

Package ‘shar’

May 4, 2021

Type Package

Title Species-Habitat Associations

Version 1.2.1

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Description Analyse species-habitat associations in R. Therefore, information about the location of the species is needed and about the environmental conditions. To test for significance habitat associations, one of the two components is randomized. Methods are mainly based on Plotkin et al. (2000) <doi:10.1006/jtbi.2000.2158> and Harms et al. (2001) <doi:10.1111/j.1365-2745.2001.00615.x>.

License GPL (>= 3)

URL <https://r-spatialecology.github.io/shar/>

BugReports <https://github.com/r-spatialecology/shar/issues/>

Depends R (>= 3.1)

Imports classInt, graphics, grDevices, methods, raster, spatstat.core, spatstat.geom, stats, utils, Rcpp

RoxygenNote 7.1.1

Suggests covr, dplyr, testthat (>= 2.1.0), knitr, rmarkdown, spatstat

VignetteBuilder knitr

Encoding UTF-8

LazyData true

LinkingTo Rcpp

NeedsCompilation yes

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Repository CRAN

Date/Publication 2021-05-04 14:50:02 UTC

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calculate_energy	<i>calculate_energy</i>
------------------	-------------------------

Description

Calculate mean energy

Usage

```
calculate_energy(
  pattern,
  weights = c(0.5, 0.5),
  return_mean = FALSE,
  comp_fast = 1000,
  verbose = TRUE
)
```

Arguments

pattern	List with reconstructed patterns.
weights	Weights used to calculate energy. The first number refers to Gest(r), the second number to pcf(r).
return_mean	Return the mean energy.
comp_fast	If pattern contains more points than threshold, summary functions are estimated in a computational fast way.
verbose	Print progress report.

Details

The function calculates the mean energy (or deviation) between the observed pattern and all reconstructed patterns (for more information see Tscheschel & Stoyan (2006) or Wiegand & Moloney (2014)). The pair correlation function and the nearest neighbour distance function are used to describe the patterns. For large patterns ‘comp_fast = TRUE’ decreases the computational demand because no edge correction is used and the pair correlation function is estimated based on Ripley’s K-function. For more information see [estimate_pcf_fast](#).

Value

numeric

References

- Tscheschel, A., & Stoyan, D. (2006). Statistical reconstruction of random point patterns. *Computational Statistics and Data Analysis*, 51(2), 859-871.
- Wiegand, T., & Moloney, K. A. (2014). *Handbook of spatial point-pattern analysis in ecology*. Boca Raton: Chapman and Hall/CRC Press.

See Also

[plot_energy](#)
[reconstruct_pattern_homo](#)
[reconstruct_pattern_hetero](#)
[reconstruct_pattern_cluster](#)
[plot_randomized_pattern](#)

Examples

```
pattern_random <- fit_point_process(species_a, n_random = 19)
calculate_energy(pattern_random)
calculate_energy(pattern_random, return_mean = TRUE)

## Not run:
marks_sub <- spatstat.geom::subset.ppp(species_a, select = dbh)
marks_recon <- reconstruct_pattern_marks(pattern_random$randomized[[1]], marks_sub,
n_random = 19, max_runs = 1000)
calculate_energy(marks_recon, return_mean = FALSE)
```

```
## End(Not run)
```

classify_habitats	<i>classify_habitats</i>
-------------------	--------------------------

Description

Classify habitats

Usage

```
classify_habitats(raster, classes = 5, style = "fisher")
```

Arguments

raster	RasterLayer.
classes	Number of classes.
style	Style of classification.

Details

Classifies a RasterLayer with continuous values into n discrete classes. Consequently, classes are non-overlapping (and left-closed). For more information see 'classIntervals'.

Value

RasterLayer

References

Armstrong, M. P., Xiao, N., Bennett, D. A., 2003. "Using genetic algorithms to create multicriteria class intervals for choropleth maps". *Annals, Association of American Geographers*, 93 (3), 595-623

Jenks, G. F., Caspall, F. C., 1971. "Error on choroplethic maps: definition, measurement, reduction". *Annals, Association of American Geographers*, 61 (2), 217-244

Dent, B. D., 1999, *Cartography: thematic map design*. McGraw-Hill, Boston, 417 pp. Slocum TA, McMaster RB, Kessler FC, Howard HH 2005 *Thematic Cartography and Geographic Visualization*, Prentice Hall, Upper Saddle River NJ.

