# Package 'secrlinear'

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Type Package

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<b>Description</b> Tools for spatially explicit capture- recapture analysis of animal populations in linear habitats, extending package 'secr'	
License GPL (>=2)	
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# **Description**

An **secr** add-on package for linear habitat models.

# **Details**

Package: secr Type: Package Version: 1.0.3 Date: 2014-12-02

License: GNU General Public License Version 2 or later

The important functions in **secrlinear** are:

read.linearmask
networkdistance
sim.linearpopn
make.line
import and discretize a linear habitat map (route)
compute network distances between detectors and points using a linear habitat mask
simulate population along linear mask
place detectors along (part of) a linear route

#### Other useful functions are:

checkmoves check capthist object for extreme movements showpath interactive examination of network distances clipmask drop mask points outside buffer distance combine two linear masks rbind.linearmask subset.linearmaskselect part of a linear mask asgraph convert linear mask to igraph snapPointsToLinearMask closest point on graph convert coordinates to SpatialLinesDataFrame make.sldf

Documentation is provided in a vignette .../doc/secrlinear-vignette.pdf and in the pdf version of the help pages .../doc/secrlinear-manual.pdf

The package draws on the packages **sp** (Pebesma and Bivand 2005) and **igraph** (Csardi and Nepusz 2006).

In order to fit a linear-habitat model with the **secr** function secr. fit:

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- 1. specify a linear mask for the 'mask' argument
- 2. specify details = list(userdist = networkdistance) so that secr.fit uses network distances rather than Euclidean distances.

An example dataset arvicola is drawn from trapping of water voles *Arvicola amphibius* in June 1984 along the River Glyme in the U.K. – the relevant linear mask is glymemask.

A more complex linear network (Silverstream) is provided as an ESRI shapefile.

#### Author(s)

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#### References

Csardi, G. and Nepusz, T. (2006) The igraph software package for complex network research. *InterJournal, Complex Systems* **1695**. http://igraph.org.

Pebesma, E.J. and Bivand, R. S. (2005) Classes and methods for spatial data in R. *R News* **5**(2), http://cran.r-project.org/doc/Rnews/.

#### See Also

addedges, asgraph, checkmoves, clipmask, deleteedges, linearmask, linearpopn, make.line, make.sldf, networkdistance, plot.linearmask, rbind.linearmask, read.linearmask, showpath, showedges, sim.linearpopn, snapPointsToLinearMask, subset.linearmask

## **Examples**

```
## Water voles in June 1984 on the R. Glyme in Oxfordshire, UK
## capture and trap location files are exactly as for a 2-D analysis
olddir <- setwd(system.file("extdata", package = "secrlinear"))</pre>
on.exit(setwd(olddir))
arvicola <- read.capthist("Jun84capt.txt", "glymetrap.txt",</pre>
             detector = "multi", covname = "sex")
## Import map of linear habitat
## -- from text file of x-y coordinates
glymemask <- read.linearmask(file = "glymemap.txt", spacing = 4)</pre>
## -- from a previously constructed SpatialLinesDataFrame
library(maptools)
SLDF <- readShapeSpatial("glymemap")</pre>
glymemask <- read.linearmask(data = SLDF, spacing = 4)</pre>
## display the mask and capture data
plot (glymemask)
plot(arvicola, add = TRUE, tracks = TRUE)
plot(traps(arvicola), add = TRUE)
## fit model, estimate density
linearfit <- secr.fit(arvicola, mask = glymemask, trace = FALSE,</pre>
    details = list(userdist = networkdistance))
predict(linearfit)
## NOTE : the unit of density (D) is animals / km
```

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Arvicola

Water Vole Capture Dataset

## **Description**

Data from a trapping study of water voles (Arvicola amphibius) along a river in Oxfordshire, U.K.

# Usage

arvicola

#### **Format**

secr capthist object

#### **Details**

Water voles were trapped monthly from May 1984 to May 1985 along 0.9 km of the River Glyme near Woodstock in Oxfordshire, U.K. (Efford 1985). Two sheet-aluminium traps were set at stations 20 m apart along one bank and checked morning and evening for 3 days. Traps were baited with slices of carrot and provisioned with bedding and additional carrot. Voles were marked with individual colour-coded ear tags. The dataset provided is from June 1984. This was early in the breeding season and most voles were overwintered adults; only 3 were young-of-the-year, and these were omitted.

