# Package: ENMTools (via r-universe)

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     biogeography. Warren, D.L., R.E. Glor, and M. Turelli (2008)
     <doi:10.1111/j.1558-5646.2008.00482.x> Glor, R.E., and D.L.
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Type Package

2 Contents

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# **Contents**

add.env	3
assert.extras	4
background.buffer	5
background.points.buffer	6
background.raster.buffer	6
	7
background.test	7
calc.B1	9
calc.B2	10
check.bg	10
check.clade	11
	12
	12
1	13
	13
	14
drop.species	15
	15
enmtools.bc	16
	18
	19
enmtools.dm	20
enmtools.ecospat.bg	21
1	23
6	24
enmtools.glm	26
Y	28
	29
enmtools.rf	30
$\mathcal{C}$	32
1	34
1	35
	36
	37
1	38
	39
find.extras	39
8	10
	10
	11
	12

add.env 3

add.	env	adds the env	vironmen alled by	tal pred the use	dictor ve r direct	alues to th	e occurrenc	t layers, and e data. Typ- m Elith and	
Index									<b>67</b>
	-								
	wrap_list								66
	visualize.overlap .								65
	visualize.enm								64
	unwrap_list								64
	trimdupes.by.raster								63
	threespace.plot								62
	species.from.file								62
	sim.points								61
	rasterToPoints2								61
	raster.standardize .								60
	raster.resid								59
	raster.pca								59
	raster.overlap								58
	raster.cor.plot								57
	raster.cor.matrix								57
	raster.cor								56
	raster.breadth								55
	rangebreak.ribbon .								54
	rangebreak.linear .								52
	rangebreak.blob								50 51
	node.overlap point.overlap								49 50
	multistack.pca								49 49
	moses.list								48
	marginal.plots								47
	interactive.plot.enmt	-							46
	interactive.plot.enmt								45
	interactive.plot								45
	install.extras								44
	identity.test								42

# Description

Takes an enmtools.species object and a set of environment layers, and adds the environmental predictor values to the occurrence data. Typically not called by the user directly. Code modified from Elith and Hijmans SDM with R tutorial

# Usage

```
add.env(species, env, verbose = TRUE)
```

4 assert.extras

# **Arguments**

species An enmtools.species or enmtools.clade object

env A raster stack of environmental variables

verbose Controls printing of progress messages

#### Value

An enmtools.species object with environmental data attached to it.

assert.extras Assert that the extra packages needed for an ENMTools function are installed and available

# **Description**

Triggers an error if any of the extra packages required for an ENMTools function are not available.

# Usage

```
assert.extras(funs = NULL)
```

# Arguments

funs character vector of ENMTools function names to install extra dependencies for.

If 'NULL', extras for all ENMTools functions will be installed.

#### Value

None, this function is used for its side-effects

# **Examples**

```
if(check.extras("enmtools.gam")) {
  assert.extras("enmtools.gam")
}
```

background.buffer 5

background.buffer

Takes a set of points, a buffer radius, a buffer type, and a mask and returns either a raster, a polygon, or background points representing the species background. Code modified from Elith and Hijmans SDM with R tutorial

### **Description**

Takes a set of points, a buffer radius, a buffer type, and a mask and returns either a raster, a polygon, or background points representing the species background. Code modified from Elith and Hijmans SDM with R tutorial

### Usage

```
background.buffer(
  points,
  buffer.width,
  buffer.type = "circles",
  mask = NA,
  return.type = "raster",
  n = 1000
)
```

### **Arguments**

points A two column data frame with X and Y coordinates

buffer.width Radius for circular buffers to draw around points, for buffer.type = circular. If

buffer.type = "convhull", denotes the amount to which the initial polygon should be buffered. It's worth noting that this argument may in some cases be treated as map units (e.g., lat and long), but in other cases may be treated as meters. If you're getting weird behavior, you might try changing the units you're using to

specify buffer.width.

buffer.type Which method to use for buffering species occurrence points. Currently accepts

"circles" and "convhull".

mask A raster to use as a mask for trimming the buffer if the return type is "raster" or

"points"

return. type What type of object to return. Can be "raster", "polygon", or "points".

n Sample size for number of background points to return, for return.type = "points".

### Value

Either a raster, a polygon, or a data frame of points representing the species background.

background.points.buffer

Takes a set of points, a buffer radius, a sample size, and a mask and returns randomly sampled points from within that buffer radius. Code modified from Elith and Hijmans SDM with R tutorial

# **Description**

NOTE: This function has been replaced by background.buffer.

# Usage

```
background.points.buffer(points, radius, n, mask)
```

### **Arguments**

points A two column data frame with X and Y coordinates

radius Radius for circular buffers to draw around points, in meters.

Sample size for number of background points to return

mask A raster to use as a mask for drawing points

#### Value

A data frame of points drawn at random from circular buffers around the occurrence points.

background.raster.buffer

Takes a set of points, a buffer radius, and a mask and returns a raster based on that buffer radius. Code modified from Elith and Hijmans SDM with R tutorial

#### **Description**

NOTE: This function has been replaced by background.buffer.

#### Usage

background.raster.buffer(points, radius, mask)

# Arguments

points A two column data frame with X and Y coordinates

radius Radius for circular buffers to draw around points, in meters.

mask A raster to use as a mask

#### Value

A raster object with values of 1 in every grid cell falling within the buffer.

### **Examples**

library(ENMTools)
background.raster.buffer(iberolacerta.clade\$species\$cyreni\$presence.points, 100000, euro.worldclim)

background.shape.buffer

Takes a set of points and a buffer radius, and returns a polygon. Code modified from Elith and Hijmans SDM with R tutorial

# **Description**

NOTE: This function has been replaced by background.buffer.

#### Usage

background.shape.buffer(points, radius)

#### **Arguments**

points A two column data frame with X and Y coordinates

radius Radius for circular buffers to draw around points, in meters.

#### Value

A polygon shapefile.

background.test

background.test Conduct a background test (also called a similarity test), as described in Warren et al. 2008. This test can either be run as an asymmetric test (species.1 vs background of species.2) or as a symmetric test (background of species.1 vs background of species.2). For GLM, Bioclim, and Domain models the replicates will be constructed from the background points supplied for each species. For Maxent, the replicates will be constructed from the range rasters stored in the enmtools.species objects.

8 background.test

# **Description**

background.test Conduct a background test (also called a similarity test), as described in Warren et al. 2008. This test can either be run as an asymmetric test (species.1 vs background of species.2) or as a symmetric test (background of species.1 vs background of species.2). For GLM, Bioclim, and Domain models the replicates will be constructed from the background points supplied for each species. For Maxent, the replicates will be constructed from the range rasters stored in the enmtools.species objects.

# Usage

```
background.test(
   species.1,
   species.2,
   env,
   type,
   f = NULL,
   nreps = 99,
   test.type = "asymmetric",
   nback = 1000,
   bg.source = "default",
   low.memory = FALSE,
   rep.dir = NA,
   verbose = FALSE,
   clamp = TRUE,
   ...
)
```

# **Arguments**

species.1	An emtools.species object from which presence points (asymmetric) or background (symmetric) will be sampled.
species.2	An enmtools.species object from which background will be sampled.
env	A SpatRaster object containing environmental data
type	The type of model to construct, currently accepts "glm", "mx", "bc", "gam", or "dm" $$
f	A function to use for model fitting. Only required for GLM models at the moment.
nreps	Number of replicates to perform
test.type	Controls whether the background test will be "symmetric" or "asymmetric"
nback	Number of background points for models
bg.source	Source for drawing background points. If "points", it just uses the background points that are already in the species object. If "range", it uses the range raster. If "env", it draws points at randome from the entire study area outlined by the first environmental layer.

calc.B1

low.memory	When set to TRUE, replicate models are written to disc instead of being stored in the output object. Replicate models stored in the output object contain paths to the replicate models on disk instead of the rasters themselves.
rep.dir	Directory for storing replicate models when low.memory is set to TRUE. If not specified, the working directory will be used.
verbose	Controls printing of various messages progress reports. Defaults to FALSE.
clamp	Controls whether empirical and replicate models should be clamped to the environment space used for training.
	Additional arguments to be passed to model fitting functions.

#### Value

results A list containing replicates, models for the empirical data, and summary statistics and plots.

# **Examples**

```
cyreni <- iberolacerta.clade$species$cyreni
monticola <- iberolacerta.clade$species$monticola
cyreni$range <- background.raster.buffer(cyreni$presence.points, 100000, euro.worldclim)
monticola$range <- background.raster.buffer(monticola$presence.points, 100000, euro.worldclim)
background.test(cyreni, monticola, env = euro.worldclim, type = "glm",
f = pres ~ bio1 + bio12, nreps = 10)</pre>
```

calc.B1 calc.B1, Calculates standardized version of Levins (1968) B1 measure of niche breadth given a vector of suitabilities

#### Description

calc.B1, Calculates standardized version of Levins (1968) B1 measure of niche breadth given a vector of suitabilities

### Usage

```
calc.B1(x)
```

# **Arguments**

x A numeric vector

# Value

B1 A calculation of Levins (1968) B1 metric

#### **Examples**

```
calc.B1(c(1, .001, .001))
```

10 check.bg

calc.B2

calc.B2, Calculates standardized version of Levins (1968) B2 measure of niche breadth given a vector of suitabilities

# **Description**

calc.B2, Calculates standardized version of Levins (1968) B2 measure of niche breadth given a vector of suitabilities

# Usage

```
calc.B2(x)
```

# Arguments

Х

A numeric vector

### Value

B2 A calculation of Levins (1968) B2 metric

# **Examples**

```
calc.B2(c(1, .001, .001))
```

check.bg

Takes an emtools.species object and adds background points if they're missing. Looks for range raster first, then goes for environmental layers

# **Description**

Takes an emtools.species object and adds background points if they're missing. Looks for range raster first, then goes for environmental layers.

# Usage

```
check.bg(
  species,
  env = NA,
  nback = 1000,
  bg.source = "default",
  verbose = FALSE,
  bias = NA
)
```

check.clade 11

#### **Arguments**

species An enmtools.species object

env A SpatRaster containing environmental data.

nback Number of background points to generate, if any

bg. source Source for drawing background points. If "points", it just uses the background

points that are already in the species object. If "range", it uses the range raster. If "env", it draws points at randome from the entire study area outlined by the

first environmental layer.

verbose Controls printing of various messages progress reports. Defaults to FALSE.

bias A SpatRaster representing estimated relative sampling bias. Used when bg.source

is either "range" or "env".

#### Value

An enmtools.species object with background points.

check.clade Checking for complians of an enmtools.clade object

# Description

Checking for complians of an enmtools.clade object

# Usage

```
check.clade(this.clade)
```

# **Arguments**

this.clade An enmtools.clade object

#### Value

An enmtools.clade object with appropriate formatting.

#### **Examples**

```
check.clade(iberolacerta.clade)
```

12 check.extras

check.env  Automating some basic tasks for using a raster stack for modeling.  Checks rasters for same extent and resolution, and sets values in each layer to NA if there is an NA in any other layer.	Checks rasters for same extent and resolution, and sets values in each

# **Description**

Automating some basic tasks for using a raster stack for modeling. Checks rasters for same extent and resolution, and sets values in each layer to NA if there is an NA in any other layer.

