

# Package ‘biomod2’

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**Type** Package

**Title** Ensemble Platform for Species Distribution Modeling

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**BugReports** <https://github.com/biomodhub/biomod2/issues>

**URL** <https://biomodhub.github.io/biomod2/>

**Description** Functions for species distribution modeling, calibration and evaluation, ensemble of models, ensemble forecasting and visualization. The package permits to run consistently up to 10 single models on a presence/absences (resp presences/pseudo-absences) dataset and to combine them in ensemble models and ensemble projections. Some bench of other evaluation and visualization tools are also available within the package.

**Depends** R (>= 4.1)

**Imports** stats, utils, methods, terra (>= 1.6-33), sp, reshape,  
reshape2, abind, foreach, ggplot2, gbm (>= 2.1.3), rpart, MASS,  
pROC (>= 1.15.0), PresenceAbsence, dplyr

**Suggests** Hmisc, gam, mgcv, earth, maxnet, mda, nnet, randomForest,  
xgboost, car, caret, dismo, ENMeval, doParallel, raster,  
ggpubr, testthat, knitr, markdown, tidyterra, ggtext

**License** GPL-3

**RoxygenNote** 7.3.2

**Encoding** UTF-8

**VignetteBuilder** knitr

**Collate** 'biomod2-package.R' 'biomod2\_globalVariables.R'  
 'biomod2\_classes\_0.R' 'biomod2\_classes\_1.R'  
 'biomod2\_classes\_2.R' 'biomod2\_classes\_3.R'  
 'biomod2\_classes\_4.R' 'biomod2\_classes\_5.R'  
 'biomod2\_internal.R' 'biomod2\_data.R'  
 'BIOMOD\_EnsembleForecasting.R' 'BIOMOD\_EnsembleModeling.R'  
 'BIOMOD\_FormatingData.R' 'BIOMOD\_LoadModels.R'  
 'BIOMOD\_Modeling.R' 'BIOMOD\_Projection.R' 'BIOMOD\_RangeSize.R'  
 'DEPRECATED.R' 'bm\_BinaryTransformation.R'  
 'bm\_CrossValidation.R' 'bm\_FindOptimStat.R' 'bm\_MakeFormula.R'  
 'bm\_ModelingOptions.R' 'bm\_PlotEvalBoxplot.R'  
 'bm\_PlotEvalMean.R' 'bm\_PlotRangeSize.R'  
 'bm\_PlotResponseCurves.R' 'bm\_PlotVarImpBoxplot.R'  
 'bm\_PseudoAbsences.R' 'bm\_RunModelsLoop.R' 'bm\_SRE.R'  
 'bm\_SampleBinaryVector.R' 'bm\_SampleFactorLevels.R'  
 'bm\_Tuning.R' 'bm\_VariablesImportance.R' 'zzz.R'

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bioclim\_current      *Bioclimatic variables for SDM based on current condition*

---

## Description

A [SpatRaster](#) with 5 bioclimatic variables commonly used for SDM and describing current climate. Additional information available at [worldclim](#)

## Usage

bioclim\_current

**Format**

A [SpatRaster](#) with 5 layers:

**bio3** Isothermality

**bio4** Temperature Seasonality

**bio7** Temperature Annual Range

**bio11** Mean Temperature of Coldest Quarter

**bio12** Annual Precipitation

---

bioclim_future	<i>Bioclimatic variables for SDM based on future condition</i>
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**Description**

A [SpatRaster](#) with 5 bioclimatic variables commonly used for SDM and describing future climate based on old RCP scenarios at the horizon 2080.

**Usage**

```
bioclim_future
```

**Format**

A [SpatRaster](#) with 5 layers:

**bio3** Isothermality

**bio4** Temperature Seasonality

**bio7** Temperature Annual Range

**bio11** Mean Temperature of Coldest Quarter

**bio12** Annual Precipitation

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BIOMOD.ensemble.models.out	<i>BIOMOD_EnsembleModeling() output object class</i>
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**Description**

Class returned by [BIOMOD\\_EnsembleModeling](#), and used by [BIOMOD\\_LoadModels](#), [BIOMOD\\_PresenceOnly](#) and [BIOMOD\\_EnsembleForecasting](#)

**Usage**

```
## S4 method for signature 'BIOMOD.ensemble.models.out'  
show(object)
```

**Arguments**

object a `BIOMOD.ensemble.models.out` object

**Slots**

`modeling.id` a character corresponding to the name (ID) of the simulation set  
`dir.name` a character corresponding to the modeling folder  
`sp.name` a character corresponding to the species name  
`expl.var.names` a vector containing names of explanatory variables  
`models.out` a `BIOMOD.stored.models.out-class` object containing informations from `BIOMOD_Modeling` object  
`em.by` a character corresponding to the way kept models have been combined to build the ensemble models, must be among PA+run, PA+algo, PA, algo, all  
`em.computed` a vector containing names of ensemble models  
`em.failed` a vector containing names of failed ensemble models  
`em.models_kept` a list containing single models for each ensemble model  
`models.evaluation` a `BIOMOD.stored.data.frame-class` object containing models evaluation variables.  
`importance` a `BIOMOD.stored.data.frame-class` object containing variables importance  
`models.prediction` a `BIOMOD.stored.data.frame-class` object containing models predictions  
`models.prediction.eval` a `BIOMOD.stored.data.frame-class` object containing models predictions for evaluation data  
`link` a character containing the file name of the saved object

**Author(s)**

Damien Georges

**See Also**

`BIOMOD_EnsembleModeling`, `BIOMOD_LoadModels`, `BIOMOD_PresenceOnly`, `bm_VariablesImportance`, `bm_PlotEvalMean`, `bm_PlotEvalBoxplot`, `bm_PlotVarImpBoxplot`, `bm_PlotResponseCurves`

Other Toolbox objects: `BIOMOD.formated.data`, `BIOMOD.formated.data.PA`, `BIOMOD.models.options`, `BIOMOD.models.out`, `BIOMOD.options.dataset`, `BIOMOD.options.default`, `BIOMOD.projection.out`, `BIOMOD.stored.data`, `biomod2_ensemble_model`, `biomod2_model`

**Examples**

```
showClass("BIOMOD.ensemble.models.out")

## ----- #
library(terra)

# Load species occurrences (6 species available)
data(DataSpecies)
```

```

head(DataSpecies)

# Select the name of the studied species
myRespName <- 'GuloGulo'

# Get corresponding presence/absence data
myResp <- as.numeric(DataSpecies[, myRespName])

# Get corresponding XY coordinates
myRespXY <- DataSpecies[, c('X_WGS84', 'Y_WGS84')]

# Load environmental variables extracted from BIOCLIM (bio_3, bio_4, bio_7, bio_11 & bio_12)
data(bioclim_current)
myExpl <- terra::rast(bioclim_current)

## ----- #
file.out <- paste0(myRespName, "/", myRespName, ".AllModels.models.out")
if (file.exists(file.out)) {
  myBiomodModelOut <- get(load(file.out))
} else {

  # Format Data with true absences
  myBiomodData <- BIOMOD_FormatingData(resp.var = myResp,
                                     expl.var = myExpl,
                                     resp.xy = myRespXY,
                                     resp.name = myRespName)

  # Model single models
  myBiomodModelOut <- BIOMOD_Modeling(bm.format = myBiomodData,
                                     modeling.id = 'AllModels',
                                     models = c('RF', 'GLM'),
                                     CV.strategy = 'random',
                                     CV.nb.rep = 2,
                                     CV.perc = 0.8,
                                     OPT.strategy = 'bigboss',
                                     metric.eval = c('TSS', 'ROC'),
                                     var.import = 3,
                                     seed.val = 42)
}

## ----- #
# Model ensemble models
myBiomodEM <- BIOMOD_EnsembleModeling(bm.mod = myBiomodModelOut,
                                     models.chosen = 'all',
                                     em.by = 'all',
                                     em.algo = c('EMmean', 'EMca'),
                                     metric.select = c('TSS'),
                                     metric.select.thresh = c(0.7),
                                     metric.eval = c('TSS', 'ROC'),
                                     var.import = 3,

```

```
myBiomodEM                                     seed.val = 42)
```

---

BIOMOD.formated.data BIOMOD\_FormatingData() *output object class*

---

## Description

Class returned by [BIOMOD\\_FormatingData](#), and used by [bm\\_Tuning](#), [bm\\_CrossValidation](#) and [BIOMOD\\_Modeling](#)

## Usage

```
## S4 method for signature 'numeric,data.frame'
BIOMOD.formated.data(
  sp,
  env,
  xy = NULL,
  dir.name = ".",
  sp.name = NULL,
  eval.sp = NULL,
  eval.env = NULL,
  eval.xy = NULL,
  na.rm = TRUE,
  data.mask = NULL,
  shared.eval.env = FALSE,
  filter.raster = FALSE
)

## S4 method for signature 'data.frame,ANY'
BIOMOD.formated.data(
  sp,
  env,
  xy = NULL,
  dir.name = ".",
  sp.name = NULL,
  eval.sp = NULL,
  eval.env = NULL,
  eval.xy = NULL,
  na.rm = TRUE,
  filter.raster = FALSE
)

## S4 method for signature 'numeric,matrix'
BIOMOD.formated.data(
```

```

    sp,
    env,
    xy = NULL,
    dir.name = ".",
    sp.name = NULL,
    eval.sp = NULL,
    eval.env = NULL,
    eval.xy = NULL,
    na.rm = TRUE,
    filter.raster = FALSE
)

## S4 method for signature 'numeric,SpatRaster'
BIOMOD.formated.data(
  sp,
  env,
  xy = NULL,
  dir.name = ".",
  sp.name = NULL,
  eval.sp = NULL,
  eval.env = NULL,
  eval.xy = NULL,
  na.rm = TRUE,
  shared.eval.env = FALSE,
  filter.raster = FALSE
)

## S4 method for signature 'BIOMOD.formated.data'
show(object)

```

### Arguments

sp	A vector, a <a href="#">SpatVector</a> without associated data ( <i>if presence-only</i> ), or a <a href="#">SpatVector</a> object containing binary data (0 : absence, 1 : presence, NA : indeterminate) for a single species that will be used to build the species distribution model(s) <i>Note that old format from <b>sp</b> are still supported such as SpatialPoints (if presence-only) or SpatialPointsDataFrame object containing binary data.</i>
env	a matrix, data.frame, <a href="#">SpatVector</a> or <a href="#">SpatRaster</a> object containing the explanatory variables (in columns or layers) that will be used to build the species distribution model(s). <i>Note that old format from <b>raster</b> and <b>sp</b> are still supported such as RasterStack and SpatialPointsDataFrame objects.</i>
xy	(optional, default NULL) If resp.var is a vector, a 2-columns matrix or data.frame containing the corresponding X and Y coordinates that will be used to build the species distribution model(s)
dir.name	a character corresponding to the modeling folder
sp.name	a character corresponding to the species name



eval.sp	(optional, default NULL) A vector, a <a href="#">SpatVector</a> without associated data (if <i>presence-only</i> ), or a <a href="#">SpatVector</a> object containing binary data (0 : absence, 1 : presence, NA : indeterminate) for a single species that will be used to evaluate the species distribution model(s) with independent data <i>Note that old format from <b>sp</b> are still supported such as SpatialPoints (if presence-only) or SpatialPointsDataFrame object containing binary data.</i>
eval.env	(optional, default NULL) A matrix, data.frame, <a href="#">SpatVector</a> or <a href="#">SpatRaster</a> object containing the explanatory variables (in columns or layers) that will be used to evaluate the species distribution model(s) with independent data <i>Note that old format from <b>raster</b> and <b>sp</b> are still supported such as RasterStack and SpatialPointsDataFrame objects.</i>
eval.xy	(optional, default NULL) If resp.var is a vector, a 2-columns matrix or data.frame containing the corresponding X and Y coordinates that will be used to evaluate the species distribution model(s) with independent data
na.rm	(optional, default TRUE) A logical value defining whether points having one or several missing values for explanatory variables should be removed from the analysis or not
data.mask	(optional, default NULL) A <a href="#">SpatRaster</a> object containing the mask of the studied area
shared.eval.env	(optional, default FALSE) A logical value defining whether the explanatory variables used for the evaluation dataset are the same than the ones for calibration (if eval.env not provided for example) or not
filter.raster	(optional, default FALSE) If env is of raster type, a logical value defining whether sp is to be filtered when several points occur in the same raster cell
object	a <a href="#">BIOMOD.formated.data</a> object

### Slots

dir.name	a character corresponding to the modeling folder
sp.name	a character corresponding to the species name
coord	a 2-columns data.frame containing the corresponding X and Y coordinates
data.species	a vector containing the species observations (0, 1 or NA)
data.env.var	a data.frame containing explanatory variables
data.mask	a <a href="#">SpatRaster</a> object containing the mask of the studied area
has.data.eval	a logical value defining whether evaluation data is given
eval.coord	(optional, default NULL) A 2-columns data.frame containing the corresponding X and Y coordinates for evaluation data

`eval.data.species` (*optional, default NULL*)  
 A vector containing the species observations (0, 1 or NA) for evaluation data

`eval.data.env.var` (*optional, default NULL*)  
 A data.frame containing explanatory variables for evaluation data

### Author(s)

Damien Georges

### See Also

[BIOMOD\\_FormatingData](#), [bm\\_Tuning](#), [bm\\_CrossValidation](#), [BIOMOD\\_Modeling](#), [bm\\_RunModelsLoop](#)  
 Other Toolbox objects: [BIOMOD.ensemble.models.out](#), [BIOMOD.formated.data.PA](#), [BIOMOD.models.options](#),  
[BIOMOD.models.out](#), [BIOMOD.options.dataset](#), [BIOMOD.options.default](#), [BIOMOD.projection.out](#),  
[BIOMOD.stored.data](#), [biomod2\\_ensemble\\_model](#), [biomod2\\_model](#)

### Examples

```
showClass("BIOMOD.formated.data")

## ----- #
library(terra)

# Load species occurrences (6 species available)
data(DataSpecies)
head(DataSpecies)

# Select the name of the studied species
myRespName <- 'GuloGulo'

# Get corresponding presence/absence data
myResp <- as.numeric(DataSpecies[, myRespName])

# Get corresponding XY coordinates
myRespXY <- DataSpecies[, c('X_WGS84', 'Y_WGS84')]

# Load environmental variables extracted from BIOCLIM (bio_3, bio_4, bio_7, bio_11 & bio_12)
data(bioclim_current)
myExpl <- terra::rast(bioclim_current)

## ----- #
# Format Data with true absences
myBiomodData <- BIOMOD_FormatingData(resp.var = myResp,
                                   expl.var = myExpl,
                                   resp.xy = myRespXY,
                                   resp.name = myRespName)

myBiomodData
plot(myBiomodData)
summary(myBiomodData)
```

---

 BIOMOD.formated.data.PA

 BIOMOD\_FormatingData() *output object class (with pseudo-absences)*


---

### Description

Class returned by [BIOMOD\\_FormatingData](#), and used by [bm\\_Tuning](#), [bm\\_CrossValidation](#) and [BIOMOD\\_Modeling](#)

### Usage

```
## S4 method for signature 'numeric,data.frame'
BIOMOD.formated.data.PA(
  sp,
  env,
  xy = NULL,
  dir.name = ".",
  sp.name = NULL,
  eval.sp = NULL,
  eval.env = NULL,
  eval.xy = NULL,
  PA.nb.rep = 1,
  PA.strategy = "random",
  PA.nb.absences = NULL,
  PA.dist.min = 0,
  PA.dist.max = NULL,
  PA.sre.quant = 0.025,
  PA.fact.aggr = NULL,
  PA.user.table = NULL,
  na.rm = TRUE,
  filter.raster = FALSE,
  seed.val = NULL
)

## S4 method for signature 'numeric,SpatRaster'
BIOMOD.formated.data.PA(
  sp,
  env,
  xy = NULL,
  dir.name = ".",
  sp.name = NULL,
  eval.sp = NULL,
  eval.env = NULL,
```

```

eval.xy = NULL,
PA.nb.rep = 1,
PA.strategy = "random",
PA.nb.absences = NULL,
PA.dist.min = 0,
PA.dist.max = NULL,
PA.sre.quant = 0.025,
PA.fact.aggr = NULL,
PA.user.table = NULL,
na.rm = TRUE,
filter.raster = FALSE,
seed.val = NULL
)

```

### Arguments

sp	A vector, a <a href="#">SpatVector</a> without associated data ( <i>if presence-only</i> ), or a <a href="#">SpatVector</a> object containing binary data (0 : absence, 1 : presence, NA : indeterminate) for a single species that will be used to build the species distribution model(s) <i>Note that old format from <b>sp</b> are still supported such as SpatialPoints (if presence-only) or SpatialPointsDataFrame object containing binary data.</i>
env	a matrix, data.frame, <a href="#">SpatVector</a> or <a href="#">SpatRaster</a> object containing the explanatory variables (in columns or layers) that will be used to build the species distribution model(s). <i>Note that old format from <b>raster</b> and <b>sp</b> are still supported such as RasterStack and SpatialPointsDataFrame objects.</i>
xy	(optional, default NULL) If resp.var is a vector, a 2-columns matrix or data.frame containing the corresponding X and Y coordinates that will be used to build the species distribution model(s)
dir.name	a character corresponding to the modeling folder
sp.name	a character corresponding to the species name
eval.sp	(optional, default NULL) A vector, a <a href="#">SpatVector</a> without associated data ( <i>if presence-only</i> ), or a <a href="#">SpatVector</a> object containing binary data (0 : absence, 1 : presence, NA : indeterminate) for a single species that will be used to evaluate the species distribution model(s) with independent data <i>Note that old format from <b>sp</b> are still supported such as SpatialPoints (if presence-only) or SpatialPointsDataFrame object containing binary data.</i>
eval.env	(optional, default NULL) A matrix, data.frame, <a href="#">SpatVector</a> or <a href="#">SpatRaster</a> object containing the explanatory variables (in columns or layers) that will be used to evaluate the species distribution model(s) with independent data <i>Note that old format from <b>raster</b> and <b>sp</b> are still supported such as RasterStack and SpatialPointsDataFrame objects.</i>
eval.xy	(optional, default NULL) If resp.var is a vector, a 2-columns matrix or data.frame containing the

	corresponding X and Y coordinates that will be used to evaluate the species distribution model(s) with independent data
PA.nb.rep	<i>(optional, default 0)</i> If pseudo-absence selection, an integer corresponding to the number of sets (repetitions) of pseudo-absence points that will be drawn
PA.strategy	<i>(optional, default NULL)</i> If pseudo-absence selection, a character defining the strategy that will be used to select the pseudo-absence points. Must be random, sre, disk or user.defined (see Details)
PA.nb.absences	<i>(optional, default 0)</i> If pseudo-absence selection, and PA.strategy = 'random' or PA.strategy = 'sre' or PA.strategy = 'disk', an integer (or a vector of integer the same size as PA.nb.rep) corresponding to the number of pseudo-absence points that will be selected for each pseudo-absence repetition (true absences included)
PA.dist.min	<i>(optional, default 0)</i> If pseudo-absence selection and PA.strategy = 'disk', a numeric defining the minimal distance to presence points used to make the disk pseudo-absence selection (in the same projection system units as coord, see Details)
PA.dist.max	<i>(optional, default 0)</i> If pseudo-absence selection and PA.strategy = 'disk', a numeric defining the maximal distance to presence points used to make the disk pseudo-absence selection (in the same projection system units as coord, see Details)
PA.sre.quant	<i>(optional, default 0)</i> If pseudo-absence selection and PA.strategy = 'sre', a numeric between 0 and 0.5 defining the half-quantile used to make the sre pseudo-absence selection (see Details)
PA.fact.aggr	<i>(optional, default NULL)</i> If strategy = 'random' or strategy = 'disk', a integer defining the factor of aggregation to reduce the resolution
PA.user.table	<i>(optional, default NULL)</i> If pseudo-absence selection and PA.strategy = 'user.defined', a matrix or data.frame with as many rows as resp.var values, as many columns as PA.nb.rep, and containing TRUE or FALSE values defining which points will be used to build the species distribution model(s) for each repetition (see Details)
na.rm	<i>(optional, default TRUE)</i> A logical value defining whether points having one or several missing values for explanatory variables should be removed from the analysis or not
filter.raster	<i>(optional, default FALSE)</i> If env is of raster type, a logical value defining whether sp is to be filtered when several points occur in the same raster cell
seed.val	<i>(optional, default NULL)</i> An integer value corresponding to the new seed value to be set

### Slots

dir.name a character corresponding to the modeling folder

sp.name a character corresponding to the species name  
 coord a 2-columns data.frame containing the corresponding X and Y coordinates  
 data.species a vector containing the species observations (0, 1 or NA)  
 data.env.var a data.frame containing explanatory variables  
 data.mask a [SpatRaster](#) object containing the mask of the studied area  
 has.data.eval a logical value defining whether evaluation data is given  
 eval.coord (*optional, default NULL*)  
     A 2-columns data.frame containing the corresponding X and Y coordinates for evaluation data  
 eval.data.species (*optional, default NULL*)  
     A vector containing the species observations (0, 1 or NA) for evaluation data  
 eval.data.env.var (*optional, default NULL*)  
     A data.frame containing explanatory variables for evaluation data  
 PA.strategy a character corresponding to the pseudo-absence selection strategy  
 PA.table a data.frame containing the corresponding table of selected pseudo-absences (indicated by TRUE or FALSE) from the pa.tab list element returned by the [bm\\_PseudoAbsences](#) function

### Author(s)

Damien Georges

### See Also

[BIOMOD\\_FormatingData](#), [bm\\_PseudoAbsences](#), [bm\\_Tuning](#), [bm\\_CrossValidation](#), [BIOMOD\\_Modeling](#), [bm\\_RunModelsLoop](#)

Other Toolbox objects: [BIOMOD.ensemble.models.out](#), [BIOMOD.formated.data](#), [BIOMOD.models.options](#), [BIOMOD.models.out](#), [BIOMOD.options.dataset](#), [BIOMOD.options.default](#), [BIOMOD.projection.out](#), [BIOMOD.stored.data](#), [biomod2\\_ensemble\\_model](#), [biomod2\\_model](#)

### Examples

```

showClass("BIOMOD.formated.data.PA")

## ----- #
library(terra)

# Load species occurrences (6 species available)
data(DataSpecies)
head(DataSpecies)

# Select the name of the studied species
myRespName <- 'GuloGulo'

# Keep only presence informations
DataSpecies <- DataSpecies[which(DataSpecies[, myRespName] == 1), ]

```

```

# Get corresponding presence/absence data
myResp <- as.numeric(DataSpecies[, myRespName])

# Get corresponding XY coordinates
myRespXY <- DataSpecies[, c('X_WGS84', 'Y_WGS84')]

# Load environmental variables extracted from BIOCLIM (bio_3, bio_4, bio_7, bio_11 & bio_12)
data(bioclim_current)
myExpl <- terra::rast(bioclim_current)

## ----- #
# Format Data with pseudo-absences : random method
myBiomodData <- BIOMOD_FormatingData(resp.var = myResp,
                                   expl.var = myExpl,
                                   resp.xy = myRespXY,
                                   resp.name = myRespName,
                                   PA.nb.rep = 4,
                                   PA.strategy = 'random',
                                   PA.nb.absences = 1000)

myBiomodData
plot(myBiomodData)

```

---

BIOMOD.models.options [bm\\_ModelingOptions](#) *output object class*

---

## Description

Class returned by [bm\\_ModelingOptions](#) and used by [BIOMOD\\_Modeling](#)

## Usage

```

## S4 method for signature 'BIOMOD.models.options'
show(object)

## S4 method for signature 'BIOMOD.models.options'
print(x, dataset = "_allData_allRun")

```

## Arguments

object	a <a href="#">BIOMOD.models.options</a> object
x	a <a href="#">BIOMOD.models.options</a> object
dataset	a character corresponding to the name of a dataset contained in the arg.values slot of the <a href="#">BIOMOD.options.dataset</a> object for each model

**Slots**

`models` a vector containing model names for which options have been retrieved and defined, must be `algo.datatype.package.function`

`options` a list containing `BIOMOD.options.dataset` object for each model

**Author(s)**

Maya Gueguen

**See Also**

`BIOMOD.options.default`, `BIOMOD.options.dataset`, `bm_ModelingOptions`, `bm_Tuning`, `BIOMOD_Modeling`

Other Toolbox objects: `BIOMOD.ensemble.models.out`, `BIOMOD.formated.data`, `BIOMOD.formated.data.PA`, `BIOMOD.models.out`, `BIOMOD.options.dataset`, `BIOMOD.options.default`, `BIOMOD.projection.out`, `BIOMOD.stored.data`, `biomod2_ensemble_model`, `biomod2_model`

**Examples**

```
showClass("BIOMOD.models.options")
```

---

BIOMOD.models.out	BIOMOD_Modeling() <i>output object class</i>
-------------------	--

---

**Description**

Class returned by `BIOMOD_Modeling`, and used by `BIOMOD_LoadModels`, `BIOMOD_PresenceOnly`, `BIOMOD_Projection` and `BIOMOD_EnsembleModeling`

**Usage**

```
## S4 method for signature 'BIOMOD.models.out'
show(object)
```

**Arguments**

`object` a `BIOMOD.models.out` object

**Slots**

`modeling.id` a character corresponding to the name (ID) of the simulation set

`dir.name` a character corresponding to the modeling folder

`sp.name` a character corresponding to the species name

`expl.var.names` a vector containing names of explanatory variables

`models.computed` a vector containing names of computed models



`models.failed` a vector containing names of failed models  
`has.evaluation.data` a logical value defining whether evaluation data is given  
`scale.models` a logical value defining whether models have been rescaled or not  
`formatted.input.data` a `BIOMOD.stored.formated.data-class` object containing informations from `BIOMOD_FormatingData` object  
`calib.lines` a `BIOMOD.stored.data.frame-class` object containing calibration lines  
`models.options` a `BIOMOD.stored.options-class` object containing informations from `bm_ModelingOptions` object  
`models.evaluation` a `BIOMOD.stored.data.frame-class` object containing models evaluation  
`variables.importance` a `BIOMOD.stored.data.frame-class` object containing variables importance  
`models.prediction` a `BIOMOD.stored.data.frame-class` object containing models predictions  
`models.prediction.eval` a `BIOMOD.stored.data.frame-class` object containing models predictions for evaluation data  
`link` a character containing the file name of the saved object

### Author(s)

Damien Georges

### See Also

[BIOMOD\\_Modeling](#), [BIOMOD\\_LoadModels](#), [BIOMOD\\_PresenceOnly](#), [BIOMOD\\_Projection](#), [BIOMOD\\_EnsembleModeling](#), [bm\\_VariablesImportance](#), [bm\\_PlotEvalMean](#), [bm\\_PlotEvalBoxplot](#), [bm\\_PlotVarImpBoxplot](#), [bm\\_PlotResponseCurves](#)

Other Toolbox objects: [BIOMOD.ensemble.models.out](#), [BIOMOD.formated.data](#), [BIOMOD.formated.data.PA](#), [BIOMOD.models.options](#), [BIOMOD.options.dataset](#), [BIOMOD.options.default](#), [BIOMOD.projection.out](#), [BIOMOD.stored.data](#), [biomod2\\_ensemble\\_model](#), [biomod2\\_model](#)

### Examples

```

showClass("BIOMOD.models.out")

## ----- #
library(terra)

# Load species occurrences (6 species available)
data(DataSpecies)
head(DataSpecies)

# Select the name of the studied species
myRespName <- 'GuloGulo'

# Get corresponding presence/absence data
myResp <- as.numeric(DataSpecies[, myRespName])

# Get corresponding XY coordinates
  
```

```

myRespXY <- DataSpecies[, c('X_WGS84', 'Y_WGS84')]

# Load environmental variables extracted from BIOCLIM (bio_3, bio_4, bio_7, bio_11 & bio_12)
data(bioclim_current)
myExpl <- terra::rast(bioclim_current)

## ----- #
# Format Data with true absences
myBiomodData <- BIOMOD_FormatingData(resp.var = myResp,
                                   expl.var = myExpl,
                                   resp.xy = myRespXY,
                                   resp.name = myRespName)

## ----- #
# Model single models
myBiomodModelOut <- BIOMOD_Modeling(bm.format = myBiomodData,
                                   modeling.id = 'AllModels',
                                   models = c('RF', 'GLM'),
                                   CV.strategy = 'random',
                                   CV.nb.rep = 2,
                                   CV.perc = 0.8,
                                   OPT.strategy = 'bigboss',
                                   metric.eval = c('TSS','ROC'),
                                   var.import = 3,
                                   seed.val = 42)

myBiomodModelOut

```

---

BIOMOD.options.dataset

[bm\\_ModelingOptions](#) *output object class*

---

## Description

Class returned by [bm\\_ModelingOptions](#) (a list of BIOMOD.options.dataset more exactly), and used by [BIOMOD\\_Modeling](#)

