

# Package ‘SpaDES.tools’

June 23, 2020

**Type** Package

**Title** Additional Tools for Developing Spatially Explicit Discrete Event Simulation (SpaDES) Models

**Description** Provides GIS and map utilities, plus additional modeling tools for developing cellular automata, dynamic raster models, and agent based models in 'SpaDES'.  
Included are various methods for spatial spreading, spatial agents, GIS operations, random map generation, and others.  
See 'SpaDES.tools' for an categorized overview of these additional tools.

**URL** <https://spades-tools.predictiveecology.org>,  
<https://github.com/PredictiveEcology/SpaDES.tools>

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 'distanceFromEachPoint.R' 'environment.R' 'initialize.R'  
 'mapReduce.R' 'mergeRaster.R' 'movement.R' 'neighbourhood.R'  
 'numerical-comparisons.R' 'probability.R' 'resample.R'  
 'rings.R' 'spades-tools-deprecated.R' 'spades-tools-package.R'  
 'splitRaster.R' 'spread.R' 'spread2.R' 'spread3.R'  
 'studyArea.R' 'zzz.R'

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SpaDES.tools-package    *Categorized overview of the SpaDES.tools package*

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## Description



### 1 Spatial spreading/distances methods

Spatial contagion is a key phenomenon for spatially explicit simulation models. Contagion can be modelled using discrete approaches or continuous approaches. Several functions assist with these:

<a href="#">adj</a>	An optimized (i.e., faster) version of <a href="#">adjacent</a>
<a href="#">cir</a>	Identify pixels in a circle around a <a href="#">SpatialPoints*</a> object
<a href="#">directionFromEachPoint</a>	Fast calculation of direction and distance surfaces
<a href="#">distanceFromEachPoint</a>	Fast calculation of distance surfaces
<a href="#">rings</a>	Identify rings around focal cells (e.g., buffers and donuts)
<a href="#">spokes</a>	TO DO: need description
<a href="#">spread</a>	Contagious cellular automata
<a href="#">wrap</a>	Create a torus from a grid

### 2 Spatial agent methods

Agents have several methods and functions specific to them:

<a href="#">crw</a>	Simple correlated random walk function
<a href="#">heading</a>	Determines the heading between <a href="#">SpatialPoints*</a>
<a href="#">makeLines</a>	Makes <a href="#">SpatialLines</a> object for, e.g., drawing arrows
<a href="#">move</a>	A meta function that can currently only take "crw"
<a href="#">specificNumPerPatch</a>	Initiate a specific number of agents per patch

### 3 GIS operations

In addition to the vast amount of GIS operations available in R (mostly from contributed packages such as `sp`, `raster`, `maps`, `maptools` and many others), we provide the following GIS-related functions:

`equalExtent` Assess whether a list of extents are all equal

### 4 Map-reduce - type operations

These functions convert between reduced and mapped representations of the same data. This allows compact representation of, e.g., rasters that have many individual pixels that share identical information.

`rasterizeReduced` Convert reduced representation to full raster

### 5 Random Map Generation

It is often useful to build dummy maps with which to build simulation models before all data are available. These dummy maps can later be replaced with actual data maps.

`gaussMap` Creates a random map using Gaussian random fields  
`randomPolygons` Creates a random polygon with specified number of classes

### 6 SELES-type approach to simulation

These functions are essentially skeletons and are not fully implemented. They are intended to make translations from **SELES**. You must know how to use SELES for these to be useful:

`agentLocation` Agent location  
`initiateAgents` Initiate agents into a `SpatialPointsDataFrame`  
`numAgents` Number of agents  
`probInit` Probability of initiating an agent or event  
`transitions` Transition probability

### 7 Package options

SpaDES packages use the following `options` to configure behaviour:

- `spades.lowMemory`: If true, some functions will use more memory efficient (but slower) algorithms. Default FALSE.

### Author(s)

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### See Also

Useful links:

- <https://spades-tools.predictiveecology.org>
- <https://github.com/PredictiveEcology/SpaDES.tools>
- Report bugs at <https://github.com/PredictiveEcology/SpaDES.tools/issues>

---

adj

*Fast 'adjacent' function, and Just In Time compiled version*

---

### Description

Faster function for determining the cells of the 4, 8 or bishop neighbours of the cells. This is a hybrid function that uses matrix for small numbers of loci (<1e4) and data.table for larger numbers of loci

### Usage

```
adj(  
  x = NULL,  
  cells,  
  directions = 8,  
  sort = FALSE,  
  pairs = TRUE,  
  include = FALSE,  
  target = NULL,  
  numCol = NULL,  
  numCell = NULL,  
  match.adjacent = FALSE,  
  cutoff.for.data.table = 2000,  
  torus = FALSE,  
  id = NULL,  
  numNeighs = NULL,  
  returnDT = FALSE  
)
```

## Arguments

<code>x</code>	Raster* object for which adjacency will be calculated.
<code>cells</code>	vector of cell numbers for which adjacent cells should be found. Cell numbers start with 1 in the upper-left corner and increase from left to right and from top to bottom.
<code>directions</code>	the number of directions in which cells should be connected: 4 (rook's case), 8 (queen's case), or "bishop" to connect cells with one-cell diagonal moves. Or a neighbourhood matrix (see Details).
<code>sort</code>	logical. Whether the outputs should be sorted or not, using cell ids of the from cells (and to cells, if <code>match.adjacent</code> is TRUE).
<code>pairs</code>	logical. If TRUE, a matrix of pairs of adjacent cells is returned. If FALSE, a vector of cells adjacent to cells is returned
<code>include</code>	logical. Should the focal cells be included in the result?
<code>target</code>	a vector of cells that can be spread to. This is the inverse of a mask.
<code>numCol</code>	numeric indicating number of columns in the raster. Using this with <code>numCell</code> is a bit faster execution time.
<code>numCell</code>	numeric indicating number of cells in the raster. Using this with <code>numCol</code> is a bit faster execution time.
<code>match.adjacent</code>	logical. Should the returned object be the same as <code>raster::adjacent</code> . Default FALSE, which is faster.
<code>cutoff.for.data.table</code>	numeric. If the number of cells is above this value, the function uses <code>data.table</code> which is faster with large numbers of cells. Default is 5000, which appears to be the turning point where <code>data.table</code> becomes faster.
<code>torus</code>	Logical. Should the spread event wrap around to the other side of the raster? Default is FALSE.
<code>id</code>	numeric If not NULL (default), then function will return "id" column.
<code>numNeighs</code>	A numeric scalar, indicating how many neighbours to return. Must be less than or equal to <code>directions</code> ; which neighbours are random with equal probabilities.
<code>returnDT</code>	A logical. If TRUE, then the function will return the result as a <code>data.table</code> , if the internals used <code>data.table</code> , i.e., if number of cells is greater than <code>cutoff.for.data.table</code> . User should be warned that this will therefore cause the output format to change depending <code>cutoff.for.data.table</code> . This will be faster for situations where <code>cutoff.for.data.table = TRUE</code> .

## Details

Between 4x (large number loci) to 200x (small number loci) speed gains over `adjacent` in raster package. There is some extra speed gain if `NumCol` and `NumCells` are passed rather than a raster. Efficiency gains come from: 1. use `data.table` internally - no need to remove NAs because wrapped or outside points are just removed directly with `data.table` - use `data.table` to sort and fast select (though not fastest possible) 2. don't make intermediate objects; just put calculation into return statement

The steps used in the algorithm are: 1. Calculate indices of neighbouring cells 2. Remove "to" cells that are  $<1$  or  $>\text{numCells}$  (i.e., they are above or below raster), using a single modulo calculation - where the modulo of "to" cells is equal to 1 if "from" cells are 0 (wrapped right to left) - or where the modulo of the "to" cells is equal to 0 if "from" cells are 1 (wrapped left to right)

### Value

Either a matrix (if more than 1 column, i.e., `pairs = TRUE`, and/or `id` is provided), a vector (if only one column), or a `data.table` (if `cutoff.for.data.table` is less than `length(cells)` and `returnDT` is `TRUE`). To get a consistent output, say a matrix, it would be wise to test the output for its class. The variable output is done to minimize coercion to maintain speed. The columns will be one or more of `id`, `from`, `to`.

### Author(s)

Eliot McIntire

### See Also

[adjacent](#)

### Examples

```
library(raster)
a <- raster(extent(0, 1000, 0, 1000), res = 1)
sam <- sample(1:length(a), 1e4)
numCol <- ncol(a)
numCell <- ncell(a)
adj.new <- adj(numCol = numCol, numCell = numCell, cells = sam, directions = 8)
adj.new <- adj(numCol = numCol, numCell = numCell, cells = sam, directions = 8,
              include = TRUE)
```

---

agentLocation

SELES - *Agent Location at initiation*

---

### Description

Sets the the location of the initiating agents. NOT YET FULLY IMPLEMENTED.

A SELES-like function to maintain conceptual backwards compatibility with that simulation tool. This is intended to ease transitions from [SELES](#).

You must know how to use SELES for these to be useful.

### Usage

```
agentLocation(map)
```

**Arguments**

map                    A `SpatialPoints*`, `SpatialPolygons*`, or `Raster*` object.

**Value**

Object of same class as provided as input. If a `Raster*`, then zeros are converted to NA.

**Author(s)**

Eliot McIntire

---

cir                    *Identify pixels in a circle or ring (donut) around an object.*

---

**Description**

Identify the pixels and coordinates that are at a (set of) buffer distance(s) of the objects passed into `coords`. This is similar to `rgeos::gBuffer` but much faster and without the geo referencing information. In other words, it can be used for similar problems, but where speed is important. This code is substantially adapted from `PlotRegionHighlighter::createCircle`.

**Usage**

```
cir(
  landscape,
  coords,
  loci,
  maxRadius = ncol(landscape)/4,
  minRadius = maxRadius,
  allowOverlap = TRUE,
  allowDuplicates = FALSE,
  includeBehavior = "includePixels",
  returnDistances = FALSE,
  angles = NA_real_,
  returnAngles = FALSE,
  returnIndices = TRUE,
  closest = FALSE,
  simplify = TRUE
)
```

**Arguments**

landscape            Raster on which the circles are built.

coords                Either a matrix with 2 (or 3) columns, x and y (and id), representing the coordinates (and an associated id, like cell index), or a `SpatialPoints*` object around which to make circles. Must be same coordinate system as the landscape argument. Default is missing, meaning it uses the default to `loci`



loci	Numeric. An alternative to coords. These are the indices on landscape to initiate this function. See coords. Default is one point in centre of landscape..
maxRadius	Numeric vector of length 1 or same length as coords
minRadius	Numeric vector of length 1 or same length as coords. Default is maxRadius, meaning return all cells that are touched by the narrow ring at that exact radius. If smaller than maxRadius, then this will create a buffer or donut or ring.
allowOverlap	Logical. Should duplicates across id be removed or kept. Default TRUE.
allowDuplicates	Logical. Should duplicates within id be removed or kept. Default FALSE. This is useful if the actual x, y coordinates are desired, rather than the cell indices. This will increase the size of the returned object.
includeBehavior	Character string. Currently accepts only "includePixels", the default, and "excludePixels". See details.
returnDistances	Logical. If TRUE, then a column will be added to the returned data.table that reports the distance from coords to every point that was in the circle/donut surrounding coords. Default FALSE, which is faster.
angles	Numeric. Optional vector of angles, in radians, to use. This will create "spokes" outward from coords. Default is NA, meaning, use internally derived angles that will "fill" the circle.
returnAngles	Logical. If TRUE, then a column will be added to the returned data.table that reports the angle from coords to every point that was in the circle/donut surrounding coords. Default FALSE.
returnIndices	Logical or numeric. If 1 or TRUE, will return a data.table with indices and values of successful spread events. If 2, it will simply return a vector of pixel indices of all cells that were touched. This will be the fastest option. If FALSE, then it will return a raster with values. See Details.
closest	Logical. When determining non-overlapping circles, should the function give preference to the closest loci or the first one (much faster). Default is FALSE, meaning the faster, though maybe not desired behaviour.
simplify	logical. If TRUE, then all duplicate pixels are removed. This means that some x, y combinations will disappear.

### Details

This function identifies all the pixels as defined by a donut with inner radius `minRadius` and outer radius of `maxRadius`. The `includeBehavior` defines whether the cells that intersect the radii but whose centres are not inside the donut are included `includePixels` or not `excludePixels` in the returned pixels identified. If this is `excludePixels`, and if a `minRadius` and `maxRadius` are equal, this will return no pixels.

### Value

A matrix with 4 columns, `id`, `indices`, `x`, `y`. The `x` and `y` indicate the exact coordinates of the indices (i.e., cell number) of the landscape associated with the ring or circle being identified by this function.

**See Also**

[rings](#) which uses `spread` internally. `cir` tends to be faster when there are few starting points, `rings` tends to be faster when there are many starting points. `cir` scales with  $\text{maxRadius}^2$  and `coords`. Another difference between the two functions is that `rings` takes the centre of the pixel as the centre of a circle, whereas `cir` takes the exact coordinates. See example. For the specific case of creating distance surfaces from specific points, see [distanceFromEachPoint](#), which is often faster. For the more general GIS buffering, see `rgeos::gBuffer`.

