# Package 'nimbleSCR'

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Type Package

Title Spatial Capture-Recapture (SCR) Methods Using 'nimble'

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**Description** Provides utility functions, distributions, and fitting methods for Bayesian Spatial Capture-Recapture (SCR) and Open Population Spatial Capture-Recapture (OPSCR) modelling using the nimble package (de Valpine et al. 2017 <doi:10.1080/10618600.2016.1172487 >). Development of the package was motivated primarily by the need for flexible and efficient analysis of large-scale SCR data (Bischof et al. 2020 <doi:10.1073/pnas.2011383117 >). Computational methods and techniques implemented in nimbleSCR include those discussed in Turek et al. 2021 <doi:10.1002/ecs2.3385>; among others. For a recent application of nimbleSCR, see Milleret et al. (2021) <doi:10.1098/rsbl.2021.0128>.

License GPL-3

**Depends** R (>= 3.5.0), nimble

**Imports** methods **Encoding** UTF-8

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Suggests knitr, rmarkdown, testthat (>= 3.0.0), coda, basicMCMCplots

**VignetteBuilder** knitr **Config/testthat/edition** 3

Collate calcWindowSizes.R getWindowIndex.R

integrateIntensityLocal\_normal.R integrateIntensityLocal\_exp.R integrateIntensity\_normal.R integrateIntensity\_exp.R stratRejectionSampler\_normal.R stratRejectionSampler\_exp.R dDispersal\_exp.R dHabitatMask.R dbernppAC.R dbernppACmovement\_normal.R dbernppACmovement\_exp.R dbernppDetection\_normal.R dbernppLocalACmovement\_normal.R dbernppLocalACmovement\_normal.R dbinomLocalACmovement\_exp.R dbinomLocalDetection\_normal.R dbinomLocal\_normal.R dbinomLocal\_normal.R dbinomLocal\_normalPlateau.R dbinomLocal\_exp.R dpoisppAC.R dpoisppDetection\_normal.R

2 R topics documented:

dpoisppLocalDetection_normal.R getLocalObjects.R
getMidPointNodes.R getWindowCoords.R getSparseY.R
getHomeRangeArea.R localTrapCalculations.R
makeConstantNimbleFunction.R marginalVoidProbIntegrand.R
$marginal Void Prob Num Integration. R\ scale Coords To Habitat Grid. R$
dcatState1Alive1Dead.R dcatState1Alive2Dead.R
dcatState2Alive2Dead.R sampler_categorical_general.R
calculateDensity.R zzz.R

# NeedsCompilation no

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# **R** topics documented:

calculateDensity	3
calcWindowSizes	4
dbernppAC	5
dbernppACmovement_exp	7
dbernppACmovement_normal	)
dbernppDetection_normal	1
dbernppLocalACmovement_exp	4
dbernppLocalACmovement_normal	3
dbernppLocalDetection_normal	1
dbinomLocal_exp	5
dbinomLocal_normal	)
dbinomLocal_normalPlateau	4
dbinom_vector	)
dcatState1Alive1Dead	1
dcatState1Alive2Dead	3
dcatState2Alive2Dead	5
dDispersal_exp	)
dHabitatMask	1
dmultiLocal_normal	3
dnormalizer	3
dpoisLocal_normal	)
dpoisppAC	3
dpoisppDetection_normal	5
dpoisppLocalDetection_normal68	3

calculateDensity 3

getHomeRangeArea	a	71
getLocalObjects		<b>7</b> 4
getMidPointNodes		76
getWindowCoords		78
getWindowIndex .		79
_	ocal_normal	
	ormal	
	ns	
_	ntegrand	
_	IumIntegration	
_	l_general	
	tatGrid	
	ler_exp	
	ler_normal	
structegeetionsumpi	ioi_normai	
Index		97
calculateDensity	NIMBLE function to calculate the density of individuals alive	in each
	habitat cell.	

# Description

 ${\tt calculateDensity} \ is \ a \ NIMBLE \ function \ to \ calculate \ number \ of \ individual \ activity \ centers \ (s) \ in \ each \ habitat \ cell.$ 

# Usage

calculateDensity(s, habitatGrid, indicator, numWindows, nIndividuals)

# Arguments

S	Matrix of x- and y-coordinates of individual AC locations.
habitatGrid	Matrix of habitat window indices. Cell values should correspond to the order of habitat windows in spatial probabilities (e.g. prob1To2Hab as used in the function dcatState1Alive2Dead) or in lowerCoords and upperCoords as used in the dbernppAC function. #' @param indicator Vector of binary arguments specifying whether the individuals are considered alive (indicator = 1) or not (indicator = 0).
indicator	Vector of binary arguments specifying whether the individuals are considered alive (indicator = $1$ ) or not (indicator = $0$ ).
numWindows	Scalar Number of habitat windows.
nIndividuals	Scalar Number of individuals.

4 calc Window Sizes

## Author(s)

Cyril Milleret

# **Examples**

```
lowerCoords <- matrix(c(0, 0, 1, 0, 0, 1, 1, 1), nrow = 4, byrow = TRUE)
upperCoords <- matrix(c(1, 1, 2, 1, 1, 2, 2, 2), nrow = 4, byrow = TRUE)
logIntensities <- log(rep(1,4))</pre>
logSumIntensity <- log(sum(c(1:4)))</pre>
habitatGrid <- matrix(c(1:4), nrow = 2, byrow = TRUE)
numGridRows <- nrow(habitatGrid)</pre>
numGridCols <- ncol(habitatGrid)</pre>
s <- matrix(NA,nrow=10,ncol=2)</pre>
for(i in 1:10){
  s[i,] <- rbernppAC(n=1, lowerCoords, upperCoords, logIntensities, logSumIntensity,</pre>
                      habitatGrid, numGridRows, numGridCols)
}
calculateDensity(s = s,
                  habitatGrid = habitatGrid,
                  indicator = rep(1, 10),
                  numWindows = prod(dim(habitatGrid)),
                  nIndividuals = 10
)
```

calcWindowSizes

Window size calculation

# **Description**

Calculates the sizes of a set of windows based on their lower and upper coordinates of each dimension. Can be applied to detection and habitat windows.

## Usage

```
calcWindowSizes(lowerCoords, upperCoords)
```

## **Arguments**

lowerCoords Matrix of lower x- and y-coordinates of all windows. One row for each window. upperCoords Matrix of upper x- and y-coordinates of all windows. One row for each window.

#### Value

A vector of window sizes.

dbernppAC 5

# Author(s)

Wei Zhang

# **Examples**

```
lowerCoords <- matrix(c(0, 0, 1, 0, 0, 1, 1, 1), nrow = 4, byrow = TRUE) upperCoords <- matrix(c(1, 1, 3, 1, 1, 4, 3, 4), nrow = 4, byrow = TRUE) calcWindowSizes(lowerCoords, upperCoords)
```

dbernppAC

Bernoulli point process for the distribution of activity centers

# **Description**

Density and random generation functions of the Bernoulli point process for the distribution of activity centers.

```
dbernppAC(
  Х,
  lowerCoords,
  upperCoords,
  logIntensities,
  logSumIntensity,
 habitatGrid,
 numGridRows,
 numGridCols,
 log = 0
)
rbernppAC(
  n,
  lowerCoords,
  upperCoords,
  logIntensities,
  logSumIntensity,
 habitatGrid,
 numGridRows,
  numGridCols
)
```

6 dbernppAC

#### **Arguments**

vector of x- and y-coordinates of a single spatial point (i.e. AC location) scaled to the habitat (see (scaleCoordsToHabitatGrid).

lowerCoords, upperCoords

Matrices of lower and upper x- and y-coordinates of all habitat windows scaled to the habitat (see (scaleCoordsToHabitatGrid). One row for each window. Each window should be of size 1x1.

logIntensities Vector of log habitat intensities for all habitat windows.

logSumIntensity

Log of the sum of habitat intensities over all windows.

habitatGrid Matrix of habitat window indices. Cell values should correspond to the order of

habitat windows in lowerCoords, upperCoords, and logIntensities. When the habitat grid only consists of a single row or column of windows, an additional row or column of dummy indices has to be added because the nimble

model code requires a matrix.

numGridRows, numGridCols

Numbers of rows and columns of the habitat grid.

log Logical argument, specifying whether to return the log-probability of the distri-

bution.

n Integer specifying the number of realisations to generate. Only n = 1 is sup-

ported.

#### **Details**

The dbernppAC distribution is a NIMBLE custom distribution which can be used to model and simulate the activity center location (x) of a single individual in continuous space over a set of habitat windows defined by their upper and lower coordinates (lowerCoords, upperCoords). The distribution assumes that the activity center follows a Bernoulli point process with intensity = exp(logIntensities).

# Value

dbernppAC gives the (log) probability density of the observation vector x. rbernppAC gives coordinates of a randomly generated spatial point.

#### Author(s)

Wei Zhang

#### References

W. Zhang, J. D. Chipperfield, J. B. Illian, P. Dupont, C. Milleret, P. de Valpine and R. Bischof. 2020. A hierarchical point process model for spatial capture-recapture data. bioRxiv. DOI 10.1101/2020.10.06.325035

## **Examples**

dbernppACmovement\_exp Bernoulli point process for activity center movement (exponential kernel)

# **Description**

Density and random generation functions of the Bernoulli point process for activity center movement between occasions based on a bivariate exponential distribution.

```
dbernppACmovement_exp(
  lowerCoords,
  upperCoords,
  s,
  lambda = -999,
  rate,
  baseIntensities,
 habitatGrid,
  numGridRows,
  numGridCols,
  numWindows,
 log = 0
)
rbernppACmovement_exp(
  lowerCoords,
  upperCoords,
  s,
  lambda = -999,
  rate,
  baseIntensities,
  habitatGrid,
```

```
numGridRows,
numGridCols,
numWindows
)
```

#### **Arguments**

v Vector of x- and y-coordinates of a single spatial point (typically AC location at time t+1) scaled to the habitat (see (scaleCoordsToHabitatGrid).

lowerCoords, upperCoords

Matrices of lower and upper x- and y-coordinates of all habitat windows. One row for each window. Each window should be of size 1x1 (after rescaling if

necessary).

s Vector of x- and y-coordinates of the isotropic multivariate exponential distribu-

tion mean (AC location at time t).

lambda Rate parameter of the isotropic bivariate exponential distribution. Soon depre-

cated, use argument "rate" instead.

rate Rate parameter of the isotropic multivariate exponential distribution.

baseIntensities

Vector of baseline habitat intensities for all habitat windows.

habitatGrid Matrix of habitat window indices. Cell values should correspond to the order of

habitat windows in lowerCoords and upperCoords. When the habitat grid only consists of a single row or column of windows, an additional row or column of dummy indices has to be added because the nimble model code requires a

matrix.

numGridRows, numGridCols

Numbers of rows and columns of the habitat grid.

numWindows Number of habitat windows. This value (positive integer) can be used to trun-

cate lowerCoords and upperCoords so that extra rows beyond numWindows are

ignored.

log Logical argument, specifying whether to return the log-probability of the distri-

bution.

n Integer specifying the number of realisations to generate. Only n = 1 is sup-

ported.

#### **Details**

The dbernppACmovement\_exp distribution is a NIMBLE custom distribution which can be used to model and simulate movement of activity centers between consecutive occasions in open population models. The distribution assumes that the new individual activity center location (x) follows an isotropic exponential normal centered on the previous activity center (s) with rate (lambda).

# Value

dbernppACmovement\_exp gives the (log) probability density of the observation vector x. rbernppACmovement\_exp gives coordinates of a randomly generated spatial point.

## Author(s)

Wei Zhang and Cyril Milleret

#### References

W. Zhang, J. D. Chipperfield, J. B. Illian, P. Dupont, C. Milleret, P. de Valpine and R. Bischof. 2020. A hierarchical point process model for spatial capture-recapture data. bioRxiv. DOI 10.1101/2020.10.06.325035

# **Examples**

```
# Use the distribution in R
lowerCoords <- matrix(c(0, 0, 1, 0, 0, 1, 1, 1), nrow = 4, byrow = TRUE)
upperCoords <- matrix(c(1, 1, 2, 1, 1, 2, 2, 2), nrow = 4, byrow = TRUE)
s \leftarrow c(1, 1) \# Currrent activity center location
rate <- 0.1
baseIntensities <- c(1:4)</pre>
habitatGrid \leftarrow matrix(c(1:4), nrow = 2, byrow = TRUE)
numRows <- nrow(habitatGrid)</pre>
numCols <- ncol(habitatGrid)</pre>
numWindows <- 4
# The log probability density of moving from (1,1) to (1.2, 0.8)
dbernppACmovement_exp(x = c(1.2, 0.8),
lowerCoords = lowerCoords,
upperCoords = upperCoords,
s = s,
rate = rate,
baseIntensities = baseIntensities,
habitatGrid = habitatGrid,
numGridRows = numRows,
numGridCols = numCols,
numWindows = numWindows,
log = TRUE)
```

 ${\tt dbernppACmovement\_normal}$ 

Bernoulli point process for activity center movement (normal kernel)

## **Description**

Density and random generation functions of the Bernoulli point process for activity center movement between occasions based on a bivariate normal distribution.

```
dbernppACmovement_normal(
    x,
    lowerCoords,
    upperCoords,
    s,
```

```
sd,
  baseIntensities,
  habitatGrid,
  numGridRows,
  numGridCols,
  numWindows,
  log = 0
rbernppACmovement_normal(
  lowerCoords,
  upperCoords,
  s,
  sd,
  baseIntensities,
  habitatGrid,
  numGridRows,
  numGridCols,
  numWindows
)
```

## **Arguments**

Х

Vector of x- and y-coordinates of a single spatial point (typically AC location at time t+1) scaled to the habitat (see (scaleCoordsToHabitatGrid).

lowerCoords, upperCoords

Matrices of lower and upper x- and y-coordinates of all habitat windows scaled to the habitat (see (scaleCoordsToHabitatGrid). One row for each window. Each window should be of size 1x1.

S

Vector of x- and y-coordinates of the isotropic bivariate normal distribution mean (AC location at time t).

sd

Standard deviation of the isotropic bivariate normal distribution..

baseIntensities

Vector of baseline habitat intensities for all habitat windows.

habitatGrid

Matrix of habitat window indices. Cell values should correspond to the order of habitat windows in lowerCoords and upperCoords. When the habitat grid only consists of a single row or column of windows, an additional row or column of dummy indices has to be added because the nimble model code requires a matrix.

numGridRows, numGridCols

Numbers of rows and columns of the habitat grid.

numWindows

Number of habitat windows. This value (positive integer) can be used to truncate lowerCoords and upperCoords so that extra rows beyond numWindows are ignored.

log

Logical argument, specifying whether to return the log-probability of the distribution.

Integer specifying the number of realisations to generate. Only n = 1 is supported.

#### **Details**

The dbernppACmovement\_normal distribution is a NIMBLE custom distribution which can be used to model and simulate movement of activity centers between consecutive occasions in open population models. The distribution assumes that the new individual activity center location (x) follows an isotropic multivariate normal centered on the previous activity center (s) with standard deviation (sd).

#### Value

dbernppACmovement\_normal gives the (log) probability density of the observation vector x. rbernppACmovement\_normal gives coordinates of a randomly generated spatial point.

## Author(s)

Wei Zhang and Cyril Milleret

#### References

W. Zhang, J. D. Chipperfield, J. B. Illian, P. Dupont, C. Milleret, P. de Valpine and R. Bischof. 2020. A hierarchical point process model for spatial capture-recapture data. bioRxiv. DOI 10.1101/2020.10.06.325035

## **Examples**

```
# Use the distribution in R lowerCoords <- matrix(c(0, 0, 1, 0, 0, 1, 1, 1), nrow = 4, byrow = TRUE) upperCoords <- matrix(c(1, 1, 2, 1, 1, 2, 2, 2), nrow = 4, byrow = TRUE) s <- c(1, 1) # Current activity center location sd <- 0.1 baseIntensities <- c(1:4) habitatGrid <- matrix(c(1:4), nrow = 2, byrow = TRUE) numRows <- nrow(habitatGrid) numCols <- ncol(habitatGrid) numWindows <- 4 # The log probability density of moving from (1,1) to (1.2, 0.8) dbernppACmovement_normal(c(1:2, 0.8), lowerCoords, upperCoords, s, sd, baseIntensities, habitatGrid, numRows, numCols, numWindows, log = TRUE)
```

dbernppDetection\_normal

Bernoulli point process detection model

# Description

Density and random generation functions of the Bernoulli point process for detection.

## Usage

```
dbernppDetection_normal(
  lowerCoords,
  upperCoords,
  s,
  sd,
  baseIntensities,
  numWindows,
  indicator,
  log = 0
)
rbernppDetection_normal(
  n,
  lowerCoords,
  upperCoords,
  s,
  sd,
  baseIntensities,
  numWindows,
  indicator
```

# **Arguments**

Х

sd

Vector with three elements representing the x- and y-coordinates and the id of the corresponding detection window for a single spatial point (detection location) scaled to the habitat (see (scaleCoordsToHabitatGrid).

lowerCoords, upperCoords

Matrices of lower and upper x- and y-coordinates of all detection windows scaled to the habitat (see (scaleCoordsToHabitatGrid). One row for each window. Each window should be of size 1x1.

S VVector of x- and y-coordinates of the isotropic bivariate normal distribution mean (i.e. the AC location)..

Standard deviation of the isotropic bivariate normal distribution.

baseIntensities

Vector of baseline detection intensities for all detection windows.

numWindows Number of detection windows. This value (positive integer) is used to truncate

lowerCoords and upperCoords so that extra rows beyond numWindows are ig-

nored.

indicator Binary argument specifying whether the individual is available for detection (in-

dicator = 1) or not (indicator = 0).

log Logical argument, specifying whether to return the log-probability of the distri-

bution.

n Integer specifying the number of realisations to generate. Only n = 1 is sup-

ported.

#### **Details**

The dbernppDetection\_normal distribution is a NIMBLE custom distribution which can be used to model and simulate Bernoulli observations (x) of a single individual in continuous space over a set of detection windows defined by their upper and lower coordinates (lowerCoords, upperCoords). The distribution assumes that an individual's detection probability follows an isotropic multivariate normal centered on the individual's activity center (s) with standard deviation (sd).

## Value

dbernppDetection\_normal gives the (log) probability density of the observation vector x. rbernppDetection\_normal gives coordinates of a randomly generated spatial point.

#### Author(s)

Wei Zhang and Cyril Milleret

#### References

W. Zhang, J. D. Chipperfield, J. B. Illian, P. Dupont, C. Milleret, P. de Valpine and R. Bischof. 2020. A hierarchical point process model for spatial capture-recapture data. bioRxiv. DOI 10.1101/2020.10.06.325035

# **Examples**

```
coordsHabitatGridCenter <- matrix(c(0.5, 3.5,</pre>
                                      1.5, 3.5,
                                      2.5, 3.5,
                                      3.5, 3.5,
                                      0.5, 2.5,
                                      1.5, 2.5,
                                      2.5, 2.5,
                                      3.5, 2.5,
                                      0.5, 1.5,
                                      1.5, 1.5,
                                      2.5, 1.5,
                                      3.5, 1.5,
                                      0.5, 0.5,
                                     1.5, 0.5,
                                      2.5, 0.5,
                                      3.5, 0.5), ncol = 2, byrow = TRUE)
colnames(coordsHabitatGridCenter) <- c("x","y")</pre>
# Create observation windows
 lowerCoords <- matrix(c(0, 0, 1, 0, 0, 1, 1, 1), nrow = 4, byrow = TRUE)
 upperCoords <- matrix(c(1, 1, 2, 1, 1, 2, 2, 2), nrow = 4, byrow = TRUE)
 colnames(lowerCoords) <- colnames(upperCoords) <- c("x","y")</pre>
# Rescale coordinates
ScaledLowerCoords <- scaleCoordsToHabitatGrid(coordsData = lowerCoords,</pre>
                                        coordsHabitatGridCenter = coordsHabitatGridCenter)
ScaledUpperCoords <- scaleCoordsToHabitatGrid(coordsData = upperCoords,</pre>
                                       coordsHabitatGridCenter = coordsHabitatGridCenter)
ScaledUpperCoords$coordsDataScaled[,2] <- ScaledUpperCoords$coordsDataScaled[,2] + 1.5</pre>
ScaledLowerCoords$coordsDataScaled[,2] <- ScaledLowerCoords$coordsDataScaled[,2] - 1.5
```

```
s <- c(1, 1)
sd <- 0.1
baseIntensities <- c(1:4)</pre>
windowIndex <- 4
numPoints <- 1
numWindows <- 4
indicator <- 1
x < -c(0.5, 2)
windowIndex <- getWindowIndex(curCoords = x,</pre>
                                lowerCoords = ScaledLowerCoordsScoordsDataScaled,
                                upperCoords =ScaledUpperCoords$coordsDataScaled)
x <- c(x, windowIndex)</pre>
dbernppDetection_normal(x, lowerCoords, upperCoords,
                         s, sd, baseIntensities
                          , numWindows,
                         indicator, log = TRUE)
```

dbernppLocalACmovement\_exp

Local evaluation of a Bernoulli point process for activity center movement (exponential kernel)

# **Description**

Density and random generation functions of the Bernoulli point process for activity center movement between occasions based on a bivariate exponential distribution and local evaluation.

```
dbernppLocalACmovement_exp(
    x,
    lowerCoords,
    upperCoords,
    s,
    lambda = -999,
    rate,
    baseIntensities,
    habitatGrid,
    habitatGridLocal,
    resizeFactor = 1,
    localHabWindowIndices,
    numLocalHabWindows,
    numGridRows,
    numGridCols,
    numWindows,
```

```
log = 0
rbernppLocalACmovement_exp(
  n,
  lowerCoords,
 upperCoords,
 lambda = -999,
  rate,
 baseIntensities,
 habitatGrid,
  habitatGridLocal,
  resizeFactor = 1,
  localHabWindowIndices,
  numLocalHabWindows,
  numGridRows,
 numGridCols,
  numWindows
)
```

## **Arguments**

s

lambda

rate

x Vector of x- and y-coordinates of a single spatial point (typically AC location at time t+1) scaled to the habitat (see (scaleCoordsToHabitatGrid).

lowerCoords, upperCoords

Matrices of lower and upper x- and y-coordinates of all habitat windows. One row for each window. Each window should be of size 1x1 (after rescaling if necessary).