Raw data files "Jun84capt.txt" and "glymetrap.txt" are provided in the 'extdata' folder of the **secrlinear** installation. The vignette ../doc/secrlinear-vignette.pdf shows how to import the raw data.

The data comprise detections of 26 voles at 44 stations on 6 occasions. The two traps at each station were notionally labelled 'A' and 'B', but captures were recorded only by the station at which they occurred: captures were assigned label 'A' or 'B' effectively at random. Each trap could catch a single adult vole, but trap saturation was low (maximum 20.5% of traps caught a vole). No voles died in traps in June 1984.

Water voles in the U.K. restrict their activity to waterways and their immediate banks, except for some populations in more extensive (2-dimensional) wetlands. It is therefore natural to treat their habitat as linear in a spatially explicit capture—recapture model of these data. A suitable linear habitat mask is provided in the accompanying dataset glymemask.

See ../doc/secrlinear-vignette.pdf for more analysis of this dataset.

# Source

M. G. Efford unpublished data.

# References

Efford, M. G. (1985) *The structure and dynamics of water vole populations*. D.Phil thesis, University of Oxford.

## See Also

glymemask

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## **Examples**

```
summary(arvicola)
head(traps(arvicola))
## for speed, pre-compute distance matrix
userd <- networkdistance (traps(arvicola), glymemask, glymemask)</pre>
## fit model
glymefit <- secr.fit(arvicola, mask = glymemask, trace = FALSE,</pre>
                     details = list(userdist = userd))
## estimates of 'real' parameters
predict(glymefit)
## Not run:
tmp <- secr.test(glymefit, nsim = 1000)</pre>
tmp
plot(tmp)
## More voles were caught only once than is predicted by the model.
## This is probably due to within-population variation in movement or
## capture probability.
## End(Not run)
```

checkmoves

Check Movements

# Description

This function reports problems indicated by extreme movements in a capthist object. At present it is configured for distances measured along a linearmask only.

# Usage

```
checkmoves (CH, accept = c(0, 1000), userdist, mask, showall = TRUE, silent = FALSE)
```

## **Arguments**

CH	single-session secr capthist object
accept	range of accepted movements in metres; may include Inf
userdist	matrix of distances between all traps(CH) (optional)
mask	linearmask object (optional)
showall	logical; if TRUE the output dataframe includes all detections of each individual with at least one extreme movement
silent	logical; if TRUE then console output is suppressed

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#### **Details**

This function works with Euclidean distances (the default if neither mask nor userdist is specified), non-Euclidean network distances implied by a linear habitat mask, or any distances presented in userdist.

Linear habitat masks are prone to breaks (missing edges in the graph representation). If an animal is 'recaptured' on both sides of a break then the network distance for that 'movement' will be infinite.

checkmoves displays the message 'All OK' if there are no out-of-range movements, and otherwise displays the number of individuals with out-of-range movements.

#### Value

Invisibly returns a list with components

badmoves logical vector with one element for each animal (i.e. each row in CH)

CH subset of the input CH containing the faulty detection histories

df dataframe detailing the faulty detections

# See Also

linearmask

## **Examples**

```
out <- checkmoves(captdata, accept = c(0,100)) out
```

clipmask

Drop Points Distant From Detectors

# **Description**

clipmask may be used to drop superfluous points from a mask (those further than buffer from any detector).

# Usage

```
clipmask(mask, traps, buffer = 100, clipvert = FALSE)
```

## **Arguments**

traps 'traps' object from **secr**mask 'linearmask' object

buffer network distance for retaining mask points (m)

clipvert logical; if TRUE the underlying lines are also clipped

## **Details**

None yet.

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#### Value

'linearmask' object retaining only points within buffer m of any trap. By default, the 'SLDF' attribute is unchanged.

clipvert = TRUE causes the attribute "SLDF" (a SpatialLinesDataFrame) to be clipped to within spacing/2 of detectors, using functions from **rgeos** (Bivand and Rundel 2014).