### Usage

```
check.env(env, verbose = FALSE)
```

### Arguments

env A stack of environmental rasters

verbose Controls printing of progress messages

#### Value

A raster stack.

### **Examples**

```
check.env(euro.worldclim)
```

check.extras

Check that the extra packages needed for an ENMTools function are installed and available

# Description

Check that the extra packages needed for an ENMTools function are installed and available

#### Usage

```
check.extras(funs = NULL)
```

# **Arguments**

funs

character vector of ENMTools function names to install extra dependencies for. If 'NULL', extras for all ENMTools functions will be installed.

#### Value

Logical, 'TRUE' if all required extra packages are available, 'FALSE' otherwise

check.species 13

### **Examples**

```
check.extras("enmtools.gam")
```

check.species

Checking compliance for an object of class enmtools.species.

# Description

Checking compliance for an object of class enmtools.species.

# Usage

```
check.species(this.species, env = NA, trim.dupes = FALSE)
```

#### **Arguments**

this.species An enmtools.species object to be checked.

env Environmental rasters that will be used for modeling. If provided to check.species,

ENMTools will remove occurrence points that have NA values for any layer in

env.

trim.dupes Controls whether to trim duplicate occurrence points from the presence data.

Defaults to FALSE, which leaves duplicates in place. Alternatives are "exact", which will remove points with the same lat/long as another point, or "grid", which will trim data so that there is at most one point per grid cell for the rasters

in env, and centers those points in the cells.

#### Value

An enmtools.species object with appropriate formatting.

# **Examples**

check.species(iberolacerta.clade\$species\$monticola)

clamp.env

Takes an emtools.model object and a set of environment layers and clamps the environment layers so that no variable falls outside of the range available in the training data.

### **Description**

Takes an emtools.model object and a set of environment layers and clamps the environment layers so that no variable falls outside of the range available in the training data.

14 combine.species

#### Usage

```
clamp.env(model, env)
```

### **Arguments**

model An enmtools.model object. Alternatively the analysis.df component of an enm-

tools.model object.

env A SpatRaster of environmental data.

#### Value

An enmtools model object containing species name, model formula (if any), model object, suitability raster, marginal response plots, and any evaluation objects that were created.

### **Examples**

```
monticola.gam <- enmtools.gam(iberolacerta.clade$species$monticola, euro.worldclim[[c(1,5,9,13)]])
euro.clamped <- clamp.env(monticola.gam, euro.worldclim)
clamped.prediction <- predict(monticola.gam, euro.clamped)
terra::plot(clamped.prediction$suitability - monticola.gam$suitability)</pre>
```

combine.species

Takes a list of enmtools.species objects and combines them into a single enmtools.species object

# Description

Takes a list of enmtools.species objects and combines them into a single enmtools.species object

#### Usage

```
combine.species(species.list)
```

# **Arguments**

species.list List of enmtools.species objects that you want to combine together

# Value

An enmtools species object with the occurrence data, names, and range rasters for the species list combined into one.

# **Examples**

```
combine.species(iberolacerta.clade$species)
```

drop.species 15

drop.species	Takes a an ENMTools clade object and a vector of species names. Drops the species from the tree and removes data from the clade object.

# **Description**

Takes a an ENMTools clade object and a vector of species names. Drops the species from the tree and removes data from the clade object.

# Usage

```
drop.species(clade, species)
```

# **Arguments**

clade An ENMTools clade object

species A name or vector of names of species within the enmtools.clade object.

#### Value

An enmtools.clade object with the provided species dropped both from the tree and from the set of enmtools.species objects.

# **Examples**

```
if(requireNamespace("ape", quietly = TRUE)) {
    new.clade <- drop.species(iberolacerta.clade, c("cyreni", "monticola"))
}</pre>
```

enmtools.aoc

Takes an overlap matrix and a tree and runs permutation tests to determine the statistical significance of the relationship between overlap and time

# **Description**

Takes an overlap matrix and a tree and runs permutation tests to determine the statistical significance of the relationship between overlap and time

16 enmtools.bc

#### Usage

```
enmtools.aoc(
  clade,
  env = NULL,
  overlap.source,
  nreps = 100,
  f = NULL,
  overlap.matrix = NULL,
  metric = "D",
   ...
)
```

#### **Arguments**

clade An enmtools.clade object containing species data and a phylogeny

env Environmental layers for use when overlap is calculated using niche models.

overlap.source The source of the overlaps to calculate. Choices are "bc", "dm", "gam", "glm",

"mx", "range", and "point"

nreps A number of reps to do

f The model to be used for GLM and GAM comparisons

overlap.matrix A matrix of overlaps to use, for option overlap.source = "matrix"

metric The overlap metric to use. For ENM sources, this can be any combination of

"D", "I", "cor", "env.D", "env.I", and "env.cor". for range and point overlap this

argument is ignored.

... Arguments to be passed to modeling functions for ENM-based overlaps.

#### Value

A list containing a data frame of coefficients from the empirical regression of overlap on time along with the coefficients from all Monte Carlo replicates, along with plots and p values for the accompanying statistical tests.

enmtools.bc Takes an emtools.species object with presence and background points, and builds a Bioclim model

#### **Description**

Takes an emtools species object with presence and background points, and builds a Bioclim model

enmtools.bc 17

#### Usage

```
enmtools.bc(
   species,
   env = NA,
   test.prop = 0,
   report = NULL,
   overwrite = FALSE,
   nback = 1000,
   env.nback = 10000,
   rts.reps = 0,
   bg.source = "default",
   verbose = FALSE,
   clamp = TRUE,
   corner = NA,
   bias = NA,
   ...
)
```

#### Arguments

species An enmtools.species object

env A SpatRaster of environmental data.

test.prop Proportion of data to withhold randomly for model evaluation, or "block" for

spatially structured evaluation.

report Optional name of an html file for generating reports

overwrite TRUE/FALSE whether to overwrite a report file if it already exists

nback Number of background points for models. In the case of bioclim models these

are only used for evaluation.

env.nback Number of points to draw from environment space for environment space dis-

crimination metrics.

rts.reps The number of replicates to do for a Raes and ter Steege-style test of significance

bg. source Source for drawing background points. If "points", it just uses the background

points that are already in the species object. If "range", it uses the range raster. If "env", it draws points at randome from the entire study area outlined by the

first environmental layer.

verbose Controls printing of various messages progress reports. Defaults to FALSE.

clamp When set to TRUE, clamps the environmental layers so that predictions made

outside the min/max of the training data for each predictor are set to the value for the min/max for that predictor. Prevents the model from extrapolating beyond the min/max bounds of the predictor space the model was trained in, although there could still be projections outside the multivariate training space if predic-

tors are strongly correlated.

corner An integer from 1 to 4. Selects which corner to use for "block" test data. By

default the corner is selected randomly.

18 enmtools.calibrate

bias	An optional raster estimating relative sampling effort per grid cell. Will be used
	for drawing background data.
	Arguments to be passed to bioclim()

# Value

An enmtools model object containing species name, model formula (if any), model object, suitability raster, marginal response plots, and any evaluation objects that were created.

### **Examples**

```
enmtools.bc(iberolacerta.clade$species$monticola, env = euro.worldclim)
```

enmtools.calibrate

Takes an emtools.model object, and reformats it to run through the CalibratR package, calculates Continuous Boyce Index, and runs a Hosmer-Lemeshow goodness-of-fit test. Can either do a full CalibratR run or just return ECE/MCE statistics and plots.

# Description

Takes an emtools.model object, and reformats it to run through the CalibratR package, calculates Continuous Boyce Index, and runs a Hosmer-Lemeshow goodness-of-fit test. Can either do a full CalibratR run or just return ECE/MCE statistics and plots.

# Usage

```
enmtools.calibrate(
  model,
  recalibrate = FALSE,
  cuts = 11,
  env = NA,
  n.background = 10000,
  recal.methods = c(1, 2, 3, 4),
  ...
)
```

#### **Arguments**

model	An enmtools.model object
recalibrate	When TRUE, does a full CalibratR "calibrate" run to recalibrate the model. When FALSE, just returns metrics and plots measuring calibration of the model as is.
cuts	The number of bins to split suitability scores into for calculating calibration.
env	A set of environment layers to be used for optional env space metrics

enmtools.clade 19

n. background Number of background points to be used for env space metrics

Numeric or vector specifying which methods the CalibratR package should use for recalibration. 1=hist\_scaled, 2=hist\_transformed, 3=BBQ\_scaled, 4=BBQ\_transformed. Option 5 is disabled by default due to errors.

... Further arguments to be passed to CalibratR's "calibrate" function.

#### Value

An enmtools.calibrate object containing calibration and classificaction plots, metrics of model calibration, and (optionally) versions of the model that were recalibrated using various methods.

# **Examples**

enmtools.clade

Defining a class for enmtools.clade. Each clade gets:

# **Description**

Defining a class for enmtools.clade. Each clade gets:

### Usage

```
enmtools.clade(species = NA, tree = NA)
save.enmtools.clade(x, ...)
load.enmtools.clade(x, ...)
```

#### **Arguments**

species A list of enmtools.species objects

tree A tree showing the relationships between the species

x Path to an enmtools.species file

Other arguments passed to R's load function

20 enmtools.dm

#### Value

An enmtools.clade object, either empty or containing a formatted version of the data that was passed into the function.

# **Functions**

- save.enmtools.clade(): Save an ENMTools clade object
- load.enmtools.clade(): Load an ENMTools clade object

enmtools.dm Takes an emtools.species object with presence and background points, and builds a Domain model

# Description

Takes an emtools.species object with presence and background points, and builds a Domain model

# Usage

```
enmtools.dm(
   species,
   env = NA,
   test.prop = 0,
   report = NULL,
   nback = 10000,
   env.nback = 100000,
   overwrite = FALSE,
   rts.reps = 0,
   bg.source = "default",
   verbose = FALSE,
   clamp = TRUE,
   corner = NA,
   bias = NA,
   ...
)
```

#### **Arguments**

species	An enmtools.species object
env	A SpatRaster of environmental data.
test.prop	Proportion of data to withhold randomly for model evaluation, or "block" for spatially structured evaluation.
report	Optional name of an html file for generating reports
nback	Number of background points for models. In the case of Domain, these are only used for evaluation.

enmtools.ecospat.bg 21

env.nback	Number of points to draw from environment space for environment space discrimination metrics.
overwrite	TRUE/FALSE whether to overwrite a report file if it already exists
rts.reps	The number of replicates to do for a Raes and ter Steege-style test of significance
bg.source	Source for drawing background points. If "points", it just uses the background points that are already in the species object. If "range", it uses the range raster. If "env", it draws points at randome from the entire study area outlined by the first environmental layer.
verbose	Controls printing of various messages progress reports. Defaults to FALSE.
clamp	When set to TRUE, clamps the environmental layers so that predictions made outside the min/max of the training data for each predictor are set to the value for the min/max for that predictor. Prevents the model from extrapolating beyond the min/max bounds of the predictor space the model was trained in, although there could still be projections outside the multivariate training space if predictors are strongly correlated.
corner	An integer from 1 to 4. Selects which corner to use for "block" test data. By default the corner is selected randomly.
bias	An optional raster estimating relative sampling effort per grid cell. Will be used for drawing background data.
	Arguments to be passed to domain()

# Value

An enmtools model object containing species name, model formula (if any), model object, suitability raster, marginal response plots, and any evaluation objects that were created.