Vector of x- and y-coordinates of the isotropic multivariate exponential distribution mean (AC location at time t).

Rate parameter of the isotropic bivariate exponential distribution. Soon deprecated, use argument "rate" instead.

Rate parameter of the isotropic bivariate exponential distribution.

baseIntensities

Vector of baseline habitat intensities for all habitat windows.

habitatGrid Matrix of habitat window indices. Cell values should correspond to the order of habitat windows in lowerCoords and upperCoords. When the habitat grid only consists of a single row or column of windows, an additional row or column of dummy indices has to be added because the nimble model code requires a

matrix.

habitatGridLocal

Matrix of rescaled habitat grid cells indices, as returned by the getLocalObjects function (object named habitatGrid).

resizeFactor Aggregation factor used in the getLocalObjects function to reduce the number of habitat grid cells.

#### localHabWindowIndices

Matrix of indices of local habitat windows around each local habitat grid cell (habitatGridLocal) from localIndices returned by getLocalObjects function.

#### numLocalHabWindows

Vector of numbers of local habitat windows around all habitat grid cells, from numLocalIndices returned by the getLocalObjects function. The ith number gives the number of local (original) habitat windows for the ith local habitat grid cell habitatGridLocal.

numGridRows, numGridCols

Numbers of rows and columns of the habitatGrid.

numWindows Number of habitat windows. This value (positive integer) is used to truncate

lowerCoords and upperCoords so that extra rows beyond numWindows are ig-

nored.

log Logical argument, specifying whether to return the log-probability of the distri-

bution.

n Integer specifying the number of realisations to generate. Only n = 1 is sup-

ported.

#### **Details**

The dbernppLocalACmovement\_exp distribution is a NIMBLE custom distribution which can be used to model and simulate movement of activity centers between consecutive occasions in open population models. The distribution assumes that the new individual activity center location (x) follows an isotropic exponential normal centered on the previous activity center (s) with rate (lambda). The local evaluation approach is implemented.

## Value

The (log) probability density of the observation vector x.

# Author(s)

Wei Zhang and Cyril Milleret

## References

W. Zhang, J. D. Chipperfield, J. B. Illian, P. Dupont, C. Milleret, P. de Valpine and R. Bischof. 2020. A hierarchical point process model for spatial capture-recapture data. bioRxiv. DOI 10.1101/2020.10.06.325035

C. Milleret, P. Dupont, C. Bonenfant, H. Broseth, O. Flagstad, C. Sutherland and R. Bischof. 2019. A local evaluation of the individual state-space to scale up Bayesian spatial capture-recapture. Ecology and Evolution 9:352-363

## **Examples**

```
# Creat habitat grid habitatGrid <- matrix(c(1:(4^2)), nrow = 4, ncol=4, byrow = TRUE) coordsHabitatGridCenter <- matrix(c(0.5, 3.5,
```

```
1.5, 3.5,
                                      2.5, 3.5,
                                      3.5, 3.5,
                                      0.5, 2.5,
                                      1.5, 2.5,
                                      2.5, 2.5,
                                      3.5, 2.5,
                                      0.5, 1.5,
                                      1.5, 1.5,
                                      2.5, 1.5,
                                      3.5, 1.5,
                                      0.5, 0.5,
                                      1.5, 0.5,
                                      2.5, 0.5,
                                      3.5, 0.5), ncol = 2, byrow = TRUE)
colnames(coordsHabitatGridCenter) <- c("x","y")</pre>
# Create habitat windows
lowerCoords <- coordsHabitatGridCenter-0.5</pre>
upperCoords <- coordsHabitatGridCenter+0.5</pre>
colnames(lowerCoords) <- colnames(upperCoords) <- c("x","y")</pre>
# Plot check
plot(lowerCoords[,"y"]~lowerCoords[,"x"],pch=16, xlim=c(0,4), ylim=c(0,4),col="red")
points(upperCoords[,"y"]~upperCoords[,"x"],col="red",pch=16)
points(coordsHabitatGridCenter[,"y"]~coordsHabitatGridCenter[,"x"],pch=16)
# Rescale coordinates
ScaledLowerCoords <- scaleCoordsToHabitatGrid(coordsData = lowerCoords,</pre>
                                        coordsHabitatGridCenter = coordsHabitatGridCenter)
ScaledUpperCoords <- scaleCoordsToHabitatGrid(coordsData = upperCoords,</pre>
                                        coordsHabitatGridCenter = coordsHabitatGridCenter)
ScaledUpperCoords$coordsDataScaled[,2] <- ScaledUpperCoords$coordsDataScaled[,2] + 1</pre>
ScaledLowerCoordsScoordsDataScaled[,2] <- ScaledLowerCoordsScoordsDataScaled[,2] - 1
habitatMask <- matrix(1, nrow = 4, ncol=4, byrow = TRUE)</pre>
# Create local objects
HabWindowsLocal <- getLocalObjects(habitatMask = habitatMask,</pre>
                                     coords = coordsHabitatGridCenter,
                                     dmax=4,
                                     resizeFactor = 1,
                                     plot.check = TRUE
)
s \leftarrow c(1, 1) # Currrent activity center location
rate <- 0.1
numWindows <- nrow(coordsHabitatGridCenter)</pre>
baseIntensities <- rep(1,numWindows)</pre>
numRows <- nrow(habitatGrid)</pre>
numCols <- ncol(habitatGrid)</pre>
# The log probability density of moving from (1,1) to (1.2, 0.8)
dbernppLocalACmovement_exp(x = c(1.2, 0.8),
lowerCoords =lowerCoords,
 upperCoords = upperCoords,
 s = s,
```

```
rate = rate,
baseIntensities = baseIntensities,
habitatGrid = habitatGrid,
habitatGridLocal = HabWindowsLocal$habitatGrid,
resizeFactor = HabWindowsLocal$resizeFactor,
localHabWindowIndices = HabWindowsLocal$localIndices,
numLocalHabWindows = HabWindowsLocal$numLocalIndices,
numGridRows = numRows,
numGridCols = numCols,
numWindows = numWindows,
log = TRUE)
```

dbernppLocalACmovement\_normal

Local evaluation of a Bernoulli point process for activity center movement (normal kernel)

# **Description**

Density and random generation functions of the Bernoulli point process for activity center movement between occasions based on a bivariate normal distribution and local evaluation.

```
dbernppLocalACmovement_normal(
  Х,
  lowerCoords,
  upperCoords,
  s,
  sd,
 baseIntensities,
 habitatGrid,
 habitatGridLocal,
  resizeFactor = 1,
  localHabWindowIndices,
  numLocalHabWindows,
  numGridRows,
  numGridCols,
  numWindows,
  log = 0
)
rbernppLocalACmovement_normal(
  lowerCoords,
```

```
upperCoords,
s,
sd,
baseIntensities,
habitatGrid,
habitatGridLocal,
resizeFactor = 1,
localHabWindowIndices,
numLocalHabWindows,
numGridRows,
numGridCols,
numWindows
)
```

#### **Arguments**

Χ

Vector of x- and y-coordinates of a single spatial point (typically AC location at time t+1) scaled to the habitat (see (scaleCoordsToHabitatGrid).

# lowerCoords, upperCoords

Matrices of lower and upper x- and y-coordinates of all habitat windows. One row for each window. Each window should be of size 1x1 (after rescaling if necessary).

S

Vector of x- and y-coordinates of the isotropic bivariate normal distribution mean (i.e. the AC location).

sd

Standard deviation of the isotropic bivariate normal distribution.

#### baseIntensities

Vector of baseline habitat intensities for all habitat windows.

habitatGrid

Matrix of habitat window indices. Cell values should correspond to the order of habitat windows in lowerCoords and upperCoords. When the habitat grid only consists of a single row or column of windows, an additional row or column of dummy indices has to be added because the nimble model code requires a matrix.

## habitatGridLocal

Matrix of rescaled habitat grid cells indices, from localIndices returned by the getLocalObjects function (

resizeFactor

Aggregation factor used in the getLocalObjects function to reduce the number of habitat grid cells.

#### localHabWindowIndices

Matrix of indices of local habitat windows around each local habitat grid cell (habitatGridLocal), from localIndices returned by the getLocalObjects function.

#### numLocalHabWindows

Vector of numbers of local habitat windows around all habitat grid cells, as returned by the getLocalObjects function (object named numLocalIndices). The ith number gives the number of local (original) habitat windows for the ith (rescaled) habitat window.

numGridRows, numGridCols

Numbers of rows and columns of the habitatGrid.

numWindows Number of habitat windows. This value (positive integer) is used to truncate

lowerCoords and upperCoords so that extra rows beyond numWindows are ig-

nored.

log Logical argument, specifying whether to return the log-probability of the distri-

bution.

n Integer specifying the number of realisations to generate. Only n = 1 is sup-

ported.

## **Details**

The dbernppLocalACmovement\_normal distribution is a NIMBLE custom distribution which can be used to model and simulate movement of activity centers between consecutive occasions in open population models. The distribution assumes that the new individual activity center location (x) follows an isotropic multivariate normal centered on the previous activity center (s) with standard deviation (sd). The local evaluation technique is implemented.

#### Value

The (log) probability density of the observation vector x.

#### Author(s)

Wei Zhang and Cyril Milleret

#### References

W. Zhang, J. D. Chipperfield, J. B. Illian, P. Dupont, C. Milleret, P. de Valpine and R. Bischof. 2020. A hierarchical point process model for spatial capture-recapture data. bioRxiv. DOI 10.1101/2020.10.06.325035

C. Milleret, P. Dupont, C. Bonenfant, H. Brøseth, Ø. Flagstad, C. Sutherland and R. Bischof. 2019. A local evaluation of the individual state-space to scale up Bayesian spatial capture-recapture. Ecology and Evolution 9:352-363

## **Examples**

```
3.5, 1.5,
                                      0.5, 0.5,
                                      1.5, 0.5,
                                      2.5, 0.5,
                                      3.5, 0.5), ncol = 2, byrow = TRUE)
colnames(coordsHabitatGridCenter) <- c("x","y")</pre>
# Create habitat windows
lowerCoords <- coordsHabitatGridCenter-0.5</pre>
upperCoords <- coordsHabitatGridCenter+0.5</pre>
colnames(lowerCoords) <- colnames(upperCoords) <- c("x","y")</pre>
# Plot check
plot(lowerCoords[,"y"]~lowerCoords[,"x"],pch=16, xlim=c(0,4), ylim=c(0,4),col="red")
points(upperCoords[,"y"]~upperCoords[,"x"],col="red",pch=16)
points(coordsHabitatGridCenter[,"y"]~coordsHabitatGridCenter[,"x"],pch=16)
# Rescale coordinates
ScaledLowerCoords <- scaleCoordsToHabitatGrid(coordsData = lowerCoords,</pre>
                                        coordsHabitatGridCenter = coordsHabitatGridCenter)
ScaledUpperCoords <- scaleCoordsToHabitatGrid(coordsData = upperCoords,</pre>
                                        coordsHabitatGridCenter = coordsHabitatGridCenter)
ScaledUpperCoords$coordsDataScaled[,2] <- ScaledUpperCoords$coordsDataScaled[,2] + 1</pre>
ScaledLowerCoords$coordsDataScaled[,2] <- ScaledLowerCoords$coordsDataScaled[,2] - 1
habitatMask <- matrix(1, nrow = 4, ncol=4, byrow = TRUE)</pre>
# Create local objects
HabWindowsLocal <- getLocalObjects(habitatMask = habitatMask,</pre>
                                     coords = coordsHabitatGridCenter,
                                     dmax=4,
                                     resizeFactor = 1,
                                     plot.check = TRUE
)
s \leftarrow c(1, 1) # Currrent activity center location
sd <- 0.1
numWindows <- nrow(coordsHabitatGridCenter)</pre>
baseIntensities <- rep(1,numWindows)</pre>
numRows <- nrow(habitatGrid)</pre>
numCols <- ncol(habitatGrid)</pre>
# The log probability density of moving from (1,1) to (1.2, 0.8)
dbernppLocalACmovement_normal(x = c(1.2, 0.8), lowerCoords, upperCoords, s,
                               sd, baseIntensities, habitatGrid,
                               HabWindowsLocal$habitatGrid, HabWindowsLocal$resizeFactor,
                           HabWindowsLocal$localIndices, HabWindowsLocal$numLocalIndices,
                               numRows, numCols, numWindows, log = TRUE)
```

dbernppLocalDetection\_normal

Local evaluation for a Bernoulli point process detection model

# **Description**

Density and random generation functions of the Bernoulli point process for detection based on a bivariate normal distribution.

# Usage

```
dbernppLocalDetection_normal(
  х,
 lowerCoords,
  upperCoords,
  s,
  sd,
  baseIntensities,
  habitatGridLocal,
  resizeFactor = 1,
  localObsWindowIndices,
  numLocalObsWindows,
  numWindows,
  indicator,
  log = 0
rbernppLocalDetection_normal(
  n,
  lowerCoords,
 upperCoords,
  s,
  sd,
  baseIntensities,
  habitatGridLocal,
  resizeFactor = 1,
  localObsWindowIndices,
  numLocalObsWindows,
  numWindows,
  indicator
)
```

# Arguments

Χ

Vector with three elements representing the x- and y-coordinates (x[1:2]), and the corresponding id the detection window (x[3]) of a single spatial point (detection location) scaled to the habitat (see scaleCoordsToHabitatGrid).

lowerCoords, upperCoords

Matrices of lower and upper x- and y-coordinates of all detection windows. One row for each window. Each window should be of size 1x1 (after rescaling if necessary).

s Vector of x- and y-coordinates of the isotropic bivariate normal distribution mean (i.e. the AC location).

sd Standard deviation of the bivariate normal distribution.

baseIntensities

Vector of baseline detection intensities for all detection windows.

habitatGridLocal

Matrix of rescaled habitat grid cells indices, as returned by the getLocalObjects function (object named habitatGrid).

resizeFactor A

Aggregation factor used in the getLocalObjects function to reduce the number of habitat grid cells.

localObsWindowIndices

Matrix of indices of local observation windows around each local habitat grid cell (habitatGridLocal), from localIndices returned by the getLocalObjects function.

numLocalObsWindows

Vector of numbers of local observation windows around all habitat grid cells, as returned by the getLocalObjects function (object named numLocalIndices). The ith number gives the number of local (original) observation windows for the ith (rescaled) habitat window.

numWindows Number of detection windows. This value (positive integer) is used to truncate

lowerCoords and upperCoords so that extra rows beyond numWindows are ig-

nored.

indicator Binary argument specifying whether the individual is available for detection (in-

dicator = 1) or not (indicator = 0).

log Logical argument, specifying whether to return the log-probability of the distri-

bution.

n Integer specifying the number of realizations to generate. Only n = 1 is sup-

ported.

#### **Details**

The dbernppDetection\_normal distribution is a NIMBLE custom distribution which can be used to model and simulate Bernoulli observations (x) of a single individual in continuous space over a set of detection windows defined by their upper and lower coordinates (lowerCoords, upperCoords). The distribution assumes that an individual's detection probability follows an isotropic multivariate normal centered on the individual's activity center (s) with standard deviation (sd). The local evaluation approach is implemented.

## Value

The (log) probability density of the observation vector x.

#### Author(s)

Wei Zhang and Cyril Milleret

#### References

W. Zhang, J. D. Chipperfield, J. B. Illian, P. Dupont, C. Milleret, P. de Valpine and R. Bischof. 2020. A hierarchical point process model for spatial capture-recapture data. bioRxiv. DOI 10.1101/2020.10.06.325035

C. Milleret, P. Dupont, C. Bonenfant, H. Brøseth, Ø. Flagstad, C. Sutherland and R. Bischof. 2019. A local evaluation of the individual state-space to scale up Bayesian spatial capture-recapture. Ecology and Evolution 9:352-363

# **Examples**

```
# Create habitat grid
coordsHabitatGridCenter <- matrix(c(0.5, 3.5,</pre>
                                     1.5, 3.5,
                                     2.5, 3.5,
                                     3.5, 3.5,
                                     0.5, 2.5,
                                     1.5, 2.5,
                                     2.5, 2.5,
                                     3.5, 2.5,
                                     0.5, 1.5,
                                     1.5, 1.5,
                                     2.5, 1.5,
                                     3.5, 1.5,
                                     0.5, 0.5,
                                     1.5, 0.5,
                                     2.5, 0.5,
                                      3.5, 0.5), ncol = 2, byrow = TRUE)
colnames(coordsHabitatGridCenter) <- c("x","y")</pre>
# Create observation windows
lowerCoords <- matrix(c(1, 1, 2, 1, 1, 2, 2, 2), nrow = 4, byrow = TRUE)
upperCoords <- matrix(c(2, 2, 3, 2, 2, 3, 3, 3), nrow = 4, byrow = TRUE)
colnames(lowerCoords) <- colnames(upperCoords) <- c("x","y")</pre>
plot(coordsHabitatGridCenter[,"y"]~coordsHabitatGridCenter[,"x"],pch=16)
points(lowerCoords[,"y"]~lowerCoords[,"x"],col="red",pch=16)
points(upperCoords[,"y"]~upperCoords[,"x"],col="red",pch=16)
#'
s < -c(1, 1)
sd <- 0.1
baseIntensities <- c(1:4)</pre>
windowIndex <- 4</pre>
numPoints <- 1
numWindows <- 4
indicator <- 1
# Rescale coordinates
ScaledLowerCoords <- scaleCoordsToHabitatGrid(coordsData = lowerCoords,</pre>
                                        coordsHabitatGridCenter = coordsHabitatGridCenter)
ScaledUpperCoords <- scaleCoordsToHabitatGrid(coordsData = upperCoords,</pre>
                                        coordsHabitatGridCenter = coordsHabitatGridCenter)
ScaledUpperCoords$coordsDataScaled[,2] <- ScaledUpperCoords$coordsDataScaled[,2] + 1.5
```

```
ScaledLowerCoords$coordsDataScaled[,2] <- ScaledLowerCoords$coordsDataScaled[,2] - 1.5
habitatMask <- matrix(1, nrow = 4, ncol=4, byrow = TRUE)</pre>
# Create local objects
ObsWindowsLocal <- getLocalObjects(habitatMask = habitatMask,</pre>
                                     coords = ScaledLowerCoords$coordsDataScaled,
                                     dmax=3,
                                     resizeFactor = 1,
                                     plot.check = TRUE
x \leftarrow c(1.1, 1.2)
windowIndex <- getWindowIndex(curCoords = x,</pre>
                               lowerCoords = ScaledLowerCoords$coordsDataScaled,
                               upperCoords =ScaledUpperCoords$coordsDataScaled)
x <- c(x, windowIndex)</pre>
dbernppLocalDetection_normal(x, ScaledLowerCoords$coordsDataScaled,
                              ScaledUpperCoords$coordsDataScaled,
                              s, sd, baseIntensities,
                              ObsWindowsLocal$habitatGrid, ObsWindowsLocal$resizeFactor,
                            ObsWindowsLocal$localIndices,ObsWindowsLocal$numLocalIndices,
                              numWindows, indicator, log = TRUE)
```

dbinomLocal\_exp

Local evaluation of a binomial SCR observation process

# Description

The dbinomLocal\_exp distribution is a NIMBLE custom distribution which can be used to model and simulate binomial observations (x) of a single individual over a set of traps defined by their coordinates *trapCoords* the distribution assumes that an individual's detection probability at any trap follows an exponential function of the distance between the individual's activity center (s) and the trap location. All coordinates (s and trapCoords) should be scaled to the habitat (see scaleCoordsToHabitatGrid)

```
dbinomLocal_exp(
    x,
    detNums = -999,
    detIndices,
    size,
    p0 = -999,
    p0Traps,
    rate,
    s,
    trapCoords,
    localTrapsIndices,
    localTrapsNum,
    resizeFactor = 1,
    habitatGrid,
```

```
indicator,
  lengthYCombined = 0,
  log = 0
)
rbinomLocal_exp(
  n = 1,
  detNums = -999,
  detIndices,
  size,
 p0 = -999,
 p0Traps,
  rate,
  s,
  trapCoords,
  localTrapsIndices,
  localTrapsNum,
  resizeFactor = 1,
  habitatGrid,
  indicator,
  lengthYCombined = 0
)
```

## **Arguments**

Х

Vector of individual detection frequencies. This argument can be provided in two formats: (i) with the *y* object as returned by getSparseY; (ii) with the *yCombined* object as returned by getSparseY Note that when the random generation functionality is used (rbinomLocal\_normal), only the yCombined format can be used. The *yCombined* object combines *detNums*, *x*, and *detIndices* (in that order). When such consolidated representation of the detection data x is used, *detIndices* and *detNums* arguments should not be specified.

detNums

Number of traps with at least one detection recorded in *x*; from the *detNums* object returned by the getSparseY function. This argument should not be specified when the *yCombined* object (returned by getSparseY) is provided as *x* and when detection data are simulated.

detIndices

Vector of indices of traps where the detections in *x* were recorded; from the *detIndices* object returned by the getSparseY function. This argument should not be specified when x is provided as the *yCombined* object (returned by getSparseY) and when detection data are simulated.

size

Vector of the number of trials (zero or more) for each trap (trapCoords).

p0

Baseline detection probability (scalar) used in the half-normal detection function. For trap-specific baseline detection probabilities use argument *p0Traps* 

(vector) instead.

p0Traps

Vector of baseline detection probabilities for each trap used in the half-normal detection function. When p0Traps is used, p0 should not be provided.

rate

Rate parameter of the exponential detection function.

