Package ‘Hmsc’

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alignPosterior alignPosterior

Description

Aligns posterior in terms of variables susceptible to label switching

Usage

alignPosterior(hM)
Arguments

- `hM`: a fitted Hmsc model object

Value

an Hmsc model object that is identical to the input except the posterior being aligned

Examples

```r
# Align the posterior for a previously fitted HMSC model
m = alignPosterior(TD$m)
```

Description

Constructs an ordination biplot based on the fitted model

Usage

```r
biPlot(
    hM, 
    etaPost, 
    lambdaPost, 
    factors = c(1, 2), 
    colVar = NULL, 
    colors = NULL, 
    spNames = hM$spNames, 
    ... 
)
```

Arguments

- `hM`: a fitted Hmsc model object
- `etaPost`: posterior distribution of site loadings (Eta)
- `lambdaPost`: posterior distribution of species loadings (Lambda)
- `factors`: indices of the two factors to be plotted
- `colVar`: the environmental covariate from XData according to which the sites are to be coloured
- `colors`: controls the colors of the heatmap. For continuous covariates, colors should be given as a name of palette, with default value `colorRampPalette(c("blue","white","red"))`, or as a vector of colours. For factors, colors should be given as a vector of colours, e.g. c("blue","red").
- `spNames`: a vector of species names to be added to the ordination diagram
- `...`: other parameters passed to the function.
Examples

# Construct an ordination biplot using two chosen latent factors from a previously fitted HMSC model
etaPost = getPostEstimate(TD$m, "Eta")
lambdaPost=getPostEstimate(TD$m, "Lambda")
biPlot(TD$m, etaPost = etaPost, lambdaPost=lambdaPost, factors=c(1,2))

---

c.Hmsc

Combine Posterior Samples of Several Hmsc Models

Description

Function combines posterior samples of several sampled Hmsc models (see sampleMcmc) as new chains in the first fitted model. The combined models must be comparable, and there are some tests for detecting non-equal models. These tests will only give warning, and it is at user deliberation to decide which models and which posterior samples can be combined. You should be careful not start two models from the same random number seed, because these will only duplicate your data instead of providing new independent samples.

Usage

## S3 method for class 'Hmsc'
c(...)

Arguments

... Sampled Hmsc models with posterior samples that will be added as new chains in the first listed model.

Value

An Hmsc model with chains of posterior samples.

Examples

## Fit a toy model with two chains
m1 <- sampleMcmc(TD$m, samples=10, transient=5, nChains=2, verbose=0)
## Need more data? Add chains: check carefully that these are sampled exactly like the previous model
m2 <- sampleMcmc(TD$m, nChains=2, samples=10, transient=5, verbose=0)
## Now four chains
m4 <- c(m1, m2)
m4
**computeAssociations**

Description

Computes the species association matrices associated with each random level

Usage

`computeAssociations(hM, start = 1, thin = 1)`

Arguments

- `hM`: a fitted `Hmsc` model object
- `start`: index of first MCMC sample included
- `thin`: thinning interval of posterior distribution

Value

list of association matrices (ω) corresponding to each random level in the model

Examples

```r
# Compute the associations (residual correlations) between species from a HMSC model
assoc = computeAssociations(TD$m)
```

**computeDataParameters**

Description

Computes initial values before the sampling starts

Usage

`computeDataParameters(hM)`

Arguments

- `hM`: a fitted `Hmsc` model object

Value

a list including pre-computed matrix inverses and determinants (for phylogenetic and spatial random effects) needed in MCMC sampling
computeInitialParameters

Description
Computes initial parameter values before the sampling starts

Usage
computeInitialParameters(hM, initPar)

Arguments
- hM: a fitted Hmsc model object
- initPar: a list of initial parameter values

Value
a list of Hmsc model parameters

computePredictedValues

Description
Computes predicted values from the fitted Hmsc model

Usage
computePredictedValues(hM, partition = NULL, partition.sp = NULL, start = 1, thin = 1, Yc = NULL, mcmcStep = 1, expected = TRUE, initPar = NULL, nParallel = 1, nChains = length(hM$postList), updater = list(), verbose = hM$verbose,


```r

pcomputePredictedValues(
  hM,
  partition = NULL,
  partition.sp = NULL,
  start = 1,
  thin = 1,
  Yc = NULL,
  mcmcStep = 1,
  expected = TRUE,
  initPar = NULL,
  nParallel = 1,
  useSocket = .Platform$OS.type == "windows",
  nChains = length(hM$postList),
  updater = list(),
  verbose = nParallel == 1,
  alignPost = TRUE
)
```

Arguments

- **hM**: a fitted Hmsc model object
- **partition**: partition vector for cross-validation created by `createPartition`
- **partition.sp**: species partitioning vector for conditional cross-validation
- **start**: index of first MCMC sample included
- **thin**: thinning interval of posterior distribution
- **Yc**: response matrix on which the predictions are to be conditioned
- **mcmcStep**: number of MCMC steps used to make conditional predictions
- **expected**: whether expected values (TRUE) or realizations (FALSE) are to be predicted
- **initPar**: a named list of parameter values used for initialization of MCMC states
- **nParallel**: number of parallel processes by which the chains are executed
- **nChains**: number of independent MCMC chains to be run
- **updater**: a named list, specifying which conditional updaters should be omitted
- **verbose**: the interval between MCMC steps printed to the console
- **alignPost**: boolean flag indicating whether the posterior of each chains should be aligned
- **useSocket**: (logical) use socket clusters in parallel processing; these can be used in all operating systems, but they are usually slower than forking which can only be used in non-Windows operating systems (macOS, Linux, unix-like systems).

Details

There are two alternative functions `computePredictedValues` and `pcomputePredictedValues`. Function `pcomputePredictedValues` uses more aggressive parallelization and can be much faster.
when partition is used. Function `computePredictedValues` can run chains of each sampleMcmc partition in parallel, but `pcomputePredictedValues` can run each partition fold times chain in parallel (if hardware and operating systems permit). Function `pcomputePredictedValues` is still experimental, and therefore we provide both the old and new functions, but the old functions is scheduled to be removed in the future. Species partitions are not yet parallelized, and they can be very slow, especially with many mcmcSteps.