**Examples**

```
library(data.table)
library(sp)
library(raster)
library(quickPlot)

set.seed(1642)

# circle centred
ras <- raster(extent(0, 15, 0, 15), res = 1, val = 0)
middleCircle <- cir(ras)
ras[middleCircle[, "indices"]] <- 1
circlePoints <- SpatialPoints(middleCircle[, c("x", "y")])
if (interactive()) {
  clearPlot()
  Plot(ras)
  Plot(circlePoints, addTo = "ras")
}

# circles non centred
ras <- randomPolygons(ras, numTypes = 4)
n <- 2
agent <- SpatialPoints(coords = cbind(x = stats::runif(n, xmin(ras), xmax(ras)),
                                     y = stats::runif(n, xmin(ras), xmax(ras))))

cirs <- cir(ras, agent, maxRadius = 15, simplify = TRUE)
cirsSP <- SpatialPoints(coords = cirs[, c("x", "y")])
cirsRas <- raster(ras)
cirsRas[] <- 0
cirsRas[cirs[, "indices"]] <- 1

if (interactive()) {
  clearPlot()
  Plot(ras)
  Plot(cirsRas, addTo = "ras", cols = c("transparent", "#00000055"))
  Plot(agent, addTo = "ras")
  Plot(cirsSP, addTo = "ras")
}

# Example comparing rings and cir
a <- raster(extent(0, 30, 0, 30), res = 1)
hab <- gaussMap(a, speedup = 1) # if raster is large (>1e6 pixels) use speedup > 1
radius <- 4
```

```

n <- 2
coords <- SpatialPoints(coords = cbind(x = stats::runif(n, xmin(hab), xmax(hab)),
                                       y = stats::runif(n, xmin(hab), xmax(hab))))

# cirs
cirs <- cir(hab, coords, maxRadius = rep(radius, length(coords)), simplify = TRUE)

# rings
loci <- cellFromXY(hab, coordinates(coords))
cirs2 <- rings(hab, loci, maxRadius = radius, minRadius = radius - 1, returnIndices = TRUE)

# Plot both
ras1 <- raster(hab)
ras1[] <- 0
ras1[cirs[, "indices"]] <- cirs[, "id"]

ras2 <- raster(hab)
ras2[] <- 0
ras2[cirs2$indices] <- cirs2$id
if (interactive()) {
  clearPlot()
  Plot(ras1, ras2)
}

a <- raster(extent(0, 100, 0, 100), res = 1)
hab <- gaussMap(a, speedup = 1)
cirs <- cir(hab, coords, maxRadius = 44, minRadius = 0)
ras1 <- raster(hab)
ras1[] <- 0
cirsOverlap <- data.table(cirs)[, list(sumIDs = sum(id)), by = indices]
ras1[cirsOverlap$indices] <- cirsOverlap$sumIDs
if (interactive()) {
  clearPlot()
  Plot(ras1)
}

# Provide a specific set of angles
ras <- raster(extent(0, 330, 0, 330), res = 1)
ras[] <- 0
n <- 2
coords <- cbind(x = stats::runif(n, xmin(ras), xmax(ras)),
               y = stats::runif(n, xmin(ras), xmax(ras)))
circ <- cir(ras, coords, angles = seq(0, 2 * pi, length.out = 21),
           maxRadius = 200, minRadius = 0, returnIndices = FALSE,
           allowOverlap = TRUE, returnAngles = TRUE)

```

---

`cirSpecialQuick`      *This is a very fast version of `cir` with `allowOverlap = TRUE`, `allowDuplicates = FALSE`, `returnIndices = TRUE`, `returnDistances = TRUE`, and `includeBehavior = "excludePixels"`. It is used inside `spread2`, when `asymmetry` is active. The basic algorithm is to run `cir` just once, then add to the `x,y` coordinates of every locus.*

---

### Description

This is a very fast version of `cir` with `allowOverlap = TRUE`, `allowDuplicates = FALSE`, `returnIndices = TRUE`, `returnDistances = TRUE`, and `includeBehavior = "excludePixels"`. It is used inside `spread2`, when `asymmetry` is active. The basic algorithm is to run `cir` just once, then add to the `x,y` coordinates of every locus.

### Usage

```
.cirSpecialQuick(landscape, loci, maxRadius, minRadius)
```

### Arguments

<code>landscape</code>	Raster on which the circles are built.
<code>loci</code>	Numeric. An alternative to <code>coords</code> . These are the indices on <code>landscape</code> to initiate this function. See <code>coords</code> . Default is one point in centre of <code>landscape</code> .
<code>maxRadius</code>	Numeric vector of length 1 or same length as <code>coords</code>
<code>minRadius</code>	Numeric vector of length 1 or same length as <code>coords</code> . Default is <code>maxRadius</code> , meaning return all cells that are touched by the narrow ring at that exact radius. If smaller than <code>maxRadius</code> , then this will create a buffer or donut or ring.

---

`distanceFromEachPoint`      *Calculate distances and directions between many points and many grid cells*

---

### Description

This is a modification of [distanceFromPoints](#) for the case of many points. This version can often be faster for a single point because it does not return a `RasterLayer`. This is different than [distanceFromPoints](#) because it does not take the minimum distance from the set of points to all cells. Rather this returns the every pair-wise point distance. As a result, this can be used for doing inverse distance weightings, seed rain, cumulative effects of distance-based processes etc. If memory limitation is an issue, `maxDistance` will keep memory use down, but with the consequences that there will be a maximum distance returned. This function has the potential to use a lot of memory if there are a lot of from and to points.

**Usage**

```

distanceFromEachPoint(
  from,
  to = NULL,
  landscape,
  angles = NA_real_,
  maxDistance = NA_real_,
  cumulativeFn = NULL,
  distFn = function(dist) 1/(1 + dist),
  cl,
  ...
)

```

**Arguments**

from	Numeric matrix with 2 or 3 or more columns. They must include x and y, representing x and y coordinates of "from" cell. If there is a column named "id", it will be "id" from to, i.e., specific pair distances. All other columns will be included in the return value of the function.
to	Numeric matrix with 2 or 3 columns (or optionally more, all of which will be returned), x and y, representing x and y coordinates of "to" cells, and optional "id" which will be matched with "id" from from. Default is all cells.
landscape	RasterLayer. optional. This is only used if to is NULL, in which case all cells are considered to.
angles	Logical. If TRUE, then the function will return angles in radians, as well as distances.
maxDistance	Numeric in units of number of cells. The algorithm will build the whole surface (from from to to), but will remove all distances that are above this distance. Using this will keep memory use down.
cumulativeFn	A function that can be used to incrementally accumulate values in each to location, as the function iterates through each from. See Details.
distFn	A function. This can be a function of landscape, fromCells, toCells, and dist. If cumulativeFn is supplied, this will be used to convert the distances to some other set of units that will be accumulated by the cumulativeFn. See Details and examples.
cl	A cluster object. Optional. This would generally be created using parallel::makeCluster or equivalent. This is an alternative way, instead of beginCluster(), to use parallelism for this function, allowing for more control over cluster use.
...	Any additional objects needed for distFn.

**Details**

This function is cluster aware. If there is a cluster running, it will use it. To start a cluster use [beginCluster](#), with N being the number of cores to use. See examples in `SpaDES.core::experiment`.

If the user requires an id (indicating the from cell for each to cell) to be returned with the function, the user must add an identifier to the from matrix, such as "id". Otherwise, the function will only return the coordinates and distances.

`distanceFromEachPoint` calls `.pointDistance`, which is not intended to be called directly by the user.

This function has the potential to return a very large object, as it is doing pairwise distances (and optionally directions) between from and to. If there are memory limitations because there are many from and many to points, then `cumulativeFn` and `distFn` can be used. These two functions together will be used iteratively through the from points. The `distFn` should be a transformation of distances to be used by the `cumulativeFn` function. For example, if `distFn` is  $1 / (1+x)$ , the default, and `cumulativeFn` is ``+``, then it will do a sum of inverse distance weights. See examples.

### Value

A sorted matrix on `id` with same number of rows as `to`, but with one extra column, "dists", indicating the distance between from and to.

### See Also

[rings](#), [cir](#), [distanceFromPoints](#), which can all be made to do the same thing, under specific combinations of arguments. But each has different primary use cases. Each is also faster under different conditions. For instance, if `maxDistance` is relatively small compared to the number of cells in the landscape, then `cir` will likely be faster. If a minimum distance from all cells in the landscape to any cell in from, then `distanceFromPoints` will be fastest. This function scales best when there are many to points or all cells are used to = NULL (which is default).

### Examples

```
library(raster)
library(quickPlot)

n <- 2
distRas <- raster(extent(0, 40, 0, 40), res = 1)
coords <- cbind(x = round(runif(n, xmin(distRas), xmax(distRas))) + 0.5,
               y = round(runif(n, xmin(distRas), xmax(distRas))) + 0.5)

# inverse distance weights
dists1 <- distanceFromEachPoint(coords, landscape = distRas)
indices <- cellFromXY(distRas, dists1[, c("x", "y")])
invDist <- tapply(dists1[, "dists"], indices, function(x) sum(1 / (1 + x))) # idw function
distRas[] <- as.vector(invDist)
if (interactive()) {
  clearPlot()
  Plot(distRas)
}

# With iterative summing via cumulativeFn to keep memory use low, with same result
dists1 <- distanceFromEachPoint(coords[, c("x", "y")], drop = FALSE,
                               landscape = distRas, cumulativeFn = `+`)
idwRaster <- raster(distRas)
```

```

idwRaster[] <- dists1[, "val"]
if (interactive()) Plot(idwRaster)

all(idwRaster[] == distRas[]) # TRUE

# A more complex example of cumulative inverse distance sums, weighted by the value
# of the origin cell
ras <- raster(extent(0, 34, 0, 34), res = 1, val = 0)
rp <- randomPolygons(ras, numTypes = 10) ^ 2
n <- 15
cells <- sample(ncell(ras), n)
coords <- xyFromCell(ras, cells)
distFn <- function(landscape, fromCell, dist) landscape[fromCell] / (1 + dist)

#beginCluster(3) # can do parallel
dists1 <- distanceFromEachPoint(coords[, c("x", "y"), drop = FALSE],
                                landscape = rp, distFn = distFn, cumulativeFn = `+`)
#endCluster() # if beginCluster was run

idwRaster <- raster(ras)
idwRaster[] <- dists1[, "val"]
if (interactive()) {
  clearPlot()
  Plot(rp, idwRaster)
  sp1 <- SpatialPoints(coords)
  Plot(sp1, addTo = "rp")
  Plot(sp1, addTo = "idwRaster")
}

```

---

duplicatedInt

*Rcpp duplicated on integers using Rcpp Sugar*


---

### Description

.duplicatedInt does same as duplicated in R, but only on integers, and faster. It uses Rcpp sugar

### Usage

```
duplicatedInt(x)
```

### Arguments

x                    Integer Vector

### Value

A logical vector, as per duplicated

---

`dwrpnorm2`*Vectorized wrapped normal density function*

---

### Description

This is a modified version of `dwrpnorm` found in `CircStats` to allow for multiple angles at once (i.e., vectorized on `theta` and `mu`).

### Usage

```
dwrpnorm2(theta, mu, rho, sd = 1, acc = 1e-05, tol = acc)
```

### Arguments

<code>theta</code>	value at which to evaluate the density function, measured in radians.
<code>mu</code>	mean direction of distribution, measured in radians.
<code>rho</code>	mean resultant length of distribution.
<code>sd</code>	different way of select <code>rho</code> , see details below.
<code>acc</code>	parameter defining the accuracy of the estimation of the density. Terms are added to the infinite summation that defines the density function until successive estimates are within <code>acc</code> of each other.
<code>tol</code>	the same as <code>acc</code> .

### Author(s)

Eliot McIntire

### Examples

```
# Values for which to evaluate density
theta <- c(1:500) * 2 * pi / 500
# Compute wrapped normal density function
density <- c(1:500)
for(i in 1:500) density[i] <- dwrpnorm2(theta[i], pi, .75)
if (interactive()) plot(theta, density)
# Approximate area under density curve
sum(density * 2 * pi / 500)
```



---

fastCrop	<i>fastCrop is a wrapper around <code>velox::VeloxRaster_crop</code>, though <code>raster::crop</code> is faster under many tests.</i>
----------	--

---

### Description

fastCrop is a wrapper around `velox::VeloxRaster_crop`, though `raster::crop` is faster under many tests.

### Usage

```
fastCrop(x, y, ...)
```

### Arguments

x	Raster to crop
y	Extent object, or any object from which an Extent object can be extracted (see Details)
...	Additional arguments as for <a href="#">writeRaster</a>

### See Also

`velox::VeloxRaster_crop`

---

gaussMap	<i>Produce a raster of a random Gaussian process.</i>
----------	---

---

### Description

This is a wrapper for the `RFsimulate` function in the `RandomFields` package. The main addition is the `speedup` argument which allows for faster map generation. A `speedup` of 1 is normal and will get progressively faster as the number increases, at the expense of coarser pixel resolution of the pattern generated.

### Usage

```
gaussMap(  
  x,  
  scale = 10,  
  var = 1,  
  speedup = 1,  
  method = "RMexp",  
  alpha = 1,  
  inMemory = FALSE,  
  ...  
)
```

**Arguments**

x	A spatial object (e.g., a RasterLayer).
scale	The spatial scale in map units of the Gaussian pattern.
var	Spatial variance.
speedup	An numeric value indicating how much faster than 'normal' to generate maps. It may be necessary to give a value larger than 1 for large maps. Default is 1.
method	The type of model used to produce the Gaussian pattern. Should be one of "RMgauss" (Gaussian covariance model), "RMstable" (the stable powered exponential model), or the default, "RMexp" (exponential covariance model).
alpha	A required parameter of the 'RMstable' model. Should be in the interval [0,2] to provide a valid covariance function. Default is 1.
inMemory	Should the RasterLayer be forced to be in memory? Default FALSE.
...	Additional arguments to raster.

**Value**

A raster map with same extent as x, with a Gaussian random pattern.

**See Also**

RFsimulate and [extent](#)

**Examples**

```
## Not run:
if (require(RandomFields)) {
  library(raster)
  nx <- ny <- 100L
  r <- raster(nrows = ny, ncols = nx, xmn = -nx/2, xmx = nx/2, ymn = -ny/2, ymx = ny/2)
  speedup <- max(1, nx/5e2)
  map1 <- gaussMap(r, scale = 300, var = 0.03, speedup = speedup, inMemory = TRUE)
  if (interactive()) Plot(map1)

  # with non-default method
  map1 <- gaussMap(r, scale = 300, var = 0.03, method = "RMgauss")
}

## End(Not run)
```

---

heading	<i>Heading between spatial points.</i>
---------	--

---

**Description**

Determines the heading between spatial points.

**Usage**

```
heading(from, to)

## S4 method for signature 'SpatialPoints,SpatialPoints'
heading(from, to)

## S4 method for signature 'matrix,matrix'
heading(from, to)

## S4 method for signature 'matrix,SpatialPoints'
heading(from, to)

## S4 method for signature 'SpatialPoints,matrix'
heading(from, to)
```

**Arguments**

from	The starting position; an object of class SpatialPoints.
to	The ending position; an object of class SpatialPoints.

**Value**

The heading between the points, in degrees.

**Author(s)**

Eliot McIntire

**Examples**

```
library(sp)
N <- 10L # number of agents
x1 <- stats::runif(N, -50, 50) # previous X location
y1 <- stats::runif(N, -50, 50) # previous Y location
x0 <- stats::rnorm(N, x1, 5) # current X location
y0 <- stats::rnorm(N, y1, 5) # current Y location

# using SpatialPoints
prev <- SpatialPoints(cbind(x = x1, y = y1))
curr <- SpatialPoints(cbind(x = x0, y = y0))
```

```

heading(prev, curr)

# using matrix
prev <- matrix(c(x1, y1), ncol = 2, dimnames = list(NULL, c("x", "y")))
curr <- matrix(c(x0, y0), ncol = 2, dimnames = list(NULL, c("x", "y")))
heading(prev, curr)

#using both
prev <- SpatialPoints(cbind(x = x1, y = y1))
curr <- matrix(c(x0, y0), ncol = 2, dimnames = list(NULL, c("x", "y")))
heading(prev, curr)

prev <- matrix(c(x1, y1), ncol = 2, dimnames = list(NULL, c("x", "y")))
curr <- SpatialPoints(cbind(x = x0, y = y0))
heading(prev, curr)

```

---

initiateAgents

SELES - *Initiate agents*


---

## Description

Sets the the number of agents to initiate. THIS IS NOT FULLY IMPLEMENTED.