## References

Bivand, R. and Rundel, C. (2014) rgeos: Interface to Geometry Engine - Open Source (GEOS). R package version 0.3-4. http://CRAN.R-project.org/package=rgeos

#### See Also

```
make.line
```

# **Examples**

```
x <- seq(0, 4*pi, length = 200)
xy <- data.frame(x = x*100, y = sin(x)*300)
mask <- read.linearmask(data = xy, spacing = 20)

## clipmask
trps <- make.line(mask, n = 15, startbuffer = 1000, by = 30)
newmask <- clipmask(mask, trps, buffer = 200)
plot(newmask)
plot(trps, add = TRUE)

newmask <- clipmask(mask, trps, buffer = 200, clipvert = TRUE)
plot(newmask)
plot(trps, add = TRUE)</pre>
```

edges

Interactive Edge Edit

## **Description**

A 'skip' is defined here as a graph edge between two input lines. Many skips are legitimate, but some 'shortcut' skips are artefacts of the simple algorithm used by asgraph to construct the graph topology. These are better removed. These functions allow skips in a linearmask to be visualised and edited.

addedges may be used to bridge artefactual gaps in a network, or to correct editing mistakes. The edge weight of a new edge is its Euclidean length.

## Usage

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```
addedges(mask, replot = TRUE, ...)
deleteedges(mask, replot = TRUE, ...)
cleanskips(mask)
```

## **Arguments**

mask	linearmask object with 'graph' attribute
lengths	numeric vector of minimum and maximum edge length (weight)
plt	logical; if TRUE the edges are plotted
add	logical; if TRUE the edges will be added to an existing plot
type	character value for subset of edges to show
xlim	vector of min and max x-values for new plot
ylim	vector of min and max y-values for new plot
replot	logical; if TRUE the plot will be re-done using replot
	other arguments (see details)

#### **Details**

showedges is a general-purpose function for displaying or extracting edge information from the graph attribute of a linear mask.

cleanskips removes any between-line 'skips' that are longer than the shortest 'skip'.

Other functions here are used to manually edit a mask graph.

index of start vertex

replot is used to zoom in on part of a linear mask (often to inspect 'skips').

The dots argument  $\dots$  is passed by replot to plot.linearmask and by addedges and deleteedges to showedges (if replot = TRUE).

#### Value

start

y2

showedges returns a dataframe with one row for each edge, and columns –

finish	index of finish vertex
line1	line number of start
line2	line number of finish
x1	geographic coordinates of start
y1	geographic coordinates of start
x2	geographic coordinates of finish

weight "weight" attribute of edge (its length in metres)

geographic coordinates of finish

The dataframe is returned invisibly if plt = TRUE.

cleanskips returns a linearmask object.

replot returns NULL.

addedges and deleteedges invisibly returns a linear mask object with updated graph attribute.

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#### See Also

linearmask, getLineID

# **Examples**

```
## Not run:
setwd(system.file("extdata", package = "secrlinear"))
tempmask <- read.linearmask(file = "silverstream.shp", spacing = 50)

## show all edges
plot(tempmask, linecol = 'white')
tmp <- showedges(tempmask, add = TRUE)

## select a rectangular area to zoom in
replot(tempmask)

## click on the vertices of one or more edges to delete or add
tempmask <- deleteedges(tempmask)

tempmask <- addedges(tempmask)

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

glymemask

Linear Mask for Water Vole Dataset

# Description

Habitat geometry for a trapping study of water voles (*Arvicola aquaticus*) along a river in Oxfordshire, U.K., used in conjunction with the capture dataset arvicola.

# Usage

glymemask

# **Format**

A linearmask object (see read.linearmask)

#### **Details**

Water voles (*Arvicola aquaticus*) were trapped monthly from May 1984 to May 1985 along 0.9 km of the River Glyme in Oxfordshire, U.K. (Efford 1985).

The raw data are provided both as a text file and as a shapefile in the 'extdata' folder of the **secrlinear** installation. The vignette ../doc/secrlinear-vignette.pdf shows how to import the data, which are also provided here as an **secr** object.

## **Source**

M. G. Efford unpublished data.

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#### References

Efford, M. G. (1985) *The structure and dynamics of water vole populations*. D.Phil thesis, University of Oxford.