If the option partition is not used, the posterior predictive distribution is based on the model fitted to the full data. If the option partition is used but partition.sp is not used, the posterior predictive distribution is based on cross-validation over the sampling units. If partition.sp is additionally used, then, when predictions are made for each fold of the sampling units, the predictions are done separately for each fold of species. When making the predictions for one fold of species, the predictions are conditional on known occurrences (those observed in the data) of the species belonging to the other folds. If partition.sp is used, the parameter mcmcStep should be set high enough to obtain appropriate conditional predictions. The option Yc can be used alternatively to partition.sp if the conditioning is to be done based on a fixed set of data (independently of which sampling unit and species the predictions are made for).

**Value**

an array of model predictions, made for each posterior sample

**See Also**

predict.Hmsc

**Examples**

```r
# Compute predicted values using a previously fitted HMSC model
preds = computePredictedValues(TD$m)

## Not run:
# Compute predicted values for a previously fitted HMSC model using 2 folds
partition = createPartition(TD$m, nfolds = 2)
predsCV1 = computePredictedValues(TD$m, partition=partition)

# Compute conditional predictions for a previously fitted HMSC model using 2 folds
partition = createPartition(TD$m, nfolds = 2)
predsCV2 = computePredictedValues(TD$m, partition = partition,
partition.sp = 1:TD$m$ns, mcmcStep = 100)

## End(Not run)
```
computeVariancePartitioning

Description

Computes variance components with respect to given grouping of fixed effects and levels of random effects.

Usage

computeVariancePartitioning(
  hM,
  group = NULL,
  groupnames = NULL,
  start = 1,
  na.ignore = FALSE
)

Arguments

- **hM**: a fitted Hmsc model object
- **group**: vector of numeric values corresponding to group identifiers in groupnames. If the model was defined with XData and XFormula, the default is to use model terms.
- **groupnames**: vector of names for each group of fixed effect. Should match group. If the model was defined with XData and XFormula, the default is to use the labels of model terms.
- **start**: index of first MCMC sample included
- **na.ignore**: logical. If TRUE, covariates are ignored for sites where the focal species is NA when computing variance-covariance matrices for each species

Details

The vector group has one value for each column of the matrix hM$X, describing the index of the group in which this column is to be included. The names of the group are given by groupnames. The output object VP$vals gives the variance proportion for each group and species. The output object VP$R2T gives the variance among species explained by traits, measured for species’ responses to covariates (VP$R2T$Beta) and species occurrences (VP$R2T$Y).

Value

returns an object VP with components VP$vals, VP$R2T, VP$group and VP$groupnames.

See Also

Use `plotVariancePartitioning` to display the result object.

Examples

# Partition the explained variance for a previously fitted model
# without grouping environmental covariates
VP = computeVariancePartitioning(TD$m)
# Partition the explained variance for a previously fitted model
# while grouping the two environmental variables together
VP = computeVariancePartitioning(TD$m, group=c(1,1), groupnames = c("Habitat"))

---

computeWAIC  

**Description**

Computes the value of WAIC (Widely Applicable Information Criterion) for the Hmsc model

**Usage**

```
computeWAIC(hM, ghN = 11, byColumn = FALSE)
```

**Arguments**

- `hM` a fitted Hmsc model object
- `ghN` order of Gauss-Hermite quadrature for approximate numerical integration
- `byColumn` describes whether WAIC is computed for the entire model `byColumn=FALSE` or for each column (i.e. species) `byColumn=TRUE`

**Details**

The result is exact for normal and probit observational models. For Poisson-type observational model the result is obtained through numerical integration using Gauss-Hermite quadrature.

**Value**

the scalar WAIC

**Examples**

```
# Compute WAIC of previously sampled Hmsc object
WAIC = computeWAIC(TD$m)
```
**Description**

Constructs an environmental gradient over one of the variables included in `XData`.

**Usage**

```r
constructGradient(
  hM, 
  focalVariable, 
  non.focalVariables = list(), 
  ngrid = 20, 
  coordinates = list()
)
```

**Arguments**

- `hM`: a fitted `Hmsc` model object.
- `focalVariable`: focal variable over which the gradient is constructed.
- `non.focalVariables`: list giving assumptions on how non-focal variables co-vary with the focal variable or a single number given the default type for all non-focal variables.
- `ngrid`: number of points along the gradient (for continuous focal variables).
- `coordinates`: A named list of coordinates were model is evaluated in spatial or temporal models. The name should be one of the random levels, and value can be "c" for mean of coordinates (default), "i" for infinite coordinates without effect of spatial dependence, or a numeric vector of coordinates where the model is evaluated.

**Details**

In basic form, `non.focalVariables` is a list, where each element is on the form variable=list(type,value), where variable is one of the non-focal variables, and the value is needed only if type = 3. Alternatives type = 1 sets the values of the non-focal variable to the most likely value (defined as expected value for covariates, mode for factors), type = 2 sets the values of the non-focal variable to most likely value, given the value of focal variable, based on a linear relationship, and type = 3 fixes to the value given. As a shortcut, a single number 1 or 2 can be given as a type used for all non-focal variables. If a `non.focalVariable` is not listed, type=2 is used as default. Note that if the focal variable is continuous, selecting type 2 for a non-focal categorical variable can cause abrupt changes in response.

The function needs access to the original `XData` data frame, and cannot be used if you defined your model with `X` model matrix. In that case you must construct your gradient manually.

**Value**

a named list with slots `XDataNew`, `studyDesignNew` and `rLNew`
constructKnots

See Also

plotGradient, predict.