# **Examples**

enmtools.dm(iberolacerta.clade\$species\$monticola, env = euro.worldclim)

enmtools.ecospat.bg enmtools.ecospat.bg, Runs an ecospat background/similarity test using enmtool.species objects.

# Description

enmtools.ecospat.bg, Runs an ecospat background/similarity test using enmtool.species objects.

22 enmtools.ecospat.bg

# Usage

```
enmtools.ecospat.bg(
   species.1,
   species.2,
   env,
   nreps = 99,
   layers = NULL,
   test.type = "symmetric",
   th.sp = 0,
   th.env = 0,
   R = 100,
   nback = 1000,
   bg.source = "default",
   verbose = FALSE,
   ...
)
```

# Arguments

An enmtools.species object
An enmtools.species object
A set of environmental layers
The number of pseudoreplicates to perform
A vector of length 2 containing the names of the layers to be used. If no layer names are provided and there are more than two layers in env, enmtools will perform a pca and use the top two layers from that.
Symmetric or asymmetric test. An asymmetric test is bguivalent to the "one.sided" option in the ecospat documentation, while a symmetric one would be two-sided.
Quantile of species densities used as a threshold to exclude low species density values. See documentation for ecospat.grid.clim.dyn.
Quantile of environmental densities across study sites used as threshold to exclude low environmental density values. See documentation for ecospat.grid.clim.dyn.
Resolution of the grid. See documentation for ecospat.grid.clim.dyn.
Number of background points to use for density calculations.
Source for drawing background points. If "points", it just uses the background points that are already in the species object. If "range", it uses the range raster. If "env", it draws points at randome from the entire study area outlined by the first environmental layer.
Controls printing of various messages progress reports. Defaults to FALSE.
Further arguments to be passed to check.bg

#### Value

A list containing the ecospat output kernel density estimates for each species and their background, as well as the results of hypothesis tests and their accompanying plots.

enmtools.ecospat.id 23

# **Examples**

```
#install.extras(repos='http://cran.us.r-project.org')
monticola <- iberolacerta.clade$species$monticola
cyreni <- iberolacerta.clade$species$cyreni
if(check.extras("enmtools.ecospat.bg")) {
  enmtools.ecospat.bg(monticola, cyreni, euro.worldclim[[1:2]], nback = 500)
}</pre>
```

enmtools.ecospat.id enmtools.ecospat.id, Runs an ecospat identity test using enmtools.pecies objects.

#### **Description**

enmtools.ecospat.id, Runs an ecospat identity test using enmtool.species objects.

# Usage

```
enmtools.ecospat.id(
  species.1,
  species.2,
  env,
  nreps = 99,
  layers = NULL,
  th.sp = 0,
  th.env = 0,
  R = 100,
  nback = 1000,
  bg.source = "default",
  verbose = FALSE
)
```

# Arguments

An enmtools.species object
An enmtools.species object
A set of environmental layers
The number of pseudoreplicates to perform
A vector of length 2 containing the names of the layers to be used. If no layer names are provided and there are more than two layers in env, enmtools will perform a pca and use the top two layers from that.
Quantile of species densities used as a threshold to exclude low species density values. See documentation for ecospat.grid.clim.dyn.
Quantile of environmental densities across study sites used as threshold to exclude low environmental density values. See documentation for ecospat.grid.clim.dyn.

24 enmtools.gam

R	Resolution of the grid. See documentation for ecospat.grid.clim.dyn.
nback	Number of background points to use for density calculations.
bg.source	Source for drawing background points. If "points", it just uses the background points that are already in the species object. If "range", it uses the range raster. If "env", it draws points at randome from the entire study area outlined by the first environmental layer.
verbose	Controls printing of various messages progress reports. Defaults to FALSE.

# Value

A list containing the ecospat output kernel density estimates for each species and their background, as well as the results of hypothesis tests and their accompanying plots.

# **Examples**

```
#install.extras(repos='http://cran.us.r-project.org')
monticola <- iberolacerta.clade$species$monticola
cyreni <- iberolacerta.clade$species$cyreni
if(check.extras("enmtools.ecospat.id")) {
  enmtools.ecospat.id(monticola, cyreni, euro.worldclim[[1:2]], nback = 500)
}</pre>
```

enmtools.gam

Takes an emtools.species object with presence and background points, and builds a gam

# **Description**

Takes an emtools.species object with presence and background points, and builds a gam

# Usage

```
enmtools.gam(
  species,
  env,
  f = NULL,
  test.prop = 0,
  k = 4,
  nback = 1000,
  env.nback = 10000,
  report = NULL,
  overwrite = FALSE,
  rts.reps = 0,
  weights = "equal",
  gam.method = "REML",
  gam.select = TRUE,
```

enmtools.gam 25

```
bg.source = "default",
verbose = FALSE,
clamp = TRUE,
corner = NA,
bias = NA,
...
)
```

#### Arguments

species An enmtools.species object

env A SpatRaster of environmental data.

f Standard gam formula

test.prop Proportion of data to withhold randomly for model evaluation, or "block" for

spatially structured evaluation.

k Dimension of the basis used to represent the smooth term. See documentation

for s() for details.

nback Number of background points to draw from range or env, if background points

aren't provided

env.nback Number of points to draw from environment space for environment space dis-

crimination metrics.

report Optional name of an html file for generating reports

overwrite TRUE/FALSE whether to overwrite a report file if it already exists

rts.reps The number of replicates to do for a Raes and ter Steege-style test of significance

weights If this is set to "equal", presences and background data will be assigned weights

so that the sum of all presence points weights equals the sum of all background

point weights. Otherwise, weights are not provided to the model.

gam. method Defaults to restricted maximum likelihood to facilitate predictor selection, but if

you want to use another method you can pass anything here that gam's "method"

argument understands.

gam.select Controls whether gam algorithm attempts to optimize smoothness and reduce

model complexity. See help("gam.selection") for details.

bg.source Source for drawing background points. If "points", it just uses the background

points that are already in the species object. If "range", it uses the range raster. If "env", it draws points at randome from the entire study area outlined by the

first environmental layer.

verbose Controls printing of various messages progress reports. Defaults to FALSE.

clamp When set to TRUE, clamps the environmental layers so that predictions made

outside the min/max of the training data for each predictor are set to the value for the min/max for that predictor. Prevents the model from extrapolating beyond the min/max bounds of the predictor space the model was trained in, although there could still be projections outside the multivariate training space if predic-

tors are strongly correlated.

26 enmtools.glm

corner	An integer from 1 to 4. Selects which corner to use for "block" test data. By default the corner is selected randomly.
bias	An optional raster estimating relative sampling effort per grid cell. Will be used for drawing background data.
	Arguments to be passed to gam()

#### Value

An enmtools model object containing species name, model formula (if any), model object, suitability raster, marginal response plots, and any evaluation objects that were created.

# **Examples**

```
if(requireNamespace("mgcv", quietly = TRUE)) {
    enmtools.gam(iberolacerta.clade$species$monticola, env = euro.worldclim, f = pres ~ bio1 + bio9)
}
enmtools.glm

Takes an enmtools.species object with presence and background
    points, and builds a GLM
```

# Description

Takes an enmtools.species object with presence and background points, and builds a GLM

### Usage

```
enmtools.glm(
  species,
  env,
  f = NULL
  test.prop = 0,
  eval = TRUE,
  nback = 1000,
  env.nback = 10000,
  report = NULL,
  overwrite = FALSE,
  rts.reps = 0,
  weights = "equal",
  bg.source = "default",
  verbose = FALSE,
  clamp = TRUE,
  corner = NA,
  bias = NA,
  step = FALSE,
)
```

enmtools.glm 27

#### **Arguments**

species An enmtools.species object A SpatRaster of environmental data. env Standard GLM formula test.prop Proportion of data to withhold randomly for model evaluation, or "block" for spatially structured evaluation. eval Determines whether model evaluation should be done. Turned on by default, but moses turns it off to speed things up. nback Number of background points to draw from range or env, if background points aren't provided env.nback Number of points to draw from environment space for environment space discrimination metrics. Optional name of an html file for generating reports report overwrite TRUE/FALSE whether to overwrite a report file if it already exists The number of replicates to do for a Raes and ter Steege-style test of significance rts.reps If this is set to "equal", presences and background data will be assigned weights weights so that the sum of all presence points weights equals the sum of all background point weights. Otherwise, weights are not provided to the model. bg.source Source for drawing background points. If "points", it just uses the background points that are already in the species object. If "range", it uses the range raster. If "env", it draws points at randome from the entire study area outlined by the first environmental layer. verbose Controls printing of various messages progress reports. Defaults to FALSE. clamp When set to TRUE, clamps the environmental layers so that predictions made outside the min/max of the training data for each predictor are set to the value for the min/max for that predictor. Prevents the model from extrapolating beyond the min/max bounds of the predictor space the model was trained in, although there could still be projections outside the multivariate training space if predictors are strongly correlated. corner An integer from 1 to 4. Selects which corner to use for "block" test data. By default the corner is selected randomly. bias An optional raster estimating relative sampling effort per grid cell. Will be used for drawing background data.

#### Value

step

. . .

An enmtools model object containing species name, model formula (if any), model object, suitability raster, marginal response plots, and any evaluation objects that were created.

Arguments to be passed to glm()

#### **Examples**

enmtools.glm(iberolacerta.clade\$species\$monticola, env = euro.worldclim, f = pres ~ bio1 + bio9)

Logical determining whether to do stepwise model selection or not

enmtools.hypervolume

THIS FUNCTION IS CURRENTLY DISABLED. Takes an emtools.species object and environmental layers, and constructs a hypervolume using the R package hypervolume

# **Description**

THIS FUNCTION IS CURRENTLY DISABLED. Takes an emtools.species object and environmental layers, and constructs a hypervolume using the R package hypervolume

### Usage

```
enmtools.hypervolume(
   species,
   env,
   samples.per.point = 10,
   reduction.factor = 0.1,
   method = "gaussian",
   verbose = FALSE,
   clamp = TRUE,
   ...
)
```

#### **Arguments**

species An enmtools.species object
env A stack of environmental rasters
samples.per.point
To be passed to hypervolume\_gaussian
reduction.factor

To be passed to hypervolume\_project

method Method for constructing hypervolumes, defaults to "gaussian"

verbose Controls printing of various messages progress reports. Defaults to FALSE.

clamp When set to TRUE, clamps the environmental layers so that predictions made

outside the min/max of the training data for each predictor are set to the value for the min/max for that predictor. Prevents the model from extrapolating beyond the min/max bounds of the predictor space the model was trained in, although there could still be projections outside the multivariate training space if predic-

tors are strongly correlated.

