Package: spatialwarnings (via r-universe)

November 5, 2024

Title Spatial Early Warning Signals of Ecosystem Degradation Version 3.1.0 License MIT + file LICENSE Maintainer Alexandre Genin <alexandre.genin@sete.cnrs.fr> **Description** Tools to compute and assess significance of early-warnings signals (EWS) of ecosystem degradation on raster data sets. EWS are spatial metrics derived from raster data -- e.g. spatial autocorrelation -- that increase before an ecosystem undergoes a non-linear transition (Genin et al. (2018) <doi:10.1111/2041-210X.13058>). URL https://github.com/spatial-ews/spatialwarnings BugReports https://github.com/spatial-ews/spatialwarnings/issues **Depends** R (>= 3.3.0), future Imports Rcpp, ggplot2, plyr, stats, utils, future.apply, gsl, Suggests moments, poweRlaw, reshape2, testthat, covr, acss, mgcv, gstat, sp, raster LinkingTo Rcpp, RcppArmadillo LazyData true RoxygenNote 7.2.3 **Encoding UTF-8** Config/pak/sysreqs libgs10-dev Repository https://spatial-ews.r-universe.dev RemoteUrl https://github.com/spatial-ews/spatialwarnings

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arizona

Aerial views of vegetation from Arizona, USA

Description

Aerial views of vegetation from Arizona, USA

Usage

arizona

Format

A list of logical matrices which were obtained through the classification of aerial images of vegetation taken in Arizona (USA).

Source

Derived from the images provided in the Supplementary Material of Rodriguez et al. (2017).

References

Rodriguez, F., A. G. Mayor, M. Rietkerk, and S. Bautista. 2017. A null model for assessing the cover-independent role of bare soil connectivity as indicator of dryland functioning and dynamics. Ecological Indicators.

coarse_grain

Matrix coarse-graining

Description

This function averages the spatial data locally. It divides the input matrix into submatrices of dimension subsize and averages the spatial data in these submatrices. By doing this, the dimension of resultant matrix is reduced by a factor of subsize.

Usage

```
coarse_grain(mat, subsize)
```

Arguments

mat A matrix

subsize Dimension of the submatrix. This has to be a positive integer smaller than the

dimension of input matrix.

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Details

If the data is classified into discrete units, the calculation of variance and skewness can give spurious results irrelevant to the proximity to transition. Therefore, discrete data should be 'coarse-grained' before calculating the spatial early warning signals. However, this can also be applied to continuous state data.

Value

A matrix of reduced dimension.

References

Sankaran, S., Majumder, S., Kefi, S. and Guttal, V. (2017). Implications of being discrete and spatial for detecting early warning signals of regime shifts. Ecological Indicators.

See Also

```
generic_sews
```

Examples

convert_to_matrix

Convert an object to a matrix

Description

This function is mainly for internal use by the spatialwarnings package to convert objects before they are processed by *_sews functions.

Usage

```
convert_to_matrix(object, ...)
```

Arguments

```
object An object (typically, a matrix or a list of matrices)
... Additional arguments (currently ignored)
```

create_indicator 5

Details

This generic function is here so that other packages can extend it. For example, spatialwarningsGis will provide methods so that GIS objects can be handled (e.g. RasterLayer from package raster).

Examples

```
# this does nothing
convert_to_matrix(serengeti[2:3])
```

create_indicator

Custom Spatial Early-Warning signals

Description

Computation, significance assessment and display of trends of a custom, user-defined indicator.

Usage

```
create_indicator(fun, taskname = as.character(substitute(fun)))
compute_indicator(mat, fun, taskname = as.character(substitute(fun)), ...)
```

Arguments

fun A function that takes a matrix as input and returns a vector of numerical values.

If the function returns a named vector, then the names will be used in plots and

summaries. The function may also accept extra arguments.

taskname The task name. A character string used used for plots and textual summaries

that describes the indicator (or set of indicators) being computed. If a task name

cannot be derived from fun, then a default name is used.

mat A matrix or a list of matrices.

... Additional arguments to pass to the function fun

Details

spatialwarnings provides "workflow functions", named *_sews, that assist the user in computing, displaying and assessing the significance of indicator values. The functions create_indicator and compute_indicator provides such workflow for any arbitrary function.

create_indicator takes a function 'fun' and returns another function that can be used as an indicator similar to the *_sews functions. The results of this function can be assessed for significance using indictest and trends can be displayed using plot, summary, etc. (see Examples). compute_indicator does the same but without needing an intermediate indicator function.

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Value

create_indicator returns a function that can be used in the same way than the other *_sews functions (e.g. generic_sews). This function as well as compute_indicator will return simple_sews_* objects.

See Also

```
simple_sews
```

```
\# Use the maximum patch size as indicator of degradation
maxpatchsize <- function(mat) {</pre>
  max(patchsizes(mat))
}
# Create the indicator function
maxpatch_sews <- create_indicator(maxpatchsize)</pre>
# Then work with this function as if it were a function from the *_sews
# family.
mp_indic <- maxpatch_sews(forestgap)</pre>
summary(mp_indic)
# Assess significance and display trends
mp_test <- indictest(mp_indic, nulln = 49)</pre>
plot(mp_test)
# Try spatial coefficient of variation as a spatial EWS. This function can
# have arguments.
spatial_cv <- function(mat, subsize) {</pre>
  matc <- coarse_grain(mat, subsize)</pre>
  return( sd(matc) / mean(matc) )
}
# Create indicator function
cv_sews <- create_indicator(spatial_cv)</pre>
# Compute and display trends
cv_indic <- cv_sews(serengeti, subsize = 3)</pre>
plot(cv_indic, along = serengeti.rain)
# We can do the same work in one go using compute_indicator
cv_indic2 <- compute_indicator(serengeti, spatial_cv, subsize = 3)</pre>
plot(cv_indic2, along = serengeti.rain)
indictest(cv_indic, nulln = 99)
```

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dda

Density-dependent aggregation model

Description

dda is a list of matrices representing results from the density-dependent aggregation model (Siteur et al. 2023) in dda. dda.pars is a data frame with the model parameters. Each row of dda.pars corresponds to a matrix in dda, in the same order. All parameters were maintained constant, except for tau (see model definition in Siteur et al. 2023).

Usage

dda

dda.pars

Format

An object of class list of length 4.

An object of class data. frame with 4 rows and 5 columns.

Source

Kindly provided by Koen Siteur

References

Siteur, Koen, Quan-Xing Liu, Vivi Rottschäfer, Tjisse van der Heide, Max Rietkerk, Arjen Doelman, Christoffer Boström, and Johan van de Koppel. 2023. "Phase-Separation Physics Underlies New Theory for the Resilience of Patchy Ecosystems." Proceedings of the National Academy of Sciences 120 (2): e2202683120. https://doi.org/10.1073/pnas.2202683120.

```
ddasews <- lsw_sews(dda)
plot(ddasews, along = dda.pars[ ,"tau"]) # tau is the changing parameter
display_matrix(dda[[4]])</pre>
```

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display_matrix	Plot a matrix		
----------------	---------------	--	--

Description

Display a matrix or a list of matrices in a plot

Usage

```
display_matrix(object, palette = "RdYlBu", along = NULL, ...)
```

Arguments

object	A matrix, a list of matrices, an object produced by \star _sews functions or indictest()
palette	A color palette to use in the plot. It can be any color palette understood by scale_fill_brewer.
along	A vector of values used in facet headers. If object is a matrix, this is ignored.
	Other arguments are ignored.

Details

This function will plot a matrix using ggplot2, using the provided palette. Its use-case is very much like image(), but its produces nicer plots by default (image() is much faster, however).

Value

A ggplot2 object, which is printed when this function is used interactively.

```
# display_matrix works with single matrices or lists of matrices
display_matrix(serengeti[2:3])
# display_matrix is compatible with "*_sews" objects
indics <- compute_indicator(serengeti[2:3], raw_moran)
display_matrix(indics)</pre>
```

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dLSW

The Lifshitz-Slyozov-Wagner distribution

Description

Density and distribution function for the Lifshitz-Slyozov-Wagner (LSW) distribution with mean mu.

Usage

```
dLSW(x, mu, log = FALSE)
pLSW(x, mu, lower.tail = TRUE, log.p = FALSE)
LSW_fit(x)
```

Arguments

x vector of quantiles

mu the mean of the distribution

log, log.p logical; if TRUE, probabilities p are given as log(p)

lower.tail logical; if TRUE (default), probabilities are $P[X \le x]$ otherwise, P[X > x]

Details

The LSW distribution is a continuous distribution with density

$$f(x) = \frac{4x^2}{9\mu^3} \left(\frac{3\mu}{3\mu + x}\right)^{7/3} \left(\frac{3\mu}{3\mu - 2x}\right)^{11/3} e^{\frac{2x}{2x - 3\mu}}$$

where μ is the mean of the distribution.

The functions dLSW gives the probability density, pLSW gives the distribution function. qLSW and rLSW are not implemented. You can use LSW_fit to fit an LSW distribution to a set of observations.

The length of the results is determined by the length of x, and mu can only be a single value.

Please note that this distribution has support only on the interval [0,mu*3/2). Probabilities outside this interval are returned as 0.

Value

dLSW gives the density, pLSW gives the distribution function, both as numerical vectors determined by the length of x.

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References

Siteur, Koen, Quan-Xing Liu, Vivi Rottschäfer, Tjisse van der Heide, Max Rietkerk, Arjen Doelman, Christoffer Boström, and Johan van de Koppel. 2023. "Phase-Separation Physics Underlies New Theory for the Resilience of Patchy Ecosystems." Proceedings of the National Academy of Sciences 120 (2): e2202683120. https://doi.org/10.1073/pnas.2202683120.

See Also

```
1sw_sews
```

Examples

extract_spectrum

Extract the r-spectrum from objects

Description

Extract the r-spectrum from objects produced by spectral_sews.