Examples

# Construct gradient for environmental covariate called 'x1'.
Gradient = constructGradient(TD$m, focalVariable="x1")

# Construct gradient for environmental covariate called 'x1'
# while setting the other covariate to its most likely values
Gradient = constructGradient(TD$m, focalVariable="x1", non.focalVariables=list(x2=list(1)))

constructKnots

Description

Construct a Regular Grid of Knot Locations for Spatial GPP Model

Usage

constructKnots(sData, nKnots = NULL, knotDist = NULL, minKnotDist = NULL)

Arguments

sData a dataframe containing spatial or temporal coordinates of units of the random level
nKnots the number of knots wanted on the spatial dimension with the shortest range
knotDist the distance between the wanted knots
minKnotDist the minimum distance of a knot to the nearest data point

Details

This is a helper function for spatial Hmsc models with the spatial method set to GPP where user must provide knot locations. Knot locations with a distance greater than minKnotDist to the nearest data point are dropped from the grid. If the input locations are SpatialPoints data, these are treated like Euclidean coordinates, and if the points are not projected, a warning is issued.

Only one of nKnots and minKnotDist arguments can be provided.

Value

a data frame with knot locations
Examples

```r
# Creating knots for some 2 dimensional spatial data
n = 100
xycoords = matrix(runif(2*n),ncol=2)
xyKnots = constructKnots(xycoords,knotDist = 0.2, minKnotDist = 0.5)
```

convertToCodaObject  convertToCodaObject

Description

Converts the Hmsc posterior into a named list of mcmc.list objects

Usage

```r
classToCodaObject(
  hM,  # a fitted Hmsc model object
  start = 1,  # index of first MCMC sample included
  spNamesNumbers = c(TRUE, TRUE),  # logical of length 2, where first entry controls whether species names are printed,
covNamesNumbers = c(TRUE, TRUE),  # and second entry controls whether species numbers are printed
  trNamesNumbers = c(TRUE, TRUE),  # Logical of length 2, where first entry controls whether covariate names are printed,
  Beta = TRUE,  # and second entry controls whether covariate numbers are printed
  Gamma = TRUE,
  V = TRUE,
  Sigma = TRUE,
  Rho = TRUE,
  Eta = TRUE,
  Lambda = TRUE,
  Alpha = TRUE,
  Omega = TRUE,
  Psi = TRUE,
  Delta = TRUE
)
```

Arguments

- `hM`: a fitted Hmsc model object
- `start`: index of first MCMC sample included
- `spNamesNumbers`: logical of length 2, where first entry controls whether species names are printed, and second entry controls whether species numbers are printed
- `covNamesNumbers`: Logical of length 2, where first entry controls whether covariate names are printed, and second entry controls whether covariate numbers are printed
- `trNamesNumbers`: Logical of length 2, where first entry controls whether trait names are printed, and second entry controls whether traits numbers are printed
createPartition

<table>
<thead>
<tr>
<th>Beta</th>
<th>logical indicating whether posterior of Beta is included</th>
</tr>
</thead>
<tbody>
<tr>
<td>Gamma</td>
<td>logical indicating whether posterior of Gamma is included</td>
</tr>
<tr>
<td>V</td>
<td>logical indicating whether posterior of V is included</td>
</tr>
<tr>
<td>Sigma</td>
<td>logical indicating whether posterior of Sigma is included</td>
</tr>
<tr>
<td>Rho</td>
<td>logical indicating whether posterior of Rho is included</td>
</tr>
<tr>
<td>Eta</td>
<td>logical indicating whether posterior of Eta is included</td>
</tr>
<tr>
<td>Lambda</td>
<td>logical indicating whether posterior of Lambda is included</td>
</tr>
<tr>
<td>Alpha</td>
<td>logical indicating whether posterior of Alpha is included</td>
</tr>
<tr>
<td>Omega</td>
<td>logical indicating whether posterior of Omega is included</td>
</tr>
<tr>
<td>Psi</td>
<td>logical indicating whether posterior of Psi is included</td>
</tr>
<tr>
<td>Delta</td>
<td>logical indicating whether posterior of Delta is included</td>
</tr>
</tbody>
</table>

**Value**

A named list that can be analysed with `coda` functions.

**Examples**

```r
# Convert recorded posterior samples in \code{Hmsc} object to coda object
codaObject = convertToCodaObject(TD$m)

# Convert recorded posterior samples, starting from sample 100, in m object to coda object
codaObject = convertToCodaObject(TD$m, start=100)
```

**Description**

Constructs a partition vector given the number of folds and column of study design.

**Usage**

```r
createPartition(hM, nfolds = 10, column = NULL)
```

**Arguments**

- `hM`: a fitted Hmsc model object
- `nfolds`: number of cross-validation folds
- `column`: name or index of the column in the studyDesign matrix, corresponding to the level for which units are splitted into folds

**Value**

A vector describing the fold of each sampling unit.
Examples

```r
# Create 3 folds for a HMSC object
partition = createPartition(TD$m, nfolds = 3)
```

Description

Computes measures of model fit for a Hmsc model

Usage

```r
evaluateModelFit(hM, predY)
```

Arguments

- `hM`: a fitted Hmsc model object
- `predY`: array of predictions, typically posterior sample

Details

All measures of model fit are based on comparing the posterior predictive distribution (predY) to the observed values (hM$Y). The predicted distribution is first summarized to a single matrix of predicted values by taking the posterior mean (for normal and probit models) or posterior median (for Poisson models). All measures of model fit are given as vectors with one value for each species.

The kinds of measures of model fit depend on the type of response variable. For all types of response variables, root-mean-square error (RMSE) between predicted and observed values is computed. For normal models, $R^2$ is computed as squared pearson correlation between observed and predicted values, times the sign of the correlation. For probit models, Tjur $R^2$ and AUC are computed. For Poisson models, a pseudo-$R^2$ is computed as squared spearman correlation between observed and predicted values, times the sign of the correlation (SR2). For Poisson models, the observed and predicted data are also truncated to occurrences (presence-absences), for which the same measures are given as for the probit models (O.RMSE, O.AUC and O.TjurR2). For Poisson models, the observed and predicted data are also subsetted to conditional on presence, for which the root-mean-square error and pseudo-R2 based on squared spearman correlation are computed (C.RMSE, C.SR2).

The measures O.RMSE, O.AUC, O.TjurR2, C.RMSE and C.SR2 can be computed only if the option `expected=FALSE` has been used when making the predictions.