#### See Also

arvicola

# **Examples**

```
plot(glymemask)
summary(glymemask)

## add a covariate, the distance downstream from the first mask point
downstrm <- networkdistance(glymemask, glymemask[1,], glymemask)[,1]
covariates(glymemask)<- data.frame(downstream = downstrm)</pre>
```

linearmask.object

Description of Linear Mask Objects

## **Description**

An object of class linearmask maps linear habitat, such as a river, in a way that is useful for spatially explicit capture–recapture analysis. The object contains two representations of the linear habitat: as features in a SpatialLinesDataFrame, and discretized as a set of equally spaced points, as in a mask object from **secr**.

Usually the object was created in a call to read.linearmask, which performs the discretization given linear input data and a chosen spacing.

A linearmask object does not explicitly include topological information i.e. the branching pattern of river networks. Instead, a network is constructed on-the-fly in asgraph by joining adjacent mask points (those within a certain small Euclidean distance). This results in 'shortcut' edges where lines approach at an acute angle (see the Examples for a demonstration).

A more complete, but also more cumbersome, solution for river networks might be to adapt the SpatialStreamNetwork object class of the SSN package (ver Hoef et al. 2014). Data input for SSN requires pre-processing in the proprietary software ArcGIS.

# Value

The object itself is a dataframe of coordinates for the points comprising the mask, as for a 2-dimensional habitat mask (mask).

Additional information is stored as attributes (these usually remain hidden, but are essential except for 'graph'):

spacingfactor

The spacing factor determines which points in a linear mask are joined to form a linear network, as happens automatically when computing distances with networkdistance. Points are joined when their Euclidean separation is less than spacingfactor x spacing. The default value is 1.1.

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underlying description of the linear habitat.

graph igraph network. Optionally computed by read.linearmask and saved as at-

tribute, or computed on the fly by networkdistance

.

covariates A required dataframe of covariate values, including at least the column 'LineID'

Other attributes are common to masks and linearmasks:

spacing In this context, the length of habitat in metres that is associated with each mask

point (segment length).

meanSD A matrix with mean and standard deviation of coordinates.

boundingbox In this context, a dataframe giving the coordinate ranges spanned by the SLDF.

Differs from 2-D mask in not being 'buffered' by spacing/2.

Further attributes of 2-dimensional masks (area, polygon etc.) are not relevant but may be present. The object has class c("linearmask", "mask", "data.frame") or c("list", "linearmask", "mask", "mas

#### References

Ver Hoef, J. M., Peterson, E. E., Clifford, D. and Shah, R. (2014) **SSN**: an R package for spatial statistical modeling on stream networks. *Journal of Statistical Software* **56(3)**, 1–45.

## See Also

read.linearmask, mask, showpath, networkdistance

## **Examples**

make.line

linearpopn.object

Description of Linear Population Objects

## **Description**

An object of class linearpopn, usually created in a call to sim.linearpopn and used as input to sim.capthist.

#### Value

The object itself is a dataframe of coordinates for points along the mask, as for a 2-dimensional population (popn).

The attribute model2D is always "linear".

The object has class c("linearpopn", "popn", "data.frame").

## See Also

```
plot.linearpopn, popn.
```

# **Examples**

make.line

Linear Detector Array

# **Description**

Build **secr** 'traps' object, positioning detectors along a predefined set of lines. make.line is in the same family as **secr** functions make.grid etc.

# Usage

```
make.line(SLDF, n = 10, startbuffer = 0, by = 20, endbuffer = 0, cluster
= NULL,
type = c("fixedstart", "randomstart", "centred"), detector = "multi")
```

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# **Arguments**

SLDF	linearmask object or SpatialLinesDataFrame from sp
n	maximum number of detectors (or clusters of detectors) per line
startbuffer	distance of first detector from start of line(s)
endbuffer	minimum distance of last detector from end of line(s)
by	spacing between detectors (or clusters of detectors) (m)
cluster	numeric vector of within-cluster positions (m from start)
type	character
detector	character value for detector type - "single", "multi" etc.

## **Details**

Detectors are placed independently on each line.

## Value

An object of class traps comprising a data frame of x- and y-coordinates, the detector type ("single", "multi", or "proximity" etc.), and possibly other attributes. The SpatialLinesDataFrame is retained as attribute 'SLDF'.