Usage

```
extract_spectrum(x, ...)
```

Arguments

x An object produced by spectral_sews or the result of the indictest function called on such object

... Other arguments are ignored

Value

The empirical r-spectrum as a data. frame

See Also

```
spectral_sews, rspectrum
```

extract_variogram 11

Examples

```
# Extract the r-spectrum after computing indicators
indics <- spectral_sews(serengeti[2:3])
extract_spectrum(indics)</pre>
```

extract_variogram

extract_variogram() method for variogram_sews objects

Description

Extract the empirical variogram from a variogram_sews object

Usage

```
extract_variogram(x, ...)
```

Arguments

x An object produced by variogram_sews, or indictest

... Additional arguments (ignored)

Value

A data.frame containing the variogram with the distances (column dist), the empirical semivariance values (gamma), and if object contains more than one matrix, a column named matrixn.

See Also

```
variogram_sews
```

```
vario_indics <- variogram_sews(serengeti)
predict(vario_indics)
vario_test <- indictest(vario_indics, nulln = 19)
predict(vario_test) # same result</pre>
```

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flowlength_sews	Flowlength connectivity indicator (uniform topography)	
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Description

Measures the connectivity of runoff-source areas as determined by vegetation patterns and (uniform) topography

Usage

```
flowlength_sews(mat, slope = 20, cell_size = 1)
```

Arguments

mat The input matrix (must be a logical matrix)

slope The slope of the area documented by the matrix (in degree).

cell_size The horizontal size of a cell in the matrix (as viewed from above).

Details

This function computes Flowlength, a simple metric that measures the potential hydrological connectivity of runoff-source areas (e.g., bare soil) considering vegetation cover, vegetation patterns and topography. Flowlength is defined as the average length of all the potential runoff pathways in the target area. Thus, a higher value of the index indicates a higher hydrologic connectivity of runoff source areas. This function is designed for an idealized uniform hillslope (e.g., with constant slope angle, the direction of maximum slope being from the top to the bottom of the input matrices).

The deviations of Flowlength from its expected values under random or aggregated-pattern null models can be used as an indicator of imminent transition to a degraded state (Rodriguez et al. 2017) in the context of arid drylands. An increased deviation of flowlength compared to its null values is expected as a possible transition gets closer. This deviation can be computed using any null model provided by spatialwarnings (see indictest for more details), but a specific null model is provided for Flowlength based on a much-faster analytical approximation, using the argument null_method = "approx_rand" when calling indictest (see examples below).

In general, Flowlength can be used as indicator of dryland functional status by assessing potential water and soil losses in patchy landscapes (Mayor et al. 2008, Moreno-de las Heras et al. 2012, Mayor et al. 2013, Okin et al. 2015). Finally, the combination of observed and expected Flowlength under null models for random or aggregated vegetation cover can be used for assessing the cover-independent role of bare-soil connectivity (Rodriguez et al. 2018).

Value

A 'simple_sews' object containing the flow length value, among other things, see simple_sews_object for more information.

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References

Rodriguez, F., A. G. Mayor, M. Rietkerk, and S. Bautista. 2017. A null model for assessing the cover-independent role of bare soil connectivity as indicator of dryland functioning and dynamics. Ecological Indicators.

Mayor, A.G., Bautista, S., Small, E.E., Dixon, M., Bellot, J., 2008. Measurement of the connectivity of runoff source areas as determined by vegetation pattern and topography: a tool for assessing potential water and soil losses in drylands. Water Resour. Res. 44, W10423.

Mayor, A.G., Kefi, S., Bautista, S., Rodriguez, F., Carteni, F., Rietkerk, M., 2013. Feedbacks between vegetation pattern and resource loss dramatically decrease ecosystem resilience and restoration potential in a simple dryland model. Landsc. Ecol. 28, 931-942.

Moreno-de las Heras, M., Saco, P.M., Willgoose, G.R., Tongway, D.J., 2012. Variations in hydrological connectivity of Australian semiarid landscapes indicate abrupt changes in rainfall-use efficiency of vegetation. J. Geophys. Res. 117, G03009.

Okin, G.S., Moreno-de las Heras, M., Saco, P.M., Throop, H.L., Vivoni, E.R., Parsons, A.J., Wainwright, J., Peters, D.P.C., 2015. Connectivity in dryland landscapes: shifting concepts of spatial interactions. Front. Ecol. Environ. 13 (1), 20-27.

See Also

raw_flowlength_uniform, indictest to test the significance of indicator values.

Examples

```
fl_result <- flowlength_sews(arizona, slope = 20, cell_size = 1)

# Compute the Z-score (standardized deviation to null distribution) and plot
# its variations along the gradient. This Z-score is suggested by
# Rodriguez et al. (2017) as an indicator of degradation.
fl_test <- indictest(fl_result, nulln = 19)
plot(fl_test, what = "z_score")

# Use the analytical approximation suggested in Rodriguez et al. (2017),
# instead of permuting the original values in the matrix (much faster)
fl_test <- indictest(fl_result, null_method = "approx_rand")
plot(fl_test, what = "z_score")</pre>
```

forestgap

A list of binary matrices and their associated parameters

Description

A list of binary matrices and their associated parameters

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Usage

```
forestgap.pars
```

Format

A list of logical matrices which are the end results of simulations from Kubo's Forest Gap model along a gradient of increasing values of stress (see references).

The parameters used for the simulations, as a data frame.

Details

Kubo's forest gap model has three parameters, α that controls the reproductive rate of trees, d controls the non-spatialized mortality and δ the increased mortality due to the presence of a neighboring gap.

Source

Generated using the implementation of Kubo's model in caspr 0.2.0 https://github.com/fdschneider/caspr.

References

Kubo, T., Iwasa, Y., & Furumoto, N. (1996). Forest spatial dynamics with gap expansion: Total gap area and gap size distribution. Journal of Theoretical Biology, 180(3), 229-246. doi:10.1006/jtbi.1996.0099

generic_sews

Generic Spatial Early-Warning signals

Description

Computation, significance assessment and display of spatial generic early warning signals (Moran's I, variance and skewness)

Usage

```
generic_sews(
  mat,
  subsize = 4,
  abs_skewness = FALSE,
  moranI_coarse_grain = FALSE
)
```

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Arguments

mat A matrix (quantitative data), a binary matrix (which contains TRUE or FALSE

values) or a list of those

subsize The subsize used for the coarse-graining phase (see Details)

abs_skewness Should the absolute skewness be used instead of its raw values?

moranI_coarse_grain

Should the input matrix be coarse-grained before computing the Moran's I indi-

cator value?

Details

The Generic Early warning signal are based on the property of a dynamical system to "slow down" when approaching a critical point, that is, take more time to return to equilibrium after a perturbation. This is expected to be reflected in several spatial characteristics: the variance, the spatial autocorrelation (at lag-1) and the skewness. This function provides a convenient workflow to compute these indicators, assess their significance and display the results.

Before computing the actual indicators, the matrix can be "coarse-grained". This process reduces the matrix by averaging the nearby cells using a square window defined by the subsize parameter. This makes spatial variance and skewness reflect actual spatial patterns when working with binary (TRUE/FALSE data), but is optional when using continuous data. Keep in mind that it effectively reduces the size of the matrix by approximately subsize on each dimension.

The significance of generic early-warning signals can be estimated by reshuffling the original matrix (function indictest). Indicators are then recomputed on the shuffled matrices and the values obtained are used as a null distribution. P-values are obtained based on the rank of the observed value in the null distribution. A small P-value means that the indicator is significantly above the null values, as expected before a critical point.

The plot method can displays the results graphically. A text summary can be obtained using the summary method.

Value

generic_sews returns an object of class simple_sews_single (a list) if mat is a single matrix or an object of class simple_sews_list if mat is a list. You probably want to use some of the methods written for these complicated objects instead of extracting values directly (they are displayed using print(<object>)).

indictest returns an object of class generic_test (a data.frame).

plot methods return ggplot objects, usually immediately displayed when plot is being called at the R prompt.

References

Kefi, S., Guttal, V., Brock, W.A., Carpenter, S.R., Ellison, A.M., Livina, V.N., et al. (2014). Early Warning Signals of Ecological Transitions: Methods for Spatial Patterns. PLoS ONE, 9, e92097.

Dakos, V., van Nes, E. H., Donangelo, R., Fort, H., & Scheffer, M. (2010). Spatial correlation as leading indicator of catastrophic shifts. Theoretical Ecology, 3(3), 163-174.

Guttal, V., & Jayaprakash, C. (2008). Spatial variance and spatial skewness: leading indicators of regime shifts in spatial ecological systems. Theoretical Ecology, 2(1), 3-12.

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See Also

indictest, to test the significance of indicator values. Individual indicators: raw_cg_moran raw_cg_variance, raw_cg_skewness, simple_sews

Examples

```
data(serengeti)
gen_indic <- generic_sews(serengeti, subsize = 5,</pre>
                           moranI_coarse_grain = TRUE)
# Display results
summary(gen_indic)
# Display trends along the varying model parameter
plot(gen_indic, along = serengeti.rain)
# Compute significance (long)
gen_test <- indictest(gen_indic, nulln = 199)</pre>
print(gen_test)
\# Display the trend, now with a grey ribbon indicating the 5%-95% quantile
# range of the null distribution
plot(gen_test, along = serengeti.rain)
# Display the effect size compared to null distribution
plot(gen_test, along = serengeti.rain, what = "z_score")
# Note that plot() method returns a ggplot object that can be modified
# for convenience
if ( require(ggplot2) ) {
 plot(gen_test, along = serengeti.rain) +
   geom_vline(xintercept = 733, color = "red", linetype = "dashed") +
   xlab('Annual rainfall') +
    theme_minimal()
}
```

indicator_plrange

Power-law range indicator

Description

Compute the power-law range of a matrix

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Usage

indicator_plrange(mat, merge = FALSE, xmin_bounds = NULL)

Arguments

mat A logical matrix, or a list of logical matrices

merge Controls whether the patch-size distributions of the input matrices are merged

together before computing the power-law range. Setting this value to TRUE makes the function return a single value even if multiple matrices are given as

input.

xmin_bounds A vector of two integer values, defining a range in which to search for the best

xmin (see Details).

Details

Some ecosystems show typical changes in their patch-size distribution as they become more and more degraded. In particular, an increase in the truncation of the patch-size distribution (PSD) is expected to occur. The power-law range (PLR) measures the truncation of the PSD in a single value (see also patchdistr_sews for more details).