... Extra parameters to be passed to hypervolume\_gaussian

#### Value

An enmtools hypvervolume object containing a hypervolume object, a raster of suitability scores, the species name, and the occurrence data frame.

enmtools.maxent 29

### **Examples**

```
#install.extras(repos='http://cran.us.r-project.org')
env <- euro.worldclim[[c(1,8,12,17)]]
if(requireNamespace("hypervolume", quietly = TRUE)) {
    monticola.hv <- enmtools.hypervolume(iberolacerta.clade$species$monticola, env = env)
}</pre>
```

enmtools.maxent

Takes an emtools.species object with presence and background points, and builds a maxent model

# **Description**

Takes an emtools.species object with presence and background points, and builds a maxent model

# Usage

```
enmtools.maxent(
   species,
   env,
   test.prop = 0,
   nback = 1000,
   env.nback = 10000,
   report = NULL,
   overwrite = FALSE,
   rts.reps = 0,
   bg.source = "default",
   verbose = FALSE,
   clamp = TRUE,
   corner = NA,
   bias = NA,
   ...
)
```

# **Arguments**

species	An enmtools.species object
env	A SpatRaster of environmental data.
test.prop	Proportion of data to withhold randomly for model evaluation, or "block" for spatially structured evaluation.
nback	Number of background points to draw from range or env, if background points aren't provided
env.nback	Number of points to draw from environment space for environment space discrimination metrics.
report	Optional name of an html file for generating reports

30 enmtools.rf

overwrite	TRUE/FALSE whether to overwrite a report file if it already exists
rts.reps	The number of replicates to do for a Raes and ter Steege-style test of significance
bg.source	Source for drawing background points. If "points", it just uses the background points that are already in the species object. If "range", it uses the range raster. If "env", it draws points at randome from the entire study area outlined by the first environmental layer.
verbose	Controls printing of various messages progress reports. Defaults to FALSE.
clamp	When set to TRUE, clamps the environmental layers so that predictions made outside the min/max of the training data for each predictor are set to the value for the min/max for that predictor. Prevents the model from extrapolating beyond the min/max bounds of the predictor space the model was trained in, although there could still be projections outside the multivariate training space if predictors are strongly correlated.
corner	An integer from 1 to 4. Selects which corner to use for "block" test data. By default the corner is selected randomly.
bias	An optional raster estimating relative sampling effort per grid cell. Will be used for drawing background data.
	Arguments to be passed to maxent()

#### Value

An enmtools model object containing species name, model formula (if any), model object, suitability raster, marginal response plots, and any evaluation objects that were created.

# **Examples**

```
if(check.extras("enmtools.maxent")) {
    ## maxent is not working on some platforms so use try()
    try(enmtools.maxent(iberolacerta.clade$species$monticola, env = euro.worldclim))
}
```

 ${\tt enmtools.rf}$ 

Takes an emtools.species object with presence and background points, and builds a random forest model

# **Description**

Takes an emtools.species object with presence and background points, and builds a random forest model

enmtools.rf 31

# Usage

```
enmtools.rf(
  species,
  env,
  f = NULL,
  test.prop = 0,
  eval = TRUE,
  nback = 1000,
  env.nback = 10000,
  report = NULL,
  overwrite = FALSE,
  rts.reps = 0,
  bg.source = "default",
  verbose = FALSE,
  clamp = TRUE,
  corner = NA,
  bias = NA,
)
```

# **Arguments**

species

env A SpatRaster of environmental data.  f A formula for fitting the model  test.prop Proportion of data to withhold randomly for model evaluation, or "block" for spatially structured evaluation.  eval Determines whether model evaluation should be done. Turned on by default, but moses turns it off to speed things up.  nback Number of background points to draw from range or env, if background points aren't provided  env.nback Number of points to draw from environment space for environment space discrimination metrics.  report Optional name of an html file for generating reports  overwrite TRUE/FALSE whether to overwrite a report file if it already exists  rts.reps The number of replicates to do for a Raes and ter Steege-style test of significance bg.source Source for drawing background points. If "points", it just uses the background points that are already in the species object. If "range", it uses the range raster. If "env", it draws points at randome from the entire study area outlined by the first environmental layer.  verbose Controls printing of various messages progress reports. Defaults to FALSE.  clamp When set to TRUE, clamps the environmental layers so that predictions made outside the min/max of the training data for each predictor are set to the value for	•	1 3
Proportion of data to withhold randomly for model evaluation, or "block" for spatially structured evaluation.  eval Determines whether model evaluation should be done. Turned on by default, but moses turns it off to speed things up.  Number of background points to draw from range or env, if background points aren't provided  env.nback Number of points to draw from environment space for environment space discrimination metrics.  report Optional name of an html file for generating reports  overwrite TRUE/FALSE whether to overwrite a report file if it already exists  rts.reps The number of replicates to do for a Raes and ter Steege-style test of significance bg.source Source for drawing background points. If "points", it just uses the background points that are already in the species object. If "range", it uses the range raster. If "env", it draws points at randome from the entire study area outlined by the first environmental layer.  verbose Controls printing of various messages progress reports. Defaults to FALSE.  clamp When set to TRUE, clamps the environmental layers so that predictions made	env	A SpatRaster of environmental data.
petermines whether model evaluation should be done. Turned on by default, but moses turns it off to speed things up.  Number of background points to draw from range or env, if background points aren't provided  env.nback  Number of points to draw from environment space for environment space discrimination metrics.  report  Optional name of an html file for generating reports  overwrite  TRUE/FALSE whether to overwrite a report file if it already exists  rts.reps  The number of replicates to do for a Raes and ter Steege-style test of significance bg. source  Source for drawing background points. If "points", it just uses the background points that are already in the species object. If "range", it uses the range raster. If "env", it draws points at randome from the entire study area outlined by the first environmental layer.  Verbose  Controls printing of various messages progress reports. Defaults to FALSE.  Clamp  When set to TRUE, clamps the environmental layers so that predictions made	f	A formula for fitting the model
but moses turns it off to speed things up.  Number of background points to draw from range or env, if background points aren't provided  env.nback  Number of points to draw from environment space for environment space discrimination metrics.  report  Optional name of an html file for generating reports  overwrite  TRUE/FALSE whether to overwrite a report file if it already exists  rts.reps  The number of replicates to do for a Raes and ter Steege-style test of significance  bg.source  Source for drawing background points. If "points", it just uses the background points that are already in the species object. If "range", it uses the range raster. If "env", it draws points at randome from the entire study area outlined by the first environmental layer.  verbose  Controls printing of various messages progress reports. Defaults to FALSE.  clamp  When set to TRUE, clamps the environmental layers so that predictions made	test.prop	· · · · · · · · · · · · · · · · · · ·
env.nback  Number of points to draw from environment space for environment space discrimination metrics.  report  Optional name of an html file for generating reports  overwrite  TRUE/FALSE whether to overwrite a report file if it already exists  rts.reps  The number of replicates to do for a Raes and ter Steege-style test of significance  bg.source  Source for drawing background points. If "points", it just uses the background points that are already in the species object. If "range", it uses the range raster. If "env", it draws points at randome from the entire study area outlined by the first environmental layer.  verbose  Controls printing of various messages progress reports. Defaults to FALSE.  clamp  When set to TRUE, clamps the environmental layers so that predictions made	eval	· · · · · · · · · · · · · · · · · · ·
crimination metrics.  report Optional name of an html file for generating reports  overwrite TRUE/FALSE whether to overwrite a report file if it already exists  rts.reps The number of replicates to do for a Raes and ter Steege-style test of significance  bg.source Source for drawing background points. If "points", it just uses the background points that are already in the species object. If "range", it uses the range raster. If "env", it draws points at randome from the entire study area outlined by the first environmental layer.  verbose Controls printing of various messages progress reports. Defaults to FALSE.  clamp When set to TRUE, clamps the environmental layers so that predictions made	nback	
overwrite TRUE/FALSE whether to overwrite a report file if it already exists  rts.reps The number of replicates to do for a Raes and ter Steege-style test of significance  bg.source Source for drawing background points. If "points", it just uses the background points that are already in the species object. If "range", it uses the range raster.  If "env", it draws points at randome from the entire study area outlined by the first environmental layer.  verbose Controls printing of various messages progress reports. Defaults to FALSE.  clamp When set to TRUE, clamps the environmental layers so that predictions made	env.nback	
The number of replicates to do for a Raes and ter Steege-style test of significance bg.source  Source for drawing background points. If "points", it just uses the background points that are already in the species object. If "range", it uses the range raster. If "env", it draws points at randome from the entire study area outlined by the first environmental layer.  Verbose Controls printing of various messages progress reports. Defaults to FALSE. clamp  When set to TRUE, clamps the environmental layers so that predictions made	report	Optional name of an html file for generating reports
bg.source Source for drawing background points. If "points", it just uses the background points that are already in the species object. If "range", it uses the range raster. If "env", it draws points at randome from the entire study area outlined by the first environmental layer.  Verbose Controls printing of various messages progress reports. Defaults to FALSE. clamp When set to TRUE, clamps the environmental layers so that predictions made	overwrite	TRUE/FALSE whether to overwrite a report file if it already exists
points that are already in the species object. If "range", it uses the range raster. If "env", it draws points at randome from the entire study area outlined by the first environmental layer.  Verbose Controls printing of various messages progress reports. Defaults to FALSE. clamp When set to TRUE, clamps the environmental layers so that predictions made	rts.reps	The number of replicates to do for a Raes and ter Steege-style test of significance
clamp When set to TRUE, clamps the environmental layers so that predictions made	bg.source	points that are already in the species object. If "range", it uses the range raster. If "env", it draws points at randome from the entire study area outlined by the
,	verbose	Controls printing of various messages progress reports. Defaults to FALSE.
· · · · · · · · · · · · · · · · · · ·	clamp	• • • • • • • • • • • • • • • • • • • •

the min/max for that predictor. Prevents the model from extrapolating beyond the min/max bounds of the predictor space the model was trained in, although

An enmtools.species object

32 enmtools.rf.ranger

	there could still be projections outside the multivariate training space if predictors are strongly correlated.
corner	An integer from 1 to 4. Selects which corner to use for "block" test data. By default the corner is selected randomly.
bias	An optional raster estimating relative sampling effort per grid cell. Will be used for drawing background data.
	Arguments to be passed to rf()

#### Value

An enmtools model object containing species name, model formula (if any), model object, suitability raster, marginal response plots, and any evaluation objects that were created.