## Usage

```

## S4 method for signature 'character'
BIOMOD.options.dataset(
  mod,
  typ,
  pkg,
  fun,
  strategy,

```

```

    user.val = NULL,
    user.base = NULL,
    tuning.fun = NULL,
    bm.format = NULL,
    calib.lines = NULL
  )

## S4 method for signature 'BIOMOD.options.dataset'
show(object)

## S4 method for signature 'BIOMOD.options.dataset'
print(x, dataset = "_allData_allRun")

```

### Arguments

mod	a character corresponding to the model name to be computed, must be either ANN, CTA, FDA, GAM, GBM, GLM, MARS, MAXENT, MAXNET, RF, SRE, XGBOOST
typ	a character corresponding to the data type to be used, must be either binary, binary.PA, abundance, compositional
pkg	a character corresponding to the package containing the model function to be called
fun	a character corresponding to the model function name to be called
strategy	a character corresponding to the method to select models' parameters values, must be either default, bigboss, user.defined, tuned
user.val	(optional, default NULL) A list containing parameters values
user.base	(optional, default NULL) A character, default or bigboss used when strategy = 'user.defined'. It sets the bases of parameters to be modified by user defined values.
tuning.fun	(optional, default NULL) A character corresponding to the model function name to be called through <a href="#">train</a> function for tuning parameters
bm.format	(optional, default NULL) A <a href="#">BIOMOD.formated.data</a> or <a href="#">BIOMOD.formated.data.PA</a> object returned by the <a href="#">BIOMOD_FormatingData</a> function
calib.lines	(optional, default NULL) A data.frame object returned by <a href="#">get_calib_lines</a> or <a href="#">bm_CrossValidation</a> functions, to explore the distribution of calibration and validation datasets
object	a <a href="#">BIOMOD.options.dataset</a> object
x	a <a href="#">BIOMOD.options.dataset</a> object
dataset	a character corresponding to the name of a dataset contained in the arg.values slot

**Slots**

`model` a character corresponding to the model  
`type` a character corresponding to the data type (binary, binary.PA, abundance, compositional)  
`package` a character corresponding to the package containing the model function to be called  
`func` a character corresponding to the model function name to be called  
`args.names` a vector containing character corresponding to the model function arguments  
`args.default` a list containing for each dataset the default values for all arguments listed in `args.names`  
`args.values` a list containing for each dataset the to-be-used values for all arguments listed in `args.names`

**Author(s)**

Maya Gueguen

**See Also**

[BIOMOD.options.default](#), [bm\\_ModelingOptions](#), [bm\\_Tuning](#), [BIOMOD\\_Modeling](#), [bm\\_RunModelsLoop](#)

Other Toolbox objects: [BIOMOD.ensemble.models.out](#), [BIOMOD.formated.data](#), [BIOMOD.formated.data.PA](#), [BIOMOD.models.options](#), [BIOMOD.models.out](#), [BIOMOD.options.default](#), [BIOMOD.projection.out](#), [BIOMOD.stored.data](#), [biomod2\\_ensemble\\_model](#), [biomod2\\_model](#)

**Examples**

```
showClass("BIOMOD.options.dataset")
```

---

BIOMOD.options.default

[bm\\_ModelingOptions](#) *output object class*

---

**Description**

Class returned by [bm\\_ModelingOptions](#) (a list of [BIOMOD.options.dataset](#) more exactly), and used by [BIOMOD\\_Modeling](#)

**Usage**

```
## S4 method for signature 'character,character'
BIOMOD.options.default(mod, typ, pkg, fun)
```

**Arguments**

mod	a character corresponding to the model name to be computed, must be either ANN, CTA, FDA, GAM, GBM, GLM, MARS, MAXENT, MAXNET, RF, SRE, XGBOOST
typ	a character corresponding to the data type to be used, must be either binary, binary.PA, abundance, compositional
pkg	a character corresponding to the package containing the model function to be called
fun	a character corresponding to the model function name to be called

**Slots**

model	a character corresponding to the model
type	a character corresponding to the data type (binary, binary.PA, abundance, compositional)
package	a character corresponding to the package containing the model function to be called
func	a character corresponding to the model function name to be called
args.names	a vector containing character corresponding to the model function arguments
args.default	a list containing for each dataset the default values for all arguments listed in args.names

**Author(s)**

Maya Gueguen

**See Also**

[BIOMOD.options.dataset](#), [bm\\_ModelingOptions](#), [bm\\_Tuning](#), [BIOMOD\\_Modeling](#), [bm\\_RunModelsLoop](#)

Other Toolbox objects: [BIOMOD.ensemble.models.out](#), [BIOMOD.formated.data](#), [BIOMOD.formated.data.PA](#), [BIOMOD.models.options](#), [BIOMOD.models.out](#), [BIOMOD.options.dataset](#), [BIOMOD.projection.out](#), [BIOMOD.stored.data](#), [biomod2\\_ensemble\\_model](#), [biomod2\\_model](#)

**Examples**

```
showClass("BIOMOD.options.default")
```

---

BIOMOD.projection.out [BIOMOD\\_Projection\(\)](#) *output object class*

---

**Description**

Class returned by [BIOMOD\\_Projection](#), and used by [BIOMOD\\_EnsembleForecasting](#)

**Usage**

```
## S4 method for signature 'BIOMOD.projection.out,missing'
plot(
  x,
  coord = NULL,
  plot.output,
  do.plot = TRUE,
  std = TRUE,
  scales,
  size,
  maxcell = 5e+05,
  ...
)

## S4 method for signature 'BIOMOD.projection.out'
show(object)
```

**Arguments**

x	a <a href="#">BIOMOD.projection.out</a> object
coord	a 2-columns data.frame containing the corresponding X and Y
plot.output	( <i>optional, default facet</i> ) a character determining the type of output: with <code>plot.output = 'list'</code> the function will return a list of plots (one plot per model); with <code>'facet'</code> ; with <code>plot.output = 'facet'</code> the function will return a single plot with all asked projections as facet.
do.plot	( <i>optional, default TRUE</i> ) a boolean determining whether the plot should be displayed or just returned.
std	( <i>optional, default TRUE</i> ) a boolean controlling the limits of the color scales. With <code>std = TRUE</code> color scales are displayed between 0 and 1 (or 1000). With <code>std = FALSE</code> color scales are displayed between 0 and the maximum value observed.
scales	( <i>optional, default fixed</i> ) a character determining whether x and y scales are shared among facet. Argument passed to <a href="#">facet_wrap</a> . Possible values: <code>'fixed'</code> , <code>'free_x'</code> , <code>'free_y'</code> , <code>'free'</code> .
size	( <i>optional, default 0.75</i> ) a numeric determining the size of points on the plots and passed to <a href="#">geom_point</a> .
maxcell	maximum number of cells to plot. Argument transmitted to <a href="#">plot</a> .
...	additional parameters to be passed to <a href="#">get_predictions</a> to select the models that will be plotted
object	a <a href="#">BIOMOD.projection.out</a> object

**Slots**

`modeling.id` a character corresponding to the name (ID) of the simulation set  
`proj.name` a character corresponding to the projection name  
`dir.name` a character corresponding to the modeling folder

sp.name a character corresponding to the species name  
 expl.var.names a vector containing names of explanatory variables  
 coord a 2-columns matrix or data.frame containing the corresponding X and Y coordinates used to project the species distribution model(s)  
 scale.models a logical value defining whether models have been rescaled or not  
 models.projected a vector containing names of projected models  
 models.out a [BIOMOD.stored.data](#) object  
 type a character corresponding to the class of the val slot of the proj.out slot  
 proj.out a [BIOMOD.stored.data](#) object

**Author(s)**

Damien Georges

**See Also**

[BIOMOD\\_Projection](#), [BIOMOD\\_EnsembleForecasting](#)

Other Toolbox objects: [BIOMOD.ensemble.models.out](#), [BIOMOD.formated.data](#), [BIOMOD.formated.data.PA](#), [BIOMOD.models.options](#), [BIOMOD.models.out](#), [BIOMOD.options.dataset](#), [BIOMOD.options.default](#), [BIOMOD.stored.data](#), [biomod2\\_ensemble\\_model](#), [biomod2\\_model](#)

**Examples**

```

showClass("BIOMOD.projection.out")

## ----- #
library(terra)

# Load species occurrences (6 species available)
data(DataSpecies)
head(DataSpecies)

# Select the name of the studied species
myRespName <- 'GuloGulo'

# Get corresponding presence/absence data
myResp <- as.numeric(DataSpecies[, myRespName])

# Get corresponding XY coordinates
myRespXY <- DataSpecies[, c('X_WGS84', 'Y_WGS84')]

# Load environmental variables extracted from BIOCLIM (bio_3, bio_4, bio_7, bio_11 & bio_12)
data(bioclim_current)
myExpl <- terra::rast(bioclim_current)

## ----- #
file.out <- paste0(myRespName, "/", myRespName, ".AllModels.models.out")

```

```

if (file.exists(file.out)) {
  myBiomodModelOut <- get(load(file.out))
} else {

  # Format Data with true absences
  myBiomodData <- BIOMOD_FormatingData(resp.var = myResp,
                                       expl.var = myExpl,
                                       resp.xy = myRespXY,
                                       resp.name = myRespName)

  # Model single models
  myBiomodModelOut <- BIOMOD_Modeling(bm.format = myBiomodData,
                                     modeling.id = 'AllModels',
                                     models = c('RF', 'GLM'),
                                     CV.strategy = 'random',
                                     CV.nb.rep = 2,
                                     CV.perc = 0.8,
                                     OPT.strategy = 'bigboss',
                                     metric.eval = c('TSS', 'ROC'),
                                     var.import = 3,
                                     seed.val = 42)
}

## ----- #
# Project single models
myBiomodProj <- BIOMOD_Projection(bm.mod = myBiomodModelOut,
                                 proj.name = 'Current',
                                 new.env = myExpl,
                                 models.chosen = 'all',
                                 metric.binary = 'all',
                                 metric.filter = 'all',
                                 build.clamping.mask = TRUE)

myBiomodProj
plot(myBiomodProj)

```

---

BIOMOD.stored.data      [BIOMOD\\_Modeling](#) and [BIOMOD\\_EnsembleModeling](#) output object  
class

---

## Description

Classes used by [BIOMOD\\_Modeling](#) and [BIOMOD\\_EnsembleModeling](#) to build their output object (see [BIOMOD.models.out](#) objects)

## Details

BIOMOD.stored.data is the basic object containing the slots inMemory and link.



All listed classes below are derived from `BIOMOD.stored.data`, and contain a `val` slot of specific type :

- `BIOMOD.stored.data.frame` : `val` is a `data.frame`
- `BIOMOD.stored.SpatRaster` : `val` is a [PackedSpatRaster](#)
- `BIOMOD.stored.files` : `val` is a character
- `BIOMOD.stored.formated.data` : `val` is a [BIOMOD.formated.data](#) object
- `BIOMOD.stored.options` : `val` is a [BIOMOD.models.options](#) object
- `BIOMOD.stored.models.out` : `val` is a [BIOMOD.models.out](#) object

### Slots

`inMemory` a logical defining whether the `val` slot has been loaded in memory or not

`link` a character containing the file name of the saved `val` slot

`val` an object of type depending on the `BIOMOD.stored.[...]` class (see Details)

### Author(s)

Damien Georges

### See Also

[BIOMOD.formated.data](#), [BIOMOD.models.out](#), [BIOMOD\\_Modeling](#), [BIOMOD\\_EnsembleModeling](#), [BIOMOD\\_Projection](#), [BIOMOD\\_EnsembleForecasting](#)

Other Toolbox objects: [BIOMOD.ensemble.models.out](#), [BIOMOD.formated.data](#), [BIOMOD.formated.data.PA](#), [BIOMOD.models.options](#), [BIOMOD.models.out](#), [BIOMOD.options.dataset](#), [BIOMOD.options.default](#), [BIOMOD.projection.out](#), [biomod2\\_ensemble\\_model](#), [biomod2\\_model](#)

### Examples

```
showClass("BIOMOD.stored.data")
showClass("BIOMOD.stored.data.frame")
showClass("BIOMOD.stored.SpatRaster")
showClass("BIOMOD.stored.files")
showClass("BIOMOD.stored.formated.data")
showClass("BIOMOD.stored.options")
showClass("BIOMOD.stored.models.out")
```

---

```
biomod2_ensemble_model
      Ensemble model output object class (when running
      BIOMOD_EnsembleModeling())
```

---

## Description

Class created by [BIOMOD\\_EnsembleModeling](#)

## Usage

```
## S4 method for signature 'biomod2_ensemble_model'
show(object)
```

## Arguments

object            a [biomod2\\_ensemble\\_model](#) object

## Details

biomod2\_model is the basic object for **biomod2** ensemble species distribution models. All listed classes below are derived from biomod2\_model, and have a model\_class slot specific value :

- biomod2\_ensemble\_model : model\_class is EM
- EMmean\_biomod2\_model : model\_class is EMmean
- EMmedian\_biomod2\_model : model\_class is EMmedian
- EMcv\_biomod2\_model : model\_class is EMcv
- EMci\_biomod2\_model : model\_class is EMci
- EMca\_biomod2\_model : model\_class is EMca
- EMwmean\_biomod2\_model : model\_class is EMwmean

## Slots

modeling.id a character corresponding to the name (ID) of the simulation set  
 model\_name a character corresponding to the model name  
 model\_class a character corresponding to the model class  
 model\_options a list containing the model options  
 model the corresponding model object  
 scaling\_model the corresponding scaled model object  
 dir\_name a character corresponding to the modeling folder  
 resp\_name a character corresponding to the species name  
 expl\_var\_names a vector containing names of explanatory variables

`expl_var_type` a vector containing classes of explanatory variables  
`expl_var_range` a list containing ranges of explanatory variables  
`model_evaluation` a data.frame containing the model evaluations  
`model_variables_importance` a data.frame containing the model variables importance

### Author(s)

Damien Georges

### See Also

[biomod2\\_model](#), [BIOMOD\\_EnsembleModeling](#)

Other Toolbox objects: [BIOMOD.ensemble.models.out](#), [BIOMOD.formated.data](#), [BIOMOD.formated.data.PA](#), [BIOMOD.models.options](#), [BIOMOD.models.out](#), [BIOMOD.options.dataset](#), [BIOMOD.options.default](#), [BIOMOD.projection.out](#), [BIOMOD.stored.data](#), [biomod2\\_model](#)

### Examples

```

showClass("biomod2_ensemble_model")
showClass("EMmean_biomod2_model")
showClass("EMmedian_biomod2_model")
showClass("EMcv_biomod2_model")
showClass("EMci_biomod2_model")
showClass("EMca_biomod2_model")
showClass("EMwmean_biomod2_model")
  
```

---

<code>biomod2_model</code>	<i>Single model output object class (when running <code>BIOMOD_Modeling()</code>)</i>
----------------------------	---

---

### Description

Class created by [BIOMOD\\_Modeling](#) and [bm\\_RunModel](#)

### Usage

```

## S4 method for signature 'biomod2_model'
show(object)
  
```

### Arguments

`object` a [biomod2\\_model](#) object

## Details

biomod2\_model is the basic object for **biomod2** single species distribution models.

All listed classes below are derived from biomod2\_model, and have a model\_class slot specific value :

- ANN\_biomod2\_model : model\_class is ANN
- CTA\_biomod2\_model : model\_class is CTA
- FDA\_biomod2\_model : model\_class is FDA
- GBM\_biomod2\_model : model\_class is GBM
- GLM\_biomod2\_model : model\_class is GLM
- MARS\_biomod2\_model : model\_class is MARS
- MAXENT\_biomod2\_model : model\_class is MAXENT
- MAXNET\_biomod2\_model : model\_class is MAXNET
- RF\_biomod2\_model : model\_class is RF
- RFd\_biomod2\_model : model\_class is RFd
- SRE\_biomod2\_model : model\_class is SRE

## Slots

model\_name a character corresponding to the model name

model\_class a character corresponding to the model class

model\_options a list containing the model options

model the corresponding model object

scaling\_model the corresponding scaled model object

dir\_name a character corresponding to the modeling folder

resp\_name a character corresponding to the species name

expl\_var\_names a vector containing names of explanatory variables

expl\_var\_type a vector containing classes of explanatory variables

expl\_var\_range a list containing ranges of explanatory variables

model\_evaluation a data.frame containing the model evaluations

model\_variables\_importance a data.frame containing the model variables importance

## Author(s)

Damien Georges

## See Also

[BIOMOD\\_Modeling](#), [bm\\_RunModel](#)

Other Toolbox objects: [BIOMOD.ensemble.models.out](#), [BIOMOD.formated.data](#), [BIOMOD.formated.data.PA](#), [BIOMOD.models.options](#), [BIOMOD.models.out](#), [BIOMOD.options.dataset](#), [BIOMOD.options.default](#), [BIOMOD.projection.out](#), [BIOMOD.stored.data](#), [biomod2\\_ensemble\\_model](#)

## Examples

```
showClass("biomod2_model")
showClass("ANN_biomod2_model")
showClass("CTA_biomod2_model")
showClass("FDA_biomod2_model")
showClass("GAM_biomod2_model")
showClass("GBM_biomod2_model")
showClass("GLM_biomod2_model")
showClass("MARS_biomod2_model")
showClass("MAXENT_biomod2_model")
showClass("MAXNET_biomod2_model")
showClass("RF_biomod2_model")
showClass("RFd_biomod2_model")
showClass("SRE_biomod2_model")
```

---

BIOMOD\_EnsembleForecasting

*Project ensemble species distribution models onto new environment*

---

## Description

This function allows to project ensemble models built with the [BIOMOD\\_EnsembleModeling](#) function onto new environmental data (*which can represent new areas, resolution or time scales for example*).

## Usage

```
BIOMOD_EnsembleForecasting(  
  bm.em,  
  bm.proj = NULL,  
  proj.name = NULL,  
  new.env = NULL,  
  new.env.xy = NULL,  
  models.chosen = "all",  
  metric.binary = NULL,  
  metric.filter = NULL,  
  compress = TRUE,  
  nb.cpu = 1,  
  na.rm = TRUE,  
  ...  
)
```

## Arguments

**bm.em** a `BIOMOD.ensemble.models.out` object returned by the [BIOMOD\\_EnsembleModeling](#) function

bm.proj	a <a href="#">BIOMOD.projection.out</a> object returned by the <a href="#">BIOMOD_Projection</a> function
proj.name	<i>(optional, default NULL)</i> If bm.proj = NULL, a character corresponding to the name (ID) of the projection set ( <i>a new folder will be created within the simulation folder with this name</i> )
new.env	<i>(optional, default NULL)</i> If bm.proj = NULL, a matrix, data.frame or <a href="#">SpatRaster</a> object containing the new explanatory variables (in columns or layers, with names matching the variables names given to the <a href="#">BIOMOD_FormatingData</a> function to build bm.mod) that will be used to project the species distribution model(s) <i>Note that old format from <b>raster</b> are still supported such as RasterStack objects.</i>
new.env.xy	<i>(optional, default NULL)</i> If new.env is a matrix or a data.frame, a 2-columns matrix or data.frame containing the corresponding X and Y coordinates that will be used to project the ensemble species distribution model(s)
models.chosen	a vector containing model names to be kept, must be either all or a sub-selection of model names that can be obtained with the <a href="#">get_built_models</a> function
metric.binary	<i>(optional, default NULL)</i> A vector containing evaluation metric names to be used to transform prediction values into binary values based on models evaluation scores obtained with the <a href="#">BIOMOD_Modeling</a> function. Must be among all (same evaluation metrics than those of modeling.output) or POD, FAR, POFD, SR, ACCURACY, BIAS, ROC, TSS, KAPPA, OR, ORSS, CSI, ETS, BOYCE, MPA
metric.filter	<i>(optional, default NULL)</i> A vector containing evaluation metric names to be used to transform prediction values into filtered values based on models evaluation scores obtained with the <a href="#">BIOMOD_Modeling</a> function. Must be among all (same evaluation metrics than those of modeling.output) or POD, FAR, POFD, SR, ACCURACY, BIAS, ROC, TSS, KAPPA, OR, ORSS, CSI, ETS, BOYCE, MPA
compress	<i>(optional, default TRUE)</i> A logical or a character value defining whether and how objects should be compressed when saved on hard drive, must be either TRUE, FALSE, xz or gzip (see Details)
nb.cpu	<i>(optional, default 1)</i> An integer value corresponding to the number of computing resources to be used to parallelize the single models computation
na.rm	<i>(optional, default TRUE)</i> A boolean defining whether Ensemble Model projection should ignore NA in Individual Model projection. Argument ignored by EWmean ensemble algorithm.
...	<i>(optional, see Details)</i>

## Details

If `models.chosen = 'all'`, projections are done for all calibration and pseudo absences runs if applicable.

These projections may be used later by the [BIOMOD\\_EnsembleForecasting](#) function.

If `build.clamping.mask = TRUE`, a raster file will be saved within the projection folder. This mask values will correspond to the number of variables in each pixel that are out of their calibration / validation range, identifying locations where predictions are uncertain.

... can take the following values :

- `on_0_1000` : a logical value defining whether 0 - 1 probabilities are to be converted to 0 - 1000 scale to save memory on backup
- `do.stack` : a logical value defining whether all projections are to be saved as one `SpatRaster` object or several `SpatRaster` files (*the default if projections are too heavy to be all loaded at once in memory*)
- `keep.in.memory` : a logical value defining whether all projections are to be kept loaded at once in memory, or only links pointing to hard drive are to be returned
- `output.format` : a character value corresponding to the projections saving format on hard drive, must be either `.grd`, `.img`, `.tif` or `.RData` (the default if `new.env` is given as `matrix` or `data.frame`)

## Value

A `BIOMOD.projection.out` object containing models projections, or links to saved outputs.

Models projections are stored out of R (for memory storage reasons) in `proj.name` folder created in the current working directory :

1. the output is a `data.frame` if `new.env` is a `matrix` or a `data.frame`
2. it is a `SpatRaster` if `new.env` is a `SpatRaster` (or several `SpatRaster` objects, if `new.env` is too large)
3. raw projections, as well as binary and filtered projections (if asked), are saved in the `proj.name` folder

## Author(s)

Wilfried Thuiller, Damien Georges, Robin Engler

## See Also

[BIOMOD\\_FormatingData](#), [bm\\_ModelingOptions](#), [BIOMOD\\_Modeling](#), [BIOMOD\\_EnsembleModeling](#), [BIOMOD\\_RangeSize](#)

Other Main functions: [BIOMOD\\_EnsembleModeling\(\)](#), [BIOMOD\\_FormatingData\(\)](#), [BIOMOD\\_LoadModels\(\)](#), [BIOMOD\\_Modeling\(\)](#), [BIOMOD\\_Projection\(\)](#), [BIOMOD\\_RangeSize\(\)](#)

**Examples**

```

library(terra)
# Load species occurrences (6 species available)
data(DataSpecies)
head(DataSpecies)

# Select the name of the studied species
myRespName <- 'GuloGulo'

# Get corresponding presence/absence data
myResp <- as.numeric(DataSpecies[, myRespName])

# Get corresponding XY coordinates
myRespXY <- DataSpecies[, c('X_WGS84', 'Y_WGS84')]

# Load environmental variables extracted from BIOCLIM (bio_3, bio_4, bio_7, bio_11 & bio_12)
data(bioclim_current)
myExpl <- terra::rast(bioclim_current)

# ----- #
file.out <- paste0(myRespName, "/", myRespName, ".AllModels.models.out")
if (file.exists(file.out)) {
  myBiomodModelOut <- get(load(file.out))
} else {

  # Format Data with true absences
  myBiomodData <- BIOMOD_FormatingData(resp.var = myResp,
                                     expl.var = myExpl,
                                     resp.xy = myRespXY,
                                     resp.name = myRespName)

  # Model single models
  myBiomodModelOut <- BIOMOD_Modeling(bm.format = myBiomodData,
                                     modeling.id = 'AllModels',
                                     models = c('RF', 'GLM'),
                                     CV.strategy = 'random',
                                     CV.nb.rep = 2,
                                     CV.perc = 0.8,
                                     OPT.strategy = 'bigboss',
                                     metric.eval = c('TSS', 'ROC'),
                                     var.import = 3,
                                     seed.val = 42)
}

file.proj <- paste0(myRespName, "/proj_Current/", myRespName, ".Current.projection.out")
if (file.exists(file.proj)) {
  myBiomodProj <- get(load(file.proj))
} else {

```



```

# Project single models
myBiomodProj <- BIOMOD_Projection(bm.mod = myBiomodModelOut,
                                proj.name = 'Current',
                                new.env = myExpl,
                                models.chosen = 'all',
                                build.clamping.mask = TRUE)
}

file.EM <- paste0(myRespName, "/", myRespName, ".AllModels.ensemble.models.out")
if (file.exists(file.EM)) {
  myBiomodEM <- get(load(file.EM))
} else {

# Model ensemble models
myBiomodEM <- BIOMOD_EnsembleModeling(bm.mod = myBiomodModelOut,
                                       models.chosen = 'all',
                                       em.by = 'all',
                                       em.algo = c('EMmean', 'EMca'),
                                       metric.select = c('TSS'),
                                       metric.select.thresh = c(0.7),
                                       metric.eval = c('TSS', 'ROC'),
                                       var.import = 3,
                                       seed.val = 42)
}

# ----- #
# Project ensemble models (from single projections)
myBiomodEMProj <- BIOMOD_EnsembleForecasting(bm.em = myBiomodEM,
                                             bm.proj = myBiomodProj,
                                             models.chosen = 'all',
                                             metric.binary = 'all',
                                             metric.filter = 'all')

# Project ensemble models (building single projections)
myBiomodEMProj <- BIOMOD_EnsembleForecasting(bm.em = myBiomodEM,
                                             proj.name = 'CurrentEM',
                                             new.env = myExpl,
                                             models.chosen = 'all',
                                             metric.binary = 'all',
                                             metric.filter = 'all')

myBiomodEMProj
plot(myBiomodEMProj)

```

## Description

This function allows to combine a range of models built with the [BIOMOD\\_Modeling](#) function in one (or several) ensemble model. Modeling uncertainty can be assessed as well as variables importance, ensemble predictions can be evaluated against original data, and created ensemble models can be projected over new conditions (see Details).

## Usage

```
BIOMOD_EnsembleModeling(
  bm.mod,
  models.chosen = "all",
  em.by = "PA+run",
  em.algo,
  metric.select = "all",
  metric.select.thresh = NULL,
  metric.select.table = NULL,
  metric.select.dataset = NULL,
  metric.eval = c("KAPPA", "TSS", "ROC"),
  var.import = 0,
  EMci.alpha = 0.05,
  EMwmean.decay = "proportional",
  nb.cpu = 1,
  seed.val = NULL,
  do.progress = TRUE,
  prob.mean,
  prob.median,
  prob.cv,
  prob.ci,
  committee.averaging,
  prob.mean.weight,
  prob.mean.weight.decay,
  prob.ci.alpha
)
```

## Arguments

<code>bm.mod</code>	a <a href="#">BIOMOD.models.out</a> object returned by the <a href="#">BIOMOD_Modeling</a> function
<code>models.chosen</code>	a vector containing model names to be kept, must be either <code>all</code> or a sub-selection of model names that can be obtained with the <a href="#">get_built_models</a> function
<code>em.by</code>	a character corresponding to the way kept models will be combined to build the ensemble models, must be among <code>all</code> , <code>algo</code> , <code>PA</code> , <code>PA+algo</code> , <code>PA+run</code>
<code>em.algo</code>	a vector corresponding to the ensemble models that will be computed, must be among <code>'EMmean'</code> , <code>'EMmedian'</code> , <code>'EMcv'</code> , <code>'EMci'</code> , <code>'EMca'</code> , <code>'EMwmean'</code>
<code>metric.select</code>	a vector containing evaluation metric names to be used together with <code>metric.select.thresh</code> to exclude single models based on their evaluation scores (for ensemble methods like probability weighted mean or committee averaging). Must be among <code>all</code>

	(same evaluation metrics than those of <code>bm.mod</code> ), <code>user.defined</code> (and defined through <code>metric.select.table</code> ) or <code>POD</code> , <code>FAR</code> , <code>POFD</code> , <code>SR</code> , <code>ACCURACY</code> , <code>BIAS</code> , <code>ROC</code> , <code>TSS</code> , <code>KAPPA</code> , <code>OR</code> , <code>ORSS</code> , <code>CSI</code> , <code>ETS</code> , <code>BOYCE</code> , <code>MPA</code>
<code>metric.select.thresh</code>	<i>(optional, default NULL)</i> A vector of numeric values corresponding to the minimum scores (one for each <code>metric.select</code> ) below which single models will be excluded from the ensemble model building
<code>metric.select.table</code>	<i>(optional, default NULL)</i> If <code>metric.select = 'user.defined'</code> , a data.frame containing evaluation scores calculated for each single models and that will be compared to <code>metric.select.thresh</code> values to exclude some of them from the ensemble model building, with <code>metric.select</code> rownames, and <code>models.chosen</code> colnames
<code>metric.select.dataset</code>	<i>(optional, default 'validation' if possible)</i> . A character determining which dataset should be used to filter and/or weigh the ensemble models should be among 'evaluation', 'validation' or 'calibration'.
<code>metric.eval</code>	a vector containing evaluation metric names to be used, must be among <code>POD</code> , <code>FAR</code> , <code>POFD</code> , <code>SR</code> , <code>ACCURACY</code> , <code>BIAS</code> , <code>ROC</code> , <code>TSS</code> , <code>KAPPA</code> , <code>OR</code> , <code>ORSS</code> , <code>CSI</code> , <code>ETS</code> , <code>BOYCE</code> , <code>MPA</code>
<code>var.import</code>	<i>(optional, default NULL)</i> An integer corresponding to the number of permutations to be done for each variable to estimate variable importance
<code>EMci.alpha</code>	<i>(optional, default 0.05)</i> A numeric value corresponding to the significance level to estimate confidence interval
<code>EMwmean.decay</code>	<i>(optional, default proportional)</i> A value defining the relative importance of the weights (if 'EMwmean' was given to argument <code>em.algo</code> ). A high value will strongly discriminate <i>good</i> models from the <i>bad</i> ones (see Details), while <code>proportional</code> will attribute weights proportionally to the models evaluation scores
<code>nb.cpu</code>	<i>(optional, default 1)</i> An integer value corresponding to the number of computing resources to be used to parallelize the single models predictions and the ensemble models computation
<code>seed.val</code>	<i>(optional, default NULL)</i> An integer value corresponding to the new seed value to be set
<code>do.progress</code>	<i>(optional, default TRUE)</i> A logical value defining whether the progress bar is to be rendered or not
<code>prob.mean</code>	<i>(deprecated, please use em.algo instead)</i> A logical value defining whether to compute the mean probabilities across predictions or not
<code>prob.median</code>	<i>(deprecated, please use em.algo instead)</i> A logical value defining whether to compute the median probabilities across predictions or not

prob.cv	( <i>deprecated</i> , please use em.algo instead) A logical value defining whether to compute the coefficient of variation across predictions or not
prob.ci	( <i>deprecated</i> , please use em.algo instead) A logical value defining whether to compute the confidence interval around the prob.mean ensemble model or not
committee.averaging	( <i>deprecated</i> , please use em.algo instead) A logical value defining whether to compute the committee averaging across predictions or not
prob.mean.weight	( <i>deprecated</i> , please use em.algo instead) A logical value defining whether to compute the weighted sum of probabilities across predictions or not
prob.mean.weight.decay	( <i>deprecated</i> , please use EMwmean.decay instead) old argument name for EMwmean.decay
prob.ci.alpha	( <i>deprecated</i> , please use EMci.alpha instead) old argument name for EMci.alpha

## Details

**Models sub-selection** (models.chosen) Applying [get\\_built\\_models](#) function to the `bm.mod` object gives the names of the single models created with the [BIOMOD\\_Modeling](#) function. The `models.chosen` argument can take either a sub-selection of these single model names, or the all default value, to decide which single models will be used for the ensemble model building.

