# Package 'DeadCanMove'

March 28, 2017

Type Package

<b>Title</b> Assess How Spatial Roadkill Patterns Change with Temporal Sampling Scheme
Version 0.5
<b>Date</b> 2017-03-28
Author Barbosa A.M., Marques J.T., Santos S.M., Lourenco A., Medinas D., Beja P., Mira A.
Maintainer A. Marcia Barbosa  barbosa@uevora.pt>
Suggests carcass
<ul> <li>Description From a baseline data frame of dead individuals recorded daily at different road stretches, simulate varying sub-sampling schemes, calculate and compare roadkill patterns and hotspots based on each sampling scheme.</li> <li>License GPL-3</li> </ul>
R topics documented:
DeadCanMove-package

 plotEventCorrs
 11

 repl.hs.comp
 12

 roadkills
 13

 schemeCorrs
 14

 sequential.corr
 15

 sequential.estimateN
 17

 sequential.hotspots
 18

 sequential.Nevents
 19

 sequential.posteriorN
 20

 sequential.seqsubmat
 21

sequential.submatrix	 																22
submatrix	 																24

DeadCanMove-package

Assess How Spatial Roadkill Patterns Change with Temporal Sampling Scheme

# **Description**

From a baseline data frame of dead individuals recorded daily at different road stretches, simulate varying sub-sampling schemes, calculate and compare roadkill patterns and hotspots based on each sampling scheme.

#### **Details**

Package: DeadCanMove Type: Package

Version: 0.5

Date: 2017-03-28 License: GPL-3

# Author(s)

Barbosa A.M., Marques J.T., Santos S.M., Lourenco A., Medinas D., Beja P., Mira A.

Maintainer: A. Marcia Barbosa <br/> <br/> darbosa@uevora.pt>

#### References

Santos S.M., Marques J.T., Lourenco A., Medinas D., Barbosa A.M., Beja P., Mira A. (2015) Sampling effects on the identification of roadkill hotspots: implications for survey design. Journal of Environmental Management, 162: 87-95 (DOI: 10.1016/j.jenvman.2015.07.037)

#### See Also

carcass

```
data(roadkills)
hc <- hotspots.comparison(dataset = roadkills,
sampl.columns = 4:ncol(roadkills), sampl.intervals = 1:5,
region.column = "segment", group.column = "taxon",
include.all.together = TRUE, confidence = 0.95,</pre>
```

binary.comp.methods 3

```
min.total.events = 80, min.hotspot.threshold = 2,
comp.method = "Phi", plot = TRUE, sep.plots = FALSE,
omit.baseline.interval = TRUE, ylim = c(0, 1))
hc
```

```
binary.comp.methods
```

Binary comparison methods

# Description

This function provides the methods implemented for calculating binary similarity between hotspots obtained from different sampling schemes and those obtained from the baseline (reference) sampling scheme. It is used by functions binary.comparison, sequential.corr and hotspots.comparison.

# Usage

```
binary.comp.methods()
```

# Value

A character vector naming the methods implemented.

# Author(s)

A. Marcia Barbosa

# See Also

```
binary.comparison, sequential.corr
```

```
binary.comp.methods()
```

4 binary.comparison

```
binary.comparison Binary comparison
```

# Description

Compares two binary vectors using the coefficient specified in method.

# Usage

```
binary.comparison(x, y, method)
```

# **Arguments**

```
x a binary (0-1) vector
y a binary (0-1) vector to compare with x
method the comparison measure to use. Current options are "Phi", "Mathews", "Yule",
"Jaccard", "Baroni", "kappa", "CCR", "TSS", "gain", "loss", and "balance".
```

# Value

A numeric value indicating the cofficient of association specified in method.

# Author(s)

A. Marcia Barbosa

# See Also

```
binary.comp.methods
```

```
bin1 <- sample(c(0, 1), 100, replace = TRUE)
bin2 <- sample(c(0, 1), 100, replace = TRUE)

binary.comparison(bin1, bin2, method = "Phi")
binary.comparison(bin1, bin2, method = "kappa")</pre>
```

getBoxplots 5

getBoxplots

Get boxplots

# **Description**

Get boxplots

# Usage

```
getBoxplots(corrs.list, ...)
```

# Arguments

#### Value

Box plots

# Author(s)

A. Marcia Barbosa

#### See Also

schemeCorrs

hotspot.numbers

Hotspot numbers

# Description

This function calculates the total numbers of events and the hotspot thresholds for each group and sampling interval.

