

Package ‘spatialEco’

December 21, 2018

Type Package

Title Spatial Analysis and Modelling Utilities

Version 1.1-1

Date 2018-11-17

Description Utilities to support spatial data manipulation, query, sampling and modelling. Functions include models for species population density, download utilities for climate and global deforestation spatial products, spatial smoothing, multivariate separability, point process model for creating pseudo-absences and sub-sampling, polygon and point-distance landscape metrics, auto-logistic model, sampling models, cluster optimization, statistical exploratory tools and raster-based metrics.

Depends R (>= 3.4.0)

Imports sp, sf, raster, spatstat, cluster, spdep, SDMTools, readr, RCurl, rgeos, RANN, rms, yaImpute, SpatialPack (>= 0.3), MASS, EnvStats, maptools, methods

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License GPL-3

URL <https://cran.r-project.org/package=spatialEco>

NeedsCompilation no

Repository CRAN

LazyData true

Encoding UTF-8

RoxxygenNote 6.1.1

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Date/Publication 2018-12-21 14:30:03 UTC

R topics documented:

ants	4
bearing.distance	5
breeding.density	5
class.comparison	7
classBreaks	9
concordance	10
conf.interval	11
correlogram	12
crossCorrelation	13
csi	16
curvature	17
daymet.point	19
daymet.tiles	20
DAYMET_tiles	21
dispersion	22
dissection	23
download.daymet	24
download.hansen	25
download.prism	27
effect.size	28
elev	29
erase.point	30
focal.lmetrics	31
fuzzySum	33
gaussian.kernel	34
group.pdf	35
hexagons	36
hli	37
hsp	38
hybrid.kmeans	39
idw.smoothing	40
insert.values	41
kde.2D	42
kl.divergence	42
land.metrics	43
local.min.max	46
loess.boot	47
loess.ci	48
logistic.regression	50
moments	52
morans.plot	53
mwCorr	55
nni	55
o.ring	56
oli.aw	58
optimal.k	59

optimized.sample.variance	60
outliers	62
parea.sample	63
plot.effect.size	64
plot.loess.boot	65
point.in.poly	66
poly.regression	68
polyPerimeter	69
pp.subsample	70
print.cross.cor	72
print.effect.size	72
print.loess.boot	73
pseudo.absence	73
pu	76
raster.deviation	77
raster.downscale	79
raster.entropy	80
raster.gaussian.smooth	81
raster.invert	82
raster.kendall	83
raster.mds	85
raster.modified.ttest	86
raster.moments	88
raster.transformation	89
raster.vol	90
raster.Zscore	91
rasterCorrelation	92
remove.holes	93
sa.trans	94
sample.annulus	95
sample.line	96
sample.poly	98
sampleTransect	99
sar	100
separability	101
sg.smooth	102
shannons	104
similarity	105
smooth.time.series	106
sobal	108
sp.kde	109
sp.na.omit	110
spherical.sd	111
srr	113
stratified.random	114
subsample.distance	115
summary.cross.cor	117
summary.effect.size	117

summary.loess.boot	118
swvi	118
tpi	120
trasp	121
trend.line	122
tri	123
vrm	124
winsorize	125
wt.centroid	126
zonal.stats	127

Index	130
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ants

Ant Biodiversity Data

Description

Roth et al., (1994) Costa Rican ant diversity data

Format

A data.frame with 82 rows (species) and 5 columns (covertypes):

species Ant species (family)

Primary.Forest Primary forest type

Abandoned.cacao.plantations Abandoned cacao plantations type

Productive.cacao.plantations Active cacao plantations type

Banana.plantations Active banana plantations type

Source

<http://www.tiem.utk.edu/~gross/bioed/bealsmodules/shannonDI.html>

References

Roth, D. S., I. Perfecto, and B. Rathcke (1994) The effects of management systems on ground-foraging ant diversity in Costa Rica. Ecological Applications 4(3):423-436.

bearing.distance *Bearing and Distance*

Description

Calculates a new point [X,Y] based on defined bearing and distance

Usage

```
bearing.distance(x, y, distance, azimuth, EastOfNorth = TRUE)
```

Arguments

x	x coordinate
y	y coordinate
distance	Distance to new point (in same units as x,y)
azimuth	Azimuth to new point
EastOfNorth	Specified surveying convention

Note

East of north is a surveying convention and defaults to true.

Author(s)

Jeffrey S. Evans <jeffrey_evans@tnc.org>

Examples

```
pt <- cbind( x=480933, y=4479433)
bearing.distance(pt[1], pt[2], 1000, 40)
```

breeding.density *Breeding density areas (aka, core habitat areas)*

Description

Calculates breeding density areas base on population counts and spatial point density.

Usage

```
breeding.density(x, pop, p = 0.75, bw = 6400, b = 8500,
self = TRUE)
```

Arguments

x	sp SpatialPointsDataFrame object
pop	Population count/density column in x@data
p	Target percent of population
bw	Bandwidth distance for the kernel estimate (default 8500)
b	Buffer distance (default 8500)
self	(TRUE/FALSE) Should source observations be included in density (default TRUE)

Value

A list object with:

- pop.pts sp point object with points identified within the specified p
 - pop.area sp polygon object of buffered points specified by parameter b
 - bandwidth Specified distance bandwidth used in identifying neighbour counts
 - buffer Specified buffer distance used in buffering points for pop.area
 - p Specified population percent

Note

The breeding density areas model identifies the Nth-percent population exhibiting the highest spatial density and counts/frequency. It then buffers these points by a specified distance to produce breeding area polygons. If you would like to recreate the results in Doherty et al., (2010), then define `bw = 6400m` and `b[if p < 0.75 b = 6400m, |p |>= 0.75 b = 8500m]`

Author(s)

Jeffrey S. Evans <jeffrey_evans@tnc.org>

References

Doherty, K.E., J.D. Tack, J.S. Evans, D.E. Naugle (2010) Mapping breeding densities of greater sage-grouse: A tool for range-wide conservation planning. Bureau of Land Management. Number L10PG00911

Examples

```
bd75 <- breeding.density(pop, pop='counts', p=0.75, b=8500, bw=6400)
plot(bd75$pop.area, main='75% breeding density areas')
plot(pop, pch=20, col='black', add=TRUE)
plot(bd75$pop.pts, pch=20, col='red', add=TRUE)
```

<code>class.comparison</code>	<i>Class comparison between two nominal rasters</i>
-------------------------------	---

Description

Compares two categorical rasters using Cohen's Kappa (d) or paired t-test statistic(s)

Usage

```
class.comparison(x, y, x.idx = 1, y.idx = 1, d = "AUTO",
  stat = "kappa", sub.sample = FALSE, type = "hexagon", p = 0.1,
  size = NULL)
```

Arguments

<code>x</code>	First raster for comparison, SpatialPixelsDataFrame or SpatialGridDataFrame object
<code>y</code>	Second raster for comparison, SpatialPixelsDataFrame or SpatialGridDataFrame object
<code>x.idx</code>	Index for the column in the x raster object
<code>y.idx</code>	Index for the column in the y raster object
<code>d</code>	Distance for finding neighbors, the default "AUTO" will derive a distance
<code>stat</code>	Statistic to use in comparison ("kappa", "t.test", "both")
<code>sub.sample</code>	Should a subsampling approach be employed (FALSE/TRUE)
<code>type</code>	If sub.sample = TRUE, what type of sample ("random" or "hexagon")
<code>p</code>	If sub.sample = TRUE, what proportion of population should be sampled
<code>size</code>	If sub.sample = TRUE, alternate to proportion of population (p), using fixed sample size

Value

A SpatialPixelsDataFrame or SpatialPointsDataFrame with the following attributes:

- `x` x variable used to derive Kappa (d)
- `y` y variable used to derive Kappa (d)
- `kappa` Kappa (d) statistic
- `t.test` Paired t.test statistic (if `stat = "t.test"` or "both")
- `p.value` p-value of the paired t.test statistic (if `stat = "t.test"` or "both")

Note

This function provides a Cohen's Kappa or paired t-test to compare two classified maps. Point based subsampling is provided for computation tractability. The hexagon sampling is recommended as it is good at capturing spatial process that includes nonstationarity and anisotropy.

Author(s)

Jeffrey S. Evans <jeffrey_evans@tnc.org>

References

Cohen, J. (1960). A coefficient of agreement for nominal scales. *Educational and Psychological Measurement*, 20:37-46

Examples

```
## Not run:
library(sp)
library(raster)

data(meuse.grid)
r1 <- sp::SpatialPixelsDataFrame(points = meuse.grid[c("x", "y")], data = meuse.grid)
r1@data$class1 <- round(runif(nrow(r1), 1,5),0)
r2 <- sp::SpatialPixelsDataFrame(points = meuse.grid[c("x", "y")], data = meuse.grid)
r2@data$class2 <- round(runif(nrow(r2), 1,5),0)

d <- class.comparison(r1, r2, x.idx = 8, y.idx = 8, stat="both")
par(mfrow=c(2,2))
plot(raster(d, layer=3), main="Kappa")
plot(raster(d, layer=4), main="t.test")
plot(raster(d, layer=5), main="t.test p-value")

# Hexagonal sampling
d.hex <- class.comparison(r1, r2, x.idx = 8, y.idx = 8, stat = "both",
                           sub.sample = TRUE, d = 500, size = 1000)
sp::bubble(d.hex, "kappa")
d.hex <- sp.na.omit(d.hex, col.name = "t.test")
sp::bubble(d.hex, "t.test")

# Random sampling
d.rand <- class.comparison(r1, r2, x.idx = 8, y.idx = 8, stat = "both",
                            sub.sample = TRUE, type = "random")
sp::bubble(d.rand, "kappa")

## End(Not run)
```

classBreaks	<i>Class breaks</i>
-------------	---------------------

Description

Finds class breaks in a distribution

Usage

```
classBreaks(x, n, type = c("equal", "quantile", "std", "geometric"))
```

Arguments

x	A vector to find breaks for
n	Number of breaks
type	Statistic used to find breaks c("equal", "quantile", "std", "geometric")

Value

A vector containing class break values the length is n+1 to allow for specification of ranges

Note

The robust std method uses $\text{sqrt}(\text{sum}(x^2)/(n-1))$ to center the data before deriving "pretty" breaks.

Author(s)

Jeffrey S. Evans <jeffrey_evans@tnc.org>

Examples

```
y <- rnbinom(100, 10, 0.5)
classBreaks(y, 10)
classBreaks(y, 10, type="quantile")

par(mfrow=c(2,2))
d <- density(y)
plot(d, type="n", main="Equal Area breaks")
polygon(d, col="cyan")
abline(v=classBreaks(y, 10))
plot(d, type="n", main="Quantile breaks")
polygon(d, col="cyan")
abline(v=classBreaks(y, 10, type="quantile"))
plot(d, type="n", main="Robust Standard Deviation breaks")
polygon(d, col="cyan")
abline(v=classBreaks(y, 10, type="std"))
plot(d, type="n", main="Geometric interval breaks")
polygon(d, col="cyan")
abline(v=classBreaks(y, 10, type="geometric"))
```

```
( y.breaks <- classBreaks(y, 10) )
cut(y, y.breaks, include.lowest = TRUE, labels = 1:10)
```

concordance

*Concordance test for binomial models***Description**

Performs a concordance/disconcordance (C-statistic) test on binomial models.

Usage

```
concordance(y, p)
```

Arguments

y	vector of binomial response variable used in model
p	estimated probabilities from fit binomial model

Value

A list class object with the following components:

- concordance percent of positives that are greater than probabilities of nulls
- is concordance inverse of concordance representing null class
- tied number of tied probabilities
- pairs number of pairs compared

Note

Test of binomial regression for the hypothesis that probabilities of all positives [1], are greater than the probabilities of the nulls [0]. The concordance would be 100 for a perfect model where, disconcordance is the inverse of concordance, representing the null. The C-statistic has been shown to be comparable to the area under an ROC

Author(s)

Jeffrey S. Evans <jeffrey_evans@tnc.org>

References

Austin, P.C. & E.W. Steyerberg (2012) Interpreting the concordance statistic of a logistic regression model: relation to the variance and odds ratio of a continuous explanatory variable. *BMC Medical Research Methodology*, 12:82 Harrell, F.E. (2001) Regression modelling strategies. Springer, New York, NY. Royston, P. & D.G. Altman (2010) Visualizing and assessing discrimination in the logistic regression model. *Statistics in Medicine* 29(24):2508-2520

Examples

```
data(mtcars)
dat <- subset(mtcars, select=c(mpg, am, vs))
glm.reg <- glm(vs ~ mpg, data = dat, family = binomial)
concordance(dat$vs, predict(glm.reg, type = "response"))
```

conf.interval	<i>Confidence interval for mean or median</i>
---------------	---

Description

Calculates confidence interval for the mean or median of a distribution with unknown population variance

Usage

```
conf.interval(x, cl = 0.95, stat = "mean", std.error = TRUE)
```

Arguments

x	Vector to calculate confidence interval for
cl	Percent confidence level (default = 0.95)
stat	Statistic (mean or median)
std.error	Return standard error (TRUE/FALSE)

Value

lci	Lower confidence interval value
uci	Upper confidence interval value
mean	If stat = "mean", mean value of distribution
mean	Value of the mean or median
conf.level	Confidence level used for confidence interval
std.error	If std.error = TRUE standard error of distribution

Author(s)

Jeffrey S. Evans <jeffrey_evans@tnc.org>

Examples

```

x <- runif(100)
cr <- conf.interval(x, cl = 0.97)
print(cr)

d <- density(x)
plot(d, type="n", main = "PDF with mean and 0.97 confidence interval")
polygon(d, col="cyan3")
abline(v=mean(x, na.rm = TRUE), lty = 2)
segments( x0=cr[["lci"]], y0=mean(d$y), x1=cr[["uci"]],
          y1=mean(d$y), lwd = 2.5,
          col = "black")
legend("topright", legend = c("mean", "CI"),
       lty = c(2,1), lwd = c(1,2.5))

```

correlogram

Correlogram

Description

Calculates and plots a correlogram

Usage

```
correlogram(x, v, dist = 5000, dmatrix = FALSE, ns = 99,
           latlong = FALSE, ...)
```

Arguments

x	SpatialPointsDataFrame object
v	Test variable in x@data
dist	Distance of correlation lags, if latlong=TRUE units are in kilometers
dmatrix	Should the distance matrix be include in output (TRUE/FALSE)
ns	Number of simulations to derive simulation envelope
latlong	Coordinates are in latlong (TRUE/FALSE)
...	Arguments passed to cor ('pearson', 'kendall' or 'spearman')

Value

A list object containing:

autocorrelation is a data.frame object with the following components

autocorrelation - Autocorrelation value for each distance lag

dist - Value of distance lag

lci - Lower confidence interval (p=0.025)

uci - Upper confidence interval (p=0.975)
 CorrPlot recordedplot object to recall plot
 dmatrix Distance matrix (if dmatrix=TRUE)

Note

depends: sp

Author(s)

Jeffrey S. Evans <jeffrey_evans@tnc.org>

Examples

```
library(sp)
data(meuse)
coordinates(meuse) = ~x+y
zinc.cg <- correlogram(x = meuse, v = meuse@data[, 'zinc'], dist = 250, ns = 9)
```

crossCorrelation *Spatial cross correlation*

Description

Calculates univariate or bivariate spatial cross-correlation using local Moran's-I (LISA), following Chen (2015)

Usage

```
crossCorrelation(x, y = NULL, coords = NULL, w = NULL,
  type = c("LSCI", "GSCI"), k = 1000, dist.function = "inv.power",
  scale.xy = TRUE, scale.partial = FALSE, scale.matrix = FALSE,
  alpha = 0.05, clust = FALSE, return.sims = FALSE)
```

Arguments

x	Vector of x response variables
y	Vector of y response variables, if not specified the univariate statistic is returned
coords	A matrix of coordinates corresponding to [x,y], only used if k = NULL. Can also be an sp object with relevant x,y coordinate slot (ie., points or polygons)
w	Spatial neighbors/weights in matrix format. Dimensions must match [n(x),n(y)] and be symmetrical. If w is not defined then a default method is used.
type	c("LSCI","GSCI") Return Local Spatial Cross-correlation Index (LSCI) or Global Spatial cross-correlation Index (GSCI)

k	Number of simulations for calculating permutation distribution under the null hypothesis of no spatial autocorrelation (k=1000, default)
dist.function	("inv.power", "neg.exponent") If w = NULL, the default method for deriving spatial weights matrix, options are: inverse power or negative exponent
scale.xy	(TRUE/FALSE) scale the x,y vectors, if FALSE it is assumed that they are already scaled following Chen (2015)
scale.partial	(FALSE/TRUE) rescale partial spatial autocorrelation statistics [-1 - 1]
scale.matrix	(FALSE/TRUE) If a neighbor/distance matrix is passed, should it be scaled using [w/sum(w)]?
alpha	= 0.05 confidence interval (default is 95 pct)
clust	(FALSE/TRUE) Return approximated lisa clusters
return.sims	(FALSE/TRUE) Return randomizations vector n = k

Value

When not simulated k=0, a list containing:

- I Global autocorrelation statistic
- SCI A data.frame with two columns representing the xy and yx autocorrelation
- nsim value of NULL to represent p values were derived from observed data (k=0)
- p Probability based observations above/below confidence interval
- t.test Probability based on t-test
- clusters If "clust" argument TRUE, vector representing LISA clusters

when simulated (k>0), a list containing:

- I Global autocorrelation statistic
- SCI A data.frame with two columns representing the xy and yx autocorrelation
- nsim value representing number of simulations
- global.p p-value of global autocorrelation statistic
- local.p Probability based simulated data using successful rejection of t-test
- range.p Probability based on range of probabilities resulting from paired t-test
- clusters If "clust" argument TRUE, vector representing lisa clusters

References

Chen., Y. (2015) A New Methodology of Spatial Cross-Correlation Analysis. PLoS One 10(5):e0126158. doi:10.1371/journal.pone.0126158

Examples

```

library(sp)
library(spdep)

data(meuse)
coordinates(meuse) <- ~x+y

##### Providing a neighbor contiguity spatial weights matrix
all.linked <- max(unlist(nbdist(knn2nb(knearneigh(coordinates(meuse)))),
                           coordinates(meuse))))
nb <- nb2listw(dnearneigh(meuse, 0, all.linked), style = "B", zero.policy = TRUE)
Wij <- as.matrix(as(nb, "symmetricMatrix"))
(I <- crossCorrelation(meuse$zinc, meuse$copper, w = Wij,
                       clust=TRUE, k=99))
meuse$lisa <- I$SCI[, "lsci.xy"]
meuse$lisa.clust <- as.factor(I$cluster)
spplot(meuse, "lisa")
spplot(meuse, "lisa.clust")

##### Using a default spatial weights matrix method (inverse power function)
(I <- crossCorrelation(meuse$zinc, meuse$copper, coords = coordinates(meuse),
                       clust = TRUE, k=99))
meuse$lisa <- I$SCI[, "lsci.xy"]
meuse$lisa.clust <- as.factor(I$cluster)
spplot(meuse, "lisa")
spplot(meuse, "lisa.clust")

## Not run:
##### Simulate spatially autocorrelated random normal variables
##### using eigen-decomposition, requires ncf package
library(sp)
library(ncf)
x=expand.grid(1:20, 1:20)[,1]
y=expand.grid(1:20, 1:20)[,2]
sdat <- data.frame(x = x,y=y,
                    z1=ncf::rmvn.spa(x=x, y=y, p=2, method="exp"),
                    z2=ncf::rmvn.spa(x=x, y=y, p=2, method="exp"))
coordinates(sdat) <- ~x+y
(I <- crossCorrelation(sdat$z1, sdat$z2, coords=coordinates(sdat),
                       k=9999, clust = TRUE))
sdat$lisa <- I$SCI[, "lsci.xy"]
sdat$lisa.clust <- as.factor(I$cluster)
spplot(sdat, "lisa")
spplot(sdat, "lisa.clust")

##### 1st order polygon contingency example
##### requires UScensus2000tract package
library(sp)
library(spdep)
library(UScensus2000tract)

```

```

data(oregon.tract)
nb <- nb2listw(poly2nb(oregon.tract), style = "B", zero.policy = TRUE
  Wij <- as.matrix( as(nb, "symmetricMatrix") )

X = oregon.tract$white
Y = oregon.tract$black

# Simulated bivariate lisa
I <- crossCorrelation(X, Y, w=Wij, k=999)
oregon.tract$lisa <- I$SCI[, "lsci.xy"]
oregon.tract$lisa.clust <- as.factor(I$cluster)
  spplot(oregon.tract, "lisa")
  spplot(oregon.tract, "lisa.clust")

## End(Not run)

```

csi *Cosine Similarity Index*

Description

Calculates the cosine similarity and angular similarity on two vectors or a matrix

Usage

```
csi(x, y = NULL)
```

Arguments

- | | |
|---|--|
| x | A vector or matrix object |
| y | If x is a vector, then a vector object |

Value

If x is a matrix, a list object with: similarity and angular.similarity matrices

If x and y are vectors, a vector of similarity and angular.similarity

Note

The cosine similarity index is a measure of similarity between two vectors of an inner product space. This index is best suited for high-dimensional positive variable space. One useful application of the index is to measure separability of clusters derived from algorithmic approaches (e.g., k-means). It is a good common practice to center the data before calculating the index. It should be noted that the cosine similarity index is mathematically, and often numerically, equivalent to the Pearson's correlation coefficient

cosine similarity index is derived:

$$s(xy) = x * y / \|x\| * \|y\|$$

expected 1.0 (perfect similarity) to -1.0 (perfect dissimilarity)

A normalized angle between the vectors can be used as a bounded similarity function within [0,1]
 $\text{angular similarity} = 1 - (\cos(s)^{-1}/\pi)$

Author(s)

Jeffrey S. Evans <jeffrey_evans@tnc.org>

Examples

```
# Compare two vectors (centered using scale)
x=runif(100)
y=runif(100)^2
csi(as.vector(scale(x)),as.vector(scale(y)))

#' # Compare columns (vectors) in a matrix (centered using scale)
x <- matrix(round(runif(100),0),nrow=20,ncol=5)
( s <- csi(scale(x)) )

# Compare vector (x) to each column in a matrix (y)
y <- matrix(round(runif(500),3),nrow=100,ncol=5)
x=runif(100)
csi(as.vector(scale(x)),scale(y))
```

curvature

Surface curvature

Description

Calculates Zevenbergen & Thorne, McNab's or Bolstad's curvature

Usage

```
curvature(x, s = 3, type = c("planform", "profile", "total", "mcnab",
"bolstad"), ...)
```

Arguments

x	rasterLayer object
s	Focal window size
type	Method used c("planform", "profile", "total", "mcnab", "bolstad")
...	Additional arguments passed to writeRaster

Value

raster class object of surface curvature

Note

The planform and profile curvatures are the second derivative(s) of the elevation surface, or the slope of the slope. Profile curvature is in the direction of the maximum slope, and the planform curvature is perpendicular to the direction of the maximum slope. Negative values in the profile curvature indicate the surface is upwardly convex whereas, positive values indicate that the surface is upwardly concave. Positive values in the planform curvature indicate that the surface is laterally convex whereas, negative values indicate that the surface is laterally concave. Total curvature is the sigma of the profile and planform curvatures. A value of 0 in profile, planform or total curvature, indicates the surface is flat. The planform, profile and total curvatures are derived using Zevenbergen & Thorne (1987) via a quadratic equation fit to eight neighbors as such, the s (focal window size) argument is ignored.