If the model includes a mixture of response variable types, the resulting measures of model fit contain NA's for those response variables for which they cannot be computed.

Value

a list of measures of model fit
Examples

# Evaluate model fit
preds = computePredictedValues(TD$m)
MF = evaluateModelFit(hM=TD$m, predY=preds)

# Evaluate model performance based on cross validation: this will be slow
## Not run:
partition = createPartition(TD$m, nfolds = 2)
predsCV1 = computePredictedValues(TD$m, partition=partition)
MF = evaluateModelFit(hM=TD$m, predY=predsCV1)

## End(Not run)

---

getPostEstimate

Description

Calculates mean, support and other posterior quantities for a specified model parameter

Usage

getPostEstimate(
  hM,
  parName,
  r = 1,
  x = NULL,
  q = c(),
  chainIndex = 1:length(hM$postList),
  start = 1,
  thin = 1
)

Arguments

hM          a fitted Hmsc model object
parName     name of the parameter to be summarized. Can take value of model’s baseline parameters, "Omega" or "OmegaCor".
r           the random level for which to calculate the parameter. Has effect only for Eta, Lambda, Omega and OmegaCor.
x           values of covariates for covariate dependent omega
q            vector of quantiles to calculate.
chainIndex  which posterior chains to use for summarization (defaults to all)
start       index of first MCMC sample included
thin         thinning interval of posterior distribution
Value

A named list of posterior quantities.

Examples

# Get posterior mean and support for species' responses to environmental covariates
postBeta = getPostEstimate(TD$m, parName='Beta')

# Get posterior mean and support for species' responses to latent factors for the first random level
postLambda = getPostEstimate(TD$m, parName='Lambda', r=1)

Description

Creates an Hmsc-class object

Usage

Hmsc(
  Y,
  XFormula = ~.,
  XData = NULL,
  X = NULL,
  XScale = TRUE,
  XSelect = NULL,
  XRRRData = NULL,
  XRRRFormula = ~. - 1,
  XRRR = NULL,
  ncRRR = 2,
  XRRRScale = TRUE,
  YScale = FALSE,
  studyDesign = NULL,
  ranLevels = NULL,
  ranLevelsUsed = names(ranLevels),
  TrFormula = NULL,
  TrData = NULL,
  Tr = NULL,
  TrScale = TRUE,
  phyloTree = NULL,
  C = NULL,
  distr = "normal",
  truncateNumberOfFactors = TRUE
)
Arguments

- **Y**  
  a matrix of species occurrences or abundances
- **XFormula**  
  a formula-class object for fixed effects (linear regression)
- **XData**  
  a data frame of measured covariates for fixed effects with formula-based specification
- **X**  
  a matrix of measured covariates for fixed effects with direct specification
- **XScale**  
  a boolean flag indicating whether to scale covariates for the fixed effects
- **XSelect**  
  a list describing how variable selection is to be applied
- **XRRRData**  
  a data frame of covariates for reduced-rank regression
- **XRRRFormula**  
  formula for reduced-rank regression
- **XRRR**  
  a matrix of covariates for reduced-rank regression
- **ncRRR**  
  number of covariates (linear combinations) for reduced-rank regression
- **XRRRScale**  
  a boolean flag indicating whether to scale covariates for reduced-rank regression
- **YScale**  
  a boolean flag whether to scale responses for which normal distribution is assumed
- **studyDesign**  
  a data frame of correspondence between sampling units and units on different levels of latent factors
- **ranLevels**  
  a named list of HmscRandomLevel-class objects, specifying the structure and data for random levels
- **ranLevelsUsed**  
  a vector with names of levels of latent factors that are used in the analysis
- **TrFormula**  
  a formula-class object for regression dependence of $\beta_{kj}$ coefficients on species traits
- **TrData**  
  a data frame of measured species traits for formula-based specification
- **Tr**  
  a matrix of measured traits for direct specification
- **TrScale**  
  a boolean flag whether to scale values of species traits
- **phyloTree**  
  a phylogenetic tree (object of class phylo or corPhyl) for species in Y
- **C**  
  a phylogenetic correlation matrix for species in Y
- **distr**  
  a string shortcut or $n_s \times 2$ matrix specifying the observation models
- **truncateNumberOfFactors**  
  logical, reduces the maximal number of latent factor to be at most the number of species

Details

Matrix Y may contain missing values, but it is not recommended to add a species/sampling unit with fully missing data, since those do not bring any new additional information.

Only one of XFormula-XData and X arguments can be specified. Similar requirement applies to TrFormula-TrData and Tr. It is recommended to use the specification with formula, since that information enables additional features for postprocessing of the fitted model.

As default, scaling is applied for X and Tr matrices, but not for Y matrix. If the X and/or Tr matrices are scaled, the estimated parameters are back-transformed so that the estimated parameters
correspond to the original X and Tr matrices, not the scaled ones. In contrast, if Y is scaled, the estimated parameters are not back-transformed because doing so is not possible for all model parameters. Thus, the estimated parameters correspond to the scaled Y matrix, not the original one. If the Y matrix is scaled, the predictions generated by predict are back-transformed, so that the predicted Y matrices are directly comparable to the original Y matrix. If default priors are assumed, it is recommended that all matrices (X, Tr and Y) are scaled.

The object XSelect is a list. Each object of the list Xsel = XSelect[[i]] is a named list with objects Xsel$covGroup, Xsel$spGroup and Xsel$q. The parameter covGroup is a vector containing the columns of the matrix X for which variable selection is applied. The parameter spGroup is a vector of length equal to the number of species ns, with values 1,...,ng, where ng is the number of groups of species for which variable selection is applied simultaneously. The parameter q is a vector of length ng, containing the prior probabilities by which the variables are to be included. For example, choosing covGroup = c(2,3), spGroup = rep(1,ns) and q=0.1 either includes or excludes both of the covariates 2 and 3 simultaneously for all species. For another example, choosing covGroup = c(2,3), spGroup = 1:ns and q=rep(0.1,ns) either includes or excludes both of the covariates 2 and 3 separately for each species.