# Usage

```
hotspot.numbers(hotspots.list, sampl.intervals, groups,
include.all.together = TRUE, min.total.events = 0,
min.hotspot.threshold = 2)
```

6 hotspot.numbers

# **Arguments**

```
results of the sequential.hotspots function

sampl.intervals

integer vector of the sampling intervals to analyse (at the moment, these intervals must be consecutive and start with one)

groups

taxa or groups to analyse separately (e.g. as.character(unique(dataset$group)))

include.all.together

logical, whether to run the analysis also for all groups combined

min.total.events

minimum total number of events to calculate hotspots for a group

min.hotspot.threshold

minimum number of events for a region to be considered a hotspot
```

#### Value

A list of the following matrices:

```
N.events
HS.threshold
N.hotspots
events.in.HS
```

#### Note

This function currently works only for hotspots of submats created with 'sampl.interval', not 'window.size' and 'gap.size'. See submatrix, sequential.submatrix, and check that your names(hotspots.list[[1]]) are something like "group.intv1", not "group.w1.g2.s1".

# Author(s)

A. Marcia Barbosa, J. Tiago Marques, Sara M. Santos

# See Also

```
hotspots, sequential.hotspots
```

```
data(roadkills)
submats <- sequential.submatrix(dataset = roadkills,
sampl.columns = 4:ncol(roadkills), sampl.intervals = 1:3,
group.column = "taxon", include.all.together = TRUE,
remove.zeros = TRUE, keep.nonsampl.columns = TRUE,
n.subsampl.columns = 120)
hsl <- sequential.hotspots(dataset = roadkills, submats = submats,</pre>
```

hotspots 7

```
region.column = "segment", first.subsampl.col = 4, confidence = 0.95)
hsn <- hotspot.numbers(hotspots.list = hsl, sampl.intervals = 1:3,
groups = as.character(unique(roadkills$taxon)),
include.all.together = TRUE, min.hotspot.threshold = 2)
hsn</pre>
```

hotspots

Calculate roadkill hotspots

#### **Description**

This function identifies the hotspot regions in a dataset, or in a submatrix compared to the total dataset, using an adaptation of the method of Malo et al. (2004).

# Usage

```
hotspots(dataset, submat = NULL, region.column,
subsampl.columns = NULL, n.events.column = NULL, hotspots = TRUE,
confidence = 0.95, min.total.events = 0, min.hotspot.threshold = 2)
```

#### **Arguments**

dataset name of the matrix or dataframe containing the complete data

submat name of the matrix or dataframe containing the data of the group and sampling

window/gap for which to calculate hotspots

region.column

name or index number of the column containing the regions (road sectors, sites)

to classify as hotspots or non-hotspots

subsampl.columns

index numbers of the consecutive columns of submat (or, if there is no submat,

of the dataset) containing the (daily) sampling data, e.g. 4:180

n.events.column

alternatively to subsampl.columns, the name or index number of the col-

umn containing the number of events (e.g. individual deaths) in each row

hotspots logical, whether to calculate the hotspots

confidence confidence threshold to consider hotspots

min.total.events

minimum total number of events to calculate hotspots. Not totally implemented

yet!

min.hotspot.threshold

minimum number of events for a region to be considered a hotspot. If the Malo method says that regions with less than this value are hotspots, the value returned is NA. The default threshold is 2.

8 hotspots.comparison

#### Value

A list with elements threshold (an integer value indicating the number of deaths obtained as a threshold for considering a site a roadkill hotspot) and hotspots (a data frame showing the total number of deaths per region and whether or not it was considered a hospot.)