Fisher, W. D. 1958 "On grouping for maximum homogeneity", *Journal of the American Statistical Association*, 53, 789-798

See Also

[classIntervals](#)

Examples

```
landscape_classified <- classify_habitats(landscape, classes = 5)
```

`create_neighbourhood` *create_neighbourhood*

Description

Create neighbourhood

Usage

```
create_neighbourhood(cells, matrix, directions = 4)
```

Arguments

<code>cells</code>	matrix with cell ids of focal cells.
<code>matrix</code>	matrix in which cells are located.
<code>directions</code>	Cells neighbour rule: 4 (rook's case), 8 (queen's case).

Details

Get cell ids of all neighbouring cells. The neighbourhood rule can be specified and is either rook's case (4 neighbours) or queen's case (8 neighbours).

Value

matrix

See Also

[randomize_raster](#)

Examples

```
mat <- matrix(1, nrow= 10, ncol = 10)
cell_id <- rbind(cbind(3,5), cbind(7,1))
create_neighbourhood(cell_id, mat)
```

estimate_pcf_fast *estimate_pcf_fast*

Description

Fast estimation of the pair correlation function

Usage

```
estimate_pcf_fast(pattern, ...)
```

Arguments

pattern	Point pattern.
...	Arguments passed down to 'Kest' or 'pcf.fv'.

Details

The functions estimates the pair correlation functions based on an estimation of Ripley's K-function. This makes it computationally faster than estimating the pair correlation function directly. It is a wrapper around 'Kest' and 'pcf.fv'.

Value

fv.object

References

- Ripley, B.D. (1977) Modelling spatial patterns (with discussion). *Journal of the Royal Statistical Society, Series B*, 39, 172-212.
- Stoyan, D, Kendall, W.S. and Mecke, J. (1995) *Stochastic geometry and its applications*. 2nd edition. Springer Verlag.
- Stoyan, D. and Stoyan, H. (1994) *Fractals, random shapes and point fields: methods of geometrical statistics*. John Wiley and Sons.

See Also

[Kest](#)
[pcf.fv](#)

Examples

```
pcf_species_b <- estimate_pcf_fast(species_a)
```

extract_points	<i>extract_points</i>
----------------	-----------------------

Description

Extract points

Usage

```
extract_points(raster, pattern)
```

Arguments

raster	RasterLayer.
pattern	Point pattern.

Details

The function extracts the number of points within each habitat.

Value

data.frame

Examples

```
landscape_classified <- classify_habitats(landscape, classes = 5)
extract_points(raster = landscape_classified, pattern = species_b)
```

fit_point_process	<i>fit_point_process</i>
-------------------	--------------------------

Description

Create random patterns by point process fitting

Usage

```
fit_point_process(
  pattern,
  n_random = 1,
  process = "poisson",
  return_input = TRUE,
  simplify = FALSE,
  verbose = TRUE
)
```

Arguments

pattern	List with reconstructed patterns.
n_random	Number of randomized RasterLayers.
process	What point process to use. Either 'poisson' or 'cluster'.
return_input	The original input data is returned as last list entry.
simplify	If n_random = 1 and return_input = FALSE only pattern will be returned.
verbose	Print progress report.

Details

The functions randomizes the observed pattern by fitting a point process to the data. It is possible to choose between a Poisson process or a Thomas cluster process.

Value

list

References

Plotkin, J. B., Potts, M. D., Leslie, N., Manokaran, N., LaFrankie, J. V., & Ashton, P. S. (2000). Species-area curves, spatial aggregation, and habitat specialization in tropical forests. *Journal of Theoretical Biology*, 207(1), 81-99.

Examples

```
pattern_fitted <- fit_point_process(pattern = species_a, n_random = 39)
```

gamma_test

Gamma test

Description

Randomized data for species b using the gamma test.

Usage

```
gamma_test
```

Format

rd_pat object.

landscape	<i>Example landscape (random cluster neutral landscape model).</i>
-----------	--

Description

An example map to show landscapetools functionality generated with the 'nlm_fbm()' algorithm.

