

Package ‘pgirmess’

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Title Data Analysis in Ecology

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Description

Miscellaneous functions for data analysis in ecology, with special emphasis on spatial data.

Imports boot (>= 1.3-4), maptools (>= 0.8-36), rgdal (>= 0.7-8), rgeos (>= 0.3-8), sp (>= 0.9-97), spdep (>= 0.5-43), splancs (>= 2.01-31)

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CI *Confidence interval of percentages*

Description

Computes the lower limit and upper limit of the 95 percent confidence interval of percentage estimates

Usage

```
CI(x, ...)
```

Arguments

`x` a two-dimensional table, matrix or data.frame with 2 columns, giving the counts of successes and failures, respectively

`...` other arguments to pass to [prop.test](#), eg `conf.level`

Details

Simple wrapper of [prop.test](#). The default confidence interval is 95 percent, but can be modified passing values to [prop.test](#) by the `conf.level` argument.

Value

A 3 column matrix.

- Column 1: percentage estimate
- Column 2: lower limit of the confidence interval
- column 3: upper limit of the confidence interval

See Also

[prop.test](#)

Examples

```
x<-c(2,10,7,8,7) # eg: number of positive cases
y<-c(56,22,7,20,5)# eg: number of negative cases
CI(cbind(x,y))
CI(cbind(x,y), conf.level=0.99)
```

classnum	<i>Gives an index vector of the class category of each value of a numerical vector</i>
----------	--

Description

Gives an index vector of the class category of each value of a numerical vector

Usage

```
classnum(x, breaks = "Sturges")
```

Arguments

x	a vector of values for which the indices are desired
breaks	one of: <ul style="list-style-type: none">• a vector giving the breakpoints between bins,• a single number giving the number of bins,• a character string naming an algorithm to compute the number of cells (see Details).

Details

The default for 'breaks' is "Sturges": see 'nclass.Sturges'. Other names for which algorithms are supplied are "Scott" and "FD" for "Friedman-Diaconis" (with corresponding functions 'nclass.scott' and 'nclass.FD'). Case is ignored and partial matching is used. Breaks and labels are stored as attributes.

Value

A vector of the same length as x, with the index of the class which each value of x belongs to

See Also

[cut](#), [classIntervals](#)

Examples

```
x<-rnorm(30)
classnum(x)
classnum(x,breaks="fd")
classnum(x, breaks=c(-1,0,1))
classnum(x,breaks=5)
```

cormat	<i>Gives a correlation matrix and the probability of Ho for each correlation</i>
--------	--

Description

Gives a correlation matrix and the probability of Ho for each correlation estimate

Usage

```
cormat(donnees, method = "spearman", sep = FALSE)
```

Arguments

donnees	a data frame of numerics
method	a string of characters among 'pearson', 'spearman' (default), 'kendall'
sep	If true, gives the results in two matrices (default = F)

Details

Wrapper for 'cor' and 'cor.test'. The results can be given in one or two matrices.

Value

If sep = F (default) a list including:

method	The method used
prob.cor	Upper triangle, the correlations; lower triangle, the probability of Ho

If sep = T a list including:

method	The method used
coef.estimates	The correlation matrix
p.value	The Ho probability matrix

Author(s)

Patrick Giraudoux <pgiraud@univ-fcomte.fr>

See Also

[cor](#), [cor.test](#)

Examples

```
cormat(longley)
cormat(longley, sep=TRUE)
```

correlog *Computes Moran's or Geary's coefficients on distance classes*

Description

Computes Moran's or Geary's coefficients on distance classes from a set of spatial coordinates and corresponding z values

Usage

```
correlog(coords, z, method="Moran", nbclass = NULL,...)
```

Arguments

coords	a two columns array, data.frame or matrix of spatial coordinates. Column 1 = X, Column 2 = Y.
z	a vector for the values at each location. Must have the same length as the row number of coords
method	the method used. Must be "Moran" (default) or "Geary"
nbclass	number of bins. If NULL Sturges method is used to compute an optimal number
...	further arguments to pass to e.g. moran.test or geary.test

Details

Uses the library spdep including [moran.test](#) or [geary.test](#). Distances are euclidian and in the same unit as the spatial coordinates. Moran's Ho: I values larger than 0 due to chance; Geary's Ho: C values lesser than 1 due to chance. Correlog has print and plot methods; statistically significant values ($p < 0.05$) are plotted in red.

Value

An object of class "correlog", a matrix including:

class	bin centers
I	the coefficient values
p.value	probability of Ho
n	the number of pairs

Warning

Computing can take a long time for large data sets

Author(s)

Patrick Giraudoux pgiraud@univ-fcomte.fr, Colin Beale c.beale@macaulay.ac.uk and Mike Treglia mike-treglia@utulsa.edu

References

see library spdep

See Also

[geary.test](#), [moran.test](#)

Examples

```
library(spdep)
data(oldcol)
attach(COL.OLD)
coords<-cbind(X,Y)
res<-correlog(coords,CRIME)
plot(res)

res<-correlog(coords,CRIME,method="Geary")
plot(res)
```

date2winter

Convert a POSIXt date into categories corresponding to a autumn/winter/spring sequence

Description

Convert a POSIXt date into categories corresponding to the time spanning from the late months of a year to the early months of the following year

Usage

```
date2winter(x, first = 10, last=4)
```

Arguments

x	a vector of POSIXt dates
first	number of the first month to include (default 10, October)
last	number of the last month to include (default 4, April)

Details

In ecology, time data must often be analysed on a time span category covering two successive years (eg the winter period). This function convert POSIXt dates into categories corresponding to the time span stretching from a user defined month of a given year (by default October) to a user-defined month of the following year (by default April). If date month is out of the user defined time span the value 'Excluded' is returned.

Value

A vector of the same length as x, with the time span category each value belongs to.

Examples

```
dates <- strptime(c("02/12/2002", "15/01/2003", "15/10/2003", "15/6/2003", NA), "%d/%m/%Y")
date2winter(dates)
```

diag2edge

Computes the edge of a square from its diagonal

Description

Computes the edge of a square from its diagonal.

Usage

```
diag2edge(cordseg)
```

Arguments

cordseg The diagonal coordinates. This can be a vector c(x1,y1,x2,y2), a 2 x 2 matrix or a data.frame (each line a coordinate)

Details

The first point coordinates are the left top of the diagonal. The other coordinates computed are the other top of the square edge. Can be used e.g. to pass a square edge to [pave](#) in order to compute a sampling grid.

Value

A 2x2 matrix of points coordinates

Author(s)

Patrick Giraudoux <pgiraudou@univ-fcomte.fr>

See Also

[pave](#)

Examples

```
# diagonal sloping up
coord<-matrix(c(20,20,90,90),nr=2,byrow=TRUE)
plot(coord,type="n",xlim=c(0,100),ylim=c(0,110),asp=1)
lines(coord,lty=2)
# square edge
lines(diag2edge(coord),col="red")

# diagonal sloping down
coord<-matrix(c(20,90,90,20),nr=2,byrow=TRUE)
plot(coord,type="n",xlim=c(0,100),ylim=c(0,110),asp=1)
lines(coord,lty=2)
# square edge
lines(diag2edge(coord),col="red")

# diagonal vertical
coord<-matrix(c(20,90,20,20),nr=2,byrow=TRUE)
plot(coord,type="n",xlim=c(0,100),ylim=c(0,110),asp=1)
lines(coord,lty=2)
# square edge
lines(diag2edge(coord),col="red")
```

difshannonbio

Empirical confidence interval of the bootstrap of the difference between two Shannon indices

Description

Computes the empirical confidence interval of the bootstrap of the difference between two Shannon indices

Usage

```
difshannonbio(dat1, dat2, R = 1000, probs = c(0.025, 0.975))
```

Arguments

dat1	a data.frame of two columns; column 1 = category, column 2 = biomass
dat2	a data.frame of two columns; column 1 = category, column 2 = biomass
R	number of permutations
probs	the limits of the confidence interval

Details

Designated to compare the difference between two Shannon's indices computed from two data frames. In each data frame, the first column is the category of prey item, and the second column the estimated biomass.

Value

A list with the confidence interval of H' and J'

Author(s)

patrick.giraudoux <pgiraud@univ-fcomte.fr>

See Also

[shannonbio](#)

Examples

```
data(preymbiom)
attach(preymbiom)
jackal<-preymbiom[site=="Y" & sp=="C",5:6]
genet<-preymbiom[site=="Y" & sp=="G",5:6]

difshannonbio(jackal,genet,R=150)
```

dirProj

Computes new coordinates given bearings and distances.

Description

Computes new coordinates from bearings (North = 0) and distances

Usage

```
dirProj(df,deg=TRUE)
```

Arguments

df	a matrix or data frame of 4 columns giving x, y coordinates, bearings and distances
deg	if TRUE (default) bearings are in degree, otherwise in radian

Details

Computings are based on euclidian distance. Therefore, the coordinates should be given in a projected (plan) system (e.g. UTM, Lambert, etc.) and the distance in the same units as the projection system (e.g. meters).