	S	Individual activity center x- and y-coordinates scaled to the habitat (see (scaleCoordsToHabitatGr		
	trapCoords	$Matrix\ of\ x-\ and\ y-coordinates\ of\ all\ traps\ scaled\ to\ the\ habitat\ (see\ (scaleCoordsToHabitatGrid).$		
	localTrapsIndic	res		
		Matrix of indices of local traps around each habitat grid cell, as returned by the getLocalObjects function.		
	localTrapsNum	Vector of numbers of local traps around all habitat grid cells, as returned by the getLocalObjects function.		
	resizeFactor	Aggregation factor used in the getLocalObjects function to reduce the number of habitat grid cells to retrieve local traps for.		
	habitatGrid	Matrix of local habitat grid cell indices, from <i>habitatGrid</i> returned by the getLocalObjects function.		
	indicator	Binary argument specifying whether the individual is available for detection (indicator $= 1$ ) or not (indicator $= 0$ ).		
lengthYCombined				
		The length of the x argument when the ( <i>yCombined</i> ) format of the detection data is provided; from the <i>lengthYCombined</i> object returned by getSparseY		
	log	Logical argument, specifying whether to return the log-probability of the distribution.		
	n	Integer specifying the number of realizations to generate. Only $n = 1$ is supported.		

#### **Details**

The dbinomLocal\_exp distribution incorporates three features to increase computation efficiency (see Turek et al., 2021 <doi.org/10.1002/ecs2.3385> for more details):

- 1. A local evaluation of the detection probability calculation (see Milleret et al., 2019 <doi:10.1002/ece3.4751> for more details)
- 2. A sparse matrix representation (*x*, *detIndices* and *detNums*) of the observation data to reduce the size of objects to be processed.
- 3. An indicator (indicator) to shortcut calculations for individuals unavailable for detection.

The dbinomLocal\_exp distribution requires x- and y- detector coordinates (*trapCoords*) to be scaled to the habitat grid (*habitatGrid*) using the (scaleCoordsToHabitatGrid function.)

When the aim is to simulate detection data:

- 1. *x* should be provided using the *yCombined* object as returned by getSparseY,
- 2. arguments detIndices and detNums should not be provided,
- 3. argument *lengthYCombined* should be provided using the *lengthYCombined* object as returned by getSparseY.

#### Value

The log-likelihood value associated with the vector of detections, given the location of the activity center (s), and the exponential detection function : p = p0 \* exp(-rate \* d).

#### Author(s)

Soumen Dey

#### References

Dey, S., Bischof, R., Dupont, P. P. A., & Milleret, C. (2022). Does the punishment fit the crime? Consequences and diagnosis of misspecified detection functions in Bayesian spatial capture–recapture modeling. Ecology and Evolution, 12, e8600. https://doi.org/10.1002/ece3.8600

## **Examples**

```
# I. DATA SET UP
coordsHabitatGridCenter <- matrix(c(0.5, 3.5,</pre>
                                      1.5, 3.5,
                                      2.5, 3.5,
                                      3.5, 3.5,
                                      0.5, 2.5,
                                      1.5, 2.5,
                                      2.5, 2.5,
                                      3.5, 2.5,
                                      0.5, 1.5,
                                      1.5, 1.5,
                                      2.5, 1.5,
                                      3.5, 1.5,
                                      0.5, 0.5,
                                      1.5, 0.5,
                                      2.5, 0.5,
                                      3.5, 0.5), ncol=2, byrow = TRUE)
colnames(coordsHabitatGridCenter) <- c("x","y")</pre>
# CREATE OBSERVATION WINDOWS
trapCoords \leftarrow matrix(c(1.5, 1.5, 2.5, 1.5, 1.5, 2.5, 2.5, 2.5), nrow = 4, byrow = TRUE)
colnames(trapCoords) <- c("x","y")</pre>
# PLOT CHECK
plot(coordsHabitatGridCenter[,"y"]~coordsHabitatGridCenter[,"x"],pch=16)
points(trapCoords[,"y"]~trapCoords[,"x"],col="red",pch=16)
# PARAMETERS
p0 <- 0.3
rate <- 1/1.5
indicator <- 1
# WE CONSIDER 2 INDIVIDUALS
y < - matrix(c(0, 1, 1, 0,
               0, 1, 0, 1), ncol=4, nrow=2)
s \leftarrow matrix(c(0.5, 1,
              1.6, 2.3),ncol=2,nrow=2)
# RESCALE COORDINATES
ScaledtrapCoords <- scaleCoordsToHabitatGrid(coordsData = trapCoords,</pre>
                                        coordsHabitatGridCenter = coordsHabitatGridCenter)
ScaledtrapCoords<- ScaledtrapCoords$coordsDataScaled</pre>
habitatMask <- matrix(1, nrow = 4, ncol=4, byrow = TRUE)</pre>
```

```
# CREATE LOCAL OBJECTS
TrapLocal <- getLocalObjects(habitatMask = habitatMask,</pre>
                                    coords = ScaledtrapCoords,
                                    dmax=2.5,
                                    resizeFactor = 1,
                                    plot.check = TRUE
)
# GET SPARSE MATRIX
SparseY <- getSparseY(y)</pre>
# II. USING THE DENSITY FUNCTION
 # WE TAKE THE FIRST INDIVIDUAL
i=1
  # OPTION 1: USING THE RANDOM GENERATION FUNCTIONNALITY
dbinomLocal_exp(x=SparseY$y[i,,1],
                   detNums=SparseY$detNums[i],
                   detIndices=SparseY$detIndices[i,,1],
                   size=rep(1,4),
                   p0 = p0,
                   rate= rate,
                   s=s[i,1:2],
                   trapCoords=ScaledtrapCoords,
                   localTrapsIndices=TrapLocal$localIndices,
                   localTrapsNum=TrapLocal$numLocalIndices,
                   resizeFactor=TrapLocal$resizeFactor,
                   habitatGrid=TrapLocal$habitatGrid,
                   indicator=indicator)
  # OPTION 2: USING RANDOM GENERATION FUNCTIONNALITY
  # WE DO NOT PROVIDE THE detNums AND detIndices ARGUMENTS
dbinomLocal_exp(x=SparseY$yCombined[i,,1],
                   size=rep(1,4),
                   p0 = p0,
                   rate= rate,
                   s=s[i,1:2],
                   trapCoords=ScaledtrapCoords,
                   localTrapsIndices=TrapLocal$localIndices,
                   localTrapsNum=TrapLocal$numLocalIndices,
                   resizeFactor=TrapLocal$resizeFactor,
                   habitatGrid=TrapLocal$habitatGrid,
                   indicator=indicator,
                   lengthYCombined = SparseY$lengthYCombined)
# III. USING THE RANDOM GENERATION FUNCTION
rbinomLocal_exp(n=1,
                   size=rep(1,4),
                   p0 = p0,
                   rate= rate,
                   s=s[i,1:2],
                   trapCoords=ScaledtrapCoords,
                   localTrapsIndices=TrapLocal$localIndices,
```

30 dbinomLocal\_normal

```
localTrapsNum=TrapLocal$numLocalIndices,
resizeFactor=TrapLocal$resizeFactor,
habitatGrid=TrapLocal$habitatGrid,
indicator=indicator,
lengthYCombined = SparseY$lengthYCombined)
```

dbinomLocal\_normal

Local evaluation of a binomial SCR detection process

# **Description**

The dbinomLocal\_normal distribution is a NIMBLE custom distribution which can be used to model and simulate binomial observations (x) of a single individual over a set of traps defined by their coordinates *trapCoords* the distribution assumes that an individual's detection probability at any trap follows a half-normal function of the distance between the individual's activity center (s) and the trap location. All coordinates (s and trapCoords) should be scaled to the habitat (see scaleCoordsToHabitatGrid)

```
dbinomLocal_normal(
  detNums = -999,
  detIndices,
  size,
  p0 = -999,
  p0Traps,
  sigma,
  trapCoords,
  localTrapsIndices,
  localTrapsNum,
  resizeFactor = 1,
  habitatGrid,
  indicator,
  lengthYCombined = 0,
  log = 0
)
rbinomLocal_normal(
  n = 1,
  detNums = -999,
  detIndices,
  size,
  p0 = -999,
  p0Traps,
```

dbinomLocal\_normal 31

```
sigma,
s,
trapCoords,
localTrapsIndices,
localTrapsNum,
resizeFactor = 1,
habitatGrid,
indicator,
lengthYCombined = 0
```

#### **Arguments**

x Vector of individual detection frequencies. This argument can be provided in

two formats: (i) with the y object as returned by <code>getSparseY</code>; (ii) with the yCombined object as returned by <code>getSparseY</code> Note that when the random generation functionality is used (rbinomLocal\_normal), only the yCombined format can be used. The yCombined object combines detNums, x, and detIndices (in that order). When such consolidated representation of the detection data x is

used, detIndices and detNums arguments should not be specified.

detNums Number of traps with at least one detection recorded in x; from the detNums

object returned by the getSparseY function. This argument should not be specified when the *yCombined* object (returned by getSparseY) is provided as *x* and

when detection data are simulated.

detIndices Vector of indices of traps where the detections in x were recorded; from the

*detIndices* object returned by the getSparseY function. This argument should not be specified when x is provided as the *yCombined* object (returned by getSparseY

) and when detection data are simulated.

size Vector of the number of trials (zero or more) for each trap (*trapCoords*).

p0 Baseline detection probability (scalar) used in the half-normal detection func-

tion. For trap-specific baseline detection probabilities use argument p0Traps

(vector) instead.

p@Traps Vector of baseline detection probabilities for each trap used in the half-normal

detection function. When *p0Traps* is used, *p0* should not be provided.

sigma Scale parameter of the half-normal detection function.

Individual activity center x- and y-coordinates scaled to the habitat (see (scaleCoordsToHabitatGrid).

trapCoords Matrix of x- and y-coordinates of all traps scaled to the habitat (see (scaleCoordsToHabitatGrid).

localTrapsIndices

Matrix of indices of local traps around each habitat grid cell, as returned by the

getLocalObjects function.

localTrapsNum Vector of numbers of local traps around all habitat grid cells, as returned by the

getLocalObjects function.

 ${\tt resizeFactor} \qquad {\tt Aggregation \ factor \ used \ in \ the \ getLocalObjects \ function \ to \ reduce \ the \ number}$ 

of habitat grid cells to retrieve local traps for.

habitatGrid Matrix of local habitat grid cell indices, from habitatGrid returned by the getLocalObjects

function.

32 dbinomLocal\_normal

indicator	Binary argument specifying whether the individual is available for detection (indicator $= 1$ ) or not (indicator $= 0$ ).	
lengthYCombined		
	The length of the x argument when the ( <i>yCombined</i> ) format of the detection data is provided; from the <i>lengthYCombined</i> object returned by getSparseY	
log	Logical argument, specifying whether to return the log-probability of the distribution.	
n	Integer specifying the number of realizations to generate. Only $n = 1$ is supported.	

## **Details**

The dbinomLocal\_normal distribution incorporates three features to increase computation efficiency (see Turek et al., 2021 <doi.org/10.1002/ecs2.3385> for more details):

- A local evaluation of the detection probability calculation (see Milleret et al., 2019 <doi:10.1002/ece3.4751> for more details)
- 2. A sparse matrix representation (*x*, *detIndices* and *detNums*) of the observation data to reduce the size of objects to be processed.
- 3. An indicator (indicator) to shortcut calculations for individuals unavailable for detection.

The dbinomLocal\_normal distribution requires x- and y- detector coordinates (*trapCoords*) and activity centers coordinates (*s*) to be scaled to the habitat grid (*habitatGrid*) using the (scaleCoordsToHabitatGrid function.)

When the aim is to simulate detection data:

- 1. x should be provided using the yCombined object as returned by getSparseY,
- 2. arguments detIndices and detNums should not be provided,
- 3. argument *lengthYCombined* should be provided using the *lengthYCombined* object as returned by getSparseY.

## Value

The log-likelihood value associated with the vector of detections, given the location of the activity center (s), and the half-normal detection function :  $p = p0 * exp(-d^2/2\sigma^2)$ .

#### Author(s)

Cyril Milleret, Soumen Dey

## **Examples**

```
2.5, 2.5,
                                      3.5, 2.5,
                                      0.5, 1.5,
                                      1.5, 1.5,
                                      2.5, 1.5,
                                      3.5, 1.5,
                                      0.5, 0.5,
                                      1.5, 0.5,
                                      2.5, 0.5,
                                      3.5, 0.5), ncol=2, byrow = TRUE)
colnames(coordsHabitatGridCenter) <- c("x","y")</pre>
# CREATE OBSERVATION WINDOWS
trapCoords < -matrix(c(1.5, 1.5, 2.5, 1.5, 1.5, 2.5, 2.5, 2.5), nrow = 4, byrow = TRUE)
colnames(trapCoords) <- c("x","y")</pre>
# PLOT CHECK
plot(coordsHabitatGridCenter[,"y"]~coordsHabitatGridCenter[,"x"],pch=16)
points(trapCoords[,"y"]~trapCoords[,"x"],col="red",pch=16)
# PARAMETERS
p0 <- 0.2
sigma <- 2
indicator <- 1</pre>
# WE CONSIDER 2 INDIVIDUALS
y \leftarrow matrix(c(0, 1, 1, 0,
              0, 1, 0, 1), ncol=4, nrow=2)
s \leftarrow matrix(c(0.5, 1,
              1.6, 2.3),ncol=2,nrow=2)
# RESCALE COORDINATES
ScaledtrapCoords <- scaleCoordsToHabitatGrid(coordsData = trapCoords,</pre>
                                        coordsHabitatGridCenter = coordsHabitatGridCenter)
ScaledtrapCoords<- ScaledtrapCoords$coordsDataScaled</pre>
habitatMask <- matrix(1, nrow = 4, ncol=4, byrow = TRUE)</pre>
# CREATE LOCAL OBJECTS
TrapLocal <- getLocalObjects(habitatMask = habitatMask,</pre>
                                     coords = ScaledtrapCoords,
                                     dmax=2.5,
                                     resizeFactor = 1,
                                     plot.check = TRUE
)
# GET SPARSE MATRIX
SparseY <- getSparseY(y)</pre>
# II. USING THE DENSITY FUNCTION
 # WE TAKE THE FIRST INDIVIDUAL
i=1
  # OPTION 1: USING THE RANDOM GENERATION FUNCTIONNALITY
dbinomLocal_normal(x=SparseY$y[i,,1],
                    detNums=SparseY$detNums[i],
                    detIndices=SparseY$detIndices[i,,1],
```

```
size=rep(1,4),
                   p0 = p0,
                   sigma= sigma,
                   s=s[i,1:2],
                   trapCoords=ScaledtrapCoords,
                   localTrapsIndices=TrapLocal$localIndices,
                   localTrapsNum=TrapLocal$numLocalIndices,
                   resizeFactor=TrapLocal$resizeFactor,
                   habitatGrid=TrapLocal$habitatGrid,
                   indicator=indicator)
 # OPTION 2: USING RANDOM GENERATION FUNCTIONNALITY
 # WE DO NOT PROVIDE THE detNums AND detIndices ARGUMENTS
dbinomLocal_normal(x=SparseY$yCombined[i,,1],
                   size=rep(1,4),
                   p0 = p0,
                   sigma= sigma,
                   s=s[i,1:2],
                   trapCoords=ScaledtrapCoords,
                   localTrapsIndices=TrapLocal$localIndices,
                   localTrapsNum=TrapLocal$numLocalIndices,
                   resizeFactor=TrapLocal$resizeFactor,
                   habitatGrid=TrapLocal$habitatGrid,
                   indicator=indicator,
                   lengthYCombined = SparseY$lengthYCombined)
# III. USING THE RANDOM GENERATION FUNCTION
rbinomLocal_normal(n=1,
                   size=rep(1,4),
                   p0 = p0,
                   sigma= sigma,
                   s=s[i,1:2],
                   trapCoords=ScaledtrapCoords,
                   localTrapsIndices=TrapLocal$localIndices,
                   localTrapsNum=TrapLocal$numLocalIndices,
                   resizeFactor=TrapLocal$resizeFactor,
                   habitatGrid=TrapLocal$habitatGrid,
                   indicator=indicator,
                   lengthYCombined = SparseY$lengthYCombined)
```

 ${\tt dbinomLocal\_normalPlateau}$ 

Local evaluation of a binomial SCR observation process

## **Description**

The dbinomLocal\_normalPlateau distribution is a NIMBLE custom distribution which can be used to model and simulate binomial observations (x) of a single individual over a set of traps defined by their coordinates *trapCoords* the distribution assumes that an individual's detection probability at any trap follows a half-normal plateau function of the distance between the individual's

activity center (s) and the trap location. With the half-normal plateau function, detection probability remains constant with value p0 for a plateau of width w before declining with scale sigma.

# Usage

```
dbinomLocal_normalPlateau(
 detNums = -999,
 detIndices,
  size,
 p0 = -999,
 p0Traps,
  sigma,
 w = 2,
  s,
  trapCoords,
  localTrapsIndices,
  localTrapsNum,
  resizeFactor = 1,
 habitatGrid,
  indicator,
  lengthYCombined = 0,
 log = 0
)
rbinomLocal_normalPlateau(
  n = 1,
  detNums = -999,
 detIndices,
  size,
 p0 = -999,
  p0Traps,
  sigma,
 w = 2,
  s,
  trapCoords,
  localTrapsIndices,
  localTrapsNum,
  resizeFactor = 1,
 habitatGrid,
  indicator,
  lengthYCombined = 0
)
```

## **Arguments**

Χ

Vector of individual detection frequencies. This argument can be provided in two formats: (i) with the *y* object as returned by getSparseY; (ii) with the *yCombined* object as returned by getSparseY Note that when the random gener-

ation functionality is used (rbinomLocal\_normal), only the yCombined format can be used. The yCombined object combines detNums, x, and detIndices (in that order). When such consolidated representation of the detection data x is used, detIndices and detNums arguments should not be specified.

detNums Number of traps with at least one detection recorded in x; from the detNums

object returned by the <code>getSparseY</code> function. This argument should not be specified when the yCombined object (returned by <code>getSparseY</code>) is provided as x and

when detection data are simulated.

detIndices Vector of indices of traps where the detections in x were recorded; from the

detIndices object returned by the getSparseY function. This argument should not be specified when x is provided as the yCombined object (returned by getSparseY

) and when detection data are simulated.

size Vector of the number of trials (zero or more) for each trap (*trapCoords*).

p0 Baseline detection probability (scalar) used in the half-normal detection func-

tion. For trap-specific baseline detection probabilities use argument pOTraps

(vector) instead.

p@Traps Vector of baseline detection probabilities for each trap used in the half-normal

detection function. When *p0Traps* is used, *p0* should not be provided.

sigma Scale parameter of the half-normal detection function.

w Length of plateau of the half-normal plateau detection function.

Individual activity center x- and y-coordinates scaled to the habitat (see (scaleCoordsToHabitatGrid).

trapCoords Matrix of x- and y-coordinates of all traps scaled to the habitat (see (scaleCoordsToHabitatGrid).

localTrapsIndices

Matrix of indices of local traps around each habitat grid cell, as returned by the

 ${\tt getLocalObjects}\ function.$ 

localTrapsNum Vector of numbers of local traps around all habitat grid cells, as returned by the

getLocalObjects function.

resizeFactor Aggregation factor used in the getLocalObjects function to reduce the number

of habitat grid cells to retrieve local traps for.

habitatGrid Matrix of local habitat grid cell indices, from habitatGrid returned by the getLocalObjects

function.

indicator Binary argument specifying whether the individual is available for detection (in-

dicator = 1) or not (indicator = 0).

lengthYCombined

The length of the x argument when the (yCombined) format of the detection data

is provided; from the lengthYCombined object returned by getSparseY

log Logical argument, specifying whether to return the log-probability of the distri-

oution.

n Integer specifying the number of realizations to generate. Only n = 1 is sup-

ported.