Usage

```
landscape
```

Format

A raster layer object.

Source

Simulated neutral landscape model with R. <https://github.com/ropensci/NLMR/>

plot_energy	<i>plot_energy</i>
-------------	--------------------

Description

Plot energy of pattern reconstruction

Usage

```
plot_energy(pattern, col = NULL)
```

Arguments

pattern	List with reconstructed patterns.
col	Vector with colors. Must be as long as n_random.

Details

The function plots the decrease of the energy over time, i.e. the iterations. This can help to identify if enough max_runs where chosen for the reconstruction.

See Also

[calculate_energy](#)
[reconstruct_pattern_homo](#)
[reconstruct_pattern_hetero](#)
[reconstruct_pattern_cluster](#)
[plot_randomized_pattern](#)

Examples

```
## Not run:
pattern_recon <- reconstruct_pattern_homo(species_a, n_random = 3, max_runs = 1000)
plot_energy(pattern_recon)

marks_sub <- spatstat.geom::subset.ppp(species_a, select = dbh)
marks_recon <- reconstruct_pattern_marks(pattern_recon$randomized[[1]], marks_sub,
n_random = 1, max_runs = 1000)
plot_energy(marks_recon)

## End(Not run)
```

```
plot_randomized_pattern
      plot_randomized_pattern
```

Description

Plot randomized pattern

Usage

```
plot_randomized_pattern(
  pattern,
  what = "sf",
  probs = c(0.025, 0.975),
  comp_fast = 1000,
  ask = TRUE,
  verbose = TRUE
)
```

Arguments

pattern	List with reconstructed patterns.
what	Plot summary functions of point patterns (what = "sf") or actual patterns (what = "pp").
probs	Quantiles of randomized data used for envelope construction.
comp_fast	If pattern contains more points than threshold, summary functions are estimated in a computational fast way.
ask	If TRUE the user is asked to press <RETURN> before second summary function is plotted (only has influence if what = "sf" and method = "spatial").
verbose	Print progress report.

Details

The function plots the pair correlation function and the nearest neighbour function the observed pattern and the reconstructed patterns (as "simulation envelopes"). For large patterns `comp_fast = TRUE` decreases the computational demand because no edge correction is used and the pair correlation function is estimated based on Ripley's K-function. For more information see [estimate_pcf_fast](#). It is also possible to plot 3 randomized patterns and the observed pattern using `what = "pp"`.

Examples

```
pattern_random <- fit_point_process(species_a, n_random = 19, process = "cluster")
plot_randomized_pattern(pattern_random)

plot_randomized_pattern(pattern_random, what = "pp")

## Not run:
marks_sub <- spatstat.geom::subset.ppp(species_a, select = dbh)
marks_recon <- reconstruct_pattern_marks(pattern_random$randomized[[1]],
marks_sub, n_random = 19, max_runs = 1000)
plot_randomized_pattern(marks_recon)

## End(Not run)
```

```
plot_randomized_raster
      plot_randomized_raster
```

Description

Plot randomized raster

Usage

```
plot_randomized_raster(raster, n = NULL, col, verbose = TRUE, nrow, ncol)
```

Arguments

<code>raster</code>	List with randomized raster
<code>n</code>	Number of randomized rasters to plot. See details for more information.
<code>col</code>	Color palette used for plotting.
<code>verbose</code>	Print messages.
<code>nrow, ncol</code>	Number of rows and columns.

Details

Function to plot randomized rasters. If `n` is only a single number, `n` randomized rasters will be sampled. If `n` is a vector, the corresponding rasters will be plotted.

Value

plot

Examples

```
## Not run:
landscape_classes <- classify_habitats(raster = landscape, classes = 5)
landscape_random <- randomize_raster(raster = landscape_classes, n_random = 19)

plot_randomized_raster(landscape_random)

palette <- viridis::viridis(n = 5)
plot_randomized_raster(landscape_random, n = 5, col = palette, nrow = 3, ncol = 2)

## End(Not run)
```

print.rd_mar

print.rd_mar

Description

Print method for rd_mar object

Usage

```
## S3 method for class 'rd_mar'
print(x, digits = 4, ...)
```

Arguments

x	Random patterns.
digits	Number of decimal places (round).
...	Arguments passed to cat

Details

Printing method for random patterns created with [reconstruct_pattern_marks](#).