# See Also

```
make.traps
```

# **Examples**

```
x <- seq(0, 4*pi, length = 200)
xy <- data.frame(x = x*100, y = sin(x)*300)
mask <- read.linearmask(data = xy, spacing = 20)

trps <- make.line(mask, n = 15, startbuffer = 1000, by = 50)
plot(mask)
plot(trps, add = TRUE)</pre>
```

networkdistance

Network Distances

# **Description**

Simple network computation of distance between points along linear landscape features (e.g. rivers).

# Usage

```
networkdistance(xy1, xy2, geometry)
```

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## **Arguments**

xy1	2-column matrix or dataframe
xy2	2-column matrix or dataframe
geometry	'linearmask' object

#### **Details**

networkdistance computes the distance in metres between the points in xy1 and the points in xy2. The geometry is a linearmask; if it is missing then it takes the value of xy2, which then of course must be a full linearmask. A network is taken from the 'graph' attribute of geometry or (if necessary) constructed from the linearmask 'on the fly', joining any points closer than (spacingfactor x spacing(geometry)). Points in xy1 and xy2 are first snapped to the nearest point in geometry. The computed distance is the sum of the graph's edge weights along the shortest path joining two points.

The commonest use of networkdistance is to calculate distances between detectors (traps) and points on a linear mask. In this case it is sufficient to provide two arguments only, as xy2 serves as both destination and geometry.

networkdistance meets the requirements for a user-defined distance function (userdist) in **secr**. No parameter is estimated.

Use showpath to check network distances interactively.

#### Value

```
networkdistance — given k detectors and m mask points, a k x m matrix of distances. If called with no arguments networkdistance returns a zero-length character vector.
```

# See Also

```
read.linearmask, showpath, asgraph
```

# **Examples**

```
x <- seq(0, 4*pi, length = 200)
xy <- data.frame(x = x*100, y = sin(x)*300)
mask <- read.linearmask(data = xy, spacing = 20)
trps <- make.line(mask, n = 15, startbuffer = 1000, by = 30)
## networkdistance; geometry in 'mask'
d <- networkdistance (trps, mask, mask)
dim(d)
head(d)</pre>
```

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plotmethods Plot Methods for linearmask and linearpopn objects
--

# **Description**

Custom plotting.

## Usage

# Arguments

X	linearpopn object from sim.linearpopn, or a linearmask object from read.linearmask
	For plot.linearpopn: other arguments passed to plot.popn(may include add = TRUE). For plot.linearmask: other arguments passed to plot.mask (e.g., col, cex, add), to legend (pt.cex) or to the plot method for SpatialLines (lwd, lty)
jitter	numeric value for jittering
plotline	logical; if TRUE the mask line is overplotted in grey
linecol	line colour for linear habitat (see Color Specification in par)
label	logical; if TRUE each vertex is numbered
laboffset	offset of label from point (metres)

# **Details**

The linear mask used for plotting a 'linearpopn' is the one saved as an attribute of the object.

The main plotting in plot.linearmask is done by plot.mask with dots = TRUE. See the help for plot.mask for details of options such as add. The lines of the SpatialLinesdataFrame are overplotted unless linecol = NA.

Jittering shifts points by a random uniform distance –  $(\pm 0.5 \times \text{ jitter})$  x mask spacing – on both axes. This can give a better impression of density when points coincide.

The option inplot.popn for plotting rectangular 'frame' is suppressed: do not attempt to pass this argument in . . . .

# Value

```
plot.linearpopn does not return a value.
plot.linearmask invisibly returns legend details as for plot.mask.
```

# See Also

```
sim.linearpopn, plot.mask, plot.popn
```

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#### **Examples**

```
x <- seq(0, 4*pi, length = 200)
xy <- data.frame(x = x*100, y = sin(x)*300)
mask <- read.linearmask(data = xy, spacing = 10)
linpop <- sim.linearpopn(mask, 100)
plot(linpop, jitter = 2)

plot(mask)

## thicker band of grey points, dashed line
plot(mask, cex = 2, lty = 2)

## point colour determined by a covariate
plot(glymemask, cex = 2, covariate = 'x', pt.cex = 2)

## point size determined by a covariate
plot(glymemask, cex = covariates(glymemask)$x/50, pch = 21)</pre>
```

rbind.linearmask

Combine linearmask Objects

## **Description**

Form a new linearmask object by combining the underlying vertices and drawing a new systematic sample.