#### Author(s)

A. Marcia Barbosa, J. Tiago Marques, Sara M. Santos

## References

Malo, J.E., Suarez, F., Diez, A. (2004) Can we mitigate animal-vehicle accidents using predictive models? J. Appl. Ecol. 41, 701-710 (doi: 10.1111/j.0021-8901.2004.00929.x)

#### See Also

```
sequential.hotspots
```

#### **Examples**

```
data(roadkills)
hs <- hotspots(dataset = roadkills, submat = NULL, region.column = "segment",
subsampl.columns = 4:ncol(roadkills), confidence = 0.95)
hs</pre>
```

```
hotspots.comparison
```

Hotspots comparison

# Description

This is a wrapper for most of the functions in this package (one function to rule them all). You'll probably only need to use this one, which in turn calls each of the other functions and does all the calculations in one step.

#### Usage

```
hotspots.comparison(dataset, sampl.columns, sampl.intervals, region.column, group.column, include.all.together = TRUE, confidence = 0.95, min.total.events = 80, min.hotspot.threshold = 2, comp.method = "Phi", plot = TRUE, sep.plots = FALSE, omit.baseline.interval = TRUE, ...)
```

hotspots.comparison 9

# **Arguments**

dataset name of the matrix or dataframe to analyze sampl.columns index numbers of the columns containing the (daily) sampling data, e.g. 4:180 sampl.intervals intervals at which to extract sampling data, e.g. 1:30; currently must be consecutive and start with 1 region.column name or index number of the column containing the regions (road segments, sites) to classify as hotspots or non-hotspots group.column name or index number of the column containing the taxa or groups to analyse separately, e.g. 3 or "Family"; if NULL, all records will be used together include.all.together logical, whether to get subsampling matrices also for the complete data (including all groups combined) confidence threshold to consider hotspots (see Malo et al. 2004); defaults to confidence 0.95 min.total.events minimum total number of events (e.g. deaths) to calculate hotspots for a group min.hotspot.threshold minimum number of events for a region to be considered a hotspot the method with which to compare the hotspots obtained with increasing sampl.intervals comp.method with those of the baseline scenario; type binary.comp.methods() for available options plot logical, whether to plot the correlations between subsamples and baseline for each group (may cause function to fail if sep.plots = FALSE and figure margins are too large for the number of resulting plots) logical, whether to present the plots in separate windows rather than all in the sep.plots same window omit.baseline.interval logical, whether to omit the first column (correlation of baseline hotspots with themselves) from calculations and results

additional arguments to pass to the plot function

#### Value

# A list with 9 elements:

hotspots.list

N.events
HS.threshold
N.hotspots
events.in.HS
event.corrs

10 jumping.window

```
event.loss
event.gain
event.balance
```

# Author(s)

A. Marcia Barbosa, J. Tiago Marques, Sara M. Santos

#### References

Malo, J.E., Suarez, F., Diez, A. (2004) Can we mitigate animal-vehicle accidents using predictive models? J. Appl. Ecol. 41, 701-710 (doi: 10.1111/j.0021-8901.2004.00929.x)

#### See Also

hotspots

# **Examples**

```
data(roadkills)
hc <- hotspots.comparison(dataset = roadkills,
sampl.columns = 4:ncol(roadkills), sampl.intervals = 1:5,
region.column = "segment", group.column = "taxon",
include.all.together = TRUE, confidence = 0.95,
min.total.events = 80, min.hotspot.threshold = 2,
comp.method = "Phi", plot = TRUE, sep.plots = FALSE,
omit.baseline.interval = TRUE, ylim = c(0, 1))
hc</pre>
```

jumping.window

Jumping window

# **Description**

This function extracts a moving (a.k.a. running, rolling, sliding) window but with no overlap between windows and with the option for gaps between windows.

# Usage

```
jumping.window(sampl.columns, window.size, gap.size,
start.column = 1, J = FALSE)
```

jumping.window 11

# Arguments

sampl.columns	3
	index numbers of the consecutive columns with the sampling data (e.g. 3:180) from which to extract the jumping windows. Can also be any vector from which to extract a jumpting window.
window.size	size of each sampling window/season (consecutive sampling days each time)
gap.size	size of the gap between sampling windows. Can be zero or a positive integer.
start.column	column of sampl.columns where to actually start the sampling windows. The default is 1, but e.g. with a gap size of 1 between windows, the start column can be either 1 or 2.
J	logical, whether to provide the results in the form of J for function ${\tt carcass::etterson}.$ Defaults to FALSE.