McNab's and Bolstad's variants of the surface curvature (concavity/convexity) index (McNab 1993; Bolstad & Lillesand 1992; McNab 1989). The index is based on features that confine the view from the center of a 3x3 window. In the Bolstad equation, edge correction is addressed by dividing by the radius distance to the outermost cell (36.2m).

Author(s)

Jeffrey S. Evans <jeffrey_evans@tnc.org>

References

Bolstad, P.V., and T.M. Lillesand (1992). Improved classification of forest vegetation in northern Wisconsin through a rule-based combination of soils, terrain, and Landsat TM data. *Forest Science*. 38(1):5-20. Florinsky, I.V. (1998). Accuracy of Local Topographic Variables Derived from Digital Elevation Models. *International Journal of Geographical Information Science*, 12(1):47-62. McNab, H.W. (1989). Terrain shape index: quantifying effect of minor landforms on tree height. *Forest Science*. 35(1):91-104. McNab, H.W. (1993). A topographic index to quantify the effect of mesoscale landform on site productivity. *Canadian Journal of Forest Research*. 23:1100-1107. Zevenbergen, L.W. & C.R. Thorne (1987). Quantitative Analysis of Land Surface Topography. *Earth Surface Processes and Landforms*, 12:47-56.

See Also

[writeRaster](#) For additional ... arguments passed to writeRaster

Examples

```
## Not run:
library(raster)
library(spatialEco)
data(elev)
elev <- projectRaster(elev, crs="+proj=robin +datum=WGS84",
                      res=1000, method='bilinear')

mcnab.crv <- curvature(elev, type="mcnab")
plot(mcnab.crv, main="McNab's curvature")

## End(Not run)
```

daymet.point *DAYMET point values*

Description

Downloads DAYMET climate variables for specified point and time-period

Usage

```
daymet.point(lat, long, start.year, end.year, site = NULL,  
            files = FALSE, echo = FALSE)
```

Arguments

lat	latitude of point (decimal degrees WGS84)
long	longitude pf point (decimal degrees WGS84)
start.year	First year of data
end.year	Last year of data
site	Unique identification value that is appended to data
files	(TRUE/FALSE) Write file to disk
echo	(TRUE/FALSE) Echo progress

Value

A data.frame with climate results

Note

Function uses the Single Pixel Extraction tool and returns year, yday, dayl(s), prcp (mm/day), srad (W/m^2), swe (kg/m^2), tmax (deg c), tmin (deg c), vp (Pa)

Metadata for DAYMET single pixel extraction: https://daymet.ornl.gov/files/UserGuides/current/readme_singlepointextraction.pdf

data is available for Long -131.0 W and -53.0 W; lat 52.0 N and 14.5 N

Author(s)

Jeffrey S. Evans <jeffrey_evans@tnc.org>

Examples

```
## Not run:  
( d <- daymet.point(lat = 36.0133, long = -84.2625, start.year = 2013, end.year=2014,  
#           site = "1", files = FALSE, echo = FALSE) )  
  
## End(Not run)
```

daymet.tiles*DAYMET Tile ID's***Description**

Returns a vector of DAYMET tile id's within a specified extent

Usage

```
daymet.tiles(x, tiles, ids, coords, sp = FALSE)
```

Arguments

<code>x</code>	A sp, raster or extent object (with same projection as tiles)
<code>tiles</code>	A SpatialPolygonsDataFrame tile index (see notes)
<code>ids</code>	A tile id field in the tiles index
<code>coords</code>	A vector of xmin, xmax, ymin, ymax coordinates, in same projection as tiles
<code>sp</code>	(TRUE/FALSE) Should an sp class SpatialPolygonsDataFrame object of associate tiles be returned

Value

Vector of DAYMET tile IDS or if sp = TRUE a sp class SpatialPolygonsDataFrame

Note

Function accepts sp, raster or extent class object or bounding coordinates. All input must be in the same projection as the tile index SpatialPolygonsDataFrame. The library includes the DAYMAT tile index "DAYMET_tiles" which can be add using data(), see examples.

Author(s)

Jeffrey S. Evans <jeffrey_evans@tnc.org>

Examples

```
library(sp)
library(raster)
data(DAYMET_tiles)
e <- extent(-117.2567, -104.7523, 36.62797, 47.68194)
plot(DAYMET_tiles)
  plot(e, col="red", add=TRUE)

# Using extent object
daymet.tiles(x = e, tiles = DAYMET_tiles, ids = "Id")

# Using sp object
e <- as(e, "SpatialPolygons")
```

```
daymet.tiles(e, tiles = DAYMET_tiles, ids = "Id")

# Using bounding coordinates
daymet.tiles(coords=c(-117.2567, -104.7523, 36.62797, 47.68194),
             tiles = DAYMET_tiles, ids = "Id" )

# Return sp polygons object
tiles <- daymet.tiles(x = e, tiles = DAYMET_tiles, ids = "Id", sp = TRUE)
plot(DAYMET_tiles)
  plot(tiles, col="red", add=TRUE)

## Not run:
# batch download of DAYMET tiles using function
tile.ids = daymet.tiles(e)
download.daymet(years=2010, tile=tile.ids, data.type=c('tmin'))

## End(Not run)
```

DAYMET_tiles**DAYMET climate tile index**

Description

Polygon tile index for DAYMET climate data

Format

An sp SpatialPolygonsDataFrame with 404 features (rows) and 6 columns (columns):

Id Tile Index Identification

Area Area of each tile

XMin Minimum x geographic decimal degree coordinate

XMax Maximum x geographic decimal degree coordinate

YMin Minimum y geographic decimal degree coordinate

yMax Maximum y geographic decimal degree coordinate

Source

<https://daymet.ornl.gov/>

dispersion	<i>Dispersion (H-prime)</i>
------------	-----------------------------

Description

Calculates the dispersion ("rarity") of targets associated with planning units

Usage

```
dispersion(x)
```

Arguments

x	data.frame object of target values
---	------------------------------------

Value

data.frame with columns H values for each target, H , sH, sHmax

Note

The dispersion index (H-prime) is calculated $H = \sum(\sqrt{p} / \sqrt{a})$ where; P = [sum of target in planning unit / sum of target across all planning units] and a = [count of planning units containing target / number of planning units]

Author(s)

Jeffrey S. Evans <jeffrey_evans@tnc.org>

References

Evans, J.S., S.R. Schill, G.T. Raber (2015) A Systematic Framework for Spatial Conservation Planning and Ecological Priority Design in St. Lucia, Eastern Caribbean. Chapter 26 in Central American Biodiversity : Conservation, Ecology and a Sustainable Future. F. Huettman (eds). Springer, NY.

Examples

```
library(sp)
data(pu)

d <- dispersion(pu@data[,2:ncol(pu)])

## Not run:
p <- d[, "H"]
clr <- c("#3288BD", "#99D594", "#E6F598", "#FEE08B",
         "#FC8D59", "#D53E4F")
clrs <- ifelse(p < 0.5524462, clr[1],
               ifelse(p >= 0.5524462 & p < 1.223523, clr[2],
```

```

    ifelse(p >= 1.223523 & p < 2.465613, clr[3],
    ifelse(p >= 2.465613 & p < 4.76429, clr[4],
        ifelse(p >= 4.76429 & p < 8.817699, clr[5],
            ifelse(p >= 8.817699, clr[6], NA))))))
plot(pu, col=clrs, border=NA)
legend("topleft", legend=rev(c("Very Rare", "Rare", "Moderately Rare",
    "Somewhat Common", "Common", "Over Dispersed")),
    fill=clr, cex=0.6, bty="n")
box()
## End(Not run)

```

dissection

*Dissection***Description**

Calculates the Evans (1972) Martonne's modified dissection

Usage

```
dissection(x, s = 5, ...)
```

Arguments

x	raster object
s	Focal window size
...	Additional arguments passed to raster::calc

Value

raster class object of Martonne's modified dissection

Note

Dissection is calculated as: $(z(s) - \min(z(s))) / (\max(z(s)) - \min(z(s)))$

Author(s)

Jeffrey S. Evans <jeffrey_evans@tnc.org>

Examples

```

library(raster)
data(elev)
d <- dissection(elev, s=5)
plot(d, main="dissection")

```

download.daymet *Download DAYMET*

Description

Batch download of daily gridded DAYMET climate data

Usage

```
download.daymet(years, tiles, data.type = "all",
                download.folder = getwd(),
                http = "https://thredds.daac.ornl.gov/thredds/fileServer/ornldaac/1328/tiles")
```

Arguments

years	Years to download (valid years 1980-2015)
tiles	Tile index value (see url for tile index grid in notes section)
data.type	Type of climate metric: 'all', 'vp', 'tmin', 'tmax', 'swe', 'srads', 'prcp', 'dayl'.
download.folder	local download directory, defaults to current working directory
http	option to change URL

Value

DAYMET netCDF format climate metrics

Note

Available products

vp Water Vapor Pressure Daily average partial pressure of water vapor

tmin Daily minimum (degrees C) 2-meter air temperature

tmax Daily maximum (degrees C) 2-meter air temperature

swe Snow water equivalent (kg/m²). Amount of water contained within snowpack.

srads Incident shortwave radiation flux density (W/m²), taken as average over daylight period of the day.

prcp Daily total precipitation(mm/day), sum of all forms converted to water-equivalent.

dayl Duration of the daylight period for the day (s/day). Calculation is based on the period of the day during which the sun is above a hypothetical flat horizon.

DAYMET main website: <http://daymet.ornl.gov>

Tile index information <http://daymet.ornl.gov/datasupport.html>

Data repository url: <https://thredds.daac.ornl.gov/thredds/fileServer/ornldaac/1328/tiles>

Path structure: /year/tile_year/file.nc

Author(s)

Jeffrey S. Evans <jeffrey_evans@tnc.org>

References

- Thornton P.E., S.W. Running and M.A. White (1997) Generating surfaces of daily meteorological variables over large regions of complex terrain. *Journal of Hydrology* 190: 214-251.
- Thornton, P.E. and S.W. Running (1999) An improved algorithm for estimating incident daily solar radiation from measurements of temperature, humidity, and precipitation. *Agriculture and Forest Meteorology*. 93:211-228.
- Thornton, P.E., H. Hasenauer and M.A. White (2000) Simultaneous estimation of daily solar radiation and humidity from observed temperature and precipitation: An application over complex terrain in Austria. *Agricultural and Forest Meteorology* 104:255-271.

Examples

```
## Not run:
# Download 2009-2010 min and max temp for tiles 11737 and 11738
laramie.plains <- c(11737, 11738)
my.years <- c(seq(2009,2010,1))
download.daymet(years=my.years, tiles=laramie.plains, data.type=c('tmin','tmax'))

## End(Not run)
```

download.hansen

*Download Hansen Forest 2000-2013 Change***Description**

Download of Hansen Global Forest Change 2000-2013

Usage

```
download.hansen(tile, data.type = c("loss"), download.folder = getwd())
```

Arguments

- | | |
|-----------------|--|
| tile | Granule index (See project URL for granule grid index) |
| data.type | Type of data to download options: 'treecover2000', 'loss', 'gain', 'lossyear', 'datamask', 'first', 'last' |
| download.folder | Destination folder |

Value

Downloaded Hansen forest loss tif files

Note

Project website with 10x10 degree granule index: http://earthenginepartners.appspot.com/science-2013-global-forest/download_v1.1.html

Available products: treecover2000, loss, gain, lossyear, datamask, first, or last

'treecover2000' (Tree canopy cover for year 2000) - Tree cover in the year 2000, defined as canopy closure for all vegetation taller than 5m in height. Encoded as a percentage per output grid cell, in the range 0-100.

'loss' (Global forest cover loss 2000-2013) - Forest loss during the period 2000-2013, defined as a stand-replacement disturbance, or a change from a forest to non-forest state. Encoded as either 1 (loss) or 0 (no loss).

'gain' (Global forest cover gain 2000-2012) - Forest gain during the period 2000-2012, defined as the inverse of loss, or a non-forest to forest change entirely within the study period. Encoded as either 1 (gain) or 0 (no gain).

'lossyear' (Year of gross forest cover loss event) - A disaggregation of total forest loss to annual time scales. Encoded as either 0 (no loss) or else a value in the range 1-13, representing loss detected primarily in the year 2001-2013.

'datamask' (Data mask) - Three values representing areas of no data (0), mapped land surface (1), and permanent water bodies (2).

'first' (Circa year 2000 Landsat 7 cloud-free image composite) - Reference multispectral imagery from the first available year, typically 2000. If no cloud-free observations were available for year 2000, imagery was taken from the closest year with cloud-free data, within the range 1999-2012.

'last' (Circa year 2013 Landsat cloud-free image composite) - Reference multispectral imagery from the last available year, typically 2013. If no cloud-free observations were available for year 2013, imagery was taken from the closest year with cloud-free data, within the range 2010-2012.

Author(s)

Jeffrey S. Evans <jeffrey_evans@tnc.org>

References

Hansen, M. C., P. V. Potapov, R. Moore, M. Hancher, S. A. Turubanova, A. Tyukavina, D. Thau, S. V. Stehman, S. J. Goetz, T. R. Loveland, A. Kommareddy, A. Egorov, L. Chini, C. O. Justice, and J. R. G. Townshend. (2013) High-Resolution Global Maps of 21st-Century Forest Cover Change. *Science* 342:850-53.

Examples

```
## Not run:
# Download single tile
download.hansen(tile=c('00N', '130E'), data.type=c('loss', 'lossyear'),
                download.folder=getwd())

# Batch download of multiple tiles
tiles <- list(c('00N', '140E'), c('00N', '130E'))
for( j in 1:length(tiles)){
  download.hansen(tile=tiles[[j]], data.type=c('loss'))}
```

```
}
```

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

download.prism *Download PRISM*

Description

Batch download of monthly gridded PRISM climate data

Usage

```
download.prism(data.type, date.range, time.step = "monthly",
  download.folder = getwd(), by.year = FALSE, unzip.file = TRUE,
  ftp.site = "ftp://prism.oregonstate.edu")
```

Arguments

data.type	Specify climate metric ('ppt','tmin','tmax','tmean')
date.range	A vector with start and end date in y/m/d format
time.step	Timestep of product ('daily'/'monthly')
download.folder	Local download directory, defaults to current working directory
by.year	Create a directory for each year (TRUE/FALSE)
unzip.file	Unzip file on download (TRUE/FALSE)
ftp.site	PRISM ftp address to use, default: ftp://prism.oregonstate.edu

Value

Compressed or uncompressed PRISM monthly gridded data(bil raster format)

Note

Monthly data 1895-1980 is available in a single zip file on the ftp site

PRISM URL: <http://prism.nacse.org/>

FTP download sites for 400m gridded daily/monthly climate data

<ftp://prism.oregonstate.edu/daily>

<ftp://prism.oregonstate.edu/monthly>

Naming convention: PRISM_<var>_<stability>_<scale&version>_<date>_bil.zip
i.e., 'PRISM_ppt_stable_4kmD1_20100208_bil.zip'

Data description:

http://prism.nacse.org/documents/PRISM_datasets_aug2013.pdf

Author(s)

Jeffrey S. Evans <jeffrey_evans@tnc.org>

See Also

[download.daymet](#), [download.hansen](#)

Examples

```
## Not run:
my.dates <- c('2000/1/1', '2001/12/30')
download.prism('ppt', date.range=my.dates, time.step='monthly', by.year=TRUE)

# Download monthly precipitation data Jan 1st 2000 to Feb 10th 2000 (n=41)
my.dates <- c('2000/1/1', '2000/2/10')
download.prism('ppt', date.range=my.dates, time.step='daily', by.year=TRUE)

## End(Not run)
```

effect.size

Cohen's-d effect size

Description

Cohen's-d effect size with pooled sd for a control and experimental group

Usage

```
effect.size(y, x, pooled = TRUE, conf.level = 0.95)
```

Arguments

y	A character or factor vector
x	A numeric vector, same length as y
pooled	Pooled or population standard deviation (TRUE/FALSE)
conf.level	Specified confidence interval. Default is 0.95

Value

An *effect.size* class object with x, y and a data.frame with columns for effect size, lower confidence interval, lower confidence interval. The row names of the data frame represent the levels in y

Note

This implementation will iterate through each class in y and treating a given class as the experimental group and all other classes as a control case. Each class had d and the confidence interval derived. A negative d indicate directionality with same magnitude. The expected range for d is 0 - 3
 d is derived; $(\text{mean(experimental group)} - \text{mean(control group)}) / \text{sigma(p)}$
 pooled standard deviation is derived; $\sqrt{((N_e - 1) * \text{sigma}(e)^2 + (N_c - 1) * \text{sigma}(c)^2) / (N_e + N_c - 2)}$ where; $N_e, N_c = n$ of experimental and control groups.

Author(s)

Jeffrey S. Evans <jeffrey_evans@tnc.org>

References

- Cohen, J., (1988) Statistical Power Analysis for the Behavioral Sciences (second ed.). Lawrence Erlbaum Associates.
 Cohen, J (1992) A power primer. Psychological Bulletin 112(1):155-159

Examples

```
( es <- effect.size(iris$Species, iris$Sepal.Length) )
plot(es)
```

elev

Elevation raster

Description

elevation raster of Switzerland

Format

A raster RasterLayer class object:

```
resoultion 5 arc-minute 0.00833 (10000m)
nrow 264
ncol 564
ncell 148896
xmin 5.9
xmax 10.6
ymin 45.7
ymax 47.9
proj4string +proj=longlat +ellps=WGS84
```

Source

<http://www.diva-gis.org/Data>

`erase.point`

Erase points

Description

Removes points intersecting a polygon feature class

Usage

```
erase.point(y, x, inside = TRUE)
```

Arguments

<code>y</code>	A SpatialPoints or SpatialPointsDataFrame
<code>x</code>	A SpatialPolygons or SpatialPolygonsDataFrame
<code>inside</code>	(TRUE/FALSE) Remove points inside polygon, else outside polygon

Value

A SpatialPoints or SpatialPointsDataFrame

Note

Provides the same functionality as the ESRI ArcGIS Erase Point tool

Author(s)

Jeffrey S. Evans <jeffrey_evans<at>tnc.org>

Examples

```
require(sp)
data(meuse)
coordinates(meuse) = ~x+y
poly <- SpatialPolygonsDataFrame(SpatialPolygons(list(Polygons(list(
  Polygon(cbind(c(180042, 180545, 180553, 180314, 179955,
  179142, 179437, 179524, 179979, 180042), c(332373, 332026,
  331426, 330889, 330683, 331133, 331623, 332152, 332357,
  332373)))), '1'))), data.frame(row.names=c('1'), PIDS=1))

meuse.erase <- erase.point(meuse, poly)

par(mfrow=c(1,2))
plot(poly,)
points(meuse, pch=20)
```

```
plot(poly)
  points(meuse.erase, pch=20)
```

focal.lmetrics *Focal landscape metrics*

Description

Calculates a variety of landscape metrics on integer rasters using focal approach

Usage

```
focal.lmetrics(x, w = 5, bkg = 0, land.value = 1,
  metric = "prop.landscape", latlong = FALSE)
```

Arguments

x	raster class object
w	Size of focal window, a single value or two values defining size of matrix (must be odd values)
bkg	Background value (will be ignored)
land.value	Raster cell value to calculate metrics on
metric	Name of desired metric (see available metrics)
latlong	(FALSE/TRUE) Is raster data in lat-long

Details

The following metrics are available:

- class - a particular patch type from the original input matrix (mat).
- n.patches - the number of patches of a particular patch type or in a class.
- total.area - the sum of the areas (m²) of all patches of the corresponding patch type.
- prop.landscape - the proportion of the total landscape represented by this class
- patch.density - the numbers of patches of the corresponding patch type divided by total landscape area (m²).
- total.edge - the total edge length of a particular patch type.
- edge.density - edge length on a per unit area basis that facilitates comparison among landscapes of varying size.
- landscape.shape.index - a standardized measure of total edge or edge density that adjusts for the size of the landscape.
- largest.patch.index - largest patch index quantifies the percentage of total landscape area comprised by the largest patch.
- mean.patch.area - average area of patches.

- `sd.patch.area` - standard deviation of patch areas.
- `min.patch.area` - the minimum patch area of the total patch areas.
- `max.patch.area` - the maximum patch area of the total patch areas.
- `perimeter.area.frac.dim` - perimeter-area fractal dimension equals 2 divided by the slope of regression line obtained by regressing the logarithm of patch area (m^2) against the logarithm of patch perimeter (m).
- `mean.perim.area.ratio` - the mean of the ratio patch perimeter. The perimeter-area ratio is equal to the ratio of the patch perimeter (m) to area (m^2).
- `sd.perim.area.ratio` - standard deviation of the ratio patch perimeter.
- `min.perim.area.ratio` - minimum perimeter area ratio
- `max.perim.area.ratio` - maximum perimeter area ratio.
- `mean.shape.index` - mean of shape index
- `sd.shape.index` - standard deviation of shape index.
- `min.shape.index` - the minimum shape index.
- `max.shape.index` - the maximum shape index.
- `mean.frac.dim.index` - mean of fractal dimension index.
- `sd.frac.dim.index` - standard deviation of fractal dimension index.
- `min.frac.dim.index` - the minimum fractal dimension index.
- `max.frac.dim.index` - the maximum fractal dimension index.
- `total.core.area` - the sum of the core areas of the patches (m^2).
- `prop.landscape.core` - proportional landscape core
- `mean.patch.core.area` - mean patch core area.
- `sd.patch.core.area` - standard deviation of patch core area.
- `min.patch.core.area` - the minimum patch core area.
- `max.patch.core.area` - the maximum patch core area.
- `prop.like.adjacencies` - calculated from the adjacency matrix, which shows the frequency with which different pairs of patch types (including like adjacencies between the same patch type) appear side-by-side on the map (measures the degree of aggregation of patch types).
- `aggregation.index` - computed simply as an area-weighted mean class aggregation index, where each class is weighted by its proportional area in the landscape.
- `landscape.division.index` - based on the cumulative patch area distribution and is interpreted as the probability that two randomly chosen pixels in the landscape are not situated in the same patch
- `splitting.index` - based on the cumulative patch area distribution and is interpreted as the effective mesh number, or number of patches with a constant patch size when the landscape is subdivided into S patches, where S is the value of the splitting index.
- `effective.mesh.size` - equals 1 divided by the total landscape area (m^2) multiplied by the sum of patch area (m^2) squared, summed across all patches in the landscape.
- `patch.cohesion.index` - measures the physical connectedness of the corresponding patch type.