A SELES-like function to maintain conceptual backwards compatibility with that simulation tool. This is intended to ease transitions from **SELES**.

You must know how to use SELES for these to be useful.

## Usage

```

initiateAgents(map, numAgents, probInit, asSpatialPoints = TRUE, indices)

## S4 method for signature 'Raster,missing,missing,ANY,missing'
initiateAgents(map, numAgents, probInit, asSpatialPoints)

## S4 method for signature 'Raster,missing,Raster,ANY,missing'
initiateAgents(map, probInit, asSpatialPoints)

## S4 method for signature 'Raster,numeric,missing,ANY,missing'
initiateAgents(map, numAgents, probInit, asSpatialPoints = TRUE, indices)

## S4 method for signature 'Raster,numeric,Raster,ANY,missing'
initiateAgents(map, numAgents, probInit, asSpatialPoints)

## S4 method for signature 'Raster,missing,missing,ANY,numeric'
initiateAgents(map, numAgents, probInit, asSpatialPoints = TRUE, indices)

```

**Arguments**

map	RasterLayer with extent and resolution of desired return object
numAgents	numeric resulting from a call to <code>numAgents</code>
probInit	a Raster resulting from a <code>probInit</code> call
asSpatialPoints	logical. Should returned object be RasterLayer or SpatialPointsDataFrame (default)
indices	numeric. Indices of where agents should start

**Value**

A SpatialPointsDataFrame, with each row representing an individual agent

**Author(s)**

Eliot McIntire

**Examples**

```
library(magrittr)
library(raster)
library(quickPlot)

map <- raster(xmn = 0, xmx = 10, ymn = 0, ymx = 10, val = 0, res = 1)
map <- gaussMap(map, scale = 1, var = 4, speedup = 1)
pr <- probInit(map, p = (map/maxValue(map))^2)
agents <- initiateAgents(map, 100, pr)
if (interactive()) {
  clearPlot()
  Plot(map)
  Plot(agents, addTo = "map")
}
# Test that they are indeed selecting according to probabilities in pr
library(data.table)
dt1 <- data.table(table(round(map[agents], 0)))
setnames(dt1, old = "N", new = "count")
dt2 <- data.table(table(round(map[, 0])))
setnames(dt2, old = "N", new = "available")
dt <- dt1[dt2, on = "V1"] # join the counts and available data.tables
setnames(dt, old = "V1", new = "mapValue")
dt[, selection := count/available]
dt[is.na(selection), selection := 0]
if (interactive())
  with(dt, {plot(mapValue, selection)})

# Note, can also produce a Raster representing agents,
# then the number of points produced can't be more than
# the number of pixels:
agentsRas <- initiateAgents(map, 30, pr, asSpatialPoints = FALSE)
if (interactive()) Plot(agentsRas)
```

```

if (require(dplyr) && getRversion() >= 3.4) {
  # Check that the agents are more often at the higher probability areas based on pr
  if (utils::packageVersion("raster") >= "2.8-11") {
    out <- data.frame(stats::na.omit(crosstab(agentsRas, map)), table(round(map[]))) %>%
      dplyr::mutate(selectionRatio = Freq / Freq.1) %>%
      dplyr::select(-layer.1, -Var1) %>%
      dplyr::rename(Present = Freq, Avail = Freq.1, Type = layer.2)
  } else {
    out <- data.frame(stats::na.omit(crosstab(agentsRas, map)), table(round(map[]))) %>%
      dplyr::mutate(selectionRatio = Freq/Freq.1) %>%
      dplyr::select(-Var1, -Var1.1) %>%
      dplyr::rename(Present = Freq, Avail = Freq.1, Type = Var2)
  }
  out
}

```

---

inRange

*Test whether a number lies within range [a,b]*


---

### Description

Default values of  $a=0$ ;  $b=1$  allow for quick test if  $x$  is a probability.

### Usage

```
inRange(x, a = 0, b = 1)
```

### Arguments

$x$	values to be tested
$a$	lower bound (default 0)
$b$	upper bound (default 1)

### Value

Logical vectors. NA values in  $x$  are retained.

### Author(s)

Alex Chubaty

### Examples

```

set.seed(100)
x <- stats::rnorm(4) # -0.50219235  0.13153117 -0.07891709  0.88678481
inRange(x, 0, 1)

```

mergeRaster

*Split and re-merge RasterLayer(s)***Description**

splitRaster divides up a raster into an arbitrary number of pieces (tiles). Split rasters can be recombined using do.call(merge, y) or mergeRaster(y), where y <- splitRaster(x).

**Usage**

```
mergeRaster(x, fun = NULL)

## S4 method for signature 'list'
mergeRaster(x, fun = NULL)

splitRaster(
  r,
  nx = 1,
  ny = 1,
  buffer = c(0, 0),
  path = NA,
  cl,
  rType = "FLT4S"
)

## S4 method for signature 'RasterLayer'
splitRaster(
  r,
  nx = 1,
  ny = 1,
  buffer = c(0, 0),
  path = NA,
  cl,
  rType = "FLT4S"
)
```

**Arguments**

x	A list of split raster tiles (i.e., from splitRaster).
fun	Function (e.g. mean, min, or max that accepts a na.rm argument. The default is mean.
r	The raster to be split.
nx	The number of tiles to make along the x-axis.
ny	The number of tiles to make along the y-axis.

buffer	Numeric vector of length 2 giving the size of the buffer along the x and y axes. If these values less than or equal to 1 are used, this is interpreted as the number of pixels (cells) to use as a buffer. Values between 0 and 1 are interpreted as proportions of the number of pixels in each tile (rounded up to an integer value). Default is <code>c(0, 0)</code> , which means no buffer.
path	Character specifying the directory to which the split tiles will be saved. If missing, the function will write to memory.
c1	A cluster object. Optional. This would generally be created using <code>parallel::makeCluster</code> or equivalent. This is an alternative way, instead of <code>beginCluster()</code> , to use parallelism for this function, allowing for more control over cluster use.
rType	Data type of the split rasters. Defaults to <code>FLT4S</code> .

### Details

`mergeRaster` differs from `merge` in how overlapping tile regions are handled: `merge` retains the values of the first raster in the list. This has the consequence of retaining the values from the buffered region in the first tile in place of the values from the neighbouring tile. On the other hand, `mergeRaster` retains the values of the tile region, over the values in any buffered regions. This is useful for reducing edge effects when performing raster operations involving contagious processes. To use the average of cell values, or do another computation, use `mosaic`. `mergeRaster` is also faster than `merge`. `mergeRaster` also differs from `mosaic` in speed and ability to take a raster list. It can, however, use the average of cell values, or do other computations. At last, `mergeRaster` can also merge tiles of a split raster that were resampled and, therefore, could have had different changes in the buffer sizes on each side of the raster. If the user resamples the tiles and the new resolution is not a multiple of the original one, `mergeRaster` will use `mosaic` with the `max` function to merge the tiles with a message. The user can also pass the function to be used when `mosaic` is triggered.

This function is parallel-aware, using the same mechanism as used in `raster`. Specifically, if you start a cluster using `beginCluster`, then this function will automatically use that cluster. It is always a good idea to stop the cluster when finished, using `endCluster`.

### Value

`mergeRaster` returns a `RasterLayer` object.

`splitRaster` returns a list (length `nx*ny`) of cropped raster tiles.

### Author(s)

Yong Luo, Alex Chubaty, Tati Micheletti & Ian Eddy

Alex Chubaty and Yong Luo

### See Also

[merge](#), [mosaic](#)

[do.call](#), [merge](#).



**Examples**

```

library(raster)

# an example with dimensions:
# nrow: 77
# ncol: 101
# nlayers: 3
b <- brick(system.file("external/rlogo.grd", package = "raster"))
r <- b[[1]] # use first layer only
nx <- 1
ny <- 2

tmpdir <- file.path(tempdir(), "splitRaster-example")
dir.create(tmpdir)

y0 <- splitRaster(r, nx, ny, path = file.path(tmpdir, "y0")) # no buffer

# buffer: 10 pixels along both axes
y1 <- splitRaster(r, nx, ny, c(10, 10), path = file.path(tmpdir, "y1"))

# buffer: half the width and length of each tile
y2 <- splitRaster(r, nx, ny, c(0.5, 0.5), path = file.path(tmpdir, "y2"))

# parallel cropping
if (interactive()) {
  n <- pmin(parallel::detectCores(), 4) # use up to 4 cores
  beginCluster(n)
  y3 <- splitRaster(r, nx, ny, c(0.7, 0.7), path = file.path(tmpdir, "y3"))
  endCluster()
}

# the original raster:
if (interactive()) plot(r) # may require a call to `dev()` if using RStudio

# the split raster:
layout(mat = matrix(seq_len(nx * ny), ncol = nx, nrow = ny))
plotOrder <- c(4, 8, 12, 3, 7, 11, 2, 6, 10, 1, 5, 9)
if (interactive()) invisible(lapply(y0[plotOrder], plot))

# can be recombined using `raster::merge`
m0 <- do.call(merge, y0)
all.equal(m0, r) ## TRUE

m1 <- do.call(merge, y1)
all.equal(m1, r) ## TRUE

m2 <- do.call(merge, y2)
all.equal(m2, r) ## TRUE

# or recombine using mergeRaster
n0 <- mergeRaster(y0)
all.equal(n0, r) ## TRUE

```

```
n1 <- mergeRaster(y1)
all.equal(n1, r) ## TRUE

n2 <- mergeRaster(y2)
all.equal(n2, r) ## TRUE

unlink(tmpdir, recursive = TRUE)
```

---

middlePixel	<i>Return the (approximate) middle pixel on a raster</i>
-------------	--

---

### Description

This calculation is slightly different depending on whether the `nrow(ras)` and `ncol(ras)` are even or odd. It will return the exact middle pixel if these are odd, and the pixel just left and/or above the middle pixel if either dimension is even, respectively.

### Usage

```
middlePixel(ras)
```

### Arguments

ras	A Raster
-----	----------

---

move	<i>Move</i>
------	-------------

---

### Description

Wrapper for selecting different animal movement methods.

This version uses just turn angles and step lengths to define the correlated random walk.

### Usage

```
move(hypothesis = "crw", ...)

crw(agent, extent, stepLength, stddev, lonlat, torus = FALSE)

## S4 method for signature 'SpatialPointsDataFrame'
crw(agent, extent, stepLength, stddev, lonlat, torus = FALSE)

## S4 method for signature 'SpatialPoints'
crw(agent, extent, stepLength, stddev, lonlat, torus = FALSE)
```

**Arguments**

hypothesis	Character vector, length one, indicating which movement hypothesis/method to test/use. Currently defaults to 'crw' (correlated random walk) using crw.
...	arguments passed to the function in hypothesis
agent	A SpatialPoints* object. If a SpatialPointsDataFrame, 2 of the columns must be x1 and y1, indicating the previous location. If a SpatialPoints object, then x1 and y1 will be assigned randomly.
extent	An optional Extent object that will be used for torus.
stepLength	Numeric vector of length 1 or number of agents describing step length.
stddev	Numeric vector of length 1 or number of agents describing standard deviation of wrapped normal turn angles.
lonlat	Logical. If TRUE, coordinates should be in degrees. If FALSE coordinates represent planar ('Euclidean') space (e.g. units of meters)
torus	Logical. Should the movement be wrapped to the opposite side of the map, as determined by the extent argument. Default FALSE.

**Details**

This simple version of a correlated random walk is largely the version that was presented in Turchin 1998, but it was also used with bias modifications in McIntire, Schultz, Crone 2007.

**Value**

A SpatialPointsDataFrame object with updated spatial position defined by a single occurrence of step length(s) and turn angle(s).

**Author(s)**

Eliot McIntire  
Eliot McIntire

**References**

Turchin, P. 1998. Quantitative analysis of movement: measuring and modeling population redistribution in animals and plants. Sinauer Associates, Sunderland, MA.

McIntire, E. J. B., C. B. Schultz, and E. E. Crone. 2007. Designing a network for butterfly habitat restoration: where individuals, populations and landscapes interact. *Journal of Applied Ecology* 44:725-736.

**See Also**

[pointDistance](#)

---

numAgents	<i>SELES - Number of Agents to initiate</i>
-----------	---

---

**Description**

Sets the the number of agents to initiate. THIS IS NOT YET FULLY IMPLEMENTED.

A SELES-like function to maintain conceptual backwards compatibility with that simulation tool. This is intended to ease transitions from **SELES**.

You must know how to use SELES for these to be useful.

**Usage**

```
numAgents(N, probInit)
```

**Arguments**

N	Number of agents to initiate (integer scalar).
probInit	Probability of initializing an agent at the location.

**Value**

A numeric, indicating number of agents to start

**Author(s)**

Eliot McIntire

---

patchSize	<i>Patch size</i>
-----------	-------------------

---

**Description**

Patch size

**Usage**

```
patchSize(patches)
```

**Arguments**

patches	Number of patches.
---------	--------------------

---

pointDistance2      *Point distance with C++*

---

### Description

pointDistance2 performs Pythagorean Theorem and cbinds all columns from to to new dists column. It is only defined for one point (from) to many (to) points. pointDistance3 performs Pythagorean Theorem and is to be used internally within distanceFromEachPoint as an alternative to .pointDistance, where it does many points (from) to many (to) points, one from point at a time. The results are then rbinded internally. It does not cbind extra columns from to.

### Usage

```
pointDistance2(to, from)

pointDistance3(fromX, toX, fromY, toY, maxDistance)

.pointDistance(
  from,
  to,
  angles = NA,
  maxDistance = NA_real_,
  otherFromCols = FALSE
)
```

### Arguments

to	Numeric matrix with 2 or 3 columns (or optionally more, all of which will be returned), x and y, representing x and y coordinates of "to" cells, and optional "id" which will be matched with "id" from from. Default is all cells.
from	Numeric matrix with 2 or 3 or more columns. They must include x and y, representing x and y coordinates of "from" cell. If there is a column named "id", it will be "id" from to, i.e., specific pair distances. All other columns will be included in the return value of the function.
fromX	Numeric vector of x coordinates for 'from' points
toX	Numeric vector of x coordinates for 'to' points
fromY	Numeric vector of y coordinates for 'from' points
toY	Numeric vector of y coordinates for 'to' points
maxDistance	Numeric scalar. The maximum distance cutoff for returned distances.
angles	Logical. If TRUE, then the function will return angles in radians, as well as distances.
otherFromCols	TODO: description needed

### Details

A slightly faster way to calculate distances.

