Package 'nhSDM'

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Title Tools for Natural Heritage SDMs			
Date 2018-03-14			
Version 0.0.1-1Description Miscellaneous tools for working with Natural Heritage data in Species Distribution Modelling (SDM) workflows.			
		Depends R (>= $3.4.1$)	
Imports methods, sf, raster, sp Suggests dplyr, lwgeom License GPL-3 Encoding UTF-8 LazyData true RoxygenNote 6.0.1			
		nh_burn nh_group	ited:
		nh_best	Extract areas with best (highest value) as polygons from an SDM pre- diction raster, with optional feature mask
		Description If spf is given, areas or gons.	of cells intersecting these features will not be included in the returned poly

nh_best(spf = NULL, rast, top.percent = NULL, min.size = NULL,

num.patches = NULL, rank.by = "area")

Usage

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Arguments

spf	input spatial features (sp or sf spatial object); will be erased from output
rast	input raster model output with continuous values
top.percent	numeric; percent (e.g.; $0.01 = 0.01\%$) of highest cell values in raster to extract
min.size	numeric; optional minimal area of an extracted polygon
num.patches	numeric; optional number of patches to return, using rank.by criteria
rank.by	character; used for ranking patches when num.patches is not null; either 'area'

(default) or 'value'

Details

top.percent and min.size will both be derived from spf if they are null and spf is given. In this case, top.percent will be set to be equal to spf's cell coverage (prevalence relative to non-NA areas in raster). min. size will be set to the area of the smallest feature in spf. If spf are line or point features, this will be zero.

rank.by only subsets the outputs if num.patches is set. rank.by must be set to "value" to return the mean prediction value of the patch.

For better performance, crop the raster prior to running, and/or use a very low top.percent value, to return a smaller proportion of the cells as polygons.

Value

```
sp or sf object (polygons)
```

Author(s)

David Bucklin

Examples

```
## Not run:
spf <- st_read("ambymabe/polygon_data/ambymabe.shp")</pre>
rast <- raster("ambymabe/grids/ambymabe_20171018_130837.tif")</pre>
best_polys <- nh_burn(spf, rast)</pre>
## End(Not run)
```

nh_burn Add areas represented by features ('burn-in') to a binary raster SDM output

Description

Takes spatial features, a continuous raster (values between 0 and 1), and returns a classified binary (0/1) raster. Areas greater than orig. thresh are set to 1, all others 0. Areas intersecting spf features are assigned a value of 1 in the returned classified raster. See details for default calculation of orig. thresh and usage of buffer.

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Usage

```
nh_burn(spf, rast, orig.thresh = NULL, buffer = 0, ...)
```

Arguments

spf input spatial features (sp or sf spatial object)
rast input raster model output (values between 0 and 1)
orig.thresh numeric between 0 and 1; threshold value to apply to raster
buffer numeric; spatial buffer around spf to include in burn-in
additional parameters to raster::writeRaster

Details

A minimum cell value (min.cell) across all spf is calculated. If orig. thresh is NULL, the min.cell value will be used as a threshold for the full raster.

If buffer is greater than 0 orig. thresh is not NULL, orig. thresh is used as the threshold for the full raster. Additionally, cells within buffer distance of spf which are greater than min.cell are also set to 1. If a given orig. thresh is lower than the calculated min.cell value, buffer distance will have no impact on the output.

Value

RasterLayer

Author(s)

David Bucklin

Examples

```
## Not run:
spf <- st_read("ambymabe/polygon_data/ambymabe.shp")
rast <- raster("ambymabe/grids/ambymabe_20171018_130837.tif")
class_burn <- nh_burn(spf, rast, 0.75, 250, filename = "model_classified.tif", datatype = "INT2U")
## End(Not run)</pre>
```

nh_group

Group spatial features using a defined separation distance

Description

Input features are allocated into new groups depending on whether they lie within the separation distance of another input feature.

Usage

```
nh_group(spf, sep.dist = 0, union = FALSE)
```

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Arguments

spf input spatial features (sp or sf spatial object)

sep.dist separation distance with which to define groups (see description)

union whether to union output groups into multi-features

Details

The grouping is done by input ID, so multi-features are considered one input feature (they will not be be ungrouped).

The separation distance sep.dist is numeric and in the units of spf's coordinate system, unless the coordinate system uses latitude/longitude as the unit (e.g. WGS 84). In these cases, geodesic distances will be used and sep.dist should be specified in meters.

A column 'group' will be added to the output features. Specifying union = TRUE will output one (multi)feature per group, meaning original attributes will be discarded - only 'group' and 'count' (the number of original features in the group) will be returned. This feature requires the package dplyr to be installed.

Value

```
sp or sf object (same as input)
```

Author(s)

David Bucklin

Examples

```
## Not run:
spf <- rgdal::readOGR("ambymabe/polygon_data", "ambymabe")
spg <- nh_group(spf, 1000)
## End(Not run)</pre>
```

nh_sample

Create points in features in reference raster cells

Description

For each spatial feature, a given number (num_samps) of points are created in cells that the feature intersects.

Usage

```
nh_sample(spf, rast, num.samps = NULL, replace = FALSE, force.min = FALSE)
```

Arguments

spf input spatial features (sp or sf spatial object)
rast raster dataset with extent overlapping spf

num. samps number of samples to create in each feature (see details)

replace whether to sample with or without replacement

force.min whether to force num. samps points in features, even if they are duplicates

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Details

num. samps can be a a proportion (a decimal value < 1), single integer, or vector of integers equal to length of spf indicated the number of samples to take from each feature. If left NULL, num. samps will be set to the number of cells [n] intersecting the feature. If a proportion is given (e.g., 0.5), than [n * num.samps] will be returned. If a single integer is given, num. samps points will be sampled in each feature.

When replace = FALSE and force.min = FALSE (defaults), each cell can only contain one point (across each feature, and the entire returned set of points). In this case, when the number of samples points to create exceeds the number of unique cells intersected by a given feature, the number of samples for that feature equals the number of cells. If replace = TRUE, sampling is done with replacement and duplicates may be taken. The special case replace = FALSE and force.min = TRUE will always return num.samps per feature. It only produces duplicates if num.samps exceeds the number of cells intersecting the feature, in which case it will replicate the samples until num.samps is reached.

If CRS do not match, the features will be transformed to the CRS of the raster.

A column 'feat.id' is added to the output point features to indicate the row number of the feature that the point was generated within.

Value

```
sp or sf object (points)
```

Author(s)

David Bucklin

Examples

```
## Not run:
r<-raster::raster("AnnMnTemp.tif")
spf <- rgdal::readOGR("ambymabe/polygon_data", "ambymabe")
# can also use sf: spf <- sf::st_read("ambymabe/polygon_data/ambymabe.shp")
spf.samps <- nh_sample(spf, r, num.samps)
## End(Not run)</pre>
```

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