Value

RasterLayer class object of specified landscape metric

Author(s)

Jeffrey S. Evans <jeffrey_evans@tnc.org>

See Also

[land.metrics](#)

[ConnCompLabel](#)

[PatchStat](#)

[ClassStat](#)

Examples

```
## Not run:
library(raster)
library(sp)

r <- raster(nrows=180, ncols=360, xmn=571823.6, xmx=616763.6, ymn=4423540,
            ymx=4453690, resolution=270, crs = CRS("+proj=utm +zone=12 +datum=NAD83
            +units=m +no_defs +ellps=GRS80 +towgs84=0,0,0"))

r[] <- rpois(ncell(r), lambda=1)
r <- calc(r, fun=function(x) { x[x >= 1] <- 1; return(x) } )

# proportion landscape class
pland <- focal.lmetrics(r, w=11)
plot(pland)

# Aggregation index
ai <- focal.lmetrics(r, w=11, metric = "aggregation.index")
plot(ai)

## End(Not run)
```

Description

Calculates the fuzzy sum of a vector

Usage

`fuzzySum(x)`

Arguments

x Vector of values to apply fuzzy sum

Value

Value of fuzzy sum

Note

The fuzzy sum is an increasing linear combination of values. This can be used to sum probabilities or results of multiple density functions.

Author(s)

Jeffrey S. Evans <jeffrey_evans@tnc.org>

Examples

```
p = c(0.8,0.76,0.87)
fuzzySum(p)
sum(p)
```

```
p = c(0.3,0.2,0.1)
fuzzySum(p)
sum(p)
```

gaussian.kernel *Gaussian Kernel*

Description

Creates a Gaussian Kernel of specified size and sigma

Usage

```
gaussian.kernel(sigma = 2, n = 5)
```

Arguments

sigma	sigma (standard deviation) of kernel (defaults 2)
n	size of symmetrical kernel (defaults to 5x5)

Value

Symmetrical (NxN) matrix of a Gaussian distribution

Author(s)

Jeffrey S. Evans <jeffrey_evans@tnc.org>

Examples

```
par(mfrow=c(2,2))
persp(gaussian.kernel(sigma=1, n=27), theta = 135, phi = 30, col = "grey",
       ltheta = -120, shade = 0.6, border=NA )
persp(gaussian.kernel(sigma=2, n=27), theta = 135, phi = 30, col = "grey",
       ltheta = -120, shade = 0.6, border=NA )
persp(gaussian.kernel(sigma=3, n=27), theta = 135, phi = 30, col = "grey",
       ltheta = -120, shade = 0.6, border=NA )
persp(gaussian.kernel(sigma=4, n=27), theta = 135, phi = 30, col = "grey",
       ltheta = -120, shade = 0.6, border=NA )
```

group.pdf

Probability density plot by group

Description

Creates a probability density plot of y for each group of x

Usage

```
group.pdf(x, y, col = NULL, lty = NULL, lwd = NULL, lx = "topleft",
          ly = NULL, ...)
```

Arguments

x	Numeric, character or factorial vector of grouping variable (must be same length as y)
y	Numeric vector (density variable)
col	Optional line colors (see par, col)
lty	Optional line types (see par, lty)
lwd	Optional line widths (see par, lwd)
lx	Position of legend (x coordinate or 'topright', 'topleft', 'bottomright', 'bottomleft')
ly	Position of legend (y coordinate)
...	Additional arguments passed to plot

Author(s)

Jeffrey S. Evans <jeffrey_evans<at>tnc.org>

References

Simonoff, J. S. (1996). Smoothing Methods in Statistics. Springer-Verlag, New York.

Examples

```
y=dnorm(runif(100))
x=rep(c(1,2,3), length.out=length(y))
group.pdf(x=as.factor(x), y=y, main='Probability Density of y by group(x)',
ylab='PDF', xlab='Y', lty=c(1,2,3))
```

hexagons

Hexagons

Description

Create hexagon polygons

Usage

```
hexagons(x, res = 100, ...)
```

Arguments

x	sp SpatialDataFrame class object
res	Area of resulting hexagons
...	Additional arguments passed to spsample

Value

SpatialPolygonsDataFrame OBJECT

Note

depends: sp

Examples

```
require(sp)
data(meuse)
coordinates(meuse) <- ~x+y

hex.polys <- hexagons(meuse, res=100)
plot(hex.polys)
  plot(meuse,pch=20,add=TRUE)

# Points intersecting hexagons
hex.pts <- na.omit(over(meuse,hex.polys))
(hex.pts <- data.frame(PTID=rownames(hex.pts), hex.pts))
```

hli*Heat Load Index*

Description

Calculates the McCune & Keon (2002) Heat Load Index

Usage

```
hli(x, check = TRUE)
```

Arguments

x	raster object
check	(TRUE/FALSE) check for projection integrity in northern latitudes

Value

raster class object of McCune & Keon (2002) Heat Load Index

Note

Describes A southwest facing slope should have warmer temperatures than a southeast facing slope, even though the amount of solar radiation they receive is equivalent. The McCune and Keon (2002) method accounts for this by "folding" the aspect so that the highest values are southwest and the lowest values are northeast. Additionally, this method account for steepness of slope, which is not addressed in most other aspect rescaling equations. HLI values range from 0 (coolest) to 1 (hottest).

Author(s)

Jeffrey S. Evans <jeffrey_evans@tnc.org>

References

McCune, B., and D. Keon (2002) Equations for potential annual direct incident radiation and heat load index. Journal of Vegetation Science. 13:603-606.

Examples

```
library(raster)
data(elev)
heat.load <- hli(elev)
plot(heat.load, main="Heat Load Index")
```

hsp*Hierarchical Slope Position***Description**

Calculates a hierarchical scale decomposition of topographic position index

Usage

```
hsp(x, min.scale = 3, max.scale = 27, inc = 4, win = "rectangle",
normalize = FALSE)
```

Arguments

<code>x</code>	Object of class raster (requires integer raster)
<code>min.scale</code>	Minimum scale (window size)
<code>max.scale</code>	Maximum scale (window size)
<code>inc</code>	Increment to increase scales
<code>win</code>	Window type, options are "rectangle" or "circle"
<code>normalize</code>	Normalize results to 0-1 scale (FALSE TRUE)

Value

raster class object

Note

if `win = "circle"` units are distance, if `win = "rectangle"` units are number of cells

Author(s)

Jeffrey S. Evans <jeffrey_evans@tnc.org>

References

Murphy M.A., J.S. Evans, and A.S. Storfer (2010) Quantify *Bufo boreas* connectivity in Yellowstone National Park with landscape genetics. *Ecology* 91:252-261

Examples

```
## Not run:
library(raster)
data(elev)
hsp27 <- hsp(elev, 3, 27, 4, normalize = TRUE)
plot(hsp27)

## End(Not run)
```

hybrid.kmeans*Hybrid K-means*

Description

Hybrid K-means clustering using hierarchical clustering to define cluster-centers

Usage

```
hybrid.kmeans(x, k = 2, hmethod = "ward.D", stat = mean, ...)
```

Arguments

x	A data.frame or matrix with data to be clustered
k	Number of clusters
hmethod	The agglomeration method used in hclust
stat	The statistic to aggregate class centers (mean or median)
...	Additional arguments passed to kmeans

Details

This method uses hierarchical clustering to define the cluster-centers in the K-means clustering algorithm. This mitigates some of the known convergence issues in K-means.

Value

returns an object of class "kmeans" which has a print and a fitted method

Note

options for hmethod are: "ward.D", "ward.D2", "single", "complete", "average", "mcquitty", "median", "centroid"

Author(s)

Jeffrey S. Evans <jeffrey_evans@tnc.org>

References

Singh, H., & K. Kaur (2013) New Method for Finding Initial Cluster Centroids in K-means Algorithm. International Journal of Computer Application. 74(6):27-30 Ward, J.H., (1963) Hierarchical grouping to optimize an objective function. Journal of the American Statistical Association. 58:236-24

See Also

[kmeans](#) for available ... arguments and function details
[hclust](#) for details on hierarchical clustering

Examples

```
x <- rbind(matrix(rnorm(100, sd = 0.3), ncol = 2),
             matrix(rnorm(100, mean = 1, sd = 0.3), ncol = 2))

# Compare k-means to hybrid k-means with k=4
km <- kmeans(x, 4)
hkm <- hybrid.kmeans(x,k=4)

par(mfrow=c(1,2))
plot(x[,1],x[,2], col=km$cluster,pch=19, main="K-means")
plot(x[,1],x[,2], col=hkm$cluster,pch=19, main="Hybrid K-means")
```

idw.smoothing*idw.smoothing***Description**

Distance weighted smoothing of a variable in a spatial point object

Usage

```
idw.smoothing(x, y, d, k)
```

Arguments

x	Object of class SpatialPointsDataFrame
y	Numeric data in x@data
d	Distance constraint
k	Maximum number of k-nearest neighbours within d

Value

A vector, same length as nrow(x), of adjusted y values

Note

Smoothing is conducted with a weighted-mean where; weights represent inverse standardized distance lags Distance-based or neighbour-based smoothing can be specified by setting the desired neighbour smoothing method to a specified value then the other parameter to the potential maximum. For example; a constraint distance, including all neighbours within 1000 (d=1000) would require k to equal all of the potential neighbours (n-1 or k=nrow(x)-1).

Depends: sp, RANN

Examples

```
library(sp)
data(meuse)
coordinates(meuse) <- ~x+y

# Calculate distance weighted mean on cadmium variable in meuse data
cadmium.idw <- idw.smoothing(meuse, 'cadmium', k=nrow(meuse), d = 1000)
meuse@data$cadmium.wm <- cadmium.idw
opar <- par
par(mfrow=c(2,1))
plot(density(meuse@data$cadmium), main='Cadmium')
plot(density(meuse@data$cadmium.wm), main='IDW Cadmium')
par <- opar
```

insert.values

Insert Values

Description

Inserts new values into a vector at specified positions

Usage

```
insert.values(x, value, index)
```

Arguments

x	A vector to insert values
value	Values to insert into x
index	Index position(s) to insert y values into x

Value

A vector with values of y inserted into x and the position(s) defined by the index

Note

This function inserts new values at specified positions in a vector. It does not replace existing values. If a single value is provided for y and l represents multiple positions y will be replicated for the length of l. In this way you can insert the same value at multiple locations.

Author(s)

Jeffrey S. Evans <jeffrey_evans@tnc.org>

Examples

```
(x=1:10)

# Insert single value in one location
insert.values(x, 100, 2)

# Insert multiple values in multiple locations
insert.values(x, c(100,200), c(2,8))

# Insert single value in multiple locations
insert.values(x, NA, c(2,8))
```

kde.2D

2-dimensional kernel density estimate

Description

Calculates 2-dimensional kernel density estimate over specified extent

Usage

```
kde.2D(...)
```

Arguments

...	Parameters to be passed to the modern version of the function
-----	---

kl.divergence

Kullback-Leibler divergence (relative entropy)

Description

Calculates the Kullback-Leibler divergence (relative entropy) between unweighted theoretical component distributions. Divergence is calculated as: $\int [f(x) (\log f(x) - \log g(x))] dx$ for distributions with densities $f()$ and $g()$.

Usage

```
kl.divergence(object, eps = 10^-4, overlap = TRUE)
```

Arguments

object	Matrix or dataframe object with ≥ 2 columns
eps	Probabilities below this threshold are replaced by this threshold for numerical stability.
overlap	Logical, do not determine the KL divergence for those pairs where for each point at least one of the densities has a value smaller than eps .

Value

pairwise Kullback-Leibler divergence index (matrix)

Author(s)

Jeffrey S. Evans <jeffrey_evans@tnc.org>

References

Kullback S., and R. A. Leibler (1951) On information and sufficiency. *The Annals of Mathematical Statistics* 22(1):79-86

Examples

```
x <- seq(-3, 3, length=200)
y <- cbind(n=dnorm(x), t=dt(x, df=10))
  matplot(x, y, type='l')
  k1.divergence(y)

# extract value for last column
k1.divergence(y[,1:2])[3:3]
```

land.metrics

Landscape metrics for points and polygons

Description

Calculates a variety of landscape metrics, on binary rasters, for polygons or points with a buffer distance

Usage

```
land.metrics(x, y, bkgd = NA, metrics = c("prop.landscape"),
  bw = 1000, latlon = FALSE, echo = TRUE)
```

Arguments

x	SpatialPointsDataFrame or SpatialPolygonsDataFrame class object
y	raster class object
bkgd	Background value (will be ignored)
metrics	Numeric index or name(s) of desired metric (see available metrics)
bw	Buffer distance (ignored if x is SpatialPolygonsDataFrame)
latlon	(FALSE/TRUE) Is raster data in lat-long
echo	(FALSE/TRUE) Print object ID at each iteration

Details

The following metrics are available:

- class - (always included) particular patch type from the original input matrix.
- n.patches - the number of patches of a particular patch type or in a class.
- total.area - the sum of the areas (m²) of all patches of the corresponding patch type.
- prop.landscape - the proportion of the total landscape represented by this class
- patch.density - the numbers of patches of the corresponding patch type divided by total landscape area (m²).
- total.edge - the total edge length of a particular patch type.
- edge.density - edge length on a per unit area basis that facilitates comparison among landscapes of varying size.
- landscape.shape.index - a standardized measure of total edge or edge density that adjusts for the size of the landscape.
- largest.patch.index - largest patch index quantifies the percentage of total landscape area comprised by the largest patch.
- mean.patch.area - average area of patches.
- sd.patch.area - standard deviation of patch areas.
- min.patch.area - the minimum patch area of the total patch areas.
- max.patch.area - the maximum patch area of the total patch areas.
- perimeter.area.frac.dim - perimeter-area fractal dimension equals 2 divided by the slope of regression line obtained by regressing the logarithm of patch area (m²) against the logarithm of patch perimeter (m).
- mean.perim.area.ratio - the mean of the ratio patch perimeter. The perimeter-area ratio is equal to the ratio of the patch perimeter (m) to area (m²).
- sd.perim.area.ratio - standard deviation of the ratio patch perimeter.
- min.perim.area.ratio - minimum perimeter area ratio
- max.perim.area.ratio - maximum perimeter area ratio.
- mean.shape.index - mean of shape index
- sd.shape.index - standard deviation of shape index.
- min.shape.index - the minimum shape index.
- max.shape.index - the maximum shape index.
- mean.frac.dim.index - mean of fractal dimension index.
- sd.frac.dim.index - standard deviation of fractal dimension index.
- min.frac.dim.index - the minimum fractal dimension index.
- max.frac.dim.index - the maximum fractal dimension index.
- total.core.area - the sum of the core areas of the patches (m²).
- prop.landscape.core - proportional landscape core
- mean.patch.core.area - mean patch core area.

- sd.patch.core.area - standard deviation of patch core area.
- min.patch.core.area - the minimum patch core area.
- max.patch.core.area - the maximum patch core area.
- prop.like.adjacencies - calculated from the adjacency matrix, which shows the frequency with which different pairs of patch types (including like adjacencies between the same patch type) appear side-by-side on the map (measures the degree of aggregation of patch types).
- aggregation.index - computed simply as an area-weighted mean class aggregation index, where each class is weighted by its proportional area in the landscape.
- landscape.division.index - based on the cumulative patch area distribution and is interpreted as the probability that two randomly chosen pixels in the landscape are not situated in the same patch
- splitting.index - based on the cumulative patch area distribution and is interpreted as the effective mesh number, or number of patches with a constant patch size when the landscape is subdivided into S patches, where S is the value of the splitting index.
- effective.mesh.size - equals 1 divided by the total landscape area (m^2) multiplied by the sum of patch area (m^2) squared, summed across all patches in the landscape.
- patch.cohesion.index - measures the physical connectedness of the corresponding patch type.

Value

If multiple classes are evaluated a list object with a data.frame for each class containing specified metrics in columns. The data.frame is ordered and shares the same row.names as the input feature class and can be directly joined to the @data slot. For single class problems a data.frame object is returned.

Note

Modifications to the function incorporate multi-class metrics by fetching the unique values of the raster and creating a list object containing a data.frame for each class. Unfortunately, retrieving unique values is a very slow function.

It is critical to note that if using polygon units that, the resulting metric represents the spatial pattern within each unique aggregate unit and not a metric of the unit itself. That is to say, a metric such as fractal dimension represents the fractal dimension of the pattern within the unit and not of the unit.

Author(s)

Jeffrey S. Evans <jeffrey_evans@tnc.org>

See Also

[focal.lmetrics](#)
[ConnCompLabel](#)
[PatchStat](#)
[ClassStat](#)

Examples

```

library(raster)
library(sp)

r <- raster(nrows=180, ncols=360, xmn=571823.6, xmx=616763.6, ymn=4423540,
            ymx=4453690, resolution=270, crs = CRS("+proj=utm +zone=12 +datum=NAD83
            +units=m +no_defs +ellps=GRS80 +towgs84=0,0,0"))

r[] <- rpois(ncell(r), lambda=1)
r <- calc(r, fun=function(x) { x[x >= 1] <- 1; return(x) } )
x <- sampleRandom(r, 10, na.rm = TRUE, sp = TRUE)

lmet <- c("prop.landscape", "edge.density", "prop.like.adjacencies", "aggregation.index")
( class.1 <- land.metrics(x=x, y=r, bw=1000, bkgd = 0, metrics = lmet) )
( all.class <- land.metrics(x=x, y=r, bw=1000, bkgd = NA, metrics = lmet ) )

# Pull metrics associated with class "0"
all.class[["0"]]

```

local.min.max

Local minimum and maximum

Description

Calculates the local minimums and maximums in a numeric vector, indicating inflection points in the distribution.

Usage

```
local.min.max(x, dev = mean, plot = TRUE, add.points = FALSE, ...)
```

Arguments

x	A numeric vector
dev	Deviation statistic (mean or median)
plot	plot the minimum and maximum values with the distribution (TRUE/FALSE)
add.points	Should all points of x be added to plot (TRUE/FALSE)
...	Arguments passed to plot

Value

A list object with:

minima minimum local values of x

maxima maximum local values of x

mindev Absolute deviation of minimum from specified deviation statistic (dev argument)

maxdev Absolute deviation of maximum from specified deviation statistic (dev argument)

Note

Useful function for identifying inflection or enveloping points in a distribution

Author(s)

Jeffrey S. Evans <jeffrey_evans@tnc.org>

Examples

```
x <- rnorm(100,mean=1500,sd=800)
( lmm <- local.min.max(x, dev=mean, add.points=TRUE, main="Local Minima and Maxima") )

# return only local minimum values
local.min.max(x)$minima
```

loess.boot

Loess Bootstrap

Description

Bootstrap of a Local Polynomial Regression (loess)

Usage

```
loess.boot(x, y, nreps = 100, confidence = 0.95, ...)
```

Arguments

x	Independent variable
y	Dependent variable
nreps	Number of bootstrap replicates
confidence	Fraction of replicates contained in confidence region
...	Additional arguments passed to loess function

Value

nreps Number of bootstrap replicates
confidence Confidence interval (region)
span alpha (span) parameter used loess fit
degree polynomial degree used in loess fit
normalize Normalized data (TRUE/FALSE)
family Family of statistic used in fit
parametric Parametric approximation (TRUE/FALSE)

surface Surface fit, see loess.control
 data data.frame of x,y used in model
 fit data.frame including: x Equally-spaced x index (see NOTES) y.fit loess fit up.lim Upper confidence interval low.lim Lower confidence interval stddev Standard deviation of loess fit at each x value

Note

The function fits a loess curve and then calculates a symmetric nonparametric bootstrap with a confidence region. Fitted curves are evaluated at a fixed number of equally-spaced x values, regardless of the number of x values in the data. Some replicates do not include the values at the lower and upper end of the range of x values. If the number of such replicates is too large, it becomes impossible to construct a confidence region that includes a fraction "confidence" of the bootstrap replicates. In such cases, the left and/or right portion of the confidence region is truncated.

Author(s)

Jeffrey S. Evans <jeffrey_evans@tnc.org>

References

- Cleveland, WS, (1979) Robust Locally Weighted Regression and Smoothing Plots Journal of the American Statistical Association 74:829-836
- Efron, B., and R. Tibshirani (1993) An Introduction to the Bootstrap Chapman and Hall, New York
- Härdle, W., (1989) Applied Nonparametric Regression Cambridge University Press, NY.
- Tibshirani, R. (1988) Variance stabilization and the bootstrap. Biometrika 75(3):433-44.

Examples

```
n=1000
x <- seq(0, 4, length.out=n)
y <- sin(2*x)+ 0.5*x + rnorm(n, sd=0.5)
sb <- loess.boot(x, y, nreps=99, confidence=0.90, span=0.40)
plot(sb)
```

Description

Calculates a local polynomial regression fit with associated confidence intervals

Usage

```
loess.ci(y, x, p = 0.95, plot = FALSE, ...)
```

Arguments

y	Dependent variable, vector
x	Independent variable, vector
p	Percent confidence intervals (default is 0.95)
plot	Plot the fit and confidence intervals
...	Arguments passed to loess

Value

A list object with:

- loess Predicted values
- se Estimated standard error for each predicted value
- lci Lower confidence interval
- uci Upper confidence interval
- df Estimated degrees of freedom
- rs Residual scale of residuals used in computing the standard errors

Author(s)

Jeffrey S. Evans <jeffrey_evans@tnc.org>

References

W. S. Cleveland, E. Grosse and W. M. Shyu (1992) Local regression models. Chapter 8 of Statistical Models in S eds J.M. Chambers and T.J. Hastie, Wadsworth & Brooks/Cole.

Examples

```

x <- seq(-20, 20, 0.1)
y <- sin(x)/x + rnorm(length(x), sd=0.03)
p <- which(y == "NaN")
y <- y[-p]
x <- x[-p]

par(mfrow=c(2,2))
lci <- loess.ci(y, x, plot=TRUE, span=0.10)
lci <- loess.ci(y, x, plot=TRUE, span=0.30)
lci <- loess.ci(y, x, plot=TRUE, span=0.50)
lci <- loess.ci(y, x, plot=TRUE, span=0.80)

```

`logistic.regression` *Logistic and Auto-logistic regression*

Description

Performs a logistic (binomial) or auto-logistic (spatially lagged binomial) regression using maximum likelihood or penalized maximum likelihood estimation.

Usage

```
logistic.regression(ldata, y, x, penalty = TRUE, autologistic = FALSE,
  coords = NULL, bw = NULL, type = "inverse", style = "W",
  longlat = FALSE, ...)
```

Arguments

<code>ldata</code>	data.frame object containing variables
<code>y</code>	Dependent variable (<code>y</code>) in <code>ldata</code>
<code>x</code>	Independent variable(s) (<code>x</code>) in <code>ldata</code>
<code>penalty</code>	Apply regression penalty (TRUE/FALSE)
<code>autologistic</code>	Add auto-logistic term (TRUE/FALSE)
<code>coords</code>	Geographic coordinates for auto-logistic model matrix or <code>sp</code> object.
<code>bw</code>	Distance bandwidth to calculate spatial lags (if empty neighbours result, need to increase bandwidth). If not provided it will be calculated automatically based on the minimum distance that includes at least one neighbor.
<code>type</code>	Neighbour weighting scheme (see <code>autocov_dist</code>)
<code>style</code>	Type of neighbour matrix (W_{ij}), default is mean of neighbours
<code>longlat</code>	Are coordinates (<code>coords</code>) in geographic, lat/long (TRUE/FALSE)
<code>...</code>	Additional arguments passed to <code>lrm</code>

Value

A list class object with the following components:

- model `lrm` model object (`rms` class)
- bandwidth If `AutoCov` = TRUE returns the distance bandwidth used for the auto-covariance function
- `diagTable` data.frame of regression diagnostics
- `coefTable` data.frame of regression coefficients
- `Residuals` data.frame of residuals and standardized residuals
- `AutoCov` If an auto-logistic model, `AutoCov` represents lagged auto-covariance term

Note

It should be noted that the auto-logistic model (Besag 1972) is intended for exploratory analysis of spatial effects. Auto-logistic are known to underestimate the effect of environmental variables and tend to be unreliable (Dormann 2007).