Value

a matrix of two columns with the projected coordinates

See Also[distSeg](#)**Examples**

```
df<-data.frame(x1=0,y1=0,alpha=runif(3,0,360),d=runif(3,0,1))
df
plot(-1:1,-1:1,type="n")
points(0,0,pch=19)
points(dirProj(df))
text(dirProj(df)[,1],dirProj(df)[,2],1:3,pos=4)
```

dirSeg*Computes segment directions.*

Description

Computes the direction of segments from the first top clockwise (North = 0)

Usage

```
dirSeg(x,deg=TRUE)
```

Arguments

x	a matrix or data frame of 4 columns giving the coordinates of each segment tops x1, y1, x2, y2
deg	if TRUE (default) the output is in degrees, otherwise in radians

Details

The first two columns give the first top coordinates, x then y, and the next two the second top coordinates.

Value

A vector of directions

See Also[dirProj](#), [gzAzimuth](#)

Examples

```
x2<-rnorm(10)
y2<-rnorm(10)
mydata<-cbind(0,0,x2,y2)
dirs<-dirSeg(mydata)
dirs

plot(range(mydata[,c(1,3)]),range(mydata[,c(2,4)]),type="n")
Segments(mydata)
text(mydata[,3],mydata[,4],paste(round(dirs,0),"\u00b0"),cex=0.7)
```

distNNeigh*Computes distances to the nearest neighbour*

Description

Computes distances to the nearest neighbour

Usage

```
distNNeigh(db)
```

Arguments

db A matrix or data.frame of points coordinates column 1 = x,column 2 = y.

Details

Computes distances to the nearest neighbour for each line of a matrix of points coordinates

Value

A vector of distances

See Also

[knearneigh](#), [knn2nb](#), [nbdists](#)

Examples

```
distNNeigh(cbind(rnorm(30),rnorm(30)))
```

distNode	<i>Computes the distances between each nodes of a polyline.</i>
----------	---

Description

Computes the distances between each nodes of a polyline.

Usage

```
distNode(pts, decdeg=FALSE)
```

Arguments

pts	A matrix or data.frame of the node coordinates column 1 = x, column 2 = y.
decdeg	TRUE if point coordinates are longitude-latitude decimal degrees, in which case distances are measured in meters

Details

If decdeg is FALSE (default), distance computed is Euclidian. Units depends on the coordinate systems. If decdeg = TRUE, $D = 1852 * 60 * (180/\pi) * \arccos(\sin(la1) * \sin(la2) + \cos(la1) * \cos(la2) * \cos(\text{abs}(lg1 - lg2)))$. This method calculates the great circle distance, is based on spherical trigonometry, and assumes that:

- 1 minute of arc is 1 nautical mile
- 1 nautical mile is 1.852 km

Value

A vector of distances

See Also

[distTot](#), [distSeg](#)

Examples

```
x<-c(10,56,100)
y<-c(23,32,150)
distNode(cbind(x,y))
```

distSeg	<i>Computes distances between the top coordinates of segments.</i>
---------	--

Description

Computes the distances between the top coordinates of segments.

Usage

```
distSeg(mydata, decdeg=FALSE)
```

Arguments

mydata	A matrix or data frame of 4 columns giving the coordinates of each segment tops x1, y1, x2, y2
decdeg	TRUE if point coordinates are longitude-latitude decimal degrees, in which case distances are measured in meters

Details

If `decdeg` is FALSE (default), distance computed is Euclidian. Units depends on the coordinate systems. If `decdeg = TRUE`, $D = 1852 * 60 * (180/\pi) * \text{acos}(\sin(\text{la1}) * \sin(\text{la2}) + \cos(\text{la1}) * \cos(\text{la2}) * \cos(\text{abs}(\text{lg1} - \text{lg2})))$. This method calculates the great circle distance, is based on spherical trigonometry, and assumes that:

- 1 minute of arc is 1 nautical mile
- 1 nautical mile is 1.852 km

Value

A vector of distances

See Also

[distNode](#), [distTot](#)

Examples

```
x1<-rnorm(20)
y1<-rnorm(20)
x2<-rnorm(20)
y2<-rnorm(20)
mydata<-cbind(x1,y1,x2,y2)
distSeg(mydata)
```

distTot	<i>Computes the total length of a polyline.</i>
---------	---

Description

Computes the total length of a polyline.

Usage

```
distTot(pts,decdeg=FALSE)
```

Arguments

pts	A matrix or data.frame of the node coordinates column 1 = x,column 2 = y.
decdeg	TRUE if point coordinates are longitude-latitude decimal degrees, in which case distances are measured in meters

Details

If decdeg is FALSE (default), distance computed is Euclidian. Units depends on the coordinate systems. If decdeg = TRUE, $D = 1852 * 60 * (180/\pi) * \text{acos}(\sin(la1) * \sin(la2) + \cos(la1) * \cos(la2) * \cos(\text{abs}(lg1 - lg2)))$. This method calculates the great circle distance, is based on spherical trigonometry, and assumes that:

- 1 minute of arc is 1 nautical mile
- 1 nautical mile is 1.852 km

Value

A numeric distance.

See Also

, [distNode](#), [distSeg](#)

Examples

```
x<-c(10,56,100)
y<-c(23,32,150)
distTot(cbind(x,y))
```

expandpoly

Homothetia (size expansion) of a polygon

Description

Compute the new coordinates of polygon expanded by a factor.

Usage

```
expandpoly(mypol, fact)
```

Arguments

mypol	matrix or data.frame of polygon coordinates
fact	expansion factor

Details

The polygon area obtained after expansion is equal to $fact^2$ times the original polygon area

Value

A matrix of polygon coordinates

See Also

[polygon](#)

Examples

```
x<-c(-5,-4.5,0,10,5)
y<-c(-10,0,5,5,-8)
poly<-cbind(x,y)
plot(-10:20,-20:10,type="n")
polygon(poly)
polygon(expandpoly(poly,1.5),border="red")
polygon(expandpoly(poly,0.5),border="blue")
```

findR	<i>Computes the distance between the centroid and the most distant coordinate of a geographical coordinate set</i>
-------	--

Description

Computes the distance between the centroid and the most distant coordinate of a geographical coordinate set.

Usage

```
findR(coords)
```

Arguments

coords A matrix or data frame of 2 columns of geographical coordinates

Value

The distance

See Also

[polycirc](#)

Examples

```
mydata<-cbind(x=rnorm(20),y=rnorm(20))
radius<-findR(mydata)
centroid<-matrix(colMeans(mydata),ncol=2)
plot(mydata,asp=1)
points(centroid,pch=19,col="red",cex=2)
polygon(polycirc(radius,centroid),border="red")
```

friedmanmc	<i>Multiple comparisons after Friedman test</i>
------------	---

Description

Test of multiple comparison after Friedman test

Usage

```
friedmanmc(y, groups, blocks,probs=0.05)
```

Arguments

y	a numeric vector of data values, or a data matrix
groups	a vector giving the group for the corresponding elements of 'y' if this is a vector; ignored if 'y' is a matrix. If not a factor object, it is coerced to one.
blocks	a vector giving the block for the corresponding elements of 'y' if this is a vector; ignored if 'y' is a matrix. If not a factor object, it is coerced to one.
probs	a probability for the critical difference.

Details

Method for formula still not implemented. Formula 7.5a (Siegel & Castellan, 1988 p 180-181) can lead to p values larger than 1 when differences between groups are small. Eventually, they are set to NA and a warning is generated.

Value

A list of class 'mc' with the following items:

statistic	statistics used
p.value	the p value of the critical difference
dif.com	a data.frame with observed and critical differences

References

Siegel & Castellan (1988) Non parametric statistics for the behavioural sciences. Mc Graw Hill Int. Edt.