The included random levels are specified by the ranLevels and ranLevelsUsed arguments. The correspondence between units of each random level and rows of Y must be specified by a column of studyDesign, which corresponds to the name of a list item in ranLevels. It is possible to provide an arbitrary number of columns in studyDesign that are not listed in ranLevels. These do not affect the model formulation or fitting scheme, but can be utilized during certain functions postprocessing the results of statistical model fit.

The distr argument may be either a matrix, a string literal, or a vector of string literals. In the case of a matrix, the dimension must be $n_s \times 2$, where the first column defines the family of the observation model and the second argument defines the dispersion property. The elements of the first column must take values 1-normal, 2-probit and 3-Poisson with log link function. The second argument stands for the dispersion parameter being fixed (0) or estimated (1). The default fixed values of the dispersion parameters are 1 for normal and probit, and 0.01 for Poisson (implemented as a limiting case of lognormally-overdispersed Poisson). Alternatively, a string literal shortcut can be given as a value to the distr argument, simultaneously specifying similar class of observation models for all species. The available shortcuts are "normal", "probit", "poisson", "lognormal poisson". If distr is a vector of string literals, each element corresponds to one species, should be either "normal", "probit", "poisson", "lognormal poisson", and these can be abbreviated as long as they are unique strings. The matrix argument and the vector of string literals allows specifying different observation models for different species.

By default this constructor assigns default priors to the latent factors. Those priors are designed to be reasonably flat assuming that the covariates, species traits and normally distributed responses are scaled. In case when other priors needed to be specified, a call of setPriors.Hmsc methods should be made, where the particular priors may be specified.

Value

An object of Hmsc class without any posterior samples.

See Also

HmscRandomLevel, sampleMcmc, setPriors.Hmsc
Examples

# Creating a Hmsc object without phylogeny, trait data or random levels
m = Hmsc(Y=TD$Y, XData=TD$X, XFormula=~x1+x2)

# Creating a Hmsc object with phylogeny and traits
m = Hmsc(Y=TD$Y, XData=TD$X, XFormula=~x1+x2,
TrData=TD$Tr, TrFormula=~T1+T2, phyloTree=TD$phylo)

# Creating a Hmsc object with 2 nested random levels (50 sampling units in 20 plots)
studyDesign = data.frame(sample = as.factor(1:50), plot = as.factor(sample(1:20,50,replace=TRUE)))
rL1 = HmscRandomLevel(units=levels(TD$studyDesign$plot))
rL2 = HmscRandomLevel(units=levels(TD$studyDesign$sample))
m = Hmsc(Y=TD$Y, XData=TD$X, XFormula=~x1+x2,
studyDesign=studyDesign, ranLevels=list("sample"=rL1,"plot"=rL2))

HmscRandomLevel

Create an Hmsc random level

Description

Specifies the structure of a random factor, including whether the random factor is assumed to be spatially explicit or not, the spatial coordinates and the potential structure of covariate-dependent random factors.

Usage

HmscRandomLevel(
  sData = NULL,
  sMethod = "Full",
  distMat = NULL,
  xData = NULL,
  units = NULL,
  N = NULL,
  nNeighbours = 10,
  sKnot = NULL,
  longlat = FALSE
)

Arguments

sData a matrix or a dataframe containing spatial or temporal coordinates of units of the random level, or a similar SpatialPoints structure of the sp package. If spatial coordinates are unprojected longitude and latitude, great circle distances will be calculated internally. All spatial locations should be unique. If you have several observations in the same point, they should be identified by the random levels.

sMethod a string specifying which spatial method to be used. Possible values are "Full", "GPP" and "NNGP"
distMat  a distance matrix containing the distances between units of the random level, with unit names as rownames, or a dist structure with location Labels. distMat cannot be used with "GPP" spatial model.

dataframe containing the covariates measured at the units of the random level for covariate-dependent associations.

units  a vector, specifying the names of the units of a non-structured level

N  number of unique units on this level

nNeighbours  a scalar specifying the number of neighbours to be used in case the spatial method is set to NNGP. Only positive values smaller than the total number of plots are allowed.

sKnot  a dataframe containing the knot locations to be used for the Gaussian predictive process if sMethod is set to "GPP". Suitable data can be produced with constructKnots. The knot locations shall not duplicate sData.

sData as longitude and latitude in decimal degrees. If this is TRUE, great circle distances will be used instead of Euclidean distances.

Details

Only one of sData, distMat, xData, units and N arguments can be provided.

As a good practice, we recommend to specify all available units for a random level, even if some of those are not used for training the model.

Value

a HmscRandomLevel-class object that can be used for Hmsc-class object construction

See Also

setPriors.Hmsc to change the default priors of an existing HmscRandomLevel object.

Examples

# Setting a random level with 50 units
rL = HmscRandomLevel(units=TD$studyDesign$sample)

# Setting a spatial random level
rL = HmscRandomLevel(sData=TD$xycoords)

# Setting a covariate-dependent random level.
rL = HmscRandomLevel(xData=data.frame(x1=rep(1,length(TD$X$x1)),x2=TD$X$x2))
plotBeta

Description

Plots heatmaps of parameter estimates or posterior support values of species’ environmental responses, i.e. how species in Y responds to covariates in X

Usage

plotBeta(
  hM,  # a fitted Hmsc model object
  post,  # posterior summary of Beta parameters obtained fromgetPostEstimate
  param = "Support",  # controls which parameter is plotted, current options include "Mean" for posterior mean estimate, "Support" for the level of statistical support measured by posterior probability for a positive or negative response, and "Sign" to indicate whether the response is positive, negative, or neither of these given the chosen supportLevel
  plotTree = FALSE,  # logical. Whether species’ environmental responses is to be mapped onto the phylogeny used in model fitting
  SpeciesOrder = "Original",  # SpVector = NULL,  # covOrder = "Original",
  covVector = NULL,  # spNamesNumbers = c(TRUE, TRUE),
  supportLevel = 0.9,  # covNamesNumbers = c(TRUE, TRUE),
  split = 0.3,  # supportLevel = 0.9,
  cex = c(0.7, 0.7, 0.8),  # colors = colorRampPalette(c("blue", "white", "red")),
  colors = colorRampPalette(c("blue", "white", "red")),  # colorLevels = NULL,
  colorLevels = NULL,  # mar = NULL,
  mar = NULL,  # marTree = c(6, 0, 2, 0),
  marTree = c(6, 0, 2, 0),  # mgp = c(3, 2, 0),
  mgp = c(3, 2, 0),  # main = NULL,
  main = NULL,  # smallplot = NULL,
  smallplot = NULL,  # bigplot = NULL,
  bigplot = NULL,
  newplot = TRUE
)