See Also

[reconstruct_pattern_marks](#)

Examples

```
## Not run:
pattern_recon <- reconstruct_pattern_homo(species_a, n_random = 1, max_runs = 1000,
simplify = TRUE, return_input = FALSE)
marks_sub <- spatstat.geom::subset.ppp(species_a, select = dbh)
marks_recon <- reconstruct_pattern_marks(pattern_recon, marks_sub, n_random = 19, max_runs = 1000)
print(marks_recon)

## End(Not run)
```

print.rd_pat	<i>print.rd_pat</i>
--------------	---------------------

Description

Print method for rd_pat object

Usage

```
## S3 method for class 'rd_pat'
print(x, digits = 4, ...)
```

Arguments

x	Random patterns.
digits	Number of decimal places (round).
...	Arguments passed to cat

Details

Printing method for random patterns created with [reconstruct_pattern_homo](#), [reconstruct_pattern_hetero](#), [reconstruct_pattern_cluster](#) or [fit_point_process](#).

See Also

[reconstruct_pattern_homo](#)
[reconstruct_pattern_hetero](#)
[reconstruct_pattern_cluster](#)
[fit_point_process](#)

Examples

```
pattern_random <- fit_point_process(species_a, n_random = 199)
print(pattern_random)

## Not run:
pattern_recon <- reconstruct_pattern_hetero(species_b, n_random = 19, max_runs = 1000)
print(pattern_recon)

## End(Not run)
```

<code>print.rd_ras</code>	<i>print.rd_ras</i>
---------------------------	---------------------

Description

Print method for `rd_ras` object

Usage

```
## S3 method for class 'rd_ras'
print(x, ...)
```

Arguments

<code>x</code>	Random patterns.
<code>...</code>	Arguments passed to <code>cat</code>

Details

Printing method for random patterns created with [randomize_raster](#).

See Also

[randomize_raster](#)

Examples

```
## Not run:
landscape_classified <- classify_habitats(landscape, classes = 5)
landscape_random <- randomize_raster(landscape_classified, n_random = 19)

print(landscape_random)

## End(Not run)
```

randomize_raster *randomize_raster*

Description

Randomization algorithm

Usage

```
randomize_raster(  
  raster,  
  n_random = 1,  
  directions = 4,  
  return_input = TRUE,  
  simplify = FALSE,  
  verbose = TRUE  
)
```

Arguments

raster	RasterLayer.
n_random	Number of randomizations.
directions	Cells neighbour rule: 4 (rook's case), 8 (queen's case).
return_input	The original input data is returned as last list entry
simplify	If n_random = 1 and return_input = FALSE only raster will be returned.
verbose	Print progress report.

Details

The function randomizes a habitat map (as RasterLayer) as proposed by Harms et al. (2001) as “randomized-habitats procedure”. The algorithm starts with an empty habitat map starts to assign random neighbouring cells to each habitat (in increasing order of abundance in observed map). We modified the procedure slightly by increasing a probability to jump to a non-neighbouring cell as the current patch becomes larger.

Value

list

References

Harms, K. E., Condit, R., Hubbell, S. P., & Foster, R. B. (2001). Habitat associations of trees and shrubs in a 50-ha neotropical forest plot. *Journal of Ecology*, 89(6), 947-959.

See Also

[translate_raster](#)
[adjacent](#)

Examples

```
## Not run:
landscape_classified <- classify_habitats(landscape, classes = 5)
landscape_random <- randomize_raster(landscape_classified, n_random = 19)

## End(Not run)
```

random_walk	<i>Random walk</i>
-------------	--------------------

Description

Randomization of the landscape data using the habitat randomization algorithm.

Usage

```
random_walk
```

Format

rd_ras object.

rcpp_sample	<i>rcpp_sample</i>
-------------	--------------------

Description

Rcpp sample function

Usage

```
rcpp_sample(x, n, replace = FALSE)
```

Arguments

x	Vector of elements to sample from.
n	Size of the sample.
replace	Sample with replacement.

Details

Rcpp implementation of the sample function.