# **Details**

This function is now integrated within submatrix to provide for additional sampling scheme options.

# Value

When J = FALSE (the default), this function returns a vector containing the elements of sampl.columns that are included in the extracted windows; elements falling within the gaps are left out.

# Author(s)

A. Marcia Barbosa, J. Tiago Marques

```
data(roadkills)
names(roadkills)
jumping.window(4:ncol(roadkills), window.size = 1, gap.size = 0)
jumping.window(4:ncol(roadkills), window.size = 1, gap.size = 1)
w3g5 <- jumping.window(4:ncol(roadkills), window.size = 3,
gap.size = 5)
w3g5</pre>
```

12 plotEventCorrs

plotEventCorrs

Plot correlations between events in each subsampling dataset and the baseline dataset

#### **Description**

This function plots the correlation with baseline against sampling interval for each group

# Usage

```
plotEventCorrs(event.corrs, sep.plots = FALSE, ...)
```

# **Arguments**

```
event.corrs a matrix of correlations resulting from the sequential.corr function sep.plots logical, whether to place each plot in a separate window additional arguments to pass to the plot function
```

#### Value

This function produces plots.

#### Author(s)

A. Marcia Barbosa

#### See Also

```
plot, binary.comparison, sequential.corr
```

```
data(roadkills)
submats <- sequential.submatrix(dataset = roadkills,
sampl.columns = 4:ncol(roadkills), sampl.intervals = 1:3,
group.column = "taxon", include.all.together = TRUE,
remove.zeros = TRUE, keep.nonsampl.columns = TRUE,
n.subsampl.columns = 80)

names(submats)

hsl <- sequential.hotspots(dataset = roadkills, submats = submats,
region.column = "segment", first.subsampl.col = 4, confidence = 0.95)

hsn <- hotspot.numbers(hotspots.list = hsl, sampl.intervals = 1:3,
groups = as.character(unique(roadkills$taxon)), include.all.together = TRUE,
min.total.events = 0, min.hotspot.threshold = 2)</pre>
```

repl.hs.comp 13

```
seqcorr <- sequential.corr(hotspots.list = hsl,
hotspots.thresholds = hsn$HS.threshold, comp.method = "Phi",
baseline.interval = 1, messages = "TRUE")

plotEventCorrs(event.corrs = seqcorr, sep.plots = FALSE, ylim = c(0, 1),
pch = 20)</pre>
```

repl.hs.comp

Replicate hotspot comparison

# **Description**

This function calculates hotspot correlation, loss, gain or balance for the different replicates per sampling scheme and taxonomic group.

#### Usage

```
repl.hs.comp(seqsubmats.hs, hs.baseline, method = "Phi",
stats = TRUE, plot = TRUE, plot.mean = TRUE, ylim = NULL,
horiz.line = NA)
```

## **Arguments**

seqsubmats.hs hotspots for the seqsubmats hs.baseline hotspots for the baseline method binary comparison method to use. See binary.comparison for available options. stats logical, whether to calculate also the stats (mean, min, max, sd) of the replicate comparison for each group. logical, whether to plot the hotspot comparison values per replicate per group. plot logical, whether to plot (with a white circle) the mean value of the replicates per plot.mean group. ylim limits for the y axis. The default is NULL for automatic limits, but you may want to use ylim = c(0,1) for e.g. phi correlations to be directly comparable among plots. horiz.line optionally, a numeric value indicating the y axis value for a horizontal threshold line to be drawn.

## Value

This function returns a list.

#### Author(s)

A. Marcia Barbosa

14 roadkills

#### See Also

```
binary.comparison
```

## **Examples**

```
## Not run:
replicate.corrs <- repl.hs.comp(seqsubmats.hs = seqsubmats.hs,
hs.baseline = hs.baseline, method = "Phi")

replicate.gains <- repl.hs.comp(seqsubmats.hs = seqsubmats.hs,
hs.baseline = hs.baseline, method = "gain")

## End(Not run)</pre>
```

roadkills

Imaginary roadkill data

#### **Description**

An imaginary dataset of roadkill data for 5 "taxonomic" groups.