Arguments

hM a fitted Hmsc model object
post posterior summary of Beta parameters obtained fromgetPostEstimate
param controls which parameter is plotted, current options include "Mean" for posterior mean estimate, "Support" for the level of statistical support measured by posterior probability for a positive or negative response, and "Sign" to indicate whether the response is positive, negative, or neither of these given the chosen supportLevel
plotTree logical. Whether species’ environmental responses is to be mapped onto the phylogeny used in model fitting
SpeciesOrder controls the ordering of species, current options are "Original", "Tree", and "Vector". If SpeciesOrder = "Vector", an ordering vector must be provided (see SpVector). If plotTree = TRUE, SpeciesOrder is ignored.

SpVector controls the ordering of species if SpeciesOrder = "Vector". If a subset of species are listed, only those will be plotted. For alphabetic ordering, try match(1:hM$ns, as.numeric(as.factor(colnames(hM$Y))))

covOrder controls the ordering of covariates, current options are "Original" and "Vector". If covOrder = "Vector", an ordering vector must be provided (see covVector)

covVector controls the ordering of covariates if covOrder = "Vector". If a subset of covariates are listed, only those will be plotted.

spNamesNumbers logical of length 2, where first entry controls whether species names are added to axes, and second entry controls whether species numbers are added.

covNamesNumbers logical of length 2, where first entry controls whether covariate names are added to axes, and second entry controls whether covariate numbers are added.

supportLevel controls threshold posterior support for plotting.

split if plotTree = TRUE, controls the division of the plotting window between the tree and the heatmap.

cex controls character expansion (font size). Three values, controlling covariate names, species names, and color legend axis labels.

colors controls the colors of the heatmap, default value colorRampPalette(c("blue","white","red"))

colorLevels number of color levels used in the heatmap.

Examples

# Plot posterior support values of species' environmental responses
betaPost=getPostEstimate(TD$m, "Beta")
plotBeta(TD$m, post=betaPost, param="Support")

# Plot parameter estimates of species' environmental responses together with the phylogenetic tree
betaPost=getPostEstimate(TD$m, "Beta")
plotBeta(TD$m, post=betaPost, param="Mean", plotTree=TRUE)
Description

Plots heatmaps of parameter estimates or posterior support values of trait effects on species’ environmental responses, i.e. how environmental responses in \( \beta \) respond to covariates in \( X \).

Usage

```r
plotGamma(
  hM,  
  post,  
  param = "Support",  
  trOrder = "Original",  
  trVector = NULL,  
  covOrder = "Original",  
  covVector = NULL,  
  trNamesNumbers = c(TRUE, TRUE),  
  covNamesNumbers = c(TRUE, TRUE),  
  supportLevel = 0.9,  
  main = NULL,  
  cex = c(0.8, 0.8, 0.8),  
  colors = colorRampPalette(c("blue", "white", "red")),  
  colorLevels = NULL,  
  mar = c(6, 9, 2, 0),  
  smallplot = NULL,  
  bigplot = NULL,  
  newplot = TRUE
)
```

Arguments

- **hM**: a fitted \( \text{Hmsc} \) model object
- **post**: posterior summary of Gamma parameters obtained from `getPostEstimate`
- **param**: controls which parameter is plotted, current options include "Mean" for posterior mean estimate, "Support" for the level of statistical support measured by posterior probability for a positive or negative response, and "Sign" to indicate whether the response is positive, negative, or neither of these given the chosen supportLevel
- **trOrder**: controls the ordering of traits, current options are "Original", and "Vector". If `trOrder = "Vector"`, an ordering vector must be provided (see `trVector`)
- **trVector**: controls the ordering of traits if `trOrder = "Vector"`. If a subset of traits are listed, only those will be plotted
- **covOrder**: controls the ordering of covariates, current options are "Original" and "Vector". If `covOrder = "Vector"`, an ordering vector must be provided (see `covVector`)
Description

Plots an environmental gradient over one of the variables included in `XData`

Usage

```r
plotGradient(
  hM,
  Gradient,
  predY,
  measure,
  xlabel = NULL,
  ylabel = NULL,
  ...)
```
index = 1,  
q = c(0.025, 0.5, 0.975),  
cicol = rgb(0, 0, 1, alpha = 0.5),  
pointcol = "lightgrey",  
pointsize = 1,  
showData = FALSE,  
jigger = 0,  
yshow = NA,  
showPosteriorSupport = TRUE,  
main,  
...

Arguments

hM a fitted Hmsc model object
Gradient an object returned by constructGradient
predY an object returned by applying the function predict to Gradient
measure whether to plot species richness ("S"), an individual species ("Y") or community-weighted mean trait values ("T")
xlabel label for x-axis
ylabel label for y-axis
index which species or trait to plot
q quantiles of the credibility interval plotted
cicol colour with which the credibility interval is plotted
pointcol colour with which the data points are plotted
pointsize size in which the data points are plotted
showData whether raw data are plotted as well
jigger the amount by which the raw data are to be jiggered in x-direction (for factors) or y-direction (for continuous covariates)
yshow scale y-axis so that these values are also visible. This can used to scale y-axis so that it includes 0 and the expected maximum values.
showPosteriorSupport add margin text on the posterior support of predicted change from gradient minimum to maximum for continuous gradients.
main main title for the plot.
... additional arguments for plot

Details

For measure="Y", index selects which species to plot from hM$spNames. For measure="T", index selects which trait to plot from hM$strNames. With measure="S" the row sum of pred is plotted, and thus the interpretation of "species richness" holds only for probit models. For Poisson models "S" shows the total count, whereas for normal models it shows the summed response. For measure="T", in probit model the weighting is over species occurrences, whereas in count models it is over individuals. In normal models, the weights are exp-transformed predictions to avoid negative weights.
Value

For the case of a continuous covariate, returns the posterior probability that the plotted variable is greater for the last sampling unit of the gradient than for the first sampling unit of the gradient. For the case of a factor, returns the plot object.