**Value**

pointDistance2: a matrix with all the to columns plus one extra dists column. pointDistance3: a matrix with x and y columns from to plus one extra dists column.

A matrix with 3 columns, x0, y0 and dists.

---

probInit	SELES - <i>Probability of Initiation</i>
----------	--

---

**Description**

Describes the probability of initiation of agents or events. *THIS IS NOT FULLY IMPLEMENTED.*

A SELES-like function to maintain conceptual backwards compatibility with that simulation tool. This is intended to ease transitions from **SELES**.

You must know how to use SELES for these to be useful.

**Usage**

```
probInit(map, p = NULL, absolute = NULL)
```

**Arguments**

map	A spatialObjects object. Currently, only provides CRS and, if p is not a raster, then all the raster dimensions.
p	probability, provided as a numeric or raster
absolute	logical. Is p absolute probabilities or relative?

**Value**

A RasterLayer with probabilities of initialization. There are several combinations of inputs possible and they each result in different behaviours.

If p is numeric or Raster and between 0 and 1, it is treated as an absolute probability, and a map will be produced with the p value(s) everywhere.

If p is numeric or Raster and not between 0 and 1, it is treated as a relative probability, and a map will be produced with p/max(p) value(s) everywhere.

If absolute is provided, it will override the previous statements, unless absolute = TRUE and p is not between 0 and 1 (i.e., is not a probability).

**Author(s)**

Eliot McIntire

---

randomPolygons	<i>Produce a RasterLayer of random polygons</i>
----------------	---

---

### Description

These are built with the [spread](#) function internally.

Produces a `SpatialPolygons` object with 1 feature that will have approximately an area equal to `area` (expecting `area` in hectares), `#'` and a centre at approximately `x`.

### Usage

```
randomPolygons(
  ras = raster(extent(0, 15, 0, 15), res = 1, vals = 0),
  numTypes = 2,
  ...
)
```

```
randomPolygon(x, hectares, area)
```

```
## S3 method for class 'SpatialPoints'
randomPolygon(x, hectares, area)
```

```
## S3 method for class 'matrix'
randomPolygon(x, hectares, area)
```

```
## S3 method for class 'SpatialPolygons'
randomPolygon(x, hectares, area)
```

### Arguments

<code>ras</code>	A raster that whose extent will be used for the <code>randomPolygons</code> .
<code>numTypes</code>	Numeric value. The number of unique polygon types to use.
<code>...</code>	Other arguments passed to <code>spread</code> . No known uses currently.
<code>x</code>	Either a <code>SpatialPoints</code> , <code>SpatialPolygons</code> , or <code>matrix</code> with two dimensions, 1 row, with the approximate centre of the new random polygon to create. If <code>matrix</code> , then longitude and latitude are assumed (epsg:4326)
<code>hectares</code>	Deprecated. Use <code>area</code> in meters squared.
<code>area</code>	A numeric, the approximate area in meters squared of the random polygon.

### Value

A map of extent `ext` with random polygons.

A `SpatialPolygons` object, with approximately the `area` request, centred approximately at the coordinates requested, in the projection of `x`

**See Also**

[spread](#), [raster](#), [randomPolygons](#)  
[gaussMap](#) and [randomPolygons](#)

**Examples**

```
library(quickPlot)

set.seed(1234)
Ras <- randomPolygons(numTypes = 5)
if (interactive()) {
  clearPlot()
  Plot(Ras, cols = c("yellow", "dark green", "blue", "dark red"))
}

library(raster)
# more complex patterning, with a range of patch sizes
a <- randomPolygons(numTypes = 400, raster(extent(0, 50, 0, 50), res = 1, vals = 0))
a[a<320] <- 0
a[a>=320] <- 1
suppressWarnings(clumped <- clump(a)) # warning sometimes occurs, but not important
aHist <- hist(table(getValues(clumped)), plot = FALSE)
if (interactive()) {
  clearPlot()
  Plot(a)
  Plot(aHist)
}

library(raster)
b <- SpatialPoints(cbind(-110, 59))
crs(b) <- sp::CRS("+init=epsg:4326")
a <- randomPolygon(b, area = 1e6)
if (interactive()) {
  plot(a)
  points(b, pch = 19)
}
```

---

randomStudyArea

*Create default study areas for use with SpaDES modules*


---

**Description**

Create default study areas for use with SpaDES modules

**Usage**

```
randomStudyArea(center = NULL, size = 10000, seed = NULL)
```



**Arguments**

center	SpatialPoints object specifying a set of coordinates and a projection. Default is an area in southern Alberta, Canada.
size	Numeric specifying the approximate size of the area in m <sup>2</sup> . Default 1e4.
seed	Numeric indicating the random seed to set internally (useful for ensuring the same study area is produced each time).

**Value**

SpatialPolygonsDataFrame

---

rasterizeReduced	<i>Convert reduced representation to full raster</i>
------------------	--

---

**Description**

Convert reduced representation to full raster

**Usage**

```
rasterizeReduced(
  reduced,
  fullRaster,
  newRasterCols,
  mapcode = names(fullRaster),
  ...
)
```

**Arguments**

reduced	data.frame or data.table that has at least one column of codes that are represented in the fullRaster.
fullRaster	RasterLayer of codes used in reduced that represents a spatial representation of the data.
newRasterCols	Character vector, length 1 or more, with the name(s) of the column(s) in reduced whose value will be returned as a Raster or list of Rasters.
mapcode	a character, length 1, with the name of the column in reduced that is represented in fullRaster.
...	Other arguments. None used yet.

**Value**

A RasterLayer or list of RasterLayer of with same dimensions as fullRaster representing newRasterCols spatially, according to the join between the mapcode contained within reduced and fullRaster

**Author(s)**

Eliot McIntire

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

```

library(data.table)
library(raster)
library(quickPlot)

ras <- raster(extent(0, 15, 0, 15), res = 1)
fullRas <- randomPolygons(ras, numTypes = 2)
names(fullRas) <- "mapcodeAll"
uniqueComms <- unique(fullRas)
reducedDT <- data.table(mapcodeAll = uniqueComms,
                        communities = sample(1:1000, length(uniqueComms)),
                        biomass = rnbinom(length(uniqueComms), mu = 4000, 0.4))
biomass <- rasterizeReduced(reducedDT, fullRas, "biomass")

# The default key is the layer name of the fullRas, so rekey incase of miskey
setkey(reducedDT, biomass)

communities <- rasterizeReduced(reducedDT, fullRas, "communities")
setColors(communities) <- c("blue", "orange", "red")
if (interactive()) {
  clearPlot()
  Plot(biomass, communities, fullRas)
}

```

rings

*Identifies all cells within a ring around the focal cells***Description**

Identifies the cell numbers of all cells within a ring defined by minimum and maximum distances from focal cells. Uses [spread](#) under the hood, with specific values set. Under many situations, this will be faster than using `rgeos::gBuffer` twice (once for smaller ring and once for larger ring, then removing the smaller ring cells).

**Usage**

```

rings(
  landscape,
  loci = NA_real_,
  id = FALSE,
  minRadius = 2,

```

```

    maxRadius = 5,
    allowOverlap = FALSE,
    returnIndices = FALSE,
    returnDistances = TRUE,
    ...
)

## S4 method for signature 'RasterLayer'
rings(
  landscape,
  loci = NA_real_,
  id = FALSE,
  minRadius = 2,
  maxRadius = 5,
  allowOverlap = FALSE,
  returnIndices = FALSE,
  returnDistances = TRUE,
  ...
)

```

### Arguments

landscape	A RasterLayer object. This defines the possible locations for spreading events to start and spread into. This can also be used as part of stopRule.
loci	A vector of locations in landscape. These should be cell indices. If user has x and y coordinates, these can be converted with <a href="#">cellFromXY</a> .
id	Logical. If TRUE, returns a raster of events ids. If FALSE, returns a raster of iteration numbers, i.e., the spread history of one or more events. NOTE: this is overridden if returnIndices is TRUE or 1 or 2.
minRadius	Numeric. Minimum radius to be included in the ring. Note: this is inclusive, i.e., >=.
maxRadius	Numeric. Maximum radius to be included in the ring. Note: this is inclusive, i.e., <=.
allowOverlap	Logical. If TRUE, then individual events can overlap with one another, i.e., they do not interact (this is slower than if allowOverlap = FALSE). Default is FALSE.
returnIndices	Logical or numeric. If 1 or TRUE, will return a data.table with indices and values of successful spread events. If 2, it will simply return a vector of pixel indices of all cells that were touched. This will be the fastest option. If FALSE, then it will return a raster with values. See Details.
returnDistances	Logical. Should the function include a column with the individual cell distances from the locus where that event started. Default is FALSE. See Details.
...	Any other argument passed to spread

### Value

This will return a data.table with columns as described in spread when returnIndices = TRUE.

**Author(s)**

Eliot McIntire

**See Also**

`cir` which uses a different algorithm. `cir` tends to be faster when there are few starting points, `rings` tends to be faster when there are many starting points. Another difference between the two functions is that `rings` takes the centre of the pixel as the centre of a circle, whereas `cir` takes the exact coordinates. See example.

`rgeos::gBuffer`

**Examples**

```
library(raster)
library(quickPlot)

# Make random forest cover map
emptyRas <- raster(extent(0, 1e2, 0, 1e2), res = 1)

# start from two cells near middle
loci <- (ncell(emptyRas) / 2 - ncol(emptyRas)) / 2 + c(-3, 3)

# Allow overlap
## TODO: need to fix a bug when allowOverlap = TRUE
# emptyRas[] <- 0
# rngs <- rings(emptyRas, loci = loci, allowOverlap = TRUE, returnIndices = TRUE)
# # Make a raster that adds together all id in a cell
# wOverlap <- rngs[, list(sumEventID = sum(id)), by = "indices"]
# emptyRas[wOverlap$indices] <- wOverlap$sumEventID
# if (interactive()) {
#   clearPlot()
#   Plot(emptyRas)
# }

# No overlap is default, occurs randomly
emptyRas[] <- 0
rngs <- rings(emptyRas, loci = loci, minRadius = 7, maxRadius = 9, returnIndices = TRUE)
emptyRas[rngs$indices] <- rngs$id
if (interactive()) {
  clearPlot()
  Plot(emptyRas)
}

# Variable ring widths, including centre cell for smaller one
emptyRas[] <- 0
rngs <- rings(emptyRas, loci = loci, minRadius = c(0, 7), maxRadius = c(8, 18),
              returnIndices = TRUE)
emptyRas[rngs$indices] <- rngs$id
if (interactive()) {
  clearPlot()
  Plot(emptyRas)
}
```

```
}

```

---

runifC	<i>Rcpp Sugar version of runif</i>
--------	------------------------------------

---

**Description**

Slightly faster than runif, and used a lot

**Usage**

```
runifC(N)
```

**Arguments**

N	Integer Vector
---	----------------

**Value**

A vector of uniform random numbers as per runif

---

specificNumPerPatch	<i>Initiate a specific number of agents in a map of patches</i>
---------------------	---

---

**Description**

Instantiate a specific number of agents per patch. The user can either supply a table of how many to initiate in each patch, linked by a column in that table called pops.

**Usage**

```
specificNumPerPatch(patches, numPerPatchTable = NULL, numPerPatchMap = NULL)
```

**Arguments**

patches	RasterLayer of patches, with some sort of a patch id.
numPerPatchTable	A data.frame or data.table with a column named pops that matches the patches patch ids, and a second column num.in.pop with population size in each patch.
numPerPatchMap	A RasterLayer exactly the same as patches but with agent numbers rather than ids as the cell values per patch.

**Value**

A raster with 0s and 1s, where the 1s indicate starting locations of agents following the numbers above.

**Examples**

```

library(data.table)
library(raster)
library(quickPlot)

set.seed(1234)
Ntypes <- 4
ras <- randomPolygons(numTypes = Ntypes)
if (interactive()) {
  clearPlot()
  Plot(ras)
}

# Use numPerPatchTable
patchDT <- data.table(pops = 1:Ntypes, num.in.pop = c(1, 3, 5, 7))
rasAgents <- specificNumPerPatch(ras, patchDT)
rasAgents[is.na(rasAgents)] <- 0

library(testthat)
expect_true(all(unname(table(ras[rasAgents])) == patchDT$num.in.pop))

# Use numPerPatchMap
rasPatches <- ras
for (i in 1:Ntypes) {
  rasPatches[rasPatches==i] <- patchDT$num.in.pop[i]
}
if (interactive()) {
  clearPlot()
  Plot(ras, rasPatches)
}
rasAgents <- specificNumPerPatch(ras, numPerPatchMap = rasPatches)
rasAgents[is.na(rasAgents)] <- 0
if (interactive()) {
  clearPlot()
  Plot(rasAgents)
}

```

---

spokes

*Identify outward radiating spokes from initial points*


---

**Description**

This is a generalized version of a notion of a viewshed. The main difference is that there can be many "viewpoints".

**Usage**

```
spokes(
```

```

landscape,
coords,
loci,
maxRadius = ncol(landscape)/4,
minRadius = maxRadius,
allowOverlap = TRUE,
stopRule = NULL,
includeBehavior = "includePixels",
returnDistances = FALSE,
angles = NA_real_,
nAngles = NA_real_,
returnAngles = FALSE,
returnIndices = TRUE,
...
)

## S4 method for signature 'RasterLayer,SpatialPoints,missing'
spokes(
  landscape,
  coords,
  loci,
  maxRadius = ncol(landscape)/4,
  minRadius = maxRadius,
  allowOverlap = TRUE,
  stopRule = NULL,
  includeBehavior = "includePixels",
  returnDistances = FALSE,
  angles = NA_real_,
  nAngles = NA_real_,
  returnAngles = FALSE,
  returnIndices = TRUE,
  ...
)

```

### Arguments

landscape	Raster on which the circles are built.
coords	Either a matrix with 2 (or 3) columns, x and y (and id), representing the coordinates (and an associated id, like cell index), or a <code>SpatialPoints*</code> object around which to make circles. Must be same coordinate system as the landscape argument. Default is missing, meaning it uses the default to <code>loci</code>
loci	Numeric. An alternative to <code>coords</code> . These are the indices on landscape to initiate this function. See <code>coords</code> . Default is one point in centre of landscape..
maxRadius	Numeric vector of length 1 or same length as <code>coords</code>
minRadius	Numeric vector of length 1 or same length as <code>coords</code> . Default is <code>maxRadius</code> , meaning return all cells that are touched by the narrow ring at that exact radius. If smaller than <code>maxRadius</code> , then this will create a buffer or donut or ring.

