the graph nodes. The names must be the same as the node names of the graph of class `igraph` (`igraph::V(graph)$name`)
  - x: A numeric or integer indicating the longitude of the graph nodes.
  - y: A numeric or integer indicating the latitude of the graph nodes.
- **mode**: A character string indicating whether the graph is spatial (`'mode = 'spatial'` (default)) or not (`'mode = 'aspatial'`)
- **weight**: A character string indicating whether the links of the graph have weights (TRUE)(default) or not (FALSE)
- **width**: A character string indicating whether the width of the link should be proportional to link weights ("w", default) or to the inverse of link weights ("inv", convenient with distances)
- **pts_col**: (optional) A character string indicating the color used to plot the nodes (default: "#F2B950"). It must be a hexadecimal color code or a color used by default in R.

Details

When the graph is not spatial (`'mode = 'aspatial'`), the nodes coordinates are calculated with Fruchterman et Reingold algorithm. The graph object x of class `igraph` must have node names (not necessarily in the same order as IDs in crds, given a merging is done).
**Value**

A ggplot2 object to plot

**Author(s)**

P. Savary

**References**


**Examples**

```r
data(pts_pop_ex)
data(data_ex_genind)
mat_w <- mat_gen_dist(data_ex_genind, dist = "DPS")
gp <- gen_graph_topo(mat_w = mat_w, topo = "mst")
g <- plot_graph_lg(graph = gp, 
  crds = pts_pop_ex, 
  mode = "spatial", 
  weight = TRUE)
```

---

**plot_graph_modul**  
*Plot the graphs making visible their partition into modules*

**Description**

The function computes a partition of the graph into modules and plots the graph colouring the nodes with colors corresponding to their respective modules.

**Usage**

```r
plot_graph_modul(
  graph, 
  algo = "fast_greedy", 
  nb_modul = NULL, 
  weight = "inv", 
  crds, 
  mode = "spatial", 
  weight_plot = TRUE, 
  width = "inv"
)
```
plot_graph_modul

Arguments

- **graph**: A graph object of class `igraph` to partition into modules and to plot. The graph must be undirected. If `crds` is not NULL, then the graph nodes must have names corresponding to the ID column of `crds`.

- **algo**: (if x and y and graph objects) A character string indicating the algorithm used to create the modules with *igraph*:  
  - If `algo = 'fast_greedy'` (default), function `cluster_fast_greedy` from *igraph* is used (Clauset et al., 2004).
  - If `algo = 'walktrap'`, function `cluster_walktrap` from *igraph* is used (Pons et Latapy, 2006) with 4 steps (default options).
  - If `algo = 'louvain'`, function `cluster_louvain` from *igraph* is used (Blondel et al., 2008). In that case, the number of modules created in each graph is imposed.
  - If `algo = 'optimal'`, function `cluster_optimal` from *igraph* is used (Brandes et al., 2008) (can be very long). In that case, the number of modules created in each graph is imposed.

- **nb_modul**: Numeric value indicating the number of modules to create.

- **weight**: (optional) A character string or character vector indicating how to weight graph links during the calculation of the modularity.  
  - If `weight = 'inv'` (default), then links are weighted with the inverse values of their initial weights.
  - If `weight = 'w'`, then links are weighted with their initial weights values.
  - If `weight = 'none'`, then links are not weighted during the calculation.

- **crds**: (if `mode = 'spatial'`) A `data.frame` with the spatial coordinates of the graph nodes. It must have three columns: 
  - ID: A character string indicating the name of the graph nodes. The names must be the same as the node names of the graph of class `igraph` (`igraph::V(graph)$name`).
  - x: A numeric or integer indicating the longitude of the graph nodes.
  - y: A numeric or integer indicating the latitude of the graph nodes.

- **mode**: A character string indicating whether the graph is spatial (`'mode = 'spatial'`) (default) or not (`'mode = 'aspatial'`).