#### **Examples**

```
enmtools.rf(iberolacerta.clade$species$monticola, env = euro.worldclim, nback = 500)
```

enmtools.rf.ranger Takes an emtools.species object with presence and background points, and builds a random forest model using the 'probability mode' in package 'ranger'

### **Description**

Takes an emtools.species object with presence and background points, and builds a random forest model using the 'probability mode' in package 'ranger'

#### Usage

```
enmtools.rf.ranger(
  species,
  env,
  f = NULL,
  test.prop = 0,
  eval = TRUE,
  nback = 1000,
  env.nback = 10000,
  report = NULL,
  overwrite = FALSE,
  rts.reps = 0,
  bg.source = "default",
  verbose = FALSE,
  clamp = TRUE,
  corner = NA,
 bias = NA,
)
```

enmtools.rf.ranger 33

# Arguments

species	An enmtools.species object
env	A SpatRaster of environmental data.
f	A formula for fitting the model
test.prop	Proportion of data to withhold randomly for model evaluation, or "block" for spatially structured evaluation.
eval	Determines whether model evaluation should be done. Turned on by default, but moses turns it off to speed things up.
nback	Number of background points to draw from range or env, if background points aren't provided
env.nback	Number of points to draw from environment space for environment space discrimination metrics.
report	Optional name of an html file for generating reports
overwrite	TRUE/FALSE whether to overwrite a report file if it already exists
rts.reps	The number of replicates to do for a Raes and ter Steege-style test of significance
bg.source	Source for drawing background points. If "points", it just uses the background points that are already in the species object. If "range", it uses the range raster. If "env", it draws points at randome from the entire study area outlined by the first environmental layer.
verbose	Controls printing of various messages progress reports. Defaults to FALSE.
clamp	When set to TRUE, clamps the environmental layers so that predictions made outside the min/max of the training data for each predictor are set to the value for the min/max for that predictor. Prevents the model from extrapolating beyond the min/max bounds of the predictor space the model was trained in, although there could still be projections outside the multivariate training space if predictors are strongly correlated.
corner	An integer from 1 to 4. Selects which corner to use for "block" test data. By default the corner is selected randomly.
bias	An optional raster estimating relative sampling effort per grid cell. Will be used
	for drawing background data.
	for drawing background data.  Arguments to be passed to ranger

# Value

An enmtools model object containing species name, model formula (if any), model object, suitability raster, marginal response plots, and any evaluation objects that were created.

# **Examples**

enmtools.rf.ranger(iberolacerta.clade\$species\$monticola, env = euro.worldclim, nback = 500)

34 enmtools.species

enmtools.species

Defining a class for enmtools.species.

# **Description**

Each species gets:

### Usage

```
enmtools.species(
  range = NA,
  presence.points = NA,
  background.points = NA,
  species.name = NA,
  models = NA
)

save.enmtools.species(x, ...)

load.enmtools.species(x, ...)
```

# **Arguments**

range A raster or SpatialPolygon with the actual range they occur in

presence.points

A data frame with sampled localities

background.points

A data frame with absence/pseudoabsence/background localities

species.name A character vector with the species name

models A list of models that are made for the species, which will be stuffed in there as

we go along to pass the check. This is used by internal enmtools functions to

make sure the necessary data is present before processing anything.

x Path to an enmtools.species file

... Other arguments passed to R's load function

#### Value

Returns an enmtools.species object, either empty or populated with the parameter values that were passed into the function.

#### **Functions**

- save.enmtools.species(): Save an ENMTools species object
- load.enmtools.species(): Load an ENMTools species object

enmtools.vip 35

enmtools.vip	Takes an enmtools.model object, and performs variable importance analyses on it using methods from the vip package
	emanyees on a usung memous from the Apparenage

# Description

Takes an enmtools.model object, and performs variable importance analyses on it using methods from the vip package

# Usage

```
enmtools.vip(
  model,
  metric = "roc_auc",
  nsim = 10,
  method = "permute",
  verbose = FALSE,
  ...
)
```

# **Arguments**

model	An enmtools.model object
metric	The metric to use for measuring how variables affect model predictions
nsim	The number of simulations to be run for method "permute"
method	A character string or vector containing any combination of "model", "permute", "shap", or "firm". For details on what these mean, see the vip package help.
verbose	Controls printing of messages
	Further arguments to be passed to vip's "vi" functions.

# Value

An enmtools.vip object

# Examples

36 env.breadth

env.breadth	Calculates breadth of a model in environment space using latin hypercube sampling

# Description

Calculates breadth of a model in environment space using latin hypercube sampling

# Usage

```
env.breadth(model, env, tolerance = 1e-04, max.reps = 10, chunk.size = 1e+05)
```

# **Arguments**

model	An enmtools.model object or a model object that can be projected using the predict() function
env	A raster, raster stack of environmental data, or a list of minima and maxima for the environmental space to evaluate models over.
tolerance	How close do successive overlap metrics have to be before we decide we're close enough to the final answer
max.reps	Maximum number of attempts that will be made to find suitable starting conditions
chunk.size	How many combinations of environmental variables to try at a time. If your niche breadth in environment space is small, increasing this value may help you get a result.

# Value

A list containing the environmental space version of the B2 metric and a plot of B2 estimates as a function of sample size, used as a convergence diagnostic.

# **Examples**

```
cyreni <- iberolacerta.clade$species$cyreni
cyreni.glm <- enmtools.glm(cyreni, euro.worldclim, test.prop = 0.2,
f = pres ~ bio1 + bio12, nback = 500)
env.breadth(cyreni.glm, euro.worldclim)</pre>
```

env.evaluate 37

env.evaluate	Calculates evaluation metrics (AUC, etc.) using latin hypercube sam-
	pling in environment space

# Description

Calculates evaluation metrics (AUC, etc.) using latin hypercube sampling in environment space

# Usage

```
env.evaluate(
  species,
  model,
  env,
  bg.source = "background",
  n.background = 10000,
  test.eval = FALSE,
  verbose = FALSE,
  ...
)
```

### **Arguments**

species	An enmtools.species object
model	An enmtools.model object or a model that can be projected using the predict() function of dismo
env	A SpatRaster of environmental data.
bg.source	Determines whether minima and maxima of the environment space should be picked using the environment layers or the background points.
n.background	The number of background points to sample from the environment space.
test.eval	When set to "true", env.evaluate evaluates the test data stored in the model object instead of the training data.
verbose	Controls printing of various messages progress reports. Defaults to FALSE.
	Arguments to be passed to other functions

# Value

A dismo evaluate object measuring the performance of model predictions in environment space.

```
cyreni <- iberolacerta.clade$species$cyreni
cyreni.glm <- enmtools.glm(cyreni, euro.worldclim, test.prop = 0.2,
f = pres ~ bio1 + bio12, nback = 500)
env.evaluate(cyreni, cyreni.glm, euro.worldclim)</pre>
```

38 env.overlap

env.overlap	Calculates overlap between models in environment space using latin
ciiv.ovci 1up	hypercube sampling

# Description

Calculates overlap between models in environment space using latin hypercube sampling

# Usage

```
env.overlap(
  model.1,
  model.2,
  env,
  tolerance = 0.001,
  max.reps = 10,
  cor.method = "spearman",
  chunk.size = 1e+05,
  recal.model.1 = NA,
  recal.model.2 = NA,
  verbose = FALSE
)
```

# Arguments

model.1	An enmtools.model object model object that can be projected using the predict() function
model.2	Another enmtools.model object or other model object that can be projected using the predict() function
env	A raster, raster stack of environmental data, or a list of minima and maxima for the environmental space to evaluate models over
tolerance	How close do successive overlap metrics have to be before we decide we're close enough to the final answer
max.reps	Maximum number of attempts that will be made to find suitable starting conditions
cor.method	Which method to use for calculating correlations between models
chunk.size	How many combinations of environmental variables to try at a time. If your niche breadth in environment space is small, increasing this value may help you get a result.
recal.model.1	Optional. The output of enmtools.recalibrate for model 1, which needs to have been run with "recalibrate = TRUE".
recal.model.2	Optional. The output of enmtools.recalibrate for model 2, which needs to have been run with "recalibrate = TRUE".
verbose	Controls printing of various messages progress reports. Defaults to FALSE.

euro.worldclim 39

#### Value

A list of values measuring the overlap between models in environment space, as well as some plots depicting change of the estimates as a function of how many samples were used, which are included as a sort of convergence diagnostic.

## **Examples**

```
cyreni <- iberolacerta.clade$species$cyreni
monticola <- iberolacerta.clade$species$monticola
cyreni.glm <- enmtools.glm(cyreni, euro.worldclim, f = pres ~ bio1 + bio12, nback = 500)
monticola.glm <- enmtools.glm(monticola, euro.worldclim, f = pres ~ bio1 + bio12, nback = 500)
env.overlap(cyreni.glm, monticola.glm, euro.worldclim)</pre>
```

euro.worldclim

Worldclim data for Europe

## **Description**

This is a fairly low-resolution set of Worldclim layers for Europe to be used in demonstrating ENMTools functions.

#### Usage

euro.worldclim

#### **Format**

A SpatRaster object with 19 worldclim layers.

find.extras

Find the extra packages needed for an ENMTools function

#### **Description**

Find the extra packages needed for an ENMTools function

## Usage

```
find.extras(funs = NULL)
```

#### **Arguments**

funs

character vector of ENMTools function names to install extra dependencies for. If 'NULL', extras for all ENMTools functions will be installed.

40 geog.range.overlap

#### Value

A character vector with the names of extra packages required by 'funs'

#### **Examples**

```
find.extras("enmtools.calibrate")
```

find.extras.missing

Find the extra packages needed for an ENMTools function that are currently missing (not available)

## **Description**

Find the extra packages needed for an ENMTools function that are currently missing (not available)

#### Usage

```
find.extras.missing(funs = NULL)
```

#### **Arguments**

funs

character vector of ENMTools function names to install extra dependencies for. If 'NULL', extras for all ENMTools functions will be installed.

#### Value

A character vector with the names of extra packages required by 'funs' that are currently not available

#### **Examples**

```
find.extras.missing("enmtools.calibrate")
```

geog.range.overlap

Takes two emtools.species objects with range rasters, calculates overlap between them as in Fitzpatrick and Turelli 2006. This metric divides the area of overlap between two species ranges by the smaller of the two areas of the species' individual ranges. It therefore ranges from 0 (no overlap) to I (ranges are the same or the smaller species' range is contained entirely within the larger).

#### **Description**

Takes two emtools.species objects with range rasters, calculates overlap between them as in Fitz-patrick and Turelli 2006. This metric divides the area of overlap between two species ranges by the smaller of the two areas of the species' individual ranges. It therefore ranges from 0 (no overlap) to 1 (ranges are the same or the smaller species' range is contained entirely within the larger).

hypervolume.overlap 41

#### Usage

```
geog.range.overlap(x, y)
```

#### **Arguments**

x An enmtools.species object containing a range raster
y An enmtools.species object containing a range raster

#### Value

A numeric value measuring range overlap.

### **Examples**

```
cyreni <- iberolacerta.clade$species$cyreni
monticola <- iberolacerta.clade$species$monticola
cyreni$range <- background.raster.buffer(cyreni$presence.points, 100000, euro.worldclim)
monticola$range <- background.raster.buffer(monticola$presence.points, 100000, euro.worldclim)
geog.range.overlap(cyreni, monticola)</pre>
```

hypervolume.overlap

Takes an emtools.species object and environmental layers, and constructs a hypervolume using the R package hypervolume

## Description

Takes an emtools.species object and environmental layers, and constructs a hypervolume using the R package hypervolume

#### Usage

```
hypervolume.overlap(
  species.1,
  species.2,
  env = NA,
  reduction.factor = 0.1,
  ...
)
```

# **Arguments**

species.1 An enmtools.species or enmtools.hypervolume object
species.2 An enmtools.species or enmtools.hypervolume object
env A stack of environmental rasters, required when enmtools.species objects are passed
reduction.factor
Passed to hypervolume functions
... Extra parameters to be passed to enmtools.hypervolume

42 identity.test

## Value

A list containing a set of hypervolume objects and the overlap that was measured between them.