**Models assembly rules** (em.by) Single models built with the [BIOMOD\\_Modeling](#) function can be combined in 5 different ways to obtain ensemble models :

- PA+run : each combination of pseudo-absence and repetition datasets is done, *merging* algorithms together
- PA+algo : each combination of pseudo-absence and algorithm datasets is done, *merging* repetitions together
- PA : pseudo-absence datasets are considered individually, *merging* algorithms and repetitions together
- algo : algorithm datasets are considered individually, *merging* pseudo-absence and repetitions together
- all : all models are combined into one

Hence, depending on the chosen method, the number of ensemble models built will vary.

*Be aware that if no evaluation data was given to the [BIOMOD\\_FormatingData](#) function, some ensemble model evaluations may be biased due to difference in data used for single model evaluations. **Be aware that all of these combinations are allowed, but some may not make sense depending mainly on how pseudo-absence datasets have been built and whether all of them have been used for all single models or not (see PA.nb.absences and models.pa parameters in [BIOMOD\\_FormatingData](#) and [BIOMOD\\_Modeling](#) functions respectively).***

**Evaluation metrics** • `metric.select` : the selected metrics must be chosen among the ones used within the [BIOMOD\\_Modeling](#) function to build the `model.output` object, unless

`metric.select` = 'user.defined' and therefore values will be provided through the `metric.select.table` parameter.

In the case of the selection of several metrics, they will be used at different steps of the ensemble modeling function :

1. remove *low quality* single models, having a score lower than `metric.select.thresh`
  2. perform the binary transformation needed if 'EMca' was given to argument `em.algo`
  3. weight models if 'EMwmean' was given to argument `em.algo`
- `metric.select.thresh` : as many values as evaluation metrics selected with the `metric.select` parameter, and defining the corresponding quality thresholds below which the single models will be excluded from the ensemble model building.
  - `metric.select.table` : a data.frame must be given if `metric.select` = 'user.defined' to allow the use of evaluation metrics other than those calculated within **biomod2**. The data.frame must contain as many columns as models.chosen with matching names, and as many rows as evaluation metrics to be used. The number of rows must match the length of the `metric.select.thresh` parameter. The values contained in the data.frame will be compared to those defined in `metric.select.thresh` to remove *low quality* single models from the ensemble model building.
  - `metric.select.dataset` : a character determining the dataset which evaluation metric should be used to filter and/or weigh the ensemble models. Should be among `evaluation`, `validation` or `calibration`. By default BIOMOD\_EnsembleModeling will use the validation dataset unless no validation is available in which case calibration dataset are used.
  - `metric.eval` : the selected metrics will be used to validate/evaluate the ensemble models built

**Ensemble-models algorithms** The set of models to be calibrated on the data.

6 modeling techniques are currently available :

- `EMmean` : Mean of probabilities over the selected models. Old name: `prob.mean`
- `EMmedian` : Median of probabilities over the selected models  
The median is less sensitive to outliers than the mean, however it requires more computation time and memory as it loads all predictions (on the contrary to the mean or the weighted mean). Old name: `prob.median`
- `EMcv` : Coefficient of variation (`sd / mean`) of probabilities over the selected models  
This model is not scaled. It will be evaluated like all other ensemble models although its interpretation will be obviously different. CV is a measure of uncertainty rather a measure of probability of occurrence. If the CV gets a high evaluation score, it means that the uncertainty is high where the species is observed (which might not be a good feature of the model). *The lower is the score, the better are the models.* CV is a nice complement to the mean probability. Old name: `prob.cv`
- `EMci` & `EMci.alpha` : Confidence interval around the mean of probabilities of the selected models

It is also a nice complement to the mean probability. It creates 2 ensemble models :

- *LOWER* : there is less than  $100 * \text{EMci.alpha} / 2$  % of chance to get probabilities lower than the given ones
- *UPPER* : there is less than  $100 * \text{EMci.alpha} / 2$  % of chance to get probabilities upper than the given ones

These intervals are calculated with the following function :

$$I_c = \left[ \bar{x} - \frac{t_\alpha \text{sd}}{\sqrt{n}}; \bar{x} + \frac{t_\alpha \text{sd}}{\sqrt{n}} \right]$$

Old parameter name: `prob.ci` & `prob.ci.alpha`

- `EMca` : Probabilities from the selected models are first transformed into binary data according to the thresholds defined when building the `model.output` object with the `BIOMOD_Modeling` function, maximizing the evaluation metric score over the testing dataset. The committee averaging score is obtained by taking the average of these binary predictions. It is built on the analogy of a simple vote :
  - each single model votes for the species being either present (1) or absent (0)
  - the sum of 1 is then divided by the number of single models *voting*

The interesting feature of this measure is that it gives both a prediction and a measure of uncertainty. When the prediction is close to 0 or 1, it means that all models agree to predict 0 or 1 respectively. When the prediction is around 0.5, it means that half the models predict 1 and the other half 0.

Old parameter name: `committee.averaging`

- `EMwmean` & `EMwmean.decay` : Probabilities from the selected models are weighted according to their evaluation scores obtained when building the `model.output` object with the `BIOMOD_Modeling` function (*better a model is, more importance it has in the ensemble*) and summed.

Old parameter name: `prob.mean.weight` & `prob.mean.weight.decay`

The `EMwmean.decay` is the ratio between a weight and the next or previous one. The formula is :  $W = W(-1) * EMwmean.decay$ . For example, with the value of 1.6 and 4 weights wanted, the relative importance of the weights will be  $1/1.6/2.56(=1.6*1.6)/4.096(=2.56*1.6)$  from the weakest to the strongest, and gives  $0.11/0.17/0.275/0.445$  considering that the sum of the weights is equal to one. The lower the `EMwmean.decay`, the smoother the differences between the weights enhancing a weak discrimination between models.

If `EMwmean.decay = 'proportional'`, the weights are assigned to each model proportionally to their evaluation scores. The discrimination is fairer than using the *decay* method where close scores can have strongly diverging weights, while the proportional method would assign them similar weights.

It is also possible to define the `EMwmean.decay` parameter as a function that will be applied to single models scores and transform them into weights. For example, if `EMwmean.decay = function(x) {x^2}`, the squared of evaluation score of each model will be used to weight the models predictions.

## Value

A `BIOMOD.ensemble.models.out` object containing models outputs, or links to saved outputs.

Models outputs are stored out of R (for memory storage reasons) in 2 different folders created in the current working directory :

1. a *models* folder, named after the `resp.name` argument of `BIOMOD_FormatingData`, and containing all ensemble models
2. a *hidden* folder, named `.BIOMOD_DATA`, and containing outputs related files (original dataset, calibration lines, pseudo-absences selected, predictions, variables importance, evaluation values...), that can be retrieved with `get_ [...]` or `load` functions, and used by other **biomod2** functions, like `BIOMOD_EnsembleForecasting`

**Author(s)**

Wilfried Thuiller, Damien Georges, Robin Engler

**See Also**

[BIOMOD\\_FormatingData](#), [bm\\_ModelingOptions](#), [bm\\_CrossValidation](#), [bm\\_VariablesImportance](#), [BIOMOD\\_Modeling](#), [BIOMOD\\_EnsembleForecasting](#), [bm\\_PlotEvalMean](#), [bm\\_PlotEvalBoxplot](#), [bm\\_PlotVarImpBoxplot](#), [bm\\_PlotResponseCurves](#)

Other Main functions: [BIOMOD\\_EnsembleForecasting\(\)](#), [BIOMOD\\_FormatingData\(\)](#), [BIOMOD\\_LoadModels\(\)](#), [BIOMOD\\_Modeling\(\)](#), [BIOMOD\\_Projection\(\)](#), [BIOMOD\\_RangeSize\(\)](#)

**Examples**

```
library(terra)
# Load species occurrences (6 species available)
data(DataSpecies)
head(DataSpecies)

# Select the name of the studied species
myRespName <- 'GuloGulo'

# Get corresponding presence/absence data
myResp <- as.numeric(DataSpecies[, myRespName])

# Get corresponding XY coordinates
myRespXY <- DataSpecies[, c('X_WGS84', 'Y_WGS84')]

# Load environmental variables extracted from BIOCLIM (bio_3, bio_4, bio_7, bio_11 & bio_12)
data(bioclim_current)
myExpl <- terra::rast(bioclim_current)

## ----- #
file.out <- paste0(myRespName, "/", myRespName, ".AllModels.models.out")
if (file.exists(file.out)) {
  myBiomodModelOut <- get(load(file.out))
} else {

  # Format Data with true absences
  myBiomodData <- BIOMOD_FormatingData(resp.var = myResp,
                                     expl.var = myExpl,
                                     resp.xy = myRespXY,
                                     resp.name = myRespName)

  # Model single models
  myBiomodModelOut <- BIOMOD_Modeling(bm.format = myBiomodData,
                                     modeling.id = 'AllModels',
                                     models = c('RF', 'GLM'),
                                     CV.strategy = 'random',
                                     CV.nb.rep = 2,
```

```

CV.perc = 0.8,
OPT.strategy = 'bigboss',
metric.eval = c('TSS', 'ROC'),
var.import = 3,
seed.val = 42)
}

## ----- #
# Model ensemble models
myBiomodEM <- BIOMOD_EnsembleModeling(bm.mod = myBiomodModelOut,
                                     models.chosen = 'all',
                                     em.by = 'all',
                                     em.algo = c('EMmean', 'EMca'),
                                     metric.select = c('TSS'),
                                     metric.select.thresh = c(0.7),
                                     metric.eval = c('TSS', 'ROC'),
                                     var.import = 3,
                                     seed.val = 42)

myBiomodEM

# Get evaluation scores & variables importance
get_evaluations(myBiomodEM)
get_variables_importance(myBiomodEM)

# Represent evaluation scores
bm_PlotEvalMean(bm.out = myBiomodEM, dataset = 'calibration')
bm_PlotEvalBoxplot(bm.out = myBiomodEM, group.by = c('algo', 'algo'))

# # Represent variables importance
# bm_PlotVarImpBoxplot(bm.out = myBiomodEM, group.by = c('expl.var', 'algo', 'algo'))
# bm_PlotVarImpBoxplot(bm.out = myBiomodEM, group.by = c('expl.var', 'algo', 'merged.by.PA'))
# bm_PlotVarImpBoxplot(bm.out = myBiomodEM, group.by = c('algo', 'expl.var', 'merged.by.PA'))

# # Represent response curves
# bm_PlotResponseCurves(bm.out = myBiomodEM,
#                         models.chosen = get_built_models(myBiomodEM),
#                         fixed.var = 'median')
# bm_PlotResponseCurves(bm.out = myBiomodEM,
#                         models.chosen = get_built_models(myBiomodEM),
#                         fixed.var = 'min')
# bm_PlotResponseCurves(bm.out = myBiomodEM,
#                         models.chosen = get_built_models(myBiomodEM, algo = 'EMmean'),
#                         fixed.var = 'median',
#                         do.bivariate = TRUE)

```



## Description

This function gathers together all input data needed (*xy, presences/absences, explanatory variables, and the same for evaluation data if available*) to run **biomod2** models. It allows to select pseudo-absences if no absence data is available, with different strategies (see Details).

## Usage

```
BIOMOD_FormatingData(
  resp.name,
  resp.var,
  expl.var,
  dir.name = ".",
  resp.xy = NULL,
  eval.resp.var = NULL,
  eval.expl.var = NULL,
  eval.resp.xy = NULL,
  PA.nb.rep = 0,
  PA.nb.absences = 1000,
  PA.strategy = NULL,
  PA.dist.min = 0,
  PA.dist.max = NULL,
  PA.sre.quant = 0.025,
  PA.fact.aggr = NULL,
  PA.user.table = NULL,
  na.rm = TRUE,
  filter.raster = FALSE,
  seed.val = NULL
)
```

## Arguments

resp.name	a character corresponding to the species name
resp.var	a vector, a <a href="#">SpatVector</a> without associated data ( <i>if presence-only</i> ), or a <a href="#">SpatVector</a> object containing binary data (0 : absence, 1 : presence, NA : indeterminate) for a single species that will be used to build the species distribution model(s) <i>Note that old format from <b>sp</b> are still supported such as SpatialPoints (if presence-only) or SpatialPointsDataFrame object containing binary data.</i>
expl.var	a matrix, data.frame, <a href="#">SpatVector</a> or <a href="#">SpatRaster</a> object containing the explanatory variables (in columns or layers) that will be used to build the species distribution model(s) <i>Note that old format from <b>raster</b> and <b>sp</b> are still supported such as RasterStack and SpatialPointsDataFrame objects.</i>
dir.name	(optional, default .) A character corresponding to the modeling folder
resp.xy	(optional, default NULL) If resp.var is a vector, a 2-columns matrix or data.frame containing the corresponding X and Y coordinates that will be used to build the species distribution model(s)

eval.resp.var	<p><i>(optional, default NULL)</i></p> <p>A vector, a <code>SpatVector</code> without associated data (<i>if presence-only</i>), or a <code>SpatVector</code> object containing binary data (0 : absence, 1 : presence, NA : indeterminate) for a single species that will be used to evaluate the species distribution model(s) with independent data</p> <p><i>Note that old format from <b>sp</b> are still supported such as <code>SpatialPoints</code> (if presence-only) or <code>SpatialPointsDataFrame</code> object containing binary data.</i></p>
eval.expl.var	<p><i>(optional, default NULL)</i></p> <p>A matrix, data.frame, <code>SpatVector</code> or <code>SpatRaster</code> object containing the explanatory variables (in columns or layers) that will be used to evaluate the species distribution model(s) with independent data.</p> <p><i>Note that old format from <b>raster</b> and <b>sp</b> are still supported such as <code>RasterStack</code> and <code>SpatialPointsDataFrame</code> objects.</i></p>
eval.resp.xy	<p><i>(optional, default NULL)</i></p> <p>If resp.var is a vector, a 2-columns matrix or data.frame containing the corresponding X and Y coordinates that will be used to evaluate the species distribution model(s) with independent data</p>
PA.nb.rep	<p><i>(optional, default 0)</i></p> <p>If pseudo-absence selection, an integer corresponding to the number of sets (repetitions) of pseudo-absence points that will be drawn</p>
PA.nb.absences	<p><i>(optional, default 0)</i></p> <p>If pseudo-absence selection, and PA.strategy = 'random' or PA.strategy = 'sre' or PA.strategy = 'disk', an integer corresponding to the number of pseudo-absence points that will be selected for each pseudo-absence repetition (true absences included).</p> <p>It can also be a vector of the same length as PA.nb.rep containing integer values corresponding to the different numbers of pseudo-absences to be selected</p>
PA.strategy	<p><i>(optional, default NULL)</i></p> <p>If pseudo-absence selection, a character defining the strategy that will be used to select the pseudo-absence points. Must be random, sre, disk or user.defined (see Details)</p>
PA.dist.min	<p><i>(optional, default 0)</i></p> <p>If pseudo-absence selection and PA.strategy = 'disk', a numeric defining the minimal distance to presence points used to make the disk pseudo-absence selection (in the same projection system units as resp.xy and expl.var, see Details)</p>
PA.dist.max	<p><i>(optional, default 0)</i></p> <p>If pseudo-absence selection and PA.strategy = 'disk', a numeric defining the maximal distance to presence points used to make the disk pseudo-absence selection (in the same projection system units as resp.xy and expl.var, see Details)</p>
PA.sre.quant	<p><i>(optional, default 0)</i></p> <p>If pseudo-absence selection and PA.strategy = 'sre', a numeric between 0 and 0.5 defining the half-quantile used to make the sre pseudo-absence selection (see Details)</p>

PA.fact.aggr	(optional, default NULL) If strategy = 'random' or strategy = 'disk', a integer defining the factor of aggregation to reduce the resolution
PA.user.table	(optional, default NULL) If pseudo-absence selection and PA.strategy = 'user.defined', a matrix or data.frame with as many rows as resp.var values, as many columns as PA.nb.rep, and containing TRUE or FALSE values defining which points will be used to build the species distribution model(s) for each repetition (see Details)
na.rm	(optional, default TRUE) A logical value defining whether points having one or several missing values for explanatory variables should be removed from the analysis or not
filter.raster	(optional, default FALSE) If expl.var is of raster type, a logical value defining whether resp.var is to be filtered when several points occur in the same raster cell
seed.val	(optional, default NULL) An integer value corresponding to the new seed value to be set

### Details

This function gathers and formats all input data needed to run **biomod2** models. It supports different kind of inputs (e.g. matrix, [SpatVector](#), [SpatRaster](#)) and provides different methods to select pseudo-absences if needed.

#### Concerning explanatory variables and XY coordinates :

- if [SpatRaster](#), RasterLayer or RasterStack provided for expl.var or eval.expl.var, **biomod2** will extract the corresponding values from XY coordinates provided :
  - either through resp.xy or eval.resp.xy respectively
  - or resp.var or eval.resp.var, if provided as [SpatVector](#) or SpatialPointsDataFrame

*Be sure to give the objects containing XY coordinates in the same projection system than the raster objects !*

- if data.frame or matrix provided for expl.var or eval.expl.var, **biomod2** will simply merge it (cbind) with resp.var without considering XY coordinates.  
*Be sure to give explanatory and response values in the same row order !*

#### Concerning pseudo-absence selection (see [bm\\_PseudoAbsences](#)) :

- if both presence and absence data are available, and there is enough absences : set PA.nb.rep = 0 and no pseudo-absence will be selected.
- if no absence data is available, several pseudo-absence repetitions are recommended (to estimate the effect of pseudo-absence selection), as well as high number of pseudo-absence points.

*Be sure not to select more pseudo-absence points than maximum number of pixels in the studied area !*

- it is possible now to create several pseudo-absence repetitions with different number of points, BUT with the same sampling strategy.

**Response variable biomod2** models single species at a time (no multi-species). Hence, `resp.var` must be a uni-dimensional object (either a vector, a one-column matrix, `data.frame`, a [SpatVector](#) (without associated data - if presence-only), a `SpatialPoints` (if presence-only), a `SpatialPointsDataFrame` or [SpatVector](#) object), containing values among :

- 1 : presences
- 0 : true absences (if any)
- NA : no information point (might be used to select pseudo-absences if any)

If no true absences are available, pseudo-absence selection must be done.

If `resp.var` is a non-spatial object (vector, matrix or `data.frame`), XY coordinates must be provided through `resp.xy`.

If pseudo-absence points are to be selected, NA points must be provided in order to select pseudo-absences among them.

**Explanatory variables** Factorial variables are allowed, but might lead to some pseudo-absence strategy or models omissions (e.g. `sre`).

**Evaluation data** Although **biomod2** provides tools to automatically divide dataset into calibration and validation parts through the modeling process (see `CV.[. .]` parameters in [BIOMOD\\_Modeling](#) function ; or [bm\\_CrossValidation](#) function), it is also possible (and strongly advised) to directly provide two independent datasets, one for calibration/validation and one for evaluation

**Pseudo-absence selection** (see [bm\\_PseudoAbsences](#)) If no true absences are available, pseudo-absences must be selected from the *background data*, meaning data there is no information whether the species of interest occurs or not. It corresponds either to the remaining pixels of the `expl.var` (if provided as a [SpatRaster](#) or `RasterSattck`) or to the points identified as NA in `resp.var` (if `expl.var` provided as a matrix or `data.frame`).

Several methods are available to do this selection :

**random** all points of initial background are pseudo-absence candidates. `PA.nb.absences` are drawn randomly, for each `PA.nb.rep` requested.

**sre** pseudo-absences have to be selected in conditions (combination of explanatory variables) that differ in a defined proportion (`PA.sre.quant`) from those of presence points. A *Surface Range Envelop* model is first run over the species of interest (see [bm\\_SRE](#)), and pseudo-absences are selected outside this envelop.

*This case is appropriate when all the species climatic niche has been sampled, otherwise it may lead to over-optimistic model evaluations and predictions !*

**disk** pseudo-absences are selected within circles around presence points defined by `PA.dist.min` and `PA.dist.max` distance values (in the same projection system units as `coord` and `expl.var`). It allows to select pseudo-absence points that are not too close to (avoid same niche and pseudo-replication) or too far (localized sampling strategy) from presences.

**user.defined** pseudo-absences are defined in advance and given as `data.frame` through the `PA.user.table` parameter.

**Value**

A `BIOMOD.formated.data` object that can be used to build species distribution model(s) with the `BIOMOD_Modeling` function.  
`print/show`, `plot` and `summary` functions are available to have a summary of the created object.

**Author(s)**

Damien Georges, Wilfried Thuiller

**See Also**

`bm_PseudoAbsences`, `BIOMOD_Modeling`

Other Main functions: `BIOMOD_EnsembleForecasting()`, `BIOMOD_EnsembleModeling()`, `BIOMOD_LoadModels()`, `BIOMOD_Modeling()`, `BIOMOD_Projection()`, `BIOMOD_RangeSize()`

**Examples**

```
library(terra)

# Load species occurrences (6 species available)
data(DataSpecies)
head(DataSpecies)

# Select the name of the studied species
myRespName <- 'GuloGulo'

# Get corresponding presence/absence data
myResp <- as.numeric(DataSpecies[, myRespName])

# Get corresponding XY coordinates
myRespXY <- DataSpecies[, c('X_WGS84', 'Y_WGS84')]

# Load environmental variables extracted from BIOCLIM (bio_3, bio_4, bio_7, bio_11 & bio_12)
data(bioclim_current)
myExpl <- terra::rast(bioclim_current)

# -----#
# Format Data with true absences
myBiomodData <- BIOMOD_FormatingData(resp.var = myResp,
                                     expl.var = myExpl,
                                     resp.xy = myRespXY,
                                     resp.name = myRespName)

myBiomodData
summary(myBiomodData)
plot(myBiomodData)

# -----#
# # Transform true absences into potential pseudo-absences
```

```

# myResp.PA <- ifelse(myResp == 1, 1, NA)
#
# # Format Data with pseudo-absences : random method
# myBiomodData.r <- BIOMOD_FormatingData(resp.var = myResp.PA,
#                                       expl.var = myExpl,
#                                       resp.xy = myRespXY,
#                                       resp.name = myRespName,
#                                       PA.nb.rep = 4,
#                                       PA.nb.absences = 1000,
#                                       PA.strategy = 'random')
#
# # Format Data with pseudo-absences : disk method
# myBiomodData.d <- BIOMOD_FormatingData(resp.var = myResp.PA,
#                                       expl.var = myExpl,
#                                       resp.xy = myRespXY,
#                                       resp.name = myRespName,
#                                       PA.nb.rep = 4,
#                                       PA.nb.absences = 500,
#                                       PA.strategy = 'disk',
#                                       PA.dist.min = 5,
#                                       PA.dist.max = 35)
#
# # Format Data with pseudo-absences : SRE method
# myBiomodData.s <- BIOMOD_FormatingData(resp.var = myResp.PA,
#                                       expl.var = myExpl,
#                                       resp.xy = myRespXY,
#                                       resp.name = myRespName,
#                                       PA.nb.rep = 4,
#                                       PA.nb.absences = 1000,
#                                       PA.strategy = 'sre',
#                                       PA.sre.quant = 0.025)
#
# # Format Data with pseudo-absences : user.defined method
# myPAtable <- data.frame(PA1 = ifelse(myResp == 1, TRUE, FALSE),
#                        PA2 = ifelse(myResp == 1, TRUE, FALSE))
# for (i in 1:ncol(myPAtable)) myPAtable[sample(which(myPAtable[, i] == FALSE), 500), i] = TRUE
# myBiomodData.u <- BIOMOD_FormatingData(resp.var = myResp.PA,
#                                       expl.var = myExpl,
#                                       resp.xy = myRespXY,
#                                       resp.name = myRespName,
#                                       PA.strategy = 'user.defined',
#                                       PA.user.table = myPAtable)
#
# myBiomodData.r
# myBiomodData.d
# myBiomodData.s
# myBiomodData.u
# plot(myBiomodData.r)
# plot(myBiomodData.d)
# plot(myBiomodData.s)
# plot(myBiomodData.u)

```

```

# -----#
# # Select multiple sets of pseudo-absences
#
# # Transform true absences into potential pseudo-absences
# myResp.PA <- ifelse(myResp == 1, 1, NA)
#
# # Format Data with pseudo-absences : random method
# myBiomodData.multi <- BIOMOD_FormatingData(resp.var = myResp.PA,
#                                           expl.var = myExpl,
#                                           resp.xy = myRespXY,
#                                           resp.name = myRespName,
#                                           PA.nb.rep = 4,
#                                           PA.nb.absences = c(1000, 500, 500, 200),
#                                           PA.strategy = 'random')
# myBiomodData.multi
# summary(myBiomodData.multi)
# plot(myBiomodData.multi)

```

---

BIOMOD\_LoadModels

*Load species distribution models built with **biomod2***


---

## Description

This function loads individual models built with [BIOMOD\\_Modeling](#) or [BIOMOD\\_EnsembleModeling](#) functions.

## Usage

```

BIOMOD_LoadModels(
  bm.out,
  full.name = NULL,
  PA = NULL,
  run = NULL,
  algo = NULL,
  merged.by.PA = NULL,
  merged.by.run = NULL,
  merged.by.algo = NULL,
  filtered.by = NULL
)

```

## Arguments

**bm.out** a [BIOMOD.models.out](#) or [BIOMOD.ensemble.models.out](#) object that can be obtained with the [BIOMOD\\_Modeling](#) or [BIOMOD\\_EnsembleModeling](#) functions

full.name	<i>(optional, default NULL)</i> A vector containing model names to be kept, must be either all or a sub-selection of model names that can be obtained with the <a href="#">get_built_models</a> function
PA	<i>(optional, default NULL)</i> A vector containing pseudo-absence set to be loaded, must be among PA1, PA2, ..., allData
run	<i>(optional, default NULL)</i> A vector containing repetition set to be loaded, must be among RUN1, RUN2, ..., allRun
algo	<i>(optional, default NULL)</i> A character containing algorithm to be loaded, must be either ANN, CTA, FDA, GAM, GBM, GLM, MARS, MAXENT, MAXNET, RF, SRE, XGBOOST
merged.by.PA	<i>(optional, default NULL)</i> A vector containing merged pseudo-absence set to be loaded, must be among PA1, PA2, ..., mergedData
merged.by.run	<i>(optional, default NULL)</i> A vector containing merged repetition set to be loaded, must be among RUN1, RUN2, ..., mergedRun
merged.by.algo	<i>(optional, default NULL)</i> A character containing merged algorithm to be loaded, must be among ANN, CTA, FDA, GAM, GBM, GLM, MARS, MAXENT, MAXNET, RF, SRE, XGBOOST, mergedAlgo
filtered.by	<i>(optional, default NULL)</i> A vector containing evaluation metric selected to filter single models to build the ensemble models, must be among POD, FAR, POFD, SR, ACCURACY, BIAS, ROC, TSS, KAPPA, OR, ORSS, CSI, ETS, BOYCE, MPA

### Details

This function might be of particular use to load models and make response plot analyses.