Value

vector

See Also

[sample](#)

reconstruction	<i>Reconstruction</i>
----------------	-----------------------

Description

Randomized data for species b using pattern reconstruction.

Usage

```
reconstruction
```

Format

rd_pat object.

reconstruct_pattern_cluster	<i>reconstruct_pattern_cluster</i>
-----------------------------	------------------------------------

Description

Pattern reconstruction for clustered patterns

Usage

```
reconstruct_pattern_cluster(  
  pattern,  
  n_random = 1,  
  e_threshold = 0.01,  
  max_runs = 1000,  
  no_change = Inf,  
  annealing = 0.01,  
  comp_fast = 1000,  
  weights = c(0.5, 0.5),  
  r_length = 250,
```

```

    return_input = TRUE,
    simplify = FALSE,
    verbose = TRUE,
    plot = FALSE
  )

```

Arguments

pattern	ppp.
n_random	Number of randomizations.
e_threshold	Minimum energy to stop reconstruction.
max_runs	Maximum number of iterations of e_threshold is not reached.
no_change	Reconstruction will stop if energy does not decrease for this number of iterations.
annealing	Probability to keep relocated point even if energy did not decrease.
comp_fast	If pattern contains more points than threshold, summary functions are estimated in a computational fast way.
weights	Weights used to calculate energy. The first number refers to Gest(r), the second number to pcf(r).
r_length	Number of intervals from r = 0 to r = rmax the summary functions are evaluated.
return_input	The original input data is returned as last list entry
simplify	If n_random = 1 and return_input = FALSE only pattern will be returned.
verbose	Print progress report.
plot	Plot pcf function during optimization.

Details

The functions randomizes the observed pattern by using pattern reconstruction as described in Tscheschel & Stoyan (2006) and Wiegand & Moloney (2014). The algorithm starts with a random but clustered pattern, shifts a point to a new location and keeps the change only, if the deviation between the observed and the reconstructed pattern decreases. The pair correlation function and the nearest neighbour distance function are used to describe the patterns.

For large patterns ($n > \text{comp_fast}$) the pair correlation function can be estimated from Ripley's K-function without edge correction. This decreases the computational time. For more information see [estimate_pcf_fast](#).

The reconstruction can be stopped automatically if for n steps the energy does not decrease. The number of steps can be controlled by no_change and is set to no_change = Inf as default to never stop automatically.

The weights must be $0 < \text{sum}(\text{weights}) \leq 1$. To weight both summary functions identical, use `weights = c(0.5, 0.5)`.

spatstat sets r_length to 513 by default. However, a lower value decreases the computational time while increasing the "bumpiness" of the summary function.

Value

list

References

Tscheschel, A., & Stoyan, D. (2006). Statistical reconstruction of random point patterns. *Computational Statistics and Data Analysis*, 51(2), 859-871.

Wiegand, T., & Moloney, K. A. (2014). *Handbook of spatial point-pattern analysis in ecology*. Boca Raton: Chapman and Hall/CRC Press.

See Also

[calculate_energy](#)
[plot_randomized_pattern](#) [reconstruct_pattern_homo](#)
[reconstruct_pattern_hetero](#)
[reconstruct_pattern_marks](#)

Examples

```
## Not run:  
pattern_recon <- reconstruct_pattern_cluster(species_b, n_random = 19, max_runs = 1000)  
  
## End(Not run)
```

reconstruct_pattern_hetero
reconstruct_pattern_hetero

Description

Pattern reconstruction for heterogenous patterns

Usage

```
reconstruct_pattern_hetero(  
  pattern,  
  n_random = 1,  
  e_threshold = 0.01,  
  max_runs = 1000,  
  no_change = Inf,  
  annealing = 0.01,  
  comp_fast = 1000,  
  weights = c(0.5, 0.5),  
  r_length = 250,  
  return_input = TRUE,  
  simplify = FALSE,
```

```

    verbose = TRUE,
    plot = FALSE
  )

```

Arguments

pattern	ppp.
n_random	Number of randomizations.
e_threshold	Minimum energy to stop reconstruction.
max_runs	Maximum number of iterations of e_threshold is not reached.
no_change	Reconstruction will stop if energy does not decrease for this number of iterations.
annealing	Probability to keep relocated point even if energy did not decrease.
comp_fast	If pattern contains more points than threshold, summary functions are estimated in a computational fast way.
weights	Weights used to calculate energy. The first number refers to Gest(r), the second number to pcf(r).
r_length	Number of intervals from $r = 0$ to $r = r_{max}$ the summary functions are evaluated.
return_input	The original input data is returned as last list entry
simplify	If n_random = 1 and return_input = FALSE only pattern will be returned.
verbose	Print progress report.
plot	Plot pcf function during optimization.