# **Examples**

```
#install.extras(repos='http://cran.us.r-project.org')
mont <- iberolacerta.clade$species$monticola
cyr <- iberolacerta.clade$species$cyreni
env <- euro.worldclim[[c(1,8,12,17)]]
if(requireNamespace("hypervolume", quietly = TRUE)) {
    hypervolume.overlap(mont, cyr, env = env)
}</pre>
```

iberolacerta.clade

GBIF data for a clade of Iberolacerta lizards from spain

# Description

This is some sample data downloaded from GBIF for the purposes of demonstrating ENMTools functionality.

## Usage

iberolacerta.clade

#### **Format**

An enmtools.clade object with seven species and a phylogeny.

identity.test

identity.test Conduct a niche identity/equivalency test as described in Warren et al. 2008.

## **Description**

identity.test Conduct a niche identity/equivalency test as described in Warren et al. 2008.

identity.test 43

## Usage

```
identity.test(
   species.1,
   species.2,
   env,
   type,
   f = NULL,
   nreps = 99,
   nback = 1000,
   bg.source = "default",
   low.memory = FALSE,
   rep.dir = NA,
   verbose = FALSE,
   clamp = TRUE,
   ...
)
```

# Arguments

species.1	An emtools.species object
species.2	An enmtools.species object
env	A SpatRaster object containing environmental data
type	The type of model to construct, currently accepts "glm", "mx", "bc", "gam", "rf", or "dm"
f	A function to use for model fitting. Only required for GLM models at the moment.
nreps	Number of replicates to perform
nback	Number of background points for models
bg.source	Source for drawing background points. If "points", it just uses the background points that are already in the species object. If "range", it uses the range raster. If "env", it draws points at randome from the entire study area outlined by the first environmental layer.
low.memory	When set to TRUE, replicate models are written to disc instead of being stored in the output object. Replicate models stored in the output object contain paths to the replicate models on disk instead of the rasters themselves.
rep.dir	Directory for storing replicate models when low.memory is set to TRUE. If not specified, the working directory will be used.
verbose	Controls printing of various messages progress reports. Defaults to FALSE.
clamp	Controls whether empirical and replicate models should be clamped to the environment space used for training.
	Additional arguments to be passed to model fitting functions.

## Value

results A list containing the replicates, models for the empirical data, and summary statistics and plots.

44 install.extras

#### **Examples**

```
cyreni <- iberolacerta.clade$species$cyreni
monticola <- iberolacerta.clade$species$monticola
cyreni$range <- background.raster.buffer(cyreni$presence.points, 100000, euro.worldclim)
monticola$range <- background.raster.buffer(monticola$presence.points, 100000, euro.worldclim)
identity.test(cyreni, monticola, env = euro.worldclim, type = "glm",
f = pres ~ bio1 + bio12, nreps = 10)</pre>
```

install.extras

install.extras

#### **Description**

Convenience function that installs all extra packages used in ENMTools. ENMTools uses functions from a lot of external packages, and due to CRAN best practices it doesn't install those packages by default. The function install.extras() calls install.packages with a list of all of the extra packages that are not already available. Use [check.extras()] to find out which packages are needed for a list of functions.

## Usage

```
install.extras(funs = NULL, install = TRUE, ...)
```

#### **Arguments**

funs character vector of ENMTools function names to install extra dependencies for.
If 'NULL', extras for all ENMTools functions will be installed.

Should the packages be installed?

Other parameters to be passed to install.packages

#### Value

The list of missing packages is return invisibly.

```
install.extras(install = FALSE)
```

interactive.plot 45

species	interactive.plot	Generic function for making interactive plots of ENMTools models and species
---------	------------------	--

## **Description**

Function that takes an enmtools.model or enmtools.species object and calls the class-appropriate interactive.plot.xx function for it. These functions use leaflet for mapping and will only function properly if you have an active internet connection.

## Usage

```
interactive.plot(x, ...)
```

#### **Arguments**

- x entools.model or species object to plot
- ... other arguments passed to interactive.plot.enmtools.model or interactive.plot.enmtools.species

#### Value

An interactive leaflet plot depicting the predictions and data from the enmtools.model object.

```
interactive.plot.enmtools.model

Plot an enmtools.model object on an interactive map
```

## **Description**

Function that take an enmtools.model object and plots an interactive map of the presence points, background points (if applicable), and species suitability map. This function uses leaflet for mapping and will only function properly if you have an active internet connection.

#### Usage

```
## S3 method for class 'enmtools.model'
interactive.plot(
    x,
    map.provider = "Esri.WorldPhysical",
    cluster.points = FALSE,
    raster.opacity = 1,
    max.bytes = 4194304,
    ...
)
```

#### **Arguments**

entools.model object to plot Χ map.provider Name of a map provider for the underlying interactive base map. Default is "Esri.WorldPhysical", and attractive topographic map with no place labels. A preview of all map provider options can be viewed at http://leaflet-extras. github.io/leaflet-providers/preview/ cluster.points Should points be clustered? If TRUE, points close together will be grouped into clusters that can be interactively expanded by clicking on them. raster.opacity Specifies the opacity level of the suitability raster. Maximum size of raster image to plot. Defaults to 4MB (4194304 bytes) but can max.bytes be overridden if you have a large raster. Be aware that the image will be knitted into an output file if you're working in R Markdown, causing your output file to be huge if the raster is huge. other arguments (not used currently)

#### Value

An interactive leaflet plot depicting the predictions and data from the enmtools.model object.

```
interactive.plot.enm tools.species\\
```

Plot an enmtools.species object on an interactive map

# Description

Function that take an enmtools.species object and plots an interactive map of the presence points, background points (if applicable), and species range raster (if applicable). This function uses leaflet for mapping and will only function properly if you have an active internet connection.

#### Usage

```
## $3 method for class 'enmtools.species'
interactive.plot(
    x,
    map.provider = "Esri.WorldPhysical",
    cluster.points = FALSE,
    max.bytes = 4194304,
    ...
)
```

marginal.plots 47

#### **Arguments**

x entools.species object to plot

map.provider Name of a map provider for the underlying interactive base map. Default is

"Esri.WorldPhysical", and attractive topographic map with no place labels. A preview of all map provider options can be viewed at http://leaflet-extras.

github.io/leaflet-providers/preview/

cluster.points Should points be clustered? If TRUE, points close together will be grouped into

clusters that can be interactively expanded by clicking on them.

max.bytes Maximum size of range raster image to plot. Defaults to 4MB (4194304 bytes)

but can be overridden if you have a large range raster. Be aware that the image will be knitted into an output file if you're working in R Markdown, causing

your output file to be huge if the raster is huge.

... other arguments (not used currently)

#### Value

An interactive leaflet plot visualizing the data present in the species object.

marginal.plots	marginal.plots Plots the marginal response of a model to an environ-
	mental variable with all other variables held at their mean in env

#### **Description**

marginal.plots Plots the marginal response of a model to an environmental variable with all other variables held at their mean in env

#### Usage

```
marginal.plots(model, env, layer, standardize = TRUE, verbose = FALSE)
```

## **Arguments**

model An enmtools model object

env A SpatRaster object containing environmental data

layer The name of the layer to plot

standardize Whether to set the maximum of the response function to 1, or to instead use the

raw values.

verbose Controls printing of messages

#### Value

results A plot of the marginal response of the model to the environmental variable.

48 moses.list

#### **Examples**

```
cyreni.glm <- enmtools.glm(iberolacerta.clade$species$cyreni,
f = pres ~ bio1 + bio12, euro.worldclim)
marginal.plots(cyreni.glm, euro.worldclim, "bio1")</pre>
```

moses.list

Takes a list of enmtools.species objects and uses model selection to ask whether they're better treated jointly or separately

## **Description**

Takes a list of enmtools.species objects and uses model selection to ask whether they're better treated jointly or separately

## Usage

```
moses.list(
  species.list,
  env,
  f = NULL,
  eval = FALSE,
  nback = 1000,
  bg.source = "default",
  verbose = FALSE,
  step = FALSE,
  ...
)
```

#### **Arguments**

species.list A list of enmtools.species objects, or an enmtools.clade object.

env A SpatRaster of environmental data.

f A GLM-style function for model fitting

eval Boolean indicating whether or not GLMs should be evaluated using AUC/TSS/etc.

nback Number of background points to generate, if any

bg. source Source for drawing background points. If "points", it just uses the background

points that are already in the species object. If "range", it uses the range raster. If "env", it draws points at randome from the entire study area outlined by the

first environmental layer.

verbose Controls printing of various messages progress reports. Defaults to FALSE.

step Logical determining whether to do stepwise model selection or not

... further arguments to be passed to enmtools.glm

#### Value

A list containing GLMs for the groups separately and together, as well as AIC values for those models.

multistack.pca 49

multistack.pca	multistack.pca, simultaneous PCA on more than one stack of environ- mental rasters

## **Description**

multistack.pca, simultaneous PCA on more than one stack of environmental rasters

## Usage

```
multistack.pca(..., n = 2)
```

## **Arguments**

... Any number of environmental raster stacks or bricks

n The number of PCA layers to return

#### Value

A list containing a stack or brick of rasters for each input set representing the top n pca axes of the initial environmental variables, as well as the pca object from the analysis that produced them and some useful plots showing the distribution of each PC in the different stacks.

#### **Examples**

```
test1 <- terra::crop(euro.worldclim, terra::ext(-10, -5, 40, 43))
test2 <- terra::crop(euro.worldclim, terra::ext(-5, 5, 40, 48))
test3 <- terra::crop(euro.worldclim, terra::ext(5, 15, 44, 48))
multistack.pca(test1, test2, test3)</pre>
```

node.overlap

Takes an overlap matrix and a tree and returns average overlap at nodes using Fitzpatrick & Turelli averaging method. Typicall called via enmtools.aoc rather than used directly.

#### **Description**

Takes an overlap matrix and a tree and returns average overlap at nodes using Fitzpatrick & Turelli averaging method. Typicall called via enmtools.aoc rather than used directly.

## Usage

```
node.overlap(overlap, tree)
```

50 point.overlap

### **Arguments**

overlap An overlap matrix

tree A tree

#### Value

A data frame of node ages and overlaps

point.overlap Takes two emtoo

Takes two emtools.species objects with range rasters, calculates overlap between them as in Cardillo and Warren 2016

# Description

Takes two emtools.species objects with range rasters, calculates overlap between them as in Cardillo and Warren 2016

## Usage

```
point.overlap(x, y)
```

## **Arguments**

x An enmtools.species object containing presence points

y An enmtools.species object containing presence points

## Value

A numeric value measuring the overlap between point distributions.

```
cyreni <- iberolacerta.clade$species$cyreni
monticola <- iberolacerta.clade$species$monticola
point.overlap(cyreni, monticola)</pre>
```

rangebreak.blob 51

rangebreak.blob Conduct a blob rangebreak test as described in Glor and Warren 2011.

# Description

rangebreak.blob Conduct a blob rangebreak test as described in Glor and Warren 2011.