- **weight_plot**: Logical indicating whether the links of the graph have to be displayed on the plot.

- **width**: Logical indicating whether the width of the links on the plot should be proportional to link weights ("w", default) or to the inverse of link weights ("inv", convenient with distances)

Details

When the graph is not spatial (`'mode = 'aspatial'`), the nodes coordinates are calculated with Fruchterman et Reingold algorithm.
Value

A ggplot2 object to plot

Author(s)

P. Savary

References


Examples

data(data_simul_genind)
data(pts_pop_simul)
mat_dist <- suppressWarnings(graph4lg::mat_geo_dist(data=pts_pop_simul,
    ID = "ID",
    x = "x",
    y = "y"))
mat_dist <- mat_dist[order(as.character(row.names(mat_dist))),
    order(as.character(colnames(mat_dist)))]
graph <- gen_graph THR(mat_w = mat_dist, mat_thr = mat_dist,
    thr = 9500, mode = "larger")
plot_graph_modul(graph = graph, crds = pts_pop_simul)

plot_w_hist

Plot histograms of link weights

Description

The function enables to plot histogram to visualize the distribution of the link weights

Usage

plot_w_hist(graph, fill = "#396D35", class_width = NULL)
pop_gen_index

Arguments

graph A graph object of class igraph whose links are weighted

fill A character string indicating the color used to fill the bars (default: 
    
"#396D35"). It must be a hexadecimal color code or a color used by default in R.

class_width (default values: NULL) A numeric or an integer specifying the width of the 
classes displayed on the histogram. When it is not specified, the width is equal 
to the difference between the minimum and maximum values divided by 80.

Value

A ggplot2 object to plot

Author(s)

P. Savary

Examples

data(data_ex_genind)
mat_w <- mat_gen_dist(data_ex_genind, dist = "DPS")
gp <- gen_graph_topo(mat_w = mat_w, topo = "gabriel")
hist <- plot_w_hist(gp)

Description

The function computes population-level genetic indices from an object of class genind.

Usage

pop_gen_index(x, pop_names = NULL, indices = c("Nb_ind", "A", "He", "Ho"))

Arguments

x An object of class genind from package adegenet.

pop_names (optional) A character vector indicating population names. It is of the same 
length as the number of populations. Without this argument, populations are 
given the names they have initially in the 'genind' object (which is sometimes 
only a number). The order of the population names must match with their order 
in the 'genind' object. The function does not reorder them. Users must be 
careful.

indices (optional) A character vector indicating the population-level indices to compute. 
These indices can be:
  • Mean allelic richness by locus by population (indices = c("A",...))
• Mean expected heterozygosity by locus by population (indices = c("He", ...))
• Mean observed heterozygosity by locus by population (indices = c("Ho", ...))
• Number of individuals by population (indices = c("Nb_ind", ...))
• Total allelic richness by population (indices = c("A_tot", ...))
By default, indices = c("Nb_ind", "A", "He", "Ho").

Value
An object of class data.frame whose rows correspond to populations and columns to population attributes (ID, size, genetic indices). By default, the first column corresponds to the population names (ID). The order of the columns depends on the vector ‘indices’.

Author(s)
P. Savary

Examples
```r
data(data_ex_genind)
x <- data_ex_genind
pop_names <- levels(x@pop)
df_pop_indices <- pop_gen_index(x = x,
    pop_names = pop_names,
    indices = c("Nb_ind", "A"))
```

Description
Simulation dataset 10 populations located on a simulated landscape

Usage
```r
pts_pop_ex
```

Format
An object of class 'data.frame' with the following columns:

<table>
<thead>
<tr>
<th>Column</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>ID</td>
<td>Population ID of the 10 populations</td>
</tr>
<tr>
<td>x</td>
<td>Site longitude (RGF93)</td>
</tr>
<tr>
<td>y</td>
<td>Site latitude (RGF93)</td>
</tr>
</tbody>
</table>