<code>allowOverlap</code>	Logical. Should duplicates across id be removed or kept. Default TRUE.
<code>stopRule</code>	A function. If the spokes are to stop. This can be a function of <code>landscape</code> , <code>fromCell</code> , <code>toCell</code> , <code>x</code> (distance from coords cell), or any other named argument passed into the <code>...</code> of this function. See examples.
<code>includeBehavior</code>	Character string. Currently accepts only "includePixels", the default, and "excludePixels". See details.
<code>returnDistances</code>	Logical. If TRUE, then a column will be added to the returned <code>data.table</code> that reports the distance from <code>coords</code> to every point that was in the circle/donut surrounding <code>coords</code> . Default FALSE, which is faster.
<code>angles</code>	Numeric. Optional vector of angles, in radians, to use. This will create "spokes" outward from <code>coords</code> . Default is NA, meaning, use internally derived angles that will "fill" the circle.
<code>nAngles</code>	Numeric, length one. Alternative to <code>angles</code> . If provided, the function will create a sequence of angles from 0 to $2\pi$ , with a length <code>nAngles</code> , and not including $2\pi$ . Will not be used if <code>angles</code> is provided, and will show warning of both are given.
<code>returnAngles</code>	Logical. If TRUE, then a column will be added to the returned <code>data.table</code> that reports the angle from <code>coords</code> to every point that was in the circle/donut surrounding <code>coords</code> . Default FALSE.
<code>returnIndices</code>	Logical or numeric. If 1 or TRUE, will return a <code>data.table</code> with indices and values of successful spread events. If 2, it will simply return a vector of pixel indices of all cells that were touched. This will be the fastest option. If FALSE, then it will return a raster with values. See Details.
<code>...</code>	Objects to be used by <code>stopRule()</code> . See examples.

### Value

A matrix containing columns `id` (representing the row numbers of `coords`), `angles` (from `coords` to each point along the spokes), `x` and `y` coordinates of each point along the spokes, the corresponding indices on the `landscape Raster`, `dist`s (the distances between each `coords` and each point along the spokes), and `stop`, indicating if it was a point that caused a spoke to stop going outwards due to `stopRule`.

### Author(s)

Eliot McIntire

### Examples

```
library(sp)
library(raster)
library(quickPlot)

set.seed(1234)

ras <- raster(extent(0, 10, 0, 10), res = 1, val = 0)
```



```

rp <- randomPolygons(ras, numTypes = 10)

clearPlot()
Plot(rp)

angles <- seq(0, pi * 2, length.out = 17)
angles <- angles[-length(angles)]
n <- 2
loci <- sample(ncell(rp), n)
coords <- SpatialPoints(xyFromCell(rp, loci))
stopRule <- function(landscape) landscape < 3
d2 <- spokes(rp, coords = coords, stopRule = stopRule,
             minRadius = 0, maxRadius = 50,
             returnAngles = TRUE, returnDistances = TRUE,
             allowOverlap = TRUE, angles = angles, returnIndices = TRUE)

# Assign values to the "patches" that were in the viewshed of a ray
rasB <- raster(ras)
rasB[] <- 0
rasB[d2[d2[, "stop"] == 1, "indices"]] <- 1

Plot(rasB, addTo = "rp", zero.color = "transparent", cols = "red")

if (NROW(d2) > 0) {
  sp1 <- SpatialPoints(d2[, c("x", "y")])
  Plot(sp1, addTo = "rp", pch = 19, size = 5, speedup = 0.1)
}
Plot(coords, addTo = "rp", pch = 19, size = 6, cols = "blue", speedup = 0.1)

clearPlot()

```

---

spread

*Simulate a spread process on a landscape.*


---

### Description

This can be used to simulate fires, seed dispersal, calculation of iterative, concentric landscape values (symmetric or asymmetric) and many other things. Essentially, it starts from a collection of cells (`loci`) and spreads to neighbours, according to the `directions` and `spreadProb` arguments. This can become quite general, if `spreadProb` is 1 as it will expand from every `loci` until all cells in the landscape have been covered. With `id` set to `TRUE`, the resulting map will be classified by the index of the cell where that event propagated from. This can be used to examine things like fire size distributions. **NOTE:** See also [spread2](#), which is more robust and can be used to build custom functions. However, under some conditions, this `spread` function is faster. The two functions can accomplish many of the same things, and key differences are internal.

### Usage

```
spread(
```

```
landscape,
loci = NA_real_,
spreadProb = 0.23,
persistence = 0,
mask = NA,
maxSize = 100000000L,
directions = 8L,
iterations = 1000000L,
lowMemory = getOption("spades.lowMemory", FALSE),
returnIndices = FALSE,
returnDistances = FALSE,
mapID = NULL,
id = FALSE,
plot.it = FALSE,
spreadProbLater = NA_real_,
spreadState = NA,
circle = FALSE,
circleMaxRadius = NA_real_,
stopRule = NA,
stopRuleBehavior = "includeRing",
allowOverlap = FALSE,
asymmetry = NA_real_,
asymmetryAngle = NA_real_,
quick = FALSE,
neighProbs = NULL,
exactSizes = FALSE,
relativeSpreadProb = FALSE,
...
)
```

```
## S4 method for signature 'RasterLayer'
spread(
  landscape,
  loci,
  spreadProb,
  persistence,
  mask,
  maxSize,
  directions,
  iterations,
  lowMemory,
  returnIndices,
  returnDistances,
  mapID,
  id,
  plot.it,
  spreadProbLater,
  spreadState,
```

```

circle,
circleMaxRadius,
stopRule,
stopRuleBehavior,
allowOverlap,
asymmetry,
asymmetryAngle,
quick,
neighProbs,
exactSizes,
relativeSpreadProb,
.zeroVector,
...
)

```

### Arguments

landscape	A RasterLayer object. This defines the possible locations for spreading events to start and spread into. This can also be used as part of stopRule.
loci	A vector of locations in landscape. These should be cell indices. If user has x and y coordinates, these can be converted with <a href="#">cellFromXY</a> .
spreadProb	Numeric, or RasterLayer. If numeric of length 1, then this is the global probability of spreading into each cell from a neighbour. If a raster (or a vector of length ncell(landscape), resolution and extent of landscape), then this will be the cell-specific probability. Default is 0.23. If a spreadProbLater is provided, then this is only used for the first iteration. Also called "escape probability". See section on "Breaking out of spread events".
persistence	A length 1 probability that an active cell will continue to burn, per time step.
mask	non-NULL, a RasterLayer object congruent with landscape whose elements are 0, 1, where 1 indicates "cannot spread to". Currently not implemented, but identical behaviour can be achieved if spreadProb has zeros in all unspreadable locations.
maxSize	Numeric. Maximum number of cells for a single or all events to be spread. Recycled to match loci length, if it is not as long as loci. See section on Breaking out of spread events.
directions	The number of adjacent cells in which to look; default is 8 (Queen case). Can only be 4 or 8.
iterations	Number of iterations to spread. Leaving this NULL allows the spread to continue until stops spreading itself (i.e., exhausts itself).
lowMemory	Logical. If true, then function uses package <b>ff</b> internally. This is slower, but much lower memory footprint.
returnIndices	Logical or numeric. If 1 or TRUE, will return a data.table with indices and values of successful spread events. If 2, it will simply return a vector of pixel indices of all cells that were touched. This will be the fastest option. If FALSE, then it will return a raster with values. See Details.

returnDistances	Logical. Should the function include a column with the individual cell distances from the locus where that event started. Default is FALSE. See Details.
mapID	Deprecated. Use id.
id	Logical. If TRUE, returns a raster of events ids. If FALSE, returns a raster of iteration numbers, i.e., the spread history of one or more events. NOTE: this is overridden if returnIndices is TRUE or 1 or 2.
plot.it	If TRUE, then plot the raster at every iteration, so one can watch the spread event grow.
spreadProbLater	Numeric, or RasterLayer. If provided, then this will become the spreadProb after the first iteration. See Details.
spreadState	data.table. This should be the output of a previous call to spread, where returnIndices was TRUE. Default NA, meaning the spread is starting from loci. See Details.
circle	Logical. If TRUE, then outward spread will be by equidistant rings, rather than solely by adjacent cells (via directions arg.). Default is FALSE. Using circle = TRUE can be dramatically slower for large problems. Note, this should usually be used with spreadProb = 1.
circleMaxRadius	Numeric. A further way to stop the outward spread of events. If circle is TRUE, then it will grow to this maximum radius. See section on Breaking out of spread events. Default is NA.
stopRule	A function which will be used to assess whether each individual cluster should stop growing. This function can be an argument of "landscape", "id", "cells", and any other named vectors, a named list of named vectors, or a named data.frame with column names passed to spread in the ... Default NA, meaning that spreading will not stop as a function of the landscape. See section on "Breaking out of spread events" and examples.
stopRuleBehavior	Character. Can be one of "includePixel", "excludePixel", "includeRing", or "excludeRing". If stopRule contains a function, this argument is used determine what to do with the cell(s) that caused the rule to be TRUE. See details. Default is "includeRing" which means to accept the entire ring of cells that caused the rule to be TRUE.
allowOverlap	Logical. If TRUE, then individual events can overlap with one another, i.e., they do not interact (this is slower than if allowOverlap = FALSE). Default is FALSE.
asymmetry	A numeric indicating the ratio of the asymmetry to be used. Default is NA, indicating no asymmetry. See details. This is still experimental. <b>Use with caution.</b>
asymmetryAngle	A numeric indicating the angle in degrees (0 is "up", as in North on a map), that describes which way the asymmetry is.
quick	Logical. If TRUE, then several potentially time consuming checking (such as inRange) will be skipped. This should only be used if there is no concern about checking to ensure that inputs are legal.

neighProbs	A numeric vector, whose sum is 1. It indicates the probabilities an individual spread iteration spreading to 1:length(neighProbs) neighbours.
exactSizes	Logical. If TRUE, then the maxSize will be treated as exact sizes, i.e., the spread events will continue until they are floor(maxSize). This is overridden by iterations, but if iterations is run, and individual events haven't reached maxSize, then the returned data.table will still have at least one active cell per event that did not achieve maxSize, so that the events can continue if passed into spread with spreadState.
relativeSpreadProb	Logical. If TRUE, then spreadProb will be rescaled *within* the directions neighbours, such that the sum of the probabilities of all neighbours will be 1. Default FALSE, unless spreadProb values are not contained between 0 and 1, which will force relativeSpreadProb to be TRUE.
...	Additional named vectors or named list of named vectors required for stopRule. These vectors should be as long as required e.g., length loci if there is one value per event.
.zeroVector	DESCRIPTION NEEDED

## Details

For large rasters, a combination of lowMemory = TRUE and returnIndices = TRUE or returnIndices = 2 will be fastest and use the least amount of memory.

This function can be interrupted before all active cells are exhausted if the iterations value is reached before there are no more active cells to spread into. If this is desired, returnIndices should be TRUE and the output of this call can be passed subsequently as an input to this same function. This is intended to be used for situations where external events happen during a spread event, or where one or more arguments to the spread function change before a spread event is completed. For example, if it is desired that the spreadProb change before a spread event is completed because, for example, a fire is spreading, and a new set of conditions arise due to a change in weather.

asymmetry is currently used to modify the spreadProb in the following way. First for each active cell, spreadProb is converted into a length 2 numeric of Low and High spread probabilities for that cell: spreadProbsLH <- (spreadProb\*2) // (asymmetry+1)\*c(1, asymmetry), whose ratio is equal to asymmetry. Then, using asymmetryAngle, the angle between the initial starting point of the event and all potential cells is found. These are converted into a proportion of the angle from -asymmetryAngle to asymmetryAngle using: angleQuality <- (cos(angles - rad(asymmetryAngle)) + 1) / 2

These are then converted to multiple spreadProbs by spreadProbs <- lowSpreadProb + (angleQuality \* diff(spreadProbsLH)) To maintain an expected spreadProb that is the same as the asymmetric spreadProbs, these are then rescaled so that the mean of the asymmetric spreadProbs is always equal to spreadProb at every iteration: spreadProbs <- spreadProbs - diff(c(spreadProb, mean(spreadProbs)))

## Value

Either a RasterLayer indicating the spread of the process in the landscape or a data.table if returnIndices is TRUE. If a RasterLayer, then it represents every cell in which a successful spread event occurred. For the case of, say, a fire this would represent every cell that burned. If allowOverlap is TRUE, This RasterLayer will represent the sum of the individual event ids (which

are numerics `seq_along(loci)`. This will generally be of minimal use because it won't be possible to distinguish if event 2 overlapped with event 5 or if it was just event 7.

If `returnIndices` is TRUE, then this function returns a `data.table` with columns:

<code>id</code>	an arbitrary ID <code>1:length(loci)</code> identifying unique clusters of spread events, i.e., all cells that have been spread
<code>initialLocus</code>	the initial cell number of that particular spread event.
<code>indices</code>	The cell indices of cells that have been touched by the spread algorithm.
<code>active</code>	a logical indicating whether the cell is active (i.e., could still be a source for spreading) or not (no spreading)

This will generally be more useful when `allowOverlap` is TRUE.

### Breaking out of spread events

There are 4 ways for the spread to "stop" spreading. Here, each "event" is defined as all cells that are spawned from a single starting loci. So, one spread call can have multiple spreading "events". The ways outlined below are all acting at all times, i.e., they are not mutually exclusive. Therefore, it is the user's responsibility to make sure the different rules are interacting with each other correctly. Using `spreadProb` or `maxSize` are computationally fastest, sometimes dramatically so.

<code>spreadProb</code>	Probabilistically, if <code>spreadProb</code> is low enough, active spreading events will stop. In practice, active spreading events will stop.
<code>maxSize</code>	This is the number of cells that are "successfully" turned on during a spreading event. This can be vector.
<code>circleMaxRadius</code>	If <code>circle</code> is TRUE, then this will be the maximum radius reached, and then the event will stop. This is vector.
<code>stopRule</code>	This is a function that can use "landscape", "id", "cells", or any named vector passed into <code>spread</code> in the <code>args</code> .

The spread function does not return the result of this `stopRule`. If, say, an event has both `circleMaxRadius` and `stopRule`, and it is the `circleMaxRadius` that caused the event spreading to stop, there will be no indicator returned from this function that indicates which rule caused the stop.

`stopRule` has many use cases. One common use case is evaluating a neighbourhood around a focal set of points. This provides, therefore, an alternative to the `buffer` function or `focal` function. In both of those cases, the window/buffer size must be an input to the function. Here, the resulting size can be emergent based on the incremental growing and calculating of the landscape values underlying the spreading event.

### stopRuleBehavior

This determines how the `stopRule` should be implemented. Because spreading occurs outwards in concentric circles or shapes, one cell width at a time, there are 4 possible ways to interpret the logical inequality defined in `stopRule`. In order of number of cells included in resulting events, from most cells to fewest cells:

"includeRing"	Will include the entire ring of cells that, as a group, caused <code>stopRule</code> to be TRUE.
"includePixel"	Working backwards from the entire ring that caused the <code>stopRule</code> to be TRUE, this will iteratively randomize the cells until the <code>stopRule</code> is no longer TRUE.
"excludePixel"	Like "includePixel", but it will not add back the cell that causes <code>stopRule</code> to be TRUE.
"excludeRing"	Analogous to "excludePixel", but for the entire final ring of cells added. This will exclude the entire ring.

