

# Package ‘bamm’

December 20, 2022

**Type** Package

**Title** Species Distribution Models as a Function of Biotic, Abiotic and Movement Factors (BAM)

**Version** 0.4.3

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**URL** <https://luismurao.github.io/bamm/>

**BugReports** <https://github.com/luismurao/bamm/issues>

**Description** Species Distribution Modeling (SDM) is a practical methodology that aims to estimate the area of distribution of a species. However, most of the work has focused on estimating static expressions of the correlation between environmental variables. The outputs of correlative species distribution models can be interpreted as maps of the suitable environment for a species but not generally as maps of its actual distribution. Soberón and Peterson (2005) <[doi:10.17161/bi.v2i0.4](https://doi.org/10.17161/bi.v2i0.4)> presented the BAM scheme, a heuristic framework that states that the occupied area of a species occurs on sites that have been accessible through dispersal (M) and have both favorable biotic (B) and abiotic conditions (A). The 'bamm' package implements classes and functions to operate on each element of the BAM and by using a cellular automata model where the occupied area of a species at time t is estimated by the multiplication of three binary matrices: one matrix represents movements (M), another abiotic -niche- tolerances (A), and a third, biotic interactions (B). The theoretical background of the package can be found in Soberón and Osorio-Olvera (2022) <[arXiv:2212.06308](https://arxiv.org/abs/2212.06308)>.

**SystemRequirements** C++11, GDAL (>= 2.2.3): gdal-bin (deb), libgdal-dev (deb) or gdal-devel (rpm), GEOS (>= 3.4.0), PROJ (>= 4.9.3): libproj-dev (deb), sqlite3, ImageMagick++: imagemagick (deb), libmagic-dev (deb), libmagick++-dev (deb) or ImageMagick-c++-devel (rpm) ImageMagick (<http://imagemagick.org>) or GraphicsMagick (<http://www.graphicsmagick.org>) or LyX (<http://www.lyx.org>) for saveGIF(); (PDF)LaTeX for saveLatex(); SWF Tools

(<http://swftools.org>) for saveSWF(); FFmpeg (<http://ffmpeg.org>) or avconv (<https://libav.org/avconv.html>) for saveVideo()

**NeedsCompilation** yes

**License** GPL (>= 3)

**Encoding** UTF-8

**Depends** R (>= 3.5.0)

**Imports** raster (>= 3.4-13), Matrix (>= 1.2.14), RSpectra (>= 0.13.1), methods (>= 3.3), magrittr (>= 1.2), dplyr (>= 0.8.0), purrr (>= 0.2), igraph (>= 1.2), leaflet (>= 2.0), Rdpack (>= 0.11.0), animation (>= 2.3), future (>= 1.18.0), furrr (>= 0.1.0), Rcpp (>= 0.12.18), sp(>= 1.3.0)

**RdMacros** Rdpack

**RoxygenNote** 7.2.2

**Suggests** knitr, crosstalk, plotly, rmarkdown, testthat (>= 3.0.0), covr

**VignetteBuilder** knitr

**LinkingTo** Rcpp, RcppArmadillo

**Config/testthat.edition** 3

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**Repository** CRAN

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.nonzero	<i>Helper function to compute the elements in g0 that have no zero values. The function is taken from the Ringo package</i>
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## Description

Helper function to compute the elements in g0 that have no zero values. The function is taken from the Ringo package

## Usage

```
.nonzero(x)
```

## Arguments

x	A matrix of class "dgCMatrix"
---	-------------------------------

**adj\_mat***adj\_mat: Function to compute the adjacency matrix of an area.***Description**

Creates an adjacency matrix of an area of interest. This could be the accessible area (M) of a species or any geographic region of interest.

**Usage**

```
adj_mat(modelsparse, ngbs = 1, eigen_sys = FALSE, which_eigs = 1)
```

**Arguments**

<code>modelsparse</code>	A <code>setA</code> object returned by the function <a href="#">model2sparse</a> .
<code>ngbs</code>	Numeric. Number of neighbors (see details).
<code>eigen_sys</code>	Logical. If TRUE the eigen analyses of the adjacency matrix will be returned.
<code>which_eigs</code>	Numeric. Which eigen value and eigen vector will be returned.

**Details**

The model is a raster object of the area where the dispersal process will occur. The number of neighbors depends on the dispersal abilities of the species and the spatial resolution of the niche model; for example, a species's with big dispersal abilities will move throughout more than 1 km<sup>2</sup> per day, so the idea is to give an approximate number of moving neighbors (pixels) per unit of time. For more information about see adjacency matrices in the context of the theory of area of distribution (Soberón and Osorio-Olvera, 2022).

**Value**

Returns an object of class `setM` with 7 slots. The first contains the adjacency matrix. A n x n sparse matrix (n=number of non-NA cells of the niche model) where connected cells are represented by 1. The second slot has the adjacency list. It is a list of matrices with four columns (FromRasCell -from cell ID of the raster-, -to cell ID of the raster-, -from non-NA cell-, -to non-NA cell-). Other slots contain information about initial coordinates where dispersal occurs (`initial_points`), number of cells used to define the neighborhood (`ngbs`), non-NA coordinates (`coordinates`), and a matrix of eigen vectors (`eigen_vec`).

**Author(s)**

Luis Osorio-Olvera & Jorge Soberón

**References**

Soberón J, Osorio-Olvera L (2022). “A Dynamic Theory of the Area of Distribution.” [doi:10.48550/ARXIV.2212.06308](#), <https://arxiv.org/abs/2212.06308..>

## Examples

```

x_coord <- c(-106.5699, -111.3737, -113.9332,
             -110.8913, -106.4262, -106.5699)
y_coord <- c(16.62661, 17.72373, 19.87618,
             22.50763, 21.37728, 16.62661)
xy <- cbind(x_coord, y_coord)
p <- sp::Polygon(xy)
ps <- sp::Polygons(list(p),1)
sps <- sp::SpatialPolygons(list(ps))
mx_grid <- bammm::shape2Grid(sps, resolution = 0.25, ones = TRUE)
mx_sparse <- bammm::model2sparse(model=mx_grid, threshold = 0.1)
adj_mx <- bammm::adj_mat(modelsparse=mx_sparse,
                           nbgs=1, eigen_sys=TRUE, which_eigs=1)
print(adj_mx)
mx_grid_eigen <- mx_grid
mx_grid_eigen[mx_sparse@cellIDs] <- adj_mx@eigen_vec
raster::plot(mx_grid_eigen)

```

bam-class

*Class bam digram*

## Description

Class `bam` digram

## Value

An object of class `bam`

## Slots

`sdm_sim` A list of sparse vectors representing the area occupied  
`palatable_matrices` A list of sparse vectors representing palatable sites.  
`sim_steps` Number of simulation steps by the species

## Author(s)

Luis Osorio-Olvera & Jorge Soberón

**bam\_clusters***bam\_clusters: Function to estimate the connectivity of suitable areas*

## Description

Function to estimate the connectivity of suitable areas given an adjacency matrix.