W_{ij} matrix options under style argument - B is the basic binary coding, W is row standardised (sums over all links to n), C is globally standardised (sums over all links to n), U is equal to C divided by the number of neighbours (sums over all links to unity) and S is variance-stabilizing.

Spatially lagged y defined as:

$$W(y)_{ij} = \sum_j (W_{ij} y_j) / \sum_j (W_{ij})$$

where; $W_{ij} = 1/\text{Euclidean}[i,j]$

If the object passed to the function is an sp class there is no need to call the data slot directly via "object@data", just pass the object name.

depends: rms, spdep

Author(s)

Jeffrey S. Evans <jeffrey_evans@tnc.org>

References

- Besag, J.E., (1972) Nearest-neighbour systems and the auto-logistic model for binary data. Journal of the Royal Statistical Society, Series B Methodological 34:75-83
- Dormann, C.F., (2007) Assessing the validity of autologistic regression. Ecological Modelling 207:234-242
- Le Cessie, S., Van Houwelingen, J.C., (1992) Ridge estimators in logistic regression. Applied Statistics 41:191-201
- Shao, J., (1993) Linear model selection by cross-validation. JASA 88:486-494

See Also

[lrm](#)
[autocov_dist](#)

Examples

```
require(sp)
require(spdep)
require(rms)
data(meuse)
coordinates(meuse) <- ~x+y
meuse@data <- data.frame(DepVar=rbinom(dim(meuse)[1], 1, 0.5), meuse@data)

#### Logistic model
lmodel <- logistic.regression(meuse, y='DepVar', x=c('dist','cadmium','copper'))
lmodel$model
lmodel$diagTable
lmodel$coefTable
```

```

##### Logistic model with factorial variable
lmodel <- logistic.regression(meuse, y='DepVar', x=c('dist','cadmium','copper', 'soil'))
lmodel$model
lmodel$diagTable
lmodel$coefTable

### Auto-logistic model using 'autocov_dist' in 'spdep' package
lmodel <- logistic.regression(meuse, y='DepVar', x=c('dist','cadmium','copper'),
                               autologistic=TRUE, coords=coordinates(meuse), bw=5000)
lmodel$model
lmodel$diagTable
lmodel$coefTable
est <- predict(lmodel$model, type='fitted.ind')

##### Add residuals, standardized residuals and estimated probabilities
VarNames <- rownames(lmodel$model$var)[-1]
meuse@data$AutoCov <- lmodel$AutoCov
meuse@data <- data.frame(meuse@data, Residual=lmodel$Residuals[,1],
                          StdResid=lmodel$Residuals[,2], Probs=predict(lmodel$model,
                          meuse@data[,VarNames],type='fitted') )

##### Plot fit and probabilities
resid(lmodel$model, "partial", pl="loess")
resid(lmodel$model, "partial", pl=TRUE)                      # plot residuals
resid(lmodel$model, "gof")                                # global test of goodness of fit
lp1 <- resid(lmodel$model, "lp1")                         # Approx. leave-out linear predictors
-2 * sum(meuse@data$DepVar * lp1 + log(1-plogis(lp1))) # Approx leave-out-1 deviance
spplot(meuse, c('Probs'))                                # plot estimated probabilities at points

```

Description

Calculate statistical moments of a distribution

Usage

```
moments(x, plot = FALSE)
```

Arguments

x	numeric vector
plot	plot of distribution (TRUE/FALSE)

Value

A vector with the following values

- min Minimum
- 25th 25th percentile
- mean Arithmetic mean
- gmean Geometric mean
- hmean Harmonic mean
- median 50th percentile
- 7th5 75th percentile
- max Maximum
- stdv Standard deviation
- var Variance
- cv Coefficient of variation (percent)
- mad Median absolute deviation
- skew Skewness
- kurt Kurtosis
- nmodes Number of modes
- mode Mode (dominate)

Author(s)

Jeffrey S. Evans <jeffrey_evans<at>tnc.org>

Examples

```
x <- runif(1000,0,100)
( d <- moments(x, plot=TRUE) )
( mode.x <- moments(x, plot=FALSE)[16] )
```

morans.plot

Autocorrelation Plot

Description

Autocorrelation plot (Anselin 1996), following Chen (2015), aka, Moran's-I plot (univariate or bivariate)

Usage

```
morans.plot(x, y = NULL, coords = NULL, type.ac = c("xy", "yx"),
dist.function = "inv.power", scale.xy = TRUE, scale.morans = FALSE,
...)
```

Arguments

x	Vector of x response variables
y	Vector of y response variables
coords	A matrix of coordinates corresponding to [x,y]
type.ac	Type of autocorrelation plot ("xy", "yx")
dist.function	("inv.power", "neg.exponent")
scale.xy	(TRUE/FALSE) scale the x,y vectors
scale.morans	(FALSE/TRUE) standardize the Moran's index to an expected [-1 to 1]?
...	Additional arguments passed to plot

Details

The argument "type" controls the plot for x influencing y (type="xy") or y influencing x (type="yx"). If y is not defined then the statistic is univariate and only the "xy" plot will be available. The linear relationship between x and its spatial lag (Wx) is indicative of the spatial autoregressive process, underlying the spatial dependence. The statistic can be autocorrelation (univariate or cross-correlation (bivariate). The quadrants are the zero intercept for random autocorrelation and the red line represents the trend in autocorrelation. The quadrants in the plot indicate the type of spatial association/interaction (Anselin 1996). For example the upper-left quadrant represents negative associations of low values surrounded by high and the lower-right quadrant represents negative associations of high values surrounded by low.

Value

A plot of the scaled variable against its spatially lagged values.

Note

if y is not specified the univariate statistic for x is returned. the coords argument is only used if k = NULL. Can also be an sp object with relevant x,y coordinate slot (ie., points or polygons). If w = NULL, the default method for deriving spatial weights matrix, options are: inverse power or negative exponent. If scale.xy = FALSE it is assumed that they are already scaled following Chen (2015).

Author(s)

Jeffrey S. Evans <jeffrey_evans@tnc.org>

References

- Chen., Y. (2015) A New Methodology of Spatial Cross-Correlation Analysis. PLoS One 10(5):e0126158. doi:10.1371/journal.pone.0126158
- Anselin, L. (1996) The Moran scatterplot as an ESDA tool to assess local instability in spatial association. pp. 111-125 in M. M. Fischer, H. J. Scholten and D. Unwin (eds) Spatial analytical perspectives on GIS, London, Taylor and Francis
- Anselin, L. (1995) Local indicators of spatial association, Geographical Analysis, 27:93-115

Examples

```
library(sp)
library(spdep)
data(meuse)
coordinates(meuse) <- ~x+y

# Autocorrelation (univariate)
morans.plot(meuse$zinc, coords = coordinates(meuse))

# Cross-correlation of: x influencing y and y influencing x
par(mfrow=c(1,2))
morans.plot(x=meuse$zinc, y=meuse$copper, coords = coordinates(meuse),
            scale.morans = TRUE)
morans.plot(x=meuse$zinc, y=meuse$copper, coords = coordinates(meuse),
            scale.morans = TRUE, type.ac="yx")
```

mwCorr

Dutilleul moving window bivariate raster correlation

Description

A bivariate raster correlation using Dutilleul's modified t-test

Usage

```
mwCorr(...)
```

Arguments

... Parameters to be passed to the modern version of the function

nni

Average Nearest Neighbour Index (NNI)

Description

Calculates the NNI as a measure of clustering or dispersal

Usage

```
nni(x, win = "hull")
```

Arguments

x	An sp point object
win	Type of window 'hull' or 'extent'

Value

list object containing NNI = nearest neighbor index, z.score = Z Score value, p = p value, expected.mean.distance = Expected meand distance, observed.mean.distance = Observed meand distance.

Note

The nearest neighbour index is expressed as the ratio of the observed distance divided by the expected distance. The expected distance is the average distance between neighbours in a hypothetical random distribution. If the index is less than 1, the pattern exhibits clustering; if the index is greater than 1, the trend is toward dispersion or competition. The Nearest Neighbour Index is calculated as: Mean Nearest Neighbour Distance (observed) $D(nn) = \text{sum}(\min(D_{ij})/N)$ Mean Random Distance (expected) $D(e) = 0.5 \sqrt{A/N}$ Nearest Neighbour Index $NNI = D(nn)/D(e)$ Where; D=neighbour distance, A=Area

Depends: sp, spatstat

Author(s)

Jeffrey S. Evans <jeffrey_evans@tnc.org>

References

- Clark, P.J., and F.C. Evans (1954) Distance to nearest neighbour as a measure of spatial relationships in populations. *Ecology* 35:445-453
 Cressie, N (1991) Statistics for spatial data. Wiley & Sons, New York.

Examples

```
require(sp)
data(meuse)
coordinates(meuse) <- ~x+y
nni(meuse)
```

Description

Calculates the inhomogeneous O-ring point pattern statistic (Wiegand & Maloney 2004)

Usage

```
o.ring(x, inhomogeneous = FALSE, ...)
```

Arguments

x	spatstat ppp object
inhomogeneous	(boolean) Run homogeneous (pcf) or inhomogeneous (pcfinhom)
...	additional arguments passed to pcf or pcfinhom

Value

plot of o-ring and data.frame with plot labels and descriptions

Note

The function $K(r)$ is the expected number of points in a circle of radius r centered at an arbitrary point (which is not counted), divided by the intensity l of the pattern. The alternative pair correlation function $g(r)$, which arises if the circles of Ripley's K -function are replaced by rings, gives the expected number of points at distance r from an arbitrary point, divided by the intensity of the pattern. Of special interest is to determine whether a pattern is random, clumped, or regular.

Using rings instead of circles has the advantage that one can isolate specific distance classes, whereas the cumulative K -function confounds effects at larger distances with effects at shorter distances. Note that the K -function and the O-ring statistic respond to slightly different biological questions. The accumulative K -function can detect aggregation or dispersion up to a given distance r and is therefore appropriate if the process in question (e.g., the negative effect of competition) may work only up to a certain distance, whereas the O-ring statistic can detect aggregation or dispersion at a given distance r . The O-ring statistic has the additional advantage that it is a probability density function (or a conditioned probability spectrum) with the interpretation of a neighborhood density, which is more intuitive than an accumulative measure.

Depends: spatstat

Author(s)

Jeffrey S. Evans <jeffrey_evans@tnc.org>

References

Wiegand T., and K. A. Moloney (2004) Rings, circles and null-models for point pattern analysis in ecology. *Oikos* 104:209-229

Examples

```
library(spatstat)
data(lansing)
x <- spatstat::unmark(split(lansing)$maple)
o.ring(x)
```

*oli.asw**Query AWS-OLI*

Description

Query of Amazon AWS OLI-Landsat 8 cloud service

Usage

```
oli.asw(path, row, dates, cloud.cover = 10, processing)
```

Arguments

path	landsat path
row	landsat row
dates	dates, single or start-stop range in YYYY-MM-DD format
cloud.cover	percent cloud cover
processing	processing level ("L1GT" or "L1T")

Value

data.frame object with:

- entityId - Granule ID
- L = Landsat
- X = Sensor
- SS = Satellite
- PPP = WRS path
- RRR = WRS row
- YYYYMMDD = Acquisition date
- yyyyymmdd = Processing date
- CC = Collection number
- TX = Collection category
- acquisitionDate - POSIXct YYYY-MM-DD (eg., 2015-01-02)
- cloudCover -
- processingLevel - USGS processing level
- path - Landsat path
- row - Landsat row

Note

Amazons AWS cloud service is hosting OLI Landsat 8 data granules <https://aws.amazon.com/public-datasets/landsat/> <https://aws.amazon.com/blogs/aws/start-using-landsat-on-aws/>

USGS Landsat collections: <https://landsat.usgs.gov/landsat-collections> Pre-collection processing levels: "L1T", "L1GT", "L1G" Collection 1 processing levels: "L1TP", "L1GT", "L1GS" "L1T" and "L1TP" - Radiometrically calibrated and orthorectified (highest level processing) "L1GT" and "L1GT" - Radiometrically calibrated and systematic geometric corrections "L1G" and "L1GS" - Radiometrically calibrated with systematic ephemeris correction

Author(s)

Jeffrey S. Evans <jeffrey_evans@tnc.org>

Examples

```
## Not run:
# Query path 126, row 59, 2013-04-15 to 2017-03-09, <20% cloud cover
( p126r59.oli <- oli.awr(path=126, row=59, dates = c("2013-04-15", "2017-03-09"),
                           cloud.cover = 20) )

# Download query images from query
for( i in 1:length(p126r59.oli$download_url)) {
  oli.url <- p126r59.oli$download_url[i]
  try(utils::download.file(url=oli.url, destfile=getwd(), mode = "wb"))
}

## End(Not run)
```

optimal.k

optimalK

Description

Find optimal k of k-Medoid partitions using silhouette widths

Usage

```
optimal.k(x, nk = 10, plot = TRUE, cluster = TRUE, clara = FALSE,
          ...)
```

Arguments

x	Numeric dataframe, matrix or vector
nk	Number of clusters to test (2:nk)
plot	Plot cluster silhouettes(TRUE/FALSE)
cluster	Create cluster object with optimal k

<code>clara</code>	Use clara model for large data
...	Additional arguments passed to clara

Note

Depends: cluster

Author(s)

Jeffrey S. Evans <jeffrey_evans<at>tnc.org>

References

Theodoridis, S. & K. Koutroumbas(2006) Pattern Recognition 3rd ed.

See Also

[pam](#) for details on Partitioning Around Medoids (PAM)

[clara](#) for details on Clustering Large Applications (clara)

Examples

```
require(cluster)
x <- rbind(cbind(rnorm(10,0,0.5), rnorm(10,0,0.5)),
            cbind(rnorm(15,5,0.5), rnorm(15,5,0.5)))

clust <- optimal.k(x, 20, plot=TRUE, cluster=TRUE)
plot(silhouette(clust), col = c('red', 'green'))
plot(clust, which.plots=1, main='K-Medoid fit')

# Extract multivariate and univariate medioids (class centres)
clust$medoids
pam(x[,1], 1)$medoids

# join clusters to data
x <- data.frame(x, k=clust$clustering)
```

optimized.sample.variance

Optimized sample variance

Description

Draws an optimal sample that minimizes or maximizes the sample variance

Usage

```
optimized.sample.variance(x, n, type = "maximized")
```

Arguments

x	A vector to draw a sample from
n	Number of samples to draw
type	Type of sample variance optimization c("maximized", "minimized")

Value

A data.frame with "idx" representing the index of the original vector and "y" is the value of the sampled data

Author(s)

Jeffrey S. Evans <jeffrey_evans@tnc.org>

Examples

```
library(sp)
  data(meuse)
  coordinates(meuse) <- ~x+y

n = 15
# Draw n samples that maximize the variance of y
( max.sv <- optimized.sample.variance(meuse$zinc, 15) )

# Draw n samples that minimize the variance of y
( min.sv <- optimized.sample.variance(meuse$zinc, 15, type="minimized") )

# Plot results
plot(meuse, pch=19, col="grey")
  plot(meuse[max.sv$idx,], col="red", add=TRUE, pch=19)
  plot(meuse[min.sv$idx,], col="blue", add=TRUE, pch=19)
  box()
  legend("topleft", legend=c("population", "maximized variance",
    "minimized variance"), col=c("grey", "red", "blue"),
    pch=c(19, 19, 19))

## Not run:
# Raster example (not memory safe)
library(raster)
  r <- raster(system.file("external/test.grd", package="raster"))

# Calculate optimal sample variance and coerce to SpatialPointDataFrame using xyFromCell
( min.sv <- optimized.sample.variance(getValues(r), n, type="minimized") )
  min.sv <- sp::SpatialPointsDataFrame(xyFromCell(r, min.sv[, "idx"]),
    spatial=TRUE), data=min.sv)
( max.sv <- optimized.sample.variance(getValues(r), n) )
  max.sv <- sp::SpatialPointsDataFrame(xyFromCell(r, max.sv[, "idx"]),
    spatial=TRUE), data=max.sv)

plot(r)
  plot(max.sv, col="blue", add=TRUE, pch=19)
```

```

plot(min.sv, col="red", add=TRUE, pch=19)
box()
legend("topleft", legend=c("maximized variance", "minimized variance"),
       col=c("red", "blue"), pch=c(19,19))

## End(Not run)

```

outliers*Outliers***Description**

Identify outliers using modified Z-score

Usage

```
outliers(x, s = 1.4826)
```

Arguments

x	A numeric vector
s	Scaling factor for mad statistic

Value

value for the modified Z-score

Author(s)

Jeffrey S. Evans <jeffrey_evans@tnc.org>

References

Iglewicz, B. & D.C. Hoaglin (1993) How to Detect and Handle Outliers, American Society for Quality Control, Milwaukee, WI.

Examples

```

# Create data with 3 outliers
x <- seq(0.1, 5, length=100)
x[98:100] <- c(100, 55, 250)

# Calculate Z score
Z <- outliers(x)

# Show number of extreme outliers using Z-score
length(Z[Z > 9.9])

```

```
# Remove extreme outliers  
x <- x[-which(Z > 9.9)]
```

parea.sample*Percent area sample*

Description

Creates a point sample of polygons where n is based on percent area

Usage

```
parea.sample(x, pct = 0.1, join = FALSE, msamp = 1, sf = 4046.86,  
stype = "hexagonal", ...)
```

Arguments

x	sp SpatialPolygonsDataFrame object
pct	Percent of area sampled
join	Join polygon attributed to point sample
msamp	Minimum samples
sf	Scaling factor (default is meters to acres conversion factor)
stype	Sampling type ('random', 'regular', 'nonaligned', 'hexagonal')
...	Additional arguments passed to spsample

Value

A SpatialPointsDataFrame with polygon samples

Note

This function results in an adaptive sample based on the area of each polygon

Author(s)

Jeffrey S. Evans <jeffrey_evans@tnc.org>

Examples

```
require(sp)
sr1=Polygons(list(Polygon(cbind(c(180114, 180553, 181127, 181477, 181294, 181007, 180409,
  180162, 180114), c(332349, 332057, 332342, 333250, 333558, 333676,
  332618, 332413, 332349)))), '1')
sr2=Polygons(list(Polygon(cbind(c(180042, 180545, 180553, 180314, 179955, 179142, 179437,
  179524, 179979, 180042), c(332373, 332026, 331426, 330889, 330683,
  331133, 331623, 332152, 332357, 332373)))), '2')
sr=SpatialPolygons(list(sr1,sr2))
srdf=SpatialPolygonsDataFrame(sr, data.frame(row.names=c('1','2'), PIDS=1:2))

ars <- parea.sample(srdf, pct=0.20, stype='random')
plot(srdf)
  plot(ars, pch=20, add=TRUE)
```

plot.effect.size *Plot effect size*

Description

Plot function for effect.size object

Usage

```
## S3 method for class 'effect.size'
plot(x, ...)
```

Arguments

x	A effect.size object
...	Additional arguments passed to plot

Author(s)

Jeffrey S. Evans <jeffrey_evans@tnc.org>

plot.loess.boot *Plot Loess Bootstrap*

Description

Plot function for loess.boot object

Usage

```
## S3 method for class 'loess.boot'  
plot(x, ...)
```

Arguments

x	A loess.boot object
...	Additional arguments passed to plot

Author(s)

Jeffrey S. Evans <jeffrey_evans@tnc.org>

References

- Cleveland, WS, (1979) Robust Locally Weighted Regression and Smoothing Plots Journal of the American Statistical Association 74:829-836
- Efron, B., and R. Tibshirani (1993) An Introduction to the Bootstrap Chapman and Hall, New York
- Härdle, W., (1989) Applied Nonparametric Regression Cambridge University Press, NY.
- Tibshirani, R. (1988) Variance stabilization and the bootstrap. Biometrika 75(3):433-44.

Examples

```
n=1000  
x <- seq(0, 4, length.out=n)  
y <- sin(2*x)+ 0.5*x + rnorm(n, sd=0.5)  
sb <- loess.boot(x, y, nreps = 99, confidence = 0.90, span = 0.40)  
plot(sb)
```

point.in.poly*Point and Polygon Intersect***Description**

Intersects point and polygon feature classes and adds polygon attributes to points

Usage

```
point.in.poly(x, y, sp = TRUE, duplicate = TRUE, ...)
```

Arguments

x	sp SpatialPointsDataFrame or SpatialPoints or sf point object
y	sp SpatialPolygonsDataFrame or sf polygon object
sp	(TRUE/FALSE) Return an sp class object, else returns sf class object
duplicate	(TRUE/FALSE) Return duplicated features with more than one polygon intersection
...	Additional arguments passed to sf::st_join

Value

A SpatialPointsDataFrame or sf

Note

If duplicate argument is TRUE and more than one polygon intersection occurs, points will be duplicated (new row added) and all attributes joined. However, if duplicate is FALSE, with duplicate intersections, a new column for each unique intersecting polygon will be returned and the points will not be duplicated. For example, if a point intersect three polygons, three new columns will be added representing the polygons ID.