References
Landguth EL, Cushman S (2010). “CDPOP: a spatially explicit cost distance population genetics program.” *Molecular Ecology Resources*, **10**(1), 156–161. There are as many rows as there are sampled populations.
Examples

data("pts_pop_ex")
str(pts_pop_ex)

Description

Simulation dataset 50 populations located on a simulated landscape

Usage

pts_pop_simul

Format

An object of class 'data.frame' with the following columns:

- ID  Population ID of the 50 populations
- x   Site longitude (RGF93)
- y   Site latitude (RGF93)

References

Landguth EL, Cushman S (2010). “CDPOP: a spatially explicit cost distance population genetics program.” Molecular Ecology Resources, 10(1), 156–161. There are as many rows as there are sampled populations.

Examples

data("pts_pop_simul")
str(pts_pop_simul)

pw_mat_to_df

Convert a pairwise matrix into an edge-list data.frame

Description

The function converts a pairwise matrix into an edge-list data.frame

Usage

pw_mat_to_df(pw_mat)
reorder_mat

Arguments

- pw_mat: A pairwise matrix of class matrix. It must have the same row names and column names. If values represent distances, diagonal elements should be equal to 0.

Value

- An object of class data.frame

Author(s)

- P. Savary

Examples

```r
data(data_tuto)
pw_mat <- data_tuto[[1]]
df <- pw_mat_to_df(pw_mat)
```

reorder_mat

Reorder the rows and columns of a symmetric matrix

Description

The function reorders the rows and columns of a symmetric matrix according to a specified order.

Usage

```r
reorder_mat(mat, order)
```

Arguments

- mat: An object of class matrix
- order: A character vector with the rows and columns names of the matrix in the order in which they will be ordered by the function. All its elements must be rows and columns names of the matrix mat.

Details

The matrix mat must be symmetric and have rows and columns names. Its values are not modified.

Value

- A reordered symmetric matrix

Author(s)

- P. Savary
Examples

mat <- matrix(rnorm(36), 6)
mat[lower.tri(mat)] <- t(mat)[lower.tri(mat)]
row.names(mat) <- colnames(mat) <- c("A", "C", "E", "B", "D", "F")
order <- c("A", "B", "C", "D", "E", "F")
mat <- reorder_mat(mat = mat, order = order)

scatter_dist(mat_gd, mat_ld, method = "loess", thr_gd = NULL, thr_ld = NULL, se = TRUE, smooth_col = "black", pts_col = "#999999")

Description

The function enables to plot scatterplots to visualize the relationship between genetic distance (or differentiation) and landscape distance (Euclidean distance, cost-distance, etc.) between populations or sample sites.

Usage

scatter_dist(mat_gd, mat_ld, method = "loess", thr_gd = NULL, thr_ld = NULL, se = TRUE, smooth_col = "black", pts_col = "#999999")

Arguments

mat_gd A symmetric matrix with pairwise genetic distances between populations or sample sites.
mat_ld A symmetric matrix with pairwise landscape distances between populations or sample sites. These distances can be Euclidean distances, cost-distances or resistance distances, among others.
method A character string indicating the smoothing method used to fit a line on the scatterplot. Possible values are the same as with function 'geom_smooth()' from ggplot2: 'lm', 'glm', 'gam', 'loess' (default).
thr_gd (optional) A numeric value used to remove values from the data before to plot. All genetic distances values above thr_gd are removed from the data.
thr_ld (optional) A numeric value used to remove values from the data before to plot. All landscape distances values above thr_ld are removed from the data.
se Logical (optional, default = TRUE) indicating whether the confidence interval around the smooth line is displayed.
**scatter_dist_g**

- **smooth_col**: (optional) A character string indicating the color used to plot the smoothing line (default: "blue"). It must be a hexadecimal color code or a color used by default in R.
- **pts_col**: (optional) Character string indicating the color used to plot the points (default: "#999999"). It must be a hexadecimal color code or a color used by default in R.