## Usage

```
bam_clusters(model, ngbs = 1, plot_model = FALSE)
```

## Arguments

<code>model</code>	A niche model in raster format or a <code>setA</code> object (see <a href="#">model2sparse</a> ).
<code>ngbs</code>	Numeric. Number of neighbors (see details).
<code>plot_model</code>	Logical. Indicates whether to plot the niche model using a leaflet map, connected suitable cells shown in the same color.

## Details

The main result of the function is the Connectivity-Suitability-Diagram (CSD). In this diagram connected suitable cells make clusters of pixels. For more details about the CSD see (Soberón and Osorio-Olvera, 2022).

## Value

An object of class `csd`. It contains three slots. 1) connections: a data.frame with three columns where first and the second represent (x and y) centroid coordinates of the niche model and the third column with the cluster ID where they belong. 2) interactive\_map: a leaflet map of connected suitable pixels shown in the same color. 3) A RasterLayer of connected suitable pixels.

## Author(s)

Luis Osorio-Olvera & Jorge Soberón

## References

Soberón J, Osorio-Olvera L (2022). “A Dynamic Theory of the Area of Distribution.” [doi:10.48550/ARXIV.2212.06308](#), <https://arxiv.org/abs/2212.06308..>

## Examples

```
set.seed(891)
model_path <- system.file("exdata/Lepus_californicus_cont.tif",
                           package = "bammm")
model <- raster::raster(model_path)
model <- model > 0.7
clusterin <- bammm::bam_clusters(model, ngbs=1, plot_model=TRUE)
```

```
raster::plot(clusterin@raster_map)

clusterin@interactive_map
```

**bam\_sim**

*bam\_sim:* Simulate dispersal dynamics using the set B of the BAM framework.

## Description

`bam_sim`: Simulate dispersal dynamics using the set B of the BAM framework.

## Usage

```
bam_sim(
  sp1,
  sp2,
  set_M,
  initial_points,
  periods_toxic,
  periods_suitable,
  nsteps,
  progress_bar = TRUE
)
```

## Arguments

<code>sp1</code>	Niche model of the focal species (the one that disperses).
<code>sp2</code>	Niche model of the species with whom <code>sp1</code> interacts (currently no dispersal dynamics for this species).
<code>set_M</code>	A setM object containing the adjacency matrix for <code>sp1</code> . See <a href="#">adj_mat</a>
<code>initial_points</code>	A sparse vector returned by the function <a href="#">occs2sparse</a>
<code>periods_toxic</code>	Time periods that <code>sp2</code> takes to develop defense mechanisms (i.e. toxic).
<code>periods_suitable</code>	This is the time that <code>sp2</code> takes to become non-toxic
<code>nsteps</code>	Number of steps to run the simulation
<code>progress_bar</code>	Show progress bar

## Details

The returned object inherits from `setA`, `setM` classes. Details about the dynamic model can be found in Soberon and Osorio-Olvera (2022). The model is cellular automata where the occupied area of a species at time  $t + 1$  is estimated by the multiplication of three binary matrices: one matrix represents movements (`M`), another abiotic -niche- tolerances (`A`), and a third, biotic interactions (`B`) (Soberon and Osorio-Olvera, 2022).

$$\mathbf{G}_j(t + 1) = \mathbf{B}_j(t)\mathbf{A}_j(t)\mathbf{M}_j\mathbf{G}_j(t)$$

**Value**

An object of class `bam`. The object contains 12 slots of information (see details) from which simulation results are stored in `sdm_sim` object, a list of sparse matrices with results of each simulation step.

**Author(s)**

Luis Osorio-Olvera & Jorge Soberón

**References**

Soberón J, Osorio-Olvera L (2022). “A Dynamic Theory of the Area of Distribution.” doi:[10.48550/ARXIV.2212.06308](https://doi.org/10.48550/ARXIV.2212.06308), [https://arxiv.org/abs/2212.06308..](https://arxiv.org/abs/2212.06308)

**Examples**

```
# Compute dispersal dynamics of Urania boisduvalii as a function of
# palatable Omphalea
urap <- system.file("extdata/urania_omph/urania_guanahacabibes.tif",
                     package = "bammm")
ura <- raster::raster(urap)
ompp <- system.file("extdata/urania_omph/omphalea_guanahacabibes.tif",
                     package = "bammm")
omp <- raster::raster(ompp)
msparse <- bammm::model2sparse(ura)
init_coordsdf <- data.frame(x=-84.38751, y= 22.02932)
initial_points <- bammm::occ2sparse(modelsparse = msparse, init_coordsdf)
set_M <- bammm::adj_mat(modelsparse = msparse, ngbs = 1)
ura_sim <- bammm::bam_sim(sp1=ura, sp2=omp, set_M=set_M,
                           initial_points=initial_points,
                           periods_toxic=5,
                           periods_suitable=1,
                           nsteps=40)
ura_omp <- bammm::sim2Raster(ura_sim)
raster::plot(ura_omp[[c(1,5,10,15,20,30,35,40)]])
```

```
if(requireNamespace("animation")){
  # Animation example
  anp <- tempfile(pattern = "simulation_results_", fileext = ".gif")
  new_sim <- bammm::sim2Animation(sdm_simul = ura_sim,
                                   which_steps = seq_len(ura_sim@sim_steps),
                                   fmt = "GIF",
                                   filename = anp)
}
```

---

**bam\_ssimm**

*bam\_ssimm: Simulate dispersal dynamics using the set B of the BAM framework.*

---

## Description

bam\_ssimm: Simulate dispersal dynamics using the set B of the BAM framework.

## Usage

```
bam_ssimm(  
  sp1,  
  sp2,  
  set_M,  
  initial_points,  
  periods_toxic,  
  periods_suitable,  
  dispersal_prob = 0.85,  
  palatable_matrices = FALSE,  
  nsteps,  
  progress_bar = TRUE  
)
```

## Arguments

sp1	Niche model of the focal species (the one that disperses).
sp2	Niche model of the species with whom sp1 interacts (currently no dispersal dynamics for this species).
set_M	A setM object containing the adjacency matrix for sp1. See <a href="#">adj_mat</a>
initial_points	A sparse vector returned by the function <a href="#">occs2sparse</a>
periods_toxic	Time periods that sps2 takes to develop defense mechanisms (i.e. toxic).
periods_suitable	This is the time that sp2 takes to become non-toxic
dispersal_prob	A numeric value indicating the probability to disperse to neighboring cells. This probability is assumed to be binomially distributed
palatable_matrices	Logical. If TRUE palatable matrices for each time will be returned.
nsteps	Number of steps to run the simulation
progress_bar	Show progress bar

## Details

The returned object inherits from [setA](#), [setM](#) classes. Details about the dynamic model can be found in Soberon and Osorio-Olvera (2022).