To compute the PLR, power-laws are fitted with a variable minimum patch size (x_{min}) and the one with the lowest Kolmogorov-Smirnov distance to the empirical distribution is retained. PLR is then computed using this best-fitting x_{min} as:

$$\frac{log(x_{max}) - log(x_{min})}{log(x_{max}) - log(x_{smallest})}$$

where x_{max} is the maximum observed patch size, and $x_{smallest}$ is the minimum observed patch size.

Value

A data frame with columns minsize, maxsize which are the observed minimum and maximum patch sizes, the estimated x_{min} as column xmin and the value of the power-law range as plrange. If multiple matrices were provided, then a list of data frames is returned

References

Clauset, A., Shalizi, C. R., & Newman, M. E. (2009). Power-law distributions in empirical data. SIAM review, 51(4), 661-703.

Berdugo, M., Kefi, S., Soliveres, S. & Maestre, F.T. (2017). Plant spatial patterns identify alternative ecosystem multifunctionality states in global drylands. Nature in Ecology and Evolution.

See Also

patchdistr_sews

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Examples

```
forestgap.plr <- indicator_plrange(forestgap)
do.call(rbind, forestgap.plr) # convert results to data.frame

# Restrict to small xmins
forestgap.plr2 <- indicator_plrange(forestgap, xmin_bounds = c(1, 10))
do.call(rbind, forestgap.plr2)</pre>
```

indicator_psdtype

Change in patch-size distributions types

Description

This functions fits different patch size distributions types (power-law, log-normal, exponential and truncated power-law) to the patches contained in a matrix. The distributions are returned with their corresponding AIC, BIC and AICc to select the best fit.

Usage

```
indicator_psdtype(
    x,
    xmin = 1,
    merge = FALSE,
    fit_lnorm = FALSE,
    xmin_bounds = NULL,
    best_by = "AIC",
    wrap = FALSE,
    nbmask = "von_neumann"
)
```

Arguments x

wrap

xmin	The xmin to be used to fit the patch size distributions. Use the special values "estimate" to use an estimated xmin for each fit
merge	The default behavior is to produce indicators values for each matrix. If this parameter is set to TRUE then the patch size distributions are pooled together for fitting.
fit_lnorm	Fit also a log-normal distribution
xmin_bounds	Restrict the possible xmins in this range (defaults to the whole range of observed patch sizes)
best_by	The criterion used to select the best distribution type (one of "AIC", "BIC" or

A logical (TRUE/FALSE values) matrix or a list of these.

Determines whether patches are considered to wrap around the matrix when reaching the side

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nbmask

Either "moore" for 8-way neighborhood, "von_neumann" for four-way neighborhood (default), or a 3x3 matrix describing which neighbors to consider around a cell. See patchsizes for details on how to specify such neighborhoods.

Details

Patterned ecosystems can exhibit a change in their spatial structure as they become more and more stressed. It has been suggested that this should be reflected in changes in the observed patch size distributions (PSD). The following sequence is expected to occur (Kefi et al. 2011) as patterned ecosystems become more and more degraded:

- Percolation of vegetation patches occurs (a patch has a width or height equal to the size of the system)
- The patch-size distribution follows a power-law
- The patch-size distribution deviates from a power-law as larger patches break down
- The patch-size distribution is closer to an exponential distribution

This indicator fits the observed patch size distribution based on maximum-likelihood (following Clauset et al. 2009 recommendations), then select the best model using AIC, BIC (default) or AICc.

Value

A data.frame (or a list of these if x is a list) with the following columns:

- method the method used for fitting (currently: only log-likelihood is implemented, "II")
- type the type of distribution
- npars the number of parameters of the distribution type
- AIC, 'AICc' and 'BIC' the values for Akaike Information Criterion (or the corrected for small samples equivalent AICc), and Bayesion Information Criterion (BIC)
- best A logical vector indicating which distribution is the best fit
- plexpo, cutoff, meanlog, sdlog the estimates for distribution parameters (see pl_fit)
- percolation A logical value indicating whether there is percolation in the system.

References

Kefi, S., Rietkerk, M., Roy, M., Franc, A., de Ruiter, P.C. & Pascual, M. (2011). Robust scaling in ecosystems and the meltdown of patch size distributions before extinction: Patch size distributions towards extinction. Ecology Letters, 14, 29-35.

Kefi, S., Rietkerk, M., Alados, C.L., Pueyo, Y., Papanastasis, V.P., ElAich, A., et al. (2007). Spatial vegetation patterns and imminent desertification in Mediterranean arid ecosystems. Nature, 449, 213-217.

Clauset, A., Shalizi, C. R., & Newman, M. E. (2009). Power-law distributions in empirical data. SIAM review, 51(4), 661-703.

See Also

```
patchdistr_sews
patchdistr_sews
```

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Examples

```
data(forestgap)
# One logical matrix only
indicator_psdtype(forestgap[[1]])
# A list of these matrices
indicator_psdtype(forestgap)
```

indictest

Significance-assessment of spatial early-warning signals

Description

Assess the significance of spatial early-warning indicators

Usage

```
indictest(x, nulln = 999, null_method = "perm", null_control = NULL, ...)
```

Arguments

X	An object such as one produced by the *_sews functions, or compute_indicator
nulln	The number of values to produce the null distribution
null_method	The method used to produce the null values (see Details)
null_control	List of arguments used to control the generation of null matrices. If NULL, then sensible defaults are chosen (see Details)
• • •	Additional arguments are ignored

Details

indictest is used to test the significance of early-warning signals against 'null matrices', which represent the expected spatial structure in the absence of the biological process of interest.

For a given indicator, a null distribution is obtained by producing a set of 'null' matrices, from which indicator values are recomputed. This produces a null distribution of nulln indicator values against which the observed value is tested.

Several methods are available to produce the set of null matrices. If null_method is set to "perm", the original matrix is reshuffled to obtain a null matrix. If null_method is set to "intercept", then a generalized linear model of the form 'y ~ 1' (where y represents the values of the matrix) is fitted, then values are drawn from this model. If null_method is set to "smooth", then a smooth surface is fitted based on a generalized additive model (using gam) to the matrix, then values are drawn from this model. When using the "intercept" or "smooth" null models, it is important to make sure the model 'family' corresponds to the type of values present in the matrix. By default, if a matrix

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contains TRUE/FALSE values, a 'binomial()' family is used, otherwise a 'gaussian()' family is used. More information about null models is available in the spatialwarnings FAQ.

Please note that specific null methods may exists for some indicators, such as flowlength. These are often based on analytical approximation and allow faster computations.

If a matrix has attributes, then these are preserved and passed to the function used to compute the indicator value, except when using the null method 'perm', in which case matrix attributes are discarded.

The list null_control can be used to adjust the computation of null matrices. It can have the following components:

family The family used in the model used to produce the null matrices. Typically, it is one of binomial(), gaussian(), etc.

qinf The lower quantile to compute from the null distribution and display in summaries/plots. A numeric value between 0 and 1.

qsup The upper quantile to compute from the null distribution and display in summaries/plots. A numeric value between 0 and 1.

Value

An object with a class ending in *_sews_test, whose exact class depends on the input object. plot, summary methods are available to display the results of computations, and additional methods may be available depending on the input object (e.g. see patchdistr_sews_plot).

References

Kefi, S., Guttal, V., Brock, W.A., Carpenter, S.R., Ellison, A.M., Livina, V.N., et al. (2014). Early Warning Signals of Ecological Transitions: Methods for Spatial Patterns. PLoS ONE, 9, e92097

See Also

generic_sews, spectral_sews, kbdm_sews, compute_indicator, flowlength_sews

kbdm_sews

Indicator based on Kolmogorov Complexity

Description

Computes the Kolmogorov Complexity on a set of matrices, using the Block Decomposition Method.

Usage

```
kbdm_sews(mat, subsize = 3)
```

Arguments

mat A logical matrix (TRUE/FALSE values) or a list of logical matrices

subsize A submatrix size to carry out the Block Decomposition Method (must be be-

tween 1 and 3)

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Details

be a useful indicator to anticipate transitions in model ecological systems (Dakos and Soler-Toscano, 2017). When close to the transition critical point, the complexity is expected to decrease.

The Kolmogorov complexity cannot be computed directly for large strings (i.e. matrices). However, the complexity of smaller submatrices can be estimated, then combined to obtain an approximation of the complexity of the whole matrix. This method, the Block Decomposition Method is implemented in this indicator following Dakos and Soler-Toscano (2017).

Value

kbdm_sews returns an object of class simple_sews_single (a list) if mat is a single matrix, and an object of class simple_sews_list if mat is a list of matrices. These objects can be used with generic methods indictest (to test significance) or plot (to display trends), see also the examples below.

References

Dakos, V., and F. Soler-Toscano. 2017. Measuring complexity to infer changes in the dynamics of ecological systems under stress. Ecological Complexity 32:144-155.

See Also

raw_kbdm, acss, indictest, to test the significance of indicator values.

Examples

```
kbdm_result <- kbdm_sews(serengeti, subsize = 3)
plot(kbdm_result, along = serengeti.rain)
kbdm_test <- indictest(kbdm_result, nulln = 49)
plot(kbdm_test, along = serengeti.rain)
# Plot deviation to null expectation
plot(kbdm_test, along = serengeti.rain, what = "z_score")</pre>
```

label

Labelling of unique patches and detection of percolation.

Description

Label each patch with a number in a binary matrix

percolation() detects whether percolation occurs in the matrix (i.e. a patch has a width or a height equal to the size of the matrix)

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Usage

```
label(mat, nbmask = "von_neumann", wrap = FALSE)
percolation(mat, nbmask = "von_neumann")
```

Arguments

nbmask Either "moore" for 8-way neighborhood, "von_neumann" for four-way neighborhood (default), or a 3x3 matrix describing which neighbors to consider around a cell. See patchsizes for details on how to specify such neighborhoods.

wrap Whether to wrap around lattice boundaries ('TRUE'/'FALSE'), effectively us-

ing periodic boundaries.

Details

The label function "labels" the patches of a binary (TRUE/FALSE) matrix. It returns a matrix of similar height and width, with integer values representing the ID of each unique patch (contiguous cells). Empty cells are labelled as NA.