## Usage

```
## S3 method for class 'linearmask'
rbind(..., cleanskips = TRUE)
```

## **Arguments**

```
... one or more linearmask objects with the same spacing cleanskips logical; passed to internal function make.linearmask
```

#### **Details**

The maptools function spRbind is used to combine the 'SLDF' attributes of the inputs.

If the input objects have a 'graph' attribute then a new graph will be included in the output.

## Value

A linearmask object

## See Also

```
spRbind, read.linearmask, linearmask, subset.linearmask
```

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## **Examples**

```
x \leftarrow seq(0, 4*pi, length = 200)

xy \leftarrow data.frame(x = x*100, y = sin(x)*300)

xy2 \leftarrow data.frame(x = x*100, y = cos(x)*300)

test \leftarrow read.linearmask(data = xy, spacing = 20)

test2 \leftarrow read.linearmask(data = xy2, spacing = 20)

test2 \leftarrow read.linearmask(data = xy2, spacing = 20)
```

read.linearmask

Import Linear Habitat Mask

# **Description**

Construct a linearmask from line data in a text file, a polyline shapefile, a dataframe or a SpatialLinesDataFrame object.

## Usage

```
read.linearmask(file = NULL, data = NULL, spacing = 10, spacingfactor =
1.5, graph = TRUE, cleanskips = TRUE, ...)
```

# **Arguments**

file	file name
data	data frame or SpatialLinesDataFrame
spacing	length of each discretized segment
spacingfactor	numeric criterion for joining mask points
graph	logical; if TRUE the <b>igraph</b> representation of the mask is saved as an attribute
cleanskips	logical; if TRUE then cleanskips is applied to the graph
	other arguments passed to read.table for file input

# **Details**

Only one of 'file' or 'data' should be specified. The file name should include the extension (e.g., "silverstream.shp").

Input from a text file, polyline shapefile or dataframe is first converted to a SpatialLinesDataFrame object. Points are spaced equally along each SpatialLine, starting at spacing / 2.

For dataframe input via data the coordinates are expected to be in columns 'x' and 'y'; if these names are missing, or input is from a file, the first two columns are used.

spacingfactor is used by asgraph to determine which points are joined when computing distances. It should be between 1 and 2.

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#### Value

A 'linearmask' object, i.e. a 'mask' object in which each point represents a line segment of length spacing, and with additional attributes 'SLDF', 'spacingfactor' and possibly 'graph' (see linearmask).

#### Note

A valid shapefile includes at least three files with extensions '.shp', '.dbf' and '.shx'. The full name of the .shp file is provided as the 'file' argument of read.linearmask. The shapefile name without any extension is used as the first argument to the **maptools** function readShapeSpatial.

Line-level covariates are copied from SpatialLinesDataFrame input. Point-specific covariates may be added later using covariates or addCovariates.

#### See Also

```
asgraph, linearmask, make.sldf, networkdistance, SpatialLinesDataFrame spsample
```

# **Examples**

```
x <- seq(0, 4*pi, length = 200)
xy <- data.frame(x = x*100, y = sin(x)*300)
test <- read.linearmask(data = xy, spacing = 20)
plot(test)

xy2 <- data.frame(x = x*100, y = cos(x)*300)
test2 <- read.linearmask(data = xy2, spacing = 20)
plot(test2, add = TRUE)</pre>
```

showpath

Display Shortest Path

# Description

An interactive function to examine shortest paths and the associated network distances within a linear mask.

# Usage

```
showpath(mask, add = FALSE, ...)
```

# Arguments

mask	linearmask object
add	logical; if TRUE the plot is added to an existing plot
	other arguments passed to lines for drawing network path

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#### **Details**

The selected points are snapped to the nearest mask point before any calculations.

The path shown is a shortest path as determined by function get.shortest.paths of the **igraph** package. There may be others.

#### Value

Invisibly returns a list comprising

paths a list of dataframes, each with the mask coordinates along a selected path results a dataframe with one row for each path giving the distances for each pair of points (columns 'from', 'to', 'Euclidean.d', 'network.d')

#### References

Csardi, G. and Nepusz, T. (2006) The igraph software package for complex network research, InterJournal, Complex Systems 1695. http://igraph.org.