---

**bioindex-class**      *Class bioindex*

---

**Description**

Class *bioindex*

**Value**

An object of class *bioindex*

**Slots**

*alpha* A matrix with the richness of species per site

*omega* A matrix with the range size of every species

*dispersion\_field* A matrix with the set of ranges of all species that occur in at each locality

**Author(s)**

Luis Osorio-Olvera & Jorge Soberón

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**bioindex\_sparse-class**      *Class bioindex\_sparse*

---

**Description**

Class *bioindex\_sparse*

**Value**

An object of class *bioindex\_sparse*

**Slots**

*alpha* A sparse matrix with the richness of species per site

*omega* A sparse matrix with the range size of every species

*dispersion\_field* A sparse matrix with the set of ranges of all species that occur in at each locality

**Author(s)**

Luis Osorio-Olvera & Jorge Soberón

`community_sim`*community\_bam: Community bam*

## Description

Estimate community dynamics using the bam framework

## Usage

```
community_sim(
  en_models,
  ngbs_vect,
  init_coords,
  nsteps,
  threshold_vec = NULL,
  stochastic_dispersal = FALSE,
  disp_prop2_suitability = TRUE,
  disper_prop = 0.5
)
```

## Arguments

<code>en_models</code>	A stack or directory with the ecological niche models for each species in the community.
<code>ngbs_vect</code>	A vector containing the number of neighbors for each adjacency matrix of each species in the community see <a href="#">adj_mat</a> .
<code>init_coords</code>	A data.frame with 3 columns: sp_name, x and y; x is the longitude and y is the latitude of initial dispersal points
<code>nsteps</code>	Number of iteration steps for the simulation.
<code>threshold_vec</code>	A vector of threshold values used to bnarize niche models.
<code>stochastic_dispersal</code>	Logical. If dispersal depends on a probability of visiting neighbor cells (Moore neighborhood).
<code>disp_prop2_suitability</code>	Logical. If probability of dispersal is proportional to the suitability of reachable cells. The proportional value must be declared in the parameter ‘disper_prop’.
<code>disper_prop</code>	Probability of dispersal to reachable cells.

## Details

Each element in `community_sim` is an object of class. For more details about the simulation see [sdm\\_sim.bam](#).

## Value

An object of class `community_sim`. The object contains simulation results for each species in the community.

**Author(s)**

Luis Osorio-Olvera & Jorge Soberón

**References**

Soberón J, Osorio-Olvera L (2022). “A Dynamic Theory of the Area of Distribution.” doi:10.48550/ARXIV.2212.06308, <https://arxiv.org/abs/2212.06308..>

**Examples**

```
lagos_path <- system.file("extdata/conejos",
                           package = "bammm")
enm_path <- list.files(lagos_path,
                       pattern = ".tif",
                       full.names = TRUE)[seq(1,10)]
en_models <- raster:::stack(enm_path)
ngbs_vect <- sample(1:2, replace = TRUE,
                     size = raster:::nlayers(en_models))
init_coords <- read.csv(file.path(lagos_path,
                                   "lagos_initit.csv"))[seq(1,10),]
nsteps <- 12
sdm_comm <- bammm::community_sim(en_models = en_models,
                                    ngbs_vect = ngbs_vect,
                                    init_coords = init_coords,
                                    nsteps = nsteps)

com_pam <- bammm::csim2pam(sdm_comm, which_steps = seq(1,nsteps))
rich_pam <- pam2richness(com_pam, which_steps = c(1,5,10))
raster:::plot(rich_pam)
```

**community\_sim-class**    *Class community\_sim digram*

**Description**

Class `community_sim` digram

**Value**

An object of class `community_sim`

**Slots**

`community_sim` A list of sparse vectors representing the area occupied by the species

**Author(s)**

Luis Osorio-Olvera & Jorge Soberón

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csd-class	<i>Class csd</i>
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---

**Description**

Class csd

**Value**

An object of class csd

**Slots**

**connections** A data.frame with four columns: x, y, clusterID and cluster\_size  
**interactive\_map** A leaflet map with markers showing the geographical clusters  
**raster\_map** A raster map with cluster IDs as values.

**Author(s)**

Luis Osorio-Olvera & Jorge Soberón

---

csd_estimate	<i>csd_estimate: Estimate the connectivity suitability and dispersal plot</i>
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---

**Description**

csd\_plot gives an estimate of the number of geographic clusters given a set of dispersal hypothesis and a suitability raster

**Usage**

```
csd_estimate(model, dispersal_steps = c(2, 4, 8, 16, 32, 64))
```

**Arguments**

<b>model</b>	A raster model or a setA object representing the suitability model
<b>dispersal_steps</b>	A numeric vector with elements representing the dispersal hypothesis to test.

**Details**

For more information about the Connectivity-Suitability-Diagram see [bam\\_clusters](#)

**Value**

A list of length three. The first element contains the Connectivity- Suitability-Diagram information estimated for each element in the vector of dispersal\_steps. The second is tbl\_df object with a summary of the number of cluster of each dispersal step and the mean number of connected clusters. The last element is base plot showing the information cointained in the tbl\_df object.

**Author(s)**

Luis Osorio-Olvera & Jorge Soberón

**References**

Soberón J, Osorio-Olvera L (2022). “A Dynamic Theory of the Area of Distribution.” doi:10.48550/ARXIV.2212.06308, <https://arxiv.org/abs/2212.06308..>

**Examples**

```
model_path <- system.file("extdata/Lepus_californicus_cont.tif",
                           package = "bammm")
model <- raster::raster(model_path)
model <- model > 0.7
csd_plot <- bammm::csd_estimate(model,
                                  dispersal_steps=c(2,4,8))
csd_plot$plot
```

**csim2pam**

*csim2pam: Converts community simulation to a Presence Absence Matrix (PAM)*

**Description**

Converts community simulation object into a Presence Absence Matrices (PAM) for a given simulation steps.

**Usage**

```
csim2pam(community_sim, which_steps)
```

**Arguments**

`community_sim` An object of class `community_bam`.

`which_steps` Steps in the simulation object to be converted into a PAM

**Details**

For details about the object `community_sim` see `community_sim`

**Value**

An object of class `pam`; it contains five slots. 1) pams: a list of sparse matrices with Presence-Absence information (PAMs). 2) which\_steps: time steps corresponding to each PAM. 3) sp\_names: a vector of species names. 4) the grid area used in the simulation. 5) Non NA cell (pixel) IDs.

**Author(s)**

Luis Osorio-Olvera & Jorge Soberón

**References**

Soberón J, Osorio-Olvera L (2022). “A Dynamic Theory of the Area of Distribution.” doi:[10.48550/ARXIV.2212.06308](https://doi.org/10.48550/ARXIV.2212.06308), <https://arxiv.org/abs/2212.06308>.