#### **Details**

All coordinates (s and trapCoords) should be scaled to the habitat (see scaleCoordsToHabitatGrid)

The dbinomLocal\_normalPlateau distribution incorporates three features to increase computation efficiency (see Turek et al., 2021 <doi.org/10.1002/ecs2.3385> for more details):

- 1. A local evaluation of the detection probability calculation (see Milleret et al., 2019 <doi:10.1002/ece3.4751> for more details)
- 2. A sparse matrix representation (*x*, *detIndices* and *detNums*) of the observation data to reduce the size of objects to be processed.
- 3. An indicator (indicator) to shortcut calculations for individuals unavailable for detection.

The dbinomLocal\_normalPlateau distribution requires x- and y- detector coordinates (*trapCo-ords*) to be scaled to the habitat grid (*habitatGrid*) using the (scaleCoordsToHabitatGrid function.)

When the aim is to simulate detection data:

- 1. x should be provided using the yCombined object as returned by getSparseY,
- 2. arguments detIndices and detNums should not be provided,
- 3. argument *lengthYCombined* should be provided using the *lengthYCombined* object as returned by getSparseY.

#### Value

The log-likelihood value associated with the vector of detections, given the location of the activity center (s), and the half-normal plateau detection function : p = p0 when d < w and  $p = p0 * exp(-(d-w)^2/\sigma^2)$  when d >= w.

#### Author(s)

Soumen Dey

#### References

Dey, S., Bischof, R., Dupont, P. P. A., & Milleret, C. (2022). Does the punishment fit the crime? Consequences and diagnosis of misspecified detection functions in Bayesian spatial capture—recapture modeling. Ecology and Evolution, 12, e8600. https://doi.org/10.1002/ece3.8600

```
1.5, 2.5,
                                      2.5, 2.5,
                                      3.5, 2.5,
                                      0.5, 1.5,
                                      1.5, 1.5,
                                      2.5, 1.5,
                                      3.5, 1.5,
                                      0.5, 0.5,
                                      1.5, 0.5,
                                      2.5, 0.5,
                                      3.5, 0.5), ncol=2, byrow = TRUE)
colnames(coordsHabitatGridCenter) <- c("x","y")
# CREATE OBSERVATION WINDOWS
trapCoords \leftarrow matrix(c(1.5, 1.5, 2.5, 1.5, 1.5, 2.5, 2.5, 2.5), nrow = 4, byrow = TRUE)
colnames(trapCoords) <- c("x","y")</pre>
# PLOT CHECK
plot(coordsHabitatGridCenter[,"y"]~coordsHabitatGridCenter[,"x"],pch=16)
points(trapCoords[,"y"]~trapCoords[,"x"],col="red",pch=16)
# PARAMETERS
p0 <- 0.25
sigma <- 1
w <- 1.5
indicator <- 1
# WE CONSIDER 2 INDIVIDUALS
y \leftarrow matrix(c(0, 1, 1, 0,
              0, 1, 0, 1), ncol=4, nrow=2)
s \leftarrow matrix(c(0.5, 1,
              1.6, 2.3),ncol=2,nrow=2)
# RESCALE COORDINATES
ScaledtrapCoords <- scaleCoordsToHabitatGrid(coordsData = trapCoords,</pre>
                                        coordsHabitatGridCenter = coordsHabitatGridCenter)
ScaledtrapCoords<- ScaledtrapCoords$coordsDataScaled</pre>
habitatMask <- matrix(1, nrow = 4, ncol=4, byrow = TRUE)</pre>
# CREATE LOCAL OBJECTS
TrapLocal <- getLocalObjects(habitatMask = habitatMask,</pre>
                                     coords = ScaledtrapCoords,
                                     dmax=2.5,
                                     resizeFactor = 1,
                                     plot.check = TRUE
)
# GET SPARSE MATRIX
SparseY <- getSparseY(y)</pre>
# II. USING THE DENSITY FUNCTION
 # WE TAKE THE FIRST INDIVIDUAL
i=1
  # OPTION 1: USING THE RANDOM GENERATION FUNCTIONNALITY
dbinomLocal_normalPlateau(x=SparseY$y[i,,1],
```

dbinom\_vector 39

```
detNums=SparseY$detNums[i],
                   detIndices=SparseY$detIndices[i,,1],
                   size=rep(1,4),
                   p0 = p0,
                   sigma= sigma,
                   w = w,
                   s=s[i,1:2],
                   trapCoords=ScaledtrapCoords,
                   localTrapsIndices=TrapLocal$localIndices,
                   localTrapsNum=TrapLocal$numLocalIndices,
                   resizeFactor=TrapLocal$resizeFactor,
                   habitatGrid=TrapLocal$habitatGrid,
                   indicator=indicator)
 # OPTION 2: USING RANDOM GENERATION FUNCTIONNALITY
 # WE DO NOT PROVIDE THE detNums AND detIndices ARGUMENTS
dbinomLocal_normalPlateau(x=SparseY$yCombined[i,,1],
                   size=rep(1,4),
                   p0 = p0,
                   sigma= sigma,
                   w = w,
                   s=s[i,1:2],
                   trapCoords=ScaledtrapCoords,
                   localTrapsIndices=TrapLocal$localIndices,
                   local Traps Num = TrapLocal \\ snum \\ Local \\ Indices,
                   resizeFactor=TrapLocal$resizeFactor,
                   habitatGrid=TrapLocal$habitatGrid,
                   indicator=indicator,
                   lengthYCombined = SparseY$lengthYCombined)
# III. USING THE RANDOM GENERATION FUNCTION
rbinomLocal_normalPlateau(n=1,
                   size=rep(1,4),
                   p0 = p0,
                   sigma= sigma,
                   w = w,
                   s=s[i,1:2],
                   trapCoords=ScaledtrapCoords,
                   localTrapsIndices=TrapLocal$localIndices,
                   localTrapsNum=TrapLocal$numLocalIndices,
                   resizeFactor=TrapLocal$resizeFactor,
                   habitatGrid=TrapLocal$habitatGrid,
                   indicator=indicator,
                   lengthYCombined = SparseY$lengthYCombined)
```

40 dbinom\_vector

### **Description**

The dbinom\_vector distribution is a vectorized version of the binomial distribution. It can be used to model a vector of binomial realizations. NB: using the vectorized version is beneficial only when the entire joint likelihood of the vector of binomial realizations (x) is calculated simultaneously.

# Usage

```
dbinom_vector(x, size, prob, log = 0)
rbinom_vector(n = 1, size, prob)
```

### **Arguments**

X	Vector of quantiles.
size	Vector of number of trials (zero or more).
prob	Vector of success probabilities on each trial
log	Logical argument, specifying whether to return the log-probability of the distribution.
n	Number of observations. Only $n = 1$ is supported.

## Value

The log-likelihood value associated with the vector of binomial observations.

## Author(s)

Pierre Dupont

dcatState1Alive1Dead 41

```
## create NIMBLE model object
Rmodel <- nimbleModel(code, constants, data, inits)</pre>
## use model object for MCMC, etc.
```

dcatState1Alive1Dead Density and random generation of a categorical distribution describing state transition with one alive and one dead states.

### **Description**

The dcatState1Alive1Dead distribution is a NIMBLE custom distribution which can be used to model and simulate individual state transition. This function can be used to model transitions from one alive and one dead state. If  $z_i$ , t = 1, individual i can be recruited (transition to state 2) with probability prob1To2\_t, so z\_i,t+1 ~ dcat(1- prob1To2\_t, prob1To2\_t, 0,0,0) where prob1To2\_t represent the probability of an unborn individual to be recruited. If  $z_i$ , t = 2, individual i can die and transition to z\_i,t+1=3 with probability prob2To3, or survive with probability 1-prob2To3 Individuals in dead states  $(z_i, t = 3)$  remain in that state with probability 1, the absorbing state. If transition probabilities are spatially variable, a probability vector containing the transition probability value in each habitat window can be provided using the "Hab" arguments (e.g. prob1To2Hab,prob2To3Hab).

```
dcatState1Alive1Dead(
  х,
 Ζ,
 prob1To2 = -999,
 prob1To2Hab,
  prob2To3 = -999,
 prob2To3Hab,
  s,
 habitatGrid,
 log = 0
)
rcatState1Alive1Dead(
  n,
  z,
  prob1To2 = -999,
  prob1To2Hab,
 prob2To3 = -999,
 prob2To3Hab,
  habitatGrid
)
```

42 dcatState1Alive1Dead

### **Arguments**

x	Scalar, individual state z_i,t+1.
z	Scalar, initial individual state z_i,t.
prob1To2	scalar, probability to transition from state 1 to 2.
prob1To2Hab	vector, Spatially-explicit probability to transition from state 2 to 3. The length of the vector should be equal the number of habitat windows in habitatGrid.
prob2To3	scalar, probability to transition from state 2 to 3.
prob2To3Hab	vector, Spatially-explicit probability to transition from state 2 to 3. The length of the vector should be equal the number of habitat windows in habitatGrid.
S	Vector of x- and y-coordinates corresponding to the AC location of the individual. Used to extract transition spatially-explicit probabilities when they are provided.
habitatGrid	Matrix of habitat window indices. Cell values should correspond to the order of habitat windows in prob1To2Hab, prob2To3Hab and in lowerCoords and upperCoords as used in the dbernppAC function.
log	Logical argument, specifying whether to return the log-probability of the distribution. dcatState1Alive1Dead gives the (log) probability density of x. rcatState1Alive2Dead gives a randomly generated individual states conditional on the initial state z.
n	Integer specifying the number of realizations to generate. Only $n=1$ is supported.

## Author(s)

Cyril Milleret

```
# Use the distribution in R
z <- 2
prob1To2 <- 0.2
prob2To3 <- 0.7
lowerCoords <- matrix(c(0, 0, 1, 0, 0, 1, 1, 1), nrow = 4, byrow = TRUE)
upperCoords <- matrix(c(1, 1, 2, 1, 1, 2, 2, 2), nrow = 4, byrow = TRUE)
logIntensities <- log(rep(1,4))</pre>
logSumIntensity <- log(sum(c(1:4)))
habitatGrid <- matrix(c(1:4), nrow = 2, byrow = TRUE)</pre>
numGridRows <- nrow(habitatGrid)</pre>
numGridCols <- ncol(habitatGrid)</pre>
s <- rbernppAC(n=1, lowerCoords, upperCoords, logIntensities, logSumIntensity,</pre>
               habitatGrid, numGridRows, numGridCols)
## No spatial mortality
zPlusOne <- rcatState1Alive1Dead( z = z</pre>
                                  , prob1To2 = prob1To2
                                   , prob2To3 = prob2To3
                                   , s = s
```

```
, habitatGrid = habitatGrid)
zPlus0ne
dcatState1Alive1Dead(x = zPlus0ne
                     , z = z
                     , prob1To2 = prob1To2
                     , prob2To3 = prob2To3
                     , s = s
                     , habitatGrid = habitatGrid)
## With spatial mortality
prob2To3Hab <- c(0.60, 0.70, 0.74, 0.65)
prob1To2Hab <- c(0.4,0.5,0.1,0.3)
zPlusOne <- rcatState1Alive1Dead( z = z</pre>
                                 , prob1To2Hab = prob1To2Hab
                                 , prob2To3Hab = prob2To3Hab
                                , s = s
                                 , habitatGrid = habitatGrid)
zPlusOne
dcatState1Alive1Dead(x = zPlus0ne
                     , z = z
                     , prob1To2Hab = prob1To2Hab
                     , prob2To3Hab = prob2To3Hab
                     , habitatGrid = habitatGrid)
```

dcatState1Alive2Dead

Density and random generation of a categorical distribution describing state transition with one alive and two dead states.

### **Description**

The dcatState1Alive2Dead distribution is a NIMBLE custom distribution which can be used to model and simulate individual state transition. This function can be used to model transitions from one alive and two dead states. If  $z_i$ ,  $t_i = 1$ , individual i can be recruited (transition to state 2) with probability prob1To2\_t, so  $z_i$ ,  $t_i = 1$ , individual i can be recruited. If  $z_i$ ,  $t_i = 2$ , individual i can die from one cause of mortality (e.g. culling) and transition to  $z_i$ ,  $t_i = 2$ , individual i can die from another cause with probability prob2To4  $z_i$ ,  $t_i = 1$ . If the individual does not die it can survive and remain in state 2 with probability (1-(prob2To3+prob2To4)). Individuals in dead states ( $z_i$ ,  $t_i = 3$  or 4) transition to  $z_i$ ,  $t_i + 1 = 4$ , the absorbing state, with probability 1. If transition probabilities are spatially variable, a probability vector containing the transition probability value in each habitat window can be provided using the "Hab" arguments (e.g. prob1To2Hab, prob2To3Hab).

# Usage

```
dcatState1Alive2Dead(
 Х,
 prob1To2 = -999,
 prob1To2Hab,
 prob2To3 = -999,
 prob2To3Hab,
 prob2To4 = -999,
 prob2To4Hab,
 habitatGrid,
 log = 0
rcatState1Alive2Dead(
 n,
  Ζ,
 prob1To2 = -999,
 prob1To2Hab,
 prob2To3 = -999,
  prob2To3Hab,
 prob2To4 = -999,
 prob2To4Hab,
 habitatGrid
)
```

# Arguments

X	Scalar, individual state z_i,t+1.
Z	Scalar, initial individual state z_i,t.
prob1To2	scalar, probability to transition from state 1 to 2.
prob1To2Hab	vector, Spatially-explicit probability to transition from state 2 to 3. The length of the vector should be equal the number of habitat windows in habitatGrid.
prob2To3	scalar, probability to transition from state 2 to 3.
prob2To3Hab	vector, Spatially-explicit probability to transition from state 2 to 3. The length of the vector should be equal the number of habitat windows in habitatGrid.
prob2To4	scalar, probability to transition from state 2 to 4.
prob2To4Hab	vector, Spatially-explicit probability to transition from state 2 to 4. The length of the vector should be equal the number of habitat windows in habitatGrid.
S	Vector of x- and y-coordinates corresponding to the AC location of the individual. Used to extract transition spatially-explicit probabilities when they are provided.

habitatGrid	Matrix of habitat window indices. Cell values should correspond to the order of habitat windows in prob1To2Hab, prob2To3Hab and in lowerCoords and upperCoords as used in the dbernppAC function.
log	Logical argument, specifying whether to return the log-probability of the distribution.
n	Integer specifying the number of realizations to generate. Only $n = 1$ is supported.

### Value

dcatState1Alive2Dead gives the (log) probability density of x. rcatState1Alive2Dead gives a randomly generated individual states conditional on the initial state z.

### Author(s)

Cyril Milleret

```
# Use the distribution in R
z <- 2
prob1To2 <- 0.2
prob2To3 <- 0.4
prob2To4 <- 0.1
lowerCoords <- matrix(c(0, 0, 1, 0, 0, 1, 1, 1), nrow = 4, byrow = TRUE)
upperCoords <- matrix(c(1, 1, 2, 1, 1, 2, 2, 2), nrow = 4, byrow = TRUE)
logIntensities <- log(rep(1,4))</pre>
logSumIntensity <- log(sum(c(1:4)))</pre>
habitatGrid <- matrix(c(1:4), nrow = 2, byrow = TRUE)</pre>
numGridRows <- nrow(habitatGrid)</pre>
numGridCols <- ncol(habitatGrid)</pre>
s <- rbernppAC(n=1, lowerCoords, upperCoords, logIntensities, logSumIntensity,</pre>
                 habitatGrid, numGridRows, numGridCols)
## No spatial mortality
zPlusOne <- rcatState1Alive2Dead( z = z
                                  , prob1To2 = prob1To2
                                  , prob2To3 = prob2To3
                                  , prob2To4 = prob2To4
                                  , s = s
                                  , habitatGrid = habitatGrid)
dcatState1Alive2Dead(x = zPlus0ne
                      z = z
                      , prob1To2 = prob1To2
                      , prob2To3 = prob2To3
                      , prob2To4 = prob2To4
                      , s = s
                      , habitatGrid = habitatGrid)
```

```
## With spatial mortality
prob2To3Hab <- c(0.10, 0.20, 0.15, 0.30)
prob2To4Hab <- c(0.13, 0.21, 0.12, 0.08)
phiSpatial <- 1-(prob2To3Hab+prob2To4Hab)</pre>
zPlusOne <- rcatState1Alive2Dead( z = z
                                 , prob1To2Hab = prob1To2Hab
                                 , prob2To3Hab = prob2To3Hab
                                 , prob2To4Hab = prob2To4Hab
                                 , s = s
                                 , habitatGrid = habitatGrid)
dcatState1Alive2Dead(x = zPlus0ne
                     , z = z
                     , prob1To2Hab = prob1To2Hab
                     , prob2To3Hab = prob2To3Hab
                     , prob2To4Hab = prob2To4Hab
                     , habitatGrid = habitatGrid)
```

dcatState2Alive2Dead

Density and random generation of a categorical distribution describing state transition with two alive and two dead states.

### **Description**

The dcatState2Alive2Dead distribution is a NIMBLE custom distribution which can be used to model and simulate individual state transition. This function can be used to model transitions from two alive and two dead states. If  $z_i$ ,  $t_i = 1$ , individual i can be recruited (transition to state 2) with probability prob1To2\_t, so  $z_i$ ,  $t_i = 1$ , individual i can be recruited. If  $z_i$ ,  $t_i = 2$ , individual i can die from one cause of mortality (e.g. culling) and transition to  $z_i$ ,  $t_i = 2$ , individual i can die from another cause with probability prob2To5  $z_i$ ,  $t_i = 1$ . If the individual does not die (1-(prob2To4+prob2To5)), it can either transition to the second state alive ( $z_i$ ,  $t_i = 3$ , individual i can die from one cause of mortality (e.g. culling) and transition to  $z_i$ ,  $t_i = 1$ , individual i can die from one cause of mortality (e.g. culling) and transition to  $z_i$ ,  $t_i = 1$ , individual does not die (1-(prob3To4+prob3To5)), the individual remain in state 3. Individuals in dead states ( $z_i$ ,  $t_i = 4$  or 5) transition to  $z_i$ ,  $t_i = 1$ , the absorbing state, with probability 1. If transition probabilities are spatially variable, a probability vector containing the transition probability value in each habitat window can be provided using the "Hab" arguments (e.g. prob1To2Hab,prob2To3Hab).

### Usage

dcatState2Alive2Dead(

```
Х,
  Ζ,
 prob1To2 = -999,
 prob1To2Hab,
 prob2To3 = -999,
 prob2To3Hab,
 prob2To4 = -999,
 prob2To4Hab,
 prob3To4 = -999,
 prob3To4Hab,
 prob2To5 = -999,
 prob2To5Hab,
  prob3To5 = -999,
  prob3To5Hab,
 habitatGrid,
  log = 0
)
rcatState2Alive2Dead(
 n,
 z,
 prob1To2 = -999,
 prob1To2Hab,
 prob2To3 = -999,
 prob2To3Hab,
 prob2To4 = -999,
 prob2To4Hab,
 prob3To4 = -999,
 prob3To4Hab,
 prob2To5 = -999,
 prob2To5Hab,
 prob3To5 = -999,
 prob3To5Hab,
 s,
 habitatGrid
)
```

# Arguments

Χ	Scalar, individual state z_i,t+1.
Z	Scalar, initial individual state z_i,t.
prob1To2	scalar, probability to transition from state 1 to 2.
prob1To2Hab	vector, Spatially-explicit probability to transition from state 2 to 3. The length of the vector should be equal the number of habitat windows in habitatGrid.
prob2To3	scalar, probability to transition from state 2 to 3.
prob2To3Hab	vector, Spatially-explicit probability to transition from state 2 to 3. The length of the vector should be equal the number of habitat windows in habitatGrid.

prob2To4	scalar, probability to transition from state 2 to 4.
prob2To4Hab	vector, Spatially-explicit probability to transition from state 2 to 4. The length of the vector should be equal the number of habitat windows in habitatGrid.
prob3To4	scalar, probability to transition from state 3 to 4.
prob3To4Hab	vector, Spatially-explicit probability to transition from state 3 to 4. The length of the vector should be equal the number of habitat windows in habitatGrid.
prob2To5	scalar, probability to transition from state 2 to 5.
prob2To5Hab	vector, Spatially-explicit probability to transition from state 2 to 5. The length of the vector should be equal the number of habitat windows in habitatGrid.
prob3To5	scalar, probability to transition from state 3 to 5.
prob3To5Hab	vector, Spatially-explicit probability to transition from state 3 to 5. The length of the vector should be equal the number of habitat windows in habitatGrid.
S	Vector of x- and y-coordinates corresponding to the AC location of the individual. Used to extract transition spatially-explicit probabilities when they are provided.
habitatGrid	Matrix of habitat window indices. Cell values should correspond to the order of habitat windows in prob1To2Hab, prob2To3Hab and in lowerCoords and upperCoords as used in the dbernppAC function.
log	Logical argument, specifying whether to return the log-probability of the distribution.
n	Integer specifying the number of realizations to generate. Only $n=1$ is supported.