#### See Also

networkdistance

## **Examples**

```
## Not run:

x <- seq(0, 4*pi, length = 200)
xy <- data.frame(x = x*100, y = sin(x)*300)
xy2 <- data.frame(x = x*100, y = cos(x)*300)
test <- read.linearmask(data = xy, spacing = 10)
test2 <- read.linearmask(data = xy2, spacing = 10)
test3 <- rbind(test, test2)
showpath(test3, lwd = 5)

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

Silverstream

Silverstream River Centrelines

# **Description**

Silverstream (45°48' S, 170°26' E) is a small river in a forested catchment near Dunedin, New Zealand, about 12 km from top to bottom. There are numerous side streams making this a good example of a dendritic network.

#### **Format**

An ESRI polyline shapefile (silverstream.shp, silverstream.shx, silverstream.dbf).

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#### **Source**

Land Information New Zealand (LINZ) 1:50 000 topographic database http://data.linz.govt.nz/layer/327-nz-mainland-river-centrelines-topo-150k/licensed by LINZ for re-use under the Creative Commons Attribution 3.0 New Zealand licence. Downloaded 2014-10-26.

## References

ESRI (1998) ESRI shapefile technical description. http://www.esri.com/library/whitepapers/pdfs/shapefile.pdf.

# **Examples**

```
## Not run:
setwd(system.file("extdata", package = "secrlinear"))
silverstreammask <- read.linearmask(file = "silverstream.shp", spacing = 50)
par(mar = c(1,1,1,1))
plot(silverstreammask)
## End(Not run)</pre>
```

sim.linearpopn

Simulate Animals on Lines

# Description

This function is a simple substitute for the **secr** function sim.popn() for the case of a linear habitat.

# Usage

```
sim.linearpopn(mask, D, N, Ndist = c('poisson', 'fixed'), ...)
```

# Arguments

mask	linearmask object
D	numeric density animals / km
N	number of individuals
Ndist	character string for distribution of total number of individuals
	other arguments passed to sim.popn

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#### **Details**

The linearmask input represents a discretized line - essentially a chain of line segments. By default, each segment is populated with a Poisson number of individuals. The user may specify D or N.

D may be a vector with one density per mask pixel, or a single number that will be applied across all pixels.

If Ndist = 'fixed' then a constant number of individuals N are simulated in each trial; otherwise N has a Poisson distribution across trials.  $N = sum(D) \times mask \ length \ if D \ is specified.$ 

This is a simplified wrapper for sim.popn called with model2D = "linear".

## Value

```
Object of class c('linearpopn', 'popn', 'data.frame').
```

#### Note

The population output from sim.linearpopn may be used unchanged with **secr** functions such as sim.capthist. However, to be faithful to the linear network you should set the 'userdist' argument of sim.capthist to networkdistance.

## See Also

```
linearpopn, sim.popn
```

# **Examples**

```
x <- seq(0, 4*pi, length = 200)
xy <- data.frame(x = x*100, y = sin(x)*300)
mask <- read.linearmask(data = xy, spacing = 10)
trps <- make.line(mask, n = 15, startbuffer = 1000, by = 30)

newmask <- clipmask(mask, trps, buffer = 200)

linpop <- sim.linearpopn(newmask, 200)
CH <- sim.capthist(trps, linpop, userdist = networkdistance)
plot(newmask)
plot(CH, add = TRUE)

secr.fit(CH, mask = mask, details = list(userdist = networkdistance))</pre>
```

subset.linearmask

Select Part of linearmask Object

## **Description**

Methods to subset linear mask objects.

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#### Usage

```
## S3 method for class 'linearmask'
subset(x, subset, LineID, droplinesbeyond = Inf, ...)
```

# **Arguments**

```
x linearmask object from read.linearmask
subset numeric or logical vector to select rows of mask
LineID vector of identifiers for lines to retain
droplinesbeyond logical; distance criterion for dropping lines (m)
... other arguments (not used)
```

## **Details**

Specify only one of 'subset' and 'LineID'.

If the input object has a 'graph' attribute then a new graph will be included in the output.

subset.linearmask leaves the underlying SpatialLinesDataFrame intact unless either (i) LineID is specified, or (ii) a finite value is provided for droplinesbeyond. droplinesbeyond retains lines that have at least one point within the specified distance of at least one mask point, after subsetting.

#### Value

A linearmask object.

#### See Also