**Author(s)**

Eliot McIntire and Steve Cumming

**See Also**

[spread2](#) for a different implementation of the same algorithm. It is more robust, meaning, there will be fewer unexplainable errors, and the behaviour has been better tested, so it is more likely to be exactly as described under all argument combinations. Also, [rings](#) which uses `spread` but with specific argument values selected for a specific purpose. [distanceFromPoints.cir](#) to create "circles"; it is fast for many small problems.

**Examples**

```
library(raster)
library(RColorBrewer)
library(quickPlot)

# Make random forest cover map
set.seed(123)
emptyRas <- raster(extent(0, 1e2, 0, 1e2), res = 1)
hab <- randomPolygons(emptyRas, numTypes = 40)
names(hab) <- "hab"
mask <- raster(emptyRas)
mask <- setValues(mask, 0)
mask[1:5000] <- 1
numCol <- ncol(emptyRas)
numCell <- ncell(emptyRas)
directions <- 8

# Can use transparent as a colour
setColors(hab) <- paste(c("transparent", brewer.pal(8, "Greys")))

# note speedup is equivalent to making pyramids, so, some details are lost
clearPlot()
Plot(hab, speedup = 3)

# initiate 10 fires
startCells <- as.integer(sample(1:ncell(emptyRas), 100))
fires <- spread(hab, loci = startCells, 0.235, persistence = 0, numNeighs = 2,
               mask = NULL, maxSize = 1e8, directions = 8, iterations = 1e6, id = TRUE)

#set colours of raster, including a transparent layer for zeros
setColors(fires, 10) <- c("transparent", brewer.pal(8, "Reds")[5:8])
Plot(fires)
Plot(fires, addTo = "hab")

#alternatively, set colours using cols= in the Plot function
clearPlot()
Plot(hab)
Plot(fires) # default colour range makes zero transparent.

# Instead, to give a colour to the zero values, use \code{zero.color=}
```

```

Plot(fires, addTo = "hab",
     cols = colorRampPalette(c("orange", "darkred"))(10), zero.color = "transparent")

hab2 <- hab
Plot(hab2)
Plot(fires, addTo = "hab2", zero.color = "transparent",
     cols = colorRampPalette(c("orange", "darkred"))(10))
# or overplot the original (NOTE: legend stays at original values)
Plot(fires, cols = topo.colors(10), new = TRUE, zero.color = "white")

##-----
## Continue event by passing interrupted object into spreadState
##-----

## Interrupt a spread event using iterations - need `returnIndices = TRUE` to
## use outputs as new inputs in next iteration
fires <- spread(hab, loci = as.integer(sample(1:ncell(hab), 10)),
               returnIndices = TRUE, 0.235, 0, NULL, 1e8, 8, iterations = 3, id = TRUE)
fires[, list(size = length(initialLocus)), by = id] # See sizes of fires

fires2 <- spread(hab, loci = NA_real_, returnIndices = TRUE, 0.235, 0, NULL,
                1e8, 8, iterations = 2, id = TRUE, spreadState = fires)
# NOTE events are assigned arbitrary IDs, starting at 1

## Add new fires to the already burning fires
fires3 <- spread(hab, loci = as.integer(sample(1:ncell(hab), 10)),
               returnIndices = TRUE, 0.235, 0, NULL, 1e8, 8, iterations = 1,
               id = TRUE, spreadState = fires)
fires3[, list(size = length(initialLocus)), by = id] # See sizes of fires
# NOTE old ids are maintained, new events get ids beginning above previous
# maximum (e.g., new fires 11 to 20 here)

## Use data.table and loci...
fires <- spread(hab, loci = as.integer(sample(1:ncell(hab), 10)),
               returnIndices = TRUE, 0.235, 0, NULL, 1e8, 8, iterations = 2, id = TRUE)
fullRas <- raster(hab)
fullRas[] <- 1:ncell(hab)
burned <- fires[active == FALSE]
burnedMap <- rasterizeReduced(burned, fullRas, "id", "indices")

clearPlot()
Plot(burnedMap, new = TRUE)

#####
## stopRule examples
#####

# examples with stopRule, which means that the eventual size is driven by the values on the raster
# passed in to the landscape argument
set.seed(1234)
stopRule1 <- function(landscape) sum(landscape) > 50
stopRuleA <- spread(hab, loci = as.integer(sample(1:ncell(hab), 10)), 1, 0, NULL,
                  maxSize = 1e6, 8, 1e6, id = TRUE, circle = TRUE, stopRule = stopRule1)

```



```

set.seed(1234)
stopRule2 <- function(landscape) sum(landscape) > 100
# using stopRuleBehavior = "excludePixel"
stopRuleB <- spread(hab, loci = as.integer(sample(1:ncell(hab), 10)), 1, 0, NULL,
                    maxSize = 1e6, 8, 1e6, id = TRUE, circle = TRUE, stopRule = stopRule2,
                    stopRuleBehavior = "excludePixel")

# using stopRuleBehavior = "includeRing", means that end result is slightly larger patches, as a
# complete "iteration" of the spread algorithm is used.
set.seed(1234)
stopRuleBNotExact <- spread(hab, loci = as.integer(sample(1:ncell(hab), 10)), 1, 0,
                            NULL, maxSize = 1e6, 8, 1e6, id = TRUE, circle = TRUE,
                            stopRule = stopRule2)

clearPlot()
Plot(stopRuleA, stopRuleB, stopRuleBNotExact)

# Test that the stopRules work
# stopRuleA was not exact, so each value will "overshoot" the stopRule, here it was hab>50
foo <- cbind(vals = hab[stopRuleA], id = stopRuleA[stopRuleA > 0]);
tapply(foo[, "vals"], foo[, "id"], sum) # Correct ... all are above 50

# stopRuleB was exact, so each value will be as close as possible while rule still is TRUE
# Because we have discrete cells, these numbers will always slightly under the rule
foo <- cbind(vals = hab[stopRuleB], id = stopRuleB[stopRuleB > 0]);
tapply(foo[, "vals"], foo[, "id"], sum) # Correct ... all are above 50

# stopRuleB_notExact will overshoot
foo <- cbind(vals = hab[stopRuleBNotExact], id = stopRuleBNotExact[stopRuleBNotExact > 0]);
tapply(foo[, "vals"], foo[, "id"], sum) # Correct ... all are above 50

# Cellular automata shapes
# Diamonds - can make them with: a boolean raster, directions = 4,
#   stopRule in place, spreadProb = 1
diamonds <- spread(hab > 0, spreadProb = 1, directions = 4, id = TRUE, stopRule = stopRule2)

clearPlot()
Plot(diamonds)

# Squares - can make them with: a boolean raster, directions = 8,
#   stopRule in place, spreadProb = 1
squares <- spread(hab > 0, spreadProb = 1, directions = 8, id = TRUE, stopRule = stopRule2)
Plot(squares)

# Interference shapes - can make them with: a boolean raster, directions = 8,
#   stopRule in place, spreadProb = 1
stopRule2 <- function(landscape) sum(landscape) > 200
squashedDiamonds <- spread(hab > 0, spreadProb = 1,
                           loci = (ncell(hab) - ncol(hab)) / 2 + c(4, -4),
                           directions = 4, id = TRUE, stopRule = stopRule2)

clearPlot()
Plot(squashedDiamonds)

```

```

# Circles with spreadProb < 1 will give "more" circular shapes, but definitely not circles
stopRule2 <- function(landscape) sum(landscape) > 200
seed <- sample(1e4, 1)
set.seed(seed)
circlish <- spread(hab > 0, spreadProb = 0.23,
                  loci = (ncell(hab) - ncol(hab)) / 2 + c(4, -4),
                  directions = 8, id = TRUE, circle = TRUE)#, stopRule = stopRule2)
set.seed(seed)
regularCA <- spread(hab > 0, spreadProb = 0.23,
                  loci = (ncell(hab) - ncol(hab)) / 2 + c(4, -4),
                  directions = 8, id = TRUE)#, stopRule = stopRule2)

clearPlot()
Plot(circlish, regularCA)

#####
# complex stopRule
#####

initialLoci <- sample(seq_len(ncell(hab)), 2)
endSizes <- seq_along(initialLoci) * 200

# Can be a function of landscape, id, and/or any other named
# variable passed into spread
stopRule3 <- function(landscape, id, endSizes) sum(landscape) > endSizes[id]

twoCirclesDiffSize <- spread(hab, spreadProb = 1, loci = initialLoci, circle = TRUE,
                            directions = 8, id = TRUE, stopRule = stopRule3,
                            endSizes = endSizes, stopRuleBehavior = "excludePixel")

# or using named list of named elements:
twoCirclesDiffSize2 <- spread(hab, spreadProb = 1, loci = initialLoci, circle = TRUE,
                             directions = 8, id = TRUE, stopRule = stopRule3,
                             vars = list(endSizes = endSizes), stopRuleBehavior = "excludePixel")

identical(twoCirclesDiffSize, twoCirclesDiffSize2) ## TRUE

clearPlot()
Plot(twoCirclesDiffSize)

cirs <- getValues(twoCirclesDiffSize)
vals <- tapply(hab[twoCirclesDiffSize], cirs[cirs > 0], sum)

# Stop if sum of landscape is big or mean of quality is too small
quality <- raster(hab)
quality[] <- runif(ncell(quality), 0, 1)
stopRule4 <- function(landscape, quality, cells) {
  (sum(landscape) > 20) | (mean(quality[cells]) < 0.3)
}

twoCirclesDiffSize <- spread(hab, spreadProb = 1, loci = initialLoci, circle = TRUE,
                            directions = 8, id = TRUE, stopRule = stopRule4,
                            quality = quality, stopRuleBehavior = "excludePixel")

```

```
#####
# allowOverlap
#####
## TODO: fix error when allowOverlap = TRUE
# set.seed(3113)
# initialLoci <- as.integer(sample(1:ncell(hab), 10))
#
# # using "landscape", "id", and a variable passed in
# maxVal <- rep(500, length(initialLoci))
#
# # define stopRule
# stopRule2 <- function(landscape, id, maxVal) sum(landscape) > maxVal[id]
# circs <- spread(hab, spreadProb = 1, circle = TRUE, loci = initialLoci, stopRule = stopRule2,
#               id = TRUE, allowOverlap = TRUE, stopRuleBehavior = "includeRing",
#               maxVal = maxVal, returnIndices = TRUE)
# (vals <- tapply(hab[circs$indices], circs$id, sum))
# vals <= maxVal ## all TRUE
# overlapEvents <- raster(hab)
# overlapEvents[] <- 0
# toMap <- circs[, sum(id), by = indices]
# overlapEvents[toMap$indices] <- toMap$V1
#
# clearPlot()
# Plot(overlapEvents)

## Using alternative algorithm, not probabilistic diffusion
## Will give exactly correct sizes, yet still with variability
## within the spreading (i.e., cells with and without successes)
seed <- sample(1e6, 1)
set.seed(seed)
startCells <- startCells[1:4]
maxSizes <- rexp(length(startCells), rate = 1 / 500)
fires <- spread(hab, loci = startCells, 1, persistence = 0,
               neighProbs = c(0.5, 0.5, 0.5) / 1.5,
               mask = NULL, maxSize = maxSizes, directions = 8,
               iterations = 1e6, id = TRUE, plot.it = FALSE, exactSizes = TRUE);
all(table(fires[fires > 0][,]) == floor(maxSizes))

dev()
clearPlot()
Plot(fires, new = TRUE, cols = c("red", "yellow"), zero.color = "white")
Plot(hist(table(fires[][fires[] > 0])), title = "fire size distribution")

## Example with relativeSpreadProb ... i.e., a relative probability spreadProb
## (shown here because because spreadProb raster is not a probability).
## Here, we force the events to grow, choosing always 2 neighbours,
## according to the relative probabilities contained on hab layer.
##
## Note: `neighProbs = c(0,1)` forces each active pixel to move to 2 new pixels
## (`prob = 0` for 1 neighbour, `prob = 1` for 2 neighbours)
##
## Note: set hab3 to be very distinct probability differences, to detect spread
```

```

## differences
hab3 <- (hab < 20) * 200 + 1
seed <- 643503
set.seed(seed)
sam <- sample(which(hab3[] == 1), 1)
set.seed(seed)
events1 <- spread(hab3, spreadProb = hab3, loci = sam, directions = 8,
                 neighProbs = c(0, 1), maxSize = c(70), exactSizes = TRUE)

# Compare to absolute probability version
set.seed(seed)
events2 <- spread(hab3, id = TRUE, loci = sam, directions = 8,
                 neighProbs = c(0, 1), maxSize = c(70), exactSizes = TRUE)

clearPlot()
Plot(events1, new = TRUE, cols = c("red", "yellow"), zero.color = "white")
Plot(events2, new = TRUE, cols = c("red", "yellow"), zero.color = "white")
Plot(hist(table(events1[][events1[] > 0]), breaks = 30), title = "Event size distribution")
# Check that events1 resulted in higher hab3 pixels overall

# Compare outputs -- should be more high value hab pixels spread to in event1
# (randomness may prevent this in all cases)
hab3[events1[] > 0]
hab3[events2[] > 0]

sum(hab3[events1[] > 0]) >= sum(hab3[events2[] > 0]) ## should be usually TRUE

```

---

spread2

*Simulate a contagious spread process on a landscape, with data.table  
internals*

---

## Description

This can be used to simulate fires, seed dispersal, calculation of iterative, concentric, symmetric (currently) landscape values and many other things. Essentially, it starts from a collection of cells (start, called "events") and spreads to neighbours, according to the directions and spreadProb with modifications due to other arguments. **NOTE:** the spread function is similar, but sometimes slightly faster, but less robust, and more difficult to use iteratively.