Value

A matrix containing ID numbers for each connected patch. Default parameters assume 4-cell neighborhood and periodic boundaries. The distribution of patch sizes is returned as the attribute "psd" and the percolation status as "percolation" (whether a TRUE patch has a width or height equal to the size of the matrix).

See Also

```
patchsizes, patchdistr_sews
```

```
data(forestgap)

rmat <- matrix(rnorm(100) > .1, ncol = 10)
display_matrix(label(rmat))

# With 8-way neighborhood mask and no wrapping around borders
display_matrix(label(rmat, "moore", wrap = FALSE))

# On real data:
display_matrix(label(forestgap[[5]], "moore", wrap = FALSE))
```

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1sw_sews

Indicators based on the LSW distribution

Description

LSW indicators for systems with density-dependent aggregation

Usage

```
lsw_sews(mat, wrap = FALSE)
raw_patch_radii_skewness(mat, wrap = FALSE)
raw_lsw_aicw(mat, wrap = FALSE)
```

Arguments

mat A logical matrix (TRUE/FALSE values) or a list of such matrices

wrap Determines whether patches are considered to wrap around the matrix when

reaching on of its edges

Details

In systems where a mobile resource or consumer can be fixed in space by a sessile species, specific patterns are expected to appear. Such systems can include situations where nutrients available in the environment are fixed by a sessile species (e.g. seagrasses or corals), or where the behavior of herbivores is altered to restrict them to certain areas (see full theoretical background in Siteur et al. 2023).

In those systems, as environmental conditions change and the global density of the sessile species decreases, its spatial structure is expected to change. The area of patches of the sessile species (as measured by their radii, which assumes circular patches), is expected to go from a log-normal to a Lifshitz–Slyozov–Wagner (LSW) distribution. Thus, measuring how close the observed distribution of radii are to those two candidate distributions can constitute an indicator of ecosystem degradation.

This function measures this through the relative support based on AIC for the two distributions (equal to 1 when the empirical distribution is best-approximated by an LSW, and 0 when it is a log-normal distribution (dlnorm), and the skewness of the observed patch radii, which should approach a value around -0.92 as conditions worsen.

Value

lsw_sews returns an object of class simple_sews_single (a list) if mat is a single matrix or an object of class simple_sews_list if mat is a list. You probably want to use some of the methods written for these complicated objects instead of extracting values directly (they are displayed using print(<object>)).

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Author(s)

This code has received contributions from Koen Siteur

References

Siteur, Koen, Quan-Xing Liu, Vivi Rottschäfer, Tjisse van der Heide, Max Rietkerk, Arjen Doelman, Christoffer Boström, and Johan van de Koppel. 2023. "Phase-Separation Physics Underlies New Theory for the Resilience of Patchy Ecosystems." Proceedings of the National Academy of Sciences 120 (2): e2202683120. https://doi.org/10.1073/pnas.2202683120.

See Also

```
dLSW, dda, raw_patch_radii_skewness, raw_lsw_aicw
```

Examples

```
data(dda)
data(dda.pars)

# Compute all indicators at once (skewness and relative AIC support)
indics <- lsw_sews(dda)
plot(indics, along = dda.pars[ ,"tau"])

# Compute individual indicators

# Skewness of the distribution of patch radii
radii_skewness <- compute_indicator(dda, raw_patch_radii_skewness)
plot(radii_skewness, along = dda.pars[ ,"tau"])

# Aic weight of LSW distribution relative to a lognormal distribution. tau here
# represents the density at equilibrium in Siteur et al's model (2023)
lsw_aicw <- compute_indicator(dda, raw_lsw_aicw)
plot(lsw_aicw, along = 1 - dda.pars[ ,"tau"])</pre>
```

patchdistr_sews

Early-warning signals based on patch size distributions

Description

Compute early-warning signals based on patch size distributions

Usage

```
patchdistr_sews(
  mat,
  merge = FALSE,
  fit_lnorm = FALSE,
```

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```
best_by = "BIC",
xmin = 1,
xmin_bounds = NULL,
wrap = FALSE,
nbmask = "von_neumann")
```

Arguments

mat A logical matrix (TRUE/FALSE values) or a list of these

merge The default behavior is to produce indicators values for each matrix. If this

parameter is set to TRUE then the patch size distributions are pooled together before fitting, yielding only one final indicator value for the set of input matrices

(argument mat).

fit_lnorm When patch size distributions are compared, should we consider lognormal type

? (see details)

best_by The criterion to use to select the best fit (one of "AIC", "BIC" or "AICc")

xmin The x_{min} to be used to fit the patch size distributions. Use the special value

"estimate" to compute first the x_{min} that produces the best power-law fit, then

use this estimated value to fit all distributions.

xmin_bounds Bounds when estimating x_{min} for power-law distributions

wrap Determines whether patches are considered to wrap around the matrix when

reaching the side

nbmask Either "moore" for 8-way neighborhood, "von_neumann" for four-way neigh-

borhood (default), or a 3x3 matrix describing which neighbors to consider around

a cell. See patchsizes for details on how to specify such neighborhoods.

Details

Patterned ecosystems can exhibit a change in their spatial structure as they become more and more stressed. It has been suggested that this should be reflected in changes in the observed patch size distributions (PSD). The following sequence is expected to occur (Kefi et al. 2011) as patterned ecosystems become more and more degraded:

- Percolation of vegetation patches occurs (a patch has a width or height equal to the size of the system)
- The patch-size distribution follows a power-law
- The patch-size distribution deviates from a power-law as larger patches break down
- The patch-size distribution is closer to an exponential distribution

Additionally, it has been suggested that these changes in patch size distribution shape should be reflected in the power-law range (PLR). This function carries out all the required computations and helps display the results in a convenient form.

The fitting of PSDs is based on maximum-likelihood following Clauset et al.'s procedure. The best discrete distribution is estimated among these candidates: a power-law x^{λ} , an exponential $exp(\alpha x)$, a truncated power-law and $x^{\lambda}exp(\alpha x)$, and optionally, a log-normal. Each distribution parameter is estimated using maximum-likelihood, with a minimum patch size (xmin) fixed to one. The best

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distribution is selected based on BIC by default. In raw results, plexpo refers to the power-law exponent (λ in the previous equations) and cutoff referes to the exponential decay rate α .

To compute the Power-law range (PLR), power-laws are fitted with a variable minimum patch size (xmin) and the one with the lowest Kolmogorov-Smirnov distance to the empirical distribution is retained. PLR is then computed using this best-fitting xmin:

$$\frac{log(x_{max}) - log(x_{min})}{log(x_{max}) - log(x_{smallest})}$$

Results can be displayed using the text-based summary and print, but graphical options are also available to plot the trends (plot) and the fitted distributions (plot_distr). Plotting functions are documented in a separate page. Observed and fitted distributions can be produced using the predict function, as documented on this page.

Value

A list object of class 'psdfit' containing among other things - the observed patch size distribution data - the model outputs for the candidate distribution fits - the power-law range values - the percolation values (if several matrices were provided and 'merge' was TRUE, then the average percolation value is returned)

References

Kefi, S., Rietkerk, M., Alados, C. L., Pueyo, Y., Papanastasis, V. P., ElAich, A., & De Ruiter, P. C. (2007). Spatial vegetation patterns and imminent desertification in Mediterranean arid ecosystems. Nature, 449(7159), 213-217.

Kefi, S., Rietkerk, M., Roy, M., Franc, A., de Ruiter, P.C. & Pascual, M. (2011). Robust scaling in ecosystems and the meltdown of patch size distributions before extinction: Patch size distributions towards extinction. Ecology Letters, 14, 29-35.

Berdugo, M, Sonia Kefi, Santiago Soliveres, and Fernando T. Maestre. (2017) Plant Spatial Patterns Identify Alternative Ecosystem Multifunctionality States in Global Drylands. Nature in Ecology and Evolution, no. 1.

Clauset, A., Shalizi, C. R., & Newman, M. E. (2009). Power-law distributions in empirical data. SIAM review, 51(4), 661-703.

See Also

```
patchsizes, plot_distr, predict, plot,
indictest, to test the significance of indicator values.
```

```
data(forestgap)
psd_indic <- patchdistr_sews(forestgap)
summary(psd_indic)</pre>
```

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```
plot(psd_indic)

# Plots can be modified using ggplot2 directives
if ( require(ggplot2) ) {
    plot(psd_indic) +
        theme_minimal()
}

# Export results to a data.frame
psd_indic_export <- as.data.frame(psd_indic)
head(psd_indic_export)</pre>
```

patchdistr_sews_plot Early-warning signals based on patch size distributions

Description

Plot early-warning signals based on patch size distributions

Usage

```
## S3 method for class 'patchdistr_sews'
plot(x, along = NULL, ...)
plot_distr(x, along = NULL, best_only = TRUE, plrange = TRUE)
```

Arguments

X	An object as produced by patchdistr_sews, or the result of indictest called on such object
along	A vector providing values along which the indicator trends will be plotted. If NULL then the values are plotted sequentially in their original order.
	Ignored
best_only	Plot only the best fit the empirical (inverse cumulative) patch-size distribution with an overlay of the estimated fits.
plrange	Plot the power-law range

Details

The plot function will produce a figure summarizing the changes in patch size distributions along a set of values. The figure has two panels:

• the upper panel shows the percolation status of empty (FALSE) and occupied cells (TRUE), and shows the mean value (proportion of TRUE values). The background shows the proportion of each type of distribution for each unique values of the along vector.

• the bottom panel displays the power-law range

The plot_distr function displays each distribution in an individual facet, with an overlay of the best distribution fit and a blue bar showing the power-law range. If appropriate, a grey ribbon is used to display the expected distribution given the null expectation (i.e. when plot_distr is called on the results of indictest(). This function can produce quite crowded graphs, but it displays in full the shape of the distributions, and can be useful e.g. to assess the quality of the fits.