**Details**

IDs in mat_gd and mat_ld must be the same and refer to the same sampling sites or populations, and both matrices must be ordered in the same way. Matrix of genetic distance mat_gd can be computed using `mat_gen_dist`. Matrix of landscape distance mat_ld can be computed using `mat_geo_dist` when the landscape distance needed is a Euclidean geographical distance.

**Value**

A ggplot2 object to plot

**Author(s)**

P. Savary

**Examples**

```r
data(data_tuto)
mat_dps <- data_tuto[[1]]
mat_dist <- suppressWarnings(mat_geo_dist(data = pts_pop_simul, 
  ID = "ID", 
  x = "x", 
  y = "y"))
mat_dist <- mat_dist[order(as.character(row.names(mat_dist))), 
  order(as.character(colnames(mat_dist)))]
scatterplot_ex <- scatter_dist(mat_gd = mat_dps, 
  mat_ld = mat_dist)
```

---

**scatter_dist_g**

*Plot scatterplots of distances to visualize the graph pruning intensity*

**Description**

The function enables to plot scatterplots of the relationship between two distances (often a genetic distance and a landscape distance between populations or sample sites), while highlighting the population pairs between which a link was conserved during the creation of a graph whose nodes are populations (or sample sites). It thereby allows to visualize the graph pruning intensity.
scatter_dist_g

Usage

```r
scatter_dist_g(
  mat_y,
  mat_x,
  graph,
  thr_y = NULL,
  thr_x = NULL,
  pts_col_1 = "#999999",
  pts_col_2 = "black"
)
```

Arguments

- **mat_y**: A symmetric (complete) matrix with pairwise (genetic or landscape) distances between populations or sample sites. These values will be the point coordinates on the y axis. `mat_y` is the matrix used to weight the links of the graph `x`, whose nodes correspond to row and column names of `mat_y`.

- **mat_x**: A symmetric (complete) matrix with pairwise (genetic or landscape) distances between populations or sample sites. These values will be the point coordinates on the x axis. `mat_x` and `mat_y` must have the same row and column names, ordered in the same way.

- **graph**: A graph object of class `igraph`. Its nodes must have the same names as the row and column of `mat_y` and `mat_x` matrices. `x` must have weighted links. Link weights have to be values from `mat_y` matrix. `graph` must be an undirected graph.

- **thr_y**: (optional) A numeric value used to remove values from the data before to plot. All values from `mat_y` above `thr_y` are removed from the data.

- **thr_x**: (optional) A numeric value used to remove values from the data before to plot. All values from `mat_x` above `thr_x` are removed from the data.

- **pts_col_1**: (optional) A character string indicating the color used to plot the points associated to all populations or sample sites pairs (default: "#999999"). It must be a hexadecimal color code or a color used by default in R.

- **pts_col_2**: (optional) A character string indicating the color used to plot the points associated to populations or sample sites pairs connected on the graph (default: "black"). It must be a hexadecimal color code or a color used by default in R.

Details

IDs in `mat_y` and `mat_x` must be the same and refer to the same sampling sites or populations, and both matrices must be ordered in the same way. Matrices of genetic distance can be computed using `mat_gen_dist`. Matrices of landscape distance can be computed using `mat_geo_dist` when the landscape distance needed is a Euclidean geographical distance. This function is based upon `scatter_dist` function.