#### Value

## Author(s)

Cyril Milleret

```
# Use the distribution in R

z <- 3
prob1To2 <- 0.2
prob2To3 <- 0.4
prob2To4 <- 0.1
prob2To5 <- 0.1

prob3To4 <- 0.2
prob3To5 <- 0.1

lowerCoords <- matrix(c(0, 0, 1, 0, 0, 1, 1, 1), nrow = 4, byrow = TRUE)</pre>
```

```
upperCoords <- matrix(c(1, 1, 2, 1, 1, 2, 2, 2), nrow = 4, byrow = TRUE)
logIntensities <- log(rep(1,4))</pre>
logSumIntensity <- log(sum(c(1:4)))</pre>
habitatGrid <- matrix(c(1:4), nrow = 2, byrow = TRUE)
numGridRows <- nrow(habitatGrid)</pre>
numGridCols <- ncol(habitatGrid)</pre>
s <- rbernppAC(n=1, lowerCoords, upperCoords, logIntensities, logSumIntensity,</pre>
              habitatGrid, numGridRows, numGridCols)
## No spatial mortality
zPlusOne <- rcatState2Alive2Dead( z = z</pre>
                                   , prob1To2 = prob1To2
                                   , prob2To3 = prob2To3
                                   , prob2To4 = prob2To4
                                   , prob2To5 = prob2To5
                                   , prob3To4 = prob3To4
                                   , prob3To5 = prob3To5
                                   , s = s
                                   , habitatGrid = habitatGrid)
dcatState2Alive2Dead(x = zPlus0ne
                       , z = z
                       , prob1To2 = prob1To2
                       , prob2To3 = prob2To3
                       , prob2To4 = prob2To4
                       , prob2To5 = prob2To5
                       , prob3To4 = prob3To4
                       , prob3To5 = prob3To5
                       , s = s
                       , habitatGrid = habitatGrid)
## With spatial mortality
prob2To3Hab <- runif(length(habitatGrid),0,0.1)</pre>
prob2To4Hab <- runif(length(habitatGrid),0,0.1)</pre>
prob2To5Hab <- runif(length(habitatGrid),0,0.1)</pre>
prob3To4Hab <- runif(length(habitatGrid),0,0.1)</pre>
prob3To5Hab <- runif(length(habitatGrid),0,0.1)</pre>
zPlusOne <- rcatState2Alive2Dead( z = z</pre>
                                   , prob1To2 = prob1To2
                                   , prob2To3Hab = prob2To3Hab
                                   , prob2To4Hab = prob2To4Hab
                                   , prob2To5Hab = prob2To5Hab
                                   , prob3To4Hab = prob3To4Hab
                                   , prob3To5Hab = prob3To5Hab
                                   , s = s
                                   , habitatGrid = habitatGrid)
dcatState2Alive2Dead(x = zPlus0ne
                       z = z
                       , prob1To2 = prob1To2
```

50 dDispersal\_exp

```
, prob2To3Hab = prob2To3Hab
, prob2To4Hab = prob2To5Hab
, prob2To5Hab = prob2To5Hab
, prob3To4Hab = prob3To4Hab
, prob3To5Hab = prob3To5Hab
, s = s
, habitatGrid = habitatGrid)
```

dDispersal\_exp

Bivariate exponential dispersal distribution for activity centers

### **Description**

This function is deprecated, and it will be removed from a future release.

## Usage

```
dDispersal_exp(x, s, rate, log)
rDispersal_exp(n, s, rate)
```

# Arguments

x	Bivariate activity center coordinates (at time t+1).
S	Current location of the bivariate activity center (at time t).
rate	Rate parameter of the exponential distribution for dispersal distance.
log	Logical argument, specifying whether to return the log-probability of the distribution.
n	Integer specifying the number of realisations to generate. Only $n = 1$ is supported.

#### **Details**

The dDispersal\_exp distribution is a bivariate distribution which can be used to model the latent bivariate activity centers (ACs) of individuals in a population. This distribution models the situation when individual AC dispersal is uniform in direction (that is, dispersal occurs in a direction theta, where theta is uniformly distributed on [-pi, pi]), and with an exponential distribution for the radial dispersal distance.

The dDispersal\_exp distribution models the location of an AC at time (t+1), conditional on the previous AC location at time (t) and the rate parameter (rate) of the exponential distribution for dispersal distance.

#### Value

The log-probability value associated with the bivariate activity center location x, given the current activity center s, and the rate parameter of the exponential dispersal distance distribution.

dHabitatMask 51

### Author(s)

Daniel Turek

# **Examples**

```
## Not run:
## define model code
code <- nimbleCode({</pre>
    lambda ~ dgamma(0.001, 0.001)
    for(i in 1:N) {
        AC[i, 1, 1] ~ dunif(0, 100)
        AC[i, 2, 1] ~ dunif(0, 100)
        for(t in 2:T) {
            AC[i, 1:2, t+1] \sim dDispersal_exp(s = AC[i, 1:2, t], rate = lambda)
    }
})
constants <- list(N = 10, T = 6)
## create NIMBLE model object
Rmodel <- nimbleModel(code, constants)</pre>
## use model object for MCMC, etc.
## End(Not run)
```

dHabitatMask

Ones trick distribution for irregular habitat shapes

### **Description**

The dHabitatMask distribution checks and ensures that the proposed activity center location (s) falls within the suitable habitat (defined in the binary matrix habitatMask).

```
dHabitatMask(x, s, xmax, xmin, ymax, ymin, habitatMask, log = 0)
rHabitatMask(n, s, xmax, xmin, ymax, ymin, habitatMask)
```

52 dHabitatMask

## **Arguments**

X	Ones trick data.
S	Bivariate activity center coordinates.
xmax	Maximum of trap location x-coordinates.
xmin	Minimum of trap location x-coordinates.
ymax	Maximum of trap location y-coordinates.
ymin	Minimum of trap location y-coordinates.
habitatMask	A binary matrix object indicating which cells are considered as suitable habitat.
log	Logical argument, specifying whether to return the log-probability of the distribution.
n	Integer specifying the number of realisations to generate. Only $n=1$ is supported.

### **Details**

The rHabitatMask function returns the value of the habitat mask cell (0 or 1) where the proposed activity center falls. See also M. Meredith: SECR in BUGS/JAGS with patchy habitat.

#### Value

The log-likelihood value associated with the bivariate activity center location s being in the suitable habitat (i.e. 0 if it falls within the habitat mask and -Inf otherwise).

# Author(s)

Daniel Turek

```
data <- list(OK = rep(1, N))
inits <- list(s = array(runif(2*N, 0, 100), c(N,2)))
## create NIMBLE model object
Rmodel <- nimbleModel(code, constants, data, inits)
## use model object for MCMC, etc.</pre>
```

dmultiLocal\_normal

Local evaluation of a multinomial SCR detection process

## **Description**

The dmultilocal\_normal distribution is a NIMBLE custom distribution which can be used to model and simulate multinomial observations (x) of a single individual over a set of traps defined by their coordinates *trapCoords* the distribution assumes that an individual's detection probability at any trap follows a half-normal function of the distance between the individual's activity center (s) and the trap location. All coordinates (s and trapCoords) should be scaled to the habitat (see scaleCoordsToHabitatGrid)

```
dmultiLocal_normal(
 detNums = -999,
 detIndices,
  size,
 p0 = -999,
  p0Traps,
  sigma,
  s,
  trapCoords,
  localTrapsIndices,
  localTrapsNum,
  resizeFactor = 1,
  habitatGrid,
  indicator,
  lengthYCombined = 0,
  log = 0
rmultiLocal_normal(
  n = 1,
 detNums = -999,
 detIndices,
```

```
size,
p0 = -999,
p0Traps,
sigma,
s,
trapCoords,
localTrapsIndices,
localTrapsNum,
resizeFactor = 1,
habitatGrid,
indicator,
lengthYCombined = 0
```

### **Arguments**

Х

Vector of individual detection frequencies. This argument can be provided in two formats: (i) with the *y* object as returned by the getSparseY function; (ii) with the *yCombined* object as returned by getSparseY. Note that when the random generation functionality is used (rmultiLocal\_normal), only the *yCombined* format can be used. The *yCombined* object combines *detNums*, *x*, and *detIndices* (in that order). When such consolidated representation of the detection data *x* is used, *detIndices* and *detNums* arguments should not be specified

detNums

umber of traps with at least one detection recorded in x; from the detNums object returned by the getSparseY function. This argument should not be specified when the yCombined object (returned by getSparseY) is provided as x and when detection data are simulated.

detIndices

Vector of indices of traps where the detections in *x* were recorded; from the *detIndices* object returned by the getSparseY function. This argument should not be specified when x is provided as the *yCombined* object (returned by getSparseY) and when detection data are simulated.

,

size Number of occasions.

p0 Baseline detection probability (scalar) used in the half-normal detection func-

tion. For trap-specific baseline detection probabilities use argument p0Traps

(vector) instead.

p0Traps Vector of baseline detection probabilities for each trap used in the half-normal

detection function. When p0Traps is used, p0 should not be provided.

sigma Scale parameter of the half-normal detection function.

Individual activity center x- and y-coordinates scaled to the habitat (see (scaleCoordsToHabitatGrid)).

trapCoords Matrix of x- and y-coordinates of all traps scaled to the habitat (see (scaleCoordsToHabitatGrid)).

localTrapsIndices

Matrix of indices of local traps around each habitat grid cell, as returned by the

getLocalObjects function.

localTrapsNum Vector of numbers of local traps around all habitat grid cells, as returned by the

getLocalObjects function.

resizeFactor	Aggregation factor used in the getLocalObjects function to reduce the number of habitat grid cells to retrieve local traps for.	
habitatGrid	Matrix of local habitat grid cell indices, from $habitatGrid$ returned by the $getLocalObjects$ function.	
indicator	Binary argument specifying whether the individual is available for detection (indicator $= 1$ ) or not (indicator $= 0$ ).	
lengthYCombined		
	The length of the x argument when the ( <i>yCombined</i> ) format of the detection data is provided; from the <i>lengthYCombined</i> object returned by getSparseY	
log	Logical argument, specifying whether to return the log-probability of the distribution.	
n	Integer specifying the number of realizations to generate. Only $n = 1$ is supported.	

### **Details**

The dmultiLocal\_normal distribution incorporates three features to increase computation efficiency (see Turek et al., 2021 <doi.org/10.1002/ecs2.3385> for more details):

- 1. A local evaluation of the detection probability calculation (see Milleret et al., 2019 <doi:10.1002/ece3.4751> for more details)
- 2. A sparse matrix representation (*x*, *detIndices* and *detNums*) of the observation data to reduce the size of objects to be processed.
- 3. An indicator (indicator) to shortcut calculations for individuals unavailable for detection.

The dmultiLocal\_normal distribution requires x- and y- detector coordinates (*trapCoords*) and activity centers coordinates (*s*) to be scaled to the habitat grid (*habitatGrid*) using the (scaleCoordsToHabitatGrid function.)

When the aim is to simulate detection data:

- 1. x should be provided using the yCombined object as returned by getSparseY,
- 2. arguments detIndices and detNums should not be provided,
- 3. argument *lengthYCombined* should be provided using the *lengthYCombined* object as returned by getSparseY.

#### Value

The log-likelihood value associated with the vector of detections, given the location of the activity center (s), and the half-normal detection function :  $p = p0 * exp(-d^2/2\sigma^2)$ .

#### Author(s)

Soumen Dey, Cyril Milleret

```
# I. DATA SET UP
coordsHabitatGridCenter <- matrix(c(0.5, 3.5,</pre>
                                     1.5, 3.5,
                                     2.5, 3.5,
                                     3.5, 3.5,
                                     0.5, 2.5,
                                     1.5, 2.5,
                                     2.5, 2.5,
                                     3.5, 2.5,
                                     0.5, 1.5,
                                     1.5, 1.5,
                                     2.5, 1.5,
                                     3.5, 1.5,
                                     0.5, 0.5,
                                     1.5, 0.5,
                                     2.5, 0.5,
                                     3.5, 0.5), ncol=2, byrow = TRUE)
colnames(coordsHabitatGridCenter) <- c("x","y")</pre>
# CREATE OBSERVATION WINDOWS
trapCoords \leftarrow matrix(c(1.5, 1.5, 2.5, 1.5, 1.5, 2.5, 2.5, 2.5), nrow = 4, byrow = TRUE)
colnames(trapCoords) <- c("x","y")</pre>
# PLOT CHECK
plot(coordsHabitatGridCenter[,"y"]~coordsHabitatGridCenter[,"x"],pch=16)
points(trapCoords[,"y"]~trapCoords[,"x"],col="red",pch=16)
# PARAMETERS
p0 < -0.2
sigma <- 2
indicator <- 1</pre>
# WE CONSIDER 2 INDIVIDUALS
y \leftarrow matrix(c(2, 1, 1, -1, 1, 4, -1, \#id\#1 detected 2 times at detector 1 and 4
              2, 2, -1, -1, 3, -1, -1), #id#2 detected 2 times at detector 3
              ncol=7, nrow=2, byrow = TRUE)
s < - matrix(c(0.5, 1,
              1.6, 2.3), ncol=2, nrow=2)
# RESCALE COORDINATES
ScaledtrapCoords <- scaleCoordsToHabitatGrid(coordsData = trapCoords,</pre>
                                        coordsHabitatGridCenter = coordsHabitatGridCenter)
ScaledtrapCoords<- ScaledtrapCoords$coordsDataScaled
habitatMask <- matrix(1, nrow = 4, ncol=4, byrow = TRUE)</pre>
# CREATE LOCAL OBJECTS
TrapLocal <- getLocalObjects(habitatMask = habitatMask,</pre>
                                    coords = ScaledtrapCoords,
```

```
dmax=2.5,
                                    resizeFactor = 1,
                                    plot.check = TRUE
)
# GET SPARSE MATRIX
#SparseY <- getSparseY(y)</pre>
# II. USING THE DENSITY FUNCTION
# WE TAKE THE FIRST INDIVIDUAL
i=1
 # OPTION 1: USING THE RANDOM GENERATION FUNCTIONNALITY
dmultiLocal_normal(x=y[i,]
                   size=3
                   p0 = p0
                   sigma= sigma
                   s=s[i,1:2]
                   trapCoords=ScaledtrapCoords
                   local Traps Indices = Trap Local \$ local Indices
                   local Traps {\tt Num=TrapLocal \$ numLocal Indices}
                   resizeFactor=TrapLocal$resizeFactor
                   habitatGrid=TrapLocal$habitatGrid
                   indicator=indicator
                   lengthYCombined = ncol(y)
                   )
# III. USING THE RANDOM GENERATION FUNCTION
rmultiLocal_normal(n=1,
                   size=3,
                   p0 = p0,
                   sigma= sigma,
                   s=s[i,1:2],
                   trapCoords=ScaledtrapCoords,
                   localTrapsIndices=TrapLocal$localIndices,
                   localTrapsNum=TrapLocal$numLocalIndices,
                   resize Factor = TrapLocal \\ \$ resize Factor,
                   habitatGrid=TrapLocal$habitatGrid,
                   indicator=indicator,
                   lengthYCombined = ncol(y))
```

58 dnormalizer

dnormalizer

Normalizing constant generator

### **Description**

A normalizer used for normalizing nimble distributions. It is particularly useful for fitting dpoisppDetection\_normal and dpoisppLocalDetection\_normal models using the semi-complete data likelihood approach.

# Usage

```
dnormalizer(x, logNormConstant, log = 0)
rnormalizer(n, logNormConstant)
```

# Arguments

x Input data, which can be any scalar and will not influence the return value.

logNormConstant

Normalizing constant on a log scale.

log Logical. If TRUE return the log normalizing constant. Otherwise return the nor-

malizing constant.

n Integer specifying the number of realisations to generate. Only n = 1 is sup-

ported.

### Value

The normalizing constant.

## Author(s)

Wei Zhang

```
dnormalizer(1, log(0.5), log = TRUE)
dnormalizer(0, log(0.5), log = FALSE)
```

dpoisLocal\_normal

Local evaluation of a Poisson SCR detection process

### **Description**

The dpoisLocal\_normal distribution is a NIMBLE custom distribution which can be used to model and simulate Poisson observations (x) of a single individual over a set of traps defined by their coordinates *trapCoords* the distribution assumes that an individual's detection probability at any trap follows a half-normal function of the distance between the individual's activity center (s) and the trap location. All coordinates (s and trapCoords) should be scaled to the habitat (see scaleCoordsToHabitatGrid)

```
dpoisLocal_normal(
  detNums = -999,
  detIndices,
  lambda = -999,
  lambdaTraps,
  sigma,
  trapCoords,
  localTrapsIndices,
  localTrapsNum,
  resizeFactor = 1,
  habitatGrid,
  indicator,
  lengthYCombined = 0,
  log = 0
)
rpoisLocal_normal(
  n = 1,
  detNums = -999,
  detIndices,
  lambda = -999,
  lambdaTraps,
  sigma,
  s,
  trapCoords,
  localTrapsIndices,
  localTrapsNum,
  resizeFactor = 1,
  habitatGrid,
  indicator,
  lengthYCombined = 0
```

)

#### **Arguments**

vector of individual detection frequencies. This argument can be provided in two formats: (i) with the y object as returned by getSparseY; (ii) with the

two formats: (1) with the y object as returned by getSparseY; (11) with the yCombined object as returned by getSparseY Note that when the random generation functionality is used (rpoisLocal\_normal), only the yCombined format can be used. The yCombined object combines detNums, x, and detIndices (in that order). When such consolidated representation of the detection data x is

used, detIndices and detNums arguments should not be specified.

detNums Number of traps with at least one detection recorded in x; from the detNums

object returned by the getSparseY function. This argument should not be specified when the *yCombined* object (returned by getSparseY) is provided as *x* and

when detection data are simulated.

detIndices Vector of indices of traps where the detections in x were recorded; from the

*detIndices* object returned by the getSparseY function. This argument should not be specified when x is provided as the *yCombined* object (returned by getSparseY

) and when detection data are simulated.

lambda Baseline detection rate used in the half-normal detection function.

lambdaTraps Vector of baseline detection rate for each trap used in the half-normal detection

function. When *lambdaTraps* is used, *lambda* should not be provided.

sigma Scale parameter of the half-normal detection function.

Individual activity center x- and y-coordinates scaled to the habitat (see (scaleCoordsToHabitatGrid).

trapCoords Matrix of x- and y-coordinates of all traps scaled to the habitat (see (scaleCoordsToHabitatGrid).

localTrapsIndices

Matrix of indices of local traps around each habitat grid cell, as returned by the

getLocalObjects function.

localTrapsNum Vector of numbers of local traps around all habitat grid cells, as returned by the

getLocalObjects function.

resizeFactor Aggregation factor used in the getLocalObjects function to reduce the number

of habitat grid cells to retrieve local traps for.

habitatGrid Matrix of local habitat grid cell indices, from habitatGrid returned by the getLocalObjects

function.

indicator Binary argument specifying whether the individual is available for detection (in-

dicator = 1) or not (indicator = 0).

lengthYCombined

The length of the x argument when the (yCombined) format of the detection data

is provided; from the *lengthYCombined* object returned by getSparseY

log Logical argument, specifying whether to return the log-probability of the distri-

bution.

n Integer specifying the number of realizations to generate. Only n = 1 is sup-

ported.

#### **Details**

The dpoisLocal\_normal distribution incorporates three features to increase computation efficiency (see Turek et al., 2021 <doi.org/10.1002/ecs2.3385> for more details):

- A local evaluation of the detection probability calculation (see Milleret et al., 2019 <doi:10.1002/ece3.4751> for more details)
- 2. A sparse matrix representation (*x*, *detIndices* and *detNums*) of the observation data to reduce the size of objects to be processed.
- 3. An indicator (indicator) to shortcut calculations for individuals unavailable for detection.

The dpoisLocal\_normal distribution requires x- and y- detector coordinates (*trapCoords*) and activity centers coordinates (*s*) to be scaled to the habitat grid (*habitatGrid*) using the (scaleCoordsToHabitatGrid function.)

When the aim is to simulate detection data:

- 1. *x* should be provided using the *yCombined* object as returned by getSparseY,
- 2. arguments detIndices and detNums should not be provided,
- 3. argument *lengthYCombined* should be provided using the *lengthYCombined* object as returned by getSparseY.

#### Value

The log-likelihood value associated with the vector of detections, given the location of the activity center (s), and the half-normal detection function :  $p = lambda * exp(-d^2/2\sigma^2)$ .