## Usage

```

spread2(
  landscape,
  start = ncell(landscape)/2 - ncol(landscape)/2,
  spreadProb = 0.23,
  persistProb = NA_real_,
  asRaster = TRUE,
  maxSize,
  exactSize,

```

```

directions = 8L,
iterations = 1000000L,
returnDistances = FALSE,
returnDirections = FALSE,
returnFrom = FALSE,
maxRetriesPerID = 10,
spreadProbRel = NA_real_,
plot.it = FALSE,
circle = FALSE,
asymmetry = NA_real_,
asymmetryAngle = NA_real_,
allowOverlap = 0,
neighProbs = NA_real_,
skipChecks = FALSE
)

```

### Arguments

landscape	Required. A RasterLayer object. This defines the possible locations for spreading events to start and spread2 into. Required.
start	Required. Either a vector of pixel numbers to initiate spreading, or a data.table that is the output of a previous spread2. If a vector, they should be cell indices (pixels) on the landscape. If user has x and y coordinates, these can be converted with <code>cellFromXY</code> .
spreadProb	Numeric of length 1 or length <code>nCell(landscape)</code> or a RasterLayer that is the identical dimensions as landscape. If numeric of length 1, then this is the global (absolute) probability of spreading into each cell from a neighbour. If a numeric of length <code>nCell(landscape)</code> or a raster, then this must be the cell-specific (absolute) probability of a "receiving" potential cell. Default is 0.23. If relative probabilities are required, use <code>spreadProbRel</code> . If used together, then the relative probabilities will be re-scaled so that the mean relative probability of potential neighbours is equal to the mean of <code>spreadProb</code> of the potential neighbours.
persistProb	Numeric of length 1 or RasterLayer. If numeric of length 1, then this is the global (absolute) probability of cell continuing to burn per time step. If a raster, then this must be the cell-specific (absolute) probability of a fire persisting. Default is NA, which is the same as 0, i.e. a cell only burns for one time step.
asRaster	Logical, length 1. If TRUE, the function will return a Raster where raster non NA values indicate the cells that were "active", and the value is the initial starting pixel.
maxSize	Numeric. Maximum number of cells for a single or all events to be spread2. Recycled to match <code>start</code> length, if it is not as long as <code>start</code> . This will be overridden if <code>exactSize</code> also provided. See section on Breaking out of spread2 events.
exactSize	Numeric vector, length 1 or <code>length(start)</code> . Similar to <code>maxSize</code> , but these will be the exact final sizes of the events. i.e., the spread2 events will continue until they are <code>floor(exactSize)</code> . This will override <code>maxSize</code> if both provided. See Details.

directions	The number adjacent cells in which to look; default is 8 (Queen case). Can only be 4 or 8.
iterations	Number of iterations to spread2. Leaving this NULL allows the spread2 to continue until stops spreading itself (i.e., exhausts itself).
returnDistances	Logical. Should the function include a column with the individual cell distances from the locus where that event started. Default is FALSE. See Details.
returnDirections	Logical. Should the function include a column with the individual directions (in radians) from the locus where that event started. Default is FALSE.
returnFrom	Logical. Should the function return a column with the source, i.e, the lag 1 "from" pixel, for each iteration.
maxRetriesPerID	Only active if exactSize is used. This is the number of attempts that will be made per event ID, before abandoning, therefore completing the spread2 for that event with a size that is smaller than exactSize. Default 10 times.
spreadProbRel	Optional RasterLayer indicating a surface of relative probabilities useful when using neighProbs (which provides a mechanism for selecting a specific number of cells at each iteration). This indicates the relative probabilities for the selection of successful neighbours. spreadProb will still be evaluated <i>after</i> the relative probabilities and neighProbs has been evaluated, i.e., potential cells will be identified, then some could be rejected via spreadProb. If absolute spreadProb is not desired, <i>be sure to set</i> spreadProb = 1. Ignored if neighProbs is not provided.
plot.it	If TRUE, then plot the raster at every iteration, so one can watch the spread2 event grow.
circle	Logical. If TRUE, then outward spread2 will be by equidistant rings, rather than solely by adjacent cells (via directions arg.). Default is FALSE. Using circle = TRUE can be dramatically slower for large problems. Note, this will likely create unexpected results if spreadProb < 1.
asymmetry	A numeric or RasterLayer indicating the ratio of the asymmetry to be used. i.e., 1 is no asymmetry; 2 means that the angles in the direction of the asymmetryAngle are 2x the spreadProb of the angles opposite to the asymmetryAngle. Default is NA, indicating no asymmetry. See details. This is still experimental. Use with caution.
asymmetryAngle	A numeric or RasterLayer indicating the angle in degrees (0 is "up", as in North on a map), that describes which way the asymmetry is.
allowOverlap	numeric (logical will work for backwards compatibility). See details. Default is 0, i.e., no overlapping.
neighProbs	An optional numeric vector, whose sum is 1. It indicates the probabilities that an individual spread iteration will spread to 1, 2, ..., length(neighProbs) neighbours, respectively. If this is used (i.e., something other than NA), circle and returnDistances will not work currently.
skipChecks	Logical. If TRUE, the argument checking (i.e., assertions) will be skipped. This should likely only be used once it is clear that the function arguments are well

understood and function speed is of the primary importance. This is likely most useful in repeated iteration cases i.e., if this call is using the previous output from this same function.

## Details

There are 2 main underlying algorithms for active cells to "spread" to nearby cells (adjacent cells): `spreadProb` and `neighProb`. Using `spreadProb`, every "active" pixel will assess all neighbours (either 4 or 8, depending on directions), and will "activate" whichever neighbours successfully pass independent calls to `runif(1,0,1)<spreadProb`. The algorithm will iterate again and again, each time starting from the newly "activated" cells. Several built-in decisions are as follows. 1. no active cell can activate a cell that was already activated by the same event (i.e., "it won't go backwards"). 2. If `allowOverlap` is `FALSE`, then the previous rule will also apply, regardless of which "event" caused the pixels to be previously active.

This function can be interrupted before all active cells are exhausted if the `iterations` value is reached before there are no more active cells to `spread2` into. The interrupted output (a `data.table`) can be passed subsequently as an input to this same function (as `start`). This is intended to be used for situations where external events happen during a `spread2` event, or where one or more arguments to the `spread2` function change before a `spread2` event is completed. For example, if it is desired that the `spreadProb` change before a `spread2` event is completed because, for example, a fire is spreading, and a new set of conditions arise due to a change in weather.

`asymmetry` here is slightly different than in the `spread` function, so that it can deal with a `RasterLayer` of `asymmetryAngle`. Here, the `spreadProb` values of a given set of neighbours around each active pixel are adjusted to create `adjustedSpreadProb` which is calculated maintain the following two qualities:

$$\text{mean}(\text{spreadProb}) = \text{mean}(\text{adjustedSpreadProb})$$

and

$$\text{max}(\text{spreadProb})/\text{min}(\text{spreadProb}) = \text{asymmetry}$$

along the axis of `asymmetryAngle`. NOTE: this means that the 8 neighbours around an active cell may not fulfill the preceding equality if `asymmetryAngle` is not exactly one of the 8 angles of the 8 neighbours. This means that

$$\text{max}(\text{spreadProb})/\text{min}(\text{spreadProb})$$

will generally be less than `asymmetry`, for the 8 neighbours. The exact adjustment to the `spreadProb` is calculated with:

$$\text{angleQuality} < -(\cos(\text{angles} - \text{rad}(\text{asymmetryAngle})) + 1)/2$$

which is multiplied to get an angle-adjusted `spreadProb`:

$$\text{spreadProbAdj} < -\text{actualSpreadProb} * \text{angleQuality}$$

which is then rescaled:

$$\text{adjustedSpreadProb} = (\text{spreadProbAdj} - \text{min}(\text{spreadProbAdj})) * \text{par2} + \text{par1}$$

, where `par1` and `par2` are parameters calculated internally to make the 2 conditions above true.

If `exactSize` or `maxSize` are used, then spreading will continue and stop before or at `maxSize` or at `exactSize`. If `iterations` is specified, then the function will end, and the returned `data.table` will still may (if `maxSize`) or will (if `exactSize`) have at least one active cell per event that did not already achieve `maxSize` or `exactSize`. This will be very useful to build new, customized higher-level wrapper functions that iteratively call `spread2`.

### Value

Either a `data.table` (`asRaster=FALSE`) or a `RasterLayer` (`asRaster=TRUE`, the default). The `data.table` will have one attribute named `spreadState`, which is a list containing a `data.table` of current cluster-level information about the spread events. If `asRaster=TRUE`, then the `data.table` (with its `spreadState` attribute) will be attached to the `Raster` as an attribute named `pixel` as it provides pixel-level information about the spread events.

The `RasterLayer` represents every cell in which a successful `spread2` event occurred. For the case of, say, a fire this would represent every cell that burned. If `allowOverlap` is `TRUE`, the return will always be a `data.table`.

If `asRaster` is `FALSE`, then this function returns a `data.table` with 3 (or 4 if `returnFrom` is `TRUE`) columns:

<code>initialPixels</code>	the initial cell number of that particular <code>spread2</code> event.
<code>pixels</code>	The cell indices of cells that have been touched by the <code>spread2</code> algorithm.
<code>state</code>	a logical indicating whether the cell is active (i.e., could still be a source for spreading) or not (no spreading)
<code>from</code>	The pixel indices that were the immediately preceding "source" for each <code>pixels</code> , i.e., the lag 1 pixels. Only

The attribute saved with the name "spreadState" (e.g., `attr(output, "spreadState")`) includes a `data.table` with columns:

<code>id</code>	An arbitrary code, from 1 to <code>length(start)</code> for each "event".
<code>initialPixels</code>	the initial cell number of that particular <code>spread2</code> event.
<code>numRetries</code>	The number of re-starts the event did because it got stuck (normally only because <code>exactSize</code> was used and
<code>maxSize</code>	The number of pixels that were provided as inputs via <code>maxSize</code> or <code>exactSize</code> .
<code>size</code>	The current size, in pixels, of each event.

and several other objects that provide significant speed ups in iterative calls to `spread2`. If the user runs `spread2` iteratively, there will likely be significant speed gains if the `data.table` passed in to `start` should have the attribute attached, or re-attached if it was lost, e.g., via `setattr(outInput, "spreadState", attr(outInput, "spreadState"))` where `out` is the returned `data.table` from the previous call to `spread2`, and `outInput` is the modified `data.table`. Currently, the modified `data.table` **must** have the same order as `out`.

### Breaking out of spread2 events

There are 3 ways for the `spread2` to "stop" spreading. Here, each "event" is defined as all cells that are spawned from each unique `start` location. The ways outlined below are all acting at all times, i.e., they are not mutually exclusive. Therefore, it is the user's responsibility to make sure the different rules are interacting with each other correctly.



spreadProb	Probabilistically, if spreadProb is low enough, active spreading events will stop. In practice, this number generally
maxSize	This is the number of cells that are "successfully" turned on during a spreading event. spreadProb will still be
exactSize	This is the number of cells that are "successfully" turned on during a spreading event. This will override an even
iterations	This is a hard cap on the number of internal iterations to complete before returning the current state of the system

### Building custom spreading events

This function can be used iteratively, with relatively little overhead compared to using it non-iteratively. In general, this function can be called with arguments set as user needs, and with specifying iterations = 1 (say). This means that the function will spread outwards 1 iteration, then stop. The returned object will be a `data.table` or `RasterLayer` that can be passed immediately back as the start argument into a subsequent call to `spread2`. This means that every argument can be updated at each iteration.

When using this function iteratively, there are several things to keep in mind. The output will likely be sorted differently than the input (i.e., the order of start, if a vector, may not be the same order as that returned). This means that when passing the same object back into the next iteration of the function call, `maxSize` or `exactSize` may not be in the same order. To get the same order, the easiest thing to do is sort the initial start objects by their pixel location, increasing. Then, of course, sorting any vectorized arguments (e.g., `maxSize`) accordingly.

**NOTE:** the `data.table` or `RasterLayer` should not use be altered when passed back into `spread2`.

### allowOverlap

If 1 (or TRUE), then individual events can overlap with one another, i.e., allow overlap between events. If 2 (or NA), then each pixel is essentially independent, allowing overlap between and within events. This likely requires a user to intervene as it is possible to spread back onto itself. If 3 (did not exist previously), individual events can overlap, and there can be overlap within an event, but only within an iteration, i.e., once an iteration is finished, and a pixel was activated, then the spreading will not return onto these pixels. If 0 (or FALSE), then once a pixel is activated, it cannot be re-activated, within or between event. This allows events to not interfere with one another i.e., they do not interact (this is slower than if `allowOverlap = FALSE`). Default is 0. In the case of 2 or 3, this would be, perhaps, useful for dispersal of, say, insect swarms.

### Note

`exactSize` may not be achieved if there aren't enough cells in the map. Also, `exactSize` may not be achieved because the active cells are "stuck", i.e., they have no unactivated cells to move to; or the `spreadProb` is low. In the latter two cases, the algorithm will retry again, but it will only re-try from the last iterations active cells. The algorithm will only retry 10 times before quitting. Currently, there will also be an attempt to "jump" up to four cells away from the active cells to try to continue spreading.

A common way to use this function is to build wrappers around this, followed by iterative calls in a while loop. See example.

### Author(s)

Eliot McIntire and Steve Cumming

**See Also**

[spread](#) for a different implementation of the same algorithm. `spread` is less robust but it is often slightly faster.