Author(s)

Jeffrey S. Evans <jeffrey_evans@tnc.org>

Examples

```
#### Simple one-to-one feature overlay.
require(sp)
data(meuse)
coordinates(meuse) = ~x+y
meuse@data$test.na <- NA

sr1=Polygons(list(Polygon(cbind(c(180114, 180553, 181127, 181477, 181294,
181007, 180409, 180162, 180114), c(332349, 332057, 332342, 333250, 333558,
333676, 332618, 332413, 332349)))), '10')
```

```

sr2=Polygons(list(Polygon(cbind(c(180042, 180545, 180553, 180314, 179955, 179142,
  179437, 179524, 179979, 180042), c(332373, 332026, 331426, 330889, 330683,
  331133, 331623, 332152, 332357, 332373)))), '20')
sr3=Polygons(list(Polygon(cbind(c(179110, 179907, 180433, 180712, 180752, 180329,
  179875, 179668, 179572, 179269, 178879, 178600, 178544, 179046, 179110),
  c(331086, 330620, 330494, 330265, 330075, 330233, 330336, 330004,
  329783, 329665, 329720, 329933, 330478, 331062, 331086)))), '30')
sr4=Polygons(list(Polygon(cbind(c(180304, 180403, 179632, 179420, 180304),
  c(332791, 333204, 333635, 333058, 332791)))), '40')
sr=SpatialPolygons(list(sr1,sr2,sr3,sr4))
polys=SpatialPolygonsDataFrame(sr, data.frame(row.names=c('10','20','30','40'),
  PIDS=1:4, y=runif(4)))
polys@data$pid <- polys@data$PIDS + 100

plot(polys)
plot(meuse, pch=19, add=TRUE)

# Point in polygon overlay
pts.poly <- point.in.poly(meuse, polys)
head(pts.poly@data)

# Count points in each polygon
tapply(pts.poly$cadmium, pts.poly$pid, FUN=length)

##### Complex many-to-one feature overlay.
require(sf)
p <- sf::st_polygon(list(rbind(c(0,0), c(1,0), c(1,1), c(0,1), c(0,0))))
polys <- sf::st_sf(sf::st_sf(p, p + c(.8, .2), p + c(.2, .8)))
pts <- sf::st_sf(sf::st_sample(polys, size=100))

# Duplicates points for each new polygon, no attributes so returns IDs for features
pts.poly.dup <- point.in.poly(pts, polys)
head(pts.poly.dup@data)

## Not run:
# **** Should throw error due to lack of attributes ****
pts.poly <- point.in.poly(pts, polys, duplicate = FALSE)

## End(Not run)

# Coerce to sp class objects
x <- as(pts, "Spatial")
x <- SpatialPointsDataFrame(x, data.frame(IDS=1:nrow(x), pty=runif(nrow(x))))
y <- as(polys, "Spatial")
y <- SpatialPolygonsDataFrame(y, data.frame(IDS=1:nrow(y), py=runif(nrow(y))))

# Returns point attributes with column for each unique polygon
pts.poly <- point.in.poly(x, y, duplicate = FALSE)
head(pts.poly@data)

# Duplicates points for each new polygon, joins all attributes
pts.poly.dup <- point.in.poly(x, y)
head(pts.poly.dup@data)

```

```
# Count points in each polygon
tapply(pts.poly.dup$IDS.x, pts.poly.dup$IDS.y, FUN=length)
```

poly.regression *Local Polynomial Regression*

Description

Calculates a Local Polynomial Regression for smoothing or imputation of missing data.

Usage

```
poly.regression(y, x = NULL, s = 0.75, impute = FALSE,
na.only = FALSE, ci = FALSE, ...)
```

Arguments

y	Vector to smooth or impute NA values
x	Optional x covariate data (must match dimensions of y)
s	Smoothing parameter (larger equates to more smoothing)
impute	(FALSE/TRUE) Should NA values be imputed
na.only	(FALSE/TRUE) Should only NA values be change in y
ci	(FALSE/TRUE) Should confidence intervals be returned
...	Additional arguments passed to loess

Value

If ci = FALSE, a vector of smoothed values otherwise a list object with:
 loess A vector, same length of y, representing the smoothed or imputed data
 lower.ci Lower confidence interval
 upper.ci Upper confidence interval

Note

This is a wrapper function for loess that simplifies data smoothing and imputation of missing values. The function allows for smoothing a vector, based on an index (derived automatically) or covariates. If the impute option is TRUE NA values are imputed, otherwise the returned vector will still have NA's present. If impute and na.only are both TRUE the vector is returned, without being smoothed but with imputed NA values filled in.

The loess weight function is defined using the tri-cube weight function $w(x) = (1-|x|^3)^3$ where; x is the distance of a data point from the point on the curve being fitted.

Author(s)

Jeffrey S. Evans <jeffrey_evans@tnc.org>

See Also

[loess](#) for loess ... model options

Examples

```
x <- seq(-20, 20, 0.1)
y <- sin(x)/x + rnorm(length(x), sd=0.03)
p <- which(y == "NaN")
y <- y[-p]
r <- poly.regression(y, ci=TRUE, s=0.30)

plot(y,type="l", lwd=0.5, main="s = 0.10")
y.polygon <- c((r$lower.ci)[1:length(y)], (r$upper.ci)[rev(1:length(y))])
x.polygon <- c(1:length(y), rev(1:length(y)))
polygon(x.polygon, y.polygon, col="#00009933", border=NA)
lines(r$loess, lwd=1.5, col="red")

# Impute NA values, replacing only NA's
y.na <- y
y.na[c(100,200,300)] <- NA
p.y <- poly.regression(y.na, s=0.10, impute = TRUE, na.only = TRUE)
y = p.y

plot(p.y,type="l", lwd=1.5, col="blue", main="s = 0.10")
lines(y, lwd=1.5, col="red")
```

polyPerimeter

Polygon perimeter

Description

Calculates the perimeter length(s) for a polygon object

Usage

```
polyPerimeter(x)
```

Arguments

x	sp class SpatialPolygonsDataFrame object
---	--

Value

A vector of polygon perimeters

Author(s)

Jeffrey S. Evans <jeffrey_evans@tnc.org>

Examples

```
library(sp)
p1 <- Polygons(list(Polygon(cbind(c(2,4,4,1,2),c(2,3,5,4,2)))), "1")
p2 <- Polygons(list(Polygon(cbind(c(5,4,2,5),c(2,3,2,2)))), "2")
p3 <- Polygons(list(Polygon(cbind(c(4,4,5,10,4),c(5,3,2,5,5)))), "3")
polys <- SpatialPolygons(list(p1,p2,p3), 1:3)

polyPerimeter(polys)
```

pp.subsample

Point process random subsample

Description

Generates random subsample based on density estimate of observations

Usage

```
pp.subsample(x, n, window = "hull", sigma = "Scott", wts = NULL,
            gradient = 1, edge = FALSE)
```

Arguments

<i>x</i>	An sp class SpatialPointsDataFrame or SpatialPoints object
<i>n</i>	Number of random samples to generate
<i>window</i>	Type of window (hull or extent)
<i>sigma</i>	Bandwidth selection method for KDE, default is 'Scott'. Options are 'Scott', 'Stoyan', 'Diggle', 'likelihood', and 'geometry'
<i>wts</i>	Optional vector of weights corresponding to point pattern
<i>gradient</i>	A scaling factor applied to the sigma parameter used to adjust the gradient decent of the density estimate. The default is 1, for no adjustment (downweight < 1 upweight > 1)
<i>edge</i>	Apply Diggle edge correction (TRUE/FALSE)

Value

sp class SpatialPointsDataFrame containing random subsamples

Note

The window type creates a convex hull by default or, optionally, uses the maximum extent (envelope).

Available bandwidth selection methods are: Scott (Scott 1992), Scott's Rule for Bandwidth Selection (1st order) Diggle (Berman & Diggle 1989), Minimise the mean-square error via cross validation (2nd order) likelihood (Loader 1999), Maximum likelihood cross validation (2nd order) geometry, Bandwidth is based on simple window geometry (1st order) Stoyan (Stoyan & Stoyan 1995), Based on pair-correlation function (strong 2nd order)

Note; resulting bandwidth can vary widely by method. the 'diggle' method is intended for selecting bandwidth representing 2nd order spatial variation whereas the 'scott' method will represent 1st order trend. the 'geometry' approach will also represent 1st order trend. for large datasets, caution should be used with the 2nd order 'likelihood' approach, as it is slow and computationally expensive. finally, the 'stoyan' method will produce very strong 2nd order results. '

Depends: sp, spatstat

Author(s)

Jeffrey S. Evans <jeffrey_evans@tnc.org>

References

- Berman, M. and Diggle, P. (1989) Estimating weighted integrals of the second-order intensity of a spatial point process. *Journal of the Royal Statistical Society, series B* 51, 81-92. Berman, M. and Diggle, P. (1989) Estimating weighted integrals of the second-order intensity of a spatial point process. *Journal of the Royal Statistical Society, series B* 51, 81-92.
- Fithian, W & T. Hastie (2013) Finite-sample equivalence in statistical models for presence-only data. *Annals of Applied Statistics* 7(4): 1917-1939
- Hengl, T., H. Sierdsema, A. Radovic, and A. Dilo (2009) Spatial prediction of species distributions from occurrence-only records: combining point pattern analysis, ENFA and regression-kriging. *Ecological Modelling*, 220(24):3499-3511
- Loader, C. (1999) Local Regression and Likelihood. Springer, New York.
- Scott, D.W. (1992) Multivariate Density Estimation. Theory, Practice and Visualization. New York, Wiley.
- Stoyan, D. and Stoyan, H. (1995) Fractals, random shapes and point fields: methods of geometrical statistics. John Wiley and Sons.
- Warton, D.i., and L.C. Shepherd (2010) Poisson Point Process Models Solve the Pseudo-Absence Problem for Presence-only Data in Ecology. *The Annals of Applied Statistics*, 4(3):1383-1402

Examples

```
require(spatstat)
require(sp)
data(bci)
trees <- as(bci, 'SpatialPoints')
n=round(length(trees) * 0.10, digits=0)
```

```

trees.wrs <- pp.subsample(trees, n=n, window='hull')
plot(trees, pch=19, col='black')
plot(trees.wrs, pch=19, col='red', add=TRUE)
box()
title('10% subsample')
legend('bottomright', legend=c('Original sample', 'Subsample'),
       col=c('black','red'), pch=c(19,19))

```

print.cross.cor *Print spatial cross correlation*

Description

print method for class "cross.cor"

Usage

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

Arguments

x	Object of class cross.cor
...	Ignored

print.effect.size *Print effect size*

Description

print method for class "effect.size"

Usage

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

Arguments

x	Object of class effect.size
...	Ignored

print.loess.boot	<i>Print Loess bootstrap model</i>
------------------	------------------------------------

Description

print method for class "loess.boot"

Usage

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

Arguments

x	Object of class loess.boot
...	Ignored

pseudo.absence	<i>Pseudo-absence random samples</i>
----------------	--------------------------------------

Description

Generates pseudo-absence samples based on density estimate of known locations

Usage

```
pseudo.absence(x, n, window = "hull", Mask = NULL, s = NULL,
  sigma = "Scott", wts = NULL, KDE = FALSE, gradient = 1,
  p = NULL, edge = FALSE)
```

Arguments

x	An sp class SpatialPointsDataFrame or SpatialPoints object
n	Number of random samples to generate
window	Type of window (hull OR extent), overridden if mask provided
Mask	Optional rasterLayer class mask raster. The resolution of the density estimate will match mask.
s	Optional resolution passed to window argument. Caution should be used due to long processing times associated with high resolution. In contrast, coarse resolution can exclude known points.
sigma	Bandwidth selection method for KDE, default is 'Scott'. Options are 'Scott', 'Stoyan', 'Diggle', 'likelihood', and 'geometry'
wts	Optional vector of weights corresponding to point pattern

KDE	save KDE raster (TRUE/FALSE)
gradient	A scaling factor applied to the sigma parameter used to adjust the gradient decent of the density estimate. The default is 1, for no adjustment (downweight < 1 upweight > 1)
p	Minimum value for probability distribution (must be > 0)
edge	Apply Diggle edge correction (TRUE/FALSE)

Details

The window type creates a convex hull by default or, optionally, uses the maximum extent (envelope). If a mask is provided the kde will represent areas defined by the mask and defines the area that pseudo absence data will be generated. Available bandwidth selection methods are:

- Scott (Scott 1992), Scott's Rule for Bandwidth Selection (1st order)
- Diggle (Berman & Diggle 1989), Minimize the mean-square error via cross validation (2nd order)
- likelihood (Loader 1999), Maximum likelihood cross validation (2nd order)
- geometry, Bandwidth is based on simple window geometry (1st order)
- Stoyan (Stoyan & Stoyan 1995), Based on pair-correlation function (strong 2nd order)

Note; resulting bandwidth can vary widely by method. the 'diggle' method is intended for selecting bandwidth representing 2nd order spatial variation whereas the 'scott' method will represent 1st order trend. the 'geometry' approach will also represent 1st order trend. for large datasets, caution should be used with the 2nd order 'likelihood' approach, as it is slow and computationally expensive. finally, the 'stoyan' method will produce very strong 2nd order results.

Value

A list class object with the following components:

- sample SpatialPointsDataFrame containing random samples
- kde sp RasterLayer class of KDE estimates (IF KDE = TRUE)
- sigma Selected bandwidth of KDE

Author(s)

Jeffrey S. Evans <jeffrey_evans@tnc.org>

References

- Berman, M. and Diggle, P. (1989) Estimating weighted integrals of the second-order intensity of a spatial point process. *Journal of the Royal Statistical Society, series B* 51, 81-92.
- Fithian, W & T. Hastie (2013) Finite-sample equivalence in statistical models for presence-only data. *Annals of Applied Statistics* 7(4): 1917-1939
- Hengl, T., H. Sierdsema, A. Radovic, and A. Dilo (2009) Spatial prediction of species distributions from occurrence-only records: combining point pattern analysis, ENFA and regression-kriging. *Ecological Modelling*, 220(24):3499-3511

- Loader, C. (1999) Local Regression and Likelihood. Springer, New York.
- Scott, D.W. (1992) Multivariate Density Estimation. Theory, Practice and Visualization. New York, Wiley.
- Stoyan, D. and Stoyan, H. (1995) Fractals, random shapes and point fields: methods of geometrical statistics. John Wiley and Sons.
- Warton, D.i., and L.C. Shepherd (2010) Poisson Point Process Models Solve the Pseudo-Absence Problem for Presence-only Data in Ecology. The Annals of Applied Statistics, 4(3):1383-1402

Examples

```

library(raster)
library(sp)
data(meuse)
data(meuse.grid)
coordinates(meuse) = ~x+y
coordinates(meuse.grid) = ~x+y
proj4string(meuse.grid) <- CRS("+init=epsg:28992")
gridded(meuse.grid) = TRUE
r <- raster(meuse.grid)

# Using a raster mask
pa <- pseudo.absence(meuse, n=100, window='hull', KDE=TRUE, Mask = r,
                      sigma='Diggle', s=50)
col.br <- colorRampPalette(c('blue','yellow'))
plot(pa$kde, col=col.br(10))
plot(meuse, pch=20, cex=1, add=TRUE)
plot(pa$sample, col='red', pch=20, cex=1, add=TRUE)
legend('top', legend=c('Presence', 'Pseudo-absence'),
       pch=c(20,20),col=c('black','red'))

# With clustered data
library(sp)
library(spatstat)
data(bel)
trees <- as(bel, 'SpatialPoints')
trees <- SpatialPointsDataFrame(coordinates(trees),
                                 data.frame(ID=1:length(trees)))
trees.abs <- pseudo.absence(trees, n=100, window='extent', KDE=TRUE)

col.br <- colorRampPalette(c('blue','yellow'))
plot(trees.abs$kde, col=col.br(10))
plot(trees, pch=20, cex=0.50, add=TRUE)
plot(trees.abs$sample, col='red', pch=20, cex=1, add=TRUE)
legend('top', legend=c('Presence', 'Pseudo-absence'),
       pch=c(20,20),col=c('black','red'))

```

pu

Biodiversity Planning Units

Description

Subset of biodiversity planning units for Haiti ecoregional spatial reserve plan

Format

A sp SpatialPolygonsDataFrame with 5919 rows and 46 variables:

UNIT_ID Unique planning unit ID
DR_Dr_A Biodiversity target
DR_Dr_L Biodiversity target
Ht_Dr_A Biodiversity target
Ht_Dr_L Biodiversity target
DR_Ms_A Biodiversity target
DR_Ms_L Biodiversity target
Ht_Ms_L Biodiversity target
DR_LM_M Biodiversity target
H_LM_M_L Biodiversity target
H_LM_R_L Biodiversity target
DR_LM_R_L Biodiversity target
DR_Rn_L Biodiversity target
DR_LM_R_S Biodiversity target
DR_Rn_S Biodiversity target
DR_Ms_S Biodiversity target
Ht_Ms_A Biodiversity target
DR_Ms_E Biodiversity target
DR_Ms_I Biodiversity target
DR_Rn_E Biodiversity target
DR_Rn_I Biodiversity target
H_LM_R_E Biodiversity target
Ht_Ms_E Biodiversity target
Ht_Rn_E Biodiversity target
DR_Rn_A Biodiversity target
Ht_Rn_A Biodiversity target
Ht_Rn_I Biodiversity target
Ht_Dr_E Biodiversity target

Ht_Ms_S Biodiversity target
Ht_Dr_S Biodiversity target
Ht_Rn_L Biodiversity target
Ht_Th_A Biodiversity target
Ht_Th_L Biodiversity target
Ht_Th_S Biodiversity target
Ht_Dr_U Biodiversity target
Ht_Dr_I Biodiversity target
Ht_Ms_I Biodiversity target
H_LM_M_A Biodiversity target
H_LM_M_E Biodiversity target
H_LM_R_A Biodiversity target
H_LM_M_S Biodiversity target
H_LM_R_I Biodiversity target
H_LM_R_S Biodiversity target
Ht_Rn_S Biodiversity target
Ht_Ms_U Biodiversity target
Ht_Rn_U Biodiversity target

Source

http://maps.tnc.org/gis_data.html

References

Evans, J.S., S.R. Schill, G.T. Raber (2015) A Systematic Framework for Spatial Conservation Planning and Ecological Priority Design in St. Lucia, Eastern Caribbean. Chapter 26 in Central American Biodiversity : Conservation, Ecology and a Sustainable Future. F. Huettman (eds). Springer, NY.

raster.deviation *Raster local deviation from the global trend*

Description

Calculates the local deviation from the raster, a specified global statistic or a polynomial trend of the raster.

Usage

```
raster.deviation(x, type = "trend", s = 3, degree = 1,  
global = FALSE)
```

Arguments

x	raster object
type	The global statistic to represent the local deviation options are: "trend", "min", "max", "mean", "median"
s	Size of matrix (focal window), not used with type="trend"
degree	The polynomial degree if type is trend, options are 1 and 2.
global	Use single global value for deviation or cell-level values (FALSE/TRUE). Argument is ignored for type="trend"

Value

raster class object of the local deviation from the raster or specified global statistic

Note

The deviation from the trend is derived as $[y\text{-hat} - y]$ where; $y\text{-hat}$ is the Nth-order polynomial. Whereas the deviation from a global statistic is $[y - y\text{-hat}]$ where; $y\text{-hat}$ is the local (focal) statistic. The global = TRUE argument allows one to evaluate the local deviation from the global statistic $[stat(x) - y\text{-hat}]$ where; stat(x) is the global value of the specified statistic and $y\text{-hat}$ is the specified focal statistic.

Author(s)

Jeffrey S. Evans <jeffrey_evans@tnc.org>

References

Magee, Lonnie (1998). Nonlocal Behavior in Polynomial Regressions. *The American Statistician*. American Statistical Association. 52(1):20-22 Fan, J. (1996). Local Polynomial Modelling and Its Applications: From linear regression to nonlinear regression. Monographs on Statistics and Applied Probability. Chapman and Hall/CRC. ISBN 0-412-98321-4

Examples

```
library(raster)
data(elev)

# local deviation from first-order trend, global mean and raw value
r.dev.trend <- raster.deviation(elev, type="trend", degree=1)
r.dev.mean <- raster.deviation(elev, type="mean", s=5)
r.gdev.mean <- raster.deviation(elev, type="mean", s=5, global=TRUE)

par(mfrow=c(2,2))
plot(elev, main="original")
plot(r.dev.trend, main="dev from trend")
plot(r.dev.mean, main="dev of mean from raw values")
plot(r.gdev.mean, main="local dev from global mean")
```

raster.downscale *Raster Downscale*

Description

Downscales a raster to a higher resolution raster using a robust regression

Usage

```
raster.downscale(x, y, p = NULL, n = NULL, filename = FALSE,  
scatter = FALSE, ...)
```

Arguments

x	Raster class object representing independent variable(s)
y	Raster class object representing dependent variable
p	Percent sample size
n	Fixed sample size
filename	Name of output raster
scatter	(FALSE/TRUE) Optional scatter plot
...	Additional arguments passed to predict

Value

A list object containing:

- downscale downscaled raster (omitted if filename is defined)
- model rlm model object
- MSE Mean Square Error
- AIC Akaike information criterion

Author(s)

Jeffrey S. Evans <jeffrey_evans@tnc.org>

Examples

```
## Not run:  
library(raster)  
elev <- raster::getData('alt', country='SWZ', mask=TRUE)  
tmax <- raster::getData('worldclim', var='tmax', res=10, lon=8.25, lat=46.8)  
tmax <- crop(tmax[[1]], extent(elev))  
  
tmax.ds <- raster.downscale(elev, tmax, scatter=TRUE)  
par(mfrow=c(2,2))  
plot(tmax, main="Temp max")
```

```

plot(elev, main="elevation")
plot(tmax.ds$downscale, main="Downscaled Temp max")

## End(Not run)

```

raster.entropy*Raster Entropy***Description**

Calculates entropy on integer raster (i.e., 8 bit 0-255)

Usage

```
raster.entropy(x, d = 5, categorical = FALSE, global = FALSE,
filename = FALSE, ...)
```

Arguments

x	Object of class raster (requires integer raster)
d	Size of matrix (window)
categorical	Is the data categorical or continuous (FALSE/TRUE)
global	Should the model use a global or local n to calculate entropy (FALSE/TRUE)
filename	Raster file written to disk
...	Optional arguments passed to writeRaster or dataType

Value

raster class object or specified format raster written to disk

Note

Entropy calculated as: $H = -\sum(P_i \ln(P_i))$ where; P_i , Proportion of one value to total values $P_i=n(p)/m$ and m , Number of unique values Expected range: 0 to $\log(m)$ $H=0$ if window contains the same value in all cells. H increases with the number of different values in the window.

Maximum entropy is reached when all values are different, same as $\log(m) \max.ent <- function(x) log(length(unique(x)))$

Depends: raster

References

Fuchs M., R. Hoffmann, F. Schwonke (2008) Change Detection with GRASS GIS - Comparison of images taken by different sensor. On line at: http://geoinformatics.fsv.cvut.cz/gwiki/Change_Detection_with_GRASS_GIS_-Comparison_of_images_taken_by_different_sensors

Examples

```
require(raster)
r <- raster(ncols=100, nrows=100)
r[] <- round(runif(ncell(r), 1,8), digits=0)

rEnt <- raster.entropy(r, d=5, categorical = TRUE, global = TRUE)
opar <- par
  par(mfcol=c(2,1))
    plot(r)
    plot(rEnt)
par(opar)
```

raster.gaussian.smooth

Gaussian smoothing of raster

Description

Applies a Gaussian smoothing kernel to smooth raster.