Value

A ggplot2 object to plot
structure_to_genind

Author(s)

P. Savary

Examples

data(data_tuto)
mat_gen <- data_tuto[[1]]
mat_dist <- suppressWarnings(mat_geo_dist(data=pts_pop_simul,
   ID = "ID",
   x = "x",
   y = "y"))
mat_dist <- mat_dist[order(as.character(row.names(mat_dist))),
    order(as.character(colnames(mat_dist)))]
x <- gen_graph_topo(mat_w = mat_gen, mat_topo = mat_dist, topo = "gabriel")
scat <- scatter_dist_g(mat_y = mat_gen, mat_x = mat_dist,
   graph = x)

structure_to_genind   Convert a file in STRUCTURE format into a genind object

Description

The function converts a text file in STRUCTURE format into a genind object to use in R

Usage

structure_to_genind(
  path,
  pop_names = NULL,
  loci_names = NULL,
  ind_names = NULL
)

Arguments

path  A character string indicating the path to the STRUCTURE file in format .txt, or alternatively the name of the file in the working directory. The STRUCTURE file must only have:
   • A first column with the IDs of the individuals (can be a simple number)
   • A second column with the IDs of the populations (can be a simple number)
   • Some loci columns : as many columns as loci in the data

The row for loci names is optional but recommended. Each individual is displayed on 2 rows.

pop_names   (optional) A character vector indicating the population names in the same order as in the STRUCTURE file. It is of the same length as the number of populations. Without this argument, populations are numbered from 1 to the total number of individuals.
loci_names: A character vector with the names of the loci if not specified in the file first row. This argument is mandatory if the STRUCTURE file does not include the names of the loci in the first row. In other cases, the names of the loci is extracted from the file first row.

ind_names: (optional) A character vector indicating the individual names in the same order as in the STRUCTURE file. It is of the same length as the number of individuals. Without this argument, individuals are numbered from 1 to the total number of individuals.

Details

The column order of the resulting object can be different from that of objects returned by `gstud_to_genind` and `genepop_to_genind`, depending on allele and loci coding. This function uses functions from the `pegas` package. For details about STRUCTURE file format: STRUCTURE user manual.

Value

An object of type genind.

Author(s)

P. Savary

Examples

```r
data("data_ex_genind")
loci_names <- levels(data_ex_genind@loc.fac)
pop_names <- levels(data_ex_genind@pop)
ind_names <- row.names(data_ex_genind@tab)
path_in <- system.file("extdata", "data_ex_str.txt", package = "graph4lg")
file_n <- file.path(tempdir(), "data_ex_str.txt")
file.copy(path_in, file_n, overwrite = TRUE)
str <- structure_to_genind(path = file_n, loci_names = loci_names, pop_names = pop_names, ind_names = ind_names)
file.remove(file_n)
```
Index

*Topic datasets
  data_ex_genind, 7
  data_ex_gstud, 8
  data_ex_loci, 8
  data_simul_genind, 9
  data_tuto, 10
  pts_pop_ex, 51
  pts_pop_simul, 52

add_nodes_attr, 3
compar_r_fisher, 4
convert_cd, 5
cor, 27

data_ex_genind, 7
data_ex_gstud, 8
data_ex_loci, 8
data_simul_genind, 9
data_tuto, 10
df_to_pw_mat, 10
dist_max_corr, 11

g_percol, 35
gen_graph_indep, 16
gen_graph_thr, 19
gen_graph_topo, 20
genepop_to_genind, 13, 16, 58
genind2genpop, 15
genind_to_genepop, 14, 15
graph_modul_compar, 23
graph_node_compar, 26
graph_plan, 28
graph_plot_compar, 29
graph_to_df, 32
graph_to_shp, 33
graph_topo_compar, 30
graphab_to_igraph, 4, 21
gstud_to_genind, 34, 58
kernel_param, 36

loci_to_genind, 37
mantel, 13
mat_gen_dist, 12, 38, 55, 56
mat_geo_dist, 6, 13, 40, 55, 56
mat_pw_d_j, 42
mat_pw_dps, 41
mat_pw_fst, 14, 16, 43
mat_pw_gst, 44
plot_graph_lg, 22, 46
plot_graph_modul, 47
plot_w_hist, 49
pop_gen_index, 50
pts_pop_ex, 51
pts_pop_simul, 52
pw_mat_to_df, 52
reorder_mat, 53
scatter_dist, 54, 56
scatter_dist_g, 55
structure_to_genind, 57

59