Running the function providing only `bm.out` argument will load all models built by the [BIOMOD\\_Modeling](#) or [BIOMOD\\_EnsembleModeling](#) function, but a subselection of models can be done using the additional arguments (`full.name`, `PA`, `run`, `algo`, `merged.by.PA`, `merged.by.run`, `merged.by.algo`, `filtered.by`).

### Value

A vector containing the names of the loaded models.

### Author(s)

Damien Georges





```
# -----
# Loading some models built
BIOMOD_LoadModels(bm.out = myBiomodModelOut, algo = 'RF')
```

---

BIOMOD\_Modeling

*Run a range of species distribution models*


---

## Description

This function allows to calibrate and evaluate a range of modeling techniques for a given species distribution. The dataset can be split up in calibration/validation parts, and the predictive power of the different models can be estimated using a range of evaluation metrics (see Details).

## Usage

```
BIOMOD_Modeling(
  bm.format,
  modeling.id = as.character(format(Sys.time(), "%s")),
  models = c("ANN", "CTA", "FDA", "GAM", "GBM", "GLM", "MARS", "MAXENT", "MAXNET", "RF",
    "SRE", "XGBOOST"),
  models.pa = NULL,
  CV.strategy = "random",
  CV.nb.rep = 1,
  CV.perc = NULL,
  CV.k = NULL,
  CV.balance = NULL,
  CV.env.var = NULL,
  CV.strat = NULL,
  CV.user.table = NULL,
  CV.do.full.models = TRUE,
  OPT.data.type = "binary",
  OPT.strategy = "default",
  OPT.user.val = NULL,
  OPT.user.base = "bigboss",
  OPT.user = NULL,
  bm.options,
  nb.rep,
  data.split.perc,
  data.split.table,
  do.full.models,
  weights = NULL,
  prevalence = NULL,
  metric.eval = c("KAPPA", "TSS", "ROC"),
```

```

var.import = 0,
scale.models = FALSE,
nb.cpu = 1,
seed.val = NULL,
do.progress = TRUE
)

```

### Arguments

bm.format	a <a href="#">BIOMOD.formated.data</a> or <a href="#">BIOMOD.formated.data.PA</a> object returned by the <a href="#">BIOMOD_FormatingData</a> function
modeling.id	a character corresponding to the name (ID) of the simulation set ( <i>a random number by default</i> )
models	a vector containing model names to be computed, must be among ANN, CTA, FDA, GAM, GBM, GLM, MARS, MAXENT, MAXNET, RF, RFd, SRE, XGBOOST
models.pa	<i>(optional, default NULL)</i> A list containing for each model a vector defining which pseudo-absence datasets are to be used, must be among <code>colnames(bm.format@PA.table)</code>
CV.strategy	a character corresponding to the cross-validation selection strategy, must be among random, kfold, block, strat, env or user.defined
CV.nb.rep	<i>(optional, default 0)</i> If strategy = 'random' or strategy = 'kfold', an integer corresponding to the number of sets (repetitions) of cross-validation points that will be drawn
CV.perc	<i>(optional, default 0)</i> If strategy = 'random', a numeric between 0 and 1 defining the percentage of data that will be kept for calibration
CV.k	<i>(optional, default 0)</i> If strategy = 'kfold' or strategy = 'strat' or strategy = 'env', an integer corresponding to the number of partitions
CV.balance	<i>(optional, default 'presences')</i> If strategy = 'strat' or strategy = 'env', a character corresponding to how data will be balanced between partitions, must be either presences or absences
CV.env.var	<i>(optional)</i> If strategy = 'env', a character corresponding to the environmental variables used to build the partition. k partitions will be built for each environmental variables. By default the function uses all environmental variables available.
CV.strat	<i>(optional, default 'both')</i> If strategy = 'env', a character corresponding to how data will be partitioned along gradient, must be among x, y, both
CV.user.table	<i>(optional, default NULL)</i> If strategy = 'user.defined', a matrix or data.frame defining for each repetition (in columns) which observation lines should be used for models calibration (TRUE) and validation (FALSE)

CV.do.full.models	<i>(optional, default TRUE)</i> A logical value defining whether models should be also calibrated and validated over the whole dataset (and pseudo-absence datasets) or not
OPT.data.type	a character corresponding to the data type to be used, must be either binary, binary.PA, abundance, compositional
OPT.strategy	a character corresponding to the method to select models' parameters values, must be either default, bigboss, user.defined, tuned
OPT.user.val	<i>(optional, default NULL)</i> A list containing parameters values for some (all) models
OPT.user.base	<i>(optional, default bigboss)</i> A character, default or bigboss used when OPT.strategy = 'user.defined'. It sets the bases of parameters to be modified by user defined values.
OPT.user	<i>(optional, default TRUE)</i> A <code>BIOMOD.models.options</code> object returned by the <code>bm_ModelingOptions</code> function
bm.options	<i>deprecated</i> , now called OPT.user
nb.rep	<i>deprecated</i> , now called CV.nb.rep
data.split.perc	<i>deprecated</i> , now called CV.perc
data.split.table	<i>deprecated</i> , now called CV.user.table
do.full.models	<i>deprecated</i> , now called CV.do.full.models
weights	<i>(optional, default NULL)</i> A vector of numeric values corresponding to observation weights (one per observation, see Details)
prevalence	<i>(optional, default NULL)</i> A numeric between 0 and 1 corresponding to the species prevalence to build 'weighted response weights' (see Details)
metric.eval	a vector containing evaluation metric names to be used, must be among POD, FAR, POFD, SR, ACCURACY, BIAS, ROC, TSS, KAPPA, OR, ORSS, CSI, ETS, BOYCE, MPA
var.import	<i>(optional, default NULL)</i> An integer corresponding to the number of permutations to be done for each variable to estimate variable importance
scale.models	<i>(optional, default FALSE)</i> A logical value defining whether all models predictions should be scaled with a binomial GLM or not
nb.cpu	<i>(optional, default 1)</i> An integer value corresponding to the number of computing resources to be used to parallelize the single models computation
seed.val	<i>(optional, default NULL)</i> An integer value corresponding to the new seed value to be set
do.progress	<i>(optional, default TRUE)</i> A logical value defining whether the progress bar is to be rendered or not

## Details

**bm.format** If pseudo absences have been added to the original dataset (see [BIOMOD\\_FormatingData](#)),  $PA.nb.rep * (nb.rep + 1)$  models will be created.

**models** The set of models to be calibrated on the data. 12 modeling techniques are currently available :

- ANN : Artificial Neural Network ([nnet](#))
- CTA : Classification Tree Analysis ([rpart](#))
- FDA : Flexible Discriminant Analysis ([fda](#))
- GAM : Generalized Additive Model ([gam](#), [gam](#) or [bam](#))  
(see [bm\\_ModelingOptions](#) for details on algorithm selection)
- GBM : Generalized Boosting Model, or usually called Boosted Regression Trees ([gbm](#))
- GLM : Generalized Linear Model ([glm](#))
- MARS : Multiple Adaptive Regression Splines ([earth](#))
- MAXENT : Maximum Entropy (see [Maxent website](#))
- MAXNET : Maximum Entropy ([maxnet](#))
- RF : Random Forest ([randomForest](#))
- RFd : Random Forest downsampled ([randomForest](#))
- SRE : Surface Range Envelop or usually called BIOCLIM ([bm\\_SRE](#))
- XGBOOST : eXtreme Gradient Boosting Training ([xgboost](#))

**models.pa** Different models might respond differently to different numbers of pseudo-absences. It is possible to create sets of pseudo-absences with different numbers of points (see [BIOMOD\\_FormatingData](#)) and to assign only some of these datasets to each single model.

**CV**[... parameters] Different methods are available to calibrate/validate the single models (see [bm\\_CrossValidation](#)).

**OPT**[... parameters] Different methods are available to parameterize the single models (see [bm\\_ModelingOptions](#) and [BIOMOD.options.dataset](#)). Note that only binary data type is allowed currently.

- `default` : only default parameter values of default parameters of the single models functions are retrieved. Nothing is changed so it might not give good results.
- `bigboss` : uses parameters pre-defined by **biomod2** team and that are available in the dataset [OptionsBigboss](#).  
*to be optimized in near future*
- `user.defined` : updates default or bigboss parameters with some parameters values defined by the user (but matching the format of a [BIOMOD.models.options](#) object)
- `tuned` : calling the [bm\\_Tuning](#) function to try and optimize some default values

**weights & prevalence** More or less weight can be given to some specific observations.

- If `weights = prevalence = NULL`, each observation (presence or absence) will have the same weight, no matter the total number of presences and absences.
- If `prevalence = 0.5`, presences and absences will be weighted equally (*i.e. the weighted sum of presences equals the weighted sum of absences*).
- If `prevalence` is set below (*above*)  $0.5$ , more weight will be given to absences (*presences*).
- If `weights` is defined, `prevalence` argument will be ignored, and each observation will have its own weight.

- If pseudo-absences have been generated ( $PA.nb.rep > 0$  in `BIOMOD_FormatingData`), weights are by default calculated such that prevalence = 0.5. *Automatically created weights will be integer values to prevent some modeling issues.*
- *NOTE THAT* MAXENT, MAXNET, RF, RFd and SRE models do not take weights into account.

**metric.eval simple** • POD : Probability of detection (hit rate)

- FAR : False alarm ratio
- POFD : Probability of false detection (fall-out)
- SR : Success ratio
- ACCURACY : Accuracy (fraction correct)
- BIAS : Bias score (frequency bias)

**complex** • ROC : Relative operating characteristic

- TSS : True skill statistic (Hanssen and Kuipers discriminant, Peirce's skill score)
- KAPPA : Cohen's Kappa (Heidke skill score)
- OR : Odds Ratio
- ORSS : Odds ratio skill score (Yule's Q)
- CSI : Critical success index (threat score)
- ETS : Equitable threat score (Gilbert skill score)

**presence-only** • BOYCE : Boyce index

- MPA : Minimal predicted area (cutoff optimizing MPA to predict 90% of presences)

Optimal value of each method can be obtained with the `get_optim_value` function. Several evaluation metrics can be selected. *Please refer to the [CAWRC website \(section "Methods for dichotomous forecasts"\)](#) to get detailed description of each metric.* Results after modeling can be obtained through the `get_evaluations` function.

Evaluation metric are calculated on the calibrating data (column calibration), on the cross-validation data (column validation) or on the evaluation data (column evaluation).

*For cross-validation data, see `CV.[...]` parameters in `BIOMOD_Modeling` function ; for evaluation data, see `eval.[...]` parameters in `BIOMOD_FormatingData`.*

**var.import** A value characterizing how much each variable has an impact on each model predictions can be calculated by randomizing the variable of interest and computing the correlation between original and shuffled variables (see `bm_VariablesImportance`).

**scale.models** **This parameter is quite experimental and it is recommended not to use it. It may lead to reduction in projection scale amplitude.** Some categorical models always have to be scaled (FDA, ANN), but it may be interesting to scale all computed models to ensure comparable predictions (0-1000 range). It might be particularly useful when doing ensemble forecasting to remove the scale prediction effect (*the more extended projections are, the more they influence ensemble forecasting results*).

## Value

A `BIOMOD.models.out` object containing models outputs, or links to saved outputs.

Models outputs are stored out of R (for memory storage reasons) in 2 different folders created in the current working directory :

1. a `models` folder, named after the `resp.name` argument of `BIOMOD_FormatingData`, and containing all calibrated models for each repetition and pseudo-absence run

2. a *hidden* folder, named `.BIOMOD_DATA`, and containing outputs related files (original dataset, calibration lines, pseudo-absences selected, predictions, variables importance, evaluation values...), that can be retrieved with `get_[...]` or `load` functions, and used by other **biomod2** functions, like [BIOMOD\\_Projection](#) or [BIOMOD\\_EnsembleModeling](#)

### Author(s)

Wilfried Thuiller, Damien Georges, Robin Engler

### See Also

[glm](#), [gam](#), [bam](#), [gbm](#), [rpart](#), [nnet](#), [fda](#), [earth](#), [randomForest](#), [maxnet](#), [xgboost](#), [BIOMOD\\_FormatingData](#), [bm\\_ModelingOptions](#), [bm\\_Tuning](#), [bm\\_CrossValidation](#), [bm\\_VariablesImportance](#), [BIOMOD\\_Projection](#), [BIOMOD\\_EnsembleModeling](#), [bm\\_PlotEvalMean](#), [bm\\_PlotEvalBoxplot](#), [bm\\_PlotVarImpBoxplot](#), [bm\\_PlotResponseCurves](#)

Other Main functions: [BIOMOD\\_EnsembleForecasting\(\)](#), [BIOMOD\\_EnsembleModeling\(\)](#), [BIOMOD\\_FormatingData\(\)](#), [BIOMOD\\_LoadModels\(\)](#), [BIOMOD\\_Projection\(\)](#), [BIOMOD\\_RangeSize\(\)](#)

### Examples

```
library(terra)

# Load species occurrences (6 species available)
data(DataSpecies)
head(DataSpecies)

# Select the name of the studied species
myRespName <- 'GuloGulo'

# Get corresponding presence/absence data
myResp <- as.numeric(DataSpecies[, myRespName])

# Get corresponding XY coordinates
myRespXY <- DataSpecies[, c('X_WGS84', 'Y_WGS84')]

# Load environmental variables extracted from BIOCLIM (bio_3, bio_4, bio_7, bio_11 & bio_12)
data(bioclim_current)
myExpl <- terra::rast(bioclim_current)

# ----- #
# Format Data with true absences
myBiomodData <- BIOMOD_FormatingData(resp.var = myResp,
                                     expl.var = myExpl,
                                     resp.xy = myRespXY,
                                     resp.name = myRespName)

# ----- #
# Model single models
```

```

myBiomodModelOut <- BIOMOD_Modeling(bm.format = myBiomodData,
                                   modeling.id = 'AllModels',
                                   models = c('RF', 'GLM'),
                                   CV.strategy = 'random',
                                   CV.nb.rep = 2,
                                   CV.perc = 0.8,
                                   OPT.strategy = 'bigboss',
                                   metric.eval = c('TSS','ROC'),
                                   var.import = 2,
                                   seed.val = 42)

myBiomodModelOut

# Get evaluation scores & variables importance
get_evaluations(myBiomodModelOut)
get_variables_importance(myBiomodModelOut)

# Represent evaluation scores
bm_PlotEvalMean(bm.out = myBiomodModelOut, dataset = 'calibration')
bm_PlotEvalMean(bm.out = myBiomodModelOut, dataset = 'validation')
bm_PlotEvalBoxplot(bm.out = myBiomodModelOut, group.by = c('algo', 'run'))

# # Represent variables importance
# bm_PlotVarImpBoxplot(bm.out = myBiomodModelOut, group.by = c('expl.var', 'algo', 'algo'))
# bm_PlotVarImpBoxplot(bm.out = myBiomodModelOut, group.by = c('expl.var', 'algo', 'run'))
# bm_PlotVarImpBoxplot(bm.out = myBiomodModelOut, group.by = c('algo', 'expl.var', 'run'))

# # Represent response curves
# mods <- get_built_models(myBiomodModelOut, run = 'RUN1')
# bm_PlotResponseCurves(bm.out = myBiomodModelOut,
#                        # models.chosen = mods,
#                        # fixed.var = 'median')
# bm_PlotResponseCurves(bm.out = myBiomodModelOut,
#                        # models.chosen = mods,
#                        # fixed.var = 'min')
# mods <- get_built_models(myBiomodModelOut, full.name = 'GuloGulo_allData_RUN2_RF')
# bm_PlotResponseCurves(bm.out = myBiomodModelOut,
#                        # models.chosen = mods,
#                        # fixed.var = 'median',
#                        # do.bivariate = TRUE)

```

---

BIOMOD\_Projection

*Project a range of calibrated species distribution models onto new environment*


---

## Description

This function allows to project a range of models built with the [BIOMOD\\_Modeling](#) function onto new environmental data (*which can represent new areas, resolution or time scales for example*).



**Usage**

```

BIOMOD_Projection(
  bm.mod,
  proj.name,
  new.env,
  new.env.xy = NULL,
  models.chosen = "all",
  metric.binary = NULL,
  metric.filter = NULL,
  compress = TRUE,
  build.clamping.mask = TRUE,
  nb.cpu = 1,
  seed.val = NULL,
  ...
)

```

**Arguments**

bm.mod	a <a href="#">BIOMOD.models.out</a> object returned by the <a href="#">BIOMOD_Modeling</a> function
proj.name	a character corresponding to the name (ID) of the projection set ( <i>a new folder will be created within the simulation folder with this name</i> )
new.env	A matrix, data.frame or <a href="#">SpatRaster</a> object containing the new explanatory variables (in columns or layers, with names matching the variables names given to the <a href="#">BIOMOD_FormatingData</a> function to build bm.mod) that will be used to project the species distribution model(s) <i>Note that old format from <b>raster</b> are still supported such as RasterStack objects.</i>
new.env.xy	(optional, default NULL) If new.env is a matrix or a data.frame, a 2-columns matrix or data.frame containing the corresponding X and Y coordinates that will be used to project the species distribution model(s)
models.chosen	a vector containing model names to be kept, must be either all or a sub-selection of model names that can be obtained with the <a href="#">get_built_models</a> function
metric.binary	(optional, default NULL) A vector containing evaluation metric names to be used to transform prediction values into binary values based on models evaluation scores obtained with the <a href="#">BIOMOD_Modeling</a> function. Must be among all (same evaluation metrics than those of bm.mod) or POD, FAR, POFD, SR, ACCURACY, BIAS, ROC, TSS, KAPPA, OR, ORSS, CSI, ETS, BOYCE, MPA
metric.filter	(optional, default NULL) A vector containing evaluation metric names to be used to transform prediction values into filtered values based on models evaluation scores obtained with the <a href="#">BIOMOD_Modeling</a> function. Must be among all (same evaluation metrics than those of bm.mod) or POD, FAR, POFD, SR, ACCURACY, BIAS, ROC, TSS, KAPPA, OR, ORSS, CSI, ETS, BOYCE, MPA

<code>compress</code>	<i>(optional, default TRUE)</i> A logical or a character value defining whether and how objects should be compressed when saved on hard drive. Must be either TRUE, FALSE, xz or gzip (see Details)
<code>build.clamping.mask</code>	<i>(optional, default TRUE)</i> A logical value defining whether a clamping mask should be built and saved on hard drive or not (see Details)
<code>nb.cpu</code>	<i>(optional, default 1)</i> An integer value corresponding to the number of computing resources to be used to parallelize the single models computation
<code>seed.val</code>	<i>(optional, default NULL)</i> An integer value corresponding to the new seed value to be set
<code>...</code>	<i>(optional, see Details)</i>

### Details

If `models.chosen = 'all'`, projections are done for all calibration and pseudo absences runs if applicable.

These projections may be used later by the [BIOMOD\\_EnsembleForecasting](#) function.

If `build.clamping.mask = TRUE`, a raster file will be saved within the projection folder. This mask values will correspond to the number of variables in each pixel that are out of their calibration / validation range, identifying locations where predictions are uncertain.

`...` can take the following values :

- `omit.na` : a logical value defining whether all not fully referenced environmental points will get NA as predictions or not
- `on_0_1000` : a logical value defining whether 0 - 1 probabilities are to be converted to 0 - 1000 scale to save memory on backup
- `do.stack` : a logical value defining whether all projections are to be saved as one [SpatRaster](#) object or several [SpatRaster](#) files (*the default if projections are too heavy to be all loaded at once in memory*)
- `keep.in.memory` : a logical value defining whether all projections are to be kept loaded at once in memory, or only links pointing to hard drive are to be returned
- `output.format` : a character value corresponding to the projections saving format on hard drive, must be either `.grd`, `.img`, `.tif` or `.RData` (the default if `new.env` is given as `matrix` or `data.frame`)

### Value

A [BIOMOD.projection.out](#) object containing models projections, or links to saved outputs.

Models projections are stored out of R (for memory storage reasons) in `proj.name` folder created in the current working directory :



```

# Model single models
myBiomodModelOut <- BIOMOD_Modeling(bm.format = myBiomodData,
                                   modeling.id = 'AllModels',
                                   models = c('RF', 'GLM'),
                                   CV.strategy = 'random',
                                   CV.nb.rep = 2,
                                   CV.perc = 0.8,
                                   OPT.strategy = 'bigboss',
                                   metric.eval = c('TSS', 'ROC'),
                                   var.import = 3,
                                   seed.val = 42)
}

# -----#
# Project single models
file.proj <- paste0(myRespName, "/proj_Current/", myRespName, ".Current.projection.out")
if (file.exists(file.proj)) {
  myBiomodProj <- get(load(file.proj))
} else {
  myBiomodProj <- BIOMOD_Projection(bm.mod = myBiomodModelOut,
                                   proj.name = 'Current',
                                   new.env = myExpl,
                                   models.chosen = 'all')
}
myBiomodProj
plot(myBiomodProj)

```

---

BIOMOD\_RangeSize

*Analyze the range size differences between projections of species distribution models*


---

### Description

This function allows to calculate the absolute number of locations (pixels) lost, stable and gained, as well as the corresponding relative proportions, between two (or more) binary projections of (ensemble) species distribution models (*which can represent new time scales or environmental scenarios for example*).

### Usage

```
BIOMOD_RangeSize(proj.current, proj.future)
```

```
## S4 method for signature 'data.frame,data.frame'
BIOMOD_RangeSize(proj.current, proj.future)
```

```
## S4 method for signature 'SpatRaster,SpatRaster'
BIOMOD_RangeSize(proj.current, proj.future)
```

## Arguments

proj.current	a data.frame, <a href="#">RasterLayer</a> or <a href="#">SpatRaster</a> object containing the initial binary projection(s) of the (ensemble) species distribution model(s)
proj.future	a data.frame, <a href="#">RasterLayer</a> or <a href="#">SpatRaster</a> object containing the final binary projection(s) of the (ensemble) species distribution model(s)

## Details

Note that **this function is only relevant to compare binary projections, made on the same area with the same resolution.**

Comparison between proj.current and proj.future depends on the number of projection in both objects:

proj.current	proj.future
<b>1 projection</b> (e.g. data.frame with 1 column, SpatRaster with 1 layer)	<b>1 projection</b> (e.g. data.frame with 1 column, SpatRaster with 1 layer)
<b>n projections</b> (e.g. data.frame with n column, SpatRaster with n layer)	<b>n projections</b> (e.g. data.frame with n column, SpatRaster with n layer)
<b>1 projection</b> (e.g. data.frame with 1 column, SpatRaster with 1 layer)	<b>n projections</b> (e.g. data.frame with n column, SpatRaster with n layer)

Diff.By.Pixel object is obtained by applying the simple following formula :

$$proj.future - 2 * proj.current$$

## Value

A list containing two objects :

**Compt.By.Species** a data.frame containing the summary of range change for each comparison

- Loss : number of pixels predicted to be lost
- Stable0 : number of pixels not currently occupied and not predicted to be
- Stable1 : number of pixels currently occupied and predicted to remain occupied
- Gain : number of pixels predicted to be gained
- PercLoss : percentage of pixels currently occupied and predicted to be lost (Loss / (Loss + Stable1))
- PercGain : percentage of pixels predicted to be gained compare to the number of pixels currently occupied (Gain / (Loss + Stable1))
- SpeciesRangeChange : percentage of pixels predicted to change (loss or gain) compare to the number of pixels currently occupied (PercGain - PercLoss)
- CurrentRangeSize : number of pixels currently occupied
- FutureRangeSize0Disp : number of pixels predicted to be occupied, assuming no migration
- FutureRangeSize1Disp : number of pixels predicted to be occupied, assuming migration

**Diff.By.Pixel** an object in the same form than the input data (proj.current and proj.future) and containing a value for each point/pixel of each comparison among :

- -2 : predicted to be lost
- -1 : predicted to remain occupied
- 0 : predicted to remain unoccupied
- 1 : predicted to be gained

### Author(s)

Wilfried Thuiller, Damien Georges, Bruno Lafourcade

### See Also

[BIOMOD\\_Projection](#), [BIOMOD\\_EnsembleForecasting](#), [bm\\_PlotRangeSize](#)

Other Main functions: [BIOMOD\\_EnsembleForecasting\(\)](#), [BIOMOD\\_EnsembleModeling\(\)](#), [BIOMOD\\_FormatingData\(\)](#), [BIOMOD\\_LoadModels\(\)](#), [BIOMOD\\_Modeling\(\)](#), [BIOMOD\\_Projection\(\)](#)

### Examples

```
library(terra)

# Load species occurrences (6 species available)
data(DataSpecies)
head(DataSpecies)

# Select the name of the studied species
myRespName <- 'GuloGulo'

# Get corresponding presence/absence data
myResp <- as.numeric(DataSpecies[, myRespName])

# Get corresponding XY coordinates
myRespXY <- DataSpecies[, c('X_WGS84', 'Y_WGS84')]

# Load environmental variables extracted from BIOCLIM (bio_3, bio_4, bio_7, bio_11 & bio_12)
data(bioclim_current)
myExpl <- terra::rast(bioclim_current)

# ----- #
file.out <- paste0(myRespName, "/", myRespName, ".AllModels.models.out")
if (file.exists(file.out)) {
  myBiomodModelOut <- get(load(file.out))
} else {

  # Format Data with true absences
  myBiomodData <- BIOMOD_FormatingData(resp.var = myResp,
                                     expl.var = myExpl,
                                     resp.xy = myRespXY,
                                     resp.name = myRespName)

  # Model single models
```

```

myBiomodModelOut <- BIOMOD_Modeling(bm.format = myBiomodData,
                                   modeling.id = 'AllModels',
                                   models = c('RF', 'GLM'),
                                   CV.strategy = 'random',
                                   CV.nb.rep = 2,
                                   CV.perc = 0.8,
                                   OPT.strategy = 'bigboss',
                                   metric.eval = c('TSS', 'ROC'),
                                   var.import = 3,
                                   seed.val = 42)
}

models.proj <- get_built_models(myBiomodModelOut, algo = "RF")
# Project single models
myBiomodProj <- BIOMOD_Projection(bm.mod = myBiomodModelOut,
                                  proj.name = 'CurrentRangeSize',
                                  new.env = myExpl,
                                  models.chosen = models.proj,
                                  metric.binary = 'all',
                                  build.clamping.mask = TRUE)

# ----- #
# Load environmental variables extracted from BIOCLIM (bio_3, bio_4, bio_7, bio_11 & bio_12)
data(bioclim_future)
myExplFuture <- terra::rast(bioclim_future)

# Project onto future conditions
myBiomodProjectionFuture <- BIOMOD_Projection(bm.mod = myBiomodModelOut,
                                              proj.name = 'FutureRangeSize',
                                              new.env = myExplFuture,
                                              models.chosen = models.proj,
                                              metric.binary = 'TSS')

# Load current and future binary projections
CurrentProj <- get_predictions(myBiomodProj,
                              metric.binary = "TSS",
                              model.as.col = TRUE)
FutureProj <- get_predictions(myBiomodProjectionFuture,
                              metric.binary = "TSS",
                              model.as.col = TRUE)

# Compute differences
myBiomodRangeSize <- BIOMOD_RangeSize(proj.current = CurrentProj, proj.future = FutureProj)

myBiomodRangeSize$Compt.By.Models
plot(myBiomodRangeSize$Diff.By.Pixel)

# Represent main results
bm_PlotRangeSize(bm.range = myBiomodRangeSize)

```

---

```
bm_BinaryTransformation
```

*Convert probability values into binary values using a predefined threshold*

---

### Description

This internal **biomod2** function allows to convert probability (not necessary between 0 and 1) values into binary presence-absence (0 or 1) values according to a predefined threshold (see Details).

### Usage

```
bm_BinaryTransformation(data, threshold, do.filtering = FALSE)
```

```
## S4 method for signature 'data.frame'
```

```
bm_BinaryTransformation(data, threshold, do.filtering = FALSE)
```

```
## S4 method for signature 'matrix'
```

```
bm_BinaryTransformation(data, threshold, do.filtering = FALSE)
```

```
## S4 method for signature 'numeric'
```

```
bm_BinaryTransformation(data, threshold, do.filtering = FALSE)
```

```
## S4 method for signature 'SpatRaster'
```

```
bm_BinaryTransformation(data, threshold, do.filtering = FALSE)
```

### Arguments

data	a vector, a matrix, data.frame, or a <a href="#">SpatRaster</a> containing the data to be converted
threshold	a numeric or a vector of numeric corresponding to the threshold used to convert the given data
do.filtering	(optional, default FALSE) A logical value defining whether filtered data should be returned, or binary one (see Details)

### Details

If data is a vector, threshold should be a single numeric value.