See Also

[friedman.test](#)

Examples

```
data(siegelp179)
attach(siegelp179)

friedman.test(score,treatment,block)
friedmanmc(score,treatment,block)
friedmanmc(score,treatment,block,probs=0.01)

mymatrix<-matrix(score,nc=3)
friedman.test(mymatrix)
friedmanmc(mymatrix)
detach(siegelp179)
```

 gps2gpx

Download waypoints or tracks from a GPS to a gpx file

Description

Download waypoints or tracks from a GPS to a gpx file or to the console gpx formatted

Usage

```
gps2gpx(filename="",i="garmin",f = "usb:", type = "w", invisible = TRUE)
```

Arguments

filename	a character string naming the file to print to. If "" (the default), prints to the standard output connection
i	INTYPE: a supported file type, default "garmin"
f	INFILE: the appropriate device interface, default "usb:", on Windows for serial interfaces commonly "com4:" or similar
type	"w" waypoints, or "t" track, or others provided in gpsbabel
invisible	Under Windows, do not open an extra window

Details

The function calls gpsbabel via the system. The gpsbabel program must be present and on the user's PATH for the function to work see <http://www.gpsbabel.org>. A .gpx suffix is added if not included in the filename. The gpx file can then be read e.g. using [readOGR](#) to a sp spatial object. Ex: `readOGR("filename.gpx", "waypoints", drop_unsupported_fields=TRUE)`, or uploaded to a GPS

See Also

[readOGR,uploadGPS](#)

Examples

```
## Not run:
# a GPS device must be connected
gps2gpx() # download waypoints and print to the console
gps2gpx(t="t") # download tracks and print to the console
gps2gpx(filename="myfile") # download waypoints and write a gpx file

## End(Not run)
```

kruskalmc

Multiple comparison test after Kruskal-Wallis

Description

Multiple comparison test between treatments or treatments versus control after Kruskal-Wallis test

Usage

```
kruskalmc(resp,...)
## Default S3 method:
kruskalmc(resp, categ, probs = 0.05, cont=NULL,...)
## S3 method for class 'formula'
kruskalmc(resp,data=NULL,...)
```

Arguments

resp	a numeric vector of data values or a formula of the type 'response~category'.
categ	a factor object giving the group for the corresponding elements of 'x'
probs	the significance level
cont	NULL (default) for multiple comparison between treatments; 'one-tailed' or 'two-tailed' for corresponding multiple comparisons treatments versus control; partial matching allowed
data	a data.frame including the variables used in the formula
...	other parameters to be passed as arguments (not used here)

Details

When the value of a Kruskal-Wallis test is significant, it indicates that at least one of the groups is different from at least one of the others. This test helps determining which groups are different with pairwise comparisons adjusted appropriately for multiple comparisons. Those pairs of groups which have observed differences higher than a critical value are considered statistically different at a given significance level. Three types of multiple comparisons are implemented: comparisons between treatments, 'one-tailed' and 'two-tailed' comparison treatments versus control. The first factor level is considered the control. NAs are omitted from data before processing.

For further details please consider the reference below where the method is fully described, or visit <http://pagesperso-orange.fr/giraudoux/#pgirmess> where a copy of the corresponding book section is downloadable.

Value

A list of class 'mc' with the following items:

statistic	statistics used
signif.level	the significance level
dif.com	a data.frame with observed and critical differences

Note

Two alternative methods are proposed in the section 'see also', on François Gillet's suggestion. The three methods do not give necessarily the same results, and the why is still to investigate

References

Siegel and Castellan (1988) Non parametric statistics for the behavioural sciences. MacGraw Hill Int., New York. pp 213-214

See Also

[kruskal.test](#); to reorder factor levels see [relevel](#); for other functions about median multiple comparison see [posthoc.kruskal.conover.test](#), [kruskal](#)

Examples

```
resp<-c(0.44,0.44,0.54,0.32,0.21,0.28,0.7,0.77,0.48,0.64,0.71,0.75,0.8,0.76,0.34,0.80,0.73,0.8)
categ<-as.factor(rep(c("A","B","C"),times=1,each=6))
kruskalmc(resp, categ)
kruskalmc(resp, categ, probs=0.01)
kruskalmc(resp, categ, cont="one-tailed")
kruskalmc(resp, categ, cont="two-tailed")

kruskalmc(resp~categ)
kruskalmc(resp~categ, probs=0.01)
kruskalmc(resp~categ, cont="one-tailed")
kruskalmc(resp~categ, cont="two-tailed")
```

ks.gof

Kolmogorof-Smirnov goodness of fit test to normal distribution

Description

Kolmogorof-Smirnov goodness of fit test to normal distribution

Usage

```
ks.gof(var)
```

Arguments

var a numeric vector

Details

A wrapper of `ks.test()`

Value

A list with class `"htest"` containing the following components:

<code>statistic</code>	the value of the test statistic.
<code>p.value</code>	a character string indicating what type of test was performed.
<code>alternative</code>	a character string describing the alternative hypothesis.
<code>method</code>	a character string indicating what type of test was performed.
<code>data.name</code>	a character string giving the name(s) of the data.

References

see `ks.test`

See Also

[ks.test](#)

Examples

```
x<-rnorm(50)
ks.gof(x)
```

<code>mergeTrackObs</code>	<i>Merge two <code>SpatialPoints</code> or <code>SpatialPointsDataFrame</code> objects, one modelling a track, the other observations</i>
----------------------------	---

Description

Merge two `SpatialPoints` or `SpatialPointsDataFrame` objects, one modelling a track, the other observations.

Usage

```
mergeTrackObs(sppdfInt, sppdfObs, obscol=NULL)
```

Arguments

<code>sppdfInt</code>	A SpatialPoints object (the track)
<code>sppdfObs</code>	A SpatialPoints or SpatialPointsDataFrame object (the observations)
<code>obscol</code>	The column number in which the number of observations at this point can be found in <code>sppdfObs</code>

Details

Road site counts or faeces collections are often carried out along tracks (paths, roads, transects, etc.). Tracks can be discretized in regular intervals e.g. with `transLines2pix` or `thintrack`, each point being an interval centre. `mergeTrackObs` uses such a discretized track and sums observations to their nearest track interval. The output is a `SpatialPointsDataFrame` where each point corresponds to the centre of one track interval. The number of observations in each interval is given in the attribute file. If the number of observations at an observation point can be 0 or any positive number, use `obscol` to identify the column of `sppdfObs` where this number is stored.

Value

A `SpatialPointsDataframe`, with the following attributes:

- ID ID number
- nObs The number of observations in the interval

See Also

[transLines2pix](#), [thintrack](#)

Examples

```
# track
library(sp)
l2 = cbind(c(1,2,3),c(1,1.5,1))
S12 = Line(l2)
S2 = Lines(list(S12), ID="b")
S1 = SpatialLines(list(S2))
plot(S1, col = "blue")
#observations
obs <- structure(list(ID = 1:15, long = c(1.04609377280342, 1.0890625305741,
1.08125002916125, 1.24921880953755, 1.34687507719818, 1.50312510545521,
1.88984392539134, 2.37812526369453, 2.39375026652023, 2.36640651157525,
2.38593776510738, 2.62031280749291, 2.69843782162142, 2.85078159917202,
2.90546910906198), lat = c(1.04062476682306, 1.05624976964876,
1.03671851611663, 1.13828103448369, 1.16562478942867, 1.26718730779574,
1.43124983746561, 1.32968731909855, 1.32187481768569, 1.30624981485999,
1.28281231062144, 1.20468729649293, 1.13828103448369, 1.08749977530016,
1.03671851611663)), .Names = c("ID", "long", "lat"), row.names = c(NA,
-15L), class = "data.frame")
points(obs[,2:3],col="red")
coordinates(obs)<-~long+lat
obs@data$n<-c(3,4,0,1,1,5,6,4,3,4,4,7,2,2,1) # possibly a count on each location

# examples

# one observation on each location
track<-transLines2pix(S1,0.1)
trackObs<-mergeTrackObs(track,obs)
```

```

par(mfrow=c(1,2))
plot(S1)
plot(track,add=TRUE,col="blue")
plot(obs,add=TRUE,col="red",pch=1)

plot(S1)
plot(track,add=TRUE,col="blue")
plot(trackObs,cex=trackObs@data$nObs,pch=19, col="red",add=TRUE)

# 0 or more observations on each location
obs@data$n<-c(3,4,0,1,1,5,6,4,3,4,4,7,2,2,1) # possibly a count on each location
obs@data
trackObs<-mergeTrackObs(track,obs,obscol=2)

par(mfrow=c(1,2))
plot(S1)
plot(track,add=TRUE,col="blue")
plot(obs,add=TRUE,col="red",pch=1)

plot(S1)
plot(track,add=TRUE,col="blue")
plot(trackObs,cex=trackObs@data$nObs/3,pch=19, col="red",add=TRUE)

```

pairsrp

Produces a matrix of scatterplot, regression coefficient and p(Ho)

Description

Produces a matrix with scatterplot, regression line and a loess smooth in the upper right panel; correlation coefficient (Pearson, Spearman or Kendall) and the probability of Ho in the lower left panel

Usage

```
pairsrp(dataframe, meth = "spearman", pansmo = FALSE, abv = FALSE, lwt.cex = NULL, ...)
```

Arguments

dataframe	a data.frame of numeric values
meth	a character string indicating which correlation coefficient is to be computed. One of 'pearson', 'kendall', or 'spearman'(default). Can be abbreviated.
pansmo	True if a loess smooth is to be plotted. Default to False.
abv	True if the variable names must be abbreviates. Default to False.
lwt.cex	character size expansion in the lower panel.
...	graphical parameters can be given as arguments to 'plot'.

Details

This function is a wrapper for `pairs()` and `cor()`

See Also

[pairs](#)

Examples

```
data(iris)
pairsrp(iris[,1:4],meth="pears",pansmo=TRUE,abv=TRUE)
```

pave

Provide square polygons or their node coordinates along a segment

Description

Provide a user-defined cellgrid of polygon squares (or square node points) along a segment. This can be used to define a sampling grid for spatial analysis.