**Examples**

```
library(data.table)
library(raster)
library(quickPlot)

data.table::setDTthreads(1)

a <- raster(extent(0, 10, 0, 10), res = 1)
sams <- sort(sample(ncell(a), 3))

# Simple use -- similar to spread(...)
out <- spread2(a, start = sams, 0.225)
if (interactive()) {
  clearPlot()
  Plot(out)
}

# Use maxSize -- this gives an upper limit
maxSizes <- sort(sample(1:10, size = length(sams)))
out <- spread2(a, start = sams, 0.225, maxSize = maxSizes, asRaster = FALSE)
# check TRUE using data.table .N
out[, .N, by = "initialPixels"]$N <= maxSizes

# Use exactSize -- gives an exact size, if there is enough space on the Raster
exactSizes <- maxSizes
out <- spread2(a, start = sams, spreadProb = 0.225,
              exactSize = exactSizes, asRaster = FALSE)
out[, .N, by = "initialPixels"]$N == maxSizes # should be TRUE TRUE TRUE

# Use exactSize -- but where it can't be achieved
exactSizes <- sort(sample(100:110, size = length(sams)))
out <- spread2(a, start = sams, 1, exactSize = exactSizes)

# Iterative calling -- create a function with a high escape probability
spreadWithEscape <- function(ras, start, escapeProb, spreadProb) {
  out <- spread2(ras, start = sams, spreadProb = escapeProb, asRaster = FALSE)
  while (any(out$state == "sourceActive")) {
    # pass in previous output as start
    out <- spread2(ras, start = out, spreadProb = spreadProb,
                  asRaster = FALSE, skipChecks = TRUE) # skipChecks for speed
  }
  out
}

set.seed(421)
out1 <- spreadWithEscape(a, sams, escapeProb = 0.25, spreadProb = 0.225)
set.seed(421)
```

```

out2 <- spread2(a, sams, 0.225, asRaster = FALSE)
# The one with high escape probability is larger (most of the time)
NROW(out1) > NROW(out2)

## Use neighProbs, with a spreadProb that is a RasterLayer
# Create a raster of different values, which will be the relative probabilities
# i.e., they are rescaled to relative probabilities within the 8 neighbour choices.
# The neighProbs below means 70% of the time, 1 neighbour will be chosen,
# 30% of the time 2 neighbours.
# The cells with spreadProb of 5 are 5 times more likely than cells with 1 to be chosen,
# when they are both within the 8 neighbours
sp <- raster(extent(0, 3, 0, 3), res = 1, vals = 1:9) #small raster, simple values
# Check neighProbs worked
out <- list()

# enough replicates to see stabilized probabilities
for (i in 1:100) {
  out[[i]] <- spread2(sp, spreadProbRel = sp, spreadProb = 1,
    start = 5, iterations = 1,
    neighProbs = c(1), asRaster = FALSE)
}
out <- data.table::rbindlist(out)[pixels != 5] # remove starting cell
table(sp[out$pixels])
# should be non-significant -- note no 5 because that was the starting cell
# This tests whether the null model is true ... there should be proportions
# equivalent to 1:2:3:4:6:7:8:9 ... i.e., cell 9 should have 9x as many events
# spread to it as cell 1. This comes from sp object above which is providing
# the relative spread probabilities
keep <- c(1:4, 6:9)
chisq.test(keep, unname(tabulate(sp[out$pixels], 9)[keep]), simulate.p.value = TRUE)

## Example showing asymmetry
sams <- ncell(a) / 4 - ncol(a) / 4 * 3
circs <- spread2(a, spreadProb = 0.213, start = sams,
  asymmetry = 2, asymmetryAngle = 135,
  asRaster = TRUE)

## ADVANCED: Estimate spreadProb when using asymmetry, such that the expected
## event size is the same as without using asymmetry
## Not run:
if (interactive()) {
  ras <- raster(a)
  ras[] <- 1

  n <- 100
  sizes <- integer(n)
  for (i in 1:n) {
    circs <- spread2(ras, spreadProb = 0.225,
      start = round(ncell(ras) / 4 - ncol(ras) / 4 * 3),
      asRaster = FALSE)
    sizes[i] <- circs[, .N]
  }
  goalSize <- mean(sizes)
}

```

```

library(parallel)
library(DEoptim)
cl <- makeCluster(pmin(10, detectCores() - 2)) # need 10 cores for 10 populations in DEoptim
parallel::clusterEvalQ(cl, {
  library(SpaDES.tools)
  library(raster)
  library(fpCompare)
})

objFn <- function(sp, n = 20, ras, goalSize) {
  sizes <- integer(n)
  for (i in 1:n) {
    circs <- spread2(ras, spreadProb = sp, start = ncell(ras) / 4 - ncol(ras) / 4 * 3,
                    asymmetry = 2, asymmetryAngle = 135,
                    asRaster = FALSE)
    sizes[i] <- circs[, .N]
  }
  abs(mean(sizes) - goalSize)
}
aa <- DEoptim(objFn, lower = 0.2, upper = 0.23,
             control = DEoptim.control(cluster = cl, NP = 10, VTR = 0.02,
                                       initialpop = as.matrix(rnorm(10, 0.213, 0.001))),
             ras = a, goalSize = goalSize)

# The value of spreadProb that will give the same expected event sizes to spreadProb = 0.225 is:
sp <- aa$optim$bestmem
circs <- spread2(ras, spreadProb = sp, start = ncell(ras) / 4 - ncol(ras) / 4 * 3,
                asymmetry = 2, asymmetryAngle = 135, asRaster = FALSE)

stopCluster(cl)
}

## End(Not run)

```

---

spread3

*An alternative spread function, conceived for insects*

---

### Description

This is built with [spread2](#) and is still experimental. This one differs from other attempts in that it treats the advection and dispersal as mathematical vectors that are added together. They are "rounded" to pixel centres.

### Usage

```

spread3(
  start,
  rasQuality,
  rasAbundance,

```

```

    advectionDir,
    advectionMag,
    meanDist,
    plot.it = 2,
    minNumAgents = 50,
    verbose = getOption("LandR.verbose", 0),
    saveStack = NULL
)

```

### Arguments

start	Raster indices from which to initiate dispersal
rasQuality	A raster with habitat quality. Currently, must be scaled from 0 to 1, i.e., a probability of "settling"
rasAbundance	A raster where each pixel represents the number of "agents" or pseudo-agents contained. This number of agents, will be spread horizontally, and distributed from each pixel that contains a non-zero non NA value.
advectionDir	A single number or RasterLayer in degrees from North = 0 (though it will use radians if all values are $\text{abs}(\text{advectionDir}) > 2 * \pi$ ). This indicates the direction of advective forcing (i.e., wind).
advectionMag	A single number or RasterLayer in distance units of the rasQuality, e.g., meters, indicating the relative forcing that will occur. It is imposed on the total event, i.e., if the meanDist is 10000, and advectionMag is 5000, then the expected distance (i.e., 63% of agents) will have settled by 15000 map units.
meanDist	A single number indicating the mean distance parameter in map units (not pixels), for a negative exponential distribution dispersal kernel (e.g., dexp). This will mean that 63 settled at this meanDist (still experimental)
plot.it	Numeric. With increasing numbers above 0, there will be plots produced during iterations. Currently, only 0, 1, or 2+ are distinct.
minNumAgents	Single numeric indicating the minimum number of agents to consider all dispersing finished. Default is 50
verbose	Numeric. With increasing numbers above 0, there will be more messages produced. Currently, only 0, 1, or 2+ are distinct.
saveStack	If provided as a character string, it will save each iteration as part of a rasterStack to disk upon exit.

### Value

A data.table with all information used during the spreading

### Examples

```

## these tests are fairly heavy, so don't run during automated tests
if (interactive()) {
#####
# Simple case, no variation in rasQuality, numeric advectionDir and advectionMag
#####
}

```

```

library(raster)
library(quickPlot)
maxDim <- 10000
ras <- raster::raster(extent(c(0, maxDim, 0, maxDim)), res = 100, vals = 0)
rasQuality <- raster(ras)
rasQuality[] <- 1
rasAbundance <- raster(rasQuality)
rasAbundance[] <- 0
# startPixel <- middlePixel(rasAbundance)
startPixel <- sample(seq(ncell(rasAbundance)), 30)
rasAbundance[startPixel] <- 1000
advectionDir <- 70
advectionMag <- 4 * res(rasAbundance)[1]
meanDist <- 2600

if (interactive()) {
  # Test the dispersal kernel -- create a function
  plotDispersalKernel <- function(out, meanAdvectionMag) {
    out[, disGroup := round(distance / 100) * 100]
    freqs <- out[, .N, by = "disGroup"]
    freqs[, `:=`(cumSum = cumsum(N), N = N)]
    Plot(freqs$disGroup, freqs$cumSum, addTo = "CumulativeNumberSettled",
         title = "Cumulative Number Settled") # can plot the distance X number
    abline(v = meanAdvectionMag + meanDist)
    newTitle <- "Number Settled By Distance"
    Plot(freqs$disGroup, freqs$N, addTo = gsub(" ", "", newTitle),
         title = newTitle) # can plot the distance X number
    abline(v = meanAdvectionMag + meanDist)
    # should be 0.63:
    freqs[disGroup == meanAdvectionMag + meanDist, cumSum] / tail(freqs,1)[, cumSum]
    mtext(side = 3, paste("Average habitat quality: ",
                        round(mean(rasQuality[], na.rm = TRUE), 2)),
          outer = TRUE, line = -2, cex = 2)
  }
  dev() # don't use Rstudio windows, which is very slow
  clearPlot()
  out <- spread3(rasAbundance = rasAbundance,
                rasQuality = rasQuality,
                advectionDir = advectionDir,
                advectionMag = advectionMag,
                meanDist = meanDist, verbose = 2,
                plot.it = 2)

  plotDispersalKernel(out, advectionMag)
}

#####
### The case of variable quality raster
#####
if (require(sf) && require(fasterize)) {
  library(SpaDES.tools) # for gaussMap
  a <- randomStudyArea(size = 1e9)
  ras <- raster(extent(a), res = 100)

```

```

mask <- fasterize(st_as_sf(a), ras) # faster than raster::rasterize
rasQuality <- gaussMap(ras)
crs(rasQuality) <- crs(a)
rasQuality[is.na(mask)] <- NA
# rescale so min is 0.75 and max is 1
rasQuality[] <- rasQuality[] / (maxValue(rasQuality) * 4 ) + 1/4
rasAbundance <- raster(rasQuality)
rasAbundance[] <- 0
startPixel <- sample(seq(ncell(rasAbundance)), 300)
rasAbundance[startPixel] <- 1000
advectionDir <- 75
advectionMag <- 4 * res(rasAbundance)[1]
meanDist <- 2600
clearPlot()
out <- spread3(rasAbundance = rasAbundance,
               rasQuality = rasQuality,
               advectionDir = advectionDir,
               advectionMag = advectionMag,
               meanDist = meanDist, verbose = 2,
               plot.it = 1)
if (interactive()) {
  plotDispersalKernel(out, advectionMag)
}
}

#####
### The case of variable quality raster, raster for advectionDir & advectionMag
#####
library(raster)
library(quickPlot)
library(SpaDES.tools)
maxDim <- 10000
ras <- raster(extent(c(0, maxDim, 0, maxDim)), res = 100, vals = 0)
rasQuality <- raster(ras)
rasQuality[] <- 1
rasAbundance <- raster(rasQuality)
rasAbundance[] <- NA
# startPixel <- middlePixel(rasAbundance)
startPixel <- sample(seq(ncell(rasAbundance)), 25)
rasAbundance[startPixel] <- 1000

# raster for advectionDir
advectionDir <- gaussMap(ras)
crs(advectionDir) <- crs(rasQuality)
# rescale so min is 0.75 and max is 1
advectionDir[] <- advectionDir[] / (maxValue(advectionDir)) * 180

# raster for advectionMag
advectionMag <- gaussMap(ras)
crs(advectionMag) <- crs(rasQuality)
# rescale so min is 0.75 and max is 1
advectionMag[] <- advectionMag[] / (maxValue(advectionMag)) * 600

```

```

out <- spread3(rasAbundance = rasAbundance,
              rasQuality = rasQuality,
              advectionDir = advectionDir,
              advectionMag = advectionMag,
              meanDist = meanDist, verbose = 2,
              plot.it = 1)

if (interactive()) {
  dev() # don't use Rstudio windows, which is very slow
  clearPlot()
  Plot(advectionDir, title = "Wind direction", cols = "Reds")
  Plot(advectionMag, title = "Wind speed", cols = "Blues")
  plotDispersalKernel(out, mean(advectionMag[]))
  Plot(rasAbundance, addTo = "rasAbundance", cols = "black", title = "")
}

#####
# save iterations to a stack to make animated GIF
#####
tmpStack <- raster::rasterTmpFile("stackToAnimate")
out <- spread3(rasAbundance = rasAbundance,
              rasQuality = rasQuality,
              advectionDir = advectionDir,
              advectionMag = advectionMag,
              meanDist = 2600, verbose = 2,
              plot.it = 0, saveStack = tmpStack)

## This animates the series of images into an animated GIF
if (require(animation, quietly = TRUE)) {
  out2 <- raster::stack(tmpStack)
  gifName <- file.path(tempdir(), "animation.gif")
  saveGIF(interval = 0.1, movie.name = gifName, expr = {
    for (i in seq(numLayers(out2))) plot(out2[[i]])
  })
}
}

```

---

testEquivalentMetadata

*Test that metadata of 2 or more objects is the same*

---

### Description

Currently, only Raster class has a useful method. Defaults to `all(sapply(list(...)[-1], function(x) identical(list(...)[1], x)))`

### Usage

```
testEquivalentMetadata(...)
```



**Arguments**

... 2 or more of the same type of object to test for equivalent metadata

---

transitions

SELES - *Transitioning to next time step*

---

**Description**

Describes the probability of an agent successfully persisting until next time step. THIS IS NOT YET FULLY IMPLEMENTED.

A SELES-like function to maintain conceptual backwards compatibility with that simulation tool. This is intended to ease transitions from **SELES**.

You must know how to use SELES for these to be useful.

**Usage**

transitions(p, agent)

**Arguments**

p realized probability of persisting (i.e., either 0 or 1).

agent SpatialPoints\* object.

**Value**

Returns new SpatialPoints\* object with potentially fewer agents.

**Author(s)**

Eliot McIntire

---

wrap

*Wrap coordinates or pixels in a torus-like fashion*

---

**Description**

Generally useful for model development purposes.

**Usage**

```

wrap(X, bounds, withHeading)

## S4 method for signature 'matrix,Extent,missing'
wrap(X, bounds)

## S4 method for signature 'SpatialPoints,ANY,missing'
wrap(X, bounds)

## S4 method for signature 'matrix,Raster,missing'
wrap(X, bounds)

## S4 method for signature 'matrix,Raster,missing'
wrap(X, bounds)

## S4 method for signature 'matrix,matrix,missing'
wrap(X, bounds)

## S4 method for signature 'SpatialPointsDataFrame,Extent,logical'
wrap(X, bounds, withHeading)

## S4 method for signature 'SpatialPointsDataFrame,Raster,logical'
wrap(X, bounds, withHeading)

## S4 method for signature 'SpatialPointsDataFrame,matrix,logical'
wrap(X, bounds, withHeading)

```

**Arguments**

X	A <code>SpatialPoints*</code> object, or matrix of coordinates.
bounds	Either a <code>Raster*</code> , <code>Extent</code> , or <code>bbox</code> object defining bounds to wrap around.
withHeading	logical. If <code>TRUE</code> , the previous points must be wrapped also so that the subsequent heading calculation will work. Default <code>FALSE</code> . See details.

**Details**

If `withHeading` used, then `X` must be a `SpatialPointsDataFrame` that contains two columns, `x1` and `y1`, with the immediately previous agent locations.

**Value**

Object of the same class as `X`, but with coordinates updated to reflect the wrapping.

**Author(s)**

Eliot McIntire

**Examples**

```
library(raster)
library(quickPlot)

xrange <- yrange <- c(-50, 50)
hab <- raster(extent(c(xrange, yrange)))
hab[] <- 0

# initialize agents
N <- 10

# previous points
x1 <- rep(0, N)
y1 <- rep(0, N)
# initial points
starts <- cbind(x = stats::runif(N, xrange[1], xrange[2]),
                y = stats::runif(N, yrange[1], yrange[2]))

# create the agent object
agent <- SpatialPointsDataFrame(coords = starts, data = data.frame(x1, y1))

ln <- rlnorm(N, 1, 0.02) # log normal step length
sd <- 30 # could be specified globally in params

if (interactive()) {
  clearPlot()
  Plot(hab, zero.color = "white", axes = "L")
}
for (i in 1:10) {
  agent <- crw(agent = agent, extent = extent(hab), stepLength = ln,
              stddev = sd, lonlat = FALSE, torus = TRUE)
  if (interactive()) Plot(agent, addTo = "hab", axes = TRUE)
}
```

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