# Usage

```
data(roadkills)
```

# **Format**

A data frame with 900 observations on the following variables:

individ an integer vector attributing an identifier to each recorded individual
segment a numeric vector identifying the road segment at which each individual was recorded
group a character vector indicating the "taxonomic" group to which each individual belongs
day1 a numeric vector indicating whether the individual was found (1) or not (0) on that sampling
day (the same for all remaining days in the data frame)

#### **Details**

Each row corresponds to an individual recorded at a particular road stretch (segment), with a 1 if it was present and a 0 if it was not present at that segment on each of the sampling days. Individuals were not manually removed from the road, so each individual has value 1 in all days in which its body was detected on the road.

# **Source**

Freely modified from data collected by: Santos S.M., Carvalho F., Mira A. (2011) How long do the dead survive on the road? Carcass persistence probability and implications for road-kill monitoring surveys. PLoS ONE 6(9), e25383 (doi:10.1371/journal.pone.0025383)

schemeCorrs 15

# **Examples**

```
data(roadkills)
roadkills[1:20, 1:10]
```

schemeCorrs

Scheme correlations

# Description

Get correlation between each sampling scheme and the corresponding baseline

# Usage

```
schemeCorrs(dataset, submats, submats.baseline, region.column,
group.column, first.subsampl.col)
```

# **Arguments**

dataset name of the matrix or dataframe containing the complete data

submats a list of the submatrices for which to calculate the correlation (result of the sequential.submatrix function)

submats.baseline a list of the submatrices corresponding to the baseline sampling scheme for each group

region.column

name or index number of the column containing the regions (road segments, sites) to classify as hotspots or non-hotspots

group.column name or index number of the column containing the taxonomic groups

first.subsampl.col

index number of the first column containing subsampling data

#### Value

This function returns a list of corrs.tables.

# Author(s)

A. Marcia Barbosa

# See Also

```
getBoxplots
```

16 sequential.corr

```
sequential.corr
```

Sequential correlation

# **Description**

This function calculates the correlation between the hotspots obtained from each submatrix and those of the baseline (sub)matrix of the corresponding group.

# Usage

```
sequential.corr(hotspots.list, hotspots.thresholds,
comp.method = "Phi", baseline.interval = 1, baseline.gap = 0,
messages = "TRUE")
```

#### **Arguments**

# Value

A matrix of correlations (or whatever index was defined in method) between the hotspots obtained for each group and sampling scheme, and the hotspots obtained from the baseline data for the group under analysis.

#### Note

This function currently works only for hotspots of submats created with 'sampl.interval', not 'window.size' and 'gap.size'. See submatrix, sequential.submatrix, and check that your names(hotspots.list[[1]]) are something like "group.intv1", not "group.w1.g2.s1".

#### Author(s)

A. Marcia Barbosa

sequential.estimateN 17

#### See Also

```
binary.comparison
```

#### **Examples**

```
data(roadkills)
submats <- sequential.submatrix(dataset = roadkills,</pre>
sampl.columns = 4:ncol(roadkills), sampl.interval = 1:3,
group.column = "taxon", include.all.together = TRUE,
remove.zeros = TRUE, keep.nonsampl.columns = TRUE,
n.subsampl.columns = 120)
hsl <- sequential.hotspots(dataset = roadkills, submats = submats,</pre>
region.column = "segment", first.subsampl.col = 4, confidence = 0.95)
hsn <- hotspot.numbers(hotspots.list = hsl, sampl.intervals = 1:3,
groups = as.character(unique(roadkills$taxon)),
include.all.together = TRUE, min.total.events = 0,
min.hotspot.threshold = 2)
seqcorr <- sequential.corr(hotspots.list = hsl,</pre>
hotspots.thresholds = hsn$HS.threshold, comp.method = "Phi",
baseline.gap = 0, messages = "TRUE")
seqcorr
```

sequential.estimateN

Sequential estimate N

# **Description**

This function estimates the actual numbers of animal casualties given the observed numbers and a set of estimators, sequentially for all given submats. Requires package **carcass**.