See Also

constructGradient, predict

Examples

# Plot response of species 2 over the gradient of environmental variable x1
Gradient = constructGradient(TD$m, focalVariable="x1")
predY = predict(TD$m, Gradient=Gradient)
plotGradient(TD$m, Gradient, pred=predY, measure="Y", index = 2, showData = TRUE, jigger = 0.05)
# Plot modelled species richness over the gradient of environmental variable x1
Gradient = constructGradient(TD$m, focalVariable="x1")
predY = predict(TD$m, Gradient=Gradient)
plotGradient(TD$m, Gradient, pred=predY, measure="S")

plotVariancePartitioning

Description

Plots the results of variance partitioning of a Hmsc model produced by computeVariancePartitioning as a barplot

Usage

plotVariancePartitioning(
  hM,  
  VP,  
  cols = NULL,  
  main = "Variance Partitioning",  
  ...  
)

Arguments

hM a fitted Hmsc model object
VP a Hmsc variance partitioning object produced by computeVariancePartitioning
cols colors of the barplot
main main title for the plot
... additional parameters passed to the barplot function
Examples

# Plot how the explained variance of a previously fitted model is partitioned
VP = computeVariancePartitioning(TD$m)
plotVariancePartitioning(TD$m, VP)

poolMcmcChains

Description

Combines a list of single or several MCMC chains into a single chain

Usage

poolMcmcChains(postList, chainIndex = 1:length(postList), start = 1, thin = 1)

Arguments

postList list of posterior chains
chainIndex index of chains to be included
start index of first MCMC sample included
thin thinning between included MCMC samples

Value

a list with combined MCMC samples

Examples

# Combine the posteriors from all chains in a Hmec object
postList = TD$m$postList
pooledPost = poolMcmcChains(postList)
predict.Hmsc

### Description

Calculates predicted values from a fitted Hmsc model.

### Usage

```r
## S3 method for class 'Hmsc'
predict(
  object,
  post = poolMcmcChains(object$postList),
  XData = NULL,
  X = NULL,
  XRRRData = NULL,
  XRRR = NULL,
  studyDesign = object$studyDesign,
  ranLevels = object$ranLevels,
  Gradient = NULL,
  Yc = NULL,
  mcmcStep = 1,
  expected = FALSE,
  predictEtaMean = FALSE,
  predictEtaMeanField = FALSE,
  nParallel = 1,
  useSocket = TRUE,
  ...
)
```

### Arguments

- `object` a fitted Hmsc model object
- `post` a list of posterior samples of the HMSC model. By default uses all samples from the pooled posterior of the hM object.
- `XData` a dataframe specifying the unpreprocessed covariates for the predictions to be made. Works only if the `XFormula` argument was specified in the Hmsc() model constructor call. Requirements are similar to those in the Hmsc model constructor.
- `X` a matrix specifying the covariates for the predictions to be made. Only one of `XData` and `X` arguments may be provided.
- `XRRRData` a dataframe of covariates for reduced-rank regression
- `XRRR` a matrix of covariates for reduced-rank regression
- `studyDesign` a matrix, specifying the structure of the study design for the prediction. Requirements are similar to those of the `Hmsc` constructor. By default this argument is assigned the study design of the training data in the fitted Hmsc model.
ranLevels a list of HmscRandomLevel objects, further specifying the structure of random levels. Requirements are similar to those of the Hmsc constructor. Each level must cover all units, specified in the correspondingly named column of studyDesign argument. By default this argument is assigned the list of HmscRandomLevel objects specified for fitting Hmsc model.

Gradient an object returned by constructGradient. Providing Gradient is an alternative for providing XData, studyDesign and ranLevels. Cannot be used together with Yc.

Yc a matrix of the outcomes that are assumed to be known for conditional predictions. Cannot be used together with Gradient.

mcmcStep the number of extra mcmc steps used for updating the random effects

expected boolean flag indicating whether to return the location parameter of the observation models or sample the values from those.

predictEtaMean boolean flag indicating whether to use the estimated mean values of posterior predictive distribution for random effects corresponding for the new units.

predictEtaMeanField boolean flag indicating whether to use draws from the mean-field of the posterior predictive distribution for random effects corresponding for the new units.

nParallel Number of parallel processes. Parallel processing is only useful with new Yc data and extra mcmcStep.

useSocket (logical) Use socket clusters in parallel processing; these are the only alternative in Windows, but in other systems this should be usually set FALSE for forking.

... other arguments passed to functions.

Details

In mcmcStep, the number of extra mcmc steps used for updating the random effects for the Eta parameters, starting from the samples of the fitted Hmsc model in order to account for the conditional information provided in the Yc argument. The higher this number is, the more the obtained updated samples are unaffected by the posterior estimates of latent factors in the model fitted to the training data and more resembles the true conditional posterior. However, the elapsed time for conditional prediction grows approximately linearly as this parameter increases. The exact number for sufficient is problem-dependent and should be assessed by e.g. gradually increasing this parameter till the stationarity of the produced predictions.