# Usage

```
rangebreak.blob(
  species.1,
  species.2,
  env,
  type,
  f = NULL,
  nreps = 99,
  nback = 1000,
  bg.source = "default",
  low.memory = FALSE,
  rep.dir = NA,
  verbose = FALSE,
  clamp = TRUE,
  ...
)
```

# Arguments

species.1	An emtools.species object
species.2	An enmtools.species object
env	A SpatRaster object containing environmental data
type	The type of model to construct, currently accepts "glm", "mx", "bc", "gam", or "dm" $$
f	A function to use for model fitting. Only required for GLM models at the moment.
nreps	Number of replicates to perform
nback	Number of background points for models
bg.source	Source for drawing background points. If "points", it just uses the background points that are already in the species object. If "range", it uses the range raster. If "env", it draws points at randome from the entire study area outlined by the first environmental layer.
low.memory	When set to TRUE, replicate models are written to disc instead of being stored in the output object. Replicate models stored in the output object contain paths to the replicate models on disk instead of the rasters themselves.

52 rangebreak.linear

rep.dir	Directory for storing replicate models when low.memory is set to TRUE. If not specified, the working directory will be used.
verbose	Controls printing of various messages progress reports. Defaults to FALSE.
clamp	Controls whether empirical and replicate models should be clamped to the environment space used for training.
	Additional arguments to be passed to model fitting functions.

#### Value

results A list containing the replicates, models for the empirical data, and summary statistics and plots.

## **Examples**

```
cyreni <- iberolacerta.clade$species$cyreni
aranica <- iberolacerta.clade$species$aranica</pre>
```

rangebreak.linear conduct a linear rangebreak test as described in Glor and Warren 2011.

# Description

rangebreak.linear Conduct a linear rangebreak test as described in Glor and Warren 2011.

# Usage

```
rangebreak.linear(
  species.1,
  species.2,
  env,
  type,
  f = NULL,
  nreps = 99,
  nback = 1000,
  bg.source = "default",
  low.memory = FALSE,
  rep.dir = NA,
  verbose = FALSE,
  clamp = TRUE,
  ...
)
```

rangebreak.linear 53

# Arguments

species.1	An emtools.species object
species.2	An enmtools.species object
env	A SpatRaster object containing environmental data
type	The type of model to construct, currently accepts "glm", "mx", "bc", "gam", or "dm" $$
f	A function to use for model fitting. Only required for GLM models at the moment.
nreps	Number of replicates to perform
nback	Number of background points for models
bg.source	Source for drawing background points. If "points", it just uses the background points that are already in the species object. If "range", it uses the range raster. If "env", it draws points at randome from the entire study area outlined by the first environmental layer.
low.memory	When set to TRUE, replicate models are written to disc instead of being stored in the output object. Replicate models stored in the output object contain paths to the replicate models on disk instead of the rasters themselves.
rep.dir	Directory for storing replicate models when low.memory is set to TRUE. If not specified, the working directory will be used.
verbose	Controls printing of various messages progress reports. Defaults to FALSE.
clamp	Controls whether empirical and replicate models should be clamped to the environment space used for training.
	Additional arguments to be passed to model fitting functions.

#### Value

results A list containing the replicates, models for the empirical data, and summary statistics and plots.

```
cyreni <- iberolacerta.clade$species$cyreni
aranica <- iberolacerta.clade$species$aranica
rangebreak.linear(cyreni, aranica, env = euro.worldclim, type = "glm",
f= pres ~ bio1 + bio12, nreps = 10)</pre>
```

54 rangebreak.ribbon

rangebreak.ribbon

rangebreak.ribbon Conduct a ribbon rangebreak test as described in Glor and Warren 2011.

# Description

rangebreak.ribbon Conduct a ribbon rangebreak test as described in Glor and Warren 2011.

# Usage

```
rangebreak.ribbon(
  species.1,
  species.2,
 ribbon,
 env,
  type,
  f = NULL,
 width = 1,
 nreps = 99,
 nback = 1000,
 bg.source = "default",
 low.memory = FALSE,
 rep.dir = NA,
 verbose = FALSE,
 clamp = TRUE,
)
```

# Arguments

species.1	An emtools.species object
species.2	An enmtools.species object
ribbon	An enmtools.species object representing the region of marginal habtiat in the overlap between the species' ranges
env	A SpatRaster object containing environmental data
type	The type of model to construct, currently accepts "glm", "mx", "bc", or "dm"
f	A function to use for model fitting. Only required for GLM models at the moment.
width	Width of the ribbon, in the same units as the occurrence points (e.g, decimal degrees)
nreps	Number of replicates to perform
nback	Number of background points for models
bg.source	Source for drawing background points. If "points", it just uses the background points that are already in the species object. If "range", it uses the range raster. If "env", it draws points at randome from the entire study area outlined by the first environmental layer.

raster.breadth 55

low.memory	When set to TRUE, replicate models are written to disc instead of being stored in the output object. Replicate models stored in the output object contain paths to the replicate models on disk instead of the rasters themselves.
rep.dir	Directory for storing replicate models when low.memory is set to TRUE. If not specified, the working directory will be used.
verbose	Controls printing of various messages progress reports. Defaults to FALSE.
clamp	Controls whether empirical and replicate models should be clamped to the environment space used for training.
	Additional arguments to be passed to model fitting functions.

#### Value

results A list containing models for the replicates, models for the empirical data, and summary statistics and plots.

## **Examples**

```
cyreni <- iberolacerta.clade$species$cyreni
aranica <- iberolacerta.clade$species$aranica

# We're just going to fudge together occurrence data from a ribbon here
# from random points between the two species' ranges
p <- data.frame(x = runif(50, -4, -2), y = runif(50, 40, 43))
p <- terra::vect(p, geom = c("x", "y"), crs = terra::crs(cyreni$presence.points))
bg <- background.buffer(p, 100000, mask = euro.worldclim[[1]], n = 100, return.type = "points")
ribbon <- enmtools.species(species.name = "ribbon", presence.points = p, background.points = bg)
rangebreak.ribbon(cyreni, aranica, ribbon = ribbon, env = euro.worldclim,
type = "glm", f= pres ~ bio1 + bio12, nreps = 10)</pre>
```

raster.breadth

raster.breadth, applies measures of niche breadth to an ENM

## **Description**

This function measures the spatial heterogeneity of the distribution of suitability scores from an ENM. It returns Levins' (1968) two metrics of niche breadth.

## Usage

```
raster.breadth(x, verbose = FALSE)
```

#### **Arguments**

X	An object of class raster or RasterLayer, or an ENMTools model object contain-
	ing a suitability raster.
verbose	Controls printing of diagnostic messages

56 raster.cor

## **Details**

Levins, R. 1968. Evolution in changing environments: some theoretical explorations. Princeton University Press.

#### Value

results A vector containing the two metrics B1 and B2 for niche breadth.

## **Examples**

```
\label{lem:condition} $$\operatorname{aurelioi.glm} \leftarrow \operatorname{enmtools.glm}(iberolacerta.clade\$species\$aurelioi, euro.worldclim, f = pres \sim bio1 + bio12)$$ raster.breadth(aurelioi.glm)
```

raster.cor

Calculates the correlation coefficient between two rasters.

# Description

Calculates the correlation coefficient between two rasters.

## Usage

```
raster.cor(x, y, method = "spearman")
```

## **Arguments**

x Either a raster or an ENMTools model object with a suitability raster.

y Either a raster or an ENMTools model object with a suitability raster.

method The method to be used for calculating correlations. Defaults to spearman, but

can take "kendall" or "pearson" as well.

## Value

A numeric correlation coefficient.

```
data(euro.worldclim)
raster.cor(euro.worldclim[[1]], euro.worldclim[[2]])
```

raster.cor.matrix 57

raster.cor.matrix	Takes a raster stack and returns a data frame containing Pearson cor- relation coefficients between the included rasters
	35

#### **Description**

Takes a raster stack and returns a data frame containing Pearson correlation coefficients between the included rasters

## Usage

```
raster.cor.matrix(env, method = "pearson")
```

## **Arguments**

env A SpatRaster object containing two or more rasters

method Type of correlation to measure. Can do "pearson", "kendall", or "spearman"

#### Value

A data frame of correlation coefficients for a set of rasters.

## **Examples**

```
data(euro.worldclim)
raster.cor.matrix(euro.worldclim)
```

raster.cor.plot

Takes a raster stack and returns a data frame containing Pearson correlation coefficients between the included rasters

# Description

Takes a raster stack and returns a data frame containing Pearson correlation coefficients between the included rasters

## Usage

```
raster.cor.plot(env, method = "pearson")
```

# Arguments

env A SpatRaster object containing two or more rasters

method Type of correlation to measure. Can do "pearson", "kendall", or "spearman"

58 raster.overlap

### Value

A list of two plots. The first maps the correlations between rasters into an MDS space, so that predictors that fall close together in that space are more correlated. The second plot is a heatmap depicting the correlations between pairs of layers.

### **Examples**

```
data(euro.worldclim)
raster.cor.plot(euro.worldclim)
```

raster.overlap

raster.overlap, measures overlap between two ENMs

## Description

This function measures similarity in the geographic distribution of suitability scores from two ENMs. It returns two metrics, I and D. These metrics are described in Warren et al. 2008.

## Usage

```
raster.overlap(x, y, verbose = FALSE)
```

# **Arguments**

X	A raster or RasterLayer object, or ENMTools model object containing a suitability raster.
У	Another raster or RasterLayer object, or ENMTools model object containing a

avitability master

suitability raster.

verbose Controls printing of diagnostic messages

#### Value

results A vector containing the three metrics (I, D, and Spearman rank correlation)

```
aurelioi.glm <- enmtools.glm(iberolacerta.clade$species$aurelioi,
euro.worldclim, f = pres ~ bio1 + bio12)
aranica.glm <- enmtools.glm(iberolacerta.clade$species$aranica,
euro.worldclim, f = pres ~ bio1 + bio12)
raster.overlap(aurelioi.glm, aranica.glm)</pre>
```

59 raster.pca

raster.pca

raster.pca, PCA on a set of environmental rasters

## Description

raster.pca, PCA on a set of environmental rasters

## Usage

```
raster.pca(env, n)
```

## Arguments

A set of environmental layers env

The number of PCA layers to return n

#### Value

A list containing a stack of rasters representing the top n pca axes of the initial environmental variables, as well as the pca object from the analysis that produced them.

## **Examples**

```
env.pca <- raster.pca(euro.worldclim, 2)</pre>
```

raster.resid

raster.resid Measure standardized residuals from a linear regression between two rasters.

# **Description**

This function builds a linear model for the relationship between two rasters, and returns the linear model and a spatial raster of the residuals from that model.

# Usage

```
raster.resid(x, y)
```

# **Arguments**

X	A raster or RasterLayer object, or ENMTools model object containing a suit-
	.1.117

Another raster or RasterLayer object, or ENMTools model object containing a У

suitability raster.

60 raster.standardize

## Value

results A list containing a raster of residuals from a linear regression between the two supplied rasters and the linear model itself.