See Also

```
patchdistr_sews
```

Examples

```
data(forestgap)
psd_indic <- patchdistr_sews(forestgap)

plot(psd_indic, along = forestgap.pars[ ,"d"])

# When along is non-numeric, bars are used for display
plot(psd_indic, along = as.factor(forestgap.pars[ ,"d"]))

# Display individual distributions
plot_distr(psd_indic, along = forestgap.pars[ ,"d"])

# We can display the distributions along with the null expectation after
# indictest() is run
psd_test <- indictest(psd_indic, nulln = 19)
plot_distr(psd_test, along = forestgap.pars[ ,"d"])</pre>
```

Description

Export the observed and fitted patch size distributions

Usage

```
## S3 method for class 'patchdistr_sews_single'
predict(object, ..., newdata = NULL, best_only = FALSE, xmin_rescale = FALSE)
```

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Arguments

object An patchdistr_sews object
... Additional arguments (ignored)

newdata A vector of patch sizes at which the fit is returned (default to 200 regularly-

spaced values).

best_only Return values for only the best fit of each element (matrix) in object, or return

the values for all fitted distribution.

xmin_rescale If the xmin value used for fits is above one, then setting this to TRUE will rescale

the predicted probabilities so that they align on the cumulative distribution of

the observed patch sizes

Details

The function patchdistr_sews fits competing distribution models to the observed patch size distributions. This functions is able to export the observed values and the fitted values altogether.

Value

A list with component obs, a data.frame containing the observed distribution values and pred, a data.frame containing the fitted values. Both data.frames have columns matrixn, the number of the matrix for which values are given, type, the fitted type of distribution, as well as patchsize and y, the patch size and value of the inverse cumulative distribution function (i.e. 1 - P(x < k)).

See Also

```
patchdistr_sews
```

Examples

```
patch_indics <- patchdistr_sews(forestgap)
predict(patch_indics)</pre>
```

patchsizes

Get patch sizes.

Description

Get the distribution of patch sizes from a logical matrix

Usage

```
patchsizes(mat, merge = FALSE, nbmask = "von_neumann", wrap = FALSE)
```

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Arguments

mat	A logical matrix or a list of such matrices.
merge	Controls whether the obtained patch size distributions are to be pooled together if mat is a list of matrices.
nbmask	Either "moore" for 8-way neighborhood, "von_neumann" for four-way neighborhood (default), or a 3x3 matrix describing which neighbors to consider around a cell. See patchsizes for details on how to specify such neighborhoods.
wrap	Whether to wrap around lattice boundaries ('TRUE'/'FALSE'), effectively using periodic boundaries.

Value

If mat is a logical matrix, then the function returns a vector of patch sizes. If mat is a list of logical matrices, then it returns a list of vectors of patch sizes: this list is flattened if merge is TRUE.

See Also

label

Examples

Description

Spatial early-warning signals: display of trends

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Usage

```
## S3 method for class 'simple_sews_test'
plot(x, along = NULL, what = "value", display_null = TRUE, ...)
## S3 method for class 'simple_sews_list'
plot(x, along = NULL, ...)
```

Arguments

X	A simple_sews_test object (as provided by **_sews functions, such as generic_sews()
along	A vector providing values over which the indicator trend will be plotted. If NULL then the values are plotted sequentially in their original order.
what	The trendline to be displayed. Defaults to the indicator's values ("value") but other metrics can be displayed. Accepted values are "value", "pval", "difference" (obs - null mean), or "z_score" ((obs - null mean) / (null sd)).
display_null	Chooses whether a grey ribbon should be added to reflect the null distribution. Note that it can not be displayed when the trend line reflects something else than the indicator values (when what is not set to "value").
	Other arguments are ignored.

Details

Note that the produced plot is adjusted depending on whether along is numeric or not.

plot_spectrum	Display the r-spectrum of a spectral_sews object	

Description

Display the r-spectrum (or multiple spectra) that are contained in an object returned by spectral_sews object (or the result of indictest applied on such object).

Usage

```
plot_spectrum(x, along = NULL, log = TRUE, display_null = TRUE, ...)
```

Arguments

X	An object produced by spectral_sews or the result returned by indictest applied on such object
along	A vector providing numerical or categorical values along which the indicator trends will be plotted. If NULL, then the indicator values are plotted sequentially in their original order.
log	Whether to use a log scale or a linear scale on the y axis

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display_null Whether to display null information. This argument is ignored if x has not been produced through indictest (and thus does not contain data regarding the null model)
... Other arguments are ignored

See Also

rspectrum, spectral_sews, extract_spectrum

pl_fit

Distribution-fitting functions

Description

These functions fit parametric distributions to a set of discrete values.

Usage

```
pl_fit(dat, xmin = 1)
exp_fit(dat, xmin = 1)
lnorm_fit(dat, xmin = 1)
tpl_fit(dat, xmin = 1)
```

Arguments

dat The set of values to which the distribution are fit

xmin The minimum possible value to consider when fitting the distribution

Details

These functions will fit distributions to a set of values using maximum-likelihood estimation. In the context of the 'spatialwarnings' package, they are most-often used to fit parametric distributions on patch size distributions. As a result, these functions assume that the data contains only integer, strictly positive values. The type of distribution depends on the prefix of the function: 'pl' for power-law, 'tpl' for truncated power-law, 'lnorm' for lognormal and 'exp' for an exponential distribution.

In the context of distribution-fitting, 'xmin' represents the minimum value that a distribution can take. It is often used to represent the minimum scale at which a power-law model is appropriate (Clauset et al. 2009), and can be estimated on an empirical distribution using xmin_estim. Again, please note that the fitting procedure assumes here that xmin is equal or grater than one.

Please note that a best effort is made to have the fit converge, but it may sometimes fail when the parameters are far from their usual range, and numerical issues may occur. It is good practice to make sure the fits are sensible when convergence warnings are reported.

For reference, the shape of the distributions is as follow:

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power-law x^{-a} where a is the power-law exponent

exponential exp(-bx) where b is the truncation rate of the exponential

truncated power-law $x^{-a}exp(-bx)$ where a and b are the exponent of the power law and the rate of truncation

The lognormal form follows the standard definition.

The following global options can be used to change the behavior of fitting functions and/or produce more verbose output:

spatialwarnings.constants.reltol the relative tolerance to use to compute the power-law normalizing constant

$$sum_{k=1}^{\infty} x^{ak} e^{-bk}$$

. Increase to increase the precision of this constant, which can be useful in some cases, typically with large sample sizes. Default is 1e-8.

spatialwarnings.constants.maxit the maximum number of iterations to compute the normalizing constant of a truncated power-law. Increase if you get a warning that the relative tolerance level (defined above) was not reached. Default is 1e8

spatialwarnings.debug.fit_warn_on_bound logical value. Warn if the fit is at the boundary of the valid range for distribution parameter

spatialwarnings.debug.fit_warn_on_NA logical value. Warn if the returned fit has NA/NaN parameters

Value

A list containing at list the following components:

type The type of distribution fitted (as a character string)

method The method used for the fit - here, maximum likelihood, 'll'

11 The log likelihood at the estimated parameter values

xmin The value of xmin used for the fit

npars The number of parameters of the distribution

Additionally, this list may have one or more of the following elements depending on the type of distribution that has been fitted:

plexpo The exponent of the power-law

cutoff The rate of truncation, for truncated power law and exponential fits

meanlog The mean of the lognormal distribution

sdlog The s.d. of the lognormal distribution

References

Clauset, Aaron, Cosma Rohilla Shalizi, and M. E. J. Newman. 2009. "Power-Law Distributions in Empirical Data." SIAM Review 51 (4): 661–703. https://doi.org/10.1137/070710111.

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See Also

```
patchdistr_sews, xmin_estim
```

Examples

```
# Fit an exponential model to patch size distribution
exp_fit(patchsizes(forestgap[[8]]))
# Use the estimated parameters as an indicator function

get_truncation <- function(mat) {
    c(exp_cutoff = exp_fit(patchsizes(mat))$cutoff)
}
trunc_indic <- compute_indicator(forestgap, get_truncation)
plot(trunc_indic)
plot(indictest(trunc_indic, nulln = 19))</pre>
```

raw_cg_moran

Moran's Index at lag of 1

Description

This functions computes the Moran's spatial correlation index (with lag one) on a matrix.

Usage

```
raw_cg_moran(mat, subsize = 1)
```

Arguments

mat A matrix

subsize logical. Dimension of the submatrix used to coarse-grain the original matrix (set

to 1 for no coarse-graining).

Value

The Moran's I index measuring autocorrelation at lag 1 as a named vector

References

Dakos, V., van Nes, E. H., Donangelo, R., Fort, H., & Scheffer, M. (2010). Spatial correlation as leading indicator of catastrophic shifts. Theoretical Ecology, 3(3), 163-174.

Legendre, P., & Legendre, L. F. J. (2012). Numerical Ecology. Elsevier Science.

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See Also

```
generic_sews
```

Examples

```
data(serengeti)
raw_cg_moran(serengeti[[1]], subsize = 1)
```

raw_cg_skewness

Skewness indicator

Description

Compute the spatial skewness of spatial data (a matrix).

Usage

```
raw_cg_skewness(mat, subsize = 5, absolute = TRUE)
```

Arguments

mat A matrix. The matrix values can be logical, with FALSE (empty) or TRUE (occu-

pied) values. The entries can also be continuous (like NDVI or EVI data).

subsize Dimension of the submatrix used to coarse-grain the original matrix. This must

be an integer less than size of the full matrix. Coarse-graining reduces the size of the matrix by a factor subsize in each dimension of the matrix. Skewness is

calculated on the coarse-grained matrix.

absolute Should the function return the absolute value or raw value of skewness?

Details

Spatial skewness is a measure of fluctuations in space; specifically, it measures if fluctuations are getting biased (skewed) in one direction. Based on the theory of critical slowing down, when systems approach critical points they are expected to show increased fluctuations in space. Thus, increasing spatial skewness is proposed as an early warning signal of impending critical transitions.

Computing spatial skewness is straightforward. However, detecting trends of skewness that correspond to critical slowing down can be tricky, especially if data come from discrete classification of state variable.

Many high resolution spatial data are classified as FALSE (empty) or TRUE (occupied by plant). In such cases, spatial skewness captures just the skewness in data, but not that of spatial structure. To resolve the issue, this function employs a method called coarse-graining, proposed in Kefi et al (2014), and described in detail in Sankaran et al. (2017). One must specify a subsize above one for binary valued data sets to obtain meaningful values.