Usage

```
pave(cordseg, yc, xc, fix.edge=NULL, ydown = TRUE, output = "list")
```

Arguments

<code>cordseg</code>	the segment coordinates. This can be a vector $c(x1,y1,x2,y2)$, a 2 x 2 matrix or a <code>data.frame</code> (each line a coordinate)
<code>yc</code>	the number of segment divisions (y cells)
<code>xc</code>	the number of columns (x cells)
<code>fix.edge</code>	the edge length of a cell (user specified, default to NULL)
<code>ydown</code>	if TRUE (default) squares are computed decreasing y
<code>output</code>	a character string indicating which output is required. One of "list", "points" or "spdf". Partial match allowed

Details

The segment must have $x1 < x2$. If not, it is automatically reordered. When "spdf" is selected the output is an object of class `SpatialPolygonsDataframe`. It has a `plot` method and can straightfully be handled by `writeShapePoly` (see [readShapePoly](#)) of the `maptools` library to write a shapefile. The value of the edge length of a cell can passed with the argument `fix.edge`. In this case, the coordinates of the segment right top are re-computed to adjust the cell edge to an user defined fixed value.

Value

According to the output selected, a list of polygon coordinates, a 2 column matrix with the nodes coordinates or a SpatialPolygonsDataframe.

Author(s)

Patrick Giraudoux <pgiraudou@univ-fcomte.fr>

See Also

[SpatialPolygonsDataFrame-class](#), [readShapePoly](#), [readOGR](#), [over](#), [diag2edge](#)

Examples

```
# segment sloping up
coord<-matrix(c(20,20,90,90),nr=2,byrow=TRUE)
plot(coord,type="n",xlim=c(0,100),ylim=c(0,110),asp=1)
lines(coord)
# point grids
gr<-pave(coord,20,4,output="points") # y decreasing
points(gr)
gr<-pave(coord,20,4,output="points",ydown=FALSE) # y increasing
points(gr,col="blue")
# square polygon grids
gr<-pave(coord,20,4) # y decreasing
for (i in 1:length(gr)) polygon(gr[[i]])
gr<-pave(coord,20,4,ydown=FALSE) # y increasing
for (i in 1:length(gr)) polygon(gr[[i]],border="blue")

# segment sloping down
coord<-matrix(c(20,90,90,20),nr=2,byrow=TRUE)
plot(coord,type="n",xlim=c(0,100),ylim=c(0,110),asp=1)
lines(coord)

# point grids
gr<-pave(coord,20,4,output="points") # y decreasing
points(gr)
gr<-pave(coord,20,4,output="points",ydown=FALSE) # y increasing
points(gr,col="blue")

# fixed edge
plot(coord,type="n",xlim=c(0,100),ylim=c(0,110),asp=1)
lines(coord)
gr<-pave(coord,20,4,fix.edge=4,output="points")
points(gr,col="blue")

plot(coord,type="n",xlim=c(0,100),ylim=c(0,110),asp=1)
lines(coord)
gr<-pave(coord,20,4,fix.edge=5.5,output="points")
points(gr,col="red")
```

```
# square polygon grids
coord<-matrix(c(20,90,90,20),nr=2,byrow=TRUE)
plot(coord,type="n",xlim=c(0,100),ylim=c(0,110),asp=1)
lines(coord,lwd=2)
gr<-pave(coord,20,4)# y decreasing
for (i in 1:length(gr)) polygon(gr[[i]])
gr<-pave(coord,20,4,ydown=FALSE) # y increasing
for (i in 1:length(gr)) polygon(gr[[i]],border="blue")

## Not run:
# Writing a polygon shapefile
gr<-pave(coord,20,4,output="spdf") # y decreasing
library(maptools)
writePolyShape(gr, "myshapefilename")

## End(Not run)
```

pclig

Compute the percentage of each cell of a matrix or data.frame by row

Description

Compute the percentage of each cells of a matrix or data.frame by row

Usage

```
pclig(matr)
```

Arguments

matr a matrix or a data.frame

Details

Compute the percentage of each cells of a matrix by row. NA are removed.

Value

Return a matrix with percentages in each cell

See Also

[prop.table](#)

Examples

```
x<-c(2,10,7,8,7)
y<-c(56,22,7,20,5)
pclig(cbind(x,y))
```

permcont	<i>Random permutation of a contingency table n row x 2 columns</i>
----------	--

Description

Return a random permutation of a contingency table n rows x 2 columns keeping the marginal totals

Usage

```
permcont(Table)
```

Arguments

Table a contingency table

Details

The contingency table is split in a two columns table of 0/1 categories, sampled and re-organised with the function table()

Value

A matrix with the permuted values

Examples

```
tab<-cbind(n1=c(10,12,8,7,5),n2=c(4,5,8,10,12))
tab
permcont(tab)
```

PermTest	<i>Permutation test for lm, lme and glm (binomial and Poisson) objects</i>
----------	--

Description

Permutation test for lm, lme and glm (binomial and Poisson) objects

Usage

```
PermTest(obj, B=1000,...)

## S3 method for class 'lm'
PermTest(obj, B=1000,...)
## S3 method for class 'lme'
PermTest(obj, B=1000,...)
## S3 method for class 'glm'
PermTest(obj, B=1000,...)
```

Arguments

obj	an object of class lm, lme, or glm
B	number of permutations, default = 1000
...	used to pass other arguments

Details

For glm, when the response is a two-column matrix with the columns giving the numbers of successes and failures, PermTest.glm uses permcont(); PermTest.lme requires the library nlme.

Value

A list object of class PermTest including:

p.value	the p value obtained
B	the number of permutations
call	the call

Warning

This generic function is implemented in R language, thus can be quite slow.

Note

The implementation of PermTest.lme has been helped by Renaud Lancelot

Examples

```
library(MASS)
mylm<-lm(Postwt~Prewt,data=anorexia)
PermTest(mylm,B=250)

## Dobson (1990) Page 93: Randomized Controlled Trial :
counts <- c(18,17,15,20,10,20,25,13,12)
outcome <- gl(3,1,9)
treatment <- gl(3,3)
glm.D93 <- glm(counts ~ outcome + treatment, family=poisson)
PermTest(glm.D93,B=100)

library(nlme)
fm2 <- lme(distance ~ age + Sex, data = Orthodont, random = ~ 1)
PermTest(fm2,B=100)
```

piankabio

Computes the Pianka's index of niche overlap

Description

Computes the Pianka's index of niche overlap

Usage

```
piankabio(dataframe1, dataframe2)
```

Arguments

dataframe1 a data frame of two columns: column 1 = dietary category, column 2 = biomass
dataframe2 a data frame of two columns: column 1 = dietary category, column 2 = biomass

Details

Computes the Pianka's index of niche overlap

Value

Return the Pianka's index

References

Pianka R.D. 1973 The structure of lizard communities. Annual Review of Ecology and Systematics, 4: 53-74.

Amroun M., Giraudoux P., Delattre P. 2006 Comparative study of the diets of two sympatric carnivores - the Jackal (*Canis aureus*) and the Genet (*Genetta genetta*) - at two sites in Kabylia, Algeria. Mammalia, 70 (3): 247-254

See Also

[piankabioboot](#)

Examples

```
data(preymbiom)
attach(preymbiom)
jackal<-preymbiom[site=="Y" & sp=="C",5:6]
genet<-preymbiom[site=="Y" & sp=="G",5:6]

piankabio(jackal,genet)
```

piankabioboot	<i>Bootstrap Pianka's index</i>
---------------	---------------------------------

Description

Bootstrap Pianka's index and return the limits of the empirical confidence interval specified with probs

Usage

```
piankabioboot(dataframe1, dataframe2, B = 1000, probs = c(0.025, 0.975))
```

Arguments

dataframe1	a data frame of two columns: column 1 = dietary category, column 2 = biomass
dataframe2	a data frame of two columns: column 1 = dietary category, column 2 = biomass
B	number of permutations
probs	the limits of the confidence interval

Details

Bootstrap Pianka's index and return the limits of the empirical confidence interval specified with probs

Value

a vector of the two CI limits

Author(s)

Patrick Giraudoux <pgiraudou@univ-fcomte.fr>

See Also

[piankabio](#)

Examples

```
data(preymbiom)
attach(preymbiom)
jackal<-preymbiom[site=="Y" & sp=="C",5:6]
genet<-preymbiom[site=="Y" & sp=="G",5:6]

piankabioboot(jackal,genet,B=100)
```

polycirc	<i>Computes the polygon coordinates of a circle</i>
----------	---

Description

Computes the polygon coordinates of a circle

Usage

```
polycirc(radius, pts = c(0, 0), nbr = 50)
```

Arguments

radius	the length of the radius.
pts	the coordinates of the center.
nbr	the number of segments required to draw the perimeter

Details

The matrix of coordinates can then be used with the function `polygon`

Value

A matrix of coordinates.