#### Usage

```
sequential.estimateN(submats, submats.N, first.subsampl.col,
region.column, persist, effic, estimators = c("korner", "huso",
"erickson", "etterson"), margin = 0.05, ...)
```

# Arguments

```
submats result of the sequential.submatrix function.
submats.N result of the sequential.Nevents function.
first.subsampl.col
```

index number of the first column containing the (sub)sampling data in submats

18 sequential.hotspots

region.colum	n
	name or index number of the column containing the regions (road segments, sites) to classify as hotspots or non-hotspots
persist	named vector of persistence per group; group names must match those in the data
effic	named vector of detection efficiency per group; group names must match those in the data
estimators	character vector of the $\operatorname{estimator}(s)$ to use. The default is all $\operatorname{estimators}$ available.
margin	proportion of each estimator to subtract from and add to it in order to get p.lower and p.upper, respectively, when using function $\texttt{estimateN}$ in package $\texttt{carcass}$
	currently not in use

#### Value

This function returns a list.

#### Note

This function currently works only for submats created with 'window.size' and 'gap.size', not with 'sampl.interval'. See submatrix, sequential.submatrix, and check that your names(submats) are something like "group.w1.g2.s1" and not "group.intv1".

# Author(s)

A. Marcia Barbosa, J. Tiago Marques, Sara Santos

# See Also

function estimateN in package carcass

```
sequential.hotspots
```

Calculate roadkill hotspots for a series of (sub)sampling datasets

# Description

This function applies hotspots sequencially to a given set of submatrices to identify the hotspot regions in each dataset, using an adaptation of the method of Malo et al. (2004).

# Usage

```
sequential.hotspots(dataset, submats, region.column,
first.subsampl.col, confidence = 0.95)
```

sequential.hotspots 19

# **Arguments**

dataset name of the matrix or dataframe containing the complete data

submats a list of the submatrices for which to calculate the hotspots (result of the sequential.submatrix function)

region.column

name or index number of the column containing the regions (road segments, sites) to classify as hotspots or non-hotspots

first.subsampl.col

index number of the first column containing subsampling data

confidence confidence threshold to consider hotspots. The default is 0.95

#### Value

#### A list of 2 elements:

hotspots.thresholds

A named integer vector

hotspots.maps

A list of data frames, each showing the total number of events (deaths) per region and whether or not it was considered a hospot.

#### Author(s)

A. Marcia Barbosa

#### References

Malo, J.E., Suarez, F., Diez, A. (2004) Can we mitigate animal-vehicle accidents using predictive models? J. Appl. Ecol. 41, 701-710 (doi: 10.1111/j.0021-8901.2004.00929.x)

#### See Also

hotspots

```
data(roadkills)
submats <- sequential.submatrix(dataset = roadkills,
sampl.columns = 4:ncol(roadkills), window.sizes = 1, gap.sizes = 1:3,
group.column = "taxon", include.all.together = TRUE,
remove.zeros = TRUE, keep.nonsampl.columns = TRUE,
n.subsampl.columns = 85)
shs <- sequential.hotspots(dataset = roadkills, submats = submats,
region.column = "segment", first.subsampl.col = 4, confidence = 0.95)
shs
str(shs)</pre>
```

20 sequential.posteriorN

```
sequential. Nevents Sequential numbers of events
```

#### **Description**

Applies function hot spots (with hotspots=FALSE) sequentially to a given set of submatrices

#### Usage

```
sequential.Nevents(dataset, submats, region.column,
first.subsampl.col, estimate = FALSE)
```

# **Arguments**

dataset name of the matrix or dataframe containing the complete data

submats a list of the submatrices for which to calculate the hotspots (result of the sequen-

tial.submatrix function)

region.column

name or index number of the column containing the regions (road sectors, sites)

to classify as hotspots or non-hotspots

first.subsampl.col

index number of the first column containing subsampling data

estimate logical, whether to add estimates from package carcass

# Author(s)

A. Marcia Barbosa

# See Also

```
hotspots, sequential.hotspots
```

```
sequential.posteriorN
```

Sequential posterior N

# Description

Applies function posteriorN of package **carcass** sequentially to a given set of submatrices, to estimate the number of (roadkill mortality) events based on Bayes' theorem using up to four different estimators (Korner-Nievergelt et al. 2015)

sequential.posteriorN 21

# Usage

```
sequential.posteriorN(submats, submats.N, first.subsampl.col,
region.column, persist, effic, estimators = c("korner", "huso",
"erickson", "etterson"), ...)
```