**Examples**

```
lagos_path <- system.file("extdata/conejos",
                           package = "bammm")
enm_path <- list.files(lagos_path,
                       pattern = ".tif",
                       full.names = TRUE)[seq(1,10)]
en_models <- raster::stack(enm_path)
ngbs_vect <- sample(1:2, replace = TRUE,
                     size = raster::nlayers(en_models))
init_coords <- read.csv(file.path(lagos_path,
                                   "lagos_initit.csv"))[seq(1,10),]
nsteps <- 10
sdm_comm <- bammm::community_sim(en_models = en_models,
                                    ngbs_vect = ngbs_vect,
                                    init_coords = init_coords,
                                    nsteps = nsteps,
                                    threshold = 0.1)

pamt10 <- bammm::csim2pam(community_sim = sdm_comm ,
                            which_steps = 10)
pams <- bammm::csim2pam(community_sim = sdm_comm ,
                          which_steps = seq_len(10))
rich_pam <- bammm::pam2richness(pams,which_steps = c(1,5))
print(rich_pam)
```

*diversity\_range-class* *Class diversity\_range*

**Description**

Class *diversity\_range*

**Value**

An object of class diversity\_range

**Slots**

alpha A column vector with species richness per site  
omega A column vector with the size of the area of distribution per species.  
alpha\_raster Species richness in raster format.  
dispersion\_field A matrix with the set of ranges of all species that occur in at each locality.  
dispersion\_field\_raster Raster object with the observed values of dispersion field.  
diversity\_range\_raster Raster object of diversity range.  
diversity\_range\_colors Colors to plot endemism levels.  
null\_dispersion\_field\_dist A matrix with dispersion field null distribution.  
xy\_coordinates A matrix of geographical coordinates  
n\_iterations Number of iterations used to estimate the dispersion field null distribution.  
nsps Number of species in the PAM.  
nsites Number of sites in the PAM.

**Author(s)**

Luis Osorio-Olvera & Jorge Soberón

---

diversity\_range\_analysis

*range\_diversity\_analysis: diversity analysis*

---

**Description**

diversity\_range\_analysis biodiversity indices related to diversity-range plots

**Usage**

```
diversity_range_analysis(  
  pam,  
  xy_mat = NULL,  
  lower_interval = 0.05,  
  upper_interval = 0.95,  
  raster_template = NULL,  
  niter = 100,  
  return_null_dfield = FALSE,  
  parallel = TRUE,  
  n_cores = 2  
)
```

## Arguments

pam	A Presence-Absence-Matrix of matrix class or sparse matrix.
xy_mat	A two dimensional matrix with longitude and latitude data.
lower_interval	Lower interval.
upper_interval	Upper interval.
raster_template	A raster template.
niter	Number of iterations to obtain the distribution.
return_null_dfield	If TRUE the null distribution of dispersal field will be returned.
parallel	If TRUE the computations will be performed in parallel.
n_cores	Number of cores for the parallel computation.

## Details

For more information about the biodiversity indices see Soberon and Cavner (2015). For detail about the diversity range analysis see Soberon et al. (2022). To plot diversity range results use `plot` method for objects of class `diversity_range`.

## Value

An object of class `diversity_range`. The main result is the diversity range analysis which shows jointly two indices describing the community composition of every cell in the grid: (1) the relative number of species, and (2) the mean dispersion field (see plot method for `plot` (Soberon et al. 2022). It contains 12 slots with different measurements of biodiversity such as alpha diversity (species richness in each site or pixel), omega (size of the area of distribution of each species), dispersion field (the standardized size of the area of distribution of all species occurring in each pixel).

## Author(s)

Luis Osorio-Olvera & Jorge Soberón

## References

- Soberón J, Cobos ME, Nuñez-Penichet C (2021). “Visualizing species richness and site similarity from presence-absence matrices.” *Biodiversity Informatics*, **16**(1), 20–27. doi:10.17161/bi.v16i1.14782, <https://journals.ku.edu/jbi/article/view/14782..>

Soberón J, Cavner J (2015). “Indices of Biodiversity Pattern Based on Presence-Absence Matrices: A GIS Implementation.” *Biodiversity Informatics*, **10**, 22–34..

## Examples

```

            niter = 10,
            return_null_dfield=TRUE)
bammm::plot(rdivan,plot_type="diversity_range")
# Lagomorphos

lagos_path <- system.file("extdata/conejos",
                           package = "bammm")
enm_path <- list.files(lagos_path,
                       pattern = ".tif",
                       full.names = TRUE)
en_models <- raster::stack(enm_path) >0.01
nonas <- which(!is.na(en_models[[1]][[]]))
xy_mat <- sp::coordinates(en_models[[1]])[ nonas,]
pam <- bammm::models2pam(en_models,sparse=FALSE)

rdivan <- bammm::diversity_range_analysis(pam=pam,
                                             xy_mat=xy_mat,
                                             raster_template = en_models[[1]],
                                             parallel=TRUE,
                                             n_cores=2,
                                             return_null_dfield=TRUE)
bammm::plot(rdivan,plot_type="diversity_range")
bammm::plot(rdivan,plot_type="diversity_range_map")
if(requireNamespace("plotly")){
  #bammm::plot(rdivan,plot_type="diversity_range_interactive")
}

```

eigen\_bam

*eigen\_bam: Compute the Eigen system of two bam objects*

## Description

Calculates the Eigen values and Eigen vectors of bam objects

## Usage

```
eigen_bam(A = NULL, M = NULL, which_eigen = 1, rmap = TRUE)
```

## Arguments

A	A bam object of class setA.
M	A bam object of class setM.
which_eigen	An integer representing the which eigen value and eigen vector will be computed.
rmap	Logical. If TRUE the function will return a map of the eigen vector of the product AM.

## Details

The eigenvector associated with the dominant eigenvalue of an adjacency matrix provides information about the number of forms in which a cell can be visited from other cells. Details about the eigen analysis in the context of the area of distribution can be found in Soberon and Osorio-Olvera (2022).

## Value

A list with four objects. 1) eigen\_values (these are indicated in which\_eigen parameter of the function), 2) eigen\_vectors (the corresponding eigen vectors of each eigen value), 3) Standardized eigen vectors (0 to 1), 4) A RasterLayer depicting the information of the first eigen vector of the system.

## Author(s)

Luis Osorio-Olvera & Jorge Soberón

## References

Soberón J, Osorio-Olvera L (2022). “A Dynamic Theory of the Area of Distribution.” doi:10.48550/ARXIV.2212.06308, <https://arxiv.org/abs/2212.06308..>

## Examples

```
model_path <- system.file("extdata/Lepus_californicus_cont.tif",
                           package = "bammm")
model <- raster::raster(model_path)
sparse_mod <- bammm::model2sparse(model = model, 0.75)
plot(sparse_mod@niche_model)
adj_mod <- bammm::adj_mat(sparse_mod, ngbs = 1, eigen_sys = TRUE)
eig_bam <- bammm::eigen_bam(A=sparse_mod, M=adj_mod)
raster::plot(eig_bam$map)
```

## Description

S4 classes to organize data and results of `bamm` objects

## Value

An object of class `g_area`

**Slots**

**coordinates** A two column matrix with coordinates  
**eigen\_vec** Eigen vector of adjacency matrix  
**eigen\_val** Eigen value of adjacency matrix slot **g\_model** A raster representing the geographic area  
slot **g\_sparse** A sparse matrix of the geographic area

**Author(s)**

Luis Osorio-Olvera & Jorge Soberón

jaccard

*jaccard: Estimates the Jaccard index for comparing two binary maps*

**Description**

Estimates the Jaccard index for comparing two binary maps

**Usage**

```
jaccard(m1, m2)
```

**Arguments**

<b>m1</b>	A binary raster A or an object of class setA returned by the function <a href="#">model2sparse</a> .
<b>m2</b>	A binary raster A or an object of class setA returned by the function <a href="#">model2sparse</a> .