See Also

[polygon](#), [findR](#)

Examples

```
plot(1:10, 1:10, type="n", asp=1)
polygon(polycirc(5), col="blue")
polygon(polycirc(2, c(5, 5)), col="red")
```

`polycirc2`*Computes the polygon coordinates of a circle sector*

Description

Computes the polygon coordinates of a circle sector

Usage

```
polycirc2(radius = 1, center = c(0, 0), edges = 50, init = pi/2, angle = pi/2)
```

Arguments

<code>radius</code>	the circle radius
<code>center</code>	the centre coordinates (default to x=0, y=0)
<code>edges</code>	the circular outline of the sector is approximated by a polygon with this many edges
<code>init</code>	number (in radian) specifying the starting angle
<code>angle</code>	number (in radian) specifying the sector angle

Details

The matrix of coordinates obtained is intended to be passed to the function [polygon](#)

Value

A matrix of coordinates

See Also

[polygon](#), [polycirc](#), [floating.pie](#)

Examples

```
plot(c(-1,+1),c(-1,+1),type="n",asp=1)
polygon(polycirc2(),col="red")
polygon(polycirc2(init=pi,angle=pi/4),col="green")
polygon(polycirc2(init=1.5*pi,angle=pi/4),col="violet")
polygon(polycirc2(radius=0.5,center=c(0.5,1)),col="blue")

polycirc2(init=pi,angle=pi/4)
```

postxt	<i>Computes coordinates defined from their relative position on x and y in the plotting region</i>
--------	--

Description

Computes coordinates defined from their relative position on x and y in the plotting region

Usage

```
postxt(cd = "ul")
```

Arguments

cd	a numerical vector of length 2, values comprised between 0 and 1, or one predefined among "ul", "bl", "ur", "br", "uc", "bc", "ml", "mc", "mr"
----	--

Details

The argument `cd` gives the relative position to be computed in ratio of the x or y axis. For instance `c(0.025,0.985)` means 2.5 percents on the maximum range of the plot region on x, and 98.5 percents on y (means: close to the upper left corner of the plotting region). Predefined positions are available: "ul", upper left, "bl" bottom left, "ur" upper right, "br" bottom right", "uc" upper center, "bc" bottom center", "ml" medium left, "mc" medium center, "mr" medium right

Value

A list:

x	coordinate on x
y	coordinate on y

Author(s)

Patrick Giraudoux, patrick.giraudoux@univ-fcomte.fr

See Also

[text](#)

Examples

```
plot(rnorm(30), rnorm(30), type="n")
text(postxt("ul"), "here", pos=4)
text(postxt("ur"), "here again", pos=2)
text(postxt("bc"), "again and again")
```

```
preybiom
```

Jackal and Genet diet in Algeria

Description

This data set gives the results of dietary analysis performed by Mansour Amroun in two sites of Kabylie, Algeria

Usage

```
data(preymiom)
```

Format

A data frame with 2196 observations on the following variables.

faeces a factor for faeces corresponding to faeces identification numbers

site a factor for study sites with levels S Sebaou Y Yacouren

saison a factor for seasons with levels H HD HP S SD SP

sp a factor for species with levels C Jackal G Genet

category a factor for dietary items with levels dech ind ins mam mol oisauv oisdome rept vege
vegn

biomasse a numeric vector for the weight of each dietary item

References

M. Amroun, P. Giraudoux and P. Delattre 2006 Comparative study of the diets of two sympatric carnivores - the Jackal (*Canis aureus*) and the Genet (*Genetta genetta*) - at two sites in Kabylia, Algeria. *Mammalia*, 70 (3/4): 247-254.

```
print.mc
```

print method for objects of class 'mc'

Description

print method for objects of class 'mc'

Usage

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

Arguments

x an object of class 'mc'
 ... further arguments to be passed to or from other methods. They are ignored in this function

See Also

[kruskalmc](#), [friedmanmc](#)

Examples

```
resp<-c(0.44,0.44,0.54,0.32,0.21,0.28,0.7,0.77,0.48,0.64,0.71,0.75,0.8,0.76,0.34,0.80,0.73,0.8)
categ<-as.factor(rep(c("A","B","C"),times=1,each=6))
kruskalmc(resp, categ)
```

QGIS2sp

Changes a copied-to-clipboard QGIS attribute table into a sp Spatial object or a data.frame

Description

Reads from the clipboard a copy of the QGIS attribute table of a spatial object (points, linestrings, polygons) and convert it into a [Spatial-class](#) or a [data.frame](#).

Usage

```
QGIS2sp(df=FALSE)
```

Arguments

df If TRUE a data.frame is generated with the coordinates of each shape centroid

Value

A [SpatialPointsDataFrame](#), [SpatialLinesDataFrame](#) or [SpatialPolygonsDataFrame](#) (df=FALSE) or a [data.frame](#) (df=TRUE) with the two first columns corresponding to the centroid coordinates.

Note

Here, [Spatial-class](#) are generated without projection attributes (Coordinate Reference System). CRS, if requested, should be added 'manually' (see [proj4string](#) and [CRS](#)).

See Also

[readWKT](#), [read.delim](#) with the argument "clipboard", [proj4string](#), [CRS](#)

Examples

```
if((.Platform$OS.type == "windows") & (interactive())) {  
  db <-c("wkt_geom\name", "POINT(104.55 34.60)\tDENG_LING", "POINT(104.45 34.49)\tDIAO_GOU")  
  writeLines(db, "clipboard")  
  QGIS2sp() # to write in the console by hand (if copied and paste, one overwrites the clipboard)  
}
```

readGDALbbox

Read a raster using rgdal within a user specified bounding box

Description

Read a raster using rgdal within a user specified bounding box

Usage

```
readGDALbbox(gdal, spo, mar,...)
```

Arguments

gdal	any raster that can be read by readGDAL
spo	spatial object whose bounding box can be retrieved using bbox
mar	user defined margin around the bounding box (default = 2 pixels)
...	further parameters to pass to readGDAL

Details

This function read a raster file using GDAL within the bounding box of a spatial objet. This permits to extract required subset areas from very large raster data sets that cannot be loaded into the workspace.

Value

returns the required data subset from the raster file as a Spatial object

See Also

[readGDAL](#), [bbox](#)

 readVista

Download waypoints and tracks from a GPS

Description

Download GPS waypoints and tracks using gpsbabel

Usage

```
readVista(i = "garmin", f = "usb:", type="w", SPDF=NULL, invisible=TRUE)
```

Arguments

i	INTYPE: a supported file type, default "garmin"
f	INFILE: the appropriate device interface, default "usb:"
type	"w" waypoints, or "t" track
SPDF	if not NULL (the default), characters: the path and filename where to download data as gpx file
invisible	Under Windows, do not open an extra window

Details

The function calls gpsbabel via the system. The gpsbabel program must be present and on the user's PATH for the function to work see <http://www.gpsbabel.org>. The function has been tested on the following Garmin GPS devices: Etrex Summit, Etrex Vista Cx and GPSmap 60CSx. On Ubuntu Linux, USB-to-RS232 converter cables were connected successfully with "/dev/ttyUSB0"; on Windows commonly "com4:" or similar.

Value

If SPDF = NULL (the default) a data frame of four columns:

ident	waypoint names or track IDs
long	longitude
lat	latitude
altitude	elevation

Information about the data type (waypoints or tracks) and the date of download are stored as attributes.

If a path and filename is specified with the argument SPDF (e.g. SPDF="/mydata.gpx"), GPS data are downloaded as gpx file.

References

<http://www.gpsbabel.org>

See Also[readGPS](#)**Examples**

```
## Not run:  
# a GPS device must be connected  
mywaypoints<-readVista() # download waypoints  
mytracks<-readVista(type="t") # download tracks  
  
## End(Not run)
```

rmls

Select objects in the parent frame and remove them.

Description

Select objects in the parent frame and remove them.

Usage

```
rmls()
```

Details

This function has no arguments. This brings up a modal dialog box with a (scrollable) list of objects available in the parent frame. They can be selected by the mouse and then removed.

See Also

ls, rm

Examples

```
toremove<-NULL  
ls()  
if(interactive()) rmls() # select the object 'toremove' and click OK  
ls()
```

rwhatbufCat	<i>Analyses the contents of a <code>SpatialPixelsDataFrame</code> or a <code>SpatialGridDataFrame</code> of categorical values within various buffer sizes centred on points</i>
-------------	--

Description

Analyses the contents of a `SpatialPixelsDataFrame` or a `SpatialGridDataFrame` of categorical values within various buffer sizes centred on points

Usage

```
rwhatbufCat(rast, sites, bufsizes, att=1)
```

Arguments

rast	object of class <code>SpatialPixelsDataFrame</code> or <code>SpatialGridDataFrame</code> to analyse
sites	object of class inheriting from <code>SpatialPoints</code> containing the points on which buffers must be centered
bufsizes	a vector of buffer radii, e.g. <code>c(500, 1000, 1500)</code>
att	column number of the attribute variable

Details

This function generates a data.frame with the frequency of each category of a raster map within various radius buffers centered on point sites.