Details

The functions randomizes the observed pattern by using pattern reconstruction as described in Tscheschel & Stoyan (2006) and Wiegand & Moloney (2014). The algorithm starts with a random but heterogenous pattern, shifts a point to a new location and keeps the change only, if the deviation between the observed and the reconstructed pattern decreases. The pair correlation function and the nearest neighbour distance function are used to describe the patterns.

For large patterns ($n > \text{comp_fast}$) the pair correlation function can be estimated from Ripley's K-function without edge correction. This decreases the computational time. For more information see [estimate_pcf_fast](#).

The reconstruction can be stopped automatically if for n steps the energy does not decrease. The number of steps can be controlled by no_change and is set to no_change = Inf as default to never stop automatically.

The weights must be $0 < \text{sum}(\text{weights}) \leq 1$. To weight both summary functions identical, use weights = c(0.5, 0.5).

spatstat sets r_length to 513 by default. However, a lower value decreases the computational time while increasing the "bumpiness" of the summary function.

Value

list

References

Tscheschel, A., & Stoyan, D. (2006). Statistical reconstruction of random point patterns. *Computational Statistics and Data Analysis*, 51(2), 859-871.

Wiegand, T., & Moloney, K. A. (2014). *Handbook of spatial point-pattern analysis in ecology*. Boca Raton: Chapman and Hall/CRC Press.

See Also

[calculate_energy](#)
[plot_randomized_pattern](#) [reconstruct_pattern_homo](#)
[reconstruct_pattern_cluster](#)
[reconstruct_pattern_marks](#)

Examples

```
## Not run:  
input_pattern <- spatstat.core::rpoispp(lambda = function(x, y) {100 * exp(-3 * x)}), nsim = 1)  
  
pattern_recon <- reconstruct_pattern_hetero(input_pattern, n_random = 19, max_runs = 1000)  
  
## End(Not run)
```

```
reconstruct_pattern_homo  
      reconstruct_pattern_homo
```

Description

Pattern reconstruction

Usage

```
reconstruct_pattern_homo(  
  pattern,  
  n_random = 1,  
  e_threshold = 0.01,  
  max_runs = 1000,  
  no_change = Inf,  
  annealing = 0.01,  
  n_points = NULL,  
  window = NULL,  
  comp_fast = 1000,  
  weights = c(0.5, 0.5),  
  r_length = 250,  
  return_input = TRUE,  
  simplify = FALSE,
```

```

    verbose = TRUE,
    plot = FALSE
)

```

Arguments

pattern	ppp.
n_random	Number of randomizations.
e_threshold	Minimum energy to stop reconstruction.
max_runs	Maximum number of iterations of e_threshold is not reached.
no_change	Reconstruction will stop if energy does not decrease for this number of iterations.
annealing	Probability to keep relocated point even if energy did not decrease.
n_points	Number of points to be simulated.
window	Window of simulated pattern.
comp_fast	If pattern contains more points than threshold, summary functions are estimated in a computational fast way.
weights	Weights used to calculate energy. The first number refers to Gest(r), the second number to pcf(r).
r_length	Number of intervals from r = 0 to r = rmax the summary functions are evaluated.
return_input	The original input data is returned as last list entry
simplify	If n_random = 1 and return_input = FALSE only pattern will be returned.
verbose	Print progress report.
plot	Plot pcf function during optimization.

Details

The functions randomizes the observed pattern by using pattern reconstruction as described in Tscheschel & Stoyan (2006) and Wiegand & Moloney (2014). The algorithm starts with a random pattern, shifts a point to a new location and keeps the change only, if the deviation between the observed and the reconstructed pattern decreases. The pair correlation function and the nearest neighbour distance function are used to describe the patterns.