**Details**

The Jaccard index is computed as follows

$$J(A, B) = \frac{|A \cap B|}{|A \cup B|} = \frac{|A \cap B|}{|A| + |B| - |A \cap B|}.$$

**Value**

Returns a data.frame with three values: 1) jaccard (Jaccard index), 2) percentage\_m1 (the percentage of m1 that the intersection  $|A \cap B|$  represents), and 3) percentage\_m2

**Author(s)**

Luis Osorio-Olvera & Jorge Soberón

## Examples

```
m1_path <- system.file("extdata/conejos/Lepus_othus_cont.tif",
                       package = "bammm")
m2_path <- system.file("extdata/conejos/Brachylagus_idahoensis_cont.tif",
                       package = "bammm")
m1 <- raster::raster(m1_path) > 0.01
m2 <- raster::raster(m2_path) > 0.01
jcc <- bammm::jaccard(m1, m2)
print(jcc)
```

**leaflet-class**

*Class leaflet leaflet*

## Description

Class leaflet leaflet

## Value

An object of class leaflet

## Author(s)

Luis Osorio-Olvera & Jorge Soberón

**model2sparse**

*model2sparse: Converts a niche model into a diagonal sparse matrix*

## Description

model2sparse: Converts a niche model into a diagonal sparse matrix

## Usage

```
model2sparse(model, threshold = NULL)
```

## Arguments

- |           |  |
|-----------|--|
| model     | A raster object representing the geographic projection of a niche model. |
| threshold | A threshold to convert a continuous model into a binary model.           |

## Details

threshold parameter represents the suitability value used to convert continuous model into a binary model.

**Value**

An object of class `setA`. The niche model is stored as diagonal sparse matrix (slot `sparse_model`).

**Author(s)**

Luis Osorio-Olvera & Jorge Soberón

**Examples**

```
model_path <- system.file("extdata/Lepus_californicus_cont.tif",
                           package = "bammm")
model <- raster::raster(model_path)

sparse_mod <- bammm::model2sparse(model, threshold=0.75)
print(sparse_mod)
raster::plot(sparse_mod@niche_model)
```

---

models2pam

*models2pam: Converts binary rasters to a PAM*

---

**Description**

Function to convert binary raster models to a Presence Absences Matrix.

**Usage**

```
models2pam(mods_stack, sparse = TRUE, parallel = FALSE, ncores = 2)
```

**Arguments**

<code>mods_stack</code>	A raster stack containing binary models of each species in the community.
<code>sparse</code>	Logical. If TRUE the PAM will be returned as a sparse matrix.
<code>parallel</code>	Logical. If TRUE computations will be done in parallel
<code>ncores</code>	Integer. Number of cores to run the parallel process.

**Details**

For more information about PAM see Soberon and Cavner (2015).

**Value**

A presence-absence matrix (PAM).

**Author(s)**

Luis Osorio-Olvera & Jorge Soberón

## References

Soberon J, Cavner J (2015). “Indices of Biodiversity Pattern Based on Presence-Absence Matrices: A GIS Implementation.” *Biodiversity Informatics*, **10**, 22–34..

## Examples

```
lagos_path <- system.file("extdata/conejos",
                           package = "bammm")
enm_path <- list.files(lagos_path,
                       pattern = ".tif",
                       full.names = TRUE)[1:10]
en_models <- raster:::stack(enm_path) >0.01
pam <- bammm::models2pam(en_models, sparse=FALSE,
                         parallel=FALSE, ncores=2)
head(pam)
```

### null\_dispersion\_field\_distribution

*null\_dispersion\_field\_distribution: Null distribution of the dispersion field*

## Description

`null_dispersion_field_distribution` estimates a random distribution of the dispersion field values.

## Usage

```
null_dispersion_field_distribution(
  pam,
  n_iter = 10,
  parallel = TRUE,
  n_cores = 2
)
```

## Arguments

<code>pam</code>	A Presence-Absence-Matrix of matrix class or sparse matrix.
<code>n_iter</code>	Number of iterations to obtain the distribution.
<code>parallel</code>	If TRUE the computations will be performed in parallel.
<code>n_cores</code>	Number of cores for the parallel computation.

## Details

Estimates a random distribution of the dispersion field values. To obtain random values it uses the function code `permute_pam` at each step of the iterations. Randomization of the PAM is performed using the Babe Ruth Algorithm see Strona et al. (2014).

**Value**

A data matrix of size nrow(pam) X n\_iter with dispersion field values.

**Author(s)**

Luis Osorio-Olvera & Jorge Soberón

**References**

- Soberon J, Cavner J (2015). “Indices of Biodiversity Pattern Based on Presence-Absence Matrices: A GIS Implementation.” *Biodiversity Informatics*, **10**, 22–34.
- Strona G, Nappo D, Boccacci F, Fattorini S, San-Miguel-Ayanz J (2014). “A fast and unbiased procedure to randomize ecological binary matrices with fixed row and column totals.” *Nature Communications*, **5**(1), 1–9. ISSN 20411723, doi:10.1038/ncomms5114, <https://www.r-project.org>.

**Examples**

```
set.seed(111)
pam <- matrix(rbinom(100,1,0.3),nrow = 10,ncol = 10)
dfield_rand <- bammm::null_dispersion_field_distribution(pam,n_iter=10,
                                                          parallel=FALSE,
                                                          n_cores = 2)
head(dfield_rand)
```

**occs2sparse**

*occs2sparse: Converts occurrence data into a sparse matrix object*

**Description**

**occs2sparse:** Converts occurrence data into a sparse matrix object

**Usage**

```
occs2sparse(modelsparse, occs)
```

**Arguments**

- |                    |   |
|--------------------|---|
| <b>modelsparse</b> | A setA object returned by the function <a href="#">model2sparse</a>   |
| <b>occs</b>        | A matrix or a data.frame containing two columns. The first one is the longitude and the second is the latitude. |

**Details**

Rows of this column vector represent non NA pixels of the niche model.

**Value**

A sparse vector of zeros (presences) and ones (absences).

**Author(s)**

Luis Osorio-Olvera & Jorge Soberón

**Examples**

```
model_path <- system.file("extdata/Lepus_californicus_cont.tif",
                          package = "bammm")
model <- raster::raster(model_path)

sparse_mod <- bammm::model2sparse(model, threshold=0.05)

occs_lep_cal <- data.frame(longitude = c(-115.10417,
                                           -104.90417),
                             latitude = c(29.61846,
                                         29.81846))

occs_sparse <- bammm::occs2sparse(modelsparse = sparse_mod,
                                    occs = occs_lep_cal)

head(occs_sparse)
```

*pam-class*

*Class pam Presence-Absence Matrix*

**Description**

Class `pam` Presence-Absence Matrix

**Value**

An object of class `pam`

**Slots**

- `pams` A list of sparse matrices representing Presence-Absence Matrix for each simulation time
- `which_steps` Simulation steps
- `sp_names` Names of species in the PAM
- `grid` Raster grid of the studied area
- `cellIDs` Cells with ids of the PAM sites

**Author(s)**

Luis Osorio-Olvera & Jorge Soberón

---

pam2bioindex	<i>pam2bioindex: PAM to biodiversity index</i>
--------------	--

---

## Description

`pam2bioindex` estimates various biodiversity indices for a certain PAM.