Value

A dataframe, with the buffer size as first column, the site ID as second column. The other columns are the pixel frequency of each category

See Also

[over](#), [rwhatbufNum](#), [rwhatbufCat2](#)

Examples

```
# raster creation
library(sp)
data(meuse.grid)
coordinates(meuse.grid) = ~x+y
gridded(meuse.grid) = TRUE

# random selection of points within the raster area
mypoints<-spsample(meuse.grid,n=10,type="random") # random points are appx 10, see spsample doc
```



```

mypoints<-SpatialPointsDataFrame(coordinates(mypoints),data.frame(id=1:nrow(mypoints@coords)))

image(meuse.grid,att=4,col=c("red","green","blue")) # soil map
plot(mypoints,add=TRUE) # points
# get the number of pixels of each category in each buffer
rwhatbufCat(meuse.grid,mypoints,c(500,1000),att=4)

```

rwhatbufCat2	<i>Analyses the contents of a raster file readable with rgdal of categorical values within various buffer sizes centred on points</i>
--------------	---

Description

Analyses the contents of a raster file readable with rgdal of categorical values within various buffer sizes centred on points

Usage

```
rwhatbufCat2(rast, sites, bufsizes, att=1, asList=FALSE)
```

Arguments

rast	name of the raster file readable with rgdal to analyse
sites	object of class inheriting from <code>SpatialPoints</code> containing the points on which buffers must be centered
bufsizes	a vector of buffer radii, e.g. <code>c(500, 1000, 1500)</code>
att	column number of the attribute variable
asList	if TRUE the output is a list else a data.frame (default)

Details

This function does not load the full raster file into the memory but loads sequentially subsets corresponding to the size of each buffer. This allows proceeding massive rasters that cannot be loaded into RAM in full. It generates either a data.frame or a list of lists giving for each buffer size (top level of the list of lists) the number of pixels of each category value within the buffer at each point site.

The function reads the raster file on the hard disk as many times as buffers to compute. Thus, computation time is about 5 times longer than `rwhatbufCat`. Empty buffer (no pixel inside) gives (so far) unavoidable topology and dimension errors at reading and stop computation. This can be avoided adjusting buffer size so that the smaller buffer size includes at least one pixel in every position.

Value

A data.frame or a list of lists giving for each buffer size (top level of the list of lists) the number of pixels of each category value within the buffer at each point site

See Also

[over](#), [rwhatbufNum](#), [rwhatbufCat](#)

Examples

```
library(sp)

myrastername<-system.file("pictures/SP27GTIF.TIF", package = "rgdal")[1]

mylocations<-structure(list(x = c(694728, 684662, 702339, 691819, 700091),
y = c(1906654, 1886491, 1884426, 1884373, 1886872)),
.Names = c("x", "y"), row.names = c(NA, -5L), class = "data.frame")

coordinates(mylocations)<-~x+y

result<-rwhatbufCat2(myrastername,mylocations,c(500,1000))
result

result<-rwhatbufCat2(myrastername,mylocations,c(500,1000),asList=TRUE)
result[[1]] # results for buffer 500 (5 buffer centers)
result[[2]] # results for buffer 1000 (5 buffer centers)
```

rwhatbufNum	<i>Analyses the contents of a <code>SpatialPixelsDataFrame</code> or a <code>SpatialGridDataFrame</code> of numerical values within various buffer sizes centred on points</i>
-------------	--

Description

Analyses the contents of a `SpatialPixelsDataFrame` or a `SpatialGridDataFrame` of numerical values within various buffer sizes centred on points

Usage

```
rwhatbufNum(rast, sites, bufsizes, att=1)
```

Arguments

rast	object of class <code>SpatialPixelsDataFrame</code> or <code>SpatialGridDataFrame</code> to analyse
sites	object of class <code>SpatialPointsDataFrame</code> containing the points on which buffers must be centered
bufsizes	a vector of buffer radii, e.g. <code>c(500, 1000, 1500)</code>
att	column number of the attribute variable

Details

This function generates a list of lists giving for each buffer size (top level in the list of lists) the values of the raster map for each point site within the buffer.

Value

A list of lists: top level, the buffer size; second level, the values of the raster map for each point site within the buffer

See Also

[over](#), [rwhatbufCat](#), [rwhatbufCat2](#)

Examples

```
library(pgirmess)
# raster creation
library(sp)
data(meuse.grid)
coordinates(meuse.grid) = ~x+y
gridded(meuse.grid) = TRUE

# random selection of points within the raster area
mypoints<-spsample(meuse.grid,n=10,type="random") # random points are appx 10, see spsample doc
mypoints<-SpatialPointsDataFrame(coordinates(mypoints),data.frame(id=1:nrow(mypoints@coords)))

image(meuse.grid,att=3) # distance to the river
plot(mypoints,add=TRUE,pch=20,cex=0.1) # points
for (i in 1:nrow(mypoints@coords)){
  polygon(polycirc(50, mypoints@coords[i,]),border="blue") # buffer 50 place
}
for (i in 1:nrow(mypoints@coords)) {
  polygon(polycirc(100, mypoints@coords[i,]),border="green") # buffer 100 place
}
mybuffers<-rwhatbufNum(meuse.grid,mypoints,c(50,100),att=3) # get the values in each buffer

names(mybuffers) # two list given

mybuffers[[1]] # list of values for each point (buffer 50)
mybuffers[[1]][[1]] # list of values for the first buffer 50 (point #1)
```

Segments

Draw line segments between pairs of points.

Description

Draw line segments between pairs of points from a vector, matrix or data frame of 4 points coordinates x0, y0, x1, y1

Usage

```
Segments(mydata, ...)
```

Arguments

```
mydata      a vector, matrix or data frame
...         further graphical parameters (from 'par')
```

Details

a wrapper to 'segments' to handle coordinates passed as vector, matrix or data frame. Any vector is turned into a matrix of four columns.

See Also

[segments](#)

Examples

```
mydata<-cbind(rnorm(20),rnorm(20),rnorm(20),rnorm(20))
plot(range(rbind(mydata[,1],mydata[,3])),range(rbind(mydata[,2],mydata[,4])),
type="n",xlab="",ylab="")
Segments(mydata,col=rainbow(20))

myvec<-rnorm(4)
plot(myvec[c(1,3)],myvec[c(2,4)],type="n",xlab="",ylab="")
Segments(myvec)

myvec<-rnorm(16)
plot(myvec,myvec,type="n",xlab="",ylab="")
Segments(myvec)
```

```
selMod
```

Model selection according to information theoretic methods

Description

Handles lm, glm and list of e.g. lm, glm, nls, lme and nlme objects and provides parameters to compare models according to Anderson et al. (2001)

Usage

```
selMod(aModel, Order = "AICc", ...)

## S3 method for class 'lm'
selMod(aModel, Order = "AICc", dropNull = FALSE, selconv=TRUE, ...)
## S3 method for class 'list'
selMod(aModel, Order = "AICc", ...)
```

Arguments

aModel	a lm or glm model or a list of relevant models (see details)
dropNull	if TRUE, drops the simplest model (e.g. y 1)
Order	if set to "AICc" (default) sort the models on this parameter, otherwise "AIC" is allowed
selconv	if TRUE (default) keep the models for which convergence is obtained (glm object only) and with no anova singularity (lm and glm)
...	other parameters to be passed as arguments (not used here)

Details

This function provides parameters used in the information theoretic methods for model comparisons.

- .lm and glm objects can be passed directly as the upper scope of term addition (all terms added). Every model from $y \sim 1$ is computed adding one term at a time until the upper scope model is derived. This is a stepwise analysis where the terms are added sequentially and this does NOT provide all combinations of terms and interactions. Offset terms cannot be proceeded here.
- .A list of user specified lm, glm, nls, lme or nlme objects (actually any object for which AIC and logLik functions are applicable) to compare can alternately be passed.

Value

A dataframe including:

LL	the maximized log-likelihood
K	the number of estimated parameters
n2K	the number of observations/K
AIC	the Akaike index criterion
deltAIC	the difference between AIC and the lowest AIC value
w_i	the Akaike weights
deltAICc	the difference between AICc and the lowest AICc value; advised to be used when $n2K < 40$
w_ic	the AICc weights

The models examined from first to last are stored as attribute

Author(s)

Patrick Giraudoux and David Pleydell: pgiraudou@univ-fcomte.fr, dpleydel@univ-fcomte.fr

References

- Anderson, D.R., Link, W.A., Johnson, D.H. and Burnham, K.P. (2001). Suggestions for presenting the results of data analyses. *Journal of Wildlife Management*, 65, 373-378
- Burnham, K.P. and Anderson, D.R. (2002) *Model Selection and Multimodel Inference: a Practical Information-Theoretic Approach*, 2nd edn., Springer-Verlag, New York. 353 pp

See Also

[AIC](#), [logLik](#), [aictab](#)

Examples

```
library(MASS)
anorex.1 <- lm(Postwt ~ Prewt*Treat, data = anorexia)
selMod(anorex.1)
anorex.2 <- glm(Postwt ~ Prewt*Treat, family=gaussian,data = anorexia)
selMod(anorex.2)
anorex.3<-lm(Postwt ~ Prewt+Treat, data = anorexia)
mycomp<-selMod(list(anorex.1,anorex.2,anorex.3))
mycomp
attributes(mycomp)$models
```

shannon

Computes Shannon's and equitability indices

Description

Computes Shannon's and equitability indices

Usage

```
shannon(vect, base=2)
```

Arguments

vect	a probability vector whose sum = 1 or a frequency vector
base	logarithm base used (default=2)

Details

Computes Shannon's and equitability indices. The vector passed can be a probability vector whose sum equal 1 or a vector of frequencies (e.g. the number of food item of each category).