Usage

```
raster.gaussian.smooth(x, sigma = 2, n = 5, type = mean, ...)
```

Arguments

x	raster object
sigma	standard deviation (sigma) of kernel (default is 2)
n	Size of the focal matrix, single value (default is 5 for 5x5 window)
type	The statistic to use in the smoothing operator (suggest mean or sd)
...	Additional arguments passed to raster::focal

Value

raster class object of the local distributional moment

Note

This is a simple wrapper for the focal function, returning local statistical moments

Author(s)

Jeffrey S. Evans <jeffrey_evans@tnc.org>

Examples

```

library(raster)
r <- raster(nrows=500, ncols=500, xmn=571823, xmx=616763,
            ymn=4423540, ymx=4453690)
proj4string(r) <- crs("+proj=utm +zone=12 +datum=NAD83 +units=m +no_defs")
r[] <- runif(ncell(r), 1000, 2500)
r <- focal(r, focalWeight(r, 150, "Gauss") )

# Calculate Gaussian smoothing with sigma(s) = 1-4
g1 <- raster.gaussian.smooth(r, sigma=1, nc=11)
g2 <- raster.gaussian.smooth(r, sigma=2, nc=11)
g3 <- raster.gaussian.smooth(r, sigma=3, nc=11)
g4 <- raster.gaussian.smooth(r, sigma=4, nc=11)

par(mfrow=c(2,2))
plot(g1, main="Gaussian smoothing sigma = 1")
plot(g2, main="Gaussian smoothing sigma = 2")
plot(g3, main="Gaussian smoothing sigma = 3")
plot(g4, main="Gaussian smoothing sigma = 4")

```

raster.invert *Invert raster*

Description

Inverts (flip) the values of a raster

Usage

```
raster.invert(x)
```

Arguments

x	raster object
---	---------------

Value

raster class object with inverted (flipped) raster values

Note

Inverts raster values using the formula: $((x - \max(x)) * -1) + \min(x)$

Author(s)

Jeffrey S. Evans <jeffrey_evans@tnc.org>

Examples

```
library(raster)
r <- raster(nrows=500, ncols=500, xmn=571823, xmx=616763,
            ymn=4423540, ymx=4453690)
r[] <- runif(ncell(r), 1, 100)
r <- focal(r, focalWeight(r, 150, "Gauss") )
r.inv <- raster.invert(r)
par(mfrow=c(1,2))
plot(r, main="original raster")
plot(r.inv, main="inverted raster")
```

raster.kendall

Kendall tau trend with continuity correction for raster time-series

Description

Calculates a nonparametric statistic for a monotonic trend based on the Kendall tau statistic and the Theil-Sen slope modification

Usage

```
raster.kendall(x, tau = FALSE, intercept = FALSE, p.value = FALSE,
                z.value = FALSE, confidence = FALSE, autocorrelation = FALSE, ...)
```

Arguments

x	A rasterStack object with at least 5 layers
tau	(FALSE/TRUE) return a raster with the pixel wise tau values
intercept	(FALSE/TRUE) return a raster with the pixel wise intercept values
p.value	(FALSE/TRUE) return a raster with the pixel wise p.values
z.value	(FALSE/TRUE) return a raster with the pixel wise z values
confidence	(FALSE/TRUE) return a raster with the pixel wise 95 pct confidence levels
autocorrelation	(FALSE/TRUE) Apply autocorrelation correction using prewhitening
...	Additional arguments passed to the raster overlay function

Details

This function implements Kendall's nonparametric test for a monotonic trend using the Theil-Sen (Theil 1950; Sen 1968; Siegel 1982) method to estimate the slope and related confidence intervals. Critical values are $Z > 1.96$ representing a significant increasing trend and a $Z < -1.96$ a significant decreasing trend ($p < 0.05$). The null hypothesis can be rejected if Tau = 0. There is also an option for autocorrelation correction using the method proposed in Yue & Wang (2002).

Value

Depending on arguments, a raster layer or rasterBrick object containing:

- raster layer 1 Theil-Sen slope, always returned
- raster layer 2 Kendall's tau two-sided test, if tau TRUE
- raster layer 3 intercept for trend if intercept TRUE, not if prewhitened
- raster layer 4 p value for trend fit if p.value TRUE
- raster layer 5 Z value for trend fit if z.value TRUE
- raster layer 6 lower confidence level at 95-pct if confidence TRUE, not if prewhitened
- raster layer 7 upper confidence level at 95-pct if confidence TRUE, not if prewhitened

Author(s)

Jeffrey S. Evans <jeffrey_evans@tnc.org>

References

- Theil, H. (1950) A rank invariant method for linear and polynomial regression analysis. Nederl. Akad. Wetensch. Proc. Ser. A 53:386-392 (Part I), 53:521-525 (Part II), 53:1397-1412 (Part III).
- Sen, P.K. (1968) Estimates of Regression Coefficient Based on Kendall's tau. Journal of the American Statistical Association. 63(324):1379-1389.
- Siegel, A.F. (1982) Robust Regression Using Repeated Medians. Biometrika, 69(1):242-244
- Yue, S., & Wang, C. Y. (2002). Applicability of prewhitening to eliminate the influence of serial correlation on the Mann-Kendall test. Water Resources Research, 38(6):41-47.

See Also

[kendallTrendTest](#) for model details

[overlay](#) for available ... arguments

Examples

```
## Not run:
library(raster)
r.logo <- stack(system.file("external/rlogo.grd", package="raster"),
                 system.file("external/rlogo.grd", package="raster"),
                 system.file("external/rlogo.grd", package="raster"))

# Calculate trend slope with p-value and confidence level(s)
logo.trend <- raster.kendall(r.logo, tau = TRUE, intercept = TRUE, p.value = TRUE,
                               z.value = TRUE, confidence = TRUE)
names(logo.trend) <- c("slope", "tau", "intercept", "p.value", "z.value", "LCI", "UCI")
plot(logo.trend)

## End(Not run)
```

raster.mds*Raster multidimensional scaling (MDS)*

Description

Multidimensional scaling of raster values within an N x N focal window

Usage

```
raster.mds(r, s = 5, window.median = FALSE, ...)
```

Arguments

r	Raster layer
s	Window size (may be a vector of 1 or 2) of n x n dimension.
window.median	(TRUE/FALSE) Return the median of the MDS matrix values.
...	Additional arguments passed to raster::focal

Value

A raster class object or raster written to disk

Note

An MDS focal function. If only one value provided for s, then a square matrix (window) will be used. If window.median = FALSE then the center value of the matrix is returned and not the median of the matrix

Author(s)

Jeffrey S. Evans <jeffrey_evans@tnc.org>

References

Quinn, G.P., & M.J. Keough (2002) Experimental design and data analysis for biologists. Cambridge University Press. Ch. 18. Multidimensional scaling and cluster analysis.

Examples

```
## Not run:  
library(raster)  
r <- raster(system.file("external/rlogo.grd", package="raster"))  
r <- r / cellStats(r, "max")  
  
diss <- raster.mds(r)  
diss.med <- raster.mds(r, window.median = TRUE)  
par(mfrow=c(2,2))  
plot(r)
```

```

    title("R logo band-1")
    plot( focal(r, w = matrix(1, nrow=5, ncol=5), fun = var) )
    title("Variance")
    plot(diss)
    title("MDS")
    plot(diss.med)
    title("Median MDS")

## End(Not run)

```

raster.modified.ttest *Dutilleul moving window bivariate raster correlation*

Description

A bivariate raster correlation using Dutilleul's modified t-test

Usage

```
raster.modified.ttest(x, y, x.idx = 1, y.idx = 1, d = "AUTO",
                      sub.sample = FALSE, type = "hexagon", p = 0.1, size = NULL)
```

Arguments

x	x raster for correlation, SpatialPixelsDataFrame or SpatialGridDataFrame object
y	y raster for correlation, SpatialPixelsDataFrame or SpatialGridDataFrame object
x.idx	Index for the column in the x raster object
y.idx	Index for the column in the y raster object
d	Distance for finding neighbors
sub.sample	Should a sub-sampling approach be employed (TRUE/FALSE)
type	If sub.sample = TRUE, what type of sample (random or hexagon)
p	If sub.sample = TRUE, what proportion of population should be sampled
size	Fixed sample size

Value

A SpatialPixelsDataFrame or SpatialPointsDataFrame with the following attributes:

- corr Correlation
- Fstat The F-statistic calculated as [degrees of freedom * unscaled F-statistic]
- p.value p-value for the test
- moran.x Moran's-I for x
- moran.y Moran's-I for y

Note

This function provides a bivariate moving window correlation using the modified t-test to account for spatial autocorrelation. Point based subsampling is provided for computation tractability. The hexagon sampling is recommended as it is good at capturing spatial process that includes nonstationarity and anisotropy.

Author(s)

Jeffrey S. Evans <jeffrey_evans@tnc.org>

References

Clifford, P., S. Richardson, D. Hemon (1989), Assessing the significance of the correlation between two spatial processes. Biometrics 45:123-134. Dutilleul, P. (1993), Modifying the t test for assessing the correlation between two spatial processes. Biometrics 49:305-314.

Examples

```
## Not run:
library(gstat)
library(sp)

data(meuse)
data(meuse.grid)
coordinates(meuse) <- ~x + y
coordinates(meuse.grid) <- ~x + y

# GRID-1 log(copper):
v1 <- variogram(log(copper) ~ 1, meuse)
x1 <- fit.variogram(v1, vgm(1, "Sph", 800, 1))
G1 <- krige(zinc ~ 1, meuse, meuse.grid, x1, nmax = 30)
gridded(G1) <- TRUE
G1@data = as.data.frame(G1@data[,-2])

# GRID-2 log(elev):
v2 <- variogram(log(elev) ~ 1, meuse)
x2 <- fit.variogram(v2, vgm(.1, "Sph", 1000, .6))
G2 <- krige(elev ~ 1, meuse, meuse.grid, x2, nmax = 30)
gridded(G2) <- TRUE
G2@data <- as.data.frame(G2@data[,-2])
G2@data[,1] <- G2@data[,1]

corr <- raster.modified.ttest(G1, G2)
plot(raster::raster(corr,1))

corr.rand <- raster.modified.ttest(G1, G2, sub.sample = TRUE, type = "random")
corr.hex <- raster.modified.ttest(G1, G2, sub.sample = TRUE, d = 500, size = 1000)
head(corr.hex@data)
bubble(corr.hex, "corr")

## End(Not run)
```

raster.moments*Raster moments***Description**

Calculates focal statistical moments of a raster

Usage

```
raster.moments(x, type = "mean", s = 3, p = 0.75)
```

Arguments

x	raster object
type	The global statistic to represent the local deviation options are: "min", "min", "mean", "median", "var", "sd", "mad", "kurt", "skew", "quantile"
s	Size of matrix (focal window), can be single value or two values defining the [x,y] dimensions of the focal matrix
p	if type="quantile", the returned percentile.

Value

raster class object of the local distributional moment

Note

This is a simple wrapper for the focal function, returning local statistical moments

Author(s)

Jeffrey S. Evans <jeffrey_evans@tnc.org>

Examples

```
library(raster)
r <- raster(nrows=100, ncols=100, xmn=571823, xmx=616763,
            ymn=4423540, ymx=4453690)
proj4string(r) <- crs("+proj=utm +zone=12 +datum=NAD83 +units=m +no_defs")
r[] <- runif(ncell(r), 1000, 2500)
r <- focal(r, focalWeight(r, 150, "Gauss") )

# Calculate 10th percentile for 3x3 window
r.p10 <- raster.moments(r, type="quantile", p=0.10)
```

raster.transformation *Statistical transformation for rasters*

Description

Transforms raster to a specified statistical transformation

Usage

```
raster.transformation(x, trans = "norm", smin = 0, smax = 255)
```

Arguments

x	raster class object
trans	Transformation method: "norm", "rstd", "std", "stretch", "nl", "slog", "sr" (please see notes)
smin	Minimum value for stretch
smax	Maximum value for stretch

Value

raster class object of transformation

Note

("norm") Normalization [0-1]: if $\min(x) < 0$ ($x - \min(x)$) / ($\max(x) - \min(x)$) ("rstd") Row standardize [0-1]: if $\min(x) \geq 0$ $x / \max(x)$ This normalizes data with negative distributions ("std") Standardize: $(x - \text{mean}(x)) / \text{sd}(x)$ ("stretch") Stretch: $((x - \min(x)) * \text{max.stretch} / (\max(x) - \min(x)) + \min.stretch)$ This will stretch values to the specified minimum and maximum values (eg., 0-255 for 8-bit) ("nl") Natural logarithms: if $\min(x) > 0$ $\log(x)$ ("slog") Signed log 10 (for skewed data): if $\min(x) \geq 0$ $\text{ifelse}(\text{abs}(x) \leq 1, 0, \text{sign}(x) * \log10(\text{abs}(x)))$ ("sr") Square-root: if $\min(x) \geq 0$ \sqrt{x}

Author(s)

Jeffrey S. Evans <jeffrey_evans@tnc.org>

Examples

```
library(raster)
r <- raster(nrows=100, ncols=100, xmn=571823, xmx=616763,
            ymn=4423540, ymx=4453690)
r[] <- runif(ncell(r), 1000, 2500)

# Positive values so, can apply any transformation
for( i in c("norm", "rstd", "std", "stretch", "nl", "slog", "sr")) {
  print( raster.transformation(r, trans = i) )
}
```

```
# Negative values so, can't transform using "nl", "slog" or "sr"
r[] <- runif(ncell(r), -1, 1)
for( i in c("norm", "rstd", "std", "stretch", "nl", "slog", "sr")) {
  try( print( raster.transformation(r, trans = i) ) )
}
```

raster.vol*Raster Percent Volume***Description**

Calculates a percent volume on a raster or based on a systematic sample

Usage

```
raster.vol(x, p = 0.95, sample = FALSE, spct = 0.05)
```

Arguments

x	raster class object
p	percent raster-value volume
sample	base volume on systematic point sample (TRUE/FALSE)
spct	sample percent, if sample (TRUE)

Value

if sample (FALSE) binary raster object with 1 representing designated percent volume

if sample (TRUE) n sp SpatialPointsDataFrame object with points that represent the percent volume of the sub-sample

Note

Since this model needs to operate on all of the raster values, it is not memory safe

Author(s)

Jeffrey S. Evans <jeffrey_evans@tnc.org>

Examples

```

require(raster)
r <- raster(ncols=100, nrows=100)
r[] <- runif(ncell(r), 0, 1)
r <- focal(r, w=focalWeight(r, 6, "Gauss"))
r[sample(1000, 1:ncell(r))] <- NA

# full raster percent volume
p30 <- raster.vol(r, p=0.30)
p50 <- raster.vol(r, p=0.50)
p80 <- raster.vol(r, p=0.80)
par(mfrow=c(2,2))
plot(r, col=cm.colors(10), main="original raster")
plot(p30, breaks=c(0,0.1,1), col=c("cyan","red"), legend=FALSE,
     main="30% volume")
plot(p50, breaks=c(0,0.1,1), col=c("cyan","red"), legend=FALSE,
     main="50% volume")
plot(p80, breaks=c(0,0.1,1), col=c("cyan","red"), legend=FALSE,
     main="80% volume")

# point sample percent volume
# p30 <- raster.vol(r, p = 0.30, sample = TRUE, spct = 0.20)
#   plot(r, main="30% volume point sample")
#   plot(p30, pch=20, cex=0.70, add=TRUE)

```

raster.Zscore

Modified z-score for a raster

Description

Calculates the modified z-score for all cells in a raster

Usage

```
raster.Zscore(x, p.value = FALSE, file.name = NULL, ...)
```

Arguments

x	A raster class object
p.value	Return p-value rather than z-score raster (FALSE/TRUE)
file.name	Name of raster written to disk
...	Additional arguments passed to writeRaster

Value

raster class object or raster written to disk

Note

Since this functions needs to operate on all of the raster values, it is not memory safe

Author(s)

Jeffrey S. Evans <jeffrey_evans@tnc.org>

Examples

```
library(raster)
r <- raster(nrows=824, ncols=767, xmn=2451905, xmx=3218905,
            ymn=-2744771, ymx=-1920771, resolution = 5000)
r[] <- runif(ncell(r), 0, 1)

# Modified z-score
z <- raster.Zscore(r)

# P-value
p <- raster.Zscore(r, p.value = TRUE)
```

rasterCorrelation *Raster correlation*

Description

Performs a simple moving window correlation between two rasters

Usage

```
rasterCorrelation(x, y, s = 3, type = "pearson", file.name = NULL,
                  ...)
```

Arguments

x	raster class object for x
y	raster class object for y
s	Scale of window. Can be a single value, two values for uneven window or a custom matrix. Must be odd number (eg., s=3, for 3x3 window or s=c(3,5) for 3 x 5 window)
type	Type of output, options are: "pearson", "spearman", "covariance"
file.name	Name of output raster (optional)
...	Additional arguments passed to writeRaster

Value

raster class object or raster written to disk

Note

Depends: raster

Author(s)

Jeffrey S. Evans <jeffrey_evans@tnc.org>

Examples

```
library(raster)
b <- brick(system.file("external/rlogo.grd", package="raster"))
x <- b[[1]]
y <- b[[3]]
r.cor <- rasterCorrelation(x, y, s = 5, type = "spearman")
plot(r.cor)
```

remove.holes *Remove polygon holes*

Description

Removes all holes (null geometry) in polygon sp class objects

Usage

```
remove.holes(x)
```

Arguments

x SpatialPolygons or SpatialPolygonsDataFrame class object

Value

SpatialPolygonsDataFrame object with all holes removed

Note

A hole is considered a polygon within a polygon representing null geometry

Author(s)

Jeffrey S. Evans <jeffrey_evans@tnc.org>

Examples

```
library(sp)
Sr1 = Polygon(cbind(c(2,4,4,1,2),c(2,3,5,4,2)))
Sr2 = Polygon(cbind(c(5,4,2,5),c(2,3,2,2)))
Sr3 = Polygon(cbind(c(4,4,5,10,4),c(5,3,2,5,5)))
Sr4 = Polygon(cbind(c(5,6,6,5,5),c(4,4,3,3,4)), hole = TRUE)
polys <- SpatialPolygons(list(Polygons(list(Sr1), "s1"),
                               Polygons(list(Sr2), "s2"),
                               Polygons(list(Sr3, Sr4), "s3/4"))), 1:3)
par(mfrow=c(1,2))
plot(polys, col = 1:3, main="with hole")
plot(remove.holes(polys), col = 1:3, main="with hole removed")
```

sa.trans

Trigonometric transformation of a slope and aspect interaction

Description

The Trigonometric Stage (1978) [slope * cos(aspect)] or [slope * sin(aspect)]

Usage

```
sa.trans(slope, aspect, type = "cos", slp.units = "degrees",
         asp.units = "degrees")
```

Arguments

slope	slope values in degrees, radians or percent
aspect	aspect values in degrees or radians
type	Type of transformation, options are: "cos", "sin",
slp.units	Units of slope values, options are: "degrees", "radians" or "percent"
asp.units	Units of aspect values, options are: "degrees" or "radians"

Value

A vector of the modeled value

Note

An a priori assumption of a maximum in the NW quadrant (45 azimuth) and a minimum in the SW quadrant can be replaced by an empirically determined location of the optimum without repeated calculations of the regression fit. In addition it is argued that expressions for the effects of aspect should always be considered as terms involving an interaction with slope (Stage, 1976)

For slopes from 0 bounded from -1 to 1. Greater than 100 out of the -1 to 1 range.

An alternative for slopes with values approaching infinity is to take the square root of slope/100 to reduce the range of values. By default this model test all values greater than 100 to 101

Author(s)

Jeffrey S. Evans <jeffrey_evans@tnc.org>

References

Stage, A. R. 1976. An Expression of the Effects of Aspect, Slope, and Habitat Type on Tree Growth. Forest Science Vol 22, No 3, 457-460.

Examples

```
sa.trans(slope = 48.146, aspect = 360.000)

library(raster)
data(elev)
sa <- raster::terrain(elev, opt=c("slope", "aspect"), unit="degrees")
scosa <- raster::overlay(sa[[1]], sa[[2]], fun = sa.trans)
```

sample.annulus

Sample annulus

Description

Creates sample points based on annulus with defined inner and outer radius

Usage

```
sample.annulus(x, r1, r2, n = 10, ...)
```

Arguments

x	sp SpatialPoints or SpatialPointsDataFrame class object
r1	Numeric value defining inner radius of annulus (in projection units)
r2	Numeric value defining outer radius of annulus (in projection units)
n	Number of samples
...	Additional arguments passed to spsample

Value

sp SpatialPointsDataFrame OBJECT

Note

Function can be used for distance based sampling. This is one sampling method to capture the spatially lagged variation.

Author(s)

Jeffrey S. Evans <jeffrey_evans@tnc.org>

Examples

```
library(sp)
library(rgeos)
data(meuse)
coordinates(meuse) <- ~x+y
proj4string(meuse) <- CRS("+init=epsg:28992")
xy <- meuse[2,]

rs100 <- sample.annulus(xy, r1=50, r2=100, n = 50, type = "random")
rs200 <- sample.annulus(xy, r1=100, r2=200, n = 50, type = "random")

plot(rs200, pch=20, col="red")
points(rs100, pch=20, col="blue")
points(xy, pch=20, cex=2, col="black")
box()
legend("topright", legend=c("50-100m", "100-200m", "source"),
       pch=c(20,20,20), col=c("blue","red","black"))
```

sample.line

Systematic or random point sample of line(s)

Description

Creates a systematic or random point sample of an sp SpatialLinesDataFrame object based on distance spacing, fixed size or proportional size

Usage

```
sample.line(x, d = 100, p = NULL, n = NULL, type = "regular",
            longlat = FALSE, min.samp = 1, ...)
```

Arguments

x	sp class SpatialLinesDataFrame object
d	Sample distance. For regular sample.
p	Proportional sample size (length * p), expected value is 0-1. For regular or random.
n	Fixed sample size. For regular or random
type	Defines sample type. Options are "regular" or "random". A regular sample results in a systematic, evenly spaced sample.
longlat	TRUE/FALSE is data in geographic units, if TRUE distance is in kilometres
min.samp	Minimal number of sample points for a given line (default is 1 point)
...	Additional argument passed to spsample

Value

sp SpatialPointsDataFrame object.

Note

The sdist argument will produce an evenly spaced sample, whereas n produces a fixed sized sample. The p (proportional) argument calculates the percent of the line-length.

The LID column in the @data slot corresponds to the row.names of the SpatialLinesDataFrame object.