## Author(s)

Cyril Milleret, Soumen Dey

```
# I. DATA SET UP
coordsHabitatGridCenter <- matrix(c(0.5, 3.5,</pre>
                                       1.5, 3.5,
                                       2.5, 3.5,
                                       3.5, 3.5,
                                       0.5, 2.5,
                                       1.5, 2.5,
                                       2.5, 2.5,
                                       3.5, 2.5,
                                       0.5, 1.5,
                                       1.5, 1.5,
                                       2.5, 1.5,
                                       3.5, 1.5,
                                       0.5, 0.5,
                                       1.5, 0.5,
                                       2.5, 0.5,
                                       3.5, 0.5), ncol=2, byrow = TRUE)
colnames(coordsHabitatGridCenter) <- c("x","y")</pre>
# CREATE OBSERVATION WINDOWS
```

```
trapCoords \leftarrow matrix(c(1.5, 1.5, 2.5, 1.5, 1.5, 2.5, 2.5, 2.5), nrow = 4, byrow = TRUE)
colnames(trapCoords) <- c("x","y")</pre>
# PLOT CHECK
plot(coordsHabitatGridCenter[,"y"]~coordsHabitatGridCenter[,"x"],pch=16)
points(trapCoords[,"y"]~trapCoords[,"x"],col="red",pch=16)
# PARAMETERS
lambda <- 0.2
sigma <- 2
indicator <- 1
# WE CONSIDER 2 INDIVIDUALS
y \leftarrow matrix(c(0, 1, 1, 0,
              0, 1, 0, 1), ncol=4, nrow=2)
s \leftarrow matrix(c(0.5, 1,
              1.6, 2.3),ncol=2,nrow=2)
# RESCALE COORDINATES
ScaledtrapCoords <- scaleCoordsToHabitatGrid(coordsData = trapCoords,</pre>
                                        coordsHabitatGridCenter = coordsHabitatGridCenter)
ScaledtrapCoords<- ScaledtrapCoords$coordsDataScaled</pre>
habitatMask <- matrix(1, nrow = 4, ncol=4, byrow = TRUE)</pre>
# CREATE LOCAL OBJECTS
TrapLocal <- getLocalObjects(habitatMask = habitatMask,</pre>
                                     coords = ScaledtrapCoords,
                                     dmax=2.5,
                                     resizeFactor = 1,
                                     plot.check = TRUE
)
# GET SPARSE MATRIX
SparseY <- getSparseY(y)</pre>
# II. USING THE DENSITY FUNCTION
 # WE TAKE THE FIRST INDIVIDUAL
i=1
  # OPTION 1: USING THE RANDOM GENERATION FUNCTIONNALITY
dpoisLocal_normal(x=SparseY$y[i,,1],
                    detNums=SparseY$detNums[i],
                    detIndices=SparseY$detIndices[i,,1],
                    lambda = lambda,
                    sigma= sigma,
                    s=s[i,1:2],
                    trapCoords=ScaledtrapCoords,
                    localTrapsIndices=TrapLocal$localIndices,
                    localTrapsNum=TrapLocal$numLocalIndices,
                    resizeFactor=TrapLocal$resizeFactor,
                    habitatGrid=TrapLocal$habitatGrid,
                    indicator=indicator)
  # OPTION 2: USING RANDOM GENERATION FUNCTIONNALITY
  # WE DO NOT PROVIDE THE detNums AND detIndices ARGUMENTS
```

dpoisppAC 63

```
dpoisLocal_normal(x=SparseY$yCombined[i,,1],
                   lambda = lambda,
                   sigma= sigma,
                   s=s[i,1:2],
                   trapCoords=ScaledtrapCoords,
                   localTrapsIndices=TrapLocal$localIndices,
                   localTrapsNum=TrapLocal$numLocalIndices,
                   resizeFactor=TrapLocal$resizeFactor,
                   habitatGrid=TrapLocal$habitatGrid,
                   indicator=indicator,
                   lengthYCombined = SparseY$lengthYCombined)
# III. USING THE RANDOM GENERATION FUNCTION
rpoisLocal_normal(n=1,
                   lambda = lambda,
                   sigma= sigma,
                   s=s[i,1:2],
                   trapCoords=ScaledtrapCoords,
                   localTrapsIndices=TrapLocal$localIndices,
                   localTrapsNum=TrapLocal$numLocalIndices,
                   resizeFactor=TrapLocal$resizeFactor,
                   habitatGrid=TrapLocal$habitatGrid,
                   indicator=indicator,
                   lengthYCombined = SparseY$lengthYCombined)
```

dpoisppAC

Poisson point process for the distribution of activity centers

### **Description**

Density and random generation functions of the Poisson point process for the distribution of activity centers. The dpoisppAC distribution is a NIMBLE custom distribution which can be used to model and simulate activity center locations (x) of multiple individual in continuous space over a set of habitat windows defined by their upper and lower coordinates (lowerCoords, upperCoords). The distribution assumes that activity centers follow a Poisson point process with intensity = exp(logIntensities). All coordinates (s and trapCoords) should be scaled to the habitat (scaleCoordsToHabitatGrid).

```
dpoisppAC(
    x,
    lowerCoords,
    upperCoords,
    logIntensities,
    sumIntensity,
    habitatGrid,
    numGridRows,
    numGridCols,
```

64 dpoisppAC

```
numPoints,
log = 0
)

rpoisppAC(
    n,
    lowerCoords,
    upperCoords,
    logIntensities,
    sumIntensity,
    habitatGrid,
    numGridRows,
    numGridCols,
    numPoints
)
```

### **Arguments**

Х

Matrix of x- and y-coordinates of a set of spatial points (AC locations) scaled to the habitat (scaleCoordsToHabitatGrid). Each row corresponds to a point.

lowerCoords, upperCoords

Matrices of lower and upper x- and y-coordinates of all detection windows scaled to the habitat (see (scaleCoordsToHabitatGrid). One row for each window, Each window should be of size 1x1.

logIntensities Vector of log habitat intensities for all habitat windows.

sumIntensity

Sum of the habitat intensities over all windows. Provided as an argument for computational speed, instead of calculating it in the function.

habitatGrid

Matrix of habitat window indices. Cell values should correspond to the order of habitat windows in lowerCoords, upperCoords, and logIntensities. When the habitat grid only consists of a single row or column of windows, an additional row or column of dummy indices has to be added because the nimble model code requires a matrix.

numGridRows, numGridCols

Numbers of rows and columns of the habitat grid.

numPoints

log

n

Number of points in the Poisson point process. This value (non-negative integer) is used to truncate x so that extra rows beyond numPoints are ignored.

Logical argument, specifying whether to return the log-probability of the distri-

bution.

Integer specifying the number of realisations to generate. Only n = 1 is supported.

#### Value

dpoisppAC gives the (log) probability density of the observation matrix x. rpoisppAC gives coordinates of a set of randomly generated spatial points.

### Author(s)

Wei Zhang

#### References

W. Zhang, J. D. Chipperfield, J. B. Illian, P. Dupont, C. Milleret, P. de Valpine and R. Bischof. 2020. A hierarchical point process model for spatial capture-recapture data. bioRxiv. DOI 10.1101/2020.10.06.325035

## **Examples**

dpoisppDetection\_normal

Poisson point process detection model

### **Description**

Density and random generation functions of the Poisson point process for detection. The dpoisppDetection\_normal distribution is a NIMBLE custom distribution which can be used to model and simulate Poisson observations (x) of a single individual in continuous space over a set of detection windows defined by their upper and lower coordinates (*lowerCoords, upperCoords*). The distribution assumes that an individual's detection intensity follows an isotropic bivariate normal function centered on the individual's activity center (s) with standard deviation (sd). All coordinates (s and trapCoords) should be scaled to the habitat (scaleCoordsToHabitatGrid).

```
dpoisppDetection_normal(
    x,
    lowerCoords,
    upperCoords,
    s,
    sd,
    baseIntensities,
```

```
numMaxPoints,
numWindows,
indicator,
log = 0
)

rpoisppDetection_normal(
    n,
    lowerCoords,
    upperCoords,
    s,
    sd,
    baseIntensities,
    numMaxPoints,
    numWindows,
    indicator
)
```

### **Arguments**

x Array containing the total number of detections (x[1,1]), the x- and y-coordinates (x[2:(x[1,1]+1),1:2]), and the corresponding detection window indices (x[2:(x[1,1]+1),3])

for a set of spatial points (detection locations).

lowerCoords, upperCoords

Matrices of lower and upper x- and y-coordinates of all detection windows scaled to the habitat (see (scaleCoordsToHabitatGrid). One row for each

window. Each window should be of size 1x1.

s Vector of x- and y-coordinates of the isotropic multivariate normal distribution

mean (the AC location).

sd Standard deviation of the isotropic multivariate normal distribution.

baseIntensities

Vector of baseline detection intensities for all detection windows.

numMaxPoints Maximum number of points. This value (non-negative integer) is only used

when simulating detections to constrain the maximum number of detections.

numWindows Number of detection windows. This value (positive integer) is used to truncate

lowerCoords and upperCoords so that extra rows beyond numWindows are ig-

nored.

indicator Binary argument specifying whether the individual is available for detection (in-

dicator = 1) or not (indicator = 0).

log Logical argument, specifying whether to return the log-probability of the distri-

bution.

n Integer specifying the number of realisations to generate. Only n = 1 is sup-

ported.

### Value

dpoisppDetection\_normal gives the (log) probability density of the observation matrix x. rpoisppDetection\_normal gives coordinates of a set of randomly generated spatial points.

### Author(s)

Wei Zhang

#### References

W. Zhang, J. D. Chipperfield, J. B. Illian, P. Dupont, C. Milleret, P. de Valpine and R. Bischof. 2020. A hierarchical point process model for spatial capture-recapture data. bioRxiv. DOI 10.1101/2020.10.06.325035

```
coordsHabitatGridCenter <- matrix(c(0.5, 3.5,</pre>
                                      1.5, 3.5,
                                      2.5, 3.5,
                                      3.5, 3.5,
                                      0.5, 2.5,
                                      1.5, 2.5,
                                      2.5, 2.5,
                                      3.5, 2.5,
                                      0.5, 1.5,
                                      1.5, 1.5,
                                      2.5, 1.5,
                                      3.5, 1.5,
                                     0.5, 0.5,
                                     1.5, 0.5,
                                      2.5, 0.5,
                                      3.5, 0.5), ncol = 2, byrow = TRUE)
colnames(coordsHabitatGridCenter) <- c("x","y")</pre>
# Create observation windows
lowerCoords <- matrix(c(0, 0, 1, 0, 0, 1, 1, 1), nrow = 4, byrow = TRUE)
upperCoords <- matrix(c(1, 1, 2, 1, 1, 2, 2, 2), nrow = 4, byrow = TRUE)
colnames(lowerCoords) <- colnames(upperCoords) <- c("x","y")</pre>
# Rescale coordinates
ScaledLowerCoords <- scaleCoordsToHabitatGrid(coordsData = lowerCoords,</pre>
                                        coordsHabitatGridCenter = coordsHabitatGridCenter)
ScaledUpperCoords <- scaleCoordsToHabitatGrid(coordsData = upperCoords,</pre>
                                        coordsHabitatGridCenter = coordsHabitatGridCenter)
ScaledUpperCoords$coordsDataScaled[,2] <- ScaledUpperCoords$coordsDataScaled[,2] + 1.5</pre>
ScaledLowerCoords$coordsDataScaled[,2] <- ScaledLowerCoords$coordsDataScaled[,2] - 1.5
# Detection locations
x \leftarrow matrix(c(1.5, 2, 1.1, 1.5, 0.6, 2.1, 0.5, 2, 1, 1.5), nrow = 5, byrow = TRUE)
# get the window indeces on the third dimension of x
windowIndexes <- 0
for(i in 1:nrow(x)){
  windowIndexes[i] <- getWindowIndex(curCoords = x[i,],</pre>
                                       lowerCoords = ScaledLowerCoords$coordsDataScaled,
                                       upperCoords = ScaledUpperCoords$coordsDataScaled)
```

dpoisppLocalDetection\_normal

Local evaluation for a Poisson point process detection model

## **Description**

Density and random generation functions of the Poisson point process for detection. The dpoisppLocalDetection\_normal distribution is a NIMBLE custom distribution which can be used to model and simulate Poisson observations (x) of a single individual in continuous space over a set of detection windows defined by their upper and lower coordinates (lowerCoords,upperCoords). The distribution assumes that an individual's detection intensity follows an isotropic bivariate normal function centered on the individual's activity center (s) with standard deviation (sd). All coordinates (s and trapCoords) should be scaled to the habitat (scaleCoordsToHabitatGrid).

```
dpoisppLocalDetection_normal(
 Х,
  lowerCoords,
  upperCoords,
  s,
  sd,
  baseIntensities,
  habitatGridLocal,
  resizeFactor = 1,
  localObsWindowIndices,
  numLocalObsWindows,
  numMaxPoints,
  numWindows,
  indicator,
  log = 0
)
```

```
rpoisppLocalDetection_normal(
    n,
    lowerCoords,
    upperCoords,
    s,
    sd,
    baseIntensities,
    habitatGridLocal,
    resizeFactor = 1,
    localObsWindowIndices,
    numLocalObsWindows,
    numMaxPoints,
    numWindows,
    indicator
)
```

#### **Arguments**

Х

sd

Matrix containing the total number of detections (x[1,1]), the x- and y-coordinates (x[2:(x[1,1]+1),1:2]), and the corresponding detection window indices (x[2:(x[1,1]+1),3]) for a set of spatial points (detection locations).

lowerCoords, upperCoords

Matrices of lower and upper x- and y-coordinates of all detection windows scaled to the habitat (see (scaleCoordsToHabitatGrid). One row for each window. Each window should be of size 1x1.

s Vector of x- and y-coordinates of the isotropic multivariate normal distribution mean (the AC location).

Standard deviation of the isotropic multivariate normal distribution.

baseIntensities

Vector of baseline detection intensities for all detection windows.

habitatGridLocal

Matrix of rescaled habitat grid cells indices, from localIndices returned by the getLocalObjects function (

resizeFactor

Aggregation factor used in the getLocalObjects function to reduce the number of habitat grid cells.

localObsWindowIndices

Matrix of indices of local observation windows around each rescaled habitat grid cell, as returned by the getLocalObjects function (object named localIndices).

numLocalObsWindows

Vector of numbers of local observation windows around all habitat grid cells, as returned by the getLocalObjects function (object named numLocalIndices). The ith number gives the number of local (original) observation windows for the ith (rescaled) habitat window.

numMaxPoints

Maximum number of points. This value (non-negative integer) is only used when simulating detections to constrain the maximum number of detections.

numWindows	Number of detection windows. This value (positive integer) is used to truncate lowerCoords and upperCoords so that extra rows beyond numWindows are ignored.
indicator	Binary argument specifying whether the individual is available for detection (indicator = $1$ ) or not (indicator = $0$ ).
log	Logical argument, specifying whether to return the log-probability of the distribution.
n	Integer specifying the number of realisations to generate. Only $n=1$ is supported.

#### Value

The (log) probability density of the observation matrix x.

#### Author(s)

Wei Zhang, Cyril Milleret and Pierre Dupont

#### References

W. Zhang, J. D. Chipperfield, J. B. Illian, P. Dupont, C. Milleret, P. de Valpine and R. Bischof. 2020. A hierarchical point process model for spatial capture-recapture data. bioRxiv. DOI 10.1101/2020.10.06.325035

C. Milleret, P. Dupont, C. Bonenfant, H. Brøseth, Ø. Flagstad, C. Sutherland and R. Bischof. 2019. A local evaluation of the individual state-space to scale up Bayesian spatial capture-recapture. Ecology and Evolution 9:352-363

s <- c(1, 1) sd <- 0.1 baseIntensities <- c(1:4) windowIndex <- 4 numPoints <- 1 numWindows <- 4 indicator <- 1

# Rescale coordinates ScaledLowerCoords <- scaleCoordsToHabitatGrid(coordsData = lowerCoords, coordsHabitatGridCenter = coordsHabitatGridCenter)\$coordsDataScaled ScaledUpperCoords <- scaleCoordsToHabitatGrid(coordsData = upperCoords, coordsHabitatGridCenter = coordsHabitatGridCenter)\$coordsDataScaled ScaledUpperCoords[,2] <- ScaledUpperCoords[,2] + 1.5 ScaledLowerCoords[,2] <- ScaledLowerCoords[,2] - 1.5 habitatMask <- matrix(1, nrow = 4, ncol=4, byrow = TRUE) # Create local objects ObsWindowsLocal <- getLocalObjects(habitatMask = habitatMask, coords = ScaledLowerCoords, dmax=3, resizeFactor = 1, plot.check = TRUE)

# Detection locations  $x \leftarrow matrix(c(1.5, 2, 1.1, 1.5, 1.4, 0.7, 2, 1.3, 1, 1.5), nrow = 5, byrow = TRUE)$ 

# get the window indeces on the third dimension of x windowIndexes <- 0 for(i in 1:nrow(x)) windowIndexes[i] <- getWindowIndex(curCoords = x[i,], lowerCoords = ScaledLowerCoords, upperCoords = ScaledUpperCoords)

getHomeRangeArea 71

x <- cbind(x, windowIndexes) # get the total number of detections on x[1,1] x <- rbind(c(length(windowIndexes),0,0), x) dpoisppLocalDetection\_normal(x, ScaledLowerCoords, ScaledUpperCoords, s, sd, baseIntensities, ObsWindowsLocal\$habitatGrid, ObsWindowsLocal\$resizeFactor, ObsWindowsLocal\$localIndices,ObsWindowsLocal\$numMaxPoints = dim(x)[1], numWindows, indicator, log = TRUE)

getHomeRangeArea

Computation of home range radius and area

### **Description**

getHomeRangeArea returns approximates estimates of home range radius and area for a given set of parameters with respect to a specified detection function using bisection algorithm. The following circular detection functions are available to use in nimbleSCR: half-normal (detFun = 0), half-normal plateau (detFun = 1), exponential (detFun = 2), asymmetric logistic (detFun = 3), bimodal (detFun = 4) and donut (detFun = 5).

### Usage

```
getHomeRangeArea(
    x = 2,
    detFun = 0,
    prob = 0.95,
    d = 6,
    xlim = c(0, 30),
    ylim = c(0, 30),
    nBreaks = 800,
    tol = 0.00001,
    nIter = 2000
)
```

## **Arguments**

x	Vector or matrix (parameters in columns) of values for different parameters corresponding to the specified detection function.
detFun	Numeric vNumeric variable denoting the type of detection function. 0 = Half-normal, 1 = Half-normal plateau, 2 = Exponential, 3 = Asymmetric logistic, 4 = Bimodal, 5 = Donut.
prob	Numeric variable denoting the quantile probability to compute the home range radius.
d	Numeric variable giving an initial value of the radius.
xlim	Vector of length 2 giving the range along x-axis.
ylim	Vector of length 2 giving the range along y-axis.
nBreaks	Numeric variable denoting the number of breaks along an axis.
tol	Numeric variable denoting the allowed tolerance in the radius estimate.
nIter	Numeric variable giving the maximum number of iterations in bisection algorithm.