Value

A vector of two values: Shannon's and equitability indices. The base logarithm used is stored as attribute

See Also[shannonbio](#)**Examples**

```
x<-c(0.1,0.5,0.2,0.1,0.1)
sum(x)
shannon(x)
```

```
x<-rpois(10,6)
shannon(x, base=exp(1))
```

shannonbio	<i>Computes Shannon's and equitability indices from a data frame of dietary analysis (n, biomass,...)</i>
------------	---

Description

Computes Shannon's and equitability indices from a data frame of two columns: column 1, dietary category; column 2, abundance (n, biomass,...)

Usage

```
shannonbio(data1)
```

Arguments

data1	a data frame of two columns: column 1, dietary category; column 2, abundance (n, biomass,...)
-------	---

Details

Computes Shannon's and equitability indices from a data frame of two columns: column 1, dietary category; column 2, abundance (n, biomass,...)

Value

A vector of two values: Shannon's and equitability indices

Author(s)

Patrick Giraudoux <pgiraud@univ-fcomte.fr>

See Also[shannon](#), [difshannonbio](#)

Examples

```
data(preymbiom)
shannonbio(preymbiom[,5:6])
```

shannonbioboot	<i>Bootstrap Shannon's and equitability indices</i>
----------------	---

Description

Bootstrap Shannon's and equitability indices and return an object of class boot. Confidence intervals can be computed with `boot.ci()`.

Usage

```
shannonbioboot(data1, B = 1000)
```

Arguments

data1	a data frame of two columns: column 1, dietary category; column 2, abundance (n, biomass,...)
B	number of permutations

Details

Bootstrap Shannon's and equitability indices and return an object of class boot. Confidence intervals can be computed with `boot.ci()`. Requires the boot library.

Value

An object of class boot including the bootstrap statistics for H' (t1*) and J' (t2*)

Author(s)

Patrick Giraudoux <pgiraudou@univ-fcomte.fr

See Also

[boot](#), [boot.ci](#), [shannonbio](#)

Examples

```
data(preymbiom)
myboot<-shannonbioboot(preymbiom[,5:6],B=100)
library(boot)
boot.ci(myboot, index=1,type=c("norm","basic","perc")) # confidence intervals for H'
boot.ci(myboot, index=2,type=c("norm","basic","perc")) # confidence intervals for J'
```

 siegelp179

Data on rats training

Description

Ranks of 18 matched groups of rats after training under three methods of reinforcement.

Usage

```
data(siegelp179)
```

Format

A data frame with 54 observations on the following 3 variables.

block Group (each of three litter mates)

treatment A factor for the type of reinforcement with levels RR RU UR

score Speed of transfer to another behaviour (the lower, the better the learning)

Details

18 blocks made of three rats of the same litter, each being given a different learning pattern (RR, RU or UR)

Source

Grosslight J.H. and Radlow R. (1956) Patterning effect of the nonreinforcement-reinforcement sequence in a discrimination situation. *Journal of Comparative and Physiological Psychology*, 49: 542-546 in Siegel & Castellan 1988. *Non parametric statistics for the behavioural sciences*. Mc Graw Hill Int. Edt.

Examples

```
data(siegelp179)
```

 tabcont2categ

Convert a contingency table (data.frame) into a presence/absence table of categories

Description

Convert a contingency table (data frame) into a data.frame of factors

Usage

```
tabcont2categ(tab)
```

Arguments

tab A data.frame (contingency table)

Details

Convert a contingency table (data frame) into a data.frame of factors

Value

A data frame

Author(s)

Patrick Giraudoux <pgiraud@univ-fcomte.fr>

Examples

```
mydata<-as.data.frame(matrix(rpois(9,5),nr=3,nc=3))
names(mydata)<-LETTERS[1:3]
row.names(mydata)<-letters[1:3]

tabcont2categ(mydata)
```

thintrack *Thin a track just keeping the points separated by a user defined minimal distance*

Description

Thin a track stored as a [SpatialPointsDataFrame](#) object, just keeping the points separated by a user defined minimal distance.

Usage

```
thintrack(spdf, mindist=100)
```

Arguments

spdf a [SpatialPointsDataFrame](#) of point tracks
mindist minimal distance requested between two points (default = 100)

Details

Tracks downloaded from GPS often provide an unnecessary large density of points at irregular distances. This function starts reading from the first point of the track and removes all points within a user specified radius (USR), then reads the closest point and removes all points within the USR, and so on...

Value

A [SpatialPoints](#) object of the track thinned.

See Also

[mergeTrackObs](#)

Examples

```
library(sp)
mySPDF<-structure(list(x = c(748775, 748807, 748834, 748854, 748871,
748873, 748880, 748890, 748919, 748917, 748921, 748923, 748924,
748921, 748921, 748921, 748922, 748915, 748616, 748613, 748612,
748613, 748613, 748615, 748613, 748616, 748615, 748618, 748615,
748619, 748618, 748620, 748586, 748553, 748494, 748444, 748424,
748366, 748305, 748305), y = c(105716, 105761, 105808, 105856,
105911, 105964, 106019, 106065, 106114, 106167, 106219, 106274,
106329, 106385, 106441, 106494, 106550, 106571, 105835, 105779,
105723, 105665, 105600, 105537, 105473, 105412, 105350, 105293,
105234, 105180, 105123, 105070, 105023, 104960, 104956, 104947,
104906, 104905, 104901, 104904), ID = 1:40), .Names = c("x",
"y", "ID"), row.names = c("1", "2", "3", "4", "5", "6", "7",
"8", "9", "10", "11", "12", "13", "14", "15", "16", "17", "18",
"19", "20", "21", "22", "23", "24", "25", "26", "27", "28", "29",
"30", "31", "32", "33", "34", "35", "36", "37", "38", "39", "40"
), class = "data.frame")
coordinates(mySPDF)<-~x+y

plot(mySPDF,pch=19,cex=0.5)
plot(thintrack(mySPDF),pch=19,cex=0.7,col="red",add=TRUE)

plot(mySPDF,pch=19,cex=0.5)
plot(thintrack(mySPDF,min=200),pch=19,cex=0.7,col="red",add=TRUE)
```

trans2pix

Convert a transect coordinate file with some landmarks into a matrix with intermediate coordinates.

Description

Convert a transect coordinate file with some landmarks and NA values in between into a matrix with intermediate coordinates.

Usage

```
trans2pix(vect)
```

Arguments

vect A two column matrix or data.frame

Details

If vect has more than two column the two first column only are read. This function computes the intermediate coordinates of each lines materialised with NA values.

Value

A matrix with the intermediate coordinates computed.

See Also

[trans2seg](#)

Examples

```
x<-c(10,NA, NA, NA,56,NA,NA,100)
y<-c(23,NA, NA, NA,32,NA,NA,150)
cols=c("red", "blue", "blue", "blue", "red", "blue", "blue", "red")
plot(x,y,col=cols,pch=19)
plot(trans2pix(cbind(x,y)),col=cols,pch=19)
```

trans2seg	<i>Convert a transect coordinate file into a matrix with segment coordinates.</i>
-----------	---

Description

Convert a transect coordinate file (eg: landmarks) into a matrix with segment coordinates.

Usage

```
trans2seg(vect)
```

Arguments

vect A two column matrix or data.frame

Details

The argument passed is a matrix or data.frame of two columns each row is a transect interval; each column must start (first row) and end (last row) with a landmark ; intermediate landmarks must have coordinates in the two columns of the row. Other rows must be NA values.

Value

A matrix of 4 columns to be passed eg to fonctions as "segments".