If data is a matrix, data.frame or [SpatRaster](#), threshold should be a vector containing as many values as the number of columns or layers contained in data. If only one numeric value is given, the same threshold will be applied to all columns or layers.

If do.filtering = FALSE, binary (0 or 1) values are returned.

If do.filtering = TRUE, values will be *filtered* according to threshold, meaning that :



- `data < threshold` will return 0
- `data >= threshold` will return the actual values of data (not transformed in 1)

### Value

An object of the same class than data and containing either binary (0 or 1) values, or filtered values.

### Author(s)

Wilfried Thuiller, Damien Georges

### See Also

[BIOMOD\\_Projection](#), [BIOMOD\\_EnsembleForecasting](#)

Other Secondary functions: [bm\\_CrossValidation\(\)](#), [bm\\_FindOptimStat\(\)](#), [bm\\_MakeFormula\(\)](#), [bm\\_ModelingOptions\(\)](#), [bm\\_PlotEvalBoxplot\(\)](#), [bm\\_PlotEvalMean\(\)](#), [bm\\_PlotRangeSize\(\)](#), [bm\\_PlotResponseCurves\(\)](#), [bm\\_PlotVarImpBoxplot\(\)](#), [bm\\_PseudoAbsences\(\)](#), [bm\\_RunModelsLoop\(\)](#), [bm\\_SRE\(\)](#), [bm\\_SampleBinaryVector\(\)](#), [bm\\_SampleFactorLevels\(\)](#), [bm\\_Tuning\(\)](#), [bm\\_VariablesImportance\(\)](#)

### Examples

```
## Generate a 0-1000 vector (normal distribution)
vec.d <- rnorm(100, 500, 100)

## From continuous to binary / filtered vector
vec.d_bin <- bm_BinaryTransformation(data = vec.d, threshold = 500)
vec.d_filt <- bm_BinaryTransformation(data = vec.d, threshold = 500, do.filtering = TRUE)
cbind(vec.d, vec.d_bin, vec.d_filt)
```

---

<code>bm_CrossValidation</code>	<i>Build cross-validation table</i>
---------------------------------	-------------------------------------

---

### Description

This internal **biomod2** function allows to build a cross-validation table according to 6 different methods : `random`, `kfold`, `block`, `strat`, `env` or `user.defined` (see [Details](#)).

### Usage

```
bm_CrossValidation(  
  bm.format,  
  strategy = "random",  
  nb.rep = 0,  
  perc = 0.8,  
  k = 0,  
  balance = "presences",
```

```
env.var = NULL,  
strat = "both",  
user.table = NULL,  
do.full.models = FALSE  
)  
  
bm_CrossValidation_user.defined(bm.format, ...)  
  
## S4 method for signature 'BIOMOD.formated.data'  
bm_CrossValidation_user.defined(bm.format, user.table)  
  
## S4 method for signature 'BIOMOD.formated.data.PA'  
bm_CrossValidation_user.defined(bm.format, user.table)  
  
bm_CrossValidation_random(bm.format, ...)  
  
## S4 method for signature 'BIOMOD.formated.data'  
bm_CrossValidation_random(bm.format, nb.rep, perc)  
  
## S4 method for signature 'BIOMOD.formated.data.PA'  
bm_CrossValidation_random(bm.format, nb.rep, perc)  
  
bm_CrossValidation_kfold(bm.format, ...)  
  
## S4 method for signature 'BIOMOD.formated.data'  
bm_CrossValidation_kfold(bm.format, nb.rep, k)  
  
## S4 method for signature 'BIOMOD.formated.data.PA'  
bm_CrossValidation_kfold(bm.format, nb.rep, k)  
  
bm_CrossValidation_block(bm.format, ...)  
  
## S4 method for signature 'BIOMOD.formated.data'  
bm_CrossValidation_block(bm.format)  
  
## S4 method for signature 'BIOMOD.formated.data.PA'  
bm_CrossValidation_block(bm.format)  
  
bm_CrossValidation_strat(bm.format, ...)  
  
## S4 method for signature 'BIOMOD.formated.data'  
bm_CrossValidation_strat(bm.format, balance, strat, k)  
  
## S4 method for signature 'BIOMOD.formated.data.PA'  
bm_CrossValidation_strat(bm.format, balance, strat, k)  
  
bm_CrossValidation_env(bm.format, ...)
```

```
## S4 method for signature 'BIOMOD.formated.data'
bm_CrossValidation_env(bm.format, balance, k, env.var)
```

```
## S4 method for signature 'BIOMOD.formated.data.PA'
bm_CrossValidation_env(bm.format, balance, k, env.var)
```

### Arguments

bm.format	a <code>BIOMOD.formated.data</code> or <code>BIOMOD.formated.data.PA</code> object returned by the <code>BIOMOD_FormatingData</code> function
strategy	a character corresponding to the cross-validation selection strategy, must be among random, kfold, block, strat, env or user.defined
nb.rep	<i>(optional, default 0)</i> If strategy = 'random' or strategy = 'kfold', an integer corresponding to the number of sets (repetitions) of cross-validation points that will be drawn
perc	<i>(optional, default 0)</i> If strategy = 'random', a numeric between 0 and 1 defining the percentage of data that will be kept for calibration
k	<i>(optional, default 0)</i> If strategy = 'kfold' or strategy = 'strat' or strategy = 'env', an integer corresponding to the number of partitions
balance	<i>(optional, default 'presences')</i> If strategy = 'strat' or strategy = 'env', a character corresponding to how data will be balanced between partitions, must be either presences or absence
env.var	<i>(optional)</i> If strategy = 'env', a character corresponding to the environmental variables used to build the partition. k partitions will be built for each environmental variables. <i>By default the function uses all environmental variables available.</i>
strat	<i>(optional, default 'both')</i> If strategy = 'env', a character corresponding to how data will be partitioned along gradient, must be among x, y, both
user.table	<i>(optional, default NULL)</i> If strategy = 'user.defined', a matrix or data.frame defining for each repetition (in columns) which observation lines should be used for models calibration (TRUE) and validation (FALSE)
do.full.models	<i>(optional, default TRUE)</i> A logical value defining whether models should be also calibrated and validated over the whole dataset (and pseudo-absence datasets) or not
...	<i>(optional, one or several of the following arguments depending on the selected method)</i>

### Details

Several parameters are available within the function and some of them can be used with different cross-validation strategies :

```
| ..... | random | kfold | block | strat | env |
-----
| nb.rep. | x..... | x.... | ..... | ..... | ... |
| perc... | x..... | ..... | ..... | ..... | ... |
| k..... | ..... | x.... | ..... | x.... | x.. |
| balance | ..... | ..... | ..... | x.... | x.. |
| strat.. | ..... | ..... | ..... | x.... | ... |
```

### Concerning column names of matrix output :

The number of columns depends on the strategy selected. The column names are given *a posteriori* of the selection, ranging from 1 to the number of columns. If `do.full.models = TRUE`, columns merging runs (and/or pseudo-absence datasets) are added at the end.

### Concerning cross-validation strategies :

**random** Most simple method to calibrate and validate a model is to split the original dataset in two datasets : one to calibrate the model and the other one to validate it. The splitting can be repeated `nb.rep` times.

**k-fold** The k-fold method splits the original dataset in k datasets of equal sizes : each part is used successively as the validation dataset while the other k-1 parts are used for the calibration, leading to k calibration/validation ensembles. This multiple splitting can be repeated `nb.rep` times.

**block** It may be used to test for model overfitting and to assess transferability in geographic space. block stratification was described in *Muscarella et al. 2014* (see References). Four bins of equal size are partitioned (bottom-left, bottom-right, top-left and top-right).

**stratified** It may be used to test for model overfitting and to assess transferability in geographic space. x and y stratification was described in *Wenger and Olden 2012* (see References). y stratification uses k partitions along the y-gradient, x stratification does the same for the x-gradient. both returns 2k partitions: k partitions stratified along the x-gradient and k partitions stratified along the y-gradient.

**environmental** It may be used to test for model overfitting and to assess transferability in environmental space. It returns k partitions for each variable given in `env.var`.

**user-defined** Allow the user to give its own crossvalidation table. For a presence-absence dataset, column names must be formatted as: `_allData_RUNx` with x an integer. For a presence-only dataset for which several pseudo-absence dataset were generated, column names must be formatted as: `_PAX_RUNy` with x an integer and PAX an existing pseudo-absence dataset and y an integer

### Concerning balance parameter :

If `balance = 'presences'`, presences are divided (balanced) equally over the partitions (e.g. *Fig. 1b in Muscarelly et al. 2014*). Absences or pseudo-absences will however be unbalanced over the partitions especially if the presences are clumped on an edge of the study area.

If `balance = 'absences'`, absences (resp. pseudo-absences or background) are divided (balanced) as equally as possible between the partitions (geographical balanced bins given that absences are spread over the study area equally, approach similar to *Fig. 1 in Wenger et Olden 2012*). Presences will however be unbalanced over the partitions especially if the presences are clumped on an edge of the study area.

### Value

A matrix or data.frame defining for each repetition (in columns) which observation lines should be used for models calibration (TRUE) and validation (FALSE).

### Author(s)

Frank Breiner, Maya Gueguen

### References

- Muscarella, R., Galante, P.J., Soley-Guardia, M., Boria, R.A., Kass, J.M., Uriarte, M. & Anderson, R.P. (2014). ENMeval: An R package for conducting spatially independent evaluations and estimating optimal model complexity for Maxent ecological niche models. *Methods in Ecology and Evolution*, **5**, 1198-1205.
- Wenger, S.J. & Olden, J.D. (2012). Assessing transferability of ecological models: an underappreciated aspect of statistical validation. *Methods in Ecology and Evolution*, **3**, 260-267.

### See Also

[get.block](#), [kfold](#), [BIOMOD\\_FormatingData](#), [BIOMOD\\_Modeling](#)

Other Secondary functions: [bm\\_BinaryTransformation\(\)](#), [bm\\_FindOptimStat\(\)](#), [bm\\_MakeFormula\(\)](#), [bm\\_ModelingOptions\(\)](#), [bm\\_PlotEvalBoxplot\(\)](#), [bm\\_PlotEvalMean\(\)](#), [bm\\_PlotRangeSize\(\)](#), [bm\\_PlotResponseCurves\(\)](#), [bm\\_PlotVarImpBoxplot\(\)](#), [bm\\_PseudoAbsences\(\)](#), [bm\\_RunModelsLoop\(\)](#), [bm\\_SRE\(\)](#), [bm\\_SampleBinaryVector\(\)](#), [bm\\_SampleFactorLevels\(\)](#), [bm\\_Tuning\(\)](#), [bm\\_VariablesImportance\(\)](#)

### Examples

```
library(terra)
# Load species occurrences (6 species available)
data(DataSpecies)
head(DataSpecies)

# Select the name of the studied species
myRespName <- 'GuloGulo'

# Get corresponding presence/absence data
myResp <- as.numeric(DataSpecies[, myRespName])

# Get corresponding XY coordinates
myRespXY <- DataSpecies[, c('X_WGS84', 'Y_WGS84')]

# Load environmental variables extracted from BIOCLIM (bio_3, bio_4, bio_7, bio_11 & bio_12)
data(bioclim_current)
```

```
myExpl <- terra::rast(bioclim_current)

# ----- #
# Format Data with true absences
myBiomodData <- BIOMOD_FormatingData(resp.var = myResp,
                                   expl.var = myExpl,
                                   resp.xy = myRespXY,
                                   resp.name = myRespName)

# ----- #
# Create the different validation datasets

# random selection
cv.r <- bm_CrossValidation(bm.format = myBiomodData,
                          strategy = "random",
                          nb.rep = 3,
                          k = 0.8)

# k-fold selection
cv.k <- bm_CrossValidation(bm.format = myBiomodData,
                          strategy = "kfold",
                          nb.rep = 2,
                          k = 3)

# block selection
cv.b <- bm_CrossValidation(bm.format = myBiomodData,
                          strategy = "block")

# stratified selection (geographic)
cv.s <- bm_CrossValidation(bm.format = myBiomodData,
                          strategy = "strat",
                          k = 2,
                          balance = "presences",
                          strat = "x")

# stratified selection (environmental)
cv.e <- bm_CrossValidation(bm.format = myBiomodData,
                          strategy = "env",
                          k = 2,
                          balance = "presences")

head(cv.r)
apply(cv.r, 2, table)
head(cv.k)
apply(cv.k, 2, table)
head(cv.b)
apply(cv.b, 2, table)
head(cv.s)
apply(cv.s, 2, table)
head(cv.e)
apply(cv.e, 2, table)
```

---

bm_FindOptimStat	<i>Calculate the best score according to a given evaluation method</i>
------------------	--

---

### Description

This internal **biomod2** function allows the user to find the threshold to convert continuous values into binary ones leading to the best score for a given evaluation metric.

### Usage

```
bm_FindOptimStat(
  metric.eval = "TSS",
  obs,
  fit,
  nb.thresh = 100,
  threshold = NULL,
  boyce.bg.env = NULL,
  mpa.perc = 0.9
)

get_optim_value(metric.eval)

bm_CalculateStat(misc, metric.eval = "TSS")
```

### Arguments

metric.eval	a character corresponding to the evaluation metric to be used, must be either POD, FAR, POFD, SR, ACCURACY, BIAS, ROC, TSS, KAPPA, OR, ORSS, CSI, ETS, BOYCE, MPA
obs	a vector of observed values (binary, 0 or 1)
fit	a vector of fitted values (continuous)
nb.thresh	an integer corresponding to the number of thresholds to be tested over the range of fitted values
threshold	<i>(optional, default NULL)</i> A numeric corresponding to the threshold used to convert the given data
boyce.bg.env	<i>(optional, default NULL)</i> A matrix, data.frame, <a href="#">SpatVector</a> or <a href="#">SpatRaster</a> object containing values of environmental variables (in columns or layers) extracted from the background <i>(if presences are to be compared to background instead of absences or pseudo-absences selected for modeling)</i> <i>Note that old format from <b>raster</b> and <b>sp</b> are still supported such as RasterStack and SpatialPointsDataFrame objects.</i>

mpa.perc	a numeric between 0 and 1 corresponding to the percentage of correctly classified presences for Minimal Predicted Area (see <code>ecospat.mpa()</code> in <b>ecospat</b> )
misc	a matrix corresponding to a contingency table

### Details

- simple**
- POD : Probability of detection (hit rate)
  - FAR : False alarm ratio
  - POFD : Probability of false detection (fall-out)
  - SR : Success ratio
  - ACCURACY : Accuracy (fraction correct)
  - BIAS : Bias score (frequency bias)

- complex**
- ROC : Relative operating characteristic
  - TSS : True skill statistic (Hanssen and Kuipers discriminant, Peirce's skill score)
  - KAPPA : Cohen's Kappa (Heidke skill score)
  - OR : Odds Ratio
  - ORSS : Odds ratio skill score (Yule's Q)
  - CSI : Critical success index (threat score)
  - ETS : Equitable threat score (Gilbert skill score)

- presence-only**
- BOYCE : Boyce index
  - MPA : Minimal predicted area (cutoff optimizing MPA to predict 90% of presences)

Optimal value of each method can be obtained with the `get_optim_value` function.

Please refer to the [CAWRC website \(section "Methods for dichotomous forecasts"\)](#) to get detailed description of each metric.

Note that if a value is given to threshold, no optimization will be done., and only the score for this threshold will be returned.

The Boyce index returns NA values for SRE models because it can not be calculated with binary predictions.

This is also the reason why some NA values might appear for GLM models if they do not converge.

### Value

A 1 row x 5 columns data.frame containing :

- `metric.eval` : the chosen evaluation metric
- `cutoff` : the associated cut-off used to transform the continuous values into binary
- `sensitivity` : the sensibility obtained on fitted values with this threshold
- `specificity` : the specificity obtained on fitted values with this threshold
- `best.stat` : the best score obtained for the chosen evaluation metric

### Note

In order to break dependency loop between packages **biomod2** and **ecospat**, code of `ecospat.boyce()` and `ecospat.mpa()` in **ecospat** functions have been copied within this file from version 3.2.2 (august 2022).



**Author(s)**

Damien Georges

**References**

- Engler, R., Guisan, A., and Rechsteiner L. 2004. An improved approach for predicting the distribution of rare and endangered species from occurrence and pseudo-absence data. *Journal of Applied Ecology*, **41(2)**, 263-274.
- Hirzel, A. H., Le Lay, G., Helfer, V., Randin, C., and Guisan, A. 2006. Evaluating the ability of habitat suitability models to predict species presences. *Ecological Modelling*, **199(2)**, 142-152.

**See Also**

`ecospat.boyce()` and `ecospat.mpa()` in **ecospat**, [BIOMOD\\_Modeling](#), [bm\\_RunModelsLoop](#), [BIOMOD\\_EnsembleModeling](#)

Other Secondary functions: [bm\\_BinaryTransformation\(\)](#), [bm\\_CrossValidation\(\)](#), [bm\\_MakeFormula\(\)](#), [bm\\_ModelingOptions\(\)](#), [bm\\_PlotEvalBoxplot\(\)](#), [bm\\_PlotEvalMean\(\)](#), [bm\\_PlotRangeSize\(\)](#), [bm\\_PlotResponseCurves\(\)](#), [bm\\_PlotVarImpBoxplot\(\)](#), [bm\\_PseudoAbsences\(\)](#), [bm\\_RunModelsLoop\(\)](#), [bm\\_SRE\(\)](#), [bm\\_SampleBinaryVector\(\)](#), [bm\\_SampleFactorLevels\(\)](#), [bm\\_Tuning\(\)](#), [bm\\_VariablesImportance\(\)](#)

**Examples**

```
## Generate a binary vector
vec.a <- sample(c(0, 1), 100, replace = TRUE)

## Generate a 0-1000 vector (random drawing)
vec.b <- runif(100, min = 0, max = 1000)

## Generate a 0-1000 vector (biased drawing)
BiasedDrawing <- function(x, m1 = 300, sd1 = 200, m2 = 700, sd2 = 200) {
  return(ifelse(x < 0.5, rnorm(1, m1, sd1), rnorm(1, m2, sd2)))
}
vec.c <- sapply(vec.a, BiasedDrawing)
vec.c[which(vec.c < 0)] <- 0
vec.c[which(vec.c > 1000)] <- 1000

## Find optimal threshold for a specific evaluation metric
bm_FindOptimStat(metric.eval = 'TSS', fit = vec.b, obs = vec.a)
bm_FindOptimStat(metric.eval = 'TSS', fit = vec.c, obs = vec.a, nb.thresh = 100)
bm_FindOptimStat(metric.eval = 'TSS', fit = vec.c, obs = vec.a, threshold = 280)
```

**Description**

This internal **biomod2** function allows the user to create easily a standardized formula that can be used later by statistical models.

**Usage**

```
bm_MakeFormula(
  resp.name,
  expl.var,
  type = "simple",
  interaction.level = 0,
  k = NULL
)
```

**Arguments**

resp.name	a character corresponding to the response variable name
expl.var	a matrix or data.frame containing the explanatory variables that will be used at the modeling step
type	a character corresponding to the wanted type of formula, must be simple, quadratic, polynomial or s_smoother
interaction.level	an integer corresponding to the interaction level depth between explanatory variables
k	(optional, default NULL) An integer corresponding to the smoothing parameter value of <code>s</code> or <code>s</code> arguments (used only if type = 's_smoother')

**Details**

It is advised to give only a subset of `expl.var` table to avoid useless memory consuming.  
If some explanatory variables are factorial, `expl.var` must be a data.frame whose corresponding columns are defined as factor.

**Value**

A `formula` class object that can be directly given to most of R statistical models.

**Author(s)**

Damien Georges

**See Also**

[formula](#), [s](#), [s](#), [bm\\_ModelingOptions](#), [bm\\_Tuning](#), [bm\\_RunModelsLoop](#)

Other Secondary functions: [bm\\_BinaryTransformation\(\)](#), [bm\\_CrossValidation\(\)](#), [bm\\_FindOptimStat\(\)](#), [bm\\_ModelingOptions\(\)](#), [bm\\_PlotEvalBoxplot\(\)](#), [bm\\_PlotEvalMean\(\)](#), [bm\\_PlotRangeSize\(\)](#), [bm\\_PlotResponseCurves\(\)](#), [bm\\_PlotVarImpBoxplot\(\)](#), [bm\\_PseudoAbsences\(\)](#), [bm\\_RunModelsLoop\(\)](#), [bm\\_SRE\(\)](#), [bm\\_SampleBinaryVector\(\)](#), [bm\\_SampleFactorLevels\(\)](#), [bm\\_Tuning\(\)](#), [bm\\_VariablesImportance\(\)](#)

**Examples**

```
## Create simple simulated data
myResp.s <- sample(c(0, 1), 20, replace = TRUE)
myExpl.s <- data.frame(var1 = sample(c(0, 1), 100, replace = TRUE),
                      var2 = rnorm(100),
                      var3 = 1:100)

## Generate automatic formula
bm_MakeFormula(resp.name = 'myResp.s',
              expl.var = head(myExpl.s),
              type = 'quadratic',
              interaction.level = 0)
```

---

bm\_ModelingOptions      *Configure the modeling options for each selected model*

---

**Description**

Parameterize and/or tune **biomod2**'s single models options.

**Usage**

```
bm_ModelingOptions(
  data.type,
  models = c("ANN", "CTA", "FDA", "GAM", "GBM", "GLM", "MARS", "MAXENT", "MAXNET", "RF",
            "RFd", "SRE", "XGBOOST"),
  strategy,
  user.val = NULL,
  user.base = "bigboss",
  bm.format = NULL,
  calib.lines = NULL
)
```

**Arguments**

data.type	a character corresponding to the data type to be used, must be either binary, binary.PA, abundance, compositional. For the moment, only binary is accept.
models	a vector containing model names to be computed, must be among ANN, CTA, FDA, GAM, GBM, GLM, MARS, MAXENT, MAXNET, RF, RFd, SRE, XGBOOST
strategy	a character corresponding to the method to select models' parameters values, must be either default, bigboss, user.defined, tuned
user.val	(optional, default NULL) A list containing parameters values for some (all) models

user.base	(optional, default bigboss) A character, default or bigboss used when strategy = 'user.defined'. It sets the bases of parameters to be modified by user defined values.
bm.format	(optional, default NULL) A <a href="#">BIOMOD.formated.data</a> or <a href="#">BIOMOD.formated.data.PA</a> object returned by the <a href="#">BIOMOD_FormattingData</a> function
calib.lines	(optional, default NULL) A data.frame object returned by <a href="#">get_calib_lines</a> or <a href="#">bm_CrossValidation</a> functions

### Details

This function creates a [BIOMOD.models.options](#) object containing parameter values for each single model that can be run within **biomod2** through [BIOMOD\\_Modeling](#) function.

12 models are currently available, and are listed within the [ModelsTable](#) dataset.

Different strategies are available to set those parameters, through the strategy argument :

**default** all parameters names and values are directly retrieve from functions to be called through [formalArgs](#) and [formals](#) functions respectively

**bigboss** default parameter values are updated with values predefined by **biomod2** team

**user.defined** default parameter values are updated with values provided by the user

**tuned** default parameter values are updated by calling [bm\\_Tuning](#) function

To define the same options for all datasets of a model, you can provide these options as a list in user.val with the names "for\_all\_datasets".

### Value

A [BIOMOD.models.options](#) of object that can be used to build species distribution model(s) with the [BIOMOD\\_Modeling](#) function.

### Note

MAXENT being the only external model (not called through a R package), default parameters, and their values, are the following :

- path\_to\_maxent.jar = getwd() : a character corresponding to path to maxent.jar file
- memory\_allocated = 512 : an integer corresponding to the amount of memory (in Mo) reserved for java to run MAXENT, must be either 64, 128, 256, 512, 1024... or NULL to use default java memory limitation parameter
- initial\_heap\_size = NULL : a character corresponding to initial heap space (shared memory space) allocated to java (argument -Xms when calling java), must be either 1024K, 4096M, 10G ... or NULL to use default java parameter. Used in [BIOMOD\\_Projection](#) but not in [BIOMOD\\_Modeling](#).
- max\_heap\_size = NULL : a character corresponding to maximum heap space (shared memory space) allocated to java (argument -Xmx when calling java), must be either 1024K, 4096M, 10G ... or NULL to use default java parameter, and must be larger than initial\_heap\_size. Used in [BIOMOD\\_Projection](#) but not in [BIOMOD\\_Modeling](#).

- `background_data_dir = 'default'` : a character corresponding to path to folder where explanatory variables are stored as ASCII files (raster format). If specified, MAXENT will generate its own background data from rasters of explanatory variables ('default' value). Otherwise **biomod2** pseudo-absences will be used (see [BIOMOD\\_FormatingData](#)).
- `visible = FALSE` : a logical value defining whether MAXENT user interface is to be used or not
- `linear = TRUE` : a logical value defining whether linear features are to be used or not
- `quadratic = TRUE` : a logical value defining whether quadratic features are to be used or not
- `product = TRUE` : a logical value defining whether product features are to be used or not
- `threshold = TRUE` : a logical value defining whether threshold features are to be used or not
- `hinge = TRUE` : a logical value defining whether hinge features are to be used or not
- `l2lqthreshold = 10` : an integer corresponding to the number of samples at which quadratic features start being used
- `lq2lqptthreshold = 80` : an integer corresponding to the number of samples at which product and threshold features start being used
- `hingethreshold = 15` : an integer corresponding to the number of samples at which hinge features start being used
- `beta_lqp = -1.0` : a numeric corresponding to the regularization parameter to be applied to all linear, quadratic and product features (*negative value enables automatic setting*)
- `beta_threshold = -1.0` : a numeric corresponding to the regularization parameter to be applied to all threshold features (*negative value enables automatic setting*)
- `beta_hinge = -1.0` : a numeric corresponding to the regularization parameter to be applied to all hinge features (*negative value enables automatic setting*)
- `beta_categorical = -1.0` : a numeric corresponding to the regularization parameter to be applied to all categorical features (*negative value enables automatic setting*)
- `betamultiplier = 1` : a numeric corresponding to the number by which multiply all automatic regularization parameters (*higher number gives a more spread-out distribution*)
- `defaultprevalence = 0.5` : a numeric corresponding to the default prevalence of the modelled species (*probability of presence at ordinary occurrence points*)

### Author(s)

Damien Georges, Wilfried Thuiller, Maya Gueguen

### See Also

[ModelsTable](#), [BIOMOD.models.options](#), [bm\\_Tuning](#), [BIOMOD\\_Modeling](#)

Other Secondary functions: [bm\\_BinaryTransformation\(\)](#), [bm\\_CrossValidation\(\)](#), [bm\\_FindOptimStat\(\)](#), [bm\\_MakeFormula\(\)](#), [bm\\_PlotEvalBoxplot\(\)](#), [bm\\_PlotEvalMean\(\)](#), [bm\\_PlotRangeSize\(\)](#), [bm\\_PlotResponseCurves\(\)](#), [bm\\_PlotVarImpBoxplot\(\)](#), [bm\\_PseudoAbsences\(\)](#), [bm\\_RunModelsLoop\(\)](#), [bm\\_SRE\(\)](#), [bm\\_SampleBinaryVector\(\)](#), [bm\\_SampleFactorLevels\(\)](#), [bm\\_Tuning\(\)](#), [bm\\_VariablesImportance\(\)](#)

**Examples**

```

library(terra)

# Load species occurrences (6 species available)
data(DataSpecies)
head(DataSpecies)

# Select the name of the studied species
myRespName <- 'GuloGulo'

# Get corresponding presence/absence data
myResp <- as.numeric(DataSpecies[, myRespName])

# Get corresponding XY coordinates
myRespXY <- DataSpecies[, c('X_WGS84', 'Y_WGS84')]

# Load environmental variables extracted from BIOCLIM (bio_3, bio_4, bio_7, bio_11 & bio_12)
data(bioclim_current)
myExpl <- terra::rast(bioclim_current)

# -----#
# Format Data with true absences
myBiomodData <- BIOMOD_FormatingData(resp.var = myResp,
                                   expl.var = myExpl,
                                   resp.xy = myRespXY,
                                   resp.name = myRespName)

# k-fold selection
cv.k <- bm_CrossValidation(bm.format = myBiomodData,
                          strategy = 'kfold',
                          nb.rep = 2,
                          k = 3)

# -----#
allModels <- c('ANN', 'CTA', 'FDA', 'GAM', 'GBM', 'GLM'
              , 'MARS', 'MAXENT', 'MAXNET', 'RF', 'SRE', 'XGBOOST')

# default parameters
opt.d <- bm_ModelingOptions(data.type = 'binary',
                           models = allModels,
                           strategy = 'default')

# providing formatted data
opt.df <- bm_ModelingOptions(data.type = 'binary',
                            models = allModels,
                            strategy = 'default',
                            bm.format = myBiomodData,
                            calib.lines = cv.k)

```

```

opt.d
opt.d@models
opt.d@options$ANN.binary.nnet.nnet
names(opt.d@options$ANN.binary.nnet.nnet@args.values)

opt.df@options$ANN.binary.nnet.nnet
names(opt.df@options$ANN.binary.nnet.nnet@args.values)

# -----#
# bigboss parameters
opt.b <- bm_ModelingOptions(data.type = 'binary',
                             models = allModels,
                             strategy = 'bigboss')

# user defined parameters
user.SRE <- list('_allData_allRun' = list(quant = 0.01))
user.XGBOOST <- list('_allData_allRun' = list(nrounds = 10))
user.val <- list(SRE.binary.biomod2.bm_SRE = user.SRE
                 , XGBOOST.binary.xgboost.xgboost = user.XGBOOST)

opt.u <- bm_ModelingOptions(data.type = 'binary',
                             models = c('SRE', 'XGBOOST'),
                             strategy = 'user.defined',
                             user.val = user.val)

opt.b
opt.u

## Not run:
# tuned parameters with formatted data
opt.t <- bm_ModelingOptions(data.type = 'binary',
                             models = c('SRE', 'XGBOOST'),
                             strategy = 'tuned',
                             bm.format = myBiomodData)

opt.t

## End(Not run)

```

---

bm\_PlotEvalBoxplot      *Plot boxplot of evaluation scores*

---

## Description

This function represents boxplot of evaluation scores of species distribution models, from [BIOMOD.models.out](#) or [BIOMOD.ensemble.models.out](#) objects that can be obtained from [BIOMOD\\_Modeling](#) or [BIOMOD\\_EnsembleModeling](#) functions. Scores are represented according to 2 grouping methods (see Details).