72 getHomeRangeArea

### Author(s)

Soumen Dey

#### References

Dey, S., Bischof, R., Dupont, P. P. A., & Milleret, C. (2022). Does the punishment fit the crime? Consequences and diagnosis of misspecified detection functions in Bayesian spatial capture–recapture modeling. Ecology and Evolution, 12, e8600. https://doi.org/10.1002/ece3.8600

```
## Not run:
# A user friendly vignette is also available on github:
# https://github.com/nimble-dev/nimbleSCR/blob/master/nimbleSCR/vignettes/
# Vignette name: Fit_with_dbinomLocal_normalPlateau_and_HomeRangeAreaComputation.rmd
# HALF-NORMAL PLATEAU FUNCTION (detFun = 1)
habitatMask <- matrix(1, nrow = 30, ncol= 30, byrow = TRUE)
prob <- 0.95
paramnames.hr <- c("HRradius", "HRarea")</pre>
sigma <- 1
w < -1.5
params <- c(sigma, w)</pre>
names(params) <- c("sigma", "w")</pre>
HRAnim \leftarrow getHomeRangeArea(x = params, detFun = 1, prob = prob, d = 6,
                      xlim = c(0, dim(habitatMask)[2]), ylim = c(0, dim(habitatMask)[1]),
                       nBreaks = 800, tol = 1E-5, nIter = 2000)
# Different values of argument "detFun"
# 0 = Half-normal, 1 = Half-normal plateau, 2 = Exponential,
# 3 = Aysmmetric logistic, 4 = Bimodal, 5 = Donut.
HR.hnp <- c(HRAnim$run())</pre>
names(HR.hnp) <- paramnames.hr</pre>
print(HR.hnp)
# FASTER HRA COMPUTATION USING NIMBLE
samples <- cbind(rgamma(n = 500, shape = 1, rate = 1), rgamma(n = 500, shape = 1.5, rate = 1))
colnames(samples) <- c("sigma", "w")</pre>
HRAnim.mat <- getHomeRangeArea(x = samples, detFun = 1, prob = prob, d = 6,
                       xlim = c(0, dim(habitatMask)[2]), ylim = c(0, dim(habitatMask)[1]),
                          nBreaks = 800, tol = 1E-5, nIter = 2000)
cHRAnim.arr <- compileNimble(HRAnim.mat, resetFunctions = TRUE)</pre>
HRA.Runtime <- system.time(</pre>
  HR.chain <- cHRAnim.arr$run()</pre>
print(HRA.Runtime)
dimnames(HR.chain)[[2]] <- paramnames.hr</pre>
HRest <- do.call(rbind, lapply(c(1:2), function(j){</pre>
```

getHomeRangeArea 73

```
c(mean(HR.chain[,j], na.rm = TRUE), sd(HR.chain[,j], na.rm = TRUE))
dimnames(HRest) <- list(paramnames.hr, c("MEAN", "SD"))</pre>
cat("Numerical estimates using MCMC samples: \n", sep = "")
print(HRest)
# HALF-NORMAL FUNCTION (detFun = 0)
sigma = 2
params <- c(sigma)</pre>
names(params) <- c("sigma")</pre>
HRAnim \leftarrow getHomeRangeArea(x = params, detFun = 0, prob = prob, d = 6,
                      xlim = c(0, dim(habitatMask)[2]), ylim = c(0, dim(habitatMask)[1]),
                       nBreaks = 800, tol = 1E-5, nIter = 2000)
HR.hn <- c(HRAnim$run())</pre>
names(HR.hn) <- paramnames.hr</pre>
print(HR.hn)
# Exponential (detFun = 2)
rate = 1/2
params <- c(rate)</pre>
names(params) <- c("rate")</pre>
HRAnim \leftarrow getHomeRangeArea(x = params, detFun = 2, prob = prob, d = 6,
                      xlim = c(0, dim(habitatMask)[2]), ylim = c(0, dim(habitatMask)[1]),
                       nBreaks = 800, tol = 1E-5, nIter = 2000)
HR.exp <- c(HRAnim$run())</pre>
names(HR.exp) <- paramnames.hr</pre>
print(HR.exp)
# Asymmetric logistic (detFun = 3)
sigma = 2
alpha.a = 5
alpha.b = 1
params <- c(sigma, alpha.a, alpha.b)</pre>
names(params) <- c("sigma", "alpha.a", "alpha.b")</pre>
HRAnim \leftarrow getHomeRangeArea(x = params, detFun = 3, prob = prob, d = 6,
                      xlim = c(0, dim(habitatMask)[2]), ylim = c(0, dim(habitatMask)[1]),
                       nBreaks = 800, tol = 1E-5, nIter = 2000)
HR.al <- c(HRAnim$run())</pre>
names(HR.al) <- paramnames.hr</pre>
print(HR.al)
# Bimodal (detFun = 4)
p0.a = 0.25
sigma.a = 0.5
p0.b = 0.15
sigma.b = 1
```

74 getLocalObjects

```
params <- c(sigma.a, sigma.b, p0.a, p0.b, w)
names(params) <- c("sigma.a", "sigma.b", "p0.a", "p0.b", "w")
HRAnim \leftarrow getHomeRangeArea(x = params, detFun = 4, prob = prob, d = 6,
                      xlim = c(0, dim(habitatMask)[2]), ylim = c(0, dim(habitatMask)[1]),
                      nBreaks = 800, tol = 1E-5, nIter = 2000)
HR.bi <- c(HRAnim$run())</pre>
names(HR.bi) <- paramnames.hr</pre>
print(HR.bi)
# Donut (detFun = 5)
sigma.a = 1.5
sigma.b = 1
w = 1
params <- c(sigma.a, sigma.b, w)</pre>
names(params) <- c("sigma.a", "sigma.b", "w")</pre>
HRAnim \leftarrow getHomeRangeArea(x = params, detFun = 5, prob = prob, d = 6,
                      xlim = c(0, dim(habitatMask)[2]), ylim = c(0, dim(habitatMask)[1]),
                      nBreaks = 800, tol = 1E-5, nIter = 2000)
HR.dn <- c(HRAnim$run())</pre>
names(HR.dn) <- paramnames.hr</pre>
print(HR.dn)
## End(Not run)
```

getLocalObjects

Local Objects Identification

## **Description**

R utility function to identify all objects (e.g. traps) within a given radius dmax of each cell in a habitat mask. Used in the implementation of the local evaluation approach in SCR models (dbinomLocal\_normal;dpoisLocal\_normal). The distance to the activity center and the detection probability are then calculated for local objects only (i.e. the detection probability is assumed to be 0 for all other objects as they are far enough from the activity center).

#### Usage

```
getLocalObjects(habitatMask, coords, dmax, resizeFactor = 1, plot.check = TRUE)
```

# Arguments

habitatMask

a binary matrix object indicating which cells are considered as suitable habitat.

coords

A matrix giving the x- and y-coordinate of each object (i.e. trap). x- and y-coordinates should be scaled to the habitat (scaleCoordsToHabitatGrid).

getLocalObjects 75

dmax	The maximal radius from a habitat cell center within which detection probability is evaluated locally for each trap.
resizeFactor	An aggregation factor to reduce the number of habitat cells to retrieve local objects for. Defaults to 1; no aggregation.
plot.check	A visualization option (if TRUE); displays which objects are considered "local objects" for a randomly chosen habitat cell.

#### **Details**

The getLocalObjects function is used in advance of model building.

## Value

This function returns a list of objects:

- localIndices: a matrix with number of rows equal to the reduced number of habitat grid cells (following aggregation). Each row gives the id numbers of the local objects associated with this grid cell.
- habitatGrid: a matrix of habitat grid cells ID corresponding to the row indices in localIndices.
- numLocalIndices: a vector of the number of local objects for each habitat grid cell in habitat-Grid.
- numLocalIndicesMax: the maximum number of local objects for any habitat grid cell; corresponds to the number of columns in habitatGrid.
- resizeFactor: the aggregation factor used to reduce the number of habitat grid cells.

#### Author(s)

Cyril Milleret and Pierre Dupont

76 getSparseY

getMidPointNodes

Generate midpoint integration nodes

# **Description**

Generate midpoint nodes and weights for integrating a function numerically over a set of windows. For each window, generate a set of equally spaced nodes and weights.

# Usage

```
getMidPointNodes(lowerCoords, upperCoords, numSubintervals = 10)
```

## **Arguments**

lowerCoords, upperCoords

Matrices of lower and upper x- and y-coordinates of a set of windows. One row for each window.

numSubintervals

Number of subintervals each dimension of a window is divided into.

#### Value

A list of midpoint nodes and weights.

#### Author(s)

Wei Zhang

# **Examples**

```
lowerCoords <- matrix(c(0, 0, 1, 0, 0, 1, 1, 1), nrow = 4, byrow = TRUE) upperCoords <- matrix(c(1, 1, 2, 1, 1, 2, 2, 2), nrow = 4, byrow = TRUE) getMidPointNodes(lowerCoords, upperCoords, 5)
```

getSparseY

Sparse Matrix Preparation

# Description

R utility function to turn a two or three-dimensional detection array into a sparse matrix representation (see Turek et al., 2021 <doi.org/10.1002/ecs2.3385> for more details). Used in the implementation of the dbinomLocal\_normal and dpoisLocal\_normal functions.

## Usage

```
getSparseY(x, noDetections = -1, nMaxTraps = NULL)
```

getSparseY 77

#### **Arguments**

Χ

A two- or three-dimensional observation data array with dimensions : number of individuals, number of traps, (and number of detection occasions/sessions).

noDetections

The value indicating no detection. Defaults to -1.

nMaxTraps

The maximum number of traps at which detections can occur. It is necessary to artificially augment the sparse detection array when using the random generation functionality of the rbinomLocal\_normal or rpoisLocal\_normal functions. When simulating detection data, augmenting the size of the detection array is necessary to avoids artificially limiting the number of detectors at which individuals can be detected. Default value is maxDetNums \* 2, which doubles the maximum number of traps at which an individual can be detected. We generally recommend using *numLocalIndicesMax* obtained from getLocalObjects when aiming at randomly generating detections from rbinomLocal\_normal or rpoisLocal\_normal.

#### **Details**

The getSparseY function is used in advance of model building to create a sparse matrix representation of the observation data. It creates and returns a list of objects:

#### Value

A list of objects which constitute a sparse representation of the observation data:

- detNums A matrix with number of traps at which each individual (in rows) was detected at each occasions/sessions (in columns).
- maxDetNums The maximum number of traps at which an individual was detected (i.e., the maximum of detNums).
- *detIndices* An array of dimensions n.individuals, maxDetNums, and number of occasions/sessions, which contains the IDs of the traps where each individual was detected.
- y An array of dimensions n.individuals, maxDetNums, and occasions/sessions, which contains the number of observations of each individual at the traps it was detected at.
- *yCombined* An array that combines *detNums*, *y*, and *detIndices* by columns (in that specific order). Note that *y*, and *detIndices* are augmented before combining, such that the maximum number of detectors at which an individual can be detected is equal to *nMaxTraps* Consequently, the number of columns of *lengthYCombined* is 2\*nMaxTraps + 1.
- *lengthYCombined* Dimension of the augmented lengthYCombined object to be specified as the argument *lengthYCombined* of the dbinomLocal\_normal or dpoisLocal\_normal functions when simulating detection data.

#### Author(s)

Cyril Milleret

78 getWindowCoords

#### **Examples**

```
y.full <- matrix(rbinom(5000, 5, 0.02), ncol = 100)
y <- getSparseY(y.full)</pre>
```

getWindowCoords

Get lower and upper windows coordinates

# **Description**

The getWindowCoords is an R utility function to create lower and upper habitat and observation windows coordinates, as well an habitat grid with cell ids. Those objects are necessary to run all point process (pp) functions. All input data should be scaled to the habitat grid using scaleCoordsToHabitatGrid. Note that we assume homogeneous window sizes.

# Usage

```
getWindowCoords(
   scaledHabGridCenter = scaledHabGridCenter,
   scaledObsGridCenter = NULL,
   plot.check = TRUE
)
```

#### **Arguments**

scaledHabGridCenter

A matrix with the scaled "x" and "y" habitat windows grid cell centers (after using scaleCoordsToHabitatGrid).

scaledObsGridCenter

A matrix with the scaled "x" and "y" observation windows grid cell centers (afer using scaleCoordsToHabitatGrid). This is an optional argument and only necessary when modelling detection as a point process (e.g. dpoisppDetection\_normal).

plot.check

A visualization option (if TRUE); displays habitat and detection windows.

#### Value

A list of objects:

- lowerHabCoords A matrix with the "x" and "y" lower habitat window coordinates.
- upperHabCoords A matrix with the "x" and "y" upper habitat window coordinates.
- habitatGrid A matrix of habitat cell ID that can be used to lookup efficiently the cell ID from a coordinate scaled to the habitat grid: habitatGrid[trunc(scaledHabGridCenter[1,"y"]) + 1, trunc(scaledHabGridCenter[1,"x"]) + 1]. See scaleCoordsToHabitatGrid for more details.
- *lowerObsCoords* A matrix with the "x" and "y" lower observation window coordinates. Only returned when *scaledObsGridCenter* is provided.
- *upperObsCoords* A matrix with the "x" and "y" upper observation window coordinates. Only returned when *scaledObsGridCenter* is provided.

getWindowIndex 79

#### Author(s)

Cyril Milleret

# **Examples**

```
coordsGridCenter <- expand.grid(list(x = seq(50.5, 100, 1),</pre>
                                      y = seq(100.5, 150, 1))
coordsData \leftarrow expand.grid(list(x = seq(60, 90, 1),
                               y = seq(110, 140, 1))
plot(coordsGridCenter[,2] ~ coordsGridCenter[,1])
points(coordsData[,2] ~ coordsData[,1], col="red")
scaled <- scaleCoordsToHabitatGrid(coordsData = coordsData</pre>
                                    , coordsHabitatGridCenter = coordsGridCenter)
\verb|plot(scaled$coordsHabitatGridCenterScaled[,2] \sim \verb|scaled$coordsHabitatGridCenterScaled[,1]||
points(scaled$coordsDataScaled[,2] ~ scaled$coordsDataScaled[,1], col="red")
LowerAndUpperCoords <- getWindowCoords(scaledHabGridCenter = scaled$coordsHabitatGridCenterScaled,
                                          scaledObsGridCenter = scaled$coordsDataScaled)
# Plot habitat window cell centers and lower/upper coordinates
plot(scaled$coordsHabitatGridCenterScaled[,2] ~
     scaled$coordsHabitatGridCenterScaled[,1],
     pch=16, cex=0.3, col=grey(0.5))
points(LowerAndUpperCoords$lowerHabCoords[,2] ~
       LowerAndUpperCoords$lowerHabCoords[,1],
       pch=16, cex=0.3, col=grey(0.1))
points(LowerAndUpperCoords$upperHabCoords[,2] ~
       LowerAndUpperCoords$upperHabCoords[,1],
       pch=16, cex=0.3, col=grey(0.1))
# Plot observation window cells center and lower/upper coordinates
points(scaled$coordsDataScaled[,2]~scaled$coordsDataScaled[,1], pch=16,
cex=0.3, col = adjustcolor("red",alpha.f = 0.8))
points(LowerAndUpperCoords$lowerObsCoords[,2] ~
       LowerAndUpperCoords$lowerObsCoords[,1],
       pch=16, cex=0.3, col = adjustcolor("red", alpha.f = 0.8))
points(LowerAndUpperCoords$upperObsCoords[,2] ~
       LowerAndUpperCoords$upperObsCoords[,1],
       pch=16, cex=0.3, col = adjustcolor("red", alpha.f = 0.8))
```

getWindowIndex

Get window index

#### **Description**

From a set of windows, find the index of the window into which a given point falls. Can be applied to detection and habitat windows.

## Usage

```
getWindowIndex(curCoords, lowerCoords, upperCoords)
```

## **Arguments**

```
curCoords Vector of coordinates of a single spatial point lowerCoords, upperCoords
```

Matrices of lower and upper x- and y-coordinates of a set of windows. One row for each window.

## Value

Index of the window where the given point falls; -1 is returned if the point does not fall in any window.

# Author(s)

Pierre Dupont

# **Examples**

```
sourceCoords \leftarrow c(1.5,2.2) \\ lowerCoords \leftarrow cbind(c(0,1,3,0),c(0,1,2,2)) \\ upperCoords \leftarrow cbind(c(1,3,5,3),c(1,2,4,4)) \\ getWindowIndex(sourceCoords, lowerCoords, upperCoords) \\
```

integrateIntensityLocal\_normal

Integrate the multivariate normal intensity with local evaluation

# Description

Calculate the integral of the intensity function with an isotropic multivariate normal kernel over a set of windows. The local evaluation technique is implemented.

# Usage

```
integrateIntensityLocal_normal(
  lowerCoords,
  upperCoords,
  s,
  baseIntensities,
  sd,
  numLocalWindows,
  localWindows)
```

# **Arguments**

lowerCoords, upperCoords

Matrices of lower and upper x- and y-coordinates of a set of windows. One row

for each window.

s Vector of x- and y-coordinates of the isotropic multivariate normal distribution

mean (AC location).

baseIntensities

Vector of baseline intensities for all windows.

sd Standard deviation of the isotropic multivariate normal distribution.

numLocalWindows

Number of windows that are close to the activity center

localWindows Vector of indices of the windows that are close to the activity center.

#### Value

A vector of integrated intensities over all local windows.

#### Author(s)

Cyril Milleret and Wei Zhang

## References

W. Zhang, J. D. Chipperfield, J. B. Illian, P. Dupont, C. Milleret, P. de Valpine and R. Bischof. 2020. A hierarchical point process model for spatial capture-recapture data. bioRxiv. DOI 10.1101/2020.10.06.325035

```
lowerCoords <- matrix(c(0, 0, 1, 0, 0, 1, 1, 1), nrow = 4, byrow = TRUE) upperCoords <- matrix(c(1, 1, 2, 1, 1, 2, 2, 2), nrow = 4, byrow = TRUE) s <- c(0.1, 0.9) sd <- 0.1 baseIntensities <- c(1:4) numLocalWindows <- 2 localWindows <- c(1, 3) integrateIntensityLocal_normal(lowerCoords, upperCoords, s, baseIntensities, sd, numLocalWindows, localWindows)
```

integrateIntensity\_exp

Integrate the multivariate exponential intensity

## **Description**

Calculate the integral of the intensity function with an isotropic multivariate exponential kernel over a set of windows.

## Usage

```
integrateIntensity_exp(
  lowerCoords,
  upperCoords,
  s,
  baseIntensities,
  lambda,
  numWindows
)
```

## **Arguments**

lowerCoords, upperCoords

Matrices of lower and upper x- and y-coordinates of a set of windows. One row

for each window.

S Vector of x- and y-coordinates of the AC location.

baseIntensities

Vector of baseline intensities for all windows.

lambda Rate parameter of the isotropic multivariate exponential distribution.

numWindows Total number of windows. This value (positive integer) is used to truncate

lowerCoords and upperCoords so that extra rows beyond numWindows are ig-

nored.

# Value

A vector of integrated intensities over all windows.

# Author(s)

Wei Zhang

# References

W. Zhang, J. D. Chipperfield, J. B. Illian, P. Dupont, C. Milleret, P. de Valpine and R. Bischof. 2020. A hierarchical point process model for spatial capture-recapture data. bioRxiv. DOI 10.1101/2020.10.06.325035

# **Examples**

```
lowerCoords <- matrix(c(0, 0, 1, 0, 0, 1, 1, 1), nrow = 4, byrow = TRUE) upperCoords <- matrix(c(1, 1, 2, 1, 1, 2, 2, 2), nrow = 4, byrow = TRUE) s <- c(1, 1) lambda <- 1.0 baseIntensities <- c(1:4) numWindows <- 4 integrateIntensity_exp(lowerCoords, upperCoords, s, baseIntensities, lambda, numWindows)
```

integrateIntensity\_normal

Integrate the multivariate normal intensity

# Description

Calculate the integral of the intensity function with an isotropic multivariate normal kernel over a set of windows.

# Usage

```
integrateIntensity_normal(
  lowerCoords,
  upperCoords,
  s,
  baseIntensities,
  sd,
  numWindows
)
```

## **Arguments**

S

lowerCoords, upperCoords

Matrices of lower and upper x- and y-coordinates of a set of windows. One row for each window

for each window.

Vector of x- and y-coordinates of the isotropic multivariate normal distribution mean (AC location).

baseIntensities

Vector of baseline intensities for all windows.

sd Standard deviation of the isotropic multivariate normal distribution.

numWindows Total number of windows. This value (positive integer) is used to truncate

lowerCoords and upperCoords so that extra rows beyond numWindows are ig-

nored.

## Value

A vector of integrated intensities over all windows.

# Author(s)

Wei Zhang

#### References

W. Zhang, J. D. Chipperfield, J. B. Illian, P. Dupont, C. Milleret, P. de Valpine and R. Bischof. 2020. A hierarchical point process model for spatial capture-recapture data. bioRxiv. DOI 10.1101/2020.10.06.325035

# **Examples**

```
lowerCoords <- matrix(c(0, 0, 1, 0, 0, 1, 1, 1), nrow = 4, byrow = TRUE) upperCoords <- matrix(c(1, 1, 2, 1, 1, 2, 2, 2), nrow = 4, byrow = TRUE) s <- c(1, 1) sd <- 0.1 baseIntensities <- c(1:4) numWindows <- 4 integrateIntensity_normal(lowerCoords, upperCoords, s, baseIntensities, sd, numWindows)
```

localTrapCalculations Local Trap Calculations

# **Description**

These functions are deprecated, and they will be removed from a future release. Utility functions to enable local trap calculations in SCR models. See details section for more information.

## Usage

```
makeGrid(xmin = 0, ymin = 0, xmax, ymax, resolution = 1, buffer = 0)
findLocalTraps(grid, trapCoords, dmax)
getNumLocalTraps(idarg, nLocalTraps, LTD1arg)
getLocalTrapIndices(MAXNUM, localTraps, n, idarg)
calcLocalTrapDists(MAXNUM, n, localTrapInd, s, trapCoords)
calcLocalTrapExposure(R, n, d, localTrapInd, sigma, p0)
```

# Arguments

xmin	Minimal value among all trap location x-coordinates.
ymin	Minimal value among all trap location y-coordinates.
xmax	Maximal value among all trap location x-coordinates.
ymax	Maximal value among all trap location y-coordinates.

85 *localTrapCalculations* 

resolution Desired resolution (in both x and y directions) of discretized grid.

buffer Horizontal and vertical buffer for discretized grid, specifying how much it should

extend (above, below, left, and right) of the maximal trap locations.

grid The grid object returned from the makeGrid function.

trapCoords An nTraps x 2 array giving giving the x- and y-coordinate locations of all traps. dmax

The maximal radius from an activity center for performing trap calculations

(dmax).

A grid id, returned from the makeID function inside model code. idarg

The number of local traps to all grid cells, which is given by the first column of nLocalTraps

the localTraps array.

LTD1arg The number of columns in the localTraps array.

**MAXNUM** The maximum number of local traps among all grid cells. This is given by the

(number of rows)-1 of the localTraps array.

localTraps The array returned from the findLocalTraps function.

The number of local traps to a specified grid cell, as return.

localTrapInd The indices of the local traps to a grid cell, as returned by the getLocalTrapIndices

function.