# Arguments

submats	a list of submatrices resulting from function sequential.submatrix							
submats.N	a list of submatrices resulting from function sequential. Nevents							
first.subsampl.col								
	index number of the first column containing subsampling data							
region.colum	n							
	name or index number of the column containing the regions (road sectors, sites) to classify as hotspots or non-hotspots							
persist	named numeric vector of persistence probability per group; names must match the names of groups in the data							
effic	named numeric vector of detection efficiency per group; names must match the names of groups in the data							
estimators	character vector specifying the estimators to calculate; the default is all available ones							
	additional arguments for function posteriorN in package carcass							

#### Value

This function returns a list

#### Author(s)

A. Marcia Barbosa, J. Tiago Marques, Sara M. Santos

# References

Korner-Nievergelt F, Behr O, Brinkmann R, Etterson MA, Huso MMP, Dalthorp D, Korner-Nievergelt P, Roth T & Niermann I (2015) Mortality estimation from carcass searches using the R-package carcass: a tutorial. Wildlife Biology 21: 30-43

# See Also

```
sequential.submatrix, sequential.Nevents
```

22 sequential.seqsubmat

```
sequential.seqsubmat
```

Sequential sequential submatrix

# **Description**

Applies sequential.submatrix sequentially to a set of gap sizes, with window.size =1

# Usage

```
sequential.seqsubmat(dataset, sampl.columns, group.column, gap.sizes,
n.replicates.limit)
```

# **Arguments**

```
dataset name of the matrix or dataframe to analyze

sampl.columns

numbers of the consecutive columns with the (daily) sampling data, e.g. 4:180

group.column name or index number of the column containing the taxa or groups to analyse separately, e.g. 3 or "Family"; if NULL, all records will be used together

gap.sizes integer vector of the size(s) of the gaps between sampling days. Must be either 0 (for no gap) or a vector of positive integers.

n.replicates.limit

Maximum number of replicates allowed
```

# Value

This function returns a list of submatrices

#### Author(s)

A. Marcia Barbosa

## See Also

```
sequential.submatrix
```

```
data(roadkills)
seqsubmats <- sequential.seqsubmat(dataset = roadkills,
sampl.columns = 4:ncol(roadkills), group.column = "taxon", gap.sizes = 1:4,
n.replicates.limit = 7)
names(seqsubmats)</pre>
```

sequential.submatrix 23

```
# seqsubmats for gap 0 (baseline scenario) must be obtained separately:
seqsubmats.baseline <- sequential.seqsubmat(dataset = roadkills,
sampl.columns = 4:ncol(roadkills), group.column = 3, gap.sizes = 0,
n.replicates.limit = 7)</pre>
names(seqsubmats.baseline)
```

sequential.submatrix

include.all.together

Extract subsampling matrices for a series of subsampling schemes

## **Description**

This function applies submatrix sequencially to all specified sampling schemes and taxonomic groups.

# Usage

```
sequential.submatrix(dataset, sampl.columns, sampl.intervals = NULL,
window.sizes = NULL, gap.sizes = NULL, start.columns = 1,
all.combinations = TRUE, group.column = NULL,
include.all.together = TRUE, remove.zeros = TRUE,
keep.nonsampl.columns = TRUE, n.subsampl.columns = NULL)
```

## **Arguments**

name of the matrix or dataframe to analyze dataset sampl.columns numbers of the consecutive columns with the (daily) sampling data, e.g. 4:180 sampl.intervals a vector of the intervals at which to extract sampling data, e.g. 5 to take one every five samples window.sizes the size (in sampling time units, e.g. days) of the sampling periods the size of the gaps between sampling periods gap.sizes start.columns vector of start columns, see submatrix all.combinations logical, whether to use all window x gap size combinations (the default, TRUE) or just the number corresponding to the length of window.sizes and gap sizes (in which case window.sizes and gap.sizes must have the same length) group.column name or index number of the column containing the taxa or groups to analyse separately, e.g. 3 or "Family"; if NULL, all records will be used together

logical, whether to get subsampling matrices also for the complete data (including all groups together)