For large patterns ($n > \text{comp_fast}$) the pair correlation function can be estimated from Ripley's K-function without edge correction. This decreases the computational time. For more information see [estimate_pcf_fast](#).

The reconstruction can be stopped automatically if for n steps the energy does not decrease. The number of steps can be controlled by no_change and is set to no_change = Inf as default to never stop automatically.

If n_points and window are not specified (default), the simulated pattern has the same number of points and window as the input pattern.

The weights must be $0 < \text{sum}(\text{weights}) \leq 1$. To weight both summary functions identical, use weights = c(0.5, 0.5).

spatstat sets r_length to 513 by default. However, a lower value decreases the computational time while increasing the "bumpiness" of the summary function.

Value

list

References

Tscheschel, A., & Stoyan, D. (2006). Statistical reconstruction of random point patterns. *Computational Statistics and Data Analysis*, 51(2), 859-871.

Wiegand, T., & Moloney, K. A. (2014). *Handbook of spatial point-pattern analysis in ecology*. Boca Raton: Chapman and Hall/CRC Press.

See Also

[calculate_energy](#)
[plot_randomized_pattern](#)
[reconstruct_pattern_hetero](#)
[reconstruct_pattern_cluster](#)
[reconstruct_pattern_marks](#)

Examples

```
## Not run:  
pattern_recon_a <- reconstruct_pattern_homo(species_a, n_random = 19,  
max_runs = 1000)  
  
pattern_recon_b <- reconstruct_pattern_homo(species_a, n_points = 70,  
n_random = 19, max_runs = 1000)  
  
## End(Not run)
```

reconstruct_pattern_marks
reconstruct_pattern_marks

Description

Pattern reconstruction of marks

Usage

```
reconstruct_pattern_marks(  
  pattern,  
  marked_pattern,  
  n_random = 1,  
  e_threshold = 0.01,  
  max_runs = 10000,  
  no_change = Inf,
```

```

    annealing = 0.01,
    r_length = 250,
    return_input = TRUE,
    simplify = FALSE,
    verbose = TRUE,
    plot = FALSE
  )

```

Arguments

pattern	ppp.
marked_pattern	ppp (marked; see details).
n_random	Number of randomizations.
e_threshold	Minimum energy to stop reconstruction.
max_runs	Maximum number of iterations of e_threshold is not reached.
no_change	Reconstruction will stop if energy does not decrease for this number of iterations.
annealing	Probability to keep relocated point even if energy did not decrease.
r_length	Number of intervals from r = 0 to r = rmax the summary functions are evaluated.
return_input	The original input data is returned as last list entry
simplify	If n_random = 1 and return_input = FALSE only pattern will be returned.
verbose	Print progress report.
plot	Plot kmmr function during optimization.

Details

The function randomizes the numeric marks of a point pattern using pattern reconstruction as described in Tscheschel & Stoyan (2006) and Wiegand & Moloney (2014). Therefore, an unmarked as well as a marked pattern must be provided. The unmarked pattern must have the spatial characteristics and the same observation window and number of points as the marked one (see 'reconstruct_pattern' or 'fit_point_process'). Marks must be numeric because the mark-correlation function is used as summary function. Two randomly chosen marks are switched each iteration and changes only kept if the deviation between the observed and the reconstructed pattern decreases.

spatstat sets r_length to 513 by default. However, a lower value decreases the computational time while increasing the "bumpiness" of the summary function.

Value

list

References

- Tscheschel, A., & Stoyan, D. (2006). Statistical reconstruction of random point patterns. *Computational Statistics and Data Analysis*, 51(2), 859-871.
- Wiegand, T., & Moloney, K. A. (2014). *Handbook of spatial point-pattern analysis in ecology*. Boca Raton: Chapman and Hall/CRC Press.