A length-2 vector giving the activity center of an indiviual.

R The total number of traps.

A vector of distances from an activity center to the local traps.

Scale of decay for detection probability. sigma

p0 Baseline detection probability.

#### **Details**

These functions are deprecated, and they will be removed from a future release.

The makeGrid function is used in advance of model building. It creates and returns a list of two objects: a table (grid) corresponding to the discretized grid, where each row gives the x-coordinate, the y-coordinate, and the id number for a grid cell; and second, a function (makeID) to be used in the model code which operates on a discretized AC location, and returns the id number of the corresponding grid cell.

The findLocalTraps function operates on the grid object returned from makeGrid, and an array of the trap location coordinates, and the desired maximal exposure radius for caluclations (dmax). It returns a array (localTraps) with number of rows equal to the number of grid cells. The first element of each row gives the number of local traps within exposure radius to that grid cell. The following elements of each row give the id numbers of those local traps.

A visualization function (plotTraps) is also provided in the example code, which displaces the discretized grid (small black points), all trap locations (green circles), a specified grid cell location (specified by i) as a large X, and the local traps to that specified grid cell (red circles).

The getNumLocalTraps function is used inside the model code. It operates on an id for a grid cell, the localTraps array (generated by findLocalTraps), and the constant value LTD1. This function returns the number of traps which are local to a specified grid cell.

86 localTrapCalculations

The getLocalTrapIndices function is used inside the model code. It returns a vector containing the ids of the local traps to a particular grid cell.

The calcLocalTrapDists function is used inside the model code. It calculates the distances from an activity center, to the local traps relative to the grid cell nearest that activity center.

The calcLocalTrapExposure function is specific to the detection probability calculations used in this example. This function should be modified specifically to the detection function, exposure function, or otherwise calculations to be done only for the traps in the vicinity of individual activity center locations

#### Author(s)

Daniel Turek

```
## Not run:
## generate random trap locations
nTraps <- 200
traps_xmin <- 0
traps_ymin <- 0
traps_xmax <- 100
traps_ymax <- 200
set.seed(0)
traps_xCoords <- round(runif(nTraps, traps_xmin, traps_xmax))</pre>
traps_yCoords <- round(runif(nTraps, traps_ymin, traps_ymax))</pre>
trap_coords <- cbind(traps_xCoords, traps_yCoords)</pre>
## buffer distance surrounding sides of rectangular discretization grid
## which overlays trap locations
buffer <- 10
## resolution of rectangular discretization grid
resolution <- 10
## creates grid and makeID function,
## for grid overlaying trap locations,
## and to lookup nearest grid cell to any AC
makeGridReturn <- makeGrid(xmin = traps_xmin, xmax = traps_xmax,</pre>
                            ymin = traps_ymin, ymax = traps_ymax,
                            buffer = buffer,
                            resolution = resolution)
grid <- makeGridReturn$grid</pre>
makeID <- makeGridReturn$makeID</pre>
## maximum radis within an individual AC to perform trap calculations,
dmax <- 30
## n = localTraps[i,1] gives the number of local traps
```

localTrapCalculations 87

```
## localTraps[i, 2:(n+1)] gives the indices of the local traps
localTraps <- findLocalTraps(grid, trap_coords, dmax)</pre>
plotTraps <- function(i, grid, trap_coords, localTraps) {</pre>
    plot(grid[,1], grid[,2], pch = '.', cex=2)
    points(trap_coords[,1], trap_coords[,2], pch=20, col='forestgreen', cex=1)
    if(!missing(i)) {
        i <- max(i %% dim(grid)[1], 1)</pre>
        n <- localTraps[i,1]</pre>
        trapInd <- numeric(0)</pre>
        if(n > 0) trapInd <- localTraps[i,2:(n+1)]</pre>
        theseTraps <- trap_coords[trapInd,, drop = FALSE]</pre>
        points(theseTraps[,1], theseTraps[,2], pch = 20, col = 'red', cex=1.5)
        points(grid[i,1], grid[i,2], pch = 'x', col = 'blue', cex=3)
    }
}
## visualise some local traps
plotTraps(10, grid, trap_coords, localTraps)
plotTraps(200, grid, trap_coords, localTraps)
plotTraps(380, grid, trap_coords, localTraps)
## example model code
## using local trap calculations
code <- nimbleCode({</pre>
    sigma \sim dunif(0, 100)
    p0 \sim dunif(0, 1)
    for(i in 1:N) {
        S[i,1] \sim dunif(0, xmax)
        S[i,2] \sim dunif(0, ymax)
        Sdiscrete[i,1] <- round(S[i,1]/res) * res</pre>
        Sdiscrete[i,2] <- round(S[i,2]/res) * res</pre>
        id[i] <- makeID( Sdiscrete[i,1:2] )</pre>
        nLocalTraps[i] <- getNumLocalTraps(id[i], localTraps[1:LTD1,1], LTD1)</pre>
        localTrapIndices[i,1:maxTraps] <-</pre>
          getLocalTrapIndices(maxTraps, localTraps[1:LTD1,1:LTD2], nLocalTraps[i], id[i])
        d[i, 1:maxTraps] <- calcLocalTrapDists(</pre>
            maxTraps, nLocalTraps[i], localTrapIndices[i,1:maxTraps],
            S[i,1:2], trap_coords[1:nTraps,1:2])
        g[i, 1:nTraps] <- calcLocalTrapExposure(</pre>
        nTraps, nLocalTraps[i], d[i,1:maxTraps], localTrapIndices[i,1:maxTraps], sigma, p0)
        y[i, 1:nTraps] ~ dbinom_vector(prob = g[i,1:nTraps], size = trials[1:nTraps])
    }
})
## generate random detection data; completely random
N <- 100
set.seed(0)
y <- array(rbinom(N*nTraps, size=1, prob=0.8), c(N, nTraps))
## generate AC location initial values
Sinit <- cbind(runif(N, traps_xmin, traps_xmax),</pre>
                runif(N, traps_ymin, traps_ymax))
```

```
constants <- list(N = N,
                  nTraps = nTraps,
                   trap_coords = trap_coords,
                   xmax = traps_xmax,
                   ymax = traps_ymax,
                   res = resolution,
                   localTraps = localTraps,
                   LTD1 = dim(localTraps)[1],
                  LTD2 = dim(localTraps)[2],
                  maxTraps = dim(localTraps)[2] - 1)
data <- list(y = y, trials = rep(1,nTraps))</pre>
inits <- list(sigma = 1,</pre>
              p0 = 0.5,
              S = Sinit)
## create NIMBLE model object
Rmodel <- nimbleModel(code, constants, data, inits,</pre>
                       calculate = FALSE, check = FALSE)
## use model object for MCMC, etc.
## End(Not run)
```

marginal Void ProbIntegrand

Integrand of the marginal void probability integral

# **Description**

Integrand of the marginal void probability integral. The domain of this function is the habitat domain.

# Usage

```
marginalVoidProbIntegrand(
    x,
    lowerCoords,
    upperCoords,
    sd,
    baseIntensities,
    numPoints,
    numWindows
)
```

#### **Arguments**

x Matrix of x- and y-coordinates of a set of spatial points. One row corresponds to one point.

lowerCoords, upperCoords

Matrices of lower and upper x- and y-coordinates of a set of detection windows.

One row for each window.

sd Standard deviation of the isotropic multivariate normal distribution.

baseIntensities

Vector of baseline detection intensities for all detection windows.

numPoints Number of points that should be considered. This value (positive integer) is used

to truncate x so that extra rows beyond numPoints are ignored.

numWindows Number of windows. This value (positive integer) is used to truncate lowerCoords

and upperCoords so that extra rows beyond numWindows are ignored.

#### Value

A vector of values of the integrand evaluated at each point of x.

# Author(s)

Wei Zhang

## References

W. Zhang, J. D. Chipperfield, J. B. Illian, P. Dupont, C. Milleret, P. de Valpine and R. Bischof. 2020. A hierarchical point process model for spatial capture-recapture data. bioRxiv. DOI 10.1101/2020.10.06.325035

# **Examples**

marginalVoidProbNumIntegration

Marginal void probability

# Description

Calculate the marginal void probability using the midpoint integration method.

## Usage

```
marginalVoidProbNumIntegration(
  quadNodes,
  quadWeights,
  numNodes,
  lowerCoords,
  upperCoords,
  sd,
  baseIntensities,
  habIntensities,
  sumHabIntensity,
  numObsWindows,
  numHabWindows
)
```

## **Arguments**

quadNodes Three-dimensional array of nodes for midpoint integration. The dimension sizes

are equal to the number of nodes per habitat window (1st), 2 (2nd), and the

number of habitat windows (3rd).

quadWeights Vector of weights for midpoint integration.

numNodes Vector of numbers of nodes for all habitat windows.

lowerCoords, upperCoords

Matrix of lower and upper x- and y-coordinates of all detection windows. One

row for each window.

sd Standard deviation of the isotropic multivariate normal distribution.

baseIntensities

Vector of baseline detection intensities for all detection windows.

habIntensities Vector of habitat intensities for all habitat windows.

sumHabIntensity

Total habitat selection intensity over all windows.

numObsWindows Number of detection windows.

numHabWindows Number of habitat windows.

#### Value

The marginal void probability.

# Author(s)

Wei Zhang

#### References

W. Zhang, J. D. Chipperfield, J. B. Illian, P. Dupont, C. Milleret, P. de Valpine and R. Bischof. 2020. A hierarchical point process model for spatial capture-recapture data. bioRxiv. DOI 10.1101/2020.10.06.325035

## **Examples**

```
lowerHabCoords <- matrix(c(0, 0, 0, 1), nrow = 2, byrow = TRUE)
upperHabCoords \leftarrow matrix(c(2, 1, 2, 2), nrow = 2, byrow = TRUE)
lowerObsCoords <- matrix(c(0, 0, 1, 0, 0, 1, 1, 1), nrow = 4, byrow = TRUE)
upperObsCoords \leftarrow matrix(c(1, 1, 2, 1, 1, 2, 2, 2), nrow = 4, byrow = TRUE)
nodesRes <- getMidPointNodes(lowerHabCoords, upperHabCoords, 10)</pre>
quadNodes <- nodesRes$quadNodes
quadWeights <- nodesRes$quadWeights
numNodes \leftarrow rep(100, 2)
sd <- 0.1
baseDetIntensities <- c(1:4)
habIntensities <- c(1:2)
sumHabIntensity <- sum(habIntensities * c(2, 2))</pre>
numObsWindows <- 4
numHabWindows <- 2
marginalVoidProbNumIntegration(quadNodes, quadWeights, numNodes,
                                lowerObsCoords, upperObsCoords, sd,
                                baseDetIntensities, habIntensities,
                                 sumHabIntensity, numObsWindows, numHabWindows)
```

sampler\_categorical\_general

nimble MCMC sampler function for general categorial distributions

# Description

The categorical\_general sampler operates within nimble's MCMC engine to perform Gibbs sampling for a single node, which must in essence follow a categorical distribution. However, the prior distribution need not be nimble's dcat distribution, but rather can be any (potentially user-defined) distribution which has the same support as a standard categorical (dcat) distribution. Specifically: the distribution must define a discrete random variable, which can only attain values from the set 1, 2, 3, ..., numCategories.

The categorical\_general sampler requires one control list argument, named numCategories, which specifies the fixed upper-bound for the range of the random variable.

The categorical\_general sampler is designed to be used in nimble's MCMC engine, and can be added to an MCMC configuration object using the addSampler method. See help(configureMCMC) for more information about MCMC configuration objects and adding custom samplers.

#### Usage

```
sampler_categorical_general(model, mvSaved, target, control)
```

# **Arguments**

```
model (uncompiled) model on which the MCMC is to be run.

mvSaved modelValues object to be used to store MCMC samples.
```

92 scaleCoordsToHabitatGrid

target node on which the sampler will operate.

control named list containing an elemente named numCategories, which specifies the

upper-bound for the range of the random variable.

#### Author(s)

Daniel Turek

# **Examples**

```
## Not run:
## define custom dmy_categorical distribution as a nimbleFunction
dmy_categorical <- nimbleFunction(...)</pre>
## nimble model code, using custom-written dmy_categorical distribution
code <- nimbleCode({</pre>
 x ~ dmy_categorical(...)
## create NIMBLE model object
Rmodel <- nimbleModel(code)</pre>
## create MCMC configuration object with no samplers
conf <- configureMCMC(Rmodel, nodes = NULL)</pre>
## add categorical_general sampler to MCMC configuration
conf$addSampler(target = 'x', type = 'categorical_general', control = list(numCategories = 10))
## build MCMC algorithm
Rmcmc <- buildMCMC(conf)</pre>
## compile model and MCMC, run MCMC algorithm
## End(Not run)
```

scaleCoordsToHabitatGrid

*Scale x- and y-coordinates to grid cells coordinates.* 

# **Description**

R utility function to scale x- and y- coordinates to the habitat grid. Scaling the coordinates to the habitat grid allows implementation of the fast look-up approach to identify the habitat grid cell in which a point is located. This technique was first applied by Mike Meredith in SCR (https://mmeredith.net/blog/2013/1309\_SECR\_in\_JAGS\_patchy\_habitat.htm). Re-scaling the entire coordinate system of the data input is a requirement to run SCR models with the local evaluation approach. This function requires square grid cells and coordinates using projection with units in meters or km (e.g., UTM but not latitude/longitude)

scaleCoordsToHabitatGrid 93

## Usage

```
scaleCoordsToHabitatGrid(
  coordsData = coordsData,
  coordsHabitatGridCenter = coordsHabitatGridCenter,
  scaleToGrid = TRUE
)
```

## **Arguments**

coordsData A matrix or array of x- and y-coordinates to be scaled to the habitat grid. x- and y- coordinates must be identified using "x" and "y" dimnames.

coordsHabitatGridCenter

A matrix of x- and y-coordinates for each habitat grid cell center.

scaleToGrid Defaults to TRUE. If FALSE, coordsData are already scaled and will be rescaled

to its original coordinates.

#### Value

This function returns a list of objects:

- coordsDataScaled: A matrix or array of scaled (rescaled if scaleToGrid==FALSE) x- and y-coordinates for coordsData.
- coordsHabitatGridCenterScaled: A matrix of scaled x- and y-cell coordinates for coordsHabitatGridCenter.

#### Author(s)

Richard Bischof, Cyril Milleret

```
stratRejectionSampler_exp
```

Stratified rejection sampler for multivariate exponential point process

#### **Description**

Simulate data using a stratified rejection sampler from a point process with an isotropic multivariate exponential decay kernel.

# Usage

```
stratRejectionSampler_exp(
  numPoints,
  lowerCoords,
  upperCoords,
  s,
  windowIntensities,
  lambda
)
```

# **Arguments**

numPoints Number of spatial points to generate.

lowerCoords, upperCoords

Matrices of lower and upper x- and y-coordinates of a set of detection windows.

One row for each window.

s Vector of x- and y-coordinates of of the isotropic multivariate exponential dis-

tribution mean.

windowIntensities

Vector of integrated intensities over all detection windows.

lambda Rate parameter of the isotropic multivariate exponential distribution.

## Value

A matrix of x- and y-coordinates of the generated points. One row corresponds to one point.

#### Author(s)

Wei Zhang

```
numPoints <- 10 lowerObsCoords <- matrix(c(0, 0, 1, 0, 0, 1, 1, 1), nrow = 4, byrow = TRUE) upperObsCoords <- matrix(c(1, 1, 2, 1, 1, 2, 2, 2), nrow = 4, byrow = TRUE) s <- c(1, 1) windowIntensities <- c(1:4)
```

```
\label{lambda} $$$ $$ \ensuremath{\mathsf{lambda}} < - 0.1 $$ $$ stratRejectionSampler_exp(numPoints, lowerObsCoords, upperObsCoords, s, windowIntensities, lambda) $$$ $$
```

```
stratRejectionSampler_normal
```

Stratified rejection sampler for multivariate normal point process

# Description

Simulate data using a stratified rejection sampler from a point process with an isotropic multivariate normal decay kernel.

# Usage

```
stratRejectionSampler_normal(
  numPoints,
  lowerCoords,
  upperCoords,
  s,
  windowIntensities,
  sd
)
```

#### **Arguments**

numPoints Number of spatial points to generate.

lowerCoords, upperCoords

Matrices of lower and upper x- and y-coordinates of a set of detection windows.

One row for each window.

s Vector of x- and y-coordinates of of the isotropic multivariate normal distribu-

tion mean.

windowIntensities

Vector of integrated intensities over all detection windows.

sd Standard deviation of the isotropic multivariate normal distribution.

# Value

A matrix of x- and y-coordinates of the generated points. One row corresponds to one point.

#### Author(s)

Joseph D. Chipperfield and Wei Zhang

```
\label{eq:numPoints} $$ - 10$ lowerObsCoords <- matrix(c(0, 0, 1, 0, 0, 1, 1, 1), nrow = 4, byrow = TRUE)$ upperObsCoords <- matrix(c(1, 1, 2, 1, 1, 2, 2, 2), nrow = 4, byrow = TRUE)$ s <- c(1, 1)$ windowIntensities <- c(1:4)$ sd <- 0.1$ set.seed(0)$ stratRejectionSampler_normal(numPoints, lowerObsCoords, upperObsCoords, s, windowIntensities, sd)$ $$
```

# **Index**

calcLocalTrapDists	getNumLocalTraps
(localTrapCalculations), 84	(localTrapCalculations), 84
calcLocalTrapExposure	getSparseY, 26, 27, 31, 32, 35–37, 54, 55, 60,
(localTrapCalculations), 84	<i>61</i> , 76
calculateDensity, 3	<pre>getWindowCoords, 78</pre>
calcWindowSizes, 4	getWindowIndex, 79
dbernppAC, 5	<pre>integrateIntensity_exp, 82</pre>
dbernppACmovement_exp, 7	<pre>integrateIntensity_normal, 83</pre>
<pre>dbernppACmovement_normal, 9</pre>	integrateIntensityLocal_normal, $80$
dbernppDetection_normal, 11	
dbernppLocalACmovement_exp, 14	localTrapCalculations, 84
dbernppLocalACmovement_normal, 18	
dbernppLocalDetection_normal, 21	makeGrid (localTrapCalculations), 84
dbinom_vector, 39	marginalVoidProbIntegrand, 88
dbinomLocal_exp, 25	marginalVoidProbNumIntegration, 89
dbinomLocal_normal, 30, 74, 76, 77	10 (11 10) 5
dbinomLocal_normalPlateau, 34	rbernppAC (dbernppAC), 5
dcatState1Alive1Dead, 41	rbernppACmovement_exp
dcatState1Alive2Dead, 43	(dbernppACmovement_exp), 7
dcatState2Alive2Dead, 46	rbernppACmovement_normal
dDispersal_exp, 50	(dbernppACmovement_normal), 9
dHabitatMask, 51	rbernppDetection_normal
dmultiLocal_normal, 53	(dbernppDetection_normal), 11
dnormalizer, 58	rbernppLocalACmovement_exp
dpoisLocal_normal, 59, 74, 76, 77	(dbernppLocalACmovement_exp),
dpoisppAC, 63	14
dpoisppDetection_normal, 65, 78	rbernppLocalACmovement_normal
dpoisppLocalDetection_normal, 68	(dbernppLocalACmovement_normal) 18
- /	rbernppLocalDetection_normal
<pre>findLocalTraps (localTrapCalculations),</pre>	(dbernppLocalDetection_normal),
84	21
	rbinom_vector (dbinom_vector), 39
getHomeRangeArea,71	rbinomLocal_exp(dbinomLocal_exp), 25
getLocalObjects, 27, 31, 36, 54, 55, 60, 74,	rbinomLocal_normal, 77
77	rbinomLocal_normal
getLocalTrapIndices	(dbinomLocal_normal), 30
(localTrapCalculations), 84	rbinomLocal_normalPlateau
getMidPointNodes, 76	(dbinomLocal_normalPlateau), 34
getriturotiitinodes, 10	(ubilibilibocai_fibi iliairiateau), 34

98 INDEX

```
rcatState1Alive1Dead
        (dcatState1Alive1Dead), 41
rcatState1Alive2Dead
        (dcatState1Alive2Dead), 43
rcatState2Alive2Dead
        (dcatState2Alive2Dead), 46
rDispersal_exp (dDispersal_exp), 50
rHabitatMask (dHabitatMask), 51
rmultiLocal_normal
        (dmultiLocal_normal), 53
rnormalizer (dnormalizer), 58
rpoisLocal_normal, 77
rpoisLocal_normal (dpoisLocal_normal),
        59
rpoisppAC (dpoisppAC), 63
rpoisppDetection_normal
        ({\tt dpoisppDetection\_normal}),\, 65
rpoisppLocalDetection_normal
        (dpoisppLocalDetection_normal),
        68
sampler_categorical_general, 91
scaleCoordsToHabitatGrid, 6, 8, 10, 12, 15,
         19, 25, 27, 30–32, 36, 37, 53–55,
        59-61, 63-66, 68, 69, 74, 78, 92
stratRejectionSampler_exp, 94
stratRejectionSampler_normal, 95
```