Depends: sp

Author(s)

Jeffrey S. Evans <jeffrey_evans@tnc.org>

Examples

```
require(sp)
sp.lines <- SpatialLines(list(Lines(list(Line(cbind(c(1,2,3),c(3,2,2)))), ID="1"),
                                Lines(list(Line(cbind(c(1,2,3),c(1,1.5,1))))), ID="2")))
sp.lines <- SpatialLinesDataFrame( sp.lines, data.frame(ID=1:2, row.names=c(1,2)) )

par(mfrow=c(2,2))
# Create systematic sample at 20 km spacing
reg.sample <- sample.line(sp.lines, d = 20, type = "regular", longlat = TRUE)
plot(sp.lines)
plot(reg.sample, pch = 20, add = TRUE)
box()
title("systematic d = 20")

# Create fixed size (n = 20) systematic sample
reg.sample <- sample.line(sp.lines, n = 20, type = "regular", longlat = TRUE)
plot(sp.lines)
plot(reg.sample, pch = 20, add = TRUE)
box()
title("systematic n = 20")

# Create fixed size (n = 20) random sample
rand.sample <- sample.line(sp.lines, n = 20, type = "random", longlat = TRUE)
plot(sp.lines)
plot(rand.sample, pch = 20, add = TRUE)
box()
title("rand n = 20")

# Create proportional (p = 0.10) random sample
rand.sample <- sample.line(sp.lines, p = 0.10, type = "random", longlat = TRUE)
plot(sp.lines)
plot(rand.sample, pch = 20, add = TRUE)
box()
title("rand p = 0.10")
```

sample.poly*Sample Polygons***Description**

Creates an equal sample of n for each polygon in an sp Polygon class object

Usage

```
sample.poly(x, n = 10, type = "random", ...)
```

Arguments

x	sp class SpatialPolygons or SpatialPolygonsDataFrame object
n	Number of random samples
type	Type of sample with options for: "random", "regular", "stratified", "nonaligned", "hexagonal", "clustered", "Fibonacci". See "spsample" for details.
...	Additional arguments passed to spsample

Value

sp SpatialPointsDataFrame object

Note

Depends: sp

Author(s)

Jeffrey S. Evans <jeffrey_evans@tnc.org>

Examples

```
library(raster)
library(sp)
p <- raster(nrow=10, ncol=10)
p[] <- runif(ncell(p)) * 10
p <- rasterToPolygons(p, fun=function(x){x > 9})
s <- sample.poly(p, n = 5, type = "random")
plot(p)
plot(s, pch = 20, add = TRUE)
box()
title("Random sample (n=5) for each polygon")
```

sampleTransect	<i>Sample transect</i>
----------------	------------------------

Description

Creates random transects from points and generates sample points along each transect

Usage

```
sampleTransect(x, min.length, max.length, id = NULL, ...)
```

Arguments

x	A sp point object
min.length	Minimum length of transect(s)
max.length	Maximum length of transect(s)
id	A unique identification column in x
...	Additional arguments passed to sample.line

Note

Function create random direction and length transects and then creates a point sample along each transect. The characteristic of the sample points are defined by arguments passed to the sample.line function

Author(s)

Jeffrey S. Evans <jeffrey_evans@tnc.org>

Examples

```
library(sp)
data(meuse)
coordinates(meuse) <- ~x+y
proj4string(meuse) <- CRS("+init=epsg:28992")
meuse <- meuse[sample(1:nrow(meuse),10),]

transects <- sampleTransect(meuse, min.length=200, max.length=500, min.samp = 3)
plot(transects$transects)
plot(transects$samples, pch=20, add=TRUE)
```

sar*Surface Area Ratio*

Description

Calculates the Berry (2002) Surface Area Ratio based on slope

Usage

```
sar(x, s = NULL, ...)
```

Arguments

x	raster object
s	cell resolution (default is NULL, not needed if projection is in planar units)
...	Additional arguments passed to raster::calc

Value

raster class object of Berry (2002) Surface Area Ratio

Note

SAR is calculated as: $\text{resolution}^2 * \cos(\text{degrees}(\text{slope}) * (\pi / 180))$

Author(s)

Jeffrey S. Evans <jeffrey_evans@tnc.org>

References

Berry, J.K. (2002). Use surface area for realistic calculations. Geoworld 15(9):20-1.

Examples

```
library(raster)
data(elev)
surface.ratio <- sar(elev, s=90)
plot(surface.ratio)
```

<i>separability</i>	<i>separability</i>
---------------------	---------------------

Description

Calculates variety of two-class sample separability metrics

Usage

```
separability(x, y, plot = FALSE, cols = c("red", "blue"),
             clabs = c("Class1", "Class2"), ...)
```

Arguments

x	X vector
y	Y vector
plot	plot separability (TRUE/FALSE)
cols	colors for plot (must be equal to number of classes)
clabs	labels for two classes
...	additional arguments passes to plot

Value

A data.frame with the following separability metrics:

- B Bhattacharyya distance statistic
- JM Jeffries-Matusita distance statistic
- M M-Statistic
- D Divergence index
- TD Transformed Divergence index

Note

M-Statistic (Kaufman & Remer 1994) - This is a measure of the difference of the distributional peaks. A large M-statistic indicates good separation between the two classes as within-class variance is minimized and between-class variance maximized (M <1 poor, M >1 good).

Bhattacharyya distance (Bhattacharyya 1943; Harold 2003) - Measures the similarity of two discrete or continuous probability distributions.

Jeffries-Matusita distance (Bruzzone et al., 2005; Swain et al., 1971) - The J-M distance is a function of separability that directly relates to the probability of how good a resultant classification will be. The J-M distance is asymptotic to v2, where values of v2 suggest complete separability

Divergence and transformed Divergence (Du et al., 2004) - Maximum likelihood approach. Transformed divergence gives an exponentially decreasing weight to increasing distances between the classes.

Author(s)

Jeffrey S. Evans <jeffrey_evans<at>tnc.org>

References

- Anderson, M. J., & Clements, A. (2000) Resolving environmental disputes: a statistical method for choosing among competing cluster models. *Ecological Applications* 10(5):1341-1355
- Bhattacharyya, A. (1943) On a measure of divergence between two statistical populations defined by their probability distributions'. *Bulletin of the Calcutta Mathematical Society* 35:99-109
- Bruzzone, L., F. Roli, S.B. Serpico (1995) An extension to multiclass cases of the Jefferys-Matusita distance. *IEEE Transactions on Pattern Analysis and Machine Intelligence* 33:1318-1321
- Du, H., C.I. Chang, H. Ren, F.M. D'Amico, J. O. Jensen, J., (2004) New Hyperspectral Discrimination Measure for Spectral Characterization. *Optical Engineering* 43(8):1777-1786.
- Kailath, T., (1967) The Divergence and Bhattacharyya measures in signal selection. *IEEE Transactions on Communication Theory* 15:52-60
- Kaufman Y., and L. Remer (1994) Detection of forests using mid-IR reflectance: An application for aerosol studies. *IEEE T. Geosci.Remote.* 32(3):672-683.

Examples

```
norm1 <- dnorm(seq(-20,20,length=5000),mean=0,sd=1)
norm2 <- dnorm(seq(-20,20,length=5000),mean=0.2,sd=2)
separability(norm1, norm2)

s1 <- c (1362,1411,1457,1735,1621,1621,1791,1863,1863,1838)
s2 <- c (1362,1411,1457,10030,1621,1621,1791,1863,1863,1838)
separability(s1, s2, plot=TRUE)
```

sg.smooth

Savitzky-Golay smoothing filter

Description

Smoothing of time-series data using Savitzky-Golay convolution smoothing

Usage

```
sg.smooth(x, f = 4, l = 51, d = 1, na.rm, ...)
```

Arguments

x	A vector to be smoothed
f	Filter type (default 4 for quartic, specify 2 for quadratic)
l	Convolution filter length, must be odd number (default 51). Defines degree of smoothing
d	First derivative (default 1)
na.rm	NA behavior
...	not used

Value

A vector of the smoothed data equal to length of x. Please note; NA values are retained

Author(s)

Jeffrey S. Evans <jeffrey_evans@tnc.org>

References

Savitzky, A., & Golay, M.J.E. (1964). Smoothing and Differentiation of Data by Simplified Least Squares Procedures. Analytical Chemistry. 36(8):1627-39

Examples

```
y <- c(0.112220988, 0.055554941, 0.013333187, 0.055554941, 0.063332640, 0.014444285,
0.015555384, 0.057777140, 0.059999339, 0.034444068, 0.058888242, 0.136665165,
0.038888458, 0.096665606, 0.141109571, 0.015555384, 0.012222088, 0.012222088,
0.072221428, 0.052221648, 0.087776810, 0.014444285, 0.033332966, 0.012222088,
0.032221869, 0.059999339, 0.011110989, 0.011110989, 0.042221759, 0.029999670,
0.018888680, 0.098887801, 0.016666483, 0.031110767, 0.061110441, 0.022221979,
0.073332526, 0.012222088, 0.016666483, 0.012222088, 0.122220881, 0.134442955,
0.094443403, 0.128887475, 0.045555055, 0.152220547, 0.071110331, 0.018888680,
0.022221979, 0.029999670, 0.035555165, 0.014444285, 0.049999449, 0.074443623,
0.068888135, 0.062221535, 0.032221869, 0.095554501, 0.143331751, 0.121109776,
0.065554835, 0.074443623, 0.043332856, 0.017777583, 0.016666483, 0.036666263,
0.152220547, 0.032221869, 0.009999890, 0.009999890, 0.021110879, 0.025555275,
0.099998899, 0.015555384, 0.086665712, 0.008888791, 0.062221535, 0.044443958,
0.081110224, 0.015555384, 0.089999005, 0.082221314, 0.056666043, 0.013333187,
0.048888352, 0.075554721, 0.025555275, 0.056666043, 0.1466665052, 0.118887581,
0.125554174, 0.024444176, 0.1244443069, 0.012222088, 0.1266665279, 0.048888352,
0.046666153, 0.141109571, 0.015555384, 0.114443190)

plot(y, type="l", lty = 3, main="Savitzky-Golay with l = 51, 25, 10")
lines(sg.smooth(y), col="red", lwd=2)
lines(sg.smooth(y, l = 25), col="blue", lwd=2)
lines(sg.smooth(y, l = 10), col="green", lwd=2)
```

shannons

Shannon's Diversity (Entropy) Index

Description

Calculates Shannon's Diversity Index and Shannon's Evenness Index

Usage

```
shannons(x, counts = TRUE, ens = FALSE, margin = "row")
```

Arguments

x	data.frame object containing counts or proportions
counts	Are data counts (TRUE) or relative proportions (FALSE)
ens	Calculate effective number of species (TRUE/FALSE)
margin	Calculate diversity for rows or columns. c("row", "col")

Value

data.frame with "H" (Shannon's diversity) and "evenness" (Shannon's evenness where H / max(sum(x))) and ESN

Note

The expected for H is 0-3+ where a value of 2 has been suggested as medium-high diversity, for evenness is 0-1 with 0 signifying no evenness and 1, complete evenness.

Author(s)

Jeffrey S. Evans <jeffrey_evans@tnc.org>

References

- Shannon, C. E. and W. Weaver (1948) A mathematical theory of communication. *The Bell System Technical Journal*, 27:379-423.
- Simpson, E. H. (1949) Measurement of diversity. *Nature* 163:688
- Roth, D. S., I. Perfecto, and B. Rathcke (1994) The effects of management systems on ground-foraging ant diversity in Costa Rica. *Ecological Applications* 4(3):423-436.

Examples

```
# Using Costa Rican ant diversity data from Roth et al. (1994)
data(ants)

# Calculate diversity for each covertype ("col")
shannons(ants[,2:ncol(ants)], ens = TRUE, counts = FALSE, margin = "col")

# Calculate diversity for each species ("row")
ant.div <- shannons(ants[,2:ncol(ants)], ens = TRUE, counts = FALSE, margin = "row")
row.names(ant.div) <- ants[,1]
ant.div
```

similarity	<i>Ecological similarity</i>
------------	------------------------------

Description

Uses row imputation to identify "k" ecological similar observations

Usage

```
similarity(x, k = 4, method = "mahalanobis", frequency = TRUE,
           scale = TRUE, ID = NULL)
```

Arguments

- x data.frame containing ecological measures
- k Number of k nearest neighbors (kNN)
- method Method to compute multivariate distances c("mahalanobis", "raw", "euclidean", "ica")
- frequency Calculate frequency of each reference row (TRUE/FALSE)
- scale Scale multivariate distances to standard range (TRUE/FALSE)
- ID Unique ID vector to use as reference ID's (rownames). Must be unique and same length as number of rows in x

Value

data.frame with k similar targets and associated distances. If frequency = TRUE the freq column represents the number of times a row (ID) was selected as a neighbor.

Note

This function uses row-based imputation to identify k similar neighbors for each observation. Has been used to identify offsets based on ecological similarity.

Author(s)

Jeffrey S. Evans <jeffrey_evans@tnc.org>

References

Evans, J.S., S.R. Schill, G.T. Raber (2015) A Systematic Framework for Spatial Conservation Planning and Ecological Priority Design in St. Lucia, Eastern Caribbean. Chapter 26 in Central American Biodiversity : Conservation, Ecology and a Sustainable Future. F. Huettman (eds). Springer, NY.

Examples

```

data(pu)
kNN <- similarity(pu@data[2:ncol(pu)], k = 4, frequency = FALSE, ID = pu@data$UNIT_ID)

## Not run:
kNN <- similarity(pu@data[2:ncol(pu)], k = 4, frequency = TRUE, ID = pu@data$UNIT_ID)
p <- kNN$freq
clr <- c("#3288BD", "#99D594", "#E6F598", "#FEE08B",
          "#FC8D59", "#D53E4F")
p <- ifelse(p <= 0, clr[1],
            ifelse(p > 0 & p < 10, clr[2],
                  ifelse(p >= 10 & p < 20, clr[3],
                        ifelse(p >= 20 & p < 50, clr[4],
                              ifelse(p >= 50 & p < 100, clr[5],
                                ifelse(p >= 100, clr[6], NA))))))
plot(pu, col=p, border=NA)
legend("topleft", legend=c("None", "<10", "10-20",
                           "20-50", "50-100", ">100"),
       fill=clr, cex=0.6, bty="n")
box()
## End(Not run)

```

Description

Smooths pixel-level data in raster time-series and can impute missing (NA) values.

Usage

```
smooth.time.series(x, f = 0.8, smooth.data = FALSE, ...)
```

Arguments

x	A raster stack or brick class object
f	Smoothing parameter (see loess span argument)
smooth.data	(FALSE/TRUE) Smooth all of the data or just impute NA values
...	Additional arguments passed to raster calc (for writing results to disk)

Details

This function uses a LOESS regression to smooth the time-series (using the smooth.data = TRUE argument). If the data is smoothed, it will be replaced by a loess estimate of the time-series (estimated distribution at the pixel-level). The results can dramatically be effected by the choice of the smoothing parameter (f) so caution is warranted and the effect of this parameter tested. Alternately, with smooth.data = FALSE, the function can be used to impute missing pixel data (NA) in raster time-series (stacks/bricks).

Value

A raster stack or brick object with imputed NA values or smoothed data.

Author(s)

Jeffrey S. Evans <jeffrey_evans@tnc.org>

See Also

[loess](#) for details on the loess regression

[calc](#) for details on additional (...) arguments

Examples

```
## Not run:  
# fill NA values  
lai.new <- smooth.time.series(lai)  
  
# Smooth time-series with a relaxed smoothing parameter  
lai.smooth <- smooth.time.series(lai, f = 0.2, smooth.data = TRUE)  
  
## End(Not run)
```

sobel*Sobel-Feldman operator***Description**

An isotropic image gradient operator using a 3x3 window

Usage

```
sobel(x, method = "intensity", ...)
```

Arguments

<code>x</code>	A raster class object
<code>method</code>	Type of operator ("intensity", "direction", "edge")
<code>...</code>	Additional arguments passed to raster::overlay or, if method="edge", raster::focal (if you want a file written to disk use filename = "" argument)

Value

A raster class object or raster written to disk

Note

The Sobel-Feldman operator is a discrete differentiation operator, deriving an approximation of the gradient of the intensity function. abrupt discontinuity in the gradient function represents edges, making this a common approach for edge detection. The Sobel-Feldman operator is based on convolving the image with a small, separable, and integer matrix in the horizontal and vertical directions. The operator uses two 3x3 kernels which are convolved with the original image to calculate approximations of the derivatives - one for horizontal changes, and one for vertical. Where `x` is defined here as increasing in the right-direction, and `y` as increasing in the down-direction. At each pixel in the raster, the resulting gradient can be combined to give the gradient intensity, using: $\text{SQRT}(G_x^2 + G_y^2)$. This can be expanded into the gradient direction using $\text{atan}(G_x/G_y)$

Author(s)

Jeffrey S. Evans <jeffrey_evans@tnc.org>

References

Sobel, I., & G. Feldman, (1969) A 3x3 Isotropic Gradient Operator for Image Processing, presented at the Stanford Artificial Intelligence Project (SAIL).

Examples

```
library(raster)
r <- brick(system.file("external/rlogo.grd", package="raster"))
s.int <- sobal(r[[1]])
s.dir <- sobal(r[[1]], method = "direction")
s.edge <- sobal(r[[1]], method = "edge")
par(mfrow=c(2,2))
plot(r[[1]])
plot(s.int, main="intensity")
plot(s.dir, main="direction")
plot(s.edge, main="edge")
```

sp.kde

Spatial kernel density estimate

Description

A weighted or unweighted Gaussian Kernel Density estimate for spatial data

Usage

```
sp.kde(x, y, bw, newdata, n, standardize = FALSE, scale.factor)
```

Arguments

x	sp SpatialPointsDataFrame object
y	Optional values, associated with x coordinates, to be used as weights
bw	Distance bandwidth of Gaussian Kernel, must be units of projection
newdata	A Rasterlayer, any sp class object or c[xmin,xmax,ymin,ymax] vector to estimate the kde extent
n	Number of cells used in creating grid. If not defined a value based on extent or existing raster will be used
standardize	Standardize results to 0-1 (FALSE/TRUE)
scale.factor	Optional numeric scaling factor for the KDE (eg., 10000), to account for small estimate values

Value

Raster class object containing kernel density estimate

Author(s)

Jeffrey S. Evans <jeffrey_evans@tnc.org>

Examples

```

## Not run:
library(sp)
data(meuse)
coordinates(meuse) <- ~x+y

# Weighted KDE using cadmium and spatial locations
cadmium.kde <- sp.kde(x = meuse, y = meuse$cadmium, bw = 1000, n = 5000,
                      standardize = TRUE, scale.factor = 10000 )

# Unweighted KDE (spatial locations only)
pt.kde <- sp.kde(x = meuse, bw = 1000, standardize = TRUE, n = 5000,
                   scale.factor = 10000 )

# Plot results
par(mfrow=c(1,2))
plot(cadmium.kde, main="weighted kde")
points(meuse, pch=20, col="red")
plot(pt.kde, main="Unweighted kde")
points(meuse, pch=20, col="red")

# Using existing raster
library(raster)
data(meuse.grid)
coordinates(meuse.grid) = ~x+y
proj4string(meuse.grid) <- CRS("+init=epsg:28992")
gridded(meuse.grid) = TRUE
meuse.grid <- raster(meuse.grid)

cadmium.kde <- sp.kde(x = meuse, y = meuse$cadmium, newdata = meuse.grid, bw = 1000,
                      standardize = TRUE, scale.factor = 10000 )
plot(cadmium.kde, main="weighted kde")
points(meuse, pch=20, cex=0.5, col="red")

## End(Not run)

```

Description

Removes row or column NA's in sp object

Usage

```
sp.na.omit(x, col.name = NULL, margin = 1)
```

Arguments

x	Object of class SpatialPointsDataFrame OR SpatialPolygonsDataFrame
col.name	The name of a specific column to remove NA's from
margin	Margin (1,2) of data.frame 1 for rows or 2 for columns

Note

This function will remove all NA's in the object or NA's associated with a specific column.

Author(s)

Jeffrey S. Evans <jeffrey_evans@tnc.org>

Examples

```
library(sp)
data(meuse)
coordinates(meuse) <- ~x+y

# Display rows with NA
meuse@data[!complete.cases(meuse@data),]

# Remove all NA's in rows (and associated points)
meuse2 <- sp.na.omit(meuse)
dim(meuse)
dim(meuse2)

# Plot deleted points in red
plot(meuse, col='red', pch=20)
plot(meuse2, col='black', pch=20, add=TRUE)

# Remove NA's associated with specific column
meuse2 <- sp.na.omit(meuse, col.name = "om")
head(meuse@data)
head(meuse2@data)
```

Description

Derives the spherical standard deviation of a raster surface

Usage

```
spherical.sd(r, d, variance = FALSE, ...)
```

Arguments

r	Raster class object
d	Size of focal window or a matrix to use in focal function
variance	(FALSE TRUE) Output spherical variance rather than standard deviation
...	Additional arguments passed to calc (can write raster to disk here)

Details

Surface variability using spherical variance/standard deviation. The variation can be assessed using the spherical standard deviation of the normal direction within a local neighborhood. This is found by expressing the normal directions on the surfaces cells in terms of their displacements in a Cartesian (x,y,z) coordinate system. Averaging the x-coordinates, y-coordinates, and z-coordinates separately gives a vector (xb, yb, zb) pointing in the direction of the average normal. This vector will be shorter when there is more variation of the normals and it will be longest–equal to unity–when there is no variation. Its squared length is (by the Pythagorean theorem) given by: $R^2 = xb^2 + yb^2 + zb^2$ where; $x = \cos(\text{aspect}) * \sin(\text{slope})$ and $xb = nX_n$ focal mean of x $y = \sin(\text{aspect}) * \sin(\text{slope})$ and $yb = nY_n$ focal mean of y $z = \cos(\text{slope})$ and $zb = nZ_n$ focal mean of z

The slope and aspect values are expected to be in radians. The value of $(1 - R^2)$, which will lie between 0 and 1, is the spherical variance. and its square root can be considered the spherical standard deviation.

Value

rasterLayer class object of the spherical standard deviation

Author(s)

Jeffrey S. Evans <jeffrey_evans<at>tnc.org>

See Also

[focal](#) for details on focal function

[calc](#) for details on ... arguments

Examples

```
## Not run:
library(raster)
data(elev)

ssd <- spherical.sd(elev, d=5)

slope <- terrain(elev, opt='slope')
aspect <- terrain(elev, opt='aspect')
hill <- hillShade(slope, aspect, 40, 270)
plot(hill, col=grey(0:100/100), legend=FALSE,
     main='terrain spherical standard deviation')
plot(ssd, col=rainbow(25, alpha=0.35), add=TRUE)
```

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

srr*Surface Relief Ratio*

Description

Calculates the Pike (1971) Surface Relief Ratio

Usage

```
srr(x, s = 5, ...)
```

Arguments

x	raster object
s	Focal window size
...	Additional arguments passed to raster::calc

Value

raster class object of Pike's (1971) Surface Relief Ratio

Note

Describes rugosity in continuous raster surface within a specified window. The implementation of SRR can be shown as: $(\text{mean}(x) - \text{min}(x)) / (\text{max}(x) - \text{min}(x))$

Author(s)

Jeffrey S. Evans <jeffrey_evans@tnc.org>

Examples

```
library(raster)
data(elev)
r.srr <- srr(elev, s=5)
plot(r.srr, main="Surface Relief Ratio")
```

stratified.random *Stratified random sample*

Description

Creates a stratified random sample of an sp class object

Usage

```
stratified.random(x, strata, n = 10, reps = 1, replace = TRUE)
```

Arguments

x	sp class SpatialDataFrame object (point, polygon, line, pixel)
strata	Column in @data slot with stratification factor
n	Number of random samples
reps	Number of replicates per strata
replace	Sampling with replacement (TRUE/FALSE)

Value

sp SpatialDataFrame object (same as input feature) containing random samples

Note

If replace=FALSE features are removed from consideration in subsequent replicates. Conversely, if replace=TRUE, a feature can be selected multiple times across replicates. Not applicable if rep=1.