See Also

[fit_point_process](#)
[reconstruct_pattern_homo](#)
[reconstruct_pattern_hetero](#)
[reconstruct_pattern_cluster](#)

Examples

```
## Not run:  
pattern_recon <- reconstruct_pattern_homo(species_a, n_random = 1, max_runs = 1000,  
simplify = TRUE, return_input = FALSE)  
marks_sub <- spatstat.geom::subset.ppp(species_a, select = dbh)  
marks_recon <- reconstruct_pattern_marks(pattern_recon, marks_sub, n_random = 19, max_runs = 1000)  
  
## End(Not run)
```

results_habitat_association
results_habitat_association

Description

Results habitat association

Usage

```
results_habitat_association(  
  pattern,  
  raster,  
  significance_level = 0.05,  
  verbose = TRUE  
)
```

Arguments

pattern	Point pattern or list with reconstructed patterns.
raster	RasterLayer or list of RasterLayers.
significance_level	Significance level
verbose	Print output

Details

The functions shows significant habitat associations by comparing the number of points within a habitat between the observed data and randomized data as described in Plotkin et al. (2000) and Harms et al. (2001). Significant positive or associations are present if the observed count in a habitat is above or below a certain threshold of the randomized count, respectively.

Value

data.frame

References

Harms, K. E., Condit, R., Hubbell, S. P., & Foster, R. B. (2001). Habitat associations of trees and shrubs in a 50-ha neotropical forest plot. *Journal of Ecology*, 89(6), 947-959.

Plotkin, J. B., Potts, M. D., Leslie, N., Manokaran, N., LaFrankie, J. V., & Ashton, P. S. (2000). Species-area curves, spatial aggregation, and habitat specialization in tropical forests. *Journal of Theoretical Biology*, 207(1), 81-99.

See Also

[randomize_raster](#)
[translate_raster](#)
[reconstruct_pattern_homo](#)
[reconstruct_pattern_hetero](#)
[reconstruct_pattern_cluster](#)

Examples

```
landscape_classified <- classify_habitats(landscape, classes = 5)
species_a_random <- fit_point_process(species_a, n_random = 199)
results_habitat_association(pattern = species_a_random, raster = landscape_classified)
```

shar

shar

Description

Analyse species-habitat associations in R. Therefore, information about the location of the species is needed and about the environmental conditions. To test for significance habitat associations, one of the two components is randomized. Methods are mainly based on Plotkin et al. (2000) <doi:10.1006/jtbi.2000.2158> and Harms et al. (2001) <doi:10.1111/j.1365-2745.2001.00615.x>.

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See Also

Useful links:

- <https://r-spatialecology.github.io/shar/>
- Report bugs at <https://github.com/r-spatialecology/shar/issues/>

species_a	<i>Species a</i>
-----------	------------------

Description

A species with negative associations to habitat 4 of 'landscape'.

Usage

species_a

Format

A spatstat ppp object.

species_b	<i>Species b</i>
-----------	------------------

Description

A species with positive associations to habitat 5 of 'landscape'.

Usage

species_b

Format

A spatstat ppp object.

torus_trans	<i>Torus trans</i>
-------------	--------------------

Description

Torus translation of the classified landscape data.

Usage

torus_trans

Format

rd_ras object.

translate_raster	<i>translate_raster</i>
------------------	-------------------------

Description

Torus translation

Usage

```
translate_raster(
  raster,
  steps_x = NULL,
  steps_y = NULL,
  return_input = TRUE,
  simplify = FALSE,
  verbose = TRUE
)
```

Arguments

raster	RasterLayer.
steps_x, steps_y	Number of steps (cells) the raster is translated into the corresponding direction. If both are null, all possible combinations are used.
return_input	The original input data is returned as last list entry.
simplify	If n_random = 1 and return_input = FALSE only raster will be returned.
verbose	Print progress report.

Details

Torus translation test as described in Harms et al. (2001). The raster is shifted in all four cardinal directions by steps equal to the raster resolution. If a cell exits the extent on one side, it enters the extent on the opposite side.

Value

list

References

Harms, K. E., Condit, R., Hubbell, S. P., & Foster, R. B. (2001). Habitat associations of trees and shrubs in a 50-ha neotropical forest plot. *Journal of Ecology*, 89(6), 947-959.

See Also

[randomize_raster](#)

Examples

```
## Not run:  
landscape_classified <- classify_habitats(landscape, classes = 5)  
  
landscape_random <- translate_raster(landscape_classified)  
landscape_random_sub <- translate_raster(landscape_classified, steps_x = 1:10, steps_y = 1:5)  
  
## End(Not run)
```

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