See Also

trans2pix

Examples

```
x<-c(10,NA, NA, NA,56,NA,NA,100)
y<-c(23,NA, NA, NA,32,NA,NA,150)
cols=c("red","blue","blue","blue","red","blue","blue","red")
plot(x,y,col=cols,pch=19)
mysegs<-trans2seg(cbind(x,y))
segments(mysegs[,1],mysegs[,2],mysegs[,3],mysegs[,4])
```

transLines2pix	<i>Convert a SpatialLines or a SpatialLinesDataFrame object into SpatialPointsDataFrame with points at regular distance along the lines</i>
----------------	---

Description

Convert a SpatialLines or a SpatialLinesDataFrame object into SptaiialPointsDataFrame with points at regular distance along the lines

Usage

```
transLines2pix(spldf,mindist=100)
```

Arguments

spldf	A SpatialLines or a SpatialLinesDataFrame
mindist	the distance between two points (default to 100)

Details

This function can be used e.g to discretize any track line (roads, paths, transects, etc.) into series of regular points. Each point may be though of as corresponding to the centre of one interval

Value

A [SpatialPointsDataFrame](#)

See Also

[trans2pix](#), [thintrack](#), [mergeTrackObs](#)

Examples

```
# from the sp vignette:
library(sp)
l1 = cbind(c(1,2,3),c(3,2,2))
l1a = cbind(l1[,1]+.05,l1[,2]+.05)
l2 = cbind(c(1,2,3),c(1,1.5,1))
S11 = Line(l1)
S11a = Line(l1a)
S12 = Line(l2)
S1 = Lines(list(S11, S11a), ID="a")
S2 = Lines(list(S12), ID="b")
S1 = SpatialLines(list(S1,S2))
plot(S1, col = c("red", "blue"))

trpt<-transLines2pix(S1,mindist=0.1)
plot(trpt,add=TRUE)
```

TukeyHSDs

Simplify the list of a TukeyHSD object keeping the significant differences only.

Description

Simplify the list of a TukeyHSD object keeping the significant differences only.

Usage

```
TukeyHSDs(TukeyHSD.object)
```

Arguments

TukeyHSD.object
An object of calls "TukeyHSD"

Details

When TukeyHSD is used on a fitted model with large numbers of categories, the number of pairwise comparisons is extremely large ($n(n-1)/2$). TukeyHSDs simplify the TukeyHSD object keeping the significant pairwise comparisons only. A plot method exists for TukeyHSD objects.

Value

An object of class "multicomp" and "TukeyHSD"

See Also

[TukeyHSD](#)

Examples

```
summary(fm1 <- aov(breaks ~ wool + tension, data = warpbreaks))
myobject<-TukeyHSD(fm1, "tension", ordered = TRUE)
myobject
TukeyHSDs(myobject)
```

uploadGPS

Upload waypoints to Garmin GPS

Description

Upload waypoints to Garmin GPS, using gpsbabel

Usage

```
uploadGPS(gpx, f = "usb:", type="w")
```

Arguments

gpx	name of the .gpx file (can be created from a data frame using writeGPX)
f	the appropriate device interface, default "usb:", see details
type	'w' for waypoints (default), 't' for track

Details

This function uploads waypoints or a track to a garmin GPS from a '.gpx' file. gpsbabel is called via the system. Therefore gpsbabel must be installed and on the user's path, see <http://www.gpsbabel.org>. If not the default, device interface should be something as "usb:", "usb:1", "com:4" or on linux "/dev/ttyUSB0", etc.

Warning

Overwrite waypoints having the same name in the GPS

See Also

[writeGPX](#)

Examples

```
## Not run:
# a GPS device must be connected
coords<-data.frame(ID=c("C18J01", "C18J02"),Long= c(-46.996602, 47.002745),
Lat=c(-6.148734, 6.14829),Alt=c(250,1230))

writeGPX(coords,"mywaypoints")
uploadGPS("mywaypoint.gpx")

## End(Not run)
```

val4symb	<i>Centres a numerical vector on a parameter position and provides absolute values and colors according to negative and positive values</i>
----------	---

Description

Centres a numerical vector on a parameter position and provides absolute values and colors according to negative and positive values

Usage

```
val4symb(x, FUN=mean, col = c("blue", "red"),...)
```

Arguments

x	a numerical vector
FUN	a function computing a position parameter, typically mean or median . Default to mean
col	a character vector of 2 values, default=c("blue","red"), blue for <0, red for >=0
...	optional arguments to 'FUN'

Value

A list with	
size	the absolute values of the difference to the position parameter (eg mean, median)
col	a character vector with 2 colors, each corresponding to positive or negative values

Author(s)

Patrick Giraudoux, pgiraudou@univ-fcomte.fr

See Also

[symbols](#), [mean](#), [median](#), [scale](#)

Examples

```
x<-rnorm(30)
y<-rnorm(30)

z<-val4symb(rnorm(30))
symbols(x,y, circle=z$size, inches=0.2, bg=z$col)

z<-val4symb(scale(rnorm(30)))
symbols(x,y, circle=z$size, inches=0.2, bg=z$col)

z<-val4symb(rnorm(30), col=c("green", "violet"))
symbols(x,y, circle=z$size, inches=0.2, bg=z$col)

z<-val4symb(rnorm(30), trim=0.025)
symbols(x,y, circle=z$size, inches=0.2, bg=z$col)

z<-val4symb(rnorm(30), median)
symbols(x,y, circle=z$size, inches=0.2, bg=z$col)

myfun<-function(x) 20 # passes an arbitrary constant
z<-val4symb(1:30, myfun)
symbols(x,y, circle=z$size, inches=0.2, bg=z$col)
```

valchisq

Values of the partial chi-square in each cell of a contingency table

Description

Computes the values of the partial chi-square in each cell of a contingency table

Usage

```
valchisq(matr)
```

Arguments

matr a matrix (contingency table)

Details

Computes the values of the chi-square in each cell of a contingency table

Value

A matrix with the chi-square values computed

Note

No correction (e.g. Yate's etc.) is done !

See Also

valat, chisq.test

Examples

```
x <- matrix(c(12, 5, 7, 7), nc = 2)
x
valchisq(x)
```

write.delim

Write a data.frame

Description

Write a simple data.frame into a text file with header, no row.names, fields separated by tab.

Usage

```
write.delim(x, file = "", row.names = FALSE, quote = FALSE, sep = "\t", ...)
```

Arguments

x	a data.frame
file	a character string for file name
row.names	either a logical value indicating whether the row names of 'x' are to be written along with 'x', or a character vector of row names to be written
quote	a logical value or a numeric vector. If 'TRUE', any character or factor columns will be surrounded by double quotes. If a numeric vector, its elements are taken as the indices of the columns to quote. In both cases, row and column names are quoted if they are written. If 'FALSE', nothing is quoted.
sep	the field separator string. Values within each row of 'x' are separated by this string.
...	additional arguments accepted by write.table

Details

Simple wrapper of write.table.

Value

An ascii text file, tab delimited.

Author(s)

Patrick Giraudoux <pgiraud@univ-fcomte.fr>

See Also

[write.table](#)

Examples

```
data(prebybiom)
write.delim(prebybiom[1:10,]) # output to the console
write.delim(prebybiom[1:10,],file="Myfile.txt") # write a file in the working directory
```

writeGPX

Convert a data frame into a GPX file of waypoints or track

Description

Convert a data frame of labels, geographical coordinates and optionally altitude into a GPX file of waypoints or track that can be uploaded to Garmin GPS

Usage

```
writeGPX(x, filename = "", type="w")
```

Arguments

x	data.frame of three (optionally four) columns (see details)
filename	a character string naming the file to print to. If '''' (the default), prints to the standard output connection, the console (unless redirected by 'sink')
type	'w' for waypoints (default) or 't' for track

Details

The data frame must have three (optionally four) columns:

1. character or integer, waypoint ID for waypoints ; column not read for track
2. numeric, longitude (decimal degrees), negative for west
3. numeric, latitude (decimal degrees), negative for south
4. numeric, elevation (meters) (optional)

A suffix '.gpx' is added to the file name if not provided by user. The file obtained can be uploaded to Garmin GPS but cannot be read eg from MapSource for some reasons.

Note

for more standard GPX file, see [writeOGR](#) with arguments like `layer="waypoints"`, `driver="GPX"`, `dataset_options="GPX_USE_EXTENSIONS=yes"` can alternately be used; [readOGR](#) with arguments like `layer="waypoints"`, `drop_unsupported_fields=TRUE`

See Also

[writeOGR](#)

Examples

```
coords<-data.frame(ID=c("C18J01", "C18J02"),Long= c(-46.996602, 47.002745),
Lat=c(-6.148734, 6.14829),Alt=c(250,1230))

writeGPX(coords) # waypoints
writeGPX(coords,type="t") # track
```

writePRJ

Write the projection file of a shapefile from a spatial object

Description

Write the projection file of a shapefile from a spatial object

Usage

```
writePRJ(spobj, filename)
```

Arguments

spobj	any spatial object having a CRS extractible with proj4string
filename	a character string naming the file to print to. If <code>''</code> (the default), prints to the standard output connection, the console (unless redirected by <code>'sink'</code>)

Details

A suffix `'.prj'` is added to the file name if not user provided.

Examples

```
library(sp)
mypoints<-data.frame(long=runif(10,-90,+90),lat=runif(10,-90,+90))
coordinates(mypoints)<--long+lat # SpatialPoints object
proj4string(mypoints)<-CRS("+proj=longlat +ellps=WGS84 +datum=WGS84") # WGS84 coordinates
writePRJ(mypoints,"")
```

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