Depends: sp

Author(s)

Jeffrey S. Evans <jeffrey_evans@tnc.org>

References

Hudak, A.T., N.L. Crookston, J.S. Evans, M.J. Falkowski, A.M.S. Smith, P. Gessler and P. Morgan. (2006) Regression modelling and mapping of coniferous forest basal area and tree density from discrete-return lidar and multispectral satellite data. Canadian Journal of Remote Sensing 32: 126-138.

Examples

```

require(sp)
data(meuse)
coordinates(meuse) <- ~x+y

# Create stratified variable using quartile breaks
x1 <- cut(meuse@data[, 'cadmium'], summary(meuse@data[, 'cadmium'])[-4], include.lowest=TRUE)
levels(x1) <- seq(1,nlevels(x1),1)
x2 <- cut(meuse@data[, 'lead'], summary(meuse@data[, 'lead'])[-4], include.lowest=TRUE)
levels(x2) <- seq(1,nlevels(x2),1)
meuse@data <- cbind(meuse@data, STRAT=paste(x1, x2, sep='.'))

# 2 replicates and replacement
ssample <- stratified.random(meuse, strata='STRAT', n=2, reps=2)

# 2 replicates and no replacement
ssample.nr <- stratified.random(meuse, strata='STRAT', n=2, reps=2, replace=FALSE)

# n=1 and reps=10 for sequential numbering of samples
ssample.ct <- stratified.random(meuse, strata='STRAT', n=1, reps=10, replace=TRUE)

# Counts for each full strata (note; 2 strata have only 1 observations)
tapply(meuse@data$STRAT, meuse@data$STRAT, length)

# Counts for each sampled strata, with replacement
tapply(ssample@data$STRAT, ssample@data$STRAT, length)

# Counts for each sampled strata, without replacement
tapply(ssample.nr@data$STRAT, ssample.nr@data$STRAT, length)

# Counts for each sampled strata, without replacement
tapply(ssample.ct@data$STRAT, ssample.ct@data$STRAT, length)

# Plot random samples colored by replacement
ssample@data$REP <- factor(ssample@data$REP)
spplot(ssample, 'REP', col.regions=c('red','blue'))

```

subsample.distance *Subsample distances*

Description

Minimum, and optional maximum, distance constrained sub-sampling

Usage

```
subsample.distance(x, n, d, d.max = NULL, replacement = FALSE,
   latlong = FALSE, trace = FALSE)
```

Arguments

x	A spatial polygons or points sp object
n	Number of random samples drawn from x
d	Minimum sample distance
d.max	Maximum sample distance (not used unless specified)
replacement	(FALSE/TRUE) sample with or without replacement
latlong	(FALSE/TRUE) is the data in a geographic projection (latitude/longitude)
trace	(FALSE/TRUE) Print min and max sample distances

Value

A sp spatial object, of the same class as x containing the random sub-samples

Note

This function provides a distance constrained subsample of existing point or polygon data

Author(s)

Jeffrey S. Evans <jeffrey_evans@tnc.org>

Examples

```

library(sp)
data(meuse)
coordinates(meuse) <- ~ x+y

# Subsample with a 500m minimum sample spread
sub.meuse <- subsample.distance(meuse, n = 10, d = 500, trace = TRUE)
plot(meuse,pch=19, main="min dist = 500")
points(sub.meuse, pch=19, col="red")

# Check distances
dm <- spDists(sub.meuse)
diag(dm) <- NA
cat("\n", "Min distance for subsample", min(dm, na.rm=TRUE), "\n")
cat("Max distance for subsample", max(dm, na.rm=TRUE), "\n")

## Not run:
# Subsample with a 500m minimum and 3500m maximum sample spread
sub.meuse <- subsample.distance(meuse, n = 10, d = 500, d.max = 3500)
plot(meuse,pch=19, main="min dist = 500, max dist = 3500")
points(sub.meuse, pch=19, col="red")

# Check distances
dm <- spDists(sub.meuse)
diag(dm) <- NA
cat("Min distance for subsample", min(dm, na.rm=TRUE), "\n")
cat("Max distance for subsample", max(dm, na.rm=TRUE), "\n")

```

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

summary.cross.cor *Summary of spatial cross correlation*

Description

summary method for class "cross.cor"

Usage

```
## S3 method for class 'cross.cor'  
summary(object, ...)
```

Arguments

object	Object of class cross.cor
...	Ignored

summary.effect.size *Summarizing effect size*

Description

Summary method for class "effect.size".

Usage

```
## S3 method for class 'effect.size'  
summary(object, ...)
```

Arguments

object	Object of class effect.size
...	Ignored

summary.loess.boot *Summarizing Loess bootstrap models*

Description

Summary method for class "loess.boot".

Usage

```
## S3 method for class 'loess.boot'
summary(object, ...)
```

Arguments

object	Object of class loess.boot
...	Ignored

swvi *Senescence weighted Vegetation Index (swvi)*

Description

Modified Soil-adjusted Vegetation Index (MSAVI) or Modified Triangular Vegetation Index 2 (MTVI) weighted by the Normalized difference senescent vegetation index (NDSVI)

Usage

```
swvi(red, nir, swir, green = NULL, mtvi = FALSE, senescence = 0,
threshold = NULL, weight.factor = NULL, ...)
```

Arguments

red	Red band (0.636 - 0.673mm), landsat 5&7 band 3, OLI (landsat 8) band 4
nir	Near infrared band (0.851 - 0.879mm) landsat 5&7 band 4, OLI (landsat 8) band 5
swir	short-wave infrared band 1 (1.566 - 1.651mm), landsat 5&7 band 5, OLI (landsat 8) band 6
green	Green band if MTVI = TRUE
mtvi	(FALSE TRUE) Use Modified Triangular Vegetation Index 2 instead of MSAVI
senescence	The critical value, in NDSVI, representing senescent vegetation
threshold	Threshold value for defining NA based on < p
weight.factor	Apply partial weights [w * weight.factor] to the NDSVI weights
...	Additional arguments passed to raster calc function

Value

rasterLayer class object of the weighted MSAVI metric

Note

The intent of this index is to correct the MSAVI or MTVI index for bias associated with senescent vegetation. This is done by: 1) deriving the NDSVI; 2) applying a threshold to limit NDSVI to values associated with senescent vegetation; 3) converting the index to inverted weights [-1*(NDSVI/sum(NDSVI))]; 4) applying weights to MSAVI or MTVI

The MSAVI formula follows the modification proposed by Qi et al. (1994), often referred to as MSAVI2. The MSAVI index reduces soil noise and increases the dynamic range of the vegetation signal. The implemented modified version (MSAVI2) is based on an inductive method that does not use a constant L value, in separating soil effects, an highlights healthy vegetation. The MTVI(2) index follows Haboudane et al., (2004) and represents the area of a hypothetical triangle in spectral space that connects (1) green peak reflectance, (2) minimum chlorophyll absorption, and (3) the NIR shoulder. When chlorophyll absorption causes a decrease of red reflectance, and leaf tissue abundance causes an increase in NIR reflectance, the total area of the triangle increases. It is good for estimating green LAI, but its sensitivity to chlorophyll increases with an increase in canopy density. The modified version of the index accounts for the background signature of soils while preserving sensitivity to LAI and resistance to the influence of chlorophyll.

The Normalized difference senescent vegetation index (NDSVI) follows methods from Qi et al., (2000). The senescence is used to threshold the NDSVI. Values less than this value will be NA. The threshold argument is used to apply a threshold to MSAVI. The default is NULL but if specified all values [MSAVI <= threshold] will be NA. Applying a weight.factor can be used to change the influence of the weights on MSAVI.

Author(s)

Jeffrey S. Evans <jeffrey_evans@tnc.org>

References

- Haboudane, D., et al. (2004) Hyperspectral Vegetation Indices and Novel Algorithms for Predicting Green LAI of Crop Canopies: Modeling and Validation in the Context of Precision Agriculture. *Remote Sensing of Environment* 90:337-352.
- Qi J., Chehbouni A., Huete A.R., Kerr Y.H., (1994). Modified Soil Adjusted Vegetation Index (MSAVI). *Remote Sens Environ* 48:119-126.
- Qi J., Kerr Y., Chehbouni A., (1994). External factor consideration in vegetation index development. *Proc. of Physical Measurements and Signatures in Remote Sensing, ISPRS*, 723-730.
- Qi, J., Marssett, R., Moran, M.S., Goodrich, D.C., Heilman, P., Kerr, Y.H., Dedieu, G., Chehbouni, A., Zhang, X.X. (2000). Spatial and temporal dynamics of vegetation in the San Pedro River basin area. *Agricultural and Forest Meteorology*. 105:55-68.

Examples

```
## Not run:  
library(raster)
```

```

library(RStoolbox)

data(lsat)
lsat <- radCor(lsat, metaData = readMeta(system.file(
    "external/landsat/LT52240631988227CUB02_MTL.txt",
    package="RStoolbox")), method = "apref")

# Using Modified Soil-adjusted Vegetation Index (MSAVI)
( wmsavi <- swvi(red = lsat[[3]], nir = lsat[[4]], swir = lsat[[5]]) )
    plotRGB(lsat, r=6,g=5,b=2, scale=1, stretch="lin")
    plot(wmsavi, legend=FALSE, col=rev(terrain.colors(100, alpha=0.35)), add=TRUE )

# Using Modified Triangular Vegetation Index 2 (MTVI)
( wmtvi <- swvi(red = lsat[[3]], nir = lsat[[4]], swir = lsat[[5]],
    green = lsat[[3]], mtvi = TRUE) )
plotRGB(lsat, r=6,g=5,b=2, scale=1, stretch="lin")
plot(wmtvi, legend=FALSE, col=rev(terrain.colors(100, alpha=0.35)), add=TRUE )

## End(Not run)

```

tpi*Topographic Position Index (tpi)***Description**

Calculates topographic position using mean deviations

Usage

```
tpi(x, scale = 3, win = "rectangle", normalize = FALSE)
```

Arguments

x	A raster class object
scale	The focal window size
win	Window type. Options are "rectangle" and "circle"
normalize	Apply deviation correction that normalizes to local surface roughness

Value

raster class object of tpi

Author(s)

Jeffrey S. Evans <jeffrey_evans@tnc.org>

References

De Reu, J., J. Bourgeois, M. Bats, A. Zwervvaegher, V. Gelorini, et al., (2014) Application of the topographic position index to heterogeneous landscapes. *Geomorphology*, 186:39-49.

Examples

```
library(raster)
data(elev)

# calculate tpi and plot
tpi9 <- tpi(elev, scale=9)
par(mfrow=c(1,2))
plot(elev, main="original raster")
plot(tpi9, main="tpi 9x9")
```

trasp

Solar-radiation Aspect Index

Description

Calculates the Roberts and Cooper (1989) Solar-radiation Aspect Index

Usage

```
trasp(x, ...)
```

Arguments

x	raster object
...	Additional arguments passed to raster::calc

Value

raster class object of Roberts and Cooper (1989) Solar-radiation Aspect Index

Note

Roberts and Cooper (1989) rotates (transforms) the circular aspect to assign a value of zero to land oriented in a north-northeast direction, (typically the coolest and wettest orientation), and a value of one on the hotter, dryer south-southwesterly slopes. The result is a continuous variable between 0 - 1. The metric is defined as: $\text{trasp} = (1 - \cos((\pi/180)(a-30))) / 2$ where; a = aspect in degrees

Author(s)

Jeffrey S. Evans <jeffrey_evans@tnc.org>

References

Roberts. D.W., and Cooper, S.V. (1989). Concepts and techniques of vegetation mapping. In Land Classifications Based on Vegetation: Applications for Resource Management. USDA Forest Service GTR INT-257, Ogden, UT, pp 90-96

Examples

```
library(raster)
data(elev)
s <- trasp(elev)
plot(s)
```

trend.line

trend.line

Description

Calculated specified trend line of x,y

Usage

```
trend.line(x, y, type = "linear", plot = TRUE, ...)
```

Arguments

x	Vector of x
y	Vector of y
type	Trend line types are: 'linear', 'exponential', 'logarithmic', 'polynomial'
plot	plot results (TRUE/FALSE)
...	Additional arguments passed to plot

Value

A list class object with the following components: for type = 'linear' x is slope and y is intercept for type = 'exponential', 'logarithmic', or 'polynomial' x is original x variable and y is vector of fit regression line

Author(s)

Jeffrey S. Evans <jeffrey_evans@tnc.org>

Examples

```
x <- 1:10
y <- jitter(x^2)
opar <- par
par(mfcol=c(2,2))
  trend.line(x,y,type='linear',plot=TRUE,pch=20,main='Linear')
  trend.line(x,y,type='exponential',plot=TRUE,pch=20,main='Exponential')
  trend.line(x,y,type='logarithmic',plot=TRUE,pch=20,main='Logarithmic')
  trend.line(x,y,type='polynomial',plot=TRUE,pch=20,main='Polynomial')
par <- opar
```

tri

Terrain Ruggedness Index

Description

Implementation of the Riley et al (1999) Terrain Ruggedness Index

Usage

```
tri(r, s = 3, exact = TRUE, file.name = NULL, ...)
```

Arguments

r	raster class object
s	Scale of window. Must be odd number, can represent 2 dimensions (eg., s=c(3,5) would represent a 3 x 5 window)
exact	Calculate (TRUE/FALSE) the exact TRI or an algebraic approximation.
file.name	Name of output raster (optional)
...	Additional arguments passed to writeRaster

Value

raster class object or raster written to disk

Note

The algebraic approximation is considerably faster. However, because inclusion of the center cell, the larger the scale the larger the divergence of the minimum value

Recommended ranges for classifying Topographic Ruggedness Index

0-80 (1) level terrain surface.

81-116 (2) nearly level surface.

117-161 (3) slightly rugged surface.

162-239 (4) intermediately rugged surface.

240-497 (5) moderately rugged surface.

498-958 (6) highly rugged surface.

>959 (7) extremely rugged surface.

Depends: raster

Author(s)

Jeffrey S. Evans <jeffrey_evans@tnc.org>

References

Riley, S.J., S.D. DeGloria and R. Elliot (1999) A terrain ruggedness index that quantifies topographic heterogeneity, Intermountain Journal of Sciences 5(1-4):23-27.

Examples

```
library(raster)
data(elev)
( tri.ext <- tri(elev) )
( tri.app <- tri(elev, exact = FALSE) )
plot(stack(tri.ext, tri.app))
```

vrm

Vector Ruggedness Measure (VRM)

Description

Implementation of the Sappington et al., (2007) vector ruggedness measure

Usage

```
vrm(x, s = 3, file.name = NULL, ...)
```

Arguments

x	Elevation raster class object
s	Scale of window. Must be odd number, can represent 2 dimensions (eg., s=c(3,5) would represent a 3 x 5 window)
file.name	Name of output raster (optional)
...	Additional arguments passed to writeRaster

Value

raster class object or raster written to disk

Note

This function measures terrain ruggedness by calculating the vector ruggedness measure

Author(s)

Jeffrey S. Evans <jeffrey_evans@tnc.org>

References

Sappington, J.M., K.M. Longshore, D.B. Thomson (2007). Quantifying Landscape Ruggedness for Animal Habitat Analysis: A case Study Using Bighorn Sheep inthe Mojave Desert. Journal of Wildlife Management. 71(5):1419-1426

Examples

```
library(raster)
data(elev)
vrm3 <- vrm(elev)
vrm5 <- vrm(elev, s=5)
plot(stack(vrm3, vrm5))
```

winsorize

Winsorize transformation

Description

Removes extreme outliers using a winsorization transformation

Usage

```
winsorize(x, min.value = NULL, max.value = NULL, p = c(0.05, 0.95),
na.rm = FALSE)
```

Arguments

x	A numeric vector
min.value	A fixed lower bounds, all values lower than this will be replaced by this value. The default is set to the 5th-quantile of x.
max.value	A fixed upper bounds, all values higher than this will be replaced by this value. The default is set to the 95th-quantile of x.
p	A numeric vector of 2 representing the probabilities used in the quantile function.
na.rm	(FALSE/TRUE) should NAs be omitted?

Value

A transformed vector the same length as *x*, unless *na.rm* is TRUE, then *x* is length minus number of NA's

Note

Winsorization is the transformation of a distribution by limiting extreme values to reduce the effect of spurious outliers. This is done by shrinking outlying observations to the border of the main part of the distribution.

Author(s)

Jeffrey S. Evans <jeffrey_evans@tnc.org>

References

Dixon, W.J. (1960) Simplified Estimation from Censored Normal Samples. Annals of Mathematical Statistics. 31(2):385-391

Examples

```
set.seed(1234)
x <- rnorm(100)
x[1] <- x[1] * 10
winsorize(x)

plot(x, type="l", main="Winsorization transformation")
lines(winsorize(x), col="red", lwd=2)
legend("bottomright", legend=c("Original distribution","With outliers removed"),
       lty=c(1,1), col=c("black","red"))
# Behavior with NA value(s)
x[4] <- NA
winsorize(x)           # returns x with original NA's
winsorize(x, na.rm=TRUE) # removes NA's
```

wt.centroid

Weighted centroid

Description

Creates centroid of [x,y] coordinates based on a weights field

Usage

```
wt.centroid(x, p, sp = TRUE)
```

Arguments

- x sp SpatialPointsDataFrame class object
- p Weights column in x@data slot
- sp Output sp SpatailPoints class object (TRUE | FALSE)

Value

A vector or an sp class SpatialPoints object of the weighted coordinate centroid

Note

The weighted centroid is calculated as: $[Xw]=[X]*[p]$, $[Yw]=[Y]*[p]$, $[sXw]=\text{SUM}[Xw]$, $[sYw]=\text{SUM}[Yw]$, $[sP]=\text{SUM}[p]$ $wX=[sXw]/[sP]$, $wY=[sYw]/[sP]$ where; X=X COORDINATE(S), Y=Y COORDINATE(S), p=WEIGHT

Depends: sp

Examples

```
require(sp)
data(meuse)
coordinates(meuse) = ~x+y
wt.copper <- wt.centroid(meuse, 'copper', sp=TRUE)
wt.zinc <- wt.centroid(meuse, 'zinc', sp=TRUE)
plot(meuse, pch=20, cex=0.75, main='Weighted centroid(s)')
points(wt.copper, pch=19, col='red', cex=1.5)
points(wt.zinc, pch=19, col='blue', cex=1.5)
box()
legend('topleft', legend=c('all','copper', 'zinc'),
      pch=c(20,19,19),col=c('black','red','blue'))
```

zonal.stats

zonal.stats

Description

Polygon zonal statistics of a raster

Usage

```
zonal.stats(x, y, stat, trace = TRUE, plot = TRUE)
```

Arguments

x	Polygon object of class SpatialPolygonsDataFrame
y	Raster object of class raster
stat	Statistic or function
trace	Should progress counter be displayed
plot	Should subset polygons/rasters be plotted (TRUE/FALSE)

Value

Vector, length of nrow(x), of function results

Note

This function iterates through a polygon object, masks the raster to each subset polygon and then coerces the subset raster to a vector object. The resulting vector is then passed to the specified statistic/function. This is much slower than zonal functions available in GIS software but has the notable advantage of being able to accept any custom function, passed to the 'stat' argument, appropriate for a vector object (see example).

Depends: sp, raster

Author(s)

Jeffrey S. Evans <jeffrey_evans@tnc.org>

Examples

```
# skewness function
skew <- function(x, na.rm=FALSE) {
  if (na.rm)
    x <- x[!is.na(x)]
  sum( (x - mean(x)) ^ 3) / ( length(x) * sd(x) ^ 3 )
}

# percent x >= p function
pct <- function(x, p=0.30) {
  if ( length(x[x >= p]) < 1 )  return(0)
  if ( length(x[x >= p]) == length(x) ) return(1)
  else return( length(x[x >= p]) / length(x) )
}

# create some example data
library(raster)
library(sp)
p <- raster(nrow=10, ncol=10)
p[] <- runif(ncell(p)) * 10
p <- rasterToPolygons(p, fun=function(x){x > 9})
r <- raster(nrow=100, ncol=100)
r[] <- runif(ncell(r))
```

```
plot(r)
plot(p, add=TRUE, lwd=4)

# run zonal statistics using skew and pct functions
z.skew <- zonal.stats(x=p, y=r, stat=skew, trace=TRUE, plot=TRUE)
z.pct <- zonal.stats(x=p, y=r, stat=pct, trace=TRUE, plot=TRUE)
( z <- data.frame(ID=as.numeric(as.character(row.names(p@data))),
SKEW=z.skew, PCT=z.pct) )
```

Index

ants, 4
autocov_dist, 51

bearing.distance, 5
breeding.density, 5

calc, 107, 112
clara, 60
class.comparison, 7
classBreaks, 9
ClassStat, 33, 45
concordance, 10
conf.interval, 11
ConnCompLabel, 33, 45
correlogram, 12
crossCorrelation, 13
csi, 16
curvature, 17

daymet.point, 19
daymet.tiles, 20
DAYMET_tiles, 21
dispersion, 22
dissection, 23
download.daymet, 24, 28
download.hansen, 25, 28
download.prism, 27

effect.size, 28
elev, 29
erase.point, 30

focal, 112
focal.lmetrics, 31, 45
fuzzySum, 33

gaussian.kernel, 34
group.pdf, 35

hclust, 40
hexagons, 36

hli, 37
hsp, 38
hybrid.kmeans, 39

idw.smoothing, 40
insert.values, 41

kde.2D, 42
kendallTrendTest, 84
kl.divergence, 42
kmeans, 39, 40

land.metrics, 33, 43
local.min.max, 46
loess, 69, 107
loess.boot, 47
loess.ci, 48
logistic.regression, 50
lrm, 51

moments, 52
morans.plot, 53
mwCorr, 55

nni, 55

o.ring, 56
oli.asw, 58
optimal.k, 59
optimized.sample.variance, 60
outliers, 62
overlay, 84

pam, 60
parea.sample, 63
PatchStat, 33, 45
plot.effect.size, 64
plot.loess.boot, 65
point.in.poly, 66
poly.regression, 68
polyPerimeter, 69

pp.subsample, 70
print.cross.cor, 72
print.effect.size, 72
print.loess.boot, 73
pseudo.absence, 73
pu, 76
raster.deviation, 77
raster.downscale, 79
raster.entropy, 80
raster.gaussian.smooth, 81
raster.invert, 82
raster.kendall, 83
raster.mds, 85
raster.modified.ttest, 86
raster.moments, 88
raster.transformation, 89
raster.vol, 90
raster.Zscore, 91
rasterCorrelation, 92
remove.holes, 93

sa.trans, 94
sample.annulus, 95
sample.line, 96
sample.poly, 98
sampleTransect, 99
sar, 100
separability, 101
sg.smooth, 102
shannons, 104
similarity, 105
smooth.time.series, 106
sobal, 108
sp.kde, 109
sp.na.omit, 110
spherical.sd, 111
srr, 113
stratified.random, 114
subsample.distance, 115
summary.cross.cor, 117
summary.effect.size, 117
summary.loess.boot, 118
swvi, 118

tpi, 120
trasp, 121
trend.line, 122
tri, 123
vrm, 124
winsorize, 125
writeRaster, 18
wt.centroid, 126
zonal.stats, 127