**Usage**

```
bm_PlotEvalBoxplot(
  bm.out,
  dataset = "calibration",
  group.by = c("algo", "run"),
  do.plot = TRUE,
  ...
)
```

**Arguments**

bm.out	a <a href="#">BIOMOD.models.out</a> or <a href="#">BIOMOD.ensemble.models.out</a> object that can be obtained with the <a href="#">BIOMOD_Modeling</a> or <a href="#">BIOMOD_EnsembleModeling</a> functions
dataset	a character corresponding to the dataset upon which evaluation metrics have been calculated and that is to be represented, must be among calibration, validation, evaluation
group.by	a 2-length vector containing the way kept models will be represented, must be among full.name, PA, run, algo (if bm.out is a <a href="#">BIOMOD.models.out</a> object), or full.name, merged.by.PA, merged.by.run, merged.by.algo (if bm.out is a <a href="#">BIOMOD.ensemble.models.out</a> object)
do.plot	( <i>optional, default TRUE</i> ) A logical value defining whether the plot is to be rendered or not
...	some additional arguments (see Details)

**Details**

... can take the following values :

- main : a character corresponding to the graphic title
- scales : a character corresponding to the scales argument of the [facet\\_wrap](#) function, must be either fixed, free\_x, free\_y or free

**Value**

A list containing a data.frame with evaluation scores and the corresponding ggplot object representing them in boxplot.

**Author(s)**

Damien Georges, Maya Gueguen

**See Also**

[BIOMOD.models.out](#), [BIOMOD.ensemble.models.out](#), [BIOMOD\\_Modeling](#), [BIOMOD\\_EnsembleModeling](#), [get\\_evaluations](#)

Other Secondary functions: [bm\\_BinaryTransformation\(\)](#), [bm\\_CrossValidation\(\)](#), [bm\\_FindOptimStat\(\)](#), [bm\\_MakeFormula\(\)](#), [bm\\_ModelingOptions\(\)](#), [bm\\_PlotEvalMean\(\)](#), [bm\\_PlotRangeSize\(\)](#), [bm\\_PlotResponseCurves\(\)](#),



bm\_PlotVarImpBoxplot(), bm\_PseudoAbsences(), bm\_RunModelsLoop(), bm\_SRE(), bm\_SampleBinaryVector(),  
 bm\_SampleFactorLevels(), bm\_Tuning(), bm\_VariablesImportance()

Other Plot functions: bm\_PlotEvalMean(), bm\_PlotRangeSize(), bm\_PlotResponseCurves(),  
 bm\_PlotVarImpBoxplot()

## Examples

```
library(terra)

# Load species occurrences (6 species available)
data(DataSpecies)
head(DataSpecies)

# Select the name of the studied species
myRespName <- 'GuloGulo'

# Get corresponding presence/absence data
myResp <- as.numeric(DataSpecies[, myRespName])

# Get corresponding XY coordinates
myRespXY <- DataSpecies[, c('X_WGS84', 'Y_WGS84')]

# Load environmental variables extracted from BIOCLIM (bio_3, bio_4, bio_7, bio_11 & bio_12)
data(bioclim_current)
myExpl <- terra::rast(bioclim_current)

# -----
file.out <- paste0(myRespName, "/", myRespName, ".AllModels.models.out")
if (file.exists(file.out)) {
  myBiomodModelOut <- get(load(file.out))
} else {

  # Format Data with true absences
  myBiomodData <- BIOMOD_FormatingData(resp.var = myResp,
                                     expl.var = myExpl,
                                     resp.xy = myRespXY,
                                     resp.name = myRespName)

  # Model single models
  myBiomodModelOut <- BIOMOD_Modeling(bm.format = myBiomodData,
                                     modeling.id = 'AllModels',
                                     models = c('RF', 'GLM'),
                                     CV.strategy = 'random',
                                     CV.nb.rep = 2,
                                     CV.perc = 0.8,
                                     OPT.strategy = 'bigboss',
                                     metric.eval = c('TSS', 'ROC'),
                                     var.import = 3,
                                     seed.val = 42)
}
```

```
# -----
# Get evaluation scores
get_evaluations(myBiomodModelOut)

# Represent evaluation scores
bm_PlotEvalBoxplot(bm.out = myBiomodModelOut, group.by = c('algo', 'run'))
```

---

bm_PlotEvalMean	<i>Plot mean evaluation scores</i>
-----------------	------------------------------------

---

### Description

This function represents mean evaluation scores (and their standard deviation) of species distribution models, from [BIOMOD.models.out](#) or [BIOMOD.ensemble.models.out](#) objects that can be obtained from [BIOMOD\\_Modeling](#) or [BIOMOD\\_EnsembleModeling](#) functions. Scores are represented according to 2 different evaluation methods, and models can be grouped (see Details).

### Usage

```
bm_PlotEvalMean(
  bm.out,
  metric.eval = NULL,
  dataset = "calibration",
  group.by = "algo",
  do.plot = TRUE,
  ...
)
```

### Arguments

bm.out	a <a href="#">BIOMOD.models.out</a> or <a href="#">BIOMOD.ensemble.models.out</a> object that can be obtained with the <a href="#">BIOMOD_Modeling</a> or <a href="#">BIOMOD_EnsembleModeling</a> functions
metric.eval	a vector containing evaluation metric names to be used, must be among POD, FAR, POFD, SR, ACCURACY, BIAS, ROC, TSS, KAPPA, OR, ORSS, CSI, ETS, BOYCE, MPA
dataset	a character corresponding to the dataset upon which evaluation metrics have been calculated and that is to be represented, must be among calibration, validation, evaluation
group.by	a character corresponding to the way kept models will be combined to compute mean and sd evaluation scores, must be among full.name, PA, run, algo (if bm.out is a <a href="#">BIOMOD.models.out</a> object), or full.name, merged.by.PA, merged.by.run, merged.by.algo (if bm.out is a <a href="#">BIOMOD.ensemble.models.out</a> object)

```
do.plot      (optional, default TRUE)
              A logical value defining whether the plot is to be rendered or not
...          some additional arguments (see Details)
```

### Details

... can take the following values :

- xlim : an integer corresponding to the x maximum limit to represent
- ylim : an integer corresponding to the y maximum limit to represent
- main : a character corresponding to the graphic title
- col : a vector containing new color values

### Value

A list containing a data.frame with mean and standard deviation of evaluation scores and the corresponding ggplot object representing them according to 2 different evaluation methods.

### Author(s)

Damien Georges, Maya Gueguen

### See Also

[BIOMOD.models.out](#), [BIOMOD.ensemble.models.out](#), [BIOMOD\\_Modeling](#), [BIOMOD\\_EnsembleModeling](#), [get\\_evaluations](#)

Other Secondary functions: [bm\\_BinaryTransformation\(\)](#), [bm\\_CrossValidation\(\)](#), [bm\\_FindOptimStat\(\)](#), [bm\\_MakeFormula\(\)](#), [bm\\_ModelingOptions\(\)](#), [bm\\_PlotEvalBoxplot\(\)](#), [bm\\_PlotRangeSize\(\)](#), [bm\\_PlotResponseCurves\(\)](#), [bm\\_PlotVarImpBoxplot\(\)](#), [bm\\_PseudoAbsences\(\)](#), [bm\\_RunModelsLoop\(\)](#), [bm\\_SRE\(\)](#), [bm\\_SampleBinaryVector\(\)](#), [bm\\_SampleFactorLevels\(\)](#), [bm\\_Tuning\(\)](#), [bm\\_VariablesImportance\(\)](#)

Other Plot functions: [bm\\_PlotEvalBoxplot\(\)](#), [bm\\_PlotRangeSize\(\)](#), [bm\\_PlotResponseCurves\(\)](#), [bm\\_PlotVarImpBoxplot\(\)](#)

### Examples

```
library(terra)

# Load species occurrences (6 species available)
data(DataSpecies)
head(DataSpecies)

# Select the name of the studied species
myRespName <- 'GuloGulo'

# Get corresponding presence/absence data
myResp <- as.numeric(DataSpecies[, myRespName])

# Get corresponding XY coordinates
myRespXY <- DataSpecies[, c('X_WGS84', 'Y_WGS84')]
```

```

# Load environmental variables extracted from BIOCLIM (bio_3, bio_4, bio_7, bio_11 & bio_12)
data(bioclim_current)
myExpl <- terra::rast(bioclim_current)

# -----
file.out <- paste0(myRespName, "/", myRespName, ".AllModels.models.out")
if (file.exists(file.out)) {
  myBiomodModelOut <- get(load(file.out))
} else {

  # Format Data with true absences
  myBiomodData <- BIOMOD_FormatingData(resp.var = myResp,
                                     expl.var = myExpl,
                                     resp.xy = myRespXY,
                                     resp.name = myRespName)

  # Model single models
  myBiomodModelOut <- BIOMOD_Modeling(bm.format = myBiomodData,
                                     modeling.id = 'AllModels',
                                     models = c('RF', 'GLM'),
                                     CV.strategy = 'random',
                                     CV.nb.rep = 2,
                                     CV.perc = 0.8,
                                     OPT.strategy = 'bigboss',
                                     metric.eval = c('TSS', 'ROC'),
                                     var.import = 3,
                                     seed.val = 42)
}

# -----
# Get evaluation scores
get_evaluations(myBiomodModelOut)

# Represent mean evaluation scores
bm_PlotEvalMean(bm.out = myBiomodModelOut)

```

---

bm\_PlotRangeSize

*Plot species range change*


---

### Description

This function represents species range change from object that can be obtained from [BIOMOD\\_RangeSize](#) function. Several graphics can be obtained, representing global counts or proportions of gains / losses, as well as spatial representations (see Details).

**Usage**

```

bm_PlotRangeSize(
  bm.range,
  do.count = TRUE,
  do.perc = TRUE,
  do.maps = TRUE,
  do.mean = TRUE,
  do.plot = TRUE,
  row.names = c("Species", "Dataset", "Run", "Algo")
)

```

**Arguments**

bm.range	an object returned by the <a href="#">BIOMOD_RangeSize</a> function
do.count	<i>(optional, default TRUE)</i> A logical value defining whether the count plot is to be computed or not
do.perc	<i>(optional, default TRUE)</i> A logical value defining whether the percentage plot is to be computed or not
do.maps	<i>(optional, default TRUE)</i> A logical value defining whether the maps plot is to be computed or not
do.mean	<i>(optional, default TRUE)</i> A logical value defining whether the mean maps plot is to be computed or not
do.plot	<i>(optional, default TRUE)</i> A logical value defining whether the plots are to be rendered or not
row.names	<i>(optional, default c('Species', 'Dataset', 'Run', 'Algo'))</i> A vector containing tags matching <code>bm.range\$Compt.By.Models</code> rownames splitted by <code>'_'</code> character

**Details**

4 plots can be obtained with this function :

**Count barplot** representing absolute number of locations (pixels) lost, stable and gained

**Percentage barplot** representing percentage of locations (pixels) lost, stable, and the corresponding Species Range Change (PercGain - PercLoss)

**SRC models maps** representing spatially locations (pixels) lost, stable and gained for each single distribution model

**SRC community averaging maps** representing spatially locations (pixels) lost, stable and gained, taking the majoritary value across single distribution models (and representing the percentage of models' agreement)

*Please see [BIOMOD\\_RangeSize](#) function for more details about the values.*

**Value**

A list containing one or several `data.frame` and the corresponding `ggplot` object representing species range change.



```

        models = c('RF', 'GLM'),
        CV.strategy = 'random',
        CV.nb.rep = 2,
        CV.perc = 0.8,
        OPT.strategy = 'bigboss',
        metric.eval = c('TSS', 'ROC'),
        var.import = 3,
        seed.val = 42)
}

models.proj <- get_built_models(myBiomodModelOut, algo = "RF")
# Project single models
myBiomodProj <- BIOMOD_Projection(bm.mod = myBiomodModelOut,
                                proj.name = 'CurrentRangeSize',
                                new.env = myExpl,
                                models.chosen = models.proj,
                                metric.binary = 'all')

# -----#
# Load environmental variables extracted from BIOCLIM (bio_3, bio_4, bio_7, bio_11 & bio_12)
data(bioclim_future)
myExplFuture <- terra::rast(bioclim_future)

# Project onto future conditions
myBiomodProjectionFuture <- BIOMOD_Projection(bm.mod = myBiomodModelOut,
                                              proj.name = 'FutureRangeSize',
                                              new.env = myExplFuture,
                                              models.chosen = models.proj,
                                              metric.binary = 'TSS')

# Load current and future binary projections
CurrentProj <- get_predictions(myBiomodProj,
                              metric.binary = "TSS",
                              model.as.col = TRUE)
FutureProj <- get_predictions(myBiomodProjectionFuture,
                             metric.binary = "TSS",
                             model.as.col = TRUE)

# Compute differences
myBiomodRangeSize <- BIOMOD_RangeSize(proj.current = CurrentProj, proj.future = FutureProj)

# -----#
myBiomodRangeSize$Compt.By.Models
plot(myBiomodRangeSize$Diff.By.Pixel)

# Represent main results
bm_PlotRangeSize(bm.range = myBiomodRangeSize)

```

---

 bm\_PlotResponseCurves *Plot response curves*


---

### Description

This function represents response curves of species distribution models, from [BIOMOD.models.out](#) or [BIOMOD.ensemble.models.out](#) objects that can be obtained from [BIOMOD\\_Modeling](#) or [BIOMOD\\_EnsembleModeling](#) functions. Response curves can be represented in either 2 or 3 dimensions (meaning 1 or 2 explanatory variables at a time, see Details).

### Usage

```
bm_PlotResponseCurves(
  bm.out,
  models.chosen = "all",
  new.env = get_formal_data(bm.out, "expl.var"),
  show.variables = get_formal_data(bm.out, "expl.var.names"),
  fixed.var = "mean",
  do.bivariate = FALSE,
  do.plot = TRUE,
  do.progress = TRUE,
  ...
)
```

### Arguments

bm.out	a <a href="#">BIOMOD.models.out</a> or <a href="#">BIOMOD.ensemble.models.out</a> object that can be obtained with the <a href="#">BIOMOD_Modeling</a> or <a href="#">BIOMOD_EnsembleModeling</a> functions
models.chosen	a vector containing model names to be kept, must be either all or a sub-selection of model names that can be obtained with the <a href="#">get_built_models</a> function
new.env	a matrix, data.frame or <a href="#">SpatRaster</a> object containing the new explanatory variables (in columns or layers, with names matching the variables names given to the <a href="#">BIOMOD_FormatingData</a> function to build bm.out) that will be used to project the species distribution model(s) <i>Note that old format from <b>raster</b> are still supported such as RasterStack objects.</i>
show.variables	a vector containing the names of the explanatory variables present into new.env parameter and to be plotted
fixed.var	a character corresponding to the statistic to be used to fix as constant the remaining variables other than the one used to predict response, must be either mean, median, min, max
do.bivariate	<i>(optional, default FALSE)</i> A logical value defining whether the response curves are to be represented in 3 dimensions (meaning 2 explanatory variables at a time) or not (meaning only 1)



do.plot	(optional, default TRUE) A logical value defining whether the plot is to be rendered or not
do.progress	(optional, default TRUE) A logical value defining whether the progress bar is to be rendered or not
...	some additional arguments (see Details)

### Details

This function is an adaptation of the Evaluation Strip method proposed by Elith et al. (2005). To build the predicted response curves :

- n-1 variables are set constant to a fixed value determined by the `fixed.var` parameter (in the case of categorical variable, the most represented class is taken)
- the remaining variable is made to vary throughout its range given by the `new.env` parameter
- predicted values are computed with these n-1 fixed variables, and this studied variable varying

If `do.bivariate = TRUE`, 2 variables are varying at the same time.

The response curves obtained show the sensibility of the model to the studied variable. Note that this method does not account for interactions between variables.

... can take the following values :

- `main` : a character corresponding to the graphic title

### Value

A list containing a data.frame with variables and predicted values and the corresponding ggplot object representing response curves.

### Author(s)

Damien Georges, Maya Gueguen

### References

- Elith, J., Ferrier, S., Huettmann, FALSE. and Leathwick, J. R. 2005. The evaluation strip: A new and robust method for plotting predicted responses from species distribution models. *Ecological Modelling*, **186**, 280-289.

### See Also

[BIOMOD.models.out](#), [BIOMOD.ensemble.models.out](#), [BIOMOD\\_Modeling](#), [BIOMOD\\_EnsembleModeling](#)

Other Secondary functions: [bm\\_BinaryTransformation\(\)](#), [bm\\_CrossValidation\(\)](#), [bm\\_FindOptimStat\(\)](#), [bm\\_MakeFormula\(\)](#), [bm\\_ModelingOptions\(\)](#), [bm\\_PlotEvalBoxplot\(\)](#), [bm\\_PlotEvalMean\(\)](#), [bm\\_PlotRangeSize\(\)](#), [bm\\_PlotVarImpBoxplot\(\)](#), [bm\\_PseudoAbsences\(\)](#), [bm\\_RunModelsLoop\(\)](#), [bm\\_SRE\(\)](#), [bm\\_SampleBinaryVector\(\)](#), [bm\\_SampleFactorLevels\(\)](#), [bm\\_Tuning\(\)](#), [bm\\_VariablesImportance\(\)](#)

Other Plot functions: [bm\\_PlotEvalBoxplot\(\)](#), [bm\\_PlotEvalMean\(\)](#), [bm\\_PlotRangeSize\(\)](#), [bm\\_PlotVarImpBoxplot\(\)](#)

**Examples**

```

library(terra)

# Load species occurrences (6 species available)
data(DataSpecies)
head(DataSpecies)

# Select the name of the studied species
myRespName <- 'GuloGulo'

# Get corresponding presence/absence data
myResp <- as.numeric(DataSpecies[, myRespName])

# Get corresponding XY coordinates
myRespXY <- DataSpecies[, c('X_WGS84', 'Y_WGS84')]

# Load environmental variables extracted from BIOCLIM (bio_3, bio_4, bio_7, bio_11 & bio_12)
data(bioclim_current)
myExpl <- terra::rast(bioclim_current)

# -----#
file.out <- paste0(myRespName, "/", myRespName, ".AllModels.models.out")
if (file.exists(file.out)) {
  myBiomodModelOut <- get(load(file.out))
} else {

  # Format Data with true absences
  myBiomodData <- BIOMOD_FormatingData(resp.var = myResp,
                                     expl.var = myExpl,
                                     resp.xy = myRespXY,
                                     resp.name = myRespName)

  # Model single models
  myBiomodModelOut <- BIOMOD_Modeling(bm.format = myBiomodData,
                                     modeling.id = 'AllModels',
                                     models = c('RF', 'GLM'),
                                     CV.strategy = 'random',
                                     CV.nb.rep = 2,
                                     CV.perc = 0.8,
                                     OPT.strategy = 'bigboss',
                                     metric.eval = c('TSS', 'ROC'),
                                     var.import = 3,
                                     seed.val = 42)
}

# -----#
# Represent response curves
mods <- get_built_models(myBiomodModelOut, run = 'RUN1')
bm_PlotResponseCurves(bm.out = myBiomodModelOut,

```

```

                                models.chosen = mods,
                                fixed.var = 'median')
## fixed.var can also be set to 'min', 'max' or 'mean'
# bm_PlotResponseCurves(bm.out = myBiomodModelOut,
#                         models.chosen = mods,
#                         fixed.var = 'min')

# Bivariate case (one model)
# variables can be selected with argument 'show.variables'
# models can be selected with argument 'models.chosen'
mods <- get_built_models(myBiomodModelOut, full.name = 'GuloGulo_allData_RUN2_RF')
bm_PlotResponseCurves(bm.out = myBiomodModelOut,
                       show.variables = c("bio4", "bio12", "bio11"),
                       models.chosen = mods,
                       fixed.var = 'median',
                       do.bivariate = TRUE)

```

---

bm\_PlotVarImpBoxplot *Plot boxplot of variables importance*

---

## Description

This function represents boxplot of variables importance of species distribution models, from [BIOMOD.models.out](#) or [BIOMOD.ensemble.models.out](#) objects that can be obtained from [BIOMOD\\_Modeling](#) or [BIOMOD\\_EnsembleModeling](#) functions. Scores are represented according to 3 grouping methods (see Details).

## Usage

```

bm_PlotVarImpBoxplot(
  bm.out,
  group.by = c("run", "expl.var", "algo"),
  do.plot = TRUE,
  ...
)

```

## Arguments

bm.out	a <a href="#">BIOMOD.models.out</a> or <a href="#">BIOMOD.ensemble.models.out</a> object that can be obtained with the <a href="#">BIOMOD_Modeling</a> or <a href="#">BIOMOD_EnsembleModeling</a> functions
group.by	a 3-length vector containing the way kept models will be represented, must be among full.name, PA, run, algo, expl.var (if bm.out is a <a href="#">BIOMOD.models.out</a> object), or full.name, merged.by.PA, merged.by.run, merged.by.algo, expl.var (if bm.out is a <a href="#">BIOMOD.ensemble.models.out</a> object)
do.plot	(optional, default TRUE) A logical value defining whether the plot is to be rendered or not
...	some additional arguments (see Details)

**Details**

... can take the following values :

- `main` : a character corresponding to the graphic title

**Value**

A list containing a `data.frame` with variables importance and the corresponding `ggplot` object representing them in boxplot.

**Author(s)**

Damien Georges, Maya Gueguen

**See Also**

[BIOMOD.models.out](#), [BIOMOD.ensemble.models.out](#), [BIOMOD\\_Modeling](#), [BIOMOD\\_EnsembleModeling](#), [get\\_variables\\_importance](#)

Other Secondary functions: [bm\\_BinaryTransformation\(\)](#), [bm\\_CrossValidation\(\)](#), [bm\\_FindOptimStat\(\)](#), [bm\\_MakeFormula\(\)](#), [bm\\_ModelingOptions\(\)](#), [bm\\_PlotEvalBoxplot\(\)](#), [bm\\_PlotEvalMean\(\)](#), [bm\\_PlotRangeSize\(\)](#), [bm\\_PlotResponseCurves\(\)](#), [bm\\_PseudoAbsences\(\)](#), [bm\\_RunModelsLoop\(\)](#), [bm\\_SRE\(\)](#), [bm\\_SampleBinaryVector\(\)](#), [bm\\_SampleFactorLevels\(\)](#), [bm\\_Tuning\(\)](#), [bm\\_VariablesImportance\(\)](#)

Other Plot functions: [bm\\_PlotEvalBoxplot\(\)](#), [bm\\_PlotEvalMean\(\)](#), [bm\\_PlotRangeSize\(\)](#), [bm\\_PlotResponseCurves\(\)](#)

**Examples**

```
library(terra)

# Load species occurrences (6 species available)
data(DataSpecies)
head(DataSpecies)

# Select the name of the studied species
myRespName <- 'GuloGulo'

# Get corresponding presence/absence data
myResp <- as.numeric(DataSpecies[, myRespName])

# Get corresponding XY coordinates
myRespXY <- DataSpecies[, c('X_WGS84', 'Y_WGS84')]

# Load environmental variables extracted from BIOCLIM (bio_3, bio_4, bio_7, bio_11 & bio_12)
data(bioclim_current)
myExpl <- terra::rast(bioclim_current)

# -----
file.out <- paste0(myRespName, "/", myRespName, ".AllModels.models.out")
if (file.exists(file.out)) {
```

```

myBiomodModelOut <- get(load(file.out))
} else {

# Format Data with true absences
myBiomodData <- BIOMOD_FormatingData(resp.var = myResp,
                                     expl.var = myExpl,
                                     resp.xy = myRespXY,
                                     resp.name = myRespName)

# Model single models
myBiomodModelOut <- BIOMOD_Modeling(bm.format = myBiomodData,
                                    modeling.id = 'AllModels',
                                    models = c('RF', 'GLM'),
                                    CV.strategy = 'random',
                                    CV.nb.rep = 2,
                                    CV.perc = 0.8,
                                    OPT.strategy = 'bigboss',
                                    metric.eval = c('TSS', 'ROC'),
                                    var.import = 3,
                                    seed.val = 42)
}

# -----
# Get variables importance
get_variables_importance(myBiomodModelOut)

# Represent variables importance
bm_PlotVarImpBoxplot(bm.out = myBiomodModelOut, group.by = c('expl.var', 'algo', 'algo'))
bm_PlotVarImpBoxplot(bm.out = myBiomodModelOut, group.by = c('expl.var', 'algo', 'PA'))
bm_PlotVarImpBoxplot(bm.out = myBiomodModelOut, group.by = c('algo', 'expl.var', 'PA'))

```

---

bm\_PseudoAbsences      *Select pseudo-absences*

---

## Description

This internal **biomod2** function allows to select pseudo-absences according to 4 different methods : random, sre, disk or user .defined (see Details).

## Usage

```

bm_PseudoAbsences(
  resp.var,
  expl.var,
  nb.rep = 1,
  strategy = "random",
  nb.absences = NULL,

```

```

    sre.quant = 0,
    dist.min = 0,
    dist.max = NULL,
    fact.aggr = NULL,
    user.table = NULL,
    seed.val = NULL
)

bm_PseudoAbsences_user.defined(resp.var, expl.var, ...)

## S4 method for signature 'ANY,SpatVector'
bm_PseudoAbsences_user.defined(resp.var, expl.var, user.table)

## S4 method for signature 'ANY,SpatRaster'
bm_PseudoAbsences_user.defined(resp.var, expl.var, user.table)

bm_PseudoAbsences_random(resp.var, expl.var, ...)

## S4 method for signature 'ANY,SpatVector'
bm_PseudoAbsences_random(resp.var, expl.var, nb.absences, nb.rep, fact.aggr)

## S4 method for signature 'ANY,SpatRaster'
bm_PseudoAbsences_random(resp.var, expl.var, nb.absences, nb.rep, fact.aggr)

bm_PseudoAbsences_sre(resp.var, expl.var, ...)

## S4 method for signature 'ANY,SpatVector'
bm_PseudoAbsences_sre(resp.var, expl.var, sre.quant, nb.absences, nb.rep)

## S4 method for signature 'ANY,SpatRaster'
bm_PseudoAbsences_sre(resp.var, expl.var, sre.quant, nb.absences, nb.rep)

bm_PseudoAbsences_disk(resp.var, expl.var, ...)

## S4 method for signature 'ANY,SpatVector'
bm_PseudoAbsences_disk(
  resp.var,
  expl.var,
  dist.min,
  dist.max,
  nb.absences,
  nb.rep,
  fact.aggr
)

## S4 method for signature 'ANY,SpatRaster'
bm_PseudoAbsences_disk(
  resp.var,

```

```

    expl.var,
    dist.min,
    dist.max,
    nb.absences,
    nb.rep,
    fact.aggr
  )

```

### Arguments

resp.var	a vector, <a href="#">SpatialPoints</a> or <a href="#">SpatialPointsDataFrame</a> object containing binary data (0 : absence, 1 : presence, NA : indeterminate) for a single species that will be used to find the pseudo-absences
expl.var	a matrix, data.frame, <a href="#">SpatialPointsDataFrame</a> or <a href="#">SpatRaster</a> object containing the explanatory variables (in columns or layers) that will be used to find the pseudo-absences
nb.rep	an integer corresponding to the number of sets (repetitions) of pseudo-absence points that will be drawn
strategy	a character corresponding to the pseudo-absence selection strategy, must be among random, sre, disk or user.defined
nb.absences	<i>(optional, default NULL)</i> If strategy = 'random' or strategy = 'sre' or strategy = 'disk', an integer corresponding to the number of pseudo-absence points that will be selected for each pseudo-absence repetition (true absences included)
sre.quant	<i>(optional, default 0)</i> If strategy = 'sre', a numeric between 0 and 0.5 defining the half-quantile used to make the sre pseudo-absence selection (see <a href="#">bm_SRE</a> )
dist.min	<i>(optional, default 0)</i> If strategy = 'disk', a numeric defining the minimal distance to presence points used to make the disk pseudo-absence selection (in the same projection system units as expl.var)
dist.max	<i>(optional, default NULL)</i> If strategy = 'disk', a numeric defining the maximal distance to presence points used to make the disk pseudo-absence selection (in the same projection system units as expl.var)
fact.aggr	<i>(optional, default NULL)</i> If strategy = 'random' or strategy = 'disk', a integer defining the factor of aggregation to reduce the resolution
user.table	<i>(optional, default NULL)</i> If strategy = 'user.defined', a matrix or data.frame with as many rows as resp.var values, as many columns as nb.rep, and containing TRUE or FALSE values defining which points will be used to build the species distribution model(s) for each repetition
seed.val	<i>(optional, default NULL)</i> An integer value corresponding to the new seed value to be set
...	<i>(optional, one or several of the above arguments depending on the selected method)</i>

## Details

### Concerning random selection :

The idea is to select pseudo-absences randomly in spatial locations where the species has not been sampled. This method is the simplest one and the most appropriate if lacking information about the presence sampling (non-exhaustive, biased sampling, etc).