```
read.linearmask,linearmask
```

# **Examples**

```
## rbind two linear masks,
x <- seq(0, 4*pi, length = 200)
xy <- data.frame(x = x*100, y = sin(x)*300)
xy2 <- data.frame(x = x*100, y = cos(x)*300)
test <- read.linearmask(data = xy, spacing = 10)
test2 <- read.linearmask(data = xy2, spacing = 10)
test3 <- rbind(test, test2)
table(covariates(test3)$LineID)

## then retrieve one...
test4 <- subset(test3, LineID = '1.1')

## ... or the other
test5 <- subset(test3, LineID = '2.1')</pre>
```

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# **Description**

Utilities for various functions

- conversion of a flat table of coordinates for a line or set of lines to a SpatialLinesDataFrame object
- snapping points from objects of various types (traps, popn, matrix etc.) to a linearmask object.
- converting distances along a linear mask to x-y coordinates
- build igraph network from linearmask object

# Usage

```
make.sldf(coord, f)
snapPointsToLinearMask(xy, mask)
asgraph(mask)
getLineID(mask, laboffset = rep(spacing(mask) * 3, 2), ...)
```

## **Arguments**

coord	dataframe of coordinates - must include columns 'x','y'
f	vector of values by which to split coord rows
ху	any object that may be coerced to a 2-column matrix (columns interpreted as $x$ -and $y$ -coordinates)
mask	'linearmask' object
laboffset	offset of label from point
	other arguments passed to plot.linearmask

## **Details**

In make.sldf, if f is missing and there is a column 'lineID' in 'coord' then this will be used instead.

 $snapPointsToLinearMask\ shifts\ each\ point\ in\ the\ input\ to\ the\ nearest\ point\ on\ the\ graph\ stored\ as\ the\ 'graph'\ attribute\ of\ mask.$ 

make.line creates a 'traps' object with equally spaced or clustered detectors.

asgraph is a utility function used internally for expressing the points in mask as an igraph network (Csardi and Nepusz 2006).

getLineID plots the mask and for each mouse click displays the corresponding point and LineID.

Other utilities specific to editing graphs are documented in edges.

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#### Value

For make.sldf, a SpatialLinesDataFrame.

For snapPointsToLinearMask, the output has the same attributes (including class) as the xy input: only the coordinates themselves may change.

For alongmask, a 2-column dataframe with the x and y coordinates of the points.

For asgraph, an object of class 'igraph'

getLineID invisibly returns a dataframe with columns 'Point' and 'LineID'.

#### References

Csardi, G. and Nepusz, T. (2006) The igraph software package for complex network research, InterJournal, Complex Systems 1695. http://igraph.org.

## See Also

checkmoves make.line, networkdistance, SpatialLinesDataFrame, showedges, deleteedges, addedges

## **Examples**

```
## test make.sldf()
coord <- data.frame(x = c(1,2,3,1.05,2.05,3.05),
    y = c(3,2,2,3.05,2.05,2.05)
sldf1 <- make.sldf(coord)</pre>
sldf2 \leftarrow make.sldf(coord, f = c(1,1,1,2,2,2))
require(sp)
plot(sldf1)
plot(sldf2)
## test snapPointsToLinearMask()
## Not run:
x <- seq(0, 4*pi, length = 200)
xy < - data.frame(x = x*100, y = sin(x)*300)
mask <- read.linearmask(data = xy, spacing = 20)</pre>
plot(mask)
## click several points, right-click and select 'stop'
pts <- locator()</pre>
pts <- do.call(cbind, pts)</pre>
points(pts)
pts1 <- snapPointsToLinearMask(pts, mask)</pre>
segments(pts[,1], pts[,2], pts1[,1], pts1[,2])
points(pts1, pch = 16, col = 'red')
df <- data.frame(pts, pts1)</pre>
names(df) \leftarrow c('from.x', 'from.y', 'to.x', 'to.y')
## End(Not run)
```

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```
## asgraph (called by networkdistance)
## see 'igraph' for further manipulation
grmask <- asgraph(glymemask)
require(igraph)
summary(grmask)</pre>
```

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