## Usage

```
pam2bioindex(pam, biodiv_index = "dispersion_field", as_sparse = FALSE)
```

## Arguments

- |                           |   |
|---------------------------|---|
| <code>pam</code>          | A Presence-Absence-Matrix of matrix class or sparse matrix. |
| <code>biodiv_index</code> | Possible values are alpha, omega, dispersion_field, all.    |
| <code>as_sparse</code>    | Return indices as sparse objects                            |

## Details

The biodiversity indices can be found in Soberón and Cavner (2015).

## Value

An object of class `bioindex` with three slots each represents a matrix of diversity indices: alpha, omega, and dispersion field.

## Author(s)

Luis Osorio-Olvera & Jorge Soberón

## References

Soberon J, Cavner J (2015). “Indices of Biodiversity Pattern Based on Presence-Absence Matrices: A GIS Implementation.” *Biodiversity Informatics*, **10**, 22–34.

## Examples

```
set.seed(111)
pam <- matrix(rbinom(100,1,0.3),nrow = 10,ncol = 10)
bioindices <- bammm::pam2bioindex(pam=pam,biodiv_index="all")
# Return results as sparse models
bioindices <- bammm::pam2bioindex(pam=pam,biodiv_index="all",as_sparse=TRUE)
bioindices@alpha
bioindices@omega
bioindices@dispersion_field
```

---

<b>pam2richness</b>	<i>pam2richness:</i> Converts Presence Absence Matrix (pam object) to richness raster
---------------------	---

---

## Description

Converts Presence Absence Matrix (pam object) to richness raster

## Usage

```
pam2richness(pamobj, which_steps)
```

## Arguments

pamobj	An object of class pam see <a href="#">csim2pam</a>
which_steps	Time steps in the pam to convert

## Value

A RasterStack richness for each simulation step

## Author(s)

Luis Osorio-Olvera & Jorge Soberón.

## Examples

```
lagos_path <- system.file("extdata/conejos",
                           package = "bammm")
enm_path <- list.files(lagos_path,
                       pattern = ".tif",
                       full.names = TRUE)[seq(1,10)]
en_models <- raster::stack(enm_path)
ngbs_vect <- sample(2,replace = TRUE,
                     size = raster::nlayers(en_models))
init_coords <- read.csv(file.path(lagos_path,
                                   "lagos_initit.csv"))[seq(1,10),]
nsteps <- 10
sdm_comm <- bammm::community_sim(en_models = en_models,
                                    ngbs_vect = ngbs_vect,
                                    init_coords = init_coords,
                                    nsteps = nsteps,
                                    threshold = 0.1)

pams <-bammm::csim2pam(community_sim = sdm_comm ,
                        which_steps = seq_len(nsteps))
richness_stack <- bammm::pam2richness(pams,which_steps=pams@which_steps)
raster::plot(richness_stack)
```

permute\_pam

*permute\_pam: Function to permute a Presence-Absence-Matrix.*

## Description

`permute_pam`: Function to permute a Presence-Absence-Matrix.

## Usage

```
permute_pam(m, niter = NULL, as_sparse = FALSE)
```

## Arguments

<code>m</code>	Presence-Absence-Matrix (PAM) or a binary matrix with columns representing species and rows sites.
<code>niter</code>	Number of iterations to permute the PAM.
<code>as_sparse</code>	If TRUE the PAM will be returned as a sparse matrix

## Details

This function is an implementation of the curve ball algorithm following Strona et al. (2014).

## Value

Returns a permuted matrix of the same dimensions of `m` (same number of rows and columns). Note that the sum of each row and column of this permuted matrix is equal to that of `m`. species.

## Author(s)

Luis Osorio-Olvera & Jorge Soberón

## References

Strona G, Nappo D, Boccacci F, Fattorini S, San-Miguel-Ayanz J (2014). “A fast and unbiased procedure to randomize ecological binary matrices with fixed row and column totals.” *Nature Communications*, **5**(1), 1–9. ISSN 20411723, doi:10.1038/ncomms5114, <https://www.r-project.org..>

## Examples

```
set.seed(111)
pam <- matrix(rbinom(100, 1, 0.3), nrow = 10, ncol = 10)
ppam <- bamr::permute_pam(m = pam, niter = NULL, as_sparse = FALSE)
# Check if matrices are different
all(pam == ppam)
# Check if row totals are the same
all(Matrix::rowSums(pam) == Matrix::rowSums(ppam))
# Check if column total are the same
all(Matrix::colSums(pam) == Matrix::colSums(ppam))
```

---

**plot,diversity\_range,ANY-method**

*Plot method for objects of class diversity\_range **bamm**.*

---

## Description

Plot method for objects of class `diversity_range` **bamm**.

## Usage

```
## S4 method for signature 'diversity_range,ANY'
plot(
  x,
  xlab = NULL,
  plot_type = "diversity_range",
  legend = TRUE,
  legend_position = "bottomright",
  ylab = NULL,
  col = NULL,
  pch = NULL,
  pch_legend = 19,
  radius = 0.5,
  ...
)
```

## Arguments

<code>x</code>	An object of class <code>diversity_range</code>
<code>xlab</code>	x label
<code>plot_type</code>	Plot type: possible options: "diversity_range" (range-diversity plot), "diversity_range_map" (a raster map with <code>diversity_range</code> categories), "alpha" (a raster map with alpha diversity values), "dispersion_field" (a raster with dispersion field)
<code>legend</code>	Logical. If TRUE the legend of the categorical diversity range values will appear.
<code>legend_position</code>	Legend position.
<code>ylab</code>	y label
<code>col</code>	Plot colors.
<code>pch</code>	Patch type.
<code>pch_legend</code>	Patch type for legends.
<code>radius</code>	Size of the patch for the interactive map.
<code>...</code>	Graphical parameters. Any argument that can be passed to 1) <code>base::plot</code> , such as <code>axes=FALSE, main='title', ylab='latitude'</code> 2) <code>leaflet::leaflet</code> or 3) <code>leaflet::addCircleMarkers</code> .

**Details**

To show interactive diversity\_range plots install the 'plotly' R package.