### Concerning SRE selection (see [bm\\_SRE](#)) :

The idea is to select pseudo-absences in spatial locations whose environmental conditions are different from those of the presence points. This method is appropriate when most of the environmental space of the species has been sampled.

### Concerning disk selection :

The idea is to select pseudo-absences, not too close from presence points, but not too far away either. This method is appropriate when most of the spatial range of the species has been sampled.

### Concerning user defined selection :

The user can provide pseudo-absences locations through a table containing spatial locations in rows, pseudo-absences repetitions in columns, and TRUE/FALSE values indicating whether each point is to be considered as pseudo-absence or not for each dataset.

## Value

A list containing the following elements :

- xy : the coordinates of the species observations
- sp : the values of the species observations (0, 1 or NA)
- env : the explanatory variables
- pa.tab : the corresponding table of selected pseudo-absences (indicated by TRUE or FALSE)

## Author(s)

Wilfried Thuiller, Damien Georges

## See Also

[bm\\_SRE](#), [BIOMOD.formated.data.PA](#), [BIOMOD\\_FormatingData](#)

Other Secondary functions: [bm\\_BinaryTransformation\(\)](#), [bm\\_CrossValidation\(\)](#), [bm\\_FindOptimStat\(\)](#), [bm\\_MakeFormula\(\)](#), [bm\\_ModelingOptions\(\)](#), [bm\\_PlotEvalBoxplot\(\)](#), [bm\\_PlotEvalMean\(\)](#), [bm\\_PlotRangeSize\(\)](#), [bm\\_PlotResponseCurves\(\)](#), [bm\\_PlotVarImpBoxplot\(\)](#), [bm\\_RunModelsLoop\(\)](#), [bm\\_SRE\(\)](#), [bm\\_SampleBinaryVector\(\)](#), [bm\\_SampleFactorLevels\(\)](#), [bm\\_Tuning\(\)](#), [bm\\_VariablesImportance\(\)](#)



**Examples**

```

library(terra)

# Load species occurrences (6 species available)
data(DataSpecies)
head(DataSpecies)

# Select the name of the studied species
myRespName <- 'GuloGulo'

# Get corresponding presence/absence data
myResp <- as.numeric(DataSpecies[, myRespName])

# Get corresponding XY coordinates
myRespXY <- DataSpecies[, c('X_WGS84', 'Y_WGS84')]

# Load environmental variables extracted from BIOCLIM (bio_3, bio_4, bio_7, bio_11 & bio_12)
data(bioclim_current)
myExpl <- terra::rast(bioclim_current)

# ----- #
# Create the different pseudo-absence datasets

# Transform true absences into potential pseudo-absences
myResp.PA <- ifelse(myResp == 1, 1, NA)
myResp.PA.vect <- vect(cbind(myRespXY, myResp.PA), geom = c("X_WGS84", "Y_WGS84"))

# random method
PA.r <- bm_PseudoAbsences(resp.var = myResp.PA.vect,
                          expl.var = myExpl,
                          nb.rep = 4,
                          nb.absences = 1000,
                          strategy = 'random')

# disk method
PA.d <- bm_PseudoAbsences(resp.var = myResp.PA.vect,
                          expl.var = myExpl,
                          nb.rep = 4,
                          nb.absences = 500,
                          strategy = 'disk',
                          dist.min = 5,
                          dist.max = 35)

# SRE method
PA.s <- bm_PseudoAbsences(resp.var = myResp.PA.vect,
                          expl.var = myExpl,
                          nb.rep = 4,
                          nb.absences = 1000,
                          strategy = 'sre',

```

```

sre.quant = 0.025)

# user.defined method
myPatable <- data.frame(PA1 = ifelse(myResp == 1, TRUE, FALSE),
                        PA2 = ifelse(myResp == 1, TRUE, FALSE))
for (i in 1:ncol(myPatable)) myPatable[sample(which(myPatable[, i] == FALSE), 500), i] = TRUE
PA.u <- bm_PseudoAbsences(resp.var = myResp.PA.vect,
                          expl.var = myExpl,
                          strategy = 'user.defined',
                          user.table = myPatable)

str(PA.r)
head(PA.r$pa.tab)
apply(PA.r$pa.tab, 2, table)

head(PA.d$pa.tab)
apply(PA.d$pa.tab, 2, table)

head(PA.s$pa.tab)
apply(PA.s$pa.tab, 2, table)

tail(PA.u$pa.tab)
apply(PA.u$pa.tab, 2, table)

# random method : different number of PA
PA.r_mult <- bm_PseudoAbsences(resp.var = myResp.PA.vect,
                              expl.var = myExpl,
                              nb.rep = 4,
                              nb.absences = c(1000, 500, 500, 200),
                              strategy = 'random')

str(PA.r_mult)
head(PA.r_mult$pa.tab)
apply(PA.r_mult$pa.tab, 2, table)

```

---

bm\_RunModelsLoop

*Loop to compute all single species distribution models*


---

## Description

This internal **biomod2** function allows the user to compute all single species distribution models (asked by the [BIOMOD\\_Modeling](#) function).

## Usage

```

bm_RunModelsLoop(
  bm.format,

```

```

    weights,
    calib.lines,
    modeling.id,
    models,
    models.pa,
    bm.options,
    metric.eval,
    var.import,
    scale.models = TRUE,
    nb.cpu = 1,
    seed.val = NULL,
    do.progress = TRUE
)

bm_RunModel(
  model,
  run.name,
  dir.name = ".",
  modeling.id = "",
  bm.options,
  Data,
  weights.vec,
  calib.lines.vec,
  eval.data = NULL,
  metric.eval = c("ROC", "TSS", "KAPPA"),
  var.import = 0,
  scale.models = TRUE,
  nb.cpu = 1,
  seed.val = NULL,
  do.progress = TRUE
)

```

### Arguments

bm.format	a <a href="#">BIOMOD.formated.data</a> or <a href="#">BIOMOD.formated.data.PA</a> object returned by the <a href="#">BIOMOD_FormatingData</a> function
weights	a matrix containing observation weights for each pseudo-absence (or allData) dataset
calib.lines	a matrix containing calibration / validation lines for each pseudo-absence (or allData) x repetition (or allRun) combination that can be obtained with the <a href="#">bm_CrossValidation</a> function
modeling.id	a character corresponding to the name (ID) of the simulation set ( <i>a random number by default</i> )
models	a vector containing model names to be computed, must be among ANN, CTA, FDA, GAM, GBM, GLM, MARS, MAXENT, MAXNET, RF, RFd, SRE, XGBOOST
models.pa	( <i>optional, default NULL</i> ) A list containing for each model a vector defining which pseudo-absence datasets are to be used, must be among <code>colnames(bm.format@PA.table)</code>

bm.options	a <code>BIOMOD.models.options</code> object returned by the <code>bm_ModelingOptions</code> function
metric.eval	a vector containing evaluation metric names to be used, must be among POD, FAR, POFD, SR, ACCURACY, BIAS, ROC, TSS, KAPPA, OR, ORSS, CSI, ETS, BOYCE, MPA
var.import	<i>(optional, default NULL)</i> An integer corresponding to the number of permutations to be done for each variable to estimate variable importance
scale.models	<i>(optional, default FALSE)</i> A logical value defining whether all models predictions should be scaled with a binomial GLM or not
nb.cpu	<i>(optional, default 1)</i> An integer value corresponding to the number of computing resources to be used to parallelize the single models computation
seed.val	<i>(optional, default NULL)</i> An integer value corresponding to the new seed value to be set
do.progress	<i>(optional, default TRUE)</i> A logical value defining whether the progress bar is to be rendered or not
model	a character corresponding to the model name to be computed, must be either ANN, CTA, FDA, GAM, GBM, GLM, MARS, MAXENT, MAXNET, RF, RFd, SRE, XGBOOST
run.name	a character corresponding to the model to be run (sp.name + pa.id + run.id)
dir.name	<i>(optional, default .)</i> A character corresponding to the modeling folder
Data	a <code>data.frame</code> containing observations, coordinates and environmental variables that can be obtained with the <code>get_species_data</code> function
weights.vec	a vector containing observation weights the concerned pseudo-absence (or <code>allData</code> ) dataset
calib.lines.vec	a vector containing calibration / validation lines for the concerned pseudo-absence (or <code>allData</code> ) x repetition (or <code>allRun</code> ) combination
eval.data	<i>(optional, default NULL)</i> A <code>data.frame</code> containing validation observations, coordinates and environmental variables that can be obtained with the <code>get_eval_data</code> function

## Value

A list containing for each model a list containing the following elements :

- `model` : the name of correctly computed model
- `calib.failure` : the name of incorrectly computed model
- `pred` : the prediction outputs for calibration data
- `pred.eval` : the prediction outputs for evaluation data
- `evaluation` : the evaluation outputs returned by the `bm_FindOptimStat` function
- `var.import` : the mean of variables importance returned by the `bm_VariablesImportance` function

**Author(s)**

Damien Georges

**See Also**

[rpart](#), [prune](#), [gbm](#), [nnet](#), [earth](#), [fda](#), [mars](#), [maxnet](#), [randomForest](#), [xgboost](#), [bm\\_ModelingOptions](#), [BIOMOD\\_Modeling](#), [bm\\_MakeFormula](#), [bm\\_SampleFactorLevels](#), [bm\\_FindOptimStat](#), [bm\\_VariablesImportance](#)

Other Secondary functions: [bm\\_BinaryTransformation\(\)](#), [bm\\_CrossValidation\(\)](#), [bm\\_FindOptimStat\(\)](#), [bm\\_MakeFormula\(\)](#), [bm\\_ModelingOptions\(\)](#), [bm\\_PlotEvalBoxplot\(\)](#), [bm\\_PlotEvalMean\(\)](#), [bm\\_PlotRangeSize\(\)](#), [bm\\_PlotResponseCurves\(\)](#), [bm\\_PlotVarImpBoxplot\(\)](#), [bm\\_PseudoAbsences\(\)](#), [bm\\_SRE\(\)](#), [bm\\_SampleBinaryVector\(\)](#), [bm\\_SampleFactorLevels\(\)](#), [bm\\_Tuning\(\)](#), [bm\\_VariablesImportance\(\)](#)

---

bm\_SampleBinaryVector *Sample binary vector*

---

**Description**

This internal **biomod2** function allows the user to sample a binary vector keeping the same proportion of 0 and 1 as the initial vector.

**Usage**

```
bm_SampleBinaryVector(obs, ratio, as.logical = FALSE, seedval = NULL)
```

**Arguments**

obs	a vector containing binary values (either 0 or 1)
ratio	a numeric between 0 and 1 corresponding to the proportion of obs values to sample
as.logical	<i>(optional, default FALSE)</i> A logical value defining whether output should be returned as a vector of TRUE/FALSE values or integer values corresponding to the indices of obs elements to be kept
seedval	<i>(optional, default NULL)</i> An integer value corresponding to the new seed value to be set

**Value**

A list containing the following elements :

- calibration : elements selected for calibration
- validation : elements selected for validation (complementary to the calibration set)

**Author(s)**

Damien Georges

**See Also**

Other Secondary functions: [bm\\_BinaryTransformation\(\)](#), [bm\\_CrossValidation\(\)](#), [bm\\_FindOptimStat\(\)](#), [bm\\_MakeFormula\(\)](#), [bm\\_ModelingOptions\(\)](#), [bm\\_PlotEvalBoxplot\(\)](#), [bm\\_PlotEvalMean\(\)](#), [bm\\_PlotRangeSize\(\)](#), [bm\\_PlotResponseCurves\(\)](#), [bm\\_PlotVarImpBoxplot\(\)](#), [bm\\_PseudoAbsences\(\)](#), [bm\\_RunModelsLoop\(\)](#), [bm\\_SRE\(\)](#), [bm\\_SampleFactorLevels\(\)](#), [bm\\_Tuning\(\)](#), [bm\\_VariablesImportance\(\)](#)

**Examples**

```
## Generate a binary vector
vec.a <- sample(c(0, 1), 100, replace = TRUE)

## Generate calibration / validation datasets
bm_SampleBinaryVector(obs = vec.a, ratio = 0.7)
```

---

bm\_SampleFactorLevels *Sample all levels of a factorial variable*

---

**Description**

This internal **biomod2** function allows the user to sample all levels of all the factorial variables contained in a `data.frame` or [SpatRaster](#) object.

**Usage**

```
bm_SampleFactorLevels(expl.var, mask.out = NULL, mask.in = NULL)
```

**Arguments**

expl.var	a <code>data.frame</code> or <a href="#">SpatRaster</a> object containing the explanatory variables (in columns or layers)
mask.out	a <code>data.frame</code> or <a href="#">SpatRaster</a> object containing the area that has already been sampled ( <i>factor levels within this mask will not be sampled</i> )
mask.in	a <code>data.frame</code> or <a href="#">SpatRaster</a> object containing areas where factor levels are to be sampled in priority. <i>Note that if after having explored these masks, some factor levels remain unsampled, they will be sampled in the reference input object expl.var.</i>

**Details**

The `expl.var`, `mask.out` and `mask.in` parameters must be coherent in terms of dimensions :

- same number of rows for `data.frame` objects
- same resolution, projection system and number of cells for [SpatRaster](#) objects

If `mask.in` contains several columns (`data.frame`) or layers (`SpatRaster`), then their order matters : they will be considered successively to sample missing factor levels.

- Values in `data.frame` will be understood as :
  - FALSE : out of mask
  - TRUE : in mask
- Values in `SpatRaster` will be understood as :
  - NA : out of mask
  - not NA : in mask

### Value

A vector of numeric values corresponding to either row (`data.frame`) or cell (`SpatRaster`) numbers, each referring to a single level of a single factorial variable.

In case no factorial variable is found in the input object, NULL is returned.

### Author(s)

Damien Georges

### See Also

[bm\\_PseudoAbsences](#), [bm\\_CrossValidation](#)

Other Secondary functions: [bm\\_BinaryTransformation\(\)](#), [bm\\_CrossValidation\(\)](#), [bm\\_FindOptimStat\(\)](#), [bm\\_MakeFormula\(\)](#), [bm\\_ModelingOptions\(\)](#), [bm\\_PlotEvalBoxplot\(\)](#), [bm\\_PlotEvalMean\(\)](#), [bm\\_PlotRangeSize\(\)](#), [bm\\_PlotResponseCurves\(\)](#), [bm\\_PlotVarImpBoxplot\(\)](#), [bm\\_PseudoAbsences\(\)](#), [bm\\_RunModelsLoop\(\)](#), [bm\\_SRE\(\)](#), [bm\\_SampleBinaryVector\(\)](#), [bm\\_Tuning\(\)](#), [bm\\_VariablesImportance\(\)](#)

### Examples

```
library(terra)

## Create raster data
ras.1 <- ras.2 <- mask.out <- rast(nrows = 10, ncols = 10)
ras.1[] <- as.factor(rep(c(1, 2, 3, 4, 5), each = 20))
ras.1 <- as.factor(ras.1)
ras.2[] <- rnorm(100)
stk <- c(ras.1, ras.2)
names(stk) <- c("varFact", "varNorm")

## define a mask for already sampled points
mask.out[1:40] <- 1

## define a list of masks where we want to sample in priority
mask.in <- list(ras.1, ras.1)
mask.in[[1]][1:80] <- NA ## only level 5 should be sampled in this mask
mask.in[[1]][21:80] <- NA ## only levels 1 and 5 should be sampled in this mask
```

```
## Sample all factor levels
samp1 <- bm_SampleFactorLevels(expl.var = stk, mask.out = mask.out)
samp2 <- bm_SampleFactorLevels(expl.var = stk, mask.in = mask.in)
samp3 <- bm_SampleFactorLevels(expl.var = stk, mask.out = mask.out, mask.in = mask.in)
```

---

 bm\_SRE
 

---

*Surface Range Envelope*


---

### Description

This internal **biomod2** function allows the user to run a rectilinear surface range envelop (SRE) (equivalent to **BIOCLIM**) using the extreme percentiles (as recommended by Nix or Busby, see References and Details).

### Usage

```
bm_SRE(
  resp.var = NULL,
  expl.var = NULL,
  new.env = NULL,
  quant = 0.025,
  do.extrem = FALSE
)
```

### Arguments

- |          |   |
|----------|---|
| resp.var | a vector, a <a href="#">SpatVector</a> without associated data ( <i>if presence-only</i> ), or a <a href="#">SpatVector</a> object containing binary data (0 : absence, 1 : presence, NA : indeterminate) for a single species that will be used to build the species distribution model(s)<br><i>Note that old format from <b>sp</b> are still supported such as SpatialPoints (if presence-only) or SpatialPointsDataFrame object containing binary data.</i> |
| expl.var | a matrix, data.frame, <a href="#">SpatVector</a> or <a href="#">SpatRaster</a> object containing the explanatory variables (in columns or layers) that will be used to build the SRE model<br><i>Note that old format from <b>raster</b> and <b>sp</b> are still supported such as RasterStack and SpatialPointsDataFrame objects.</i>  |
| new.env  | a matrix, data.frame, <a href="#">SpatVector</a> or <a href="#">SpatRaster</a> object containing the explanatory variables (in columns or layers) that will be used to predict the SRE model<br><i>Note that old format from <b>raster</b> and <b>sp</b> are still supported such as RasterStack and SpatialPointsDataFrame objects.</i>  |
| quant    | a numeric between 0 and 0.5 defining the half-quantile corresponding to the most extreme value for each variable not to be taken into account for determining the tolerance boundaries of the considered species (see Details)  |



do.extrem      (*optional, default FALSE*)  
 A logical value defining whether a matrix containing extreme conditions supported should be returned or not

### Details

*Please refer to References to get more information about surface range envelop models.*

This method is highly influenced by the extremes of the data input. Whereas a linear model can discriminate the extreme values from the main tendency, the SRE considers them as important as any other data point leading to changes in predictions.

*The more (non-collinear) variables, the more restrictive the model will be.*

Predictions are returned as binary (0 or 1) values, a site being either potentially suitable for all the variables, or out of bounds for at least one variable and therefore considered unsuitable.

quant determines the threshold from which the data will be taken into account for calibration. The default value of 0.05 induces that the 5% most extreme values will be avoided for each variable on each side of its distribution along the gradient, meaning that a total of 10% of the data will not be considered.

### Value

A vector or a [SpatRaster](#) object, containing binary (0 or 1) values.

### Author(s)

Wilfried Thuiller, Bruno Lafourcade, Damien Georges

### References

- Nix, H.A., 1986. A biogeographic analysis of Australian elapid snakes. In: *Atlas of Elapid Snakes of Australia*. (Ed.) R. Longmore, pp. 4-15. **Australian Flora and Fauna Series Number 7**. Australian Government Publishing Service: Canberra.
- Busby, Jeremy. BIOCLIM - a bioclimate analysis and prediction system. *Plant protection quarterly* **6** (1991): 8-9.

### See Also

[bm\\_PseudoAbsences](#), [BIOMOD\\_FormattingData](#), [bm\\_ModelingOptions](#), [bm\\_Tuning](#), [bm\\_RunModelsLoop](#), [BIOMOD\\_Modeling](#),

Other Secondary functions: [bm\\_BinaryTransformation\(\)](#), [bm\\_CrossValidation\(\)](#), [bm\\_FindOptimStat\(\)](#), [bm\\_MakeFormula\(\)](#), [bm\\_ModelingOptions\(\)](#), [bm\\_PlotEvalBoxplot\(\)](#), [bm\\_PlotEvalMean\(\)](#), [bm\\_PlotRangeSize\(\)](#), [bm\\_PlotResponseCurves\(\)](#), [bm\\_PlotVarImpBoxplot\(\)](#), [bm\\_PseudoAbsences\(\)](#), [bm\\_RunModelsLoop\(\)](#), [bm\\_SampleBinaryVector\(\)](#), [bm\\_SampleFactorLevels\(\)](#), [bm\\_Tuning\(\)](#), [bm\\_VariablesImportance\(\)](#)

**Examples**

```

library(terra)
## Load real data
data(DataSpecies)
myResp.r <- as.numeric(DataSpecies[, 'GuloGulo'])

data(bioclim_current)
myExpl.r <- rast(bioclim_current)

myRespXY <- DataSpecies[which(myResp.r == 1), c('X_WGS84', 'Y_WGS84')]
myResp.v <- classify(subset(myExpl.r, 1),
                    matrix(c(-Inf, Inf, 0), ncol = 3, byrow = TRUE))
myResp.v[cellFromXY(myResp.v, myRespXY)] <- 1

## Compute SRE for several quantile values
sre.100 <- bm_SRE(resp.var = myResp.v,
                 expl.var = myExpl.r,
                 new.env = myExpl.r,
                 quant = 0)
sre.095 <- bm_SRE(resp.var = myResp.v,
                 expl.var = myExpl.r,
                 new.env = myExpl.r,
                 quant = 0.025)
sre.090 <- bm_SRE(resp.var = myResp.v,
                 expl.var = myExpl.r,
                 new.env = myExpl.r,
                 quant = 0.05)

## Visualize results
res <- c(myResp.v, sre.100, sre.095, sre.090)
names(res) <- c("Original distribution", "Full data calibration"
              , "Over 95 percent", "Over 90 percent")
plot(res)

```

---

 bm\_Tuning

*Tune models parameters*


---

**Description**

This internal **biomod2** function allows to tune single model parameters and select more efficient ones based on an evaluation metric.

**Usage**

```

bm_Tuning(
  model,
  tuning.fun,

```

```

do.formula = FALSE,
do.stepAIC = FALSE,
bm.options,
bm.format,
calib.lines = NULL,
metric.eval = "TSS",
metric.AIC = "AIC",
weights = NULL,
ctrl.train = NULL,
params.train = list(ANN.size = c(2, 4, 6, 8), ANN.decay = c(0.01, 0.05, 0.1), ANN.bag =
FALSE, FDA.degree = 1:2, FDA.nprune = 2:25, GAM.select = c(TRUE, FALSE), GAM.method =
c("GCV.Cp", "GACV.Cp", "REML", "P-REML", "ML", "P-ML"), GAM.span = c(0.3, 0.5, 0.7),
GAM.degree = 1, GBM.n.trees = c(500, 1000, 2500), GBM.interaction.depth = seq(2, 8,
by = 3), GBM.shrinkage = c(0.001, 0.01, 0.1), GBM.n.minobsinnode = 10, MARS.degree =
1:2, MARS.nprune = 2:max(21, 2 * ncol(bm.format@data.env.var) + 1), MAXENT.algorithm
= "maxnet",
  MAXENT.parallel = TRUE, MAXENT.tune.args = list(rm = seq(0.5, 1,
0.5), fc = c("L")), MAXENT.partitions = "randomkfold", MAXENT.kfolds = 10,
MAXENT.user.grp = NULL, RF.mtry = 1:min(10, ncol(bm.format@data.env.var)), RFd.mtry =
1:min(10, ncol(bm.format@data.env.var)), SRE.quant = c(0, 0.0125, 0.025, 0.05, 0.1),
XGBOOST.nrounds = 50, XGBOOST.max_depth = 1, XGBOOST.eta = c(0.3, 0.4), XGBOOST.gamma
= 0, XGBOOST.colsample_bytree = c(0.6, 0.8), XGBOOST.min_child_weight = 1,
XGBOOST.subsample = 0.5)
)

```

## Arguments

model	a character corresponding to the algorithm to be tuned, must be either ANN, CTA, FDA, GAM, GBM, GLM, MARS, MAXENT, MAXNET, RF, RFd, SRE, XGBOOST
tuning.fun	a character corresponding to the model function name to be called through <a href="#">train</a> function for tuning parameters (see <a href="#">ModelsTable</a> dataset)
do.formula	<i>(optional, default FALSE)</i> A logical value defining whether formula is to be optimized or not
do.stepAIC	<i>(optional, default FALSE)</i> A logical value defining whether variables selection is to be performed for GLM and GAM models or not
bm.options	a <a href="#">BIOMOD.options.default</a> or <a href="#">BIOMOD.options.dataset</a> object returned by the <a href="#">bm_ModelingOptions</a> function
bm.format	a <a href="#">BIOMOD.formated.data</a> or <a href="#">BIOMOD.formated.data.PA</a> object returned by the <a href="#">BIOMOD_FormatingData</a> function
calib.lines	<i>(optional, default NULL)</i> A data.frame object returned by <a href="#">get_calib_lines</a> or <a href="#">bm_CrossValidation</a> functions
metric.eval	a character corresponding to the evaluation metric to be used, must be either AUC, Kappa or TSS for SRE only ; auc.val.avg, auc.diff.avg, or.mtp.avg, or.10p.avg, AICc for MAXENT only ; ROC or TSS for all other models

<code>metric.AIC</code>	a character corresponding to the AIC metric to be used, must be either AIC or BIC
<code>weights</code>	( <i>optional, default NULL</i> ) A vector of numeric values corresponding to observation weights (one per observation, see Details)
<code>ctrl.train</code>	( <i>optional, default NULL</i> ) A <a href="#">trainControl</a> object
<code>params.train</code>	a list containing values of model parameters to be tested (see Details)

### Details

#### Concerning `ctrl.train` parameter :

Set by default to :

```
ctrl.train <- caret::trainControl(method = "repeatedcv", repeats = 3, number = 10,
summaryFunction = caret::twoClassSummary,
classProbs = TRUE, returnData = FALSE)
```

#### Concerning `params.train` parameter :

All elements of the list must have names matching `model.parameter_name` format, `parameter_name` being one of the parameter of the `tuning.fun` function called by `caret` package and that can be found through the [getModelInfo](#) function.

Currently, the available parameters to be tuned are the following :

**ANN** size, decay, bag

**CTA** maxdepth

**FDA** degree, nprune

**GAM.gam** span, degree

**GAM.mgcv** select, method

**GBM** n.trees, interaction.depth, shrinkage, n.minobsinnode

**MARS** degree, nprune

**MAXENT** algorithm, tune.args, parallel, partitions, kfolds, user.grp

**RF** mtry

**RFd** mtry

**SRE** quant

**XGBOOST** nrounds, max\_depth, eta, gamma, colsampl\_bytree, min\_child\_weight, subsample

The [expand.grid](#) function is used to build a matrix containing all combinations of parameters to be tested.