Value

A list of length length(post), each element of which contains a sample from the posterior predictive distribution (given the sample of the Hmsc model parameters in the corresponding element of the post argument)

See Also

predictLatentFactor
**predictLatentFactor**

**Description**

Draws samples from the conditional predictive distribution of latent factors

**Usage**

```r
predictLatentFactor(
  unitsPred,
  units,
  postEta,
  postAlpha,
  rL,
  predictMean = FALSE,
  predictMeanField = FALSE
)
```

**Arguments**

- `unitsPred`: a factor vector with random level units for which predictions are to be made
- `units`: a factor vector with random level units that are conditioned on
- `postEta`: a list containing samples of random factors at conditioned units
- `postAlpha`: a list containing samples of range (lengthscale) parameters for latent factors
- `rL`: a `HmscRandomLevel-class` object that describes the random level structure
- `predictMean`: a boolean flag indicating whether to return the mean of the predictive Gaussian process distribution
- `predictMeanField`: a boolean flag indicating whether to return the samples from the mean-field distribution of the predictive Gaussian process distribution

**Details**

Length of `units` vector and number of rows in `postEta` matrix shall be equal. The method assumes that the i-th row of `postEta` correspond to i-th element of `units`.

This method uses only the coordinates `rL$s` field of the `rL$s` argument. This field shall be a matrix with rownames covering the union of `unitsPred` and `units` factors. Alternatively, it can use distance matrix `rL$distMat` which is a symmetric square matrix with similar row names as the coordinate data (except for the GPP models that only can use coordinates).

In case of spatial random level, the computational complexity of the generic method scales cubically as the number of unobserved units to be predicted. Both `predictMean=TRUE` and `predictMeanField=TRUE` options decrease the asymptotic complexity to linear. The `predictMeanField=TRUE` option also preserves the uncertainty in marginal distribution of predicted latent factors, but neglects the interdependence between them.
**prepareGradient**

**Value**

a list of length `length(postEta)` containing samples of random factors at `unitsPred` from their predictive distribution conditional on the values at `units`

**Description**

prepares a user-made environmental and/or spatial gradient to be used for prediction

**Usage**

```r
prepareGradient(hM, XDataNew, sDataNew)
```

**Arguments**

- `hM`: a fitted Hmsc model object.
- `XDataNew`: a dataframe of the new `XData`.
- `sDataNew`: a named list of the new `sData`, where the name gives the spatial random level.

**Details**

The dataframe `XDataNew` is the for output as for input. The main purpose of this function is to prepare the study design and random levels so that predictions can be made with the `predict` function. Note that the difference between `constructGradient` and `prepareGradient` is that while `prepareGradient` takes as input the new environmental and spatial data, `constructGradient` generates those data to represent a new environmental gradient.

**Value**

a named list with members `XDataNew`, `studyDesignNew` and `rLNew`

**See Also**

`constructGradient`, `predict`
sampleMcmc

Description

Samples the posterior with block-conditional Gibbs MCMC sampler

Usage

```r
sampleMcmc(
  hM,  
  samples,  
  transient = 0,  
  thin = 1,  
  initPar = NULL,  
  verbose,  
  adaptNf = rep(transient, hM$nr),  
  nChains = 1,  
  nParallel = 1,  
  useSocket = TRUE,  
  dataParList = NULL,  
  updater = list(),  
  fromPrior = FALSE,  
  alignPost = TRUE
)
```

Arguments

- **hM**: a fitted Hmsc model object
- **samples**: the number of MCMC samples to be obtained in each chain
- **transient**: the number of MCMC steps that are executed before starting recording posterior samples
- **thin**: the number of MCMC steps between each recording of samples from the posterior
- **initPar**: a named list of parameter values used for initialization of MCMC states, or alternatively text "fixed effects" to use linear Maximum Likelihood model instead of randomizing from prior; the "fixed effects" can shorten the transient phase of sampling, but will initialize all chains to the same starting values
- **verbose**: the interval between MCMC steps printed to the console (default is an interval that prints ca. 50 reports)
- **adaptNf**: a vector of length \( n_r \) with number of MCMC steps at which the adaptation of the number of latent factors is conducted
- **nChains**: number of independent MCMC chains to be run
- **nParallel**: number of parallel processes by which the chains are executed.
useSocket (logical) use socket clusters in parallel processing; in Windows this is the only option, but in other operating systems fork clusters are a better alternative, and this should be set FALSE.

dataParList a named list with pre-computed Qg, iQg, RQg, detQg, rLPar parameters

updater a named list, specifying which conditional updaters should be omitted

fromPrior whether prior (TRUE) or posterior (FALSE) is to be sampled

alignPost boolean flag indicating whether the posterior of each chains should be aligned

Details

The exact number of samples to be recorded in order to get a proper estimate of the full posterior with Gibbs MCMC algorithms, as well as the required thinning and cut-off of transient is very problem-specific and depends both on the model structure and the data itself. Therefore, in general it is very challenging to a priori provide an informed recommendation on what values should be used for a particular problem. A common recommended strategy involves executing the posterior sampling with MCMC with some guess of the values for these arguments, checking the properties of the obtained samples (primarily potential scale reduction factor and effective sample size), and adjusting the guess accordingly.

The value of 1 for thin argument means that at each MCMC step after the transient a sample is recorded.

Typically, the value of nParallel equal to nChains leads to most efficient usage of available parallelization capacities. However, this may be not the case if R is configured with multi-tread linear algebra libraries. For debug and test purposes, the nParallel should be set to 1, since only in this case a details of the potentially encountered errors would be available.

The dataParList argument may be handy for large problems that needs to be refitted multiple times, e.g. with different prior values. In that case, the data parameters that are precomputed for the Hmsc sampling scheme may require an undesirably lot of storage space if they are saved for each of the model. Instead, they could be computed only once and then directly reused, therefore reducing the storing redundancy.

Some of the available conditional updaters partially duplicate each other. In certain cases, the usage of all of them may lead to suboptimal performance, compared to some subset of those. Then, it is possible to manually disable some of them, by adding a $UPDATER_NAME=FALSE pair to the updater argument. Another usage of this argument involves cases when some of the model parameters are known and have to be fixed. However, such tweaks of the sampling scheme should be done with caution, as if compromized they would lead to erroneuos results.

Value

An Hmsc-class object with chains of posterior samples added to the postList field

See Also

Hmsc
Examples

```r
## you need 1000 or more samples, but that will take too long
## in an example
m = sampleMcmc(TD$m, samples=10)

## Not run:
## Record 1000 posterior samples while skipping 1 MCMC step between samples