**Value**

Plot of the results of the diversity\_range analysis

**Author(s)**

Luis Osorio-Olvera & Jorge Soberón

**predict,bam-method**     *Predict method of the package **bamm**.*

**Description**

predicts species' distribution under suitability changes

**Usage**

```
## S4 method for signature 'bam'
predict(
  object,
  niche_layers,
  nbgs_vec = NULL,
  nsteps_vec,
  stochastic_dispersal = FALSE,
  disp_prop2_suitability = TRUE,
  disper_prop = 0.5,
  animate = FALSE,
  period_names = NULL,
  fmt = "GIF",
  filename,
  bg_color = "#F6F2E5",
  suit_color = "#0076BE",
  occupied_color = "#03C33F",
  png_keyword = "sdm_sim",
  ani.width = 1200,
  ani.height = 1200,
  ani.res = 300
)
```

**Arguments**

<b>object</b>	a of class bam.
<b>niche_layers</b>	A raster or RasterStack with the niche models for each time period

<code>nbgs_vec</code>	A vector with the number of neighbors for the adjacency matrices
<code>nsteps_vec</code>	Number of simulation steps for each time period.
<code>stochastic_dispersal</code>	Logical. If dispersal depends on a probability of visiting neighbor cells (Moore neighborhood).
<code>disp_prop2_suitability</code>	Logical. If probability of dispersal is proportional to the suitability of reachable cells. The proportional value must be declared in the parameter ‘ <code>disper_prop</code> ’.
<code>disper_prop</code>	Probability of dispersal to reachable cells.
<code>animate</code>	Logical. If TRUE a dispersal animation on climate change scenarios will be created
<code>period_names</code>	Character vector with the names of periods that will be animated. Default NULL.
<code>fmt</code>	Animation format. Possible values are GIF and HTML
<code>filename</code>	File name.
<code>bg_color</code>	Color for unsuitable pixels. Default "#F6F2E5".
<code>suit_color</code>	Color for suitable pixels. Default "#0076BE".
<code>occupied_color</code>	Color for occupied pixels. Default "#03C33F".
<code>png_keyword</code>	A keyword name for the png images generated by the function
<code>ani.width</code>	Animation width unit in px
<code>ani.height</code>	Animation height unit in px
<code>ani.res</code>	Animation resolution unit in px

### Value

A RasterStack of predictions of dispersal dynamics as a function of environmental change scenarios.

### Author(s)

Luis Osorio-Olvera & Jorge Soberón

### Examples

```
# rm(list = ls())
# Read raster model for Lepus californicus
model_path <- system.file("extdata/Lepus_californicus_cont.tif",
                           package = "bammm")
model <- raster::raster(model_path)
# Convert model to sparse
sparse_mod <- bammm::model2sparse(model = model, threshold=0.1)
# Compute adjacency matrix
adj_mod <- bammm::adj_mat(sparse_mod, ngb=1)

# Initial points to start dispersal process

occ_lep_cal <- data.frame(longitude = c(-115.10417,
```

```

          -104.90417),
latitude = c(29.61846,
           29.81846))
# Convert to sparse the initial points
occs_sparse <- bammm::occ2sparse(modelsparse = sparse_mod,
                                    occs = occs_lep_cal)

# Run the bam (sdm) simulation for 100 time steps
smd_lep_cal <- bammm::sdm_sim(set_A = sparse_mod,
                               set_M = adj_mod,
                               initial_points = occs_sparse,
                               nsteps = 10)
#-----
# Predict species' distribution under suitability change
# scenarios (could be climate chage scenarios).
#-----

# Read suitability layers (two suitability change scenarios)
layers_path <- system.file("extdata/suit_change",
                           package = "bammm")
niche_mods_stack <- raster::stack(list.files(layers_path,
                                              pattern = ".tif$",
                                              full.names = TRUE))
raster::plot(niche_mods_stack)
# Predict
new_preds <- predict(object = smd_lep_cal,
                      niche_layers = niche_mods_stack,
                      nsteps_vec = c(50,100))

# Generate the dispersal animation for time period 1 and 2

if(requireNamespace("animation")){
  ani_prd <- tempfile(pattern = "prediction_",fileext = ".gif")
  new_preds <- predict(object = smd_lep_cal,
                        niche_layers = niche_mods_stack,
                        nsteps_vec = c(10,10),
                        animate=TRUE,
                        filename=ani_prd,
                        fmt="GIF")
}

}

```

---

sdm\_sim*sdm\_sim: Simulate single species dispersal dynamics using the BAM framework.*

---

## Description

sdm\_sim: Simulate single species dispersal dynamics using the BAM framework.

## Usage

```
sdm_sim(
  set_A,
  set_M,
  initial_points,
  nsteps,
  stochastic_dispersal = TRUE,
  disp_prop2_suitability = TRUE,
  disper_prop = 0.5,
  progress_bar = TRUE
)
```

## Arguments

set_A	A setA object returned by the function <a href="#">model2sparse</a>
set_M	A setM object containing the adjacency matrix of the study area. See <a href="#">adj_mat</a>
initial_points	A sparse vector returned by the function <a href="#">occs2sparse</a>
nsteps	Number of steps to run the simulation
stochastic_dispersal	Logical. If dispersal depends on a probability of visiting neighbor cells (Moore neighborhood).
disp_prop2_suitability	Logical. If probability of dispersal is proportional to the suitability of reachable cells. The proportional value must be declared in the parameter ‘disper_prop’.
disper_prop	Probability of dispersal to reachable cells.
progress_bar	Show progress bar

## Details

The model is cellular automata where the occupied area of a species at time  $t + 1$  is estimated by the multiplication of two binary matrices: one matrix represents movements (M), another abiotic -niche- tolerances (A) (Soberon and Osorio-Olvera, 2022).

$$\mathbf{G}_j(t + 1) = \mathbf{A}_j(t)\mathbf{M}_j\mathbf{G}_j(t)$$

The equation describes a very simple process: To find the occupied patches in  $t + 1$  start with those occupied at time  $t$  denoted by  $\mathbf{G}_j(t)$ , allow the individuals to disperse among adjacent patches, as defined by  $\mathbf{M}_j$ , then remove individuals from patches that are unsuitable, as defined by  $\mathbf{A}_j(t)$ .

## Value

An object of class [bam](#) with simulation results. The simulation are stored in the sdm\_sim slot (a list of sparse matrices).

## Author(s)

Luis Osorio-Olvera & Jorge Soberón

## References

Soberón J, Osorio-Olvera L (2022). “A Dynamic Theory of the Area of Distribution.” doi:10.48550/ARXIV.2212.06308, <https://arxiv.org/abs/2212.06308..>

## Examples

```
model_path <- system.file("extdata/Lepus_californicus_cont.tif",
                           package = "bammm")
model <- raster::raster(model_path)

sparse_mod <- bammm::model2sparse(model, threshold=0.05)
adj_mod <- bammm::adj_mat(sparse_mod, ngbs=1)
occs_lep_cal <- data.frame(longitude = c(-110.08880,
                                             -98.89638),
                             latitude = c(30.43455,
                                         25.19919))

occs_sparse <- bammm::occs2sparse(modelsparse = sparse_mod,
                                      occs = occs_lep_cal)
sdm_lep_cal <- bammm::sdm_sim(set_A = sparse_mod,
                                 set_M = adj_mod,
                                 initial_points = occs_sparse,
                                 nsteps = 10,
                                 stochastic_dispersal = TRUE,
                                 disp_prop2_suitability=TRUE,
                                 disper_prop=0.5,
                                 progress_bar=TRUE)

sim_res <- bammm::sim2Raster(sdm_lep_cal)
raster::plot(sim_res)
```

setA-class

*Class for the A set of the BAM diagram*

## Description

A class for the A set of the BAM diagram. It contains raster models and IDs of pixels with values different than NA.