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subsize has to be an integer. It has to be less than or equal to half of matrix size (N). subsize must also be preferably a divisor of N. If it is not a divisor of N, the remainder rows and columns are discarded when computing coarse-graining matrices.

Null model evaluations are also done on coarse-grained matrices.

Value

The spatial skewness of the matrix as a named vector

References

Guttal, V., and Jayaprakash, C. (2009). Spatial variance and spatial skewness: leading indicators of regime shifts in spatial ecological systems. Theoretical Ecology, 2(1), 3-12.

Kefi, S., Guttal, V., Brock, W.A., Carpenter, S.R., Ellison, A.M., Livina, V.N., et al. (2014). Early Warning Signals of Ecological Transitions: Methods for Spatial Patterns. PLoS ONE, 9, e92097.

Sankaran, S., Majumder, S., Kefi, S., and Guttal, V. (2017). Implication of being discrete and spatial in detecting early warning signals of regime shifts. Ecological indicators.

See Also

```
generic_sews
```

Examples

```
data(serengeti)
raw_cg_skewness(serengeti[[1]])
compute_indicator(serengeti, fun = raw_cg_skewness, subsize = 5)
```

raw_cg_variance

Spatial variance indicator

Description

This functions computes the spatial variance of a matrix.

```
raw_cg_variance(mat, subsize = 5)
```

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Arguments

mat A matrix. Its values can be logical, with FALSE (empty) or TRUE (occupied)

values. The entries can also be continuous (like NDVI or EVI data).

subsize Dimension of the submatrix used to coarse-grain the original matrix. This must

be an integer less than size of the full matrix. Coarse-graining reduces the size of the matrix by a factor subsize in each dimension of the matrix. Variance is

calculated on the coarse-grained matrix.

Details

Spatial variance is a measure of fluctuations in space. Based on the theory of critical slowing down, when systems approach critical points they are expected to show increased fluctuations in space. Thus, increasing spatial variance is proposed as an early warning signal of impending critical transitions.

Many high resolution spatial data are classified as FALSE (empty) or TRUE (occupied). In such cases, spatial variance captures just the variance in data, but not that of spatial structure. To resolve the issue, this function employs a method called coarse-graining, proposed in Kefi et al (2014), and described in detail in Sankaran et al. (2017). One must specify a subsize above one for binary valued data sets to obtain meaningful values.

subsize has to be an integer. It has to be less than or equal to half of matrix size (N). subsize must also be preferably a divisor of N. If it is not a divisor of N, the remainder rows and columns are discarded when computing coarse-graining matrices.

Null model evaluations are also done on coarse-grained matrices.

Value

The variance of the coarse-grained matrix as a named vector

References

Guttal, V., and Jayaprakash, C. (2009). Spatial variance and spatial skewness: leading indicators of regime shifts in spatial ecological systems. Theoretical Ecology, 2(1), 3-12.

Kefi, S., Guttal, V., Brock, W.A., Carpenter, S.R., Ellison, A.M., Livina, V.N., et al. (2014). Early Warning Signals of Ecological Transitions: Methods for Spatial Patterns. PLoS ONE, 9, e92097.

Sankaran, S., Majumder, S., Kefi, S., and Guttal, V. (2017). Implication of being discrete and spatial in detecting early warning signals of regime shifts. Ecological Indicators.

See Also

```
generic_sews
```

```
data(serengeti)
raw_cg_variance(serengeti[[1]])
compute_indicator(serengeti, fun = raw_cg_variance, subsize = 5)
```

raw_clustering 39

raw_clustering	Clustering of pairs

Description

Compute the number of pairs of neighbor cells in a landscape and derive from it clustering indices

Usage

```
raw_clustering(mat, wrap = TRUE, use_8_nb = FALSE)
pair_counts(mat, wrap = TRUE, use_8_nb = FALSE, prop = TRUE)
```

Arguments

mat	A matrix, usually with discrete values (logical, integers, etc.)
wrap	Whether space should be considered to wrap at the edges
use_8_nb	Set to TRUE to use an 8-way neighborhood. The default is set to FALSE, which uses a 4-way neighborhood
prop	Return the counts for all pairs in the matrix (FALSE), or the proportions for each pair (out of all possible pairs)

Details

The clustering of pairs is defined as the density of pairs, i.e. the proportion of all neighboring pairs of cells that share the same state, divided by the null expectation given a random, homogeneous spatial structure.

For example, let's consider a matrix with two states, 'a' and 'b'. raw_clustering will count all pairs of cells 'a-a' or 'b-b', and divide this by the total number of pairs. This proportion is then again divided by the probability of obtaining these proportion of pairs under the assumption of no spatial structure (random mixing of cells in the matrix).

Clustering is equal to one when there is no spatial structure. It is above one when two states are found next to each other (i.e. cluster) more than expected by chance. Values below one means that those two states tend to be neighbors less frequently than expected by chance.

If you are only interested in the proportion of pairs for each combination of states, you can use the function pair_counts, which returns a matrix with as many rows and columns as there are states in the matrix, and contains the counts for all possible pairs of cells found in the matrix (or their relative proportions).

Value

A vector with the requested clutering values for raw_clustering, whose names are equal to each state (unique value) found in the original matrix, preceded by 'clust_' (to make sure names are compatible with other functions).

pair_counts returns the counts of pairs of states in the matrix, or the proportion of each pair, depending on the value of prop

Examples

```
# The clustering of a random matrix is close to one
ls <- 100 # lattice size
mm <- matrix(sample(c("sp1", "sp2", "sp3", "sp4"), size = ls^2, replace = TRUE),</pre>
             nrow = ls, ncol = ls)
clust <- raw_clustering(mm, wrap = TRUE, use_8_nb = TRUE)</pre>
print(clust)
# Compute clustering along the gradient for the serengeti dataset
data(forestgap)
clust_indic <- compute_indicator(serengeti, raw_clustering,</pre>
                                  wrap = TRUE, use_8_nb = FALSE)
# The interesting one is the clustering of state 0 (FALSE in the original matrix),
# which corresponds to grassland pixels, which get more and more clustered with
# increasing rainfall (see also ?generic_sews for how that compares with generic
# indicators)
plot(clust_indic, along = serengeti.rain)
# Add null trend
clust_test <- indictest(clust_indic, nulln = 19)</pre>
plot(clust_test, along = serengeti.rain)
# Show the proportion of each pairs of states in the matrix...
pair_counts(serengeti[[5]])
# ... or the total count
pair_counts(serengeti[[5]], prop = FALSE)
```

```
raw_flowlength_uniform
```

Flow length (uniform slope)

Description

Compute a simple approximation of the flow length assuming a constant slope

raw_kbdm 41

Usage

```
raw_flowlength_uniform(mat, slope, cell_size)
```

Arguments

mat	The input	matrix (must	he a i	logical	matrix)
ma t	I IIC IIIDut	mauna	must	oc a	iogicai	mau i A

slope The slope of the area documented by the matrix (in degrees).

cell_size The horizontal size of a cell in the matrix (as viewed from above).

Details

This function computes the Flowlength of a given matrix, using a uniform approximation (the slope is constant across the whole matrix, with maximum slope being from the top of the matrix to its bottom), as per Rodriguez et al. (2017). See flowlength_sews for more details.

Value

A named vector of length 1 containing the flow length numerical value

References

Rodriguez, F., A. G. Mayor, M. Rietkerk, and S. Bautista. 2017. A null model for assessing the cover-independent role of bare soil connectivity as indicator of dryland functioning and dynamics. Ecological Indicators.

See Also

```
flowlength_sews indictest, to test the significance of indicator values.
```

Examples

```
raw_flowlength_uniform(arizona[[1]], slope = 20, cell_size = 1)
```

|--|

Description

Compute the Kolmogorov complexity of a matrix using the Block Decomposition Method (requires the acss package).

42 raw_moran

Usage

```
raw_kbdm(mat, subsize)
```

Arguments

mat A logical matrix (with TRUE/FALSE values)

subsize A submatrix size to carry out the Block Decomposition Method (must be be-

tween 1 and 3)

Details

The Kolmogorov complexity cannot be computed directly for large strings (i.e. matrices). However, the complexity of smaller submatrices can be estimated, then combined to obtain an approximation of the complexity of the whole matrix. This method, the Block Decomposition Method is implemented in this function. See also kbdm_sews for more details.

Value

The KBDM numeric value as a named vector

See Also

```
kbdm_sews, acss
```

Examples

```
raw_kbdm(forestgap[[1]], subsize = 3)
```

raw_moran

Spatial correlation at lag 1

Description

This function computes the Moran's I index of spatial correlation at lag 1.

Usage

```
raw_moran(mat)
```

Arguments

mat

A matrix containing logical, numeric or integer values

raw_plrange 43

Details

This function returns the spatial correlation as measured by the Moran's I index. If the variance of the matrix is zero, then NaN is returned. This function assumes a 4-way neighborhood (a.k.a. von-Neumann neighborhood), and does not wrap around at the sides of the matrix.

Value

The Moran's I numeric value as a numeric number.

See Also

```
generic_sews
```

Examples

raw_plrange

Power-law range indicator

Description

Compute the power-law range of a matrix

Usage

```
raw_plrange(mat, xmin_bounds = NULL)
```

Arguments

mat A logical matrix, or a list of logical matrices

xmin_bounds A vector of two integer values, defining a range in which to search for the best

xmin (see Details).

44 raw_sdr

Details

Some ecosystems show typical changes in their patch-size distribution as they become more and more degraded. In particular, an increase in the truncation of the patch-size distribution (PSD) is expected to occur. The power-law range (PLR) measures the truncation of the PSD in a single value (see also patchdistr_sews for more details).

To compute the PLR, power-laws are fitted with a variable minimum patch size (xmin) and the one with the lowest Kolmogorov-Smirnov distance to the empirical distribution is retained. PLR is then computed using this best-fitting xmin:

$$\frac{log(x_{max}) - log(x_{min})}{log(x_{max}) - log(x_{smallest})}$$

where x_{max} is the maximum observed patch size, and $x_{smallest}$ is the minimum observed patch size.

Value

A named vector containing the power-law range value

References

Clauset, A., Shalizi, C. R., & Newman, M. E. (2009). Power-law distributions in empirical data. SIAM review, 51(4), 661-703.