### Value

A [BIOMOD.models.options](#) object (see [bm\\_ModelingOptions](#)) with optimized parameters

**Note**

- No tuning for GLM and MAXNET
- MAXENT is tuned through [ENMevaluate](#) function which is calling either :
  - maxnet (by defining `MAXENT.algorithm = 'maxnet'`) (*default*)
  - Java version of Maxent defined in **dismo** package (by defining `MAXENT.algorithm = 'maxent.jar'`)
- SRE is tuned through [bm\\_SRE](#) function
- All other models are tuned through [train](#) function
- No optimization of formula for MAXENT, MAXNET, SRE and XGBOOST
- No interaction included in formula for CTA
- Variables selection only for `GAM.gam` and `GLM`

**Author(s)**

Frank Breiner, Maya Gueguen, Helene Blancheteau

**See Also**

[trainControl](#), [train](#), [ENMevaluate](#), [ModelsTable](#), [BIOMOD.models.options](#), [bm\\_ModelingOptions](#), [BIOMOD\\_Modeling](#)

Other Secondary functions: [bm\\_BinaryTransformation\(\)](#), [bm\\_CrossValidation\(\)](#), [bm\\_FindOptimStat\(\)](#), [bm\\_MakeFormula\(\)](#), [bm\\_ModelingOptions\(\)](#), [bm\\_PlotEvalBoxplot\(\)](#), [bm\\_PlotEvalMean\(\)](#), [bm\\_PlotRangeSize\(\)](#), [bm\\_PlotResponseCurves\(\)](#), [bm\\_PlotVarImpBoxplot\(\)](#), [bm\\_PseudoAbsences\(\)](#), [bm\\_RunModelsLoop\(\)](#), [bm\\_SRE\(\)](#), [bm\\_SampleBinaryVector\(\)](#), [bm\\_SampleFactorLevels\(\)](#), [bm\\_VariablesImportance\(\)](#)

**Examples**

```
library(terra)

# Load species occurrences (6 species available)
data(DataSpecies)
head(DataSpecies)

# Select the name of the studied species
myRespName <- 'GuloGulo'

# Get corresponding presence/absence data
myResp <- as.numeric(DataSpecies[, myRespName])

# Get corresponding XY coordinates
myRespXY <- DataSpecies[, c('X_WGS84', 'Y_WGS84')]

# Load environmental variables extracted from BIOCLIM (bio_3, bio_4, bio_7, bio_11 & bio_12)
data(bioclim_current)
myExpl <- terra::rast(bioclim_current)
```

```

# ----- #
# Format Data with true absences
myBiomodData <- BIOMOD_FormatingData(resp.var = myResp,
                                   expl.var = myExpl,
                                   resp.xy = myRespXY,
                                   resp.name = myRespName)

# ----- #
# List of all models currently available in `biomod2` (and their related package and function)
# Some of them can be tuned through the `train` function of the `caret` package
# (and corresponding training function to be used is indicated)
data(ModelsTable)
ModelsTable

allModels <- c('ANN', 'CTA', 'FDA', 'GAM', 'GBM', 'GLM'
               , 'MARS', 'MAXENT', 'MAXNET', 'RF', 'SRE', 'XGBOOST')

# default parameters
opt.d <- bm_ModelingOptions(data.type = 'binary',
                           models = allModels,
                           strategy = 'default')

# tune parameters for Random Forest model
tuned.rf <- bm_Tuning(model = 'RF',
                    tuning.fun = 'rf', ## see in ModelsTable
                    do.formula = FALSE,
                    bm.options = opt.d@options$RF.binary.randomForest.randomForest,
                    bm.format = myBiomodData)

tuned.rf

## Not run:
# tune parameters for GAM (from mgcv package) model
tuned.gam <- bm_Tuning(model = 'GAM',
                    tuning.fun = 'gam', ## see in ModelsTable
                    do.formula = TRUE,
                    do.stepAIC = TRUE,
                    bm.options = opt.d@options$GAM.binary.mgcv.gam,
                    bm.format = myBiomodData)

tuned.gam

## End(Not run)

```

## Description

This internal **biomod2** function allows the user to compute a variable importance value for each variable involved in the given model.

## Usage

```
bm_VariablesImportance(
  bm.model,
  expl.var,
  variables = NULL,
  method = "full_rand",
  nb.rep = 1,
  seed.val = NULL,
  do.progress = TRUE,
  temp.workdir = NULL
)
```

## Arguments

bm.model	a biomod2_model object (or nnet, rpart, fda, gam, glm, lm, gbm, mars, randomForest, xgb.Booster) that can be obtained with the <a href="#">get_formal_model</a> function
expl.var	a data.frame containing the explanatory variables that will be used to compute the variables importance
variables	(optional, default NULL) A vector containing the names of the explanatory variables that will be considered
method	a character corresponding to the randomization method to be used, must be full_rand (only method available so far)
nb.rep	an integer corresponding to the number of permutations to be done for each variable
seed.val	(optional, default NULL) An integer value corresponding to the new seed value to be set
do.progress	(optional, default TRUE) A logical value defining whether the progress bar is to be rendered or not
temp.workdir	(optional, default NULL) A character value corresponding to the folder name containing temporal prediction files when using MAXENT

## Details

For each variable to be evaluated :

1. shuffle the original variable
2. compute model prediction with shuffled variable
3. calculate Pearson's correlation between reference and shuffled predictions
4. return score as  $1 - cor$

The highest the value, the less reference and shuffled predictions are correlated, and the more influence the variable has on the model. A value of 0 assumes no influence of the variable on the model.

*Note that this calculation does not account for variables' interactions.*

The same principle is used in [randomForest](#).

## Value

A 3 columns data.frame containing variable's importance scores for each permutation run :

- `expl.var` : the considered explanatory variable (the one permuted)
- `rand` : the ID of the permutation run
- `var.imp` : the variable's importance score

## Author(s)

Damien Georges

## See Also

[randomForest](#), [bm\\_RunModelsLoop](#), [BIOMOD\\_Modeling](#), [BIOMOD\\_EnsembleModeling](#), [bm\\_PlotVarImpBoxplot](#), [get\\_variables\\_importance](#)

Other Secondary functions: [bm\\_BinaryTransformation\(\)](#), [bm\\_CrossValidation\(\)](#), [bm\\_FindOptimStat\(\)](#), [bm\\_MakeFormula\(\)](#), [bm\\_ModelingOptions\(\)](#), [bm\\_PlotEvalBoxplot\(\)](#), [bm\\_PlotEvalMean\(\)](#), [bm\\_PlotRangeSize\(\)](#), [bm\\_PlotResponseCurves\(\)](#), [bm\\_PlotVarImpBoxplot\(\)](#), [bm\\_PseudoAbsences\(\)](#), [bm\\_RunModelsLoop\(\)](#), [bm\\_SRE\(\)](#), [bm\\_SampleBinaryVector\(\)](#), [bm\\_SampleFactorLevels\(\)](#), [bm\\_Tuning\(\)](#)

## Examples

```
## Create simple simulated data
myResp.s <- sample(c(0, 1), 20, replace = TRUE)
myExpl.s <- data.frame(var1 = sample(c(0, 1), 100, replace = TRUE),
                      var2 = rnorm(100),
                      var3 = 1:100)

## Compute variables importance
mod <- glm(var1 ~ var2 + var3, data = myExpl.s)
bm_VariablesImportance(bm.model = mod,
                      expl.var = myExpl.s[, c('var2', 'var3')],
                      method = "full_rand",
                      nb.rep = 3)
```



---

DataSpecies	<i>Presence-Absence data to build test SDM</i>
-------------	--

---

**Description**

A dataset covering all the continent with presence/absence data for 6 mammal species. Presence/absence were derived from range maps downloaded at [IUCN](#).

**Usage**

```
DataSpecies
```

**Format**

A data.frame object with 2488 rows and 10 variables:

**X\_WGS84** Longitude

**Y\_WGS84** Latitude

**ConnochaetesGnou** Presence (1) or Absence (0) for black wildebeest

**GuloGulo** Presence (1) or Absence (0) for wolverine

**PantheraOnca** Presence (1) or Absence (0) for jaguar

**PteropusGiganteus** Presence (1) or Absence (0) for Indian flying fox

**TenrecEcaudatus** Presence (1) or Absence (0) for tailless tenrec

**VulpesVulpes** Presence (1) or Absence (0) for red fox

---

getters.bm	<i>Functions to extract informations from <a href="#">biomod2_model</a> objects</i>
------------	---

---

**Description**

These functions allow the user to easily retrieve single models (formal or scaled) from [biomod2\\_model](#) objects from the modeling step.

**Usage**

```
## S4 method for signature 'biomod2_model'
get_formal_model(object)
```

```
## S4 method for signature 'biomod2_model'
get_scaling_model(object)
```

**Arguments**

object            a [biomod2\\_model](#) object

**Value**

`get_formal_model` an object from the `model` slot of a `biomod2_model` object

`get_scaling_model` an object from the `scaling_model` slot of a `biomod2_model` object

**Author(s)**

Damien Georges

**See Also**

[biomod2\\_model](#)

Other Toolbox functions: [getters.out](#), [load\\_stored\\_object\(\)](#), [predict.bm](#), [predict.em](#), [predict2.bm](#), [predict2.em](#)

---

<code>getters.out</code>	<i>Functions to extract informations from <code>BIOMOD.models.out</code>, <code>BIOMOD.projection.out</code> or <code>BIOMOD.ensemble.models.out</code> objects</i>
--------------------------	---

---

**Description**

These functions allow the user to easily retrieve informations stored in the different **biomod2** objects from the different modeling steps, such as modeling options and formatted data, models used or not, predictions, evaluations, variables importance.

**Usage**

```
## S4 method for signature 'BIOMOD.formated.data'
get_species_data(obj)

## S4 method for signature 'BIOMOD.formated.data.PA'
get_species_data(obj)

## S4 method for signature 'BIOMOD.formated.data'
get_eval_data(obj)

## S4 method for signature 'BIOMOD.models.out'
get_options(obj)

## S4 method for signature 'BIOMOD.models.out'
get_calib_lines(obj, as.data.frame = FALSE, PA = NULL, run = NULL)

## S4 method for signature 'BIOMOD.models.out'
get_formal_data(obj, subinfo = NULL)

## S4 method for signature 'BIOMOD.models.out'
get_predictions(
```

```
    obj,  
    evaluation = FALSE,  
    full.name = NULL,  
    PA = NULL,  
    run = NULL,  
    algo = NULL,  
    model.as.col = FALSE  
  )  
  
## S4 method for signature 'BIOMOD.models.out'  
get_built_models(obj, full.name = NULL, PA = NULL, run = NULL, algo = NULL)  
  
## S4 method for signature 'BIOMOD.models.out'  
get_evaluations(  
  obj,  
  full.name = NULL,  
  PA = NULL,  
  run = NULL,  
  algo = NULL,  
  metric.eval = NULL  
)  
  
## S4 method for signature 'BIOMOD.models.out'  
get_variables_importance(  
  obj,  
  full.name = NULL,  
  PA = NULL,  
  run = NULL,  
  algo = NULL,  
  expl.var = NULL  
)  
  
## S4 method for signature 'BIOMOD.projection.out'  
get_projected_models(  
  obj,  
  full.name = NULL,  
  PA = NULL,  
  run = NULL,  
  algo = NULL,  
  merged.by.algo = NULL,  
  merged.by.run = NULL,  
  merged.by.PA = NULL,  
  filtered.by = NULL  
)  
  
## S4 method for signature 'BIOMOD.projection.out'  
free(obj)
```

```
## S4 method for signature 'BIOMOD.projection.out'
get_predictions(
  obj,
  metric.binary = NULL,
  metric.filter = NULL,
  full.name = NULL,
  PA = NULL,
  run = NULL,
  algo = NULL,
  merged.by.algo = NULL,
  merged.by.run = NULL,
  merged.by.PA = NULL,
  filtered.by = NULL,
  model.as.col = FALSE,
  ...
)

## S4 method for signature 'BIOMOD.ensemble.models.out'
get_formal_data(obj, subinfo = NULL)

## S4 method for signature 'BIOMOD.ensemble.models.out'
get_built_models(
  obj,
  full.name = NULL,
  merged.by.algo = NULL,
  merged.by.run = NULL,
  merged.by.PA = NULL,
  filtered.by = NULL,
  algo = NULL
)

## S4 method for signature 'BIOMOD.ensemble.models.out'
get_kept_models(obj)

## S4 method for signature 'BIOMOD.ensemble.models.out'
get_predictions(
  obj,
  evaluation = FALSE,
  full.name = NULL,
  merged.by.algo = NULL,
  merged.by.run = NULL,
  merged.by.PA = NULL,
  filtered.by = NULL,
  algo = NULL,
  model.as.col = FALSE
)

## S4 method for signature 'BIOMOD.ensemble.models.out'
```

```

get_evaluations(
  obj,
  full.name = NULL,
  merged.by.algo = NULL,
  merged.by.run = NULL,
  merged.by.PA = NULL,
  filtered.by = NULL,
  algo = NULL,
  metric.eval = NULL
)

## S4 method for signature 'BIOMOD.ensemble.models.out'
get_variables_importance(
  obj,
  full.name = NULL,
  merged.by.algo = NULL,
  merged.by.run = NULL,
  merged.by.PA = NULL,
  filtered.by = NULL,
  algo = NULL,
  expl.var = NULL
)

```

### Arguments

obj	a <a href="#">BIOMOD.formated.data</a> , <a href="#">BIOMOD.formated.data.PA</a> , <a href="#">BIOMOD.models.out</a> , <a href="#">BIOMOD.projection.out</a> or <a href="#">BIOMOD.ensemble.models.out</a> object
as.data.frame	a logical defining whether output should be returned as data.frame or array object
PA	<i>(optional, default NULL)</i> A vector containing pseudo-absence set to be loaded, must be among PA1, PA2, ..., allData
run	<i>(optional, default NULL)</i> A vector containing repetition set to be loaded, must be among RUN1, RUN2, ..., allRun
subinfo	a character corresponding to the information to be extracted, must be among NULL, expl.var.names, resp.var, expl.var, MinMax, eval.resp.var, eval.expl.var (see Details)
evaluation	a logical defining whether evaluation data should be used or not
full.name	<i>(optional, default NULL)</i> A vector containing model names to be kept, must be either all or a sub-selection of model names that can be obtained with the <a href="#">get_built_models</a> function
algo	<i>(optional, default NULL)</i> A character containing algorithm to be loaded, must be either ANN, CTA, FDA, GAM, GBM, GLM, MARS, MAXENT, MAXNET, RF, SRE, XGBOOST

<code>model.as.col</code>	<i>(optional, default FALSE)</i> A boolean given to <code>get_predictions</code> . If TRUE prediction are returned as a wide data.frame with each column containing predictions for a single model and corresponding to the old output given by <b>biomod2</b> in version < 4.2-2. If FALSE predictions are returned as a long data.frame with many additional informations readily available.
<code>metric.eval</code>	<i>(optional, default NULL)</i> A vector containing evaluation metric to be kept, must be among POD, FAR, POFD, SR, ACCURACY, BIAS, ROC, TSS, KAPPA, OR, ORSS, CSI, ETS, BOYCE, MPA
<code>expl.var</code>	<i>(optional, default NULL)</i> A vector containing explanatory variables to be kept, that can be obtained with the <code>get_formal_data(obj, subinfo = 'expl.var.names')</code> function
<code>merged.by.algo</code>	<i>(optional, default NULL)</i> A character containing merged algorithm to be loaded, must be among ANN, CTA, FDA, GAM, GBM, GLM, MARS, MAXENT, MAXNET, RF, SRE, XGBOOST, mergedAlgo
<code>merged.by.run</code>	<i>(optional, default NULL)</i> A vector containing merged repetition set to be loaded, must be among RUN1, RUN2, ..., mergedRun
<code>merged.by.PA</code>	<i>(optional, default NULL)</i> A vector containing merged pseudo-absence set to be loaded, must be among PA1, PA2, ..., mergedData
<code>filtered.by</code>	<i>(optional, default NULL)</i> A vector containing evaluation metric selected to filter single models to build the ensemble models, must be among POD, FAR, POFD, SR, ACCURACY, BIAS, ROC, TSS, KAPPA, OR, ORSS, CSI, ETS, BOYCE, MPA
<code>metric.binary</code>	<i>(optional, default NULL)</i> A vector containing evaluation metric selected to transform predictions into binary values, must be among POD, FAR, POFD, SR, ACCURACY, BIAS, ROC, TSS, KAPPA, OR, ORSS, CSI, ETS, BOYCE, MPA
<code>metric.filter</code>	<i>(optional, default NULL)</i> A vector containing evaluation metric to filter predictions, must be among POD, FAR, POFD, SR, ACCURACY, BIAS, ROC, TSS, KAPPA, OR, ORSS, CSI, ETS, BOYCE, MPA
<code>...</code>	<i>(optional, one or several of the following arguments depending on the selected function)</i>

## Value

<code>get_species_data</code>	a data.frame combining <code>data.species</code> , <code>coord</code> , <code>data.env.var</code> (and <code>PA.table</code> ) slots of <code>BIOMOD.formated.data</code> (or <code>BIOMOD.formated.data.PA</code> ) object
<code>get_eval_data</code>	a data.frame combining <code>eval.data.species</code> , <code>eval.coord</code> , <code>eval.data.env.var</code> slots of <code>BIOMOD.formated.data</code> or <code>BIOMOD.formated.data.PA</code> object
<code>get_options</code>	a <code>BIOMOD.stored.options-class</code> object from the <code>models.options</code> slot of a <code>BIOMOD.models.out-class</code> object
<code>get_calib_lines</code>	a <code>BIOMOD.stored.data.frame-class</code> object from the <code>calib.lines</code> slot of a <code>BIOMOD.models.out</code> object

`get_projected_models` a vector from the `models.projected` slot of a `BIOMOD.projection.out` object

`get_predictions` a `BIOMOD.stored.data` object from the `proj.out` slot of a `BIOMOD.models.out`, `BIOMOD.projection.out` or `BIOMOD.ensemble.models.out` object

`get_kept_models` a vector containing names of the kept models of a `BIOMOD.ensemble.models.out` object

`get_formal_data` depending on the `subinfo` parameter :

- NULL a `BIOMOD.stored.formated.data-class` (or `BIOMOD.stored.models.out-class`) object from the `formated.input.data` (or `models.out`) slot of a `BIOMOD.models.out` (or `BIOMOD.ensemble.models.out`) object
- `expl.var.names` a vector from the `expl.var.names` slot of a `BIOMOD.models.out` or `BIOMOD.ensemble.models.out` object
- `resp.var` a vector from the `data.species` slot of the `formated.input.data` slot of a `BIOMOD.models.out` or `BIOMOD.ensemble.models.out` object
- `expl.var` a data.frame from the `data.env.var` slot of the `formated.input.data` slot of a `BIOMOD.models.out` or `BIOMOD.ensemble.models.out` object
- MinMax a list of minimum and maximum values (or levels if factorial) of variable contained in the `data.env.var` slot of the `formated.input.data` slot of a `BIOMOD.models.out` or `BIOMOD.ensemble.models.out` object
- `eval.resp.var` a vector from the `eval.data.species` slot of the `formated.input.data` slot of a `BIOMOD.models.out` or `BIOMOD.ensemble.models.out` object
- `eval.expl.var` a data.frame from the `eval.data.env.var` slot of the `formated.input.data` slot of a `BIOMOD.models.out` or `BIOMOD.ensemble.models.out` object

`get_built_models` a vector from the `models.computed` slot (or `em.computed`) of a `BIOMOD.models.out` (or `BIOMOD.ensemble.models.out`) object

`get_evaluations` a data.frame from the `models.evaluation` slot (or `model_evaluation` of each model in `em.computed`) of a `BIOMOD.models.out` (or `BIOMOD.ensemble.models.out`) object. Contains evaluation metric for different models and dataset. Evaluation metric are calculated on the calibrating data (column calibration), on the cross-validation data (column validation) or on the evaluation data (column evaluation).  
*For cross-validation data, see CV.[...] parameters in BIOMOD\_Modeling function ; for evaluation data, see eval.[...] parameters in BIOMOD\_FormatingData.*

`get_variables_importance` a `BIOMOD.stored.data.frame-class` from the `variables.importance` slot (or `model_variables_importance` of each model in `em.models`) of a `BIOMOD.models.out` (or `BIOMOD.ensemble.models.out`) object

**Author(s)**

Damien Georges

**See Also**

`BIOMOD.models.out`, `BIOMOD.projection.out`, `BIOMOD.ensemble.models.out`

Other Toolbox functions: `getters.bm`, `load_stored_object()`, `predict.bm`, `predict.em`, `predict2.bm`, `predict2.em`

---

load\_stored\_object      *Functions to load [BIOMOD.stored.data](#) objects*

---

### Description

This functions allow the user to load [BIOMOD.stored.data](#) objects into memory.

### Usage

```
load_stored_object(obj, ...)

## S4 method for signature 'BIOMOD.stored.data'
load_stored_object(obj, layer = 1)

## S4 method for signature 'BIOMOD.stored.SpatRaster'
load_stored_object(obj, layer = 1)
```

### Arguments

obj	a <a href="#">BIOMOD.stored.data</a> object
...	additional arguments
layer	an integer corresponding to the layer ID to be extracted when multilayer object considered

### Author(s)

Damien Georges

### See Also

[BIOMOD.stored.data](#)

Other Toolbox functions: [getters.bm](#), [getters.out](#), [predict.bm](#), [predict.em](#), [predict2.bm](#), [predict2.em](#)

---

ModelsTable      *Single models package and functions*

---

### Description

A data.frame containing for each single model available in **biomod2** the package and functions to be called.

### Usage

ModelsTable



**Format**

A data.frame object with 12 rows and 5 variables:

**model** all single models that can be computed in **biomod2**

**type** data type associated to the models

**package** R package used

**func** function used in the R package

**train** function called by **caret** for the tuning

All single models available are the following :

- ANN ([nnet](#))
- CTA ([rpart](#))
- FDA ([fda](#))
- GAM ([gam](#), [gam](#) or [bam](#))
- GBM ([gbm](#))
- GLM ([glm](#))
- MARS ([earth](#))
- MAXENT ([see Maxent website](#))
- MAXNET ([maxnet](#))
- RF ([randomForest](#))
- SRE ([bm\\_SRE](#))
- XGBOOST ([xgboost](#))

---

OptionsBigboss

*Bigboss pre-defined parameter values for single models*

---

**Description**

A `BIOMOD.models.options` object containing for each single model available in **biomod2** the parameter values pre-defined by **biomod2** team.

**Usage**

OptionsBigboss

**Format**

A `BIOMOD.models.options` object with some changed values :

```
ANN.binary.nnet.nnet • size = 5
  • decay = 0.1
  • trace = FALSE
  • rang = 0.1
  • maxit = 200

CTA.binary.rpart.rpart • method = 'class'
  • control = list(xval = 5, minbucket = 5, minsplit = 5, cp = 0.001, maxdepth = 10)
  • cost = NULL

FDA.binary.mda.fda • method = 'mars'

GAM.binary.gam.gam
GAM.binary.mgcv.bam
GAM.binary.mgcv.gam • family = binomial(link = 'logit')
  • method = 'GCV.Cp'
  • control = list(epsilon = 1e-06, trace = FALSE, maxit = 100)

GBM.binary.gbm.gbm • n.trees = 2500
  • interaction.depth = 7
  • n.minobsinnode = 5
  • shrinkage = 0.001
  • cv.folds = 3
  • keep.data = FALSE
  • n.cores = 1

GLM.binary.stats.glm • family = binomial(link = 'logit')
  • mustart = 0.5
  • control = glm.control(maxit = 50)

MARS.binary.earth.earth • glm = list(family = binomial(link = 'logit'))
  • ncross = 0
  • nk = NULL
  • penalty = 2
  • thresh = 0.001
  • nprune = NULL
  • pmethod = 'backward'

MAXENT.binary.MAXENT.MAXENT • path_to_maxent.jar = '.'

RF.binary.randomForest.randomForest • type = 'classification'
  • ntree = 500
  • mtry = 2
  • strata = factor(c(0, 1))
  • sampsize = NULL
  • nodesize = 5
```

```

      • maxnodes = NULL
RFd.binary.randomForest.randomForest • type = 'classification'
      • ntree = 500
      • mtry = 2
      • strata = factor(c(0, 1))
      • sampsize = NULL
      • nodesize = 5
      • maxnodes = NULL
SRE.binary.biomod2.bm_SRE • do.extrem = TRUE
XGBOOST.binary.xgboost.xgboost • params = list(max_depth = 2, eta = 1)
      • nthread = 2
      • nrounds = 4
      • objective = 'binary:logistic'

```

---

```

plot, BIOMOD.formated.data, missing-method
      plot method for BIOMOD.formated.data object class

```

---

## Description

Plot the spatial distribution of presences, absences and pseudo-absences among the different potential dataset (calibration, validation and evaluation). Available only if coordinates were given to [BIOMOD\\_FormatingData](#).

## Usage

```

## S4 method for signature 'BIOMOD.formated.data,missing'
plot(
  x,
  calib.lines = NULL,
  plot.type,
  plot.output,
  PA,
  run,
  plot.eval,
  point.size = 1.5,
  do.plot = TRUE
)

```

## Arguments

x a [BIOMOD.formated.data](#) or [BIOMOD.formated.data.PA](#) object. Coordinates must be available to be able to use plot.

calib.lines	(optional, default NULL) an data.frame object returned by <code>get_calib_lines</code> or <code>bm_CrossValidation</code> functions, to explore the distribution of calibration and validation datasets
plot.type	a character, either 'points' (default) or 'raster' (if environmental variables were given as a raster). With <code>plot.type = 'points'</code> occurrences will be represented as points (better when using fine-grained data). With <code>plot.type = 'raster'</code> occurrences will be represented as a raster (better when using coarse-grained data)
plot.output	a character, either 'facet' (default) or 'list'. <code>plot.output</code> determines whether plots are returned as a single facet with all plots or a list of individual plots (better when there are numerous graphics)
PA	(optional, default 'all') If <code>x</code> is a <code>BIOMOD.formated.data.PA</code> object, a vector containing pseudo-absence set to be represented
run	(optional, default 'all') If <code>calib.lines</code> provided, a vector containing repetition set to be represented
plot.eval	(optional, default TRUE) A logical defining whether evaluation data should be added to the plot or not
point.size	a numeric to adjust the size of points when <code>plot.type = 'points'</code> .
do.plot	(optional, default TRUE) A logical defining whether the plot is to be rendered or not

**Value**

a list with the data used to generate the plot and a ggplot2 object

**Author(s)**

Remi Patin

**Examples**

```
library(terra)

# Load species occurrences (6 species available)
data(DataSpecies)
head(DataSpecies)

# Select the name of the studied species
myRespName <- 'GuloGulo'

# Get corresponding presence/absence data
myResp <- as.numeric(DataSpecies[, myRespName])

# Get corresponding XY coordinates
myRespXY <- DataSpecies[, c('X_WGS84', 'Y_WGS84')]

# Load environmental variables extracted from BIOCLIM (bio_3, bio_4, bio_7, bio_11 & bio_12)
```

```

data(bioclim_current)
myExpl <- terra::rast(bioclim_current)

## ----- #
# Format Data with true absences
myBiomodData <- BIOMOD_FormatingData(resp.var = myResp,
                                   expl.var = myExpl,
                                   resp.xy = myRespXY,
                                   resp.name = myRespName)

myBiomodData
plot(myBiomodData)

```

---

predict.bm

*Functions to get predictions from [biomod2\\_model](#) objects*


---

### Description

This function allows the user to predict single models from [biomod2\\_model](#) on (new) explanatory variables.

### Usage

```
## S4 method for signature 'biomod2_model'
predict(object, newdata, ...)
```

### Arguments

object	a <a href="#">biomod2_model</a> object
newdata	a data.frame or <a href="#">SpatRaster</a> object containing data for new predictions
...	(optional)

### Author(s)

Damien Georges

### See Also

[biomod2\\_model](#)

Other Toolbox functions: [getters.bm](#), [getters.out](#), [load\\_stored\\_object\(\)](#), [predict.em](#), [predict2.bm](#), [predict2.em](#)

---

predict.em *Functions to get predictions from [biomod2\\_ensemble\\_model](#) objects*

---

### Description

This function allows the user to predict single models from [biomod2\\_ensemble\\_model](#) on (new) explanatory variables.

### Arguments

object a [biomod2\\_ensemble\\_model](#) object  
 newdata a data.frame or [SpatRaster](#) object containing data for new predictions  
 ... (*optional*)

### Author(s)

Damien Georges

### See Also

[biomod2\\_ensemble\\_model](#)

Other Toolbox functions: [getters.bm](#), [getters.out](#), [load\\_stored\\_object\(\)](#), [predict.bm](#), [predict2.bm](#), [predict2.em](#)

---

summary, BIOMOD.formated.data-method  
*summary method for [BIOMOD.formated.data](#) object class*

---

### Description

Summarize the number of presences, absences and pseudo-absences among the different potential dataset (calibration, validation and evaluation).

### Usage

```
## S4 method for signature 'BIOMOD.formated.data'
summary(object, calib.lines = NULL)
```

### Arguments

object a [BIOMOD.formated.data](#) or [BIOMOD.formated.data.PA](#) object returned by the [BIOMOD\\_FormatingData](#) function  
 calib.lines (*optional, default* NULL)  
 an array object returned by [get\\_calib\\_lines](#) or [bm\\_CrossValidation](#) functions, to explore the distribution of calibration and validation datasets

**Value**

a data.frame

**Author(s)**

Remi Patin

**Examples**

```
library(terra)

# Load species occurrences (6 species available)
data(DataSpecies)
head(DataSpecies)

# Select the name of the studied species
myRespName <- 'GuloGulo'

# Get corresponding presence/absence data
myResp <- as.numeric(DataSpecies[, myRespName])

# Get corresponding XY coordinates
myRespXY <- DataSpecies[, c('X_WGS84', 'Y_WGS84')]

# Load environmental variables extracted from BIOCLIM (bio_3, bio_4, bio_7, bio_11 & bio_12)
data(bioclim_current)
myExpl <- terra::rast(bioclim_current)

## ----- #
# Format Data with true absences
myBiomodData <- BIOMOD_FormatingData(resp.var = myResp,
                                   expl.var = myExpl,
                                   resp.xy = myRespXY,
                                   resp.name = myRespName)

myBiomodData
summary(myBiomodData)
```

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