## Value

An object of class setA showClass("setA")

**Slots**

- `niche_model` A niche model in raster format. It can be a binary model or continuous. If the model is in a continuous format.
- `suit_threshold` Suitability value used to binarize continuous model
- `cellIDs` A numeric vector with the IDs of the cells with prediction values
- `suit_values` A numeric vector with suitability value of the continuous map
- `sparse_model` A niche model in sparse matrix format

**Author(s)**

Luis Osorio-Olvera & Jorge Soberón

**setM-class**

*Class for the M set of the bamM diagram*

**Description**

Class for the M set of the bamM diagram

**Value**

An object of class setM

**Slots**

- `adj_matrix` An adjacency matrix
- `adj_list` An adjacency list
- `initial_points` A presence-absence vector with species' occurrences
- `n_initial_points` Number of initial points used to start the dispersal process
- `ngbs` Number of neighbors

**Author(s)**

Luis Osorio-Olvera & Jorge Soberón

**Examples**

```
showClass("setM")
```

---

**shape2Grid***shape2Grid: Function to create a grid given a spatial polygon*

---

## Description

shape2Grid creates a raster grid given a spatial polygon and a grid resolution.

## Usage

```
shape2Grid(shpolygon, resolution, ones = TRUE)
```

## Arguments

shpolygon	A SpatialPolygon, SpatialPolygonDataFrame representing the desired shape of the grid.
resolution	Numeric. Spatial resolution of the grid.
ones	Logical. Fill with ones the values of the raster. If not the values will be written as cellID values.

## Value

Returns a raster object with the shape of 'shpolygon' of a given resolution.

## Author(s)

Luis Osorio-Olvera & Jorge Soberón

## Examples

```
x_coord <- c(-106.5699, -111.3737, -113.9332, -110.8913, -106.4262, -106.5699)
y_coord <- c(16.62661, 17.72373, 19.87618, 22.50763, 21.37728, 16.62661)
xy <- cbind(x_coord, y_coord)
p <- sp::Polygon(xy)
ps <- sp::Polygons(list(p),1)
sps <- sp::SpatialPolygons(list(ps))
r1 <- bamm::shape2Grid(sps,resolution = 0.1,ones = FALSE)
plot(r1)
sp::plot(sps,add=TRUE)
```

**show, setA-method**      *Show information in setA class **bamm**.*

### Description

Show information in setA class **bamm**.  
 Show information in csd class **bamm**.  
 Show information in pam class **bamm**.  
 Show information in pam class **bamm**.  
 Show information in setA class **bamm**.  
 Show information in diversity\_range class **bamm**.

### Usage

```
## S4 method for signature 'setA'
show(object)

## S4 method for signature 'csd'
show(object)

## S4 method for signature 'pam'
show(object)

## S4 method for signature 'bioindex_sparse'
show(object)

## S4 method for signature 'setM'
show(object)

## S4 method for signature 'diversity_range'
show(object)
```

### Arguments

**object**      An object of class diversity\_range

### Value

Display information about the setA object  
 Display information about the csd object  
 Display information about the pam object  
 Display information about the bioindex\_spars object  
 Display information about the setM object  
 Display information about the diversity\_range object

**Author(s)**

Luis Osorio-Olvera & Jorge Soberón

**sim2Animation**

*sim2Animation: Animate BAM simulation object.*

**Description**

Animates BAM simulation object.

**Usage**

```
sim2Animation(
  sdm_simul,
  which_steps,
  fmt = "GIF",
  filename,
  png_keyword = "sdm_sim",
  extra_legend = NULL,
  bg_color = "#F6F2E5",
  suit_color = "#0076BE",
  occupied_color = "#03C33F",
  gif_vel = 0.8,
  ani.width = 1200,
  ani.height = 1200,
  ani.res = 300
)
```

**Arguments**

<code>sdm_simul</code>	A bam object. See <a href="#">sdm_sim</a>
<code>which_steps</code>	A numeric vector indicating the simulation steps that are going to be converted into raster layers.
<code>fmt</code>	Animation format. Possible values are GIF and HTML
<code>filename</code>	File name.
<code>png_keyword</code>	A keyword name for the png images generated by the function
<code>extra_legend</code>	A legend to add to the animation.
<code>bg_color</code>	Color for unsuitable pixels. Default "#F6F2E5".
<code>suit_color</code>	Color for suitable pixels. Default "#0076BE".
<code>occupied_color</code>	Color for occupied pixels. Default "#03C33F".
<code>gif_vel</code>	A value that regulates the velocity of frame transitions. The bigger it is the transition will be slower default 0.8
<code>ani.width</code>	Animation width unit in px
<code>ani.height</code>	Animation height unit in px
<code>ani.res</code>	Animation resolution unit in px

## Details

The animation can be saved in a GIF or HTML format. Note that the generation of the GIF can be time consuming for large simulation (simulations with more than 60 time steps).

## Value

A RasterStack of species' distribution at each simulation step

## Author(s)

Luis Osorio-Olvera & Jorge Soberón

## Examples

```
model_path <- system.file("extdata/Lepus_californicus_cont.tif",
                           package = "bammm")
model <- raster::raster(model_path)
sparse_mod <- bammm::model2sparse(model, 0.1)
adj_mod <- bammm::adj_mat(sparse_mod, ngbns=2)
occs_lep_cal <- data.frame(longitude = c(-115.10417,
                                           -104.90417),
                             latitude = c(29.61846,
                                         29.81846))
occs_sparse <- bammm::occs2sparse(modelsparse = sparse_mod,
                                      occs = occs_lep_cal)
sdm_lep_cal <- bammm::sdm_sim(set_A = sparse_mod,
                                 set_M = adj_mod,
                                 initial_points = occs_sparse,
                                 nsteps = 50)

if(requireNamespace("animation")){
  ani_name <- tempfile(pattern = "simulation_", fileext = ".html")
  sdm_lep_cal_st <- bammm::sim2Animation(sdm_simul = sdm_lep_cal,
                                            which_steps = seq(1,50,by=1),
                                            fmt = "HTML", ani.width = 1200,
                                            ani.height = 1200,
                                            filename = ani_name)
}
```

## Description

Convert a BAM simulation object to RasterStack.

## Usage

```
sim2Raster(sdm_simul, which_steps = NULL)
```

**Arguments**

- sdm\_simul A bam object. See [sdm\\_sim](#)  
which\_steps A numeric vector indicating the simulation steps that are going to be converted into raster layers.

**Value**

A RasterStack of species' distribution at each simulation step

**Author(s)**

Luis Osorio-Olvera & Jorge Soberón

**Examples**

```
model_path <- system.file("extdata/Lepus_californicus_cont.tif",
                           package = "bammm")
model <- raster::raster(model_path)
sparse_mod <- bammm::model2sparse(model, threshold=0.1)
adj_mod <- bammm::adj_mat(sparse_mod, ngbs = 1)
occ_s_lep_cal <- data.frame(longitude = c(-115.10417,
                                             -104.90417),
                               latitude = c(29.61846,
                                           29.81846))
occ_s_p <- bammm::occ2sparse(modelsparse = sparse_mod,
                               occs = occs_lep_cal)
sdm_lep_cal <- bammm::sdm_sim(set_A = sparse_mod,
                                 set_M = adj_mod,
                                 initial_points = occs_p,
                                 nsteps = 10)
sdm_lep_cal_st <- bammm::sim2Raster(sdm_simul = sdm_lep_cal,
                                       which_steps = seq(1,10,by=1))

raster::plot(sdm_lep_cal_st)
```

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