24 submatrix

```
remove.zeros logical, whether to remove rows where all extracted samples have zero observa-
tions

keep.nonsampl.columns
logical, whether to include also the non-sampling columns of dataset in the
resulting submatrices)

n.subsampl.columns
number of subsampl.columns
```

#### Value

A list of submatrices

#### Author(s)

A. Marcia Barbosa

#### See Also

```
submatrix, subset
```

# **Examples**

```
data(roadkills)
submats1 <- sequential.submatrix(dataset = roadkills,</pre>
sampl.columns = 4:ncol(roadkills), sampl.intervals = c(1, 3),
group.column = "taxon", include.all.together = TRUE,
remove.zeros = TRUE, keep.nonsampl.columns = TRUE,
n.subsampl.columns = 85)
names(submats1)
head(submats1[[1]])
submats2 <- sequential.submatrix(dataset = roadkills,</pre>
sampl.columns = 4:ncol(roadkills), window.sizes = c(1,3,5),
gap.sizes = 1:3, start.columns = 1, all.combinations = TRUE,
group.column = "taxon", include.all.together = TRUE,
remove.zeros = TRUE, keep.nonsampl.columns = TRUE,
n.subsampl.columns = 85)
names(submats2)
head(submats2[[1]])
```

submatrix

Extract a submatrix for a given taxomic group and/or sampling scheme

submatrix 25

#### **Description**

Given a baseline dataset, this function extracts a sub-dataset for a given taxomic group and/or sampling scheme, defined either by a sampling interval (periodicity) or by a window size (consecutive sampling days each time) and a gap size (gaps between sampling windows).

## Usage

```
submatrix(dataset, sampl.columns, sampl.interval = NULL,
window.size = NULL, gap.size = NULL, start.column = 1,
group.column = NULL, group.names = NULL, remove.zeros = TRUE,
keep.nonsampl.columns = TRUE)
```

# **Arguments**

dataset name of the matrix or dataframe to analyze sampl.columns index numbers of the (consecutive) columns containing the baseline (daily) sampling data, e.g. 3:180 sampl.interval interval at which to extract sampling data, e.g. 5 (to take one every five samples) window.size instead of sampl.interval (for one sampling every so many time units), size (in sampling time units, e.g. days) of each sampling window (e.g. 3 for 3 consecutive days sampling each time); must be complemented with gap.size, for the gap between sampling windows. size (in sampling time units, e.g. days) of the gaps between sampling windows gap.size - e.g. 1 for 1 gap (non-sampled) day between sampling windows. start.column column of sampl.columns where to actually start the sampling (e.g. with a gap size of 1 between sampling windows, the start column can be either 1 or 2) group.column name or index number of the column containing the taxa or groups to analyse, e.g. 3 or "Family" name(s) of the group(a) to extract, e.g. c("Mustelidae", "Procyonidae"); if NULL group.names (the default), all groups in group.names are extracted remove.zeros logical indicating whether to remove rows where all extracted days have zero observations keep.nonsampl.columns logical indicating whether to keep the non-sampling columns in the extracted

#### Value

This function returns a subset of dataset containing the taxonomic groups and sampling columns resulting from the given sampling scheme.

#### Author(s)

A. Marcia Barbosa, J. Tiago Marques

26 submatrix

# See Also

```
sequential.submatrix, subset
```

```
data(roadkills)
submat1 <- submatrix(dataset = roadkills, sampl.columns = 4:ncol(roadkills),
sampl.interval = 3, start.column = 1, group.column = "taxon", group.names = NULL)
head(submat1)
submat2 <- submatrix(dataset = roadkills, sampl.columns = 4:ncol(roadkills),
window.size = 5, gap.size = 2, start.column = 1, group.column = "taxon",
group.names = NULL)
head(submat2)</pre>
```