# **Examples**

```
data(euro.worldclim)
raster.resid(euro.worldclim[[1]], euro.worldclim[[2]])
```

raster.standardize

raster.standardize, standardizes all values in a raster file

# Description

This function is used by a lot of the metrics calculated by ENMTools, in order to standardize suitability scores so they sum to 1 over a geographic space.

#### Usage

```
raster.standardize(x, verbose = FALSE)
```

## **Arguments**

x A raster or RasterLayer object, or an ENMTools model object containing a suit-

ability raster.

verbose Controls printing of diagnostic messages

#### Value

A new raster object, standardized so that values sum to 1

```
raster.standardize(euro.worldclim[[1]])
```

rasterToPoints2 61

rasterToPoints2

get terra to produce same output as raster::rasterToPoints

### **Description**

get terra to produce same output as raster::rasterToPoints

#### Usage

```
rasterToPoints2(rast)
```

## **Arguments**

rast

A raster to convert to points

sim.points

Simulate a point process from an enmtools.model object

#### **Description**

Function that takes an enmtools.model object and simulates points from it using a point process.

# Usage

```
sim.points(
  object,
  n.points = 1000,
  seed = NA,
  sample.type = "ppp",
  replace = FALSE,
  threshold = NA,
  ...
)
```

## **Arguments**

object entools.model object to simulate from

n.points approximate number of points to simulate. You may get small number fewer or

greater. If you need an exact number of points, generate too many, then drop the

extra ones.

seed optional seed for the random number generator

sample.type method for sampling occurrence points. Default is "ppp", which is a poisson

point process. Also accepts "binomial" for treating suitabilities as binomial trials, "thresh.pa" for sampling with equal probability every grid cell above a certain threshold value, and "thresh.con" for sampling in proportion to suitability

above a certain threshold value.

62 threespace.plot

replace whether to sample with replacement.

threshold a threshold suitability below which probability of sampling drops to zero. Used

for "thresh.pa" and "thresh.con" sampling.

... additional parameters, not currently used but included for S3 compatability

#### Value

A data frame of simulated points from the enmtools.model object

species.from.file

Takes a csv file and uses it to construct one or more enmtools.species objects. It constructs one object per unique value in the "species.col" column.

#### **Description**

Takes a csv file and uses it to construct one or more enmtools.species objects. It constructs one object per unique value in the "species.col" column.

# Usage

```
species.from.file(filename, species.col = "species")
```

## **Arguments**

filename Name of a .csv file

species.col Name of the column from the csv file to use for species names. Default is

"species".

## Value

A list containing species objects, one for each unique value in species.col.

threespace.plot	threespace.plot, A plot that compares the environmental distribution of presence points, background points, and the set of supplied environmental layers
	mental layers.

#### **Description**

threespace.plot, A plot that compares the environmental distribution of presence points, background points, and the set of supplied environmental layers.

#### Usage

```
threespace.plot(model, env, maxpts = NA)
```

trimdupes.by.raster 63

### **Arguments**

model	An enmtools.model object
env	A set of environment layers

maxpts Maximum number of points to plot from env layers

#### Value

A ggplot2 object that plots the distribution of environments in the climate layers to the distribution of environments at background and presence localities.

## **Examples**

```
my.model<- enmtools.gam(iberolacerta.clade$species$monticola, euro.worldclim)
threespace.plot(my.model, euro.worldclim)</pre>
```

trimdupes.by.raster

Takes a set of points and a raster mask and returns a data frame trimmed so that only one point is returned per grid cell in the mask raster.

# Description

Takes a set of points and a raster mask and returns a data frame trimmed so that only one point is returned per grid cell in the mask raster.

## Usage

```
trimdupes.by.raster(points, mask)
```

# Arguments

points A two column data frame with X and Y coordinates

mask A raster to use as a mask for drawing points

#### Value

A new data frame with one point per grid cell.

```
pts <- iberolacerta.clade$species$monticola$presence.points
trimdupes.by.raster(pts, euro.worldclim)</pre>
```

64 visualize.enm

unwrap\_list

export

## **Description**

export

## Usage

```
unwrap_list(l)
```

#### **Arguments**

1

A list containing species objects that need to be converted

visualize.enm

visualize.enm, Makes a heatmap of suitability of habitat in environment space according to a given model

# Description

visualize.enm, Makes a heatmap of suitability of habitat in environment space according to a given model

# Usage

```
visualize.enm(
  model,
  env,
  nbins = 100,
  layers = colnames(model$analysis.df)[1:2],
  plot.test.data = FALSE,
  plot.points = TRUE,
  minmax = "env"
)
```

## **Arguments**

model An enmtools.model object env A set of environmental layers

nbins The number of bins per layer to use for drawing environment space layers A vector of layer names to use for drawing environment space

plot.test.data Logical determining whether test data should be plotted, if present. If test data

is plotted, it will appear as translucent green triangles.

visualize.overlap 65

plot.points Logical determining whether presence points should be plotted on suitability

plot

minmax Either "env", to set the minima and maxima using the environment layers, "points"

to set the minima and maxima using the presence and background data, or a

named list of minima and maxima for each layer.

#### Value

suit.plot A list containing two dimensional plot of an ENM in environment space and a plot of the available environments.

## **Examples**

```
\label{eq:control} $$\operatorname{aurelioi.glm} <-\operatorname{enmtools.glm}(iberolacerta.clade\$species\$aurelioi,\ euro.worldclim, f = pres ~ poly(bio1, 4) + poly(bio12, 4))$$ visualize.enm(aurelioi.glm, euro.worldclim, layers = c("bio1", "bio12")) $$
```

visualize.overlap

visualize.overlap, Makes a contour map of suitability of habitat in environment space for two models

#### **Description**

visualize.overlap, Makes a contour map of suitability of habitat in environment space for two models

## Usage

```
visualize.overlap(
  model.1,
  model.2,
  env,
  nbins = 100,
  layers,
  plot.points = TRUE
)
```

## Arguments

model.1	An enmtools.model object
model.2	An enmtools.model object
env	A set of environmental layers
nbins	The number of bins per layer to use for drawing environment space
layers	A vector of layer names to use for drawing environment space
plot.points	Logical determining whether presence points should be plotted on suitability plot

66 wrap\_list

# Value

overlap.plot A two dimensional plot of an ENM

# **Examples**

```
ar <- iberolacerta.clade$species$aranica
au <- iberolacerta.clade$species$aurelioi
aranica.dm <- enmtools.dm(ar, euro.worldclim)
aurelioi.dm <- enmtools.dm(au, euro.worldclim)
visualize.overlap(aranica.dm, aurelioi.dm, euro.worldclim, layers = c("bio1", "bio9"))</pre>
```

wrap\_list

export

# Description

export

# Usage

wrap\_list(l)

# Arguments

1 A list containing species objects that need to be converted

# **Index**

* background	* equivalency
background.test,7	background.test,7
threespace.plot, 62	identity.test,42
* barrier	* extrapolation
rangebreak.blob,51	threespace.plot,62
rangebreak.linear,52	* hypothesis-testing
rangebreak.ribbon,54	identity.test,42
* biogeography	rangebreak.blob,51
rangebreak.blob,51	rangebreak.linear,52
rangebreak.linear,52	rangebreak.ribbon,54
rangebreak.ribbon,54	* hypothesis
* breadth	background.test,7
calc.B1,9	* identity
calc.B2, 10	identity.test,42
* comparison	* keywords
threespace.plot, 62	raster.breadth, 55
* correlation	raster.overlap, 58
raster.resid,59	raster.standardize, 60
* datasets	* niche
euro.worldclim,39	calc.B1,9
iberolacerta.clade,42	calc.B2, 10
* enmtools	enmtools.ecospat.bg, 21
background.test, $7$	enmtools.ecospat.id, 23
identity.test,42	visualize.enm, 64
rangebreak.blob,51	visualize.overlap,65
rangebreak.linear,52	* overlap
rangebreak.ribbon,54	visualize.overlap,65
* enm	* pca
calc.B1,9	multistack.pca, 49
calc.B2, <u>10</u>	raster.pca, 59
enmtools.ecospat.bg, 21	threespace.plot, 62
enmtools.ecospat.id, 23	* plot
marginal.plots,47	enmtools.ecospat.bg, $21$
visualize.enm, 64	enmtools.ecospat.id, 23
visualize.overlap,65	marginal.plots,47
* environment	visualize.enm, 64
multistack.pca,49	visualize.overlap,65
raster.pca, 59	* presence
threespace.plot, 62	threespace.plot, 62

68 INDEX

* rangebreak	enmtools.ecospat.bg, 21
rangebreak.blob,51	enmtools.ecospat.id, 23
rangebreak.linear,52	enmtools.gam, 24
rangebreak.ribbon,54	enmtools.glm, 26
* raster	enmtools.hypervolume, 28
multistack.pca, 49	enmtools.maxent, 29
raster.pca, 59	enmtools.rf, 30
raster.resid, 59	enmtools.rf.ranger, 32
* residuals	enmtools.species, 34, 46
raster.resid, 59	enmtools.vip, 35
* response	env.breadth, 36
marginal.plots, 47	env.evaluate, 37
* sdm	env.overlap, 38
calc.B1,9	euro.worldclim, 39
calc.B1, 9	cui o. woi ruci ini, 37
	find.extras, 39
enmtools.ecospat.bg, 21	find.extras.missing, 40
enmtools.ecospat.id, 23	1111d. CACI d3.1111331116, 10
marginal.plots, 47	geog.range.overlap,40
visualize.enm, 64	8008.1. 8.1.8010.101. 245, 10
visualize.overlap,65	hypervolume.overlap, 41
* testing	17
background.test, $7$	iberolacerta.clade, 42
	identity.test,42
add.env, 3	install.extras, 44
assert.extras, 4	interactive.plot, 45
	<pre>interactive.plot.enmtools.model, 45</pre>
background.buffer, 5	interactive.plot.enmtools.species, 46
${\sf background.points.buffer}, 6$	
background.raster.buffer, 6	leaflet, 45, 46
background.shape.buffer,7	load.enmtools.clade(enmtools.clade), 19
background.test, 7	load.enmtools.species
	(enmtools.species), 34
calc.B1,9	(01111100201060200), 0.
calc.B2, 10	marginal.plots,47
check.bg, 10	moses.list, 48
check.clade, 11	multistack.pca, 49
check.env, 12	
check.extras, 12	node.overlap, 49
check.species, 13	17
clamp.env, 13	point.overlap, 50
combine.species, 14	
	rangebreak.blob,51
drop.species, 15	rangebreak.linear,52
	rangebreak.ribbon,54
enmtools.aoc, 15	ranger, <i>33</i>
enmtools.bc, 16	raster.breadth, 55
enmtools.calibrate, 18	raster.cor, 56
enmtools.clade, 19	raster.cor.matrix, 57
enmtools.dm, 20	raster.cor.plot, 57

INDEX 69

```
raster.overlap, 58
raster.pca, 59
raster.resid, 59
raster.standardize, 60
rasterToPoints2, 61
save.enmtools.clade (enmtools.clade), 19
save.enmtools.species
        (enmtools.species), 34
sim.points, 61
species.from.file,62
threespace.plot, 62
trimdupes.by.raster, 63
unwrap_list, 64
visualize.enm, 64
visualize.overlap, 65
wrap_list, 66
```