Berdugo, M., Kefi, S., Soliveres, S. & Maestre, F.T. (2017). Plant spatial patterns identify alternative ecosystem multifunctionality states in global drylands. Nature in Ecology and Evolution.

See Also

```
patchdistr_sews
```

Examples

```
forestgap.plr <- raw_plrange(forestgap[[2]])
# Restrict to small xmins
forestgap.plr2 <- indicator_plrange(forestgap[[2]], xmin_bounds = c(1, 10))</pre>
```

raw_sdr

Spectral Density Ratio (SDR) indicator

Description

Compute the ratio of low frequencies over high frequencies of the r-spectrum.

```
raw_sdr(mat, sdr_low_range = NULL, sdr_high_range = NULL)
```

raw_sdr 45

Arguments

A matrix with continuous values, or a logical matrix (TRUE/FALSE values).

sdr_low_range

The range of values (in proportion) to use for the computation of the spectral density ratio. For example, for the lowest 20% (default value), set sdr_low_range to c(0, .2).

sdr_high_range

The range of values (in proportion) to use for the computation of the spectral density ratio. For example, for the highest 20% (default value), set sdr_high_range to c(.8, 1).

Details

SDR measures the increase in long-range correlations before a critical point. It is the ratio of the average low frequency value over high frequency values. In this implementation, an increase in SDR implies a "reddening" of the r-spectrum. See also spectral_sews for a more complete description.

Low and high frequencies are averaged in order to compute the SDR. The parameters sdr_low_range and sdr_high_range control which frequencies are selected for averaging. For example $sdr_low_range = c(0, .2)$ (default) uses the lower 20 the average of low frequencies. $sdr_high_range = c(.8, 1)$ uses the higher 20

Value

The SDR values computed on the matrix as a named vector

References

Carpenter, S.R. & Brock, W.A. (2010). Early warnings of regime shifts in spatial dynamics using the discrete Fourier transform. Ecosphere

See Also

```
indictest, rspectrum, plot_spectrum, spectral_sews, extract_spectrum
```

raw	st	rι	ıct	:va	r

Structural variance

Description

Compute the structural variance on a matrix.

Usage

```
raw_structvar(mat, model = "sph", nmax = 100000L, nbins = 32, cutoff = NULL)
```

Arguments

mat	A matrix
model	The variogram model to use, either "sph" (for a spherical model) or "exp" (for an exponential model)
nmax	The maximum number of pairs of cells to use when computing the variogram
nbins	Number of distance bins to use to compute the variogram
cutoff	Maximum distance to consider in the variogram. If NULL, then a distance equal to one third of the diagonal of the matrix is used

See Also

```
raw_variogram_metrics, variogram_sews
```

Examples

```
raw_structvar(serengeti[[5]])
```

```
{\tt raw\_variogram\_metrics} \ \ \textit{Variogram parameters}
```

Description

Compute the nugget, partial sill, correlation range and structural variance on a matrix.

```
raw_variogram_metrics(
  mat,
  model = "sph",
  nmax = 100000L,
  nbins = 32,
  cutoff = NULL
)
```

rspectrum 47

Arguments

mat	A matrix
model	The variogram model to use, either "sph" (for a spherical model) or "exp" (for an exponential model)
nmax	The maximum number of pairs of cells to use when computing the variogram
nbins	Number of distance bins to use to compute the variogram
cutoff	Maximum distance to consider in the variogram. If NULL, then a distance equal

to one third of the diagonal of the matrix is used

See Also

variogram_sews, raw_structvar

Examples

raw_variogram_metrics(serengeti[[5]])

Description

Compute the r-spectrum of a matrix

Usage

```
rspectrum(mat)
```

Arguments

mat

A matrix with logical or numeric values

Details

This functions returns a data.frame with NAs in the rspec column if the input matrix has zero variance. Note that if the matrix is not square, then only the largest square matrix fitting in the upper left corner is used.

Value

A data.frame with two columns: dist, the wave number and rspec, the normalized value of the r-spectrum

See Also

```
spectral_sews
```

48 serengeti

Examples

```
# Spectrum of white noise
rmat <- matrix(runif(100*100) > .5, ncol = 100)
spec <- rspectrum(rmat)
plot(spec, type = "l")

# Add some spatial correlation and compare the two spectra
rmat.cor <- rmat
for (i in seq(1, nrow(rmat)-1)) {
   for (j in seq(1, nrow(rmat)-1)) {
     rmat.cor[i,j] <- mean(rmat[(i-1):(i+1), (j-1):(j+1)])
   }
}
spec.cor <- rspectrum(rmat.cor)
plot(spec.cor, type = "n")
lines(spec, col = "black")
lines(spec.cor, col = "blue")</pre>
```

serengeti

Serengeti dataset

Description

Vegetation data along a rainfall gradient in Serengeti national park.

Usage

```
serengeti.rain
```

Format

A list of logical matrices

The annual rainfall corresponding to the matrices in the serengeti dataset, in the corresponding order.

Details

The data-set consists of a rectangular area of size 7.5 km x 90 km. These data are represented as a list of matrices. Each matrix is a moving window of 7.5 km x 7.5 km which moves my 2.5 km along the length of the rectangular data-set.

Each entry in the matrix is vegetation data at a resolution of 30m as classified into binary units with FALSE (grass) and TRUE (forest). The rainfall data provided here is the average rainfall (mm/yr) of a moving window of dimension 7.5km which moves my 2.5 km along the length of the rectangular data-set.

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Source

Extracted from Eby's et al (2017) supplementary material https://github.com/tee-lab/spacetime-csd/

References

Eby, S., Agrawal, A., Majumder, S., Dobson, A.P. & Guttal, V. (2017). Alternative stable states and spatial indicators of critical slowing down along a spatial gradient in a savanna ecosystem: Global Ecology and Biogeography, 26, 638-649

Reed, D. N., Anderson, T. M., Dempewolf, J., Metzger, K., & Serneels, S. (2009). The spatial distribution of vegetation types in the Serengeti ecosystem: the influence of rainfall and topographic relief on vegetation patch characteristics. Journal of Biogeography, 36(4), 770-782.

simple_sews

simple_sews objects

Description

This help page describes the structure of simple_sews_* objects, such as those defined by the classes simple_sews_single, simple_sews_list

Details

The spatialwarnings uses S3 objects (lists) internally to store indicator values, along with the necessary data to plot and display results. It is not recommended to extract data directly from these objects, as they are subject to change with different releases of the package. The preferred method is to use dedicated generic functions such as plot() or as.data.frame() to display or export the results. Nonetheless, we document the structure of these objects here for reference.

simple_sews objects are returned by all indicator functions that return numeric values. This includes for example generic_sews, flowlength_sews, compute_indicator but *not* patchdistr_sews or spectral_sews, which provide indicators that depend on non-numeric values (e.g. patch-size distribution type), or need to store more information than just a single numerical value (e.g. the spectrum of the input matrix).

simple_sews objects come in multiple variants: simple_sews_single is the result of an indicator function applied to a single matrix, and simple_sews_test_single is the result of indictest applied to a simple_sews_single object. Both these objects have list equivalents, simple_sews_list and simple_sews_test_list which are simply a collection of their 'single' equivalent. These 'list' objects are used to store the results of computations when working with multiple matrices.

A simple_sews_single object is a list with the following components

itemvalue the indicator values. A vector of length one if there is only one numeric value returned by the indicator function (e.g. flowlength_sews, or with a length above one otherwise

orig_data the original matrix on which the indicator was computed

fun.args the argument used in the call to the indicator function (the function that given a matrix, returns the spatial metrics of interest)

taskname a character string describing the current indicator(s) being computed

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indicf the indicator function, which given the matrix, returns the spatial metric(s) of interest

simple_sews_test_single have all of the above components, plus the following:

nulldistr the null distribution of values, with nulln rows and as many columns as the number of values returned by the indicator function

null_mean the mean indicator values in the null distribution

null_sd the standard deviation of the null distribution

null_qsup the upper quantile of the null distribution, by default the 95 adjust this

null_qinf the upper quantile of the null distribution, by default the 05 adjust this

z_score the z_score of the observed value relative to the null distribution, i.e. (value - null_mean) / null sd

pval the p-value of the indicator, i.e. the proporation of values of the null distribution that fall below the observed indicator value

null_method the method used to produce the null matrices. See indictest for details

nulln the number of null matrices used

get_nullmat a function that can be called to obtain a randomized matrix

matrixn the number of the matrix, can be above one if the computations have been run on a list of matrices, or non-existent if only one matrix was used

See Also

custom_indicator

spatialwarnings

Early Spatial-Warnings of Ecosystem Degradation

Description

Many dynamical systems such as ecosystems exhibit non-linear responses to changes in their external drivers, resulting in possible wide state shifts with strong ecological or economical consequences. This often happens when a system exhibit a change in its stability properties as a threshold is crossed, *e.g.* going from multiple stable states to a single stable state. For a few decades, much research has been dedicated to finding a way to anticipate these tipping points in ecological systems. This has led to the development of indicators, which are metrics based on spatial structure, that could reflect the proximity of an ecosystem to a tipping point.

This package implements many of these indicators, or *early-warning signals* (EWS), on spatial raster data. High-level functions and methods provide familiar workflows to compute several related indicators at once, and display their variations along environmental gradients or time-series. Lower-level functions are also available to integrate early-warning signals into different workflows.

Main, "workflow" functions provided by this package:

"Workflow" functions, which may compute several indicators at once:

• generic_sews: Generic spatial EWS

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- spectral_sews: Spectrum-based EWS
- patchdistr_sews: EWS based on patch-size distributions
- kbdm_sews: Kolmogorov complexity
- flowlength_sews: Flow length
- variogram_sews: Variogram-based indicators
- lsw_sews: Indicators based on the Lifshitz-Slyozov-Wagner distribution

Individual indicator functions, which may be used to compute raw indicator values

- raw_cg_moran: lag-1 spatial autocorrelation
- raw_cg_variance: Spatial variance
- raw_cg_skewness: Spatial skewness
- raw_sdr: Spectral density ratio (SDR)
- indicator_psdtype: Patch-size distribution shape
- indicator_plrange, raw_plrange : Power-law range
- raw_kbdm: Kolmogorov complexity
- raw_flowlength_uniform: Flow Length
- raw_structvar: Structural variance
- raw_lsw_aicw: Support for LSW distribution relative to lognormal
- raw_patch_radii_skewness: Skewness of patch radii

The package homepage is available at Github and a user guide is also available. Do not hesitate to get in touch if you want to make changes to the package or add another indicator.

spectral_sews

Spectrum-based spatial early-warning signals.

Description

Computation of spatial early warning signals based on spectral properties.

Usage

```
spectral_sews(mat, sdr_low_range = NULL, sdr_high_range = NULL, quiet = FALSE)
```

Arguments

mat	The input matrix or a list of matrices.
sdr_low_range	The range of values (in proportion) to use for the computation of the spectral density ratio. For example, for the lowest 20% (default value), set sdr_low_range to c(0, .2). See also the Details section.
sdr_high_range	The range of values (in proportion) to use for the computation of the spectral density ratio. For example, for the higher 20% (default value), set sdr_high_range to c(.8, 1). See also the Details section.
quiet	Do not display some warnings

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Details

Spectral early warning signals are based on the fact that some dynamical systems can exhibit an change in specific characteristics of their spatial structure when approaching a transition. In particular, long-range correlations are expected to have an increased importance. This is expected to be reflected in the spectrum of the spatial structure as an increase of the relative importance of lower frequencies over higher frequencies ("reddening" of the spatial spectrum).

This task allows computing the radial-spectrum which gives the relative importance of each space scale as a function of distance, from 1 to N/2 (N being the minimum between the number of rows and columns). If the matrix is not square, then it is cropped to biggest square that fits within the left side of the matrix.

Additionally, it summarizes this spectrum into a Spectral Density Ratio (SDR), which is the ratio of low frequencies over high frequencies of the r-spectrum. The SDR value is expected to increase before a transition point.

The significance of spectral early-warning signals can be estimated by reshuffling the original matrix (function indictest). Indicators are then recomputed on the shuffled matrices and the values obtained are used as a null distribution. P-values are obtained based on the rank of the observed value in the null distribution.

The trend of SDR values can be plotted using the plot() method. Alternatively, the spectrum itself can be plotted (with facets if multiple input matrices were used) using the plot_spectrum method.

Value

Function spectral_sews object of class spectral_sews_list or spectral_sews_single depending on whether the input was a list of matrices or a single matrix.

Function indictest

The plot methods returns a ggplot object (usually displayed immediately when called interactively).

References

Kefi, S., Guttal, V., Brock, W.A., Carpenter, S.R., Ellison, A.M., Livina, V.N., et al. (2014). Early Warning Signals of Ecological Transitions: Methods for Spatial Patterns. PLoS ONE, 9, e92097.

See Also

```
rspectrum, plot_spectrum, raw_sdr, extract_spectrum indictest, to test the significance of indicator values.
```

variogram_sews 53

```
sdr_high_range = c(.8, 1)
summary(spec_indic)
# Display trends along the varying model parameter
plot(spec_indic, along = serengeti.rain)
# Computing spectra many times is expensive, consider setting parallel
# computing using: options(mc.cores = n)
# Assess significance
spec_test <- indictest(spec_indic, nulln = 199)</pre>
summary(spec_test)
# Display the SDR trend, now with a grey ribbon representing 5%-95%
# quantiles of the null distribution
plot(spec_test, along = serengeti.rain)
# Add a line highlighting the shift
if (require(ggplot2)) {
  plot(spec_test, along = serengeti.rain) +
    geom_vline(xintercept = 590, color = "red", linetype = "dashed")
}
# Display radial-spectra
plot_spectrum(spec_indic, along = serengeti.rain)
# Graphics can be modified using ggplot2 functions
if (require(ggplot2)) {
  plot_spectrum(spec_indic, along = serengeti.rain) +
    scale_y_log10()
}
```

variogram_sews

Early-Warning signals based on variograms (EXPERIMENTAL)

Description

Compute Early-warning signals based on metrics derived form semi-variograms.

```
variogram_sews(mat, model = "sph", nmax = 1e+05, nbins = 32, cutoff = NULL)
```

54 variogram_sews

Arguments

mat A matrix (TRUE/FALSE values) or a list of matrices

model The variogram model to use, either "sph" (for a spherical model) or "exp" (for an exponential model)

nmax The maximum number of pairs of cells to use when computing the variogram

nbins Number of distance bins to use to compute the variogram

cutoff Maximum distance to consider in the variogram. If NULL, then a distance equal to one third of the diagonal of the matrix is used

Details

During ecosystem degradation and especially before a regime shift occurs in some ecosystems, spatial autocorrelation is expected to increase in a landscape. This increase can be measured based on variograms, which represent how the difference (variance) between two points in a landscape varies as a function of distance.

The approach used to derive variogram-based EWS is to compute the empirical variogram of a land-scape (represented passed as a matrix of values), then fit a variogram model to it. Three parameters are then extracted from the variogram model (see Nijp et al. 2019 for a visual description of these parameters):

- 1. The *nugget* (intercept)
- 2. The partial sill, i.e. the reduction in semivariance at distance zero
- 3. The *correlation range*, i.e. the distance at which the relationship between semivariance and distance flattens

Additionally, the *structural variance* is computed as (partial sill)/(nugget + partial sill), wich quantifies whether the data are spatially structured (structural variance of one), or completely unstructured (value of zero). Theoretical work suggests that partial sill, correlation range and structural variance should increase before a regime shift occurs in an ecosystem (Nijp et al. 2019).

This function offers to fit a spherical model or an exponential model. The best-fitting model depends on your data, you should try different options and review the fits using plot_variogram.

Please note that this part of the package is still experimental and deserves more testing.

Value

A list object of class "variogram_sews", that can be displayed using summary(), plot(), etc. Significance of values can be tested using indictest.

References

Nijp, Jelmer J., Arnaud J.A.M. Temme, George A.K. Voorn, Lammert Kooistra, Geerten M. Hengeveld, Merel B. Soons, Adriaan J. Teuling, and Jakob Wallinga. (2019) Spatial Early Warning Signals for Impending Regime Shifts: A Practical Framework for Application in Real-world Landscapes. Global Change Biology 25 (6): 1905-21. doi:10.1111/gcb.14591

variogram_sews_plot 55

See Also

```
raw_structvar, plot_variogram, extract_variogram, predict
indictest, to test the significance of indicator values.
```

Examples

variogram_sews_plot

Early-warning signals based on variograms

Description

Plot trends of indicators based on variograms

```
## S3 method for class 'variogram_sews'
plot(x, along = NULL, ...)

plot_variogram(x, along = NULL, ...)

## S3 method for class 'variogram_sews'
plot_variogram(x, along = NULL, ...)

## S3 method for class 'variogram_sews_test'
plot(x, along = NULL, what = "value", display_null = TRUE, ...)

## S3 method for class 'variogram_sews_test'
plot_variogram(x, along = NULL, what = "value", display_null = TRUE, ...)
```

variogram_sews_plot

Arguments

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X	An object produced by variogram_sews, or the result of applying indictest on such object.
along	A vector providing values along which the indicator trends will be plotted. If NULL then the indicator values are plotted sequentially in their original order.
	Other arguments are ignored.
what	The trendline to be displayed. Defaults to the indicator's values ("value") but other metrics can be displayed. Accepted values are "value", "pval", "difference" (obs - null mean), or "z_score" ((obs - null mean) / (null sd)).
display_null	Chooses whether a grey ribbon should be added to reflect the null distribution. Note that it can not be displayed when the trend line reflects something else than the indicator values (when what is not set to "value").

Details

The plot() function will display how the estimated variogram parameters change along a set of values (passed with argument along). If the object passed has been processed through indictest, then the null values are also displayed. plot_variogram() can be used to display the individual variograms that have been fit to the data.

See Also

```
variogram_sews, indictest, plot_variogram
```

```
serengeti_ews <- variogram_sews(serengeti, model ="exp")

# Display the change in variogram parameters
plot(serengeti_ews, along = serengeti.rain) +
    ggplot2::labs(x = "Rainfall (mm)")

# Visualize the fitted variograms
plot_variogram(serengeti_ews, along = serengeti.rain)

# Test the trends (nulln should be set to a higher value to obtain
    # meaningful results
    serengeti_test <- indictest(serengeti_ews, nulln = 19)
    plot(serengeti_test, along = serengeti.rain)
    plot_variogram(serengeti_test, along = serengeti.rain)</pre>
```

variogram_sews_predict

```
variogram_sews_predict
```

predict() method for variogram_sews objects

Description

Export the fitted variogram(s)

Usage

```
## S3 method for class 'variogram_sews_list'
predict(object, newdist = NULL, ...)
```

Arguments

object An object produced by variogram_sews

newdist A vector of distances at which to return the variogram fit values (defaults to 128

regularly-spaced values).

. . . Additional arguments (ignored)

Value

A data.frame with the distances (column dist), the fitted values (gamma), and if object contains more than one matrix, a column matrixn.

See Also

```
variogram_sews
```

```
vario_indics <- variogram_sews(serengeti)
predict(vario_indics)
vario_test <- indictest(vario_indics, nulln = 19)
predict(vario_test) # same result</pre>
```

58 xmin_estim

xmin_estim

Estimate the minimum patch size of a power-law distribution

Description

When fitting a power-law to a discrete distribution, it might be worth discarding points below a certain threshold (xmin) to improve the fit. This function estimates the optimal xmin based on the Kolmogorov-Smirnoff distance between the fit and the empirical distribution, as suggested by Clauset et al. (2009).

Usage

```
xmin_estim(dat, bounds = range(dat))
```

Arguments

dat A vector of integer values

bounds A vector of two values representing the bounds in which the best xmin is searched

Details

The function returns NA if dat has only three unique values or if the power-law fit failed.

Value

The estimated xmin as an integer value

References

Clauset, A., Shalizi, C. R., & Newman, M. E. (2009). Power-law distributions in empirical data. SIAM review, 51(4), 661-703.

See Also

```
patchdistr_sews
```

```
psd <- patchsizes(forestgap[[5]])
xmin_estim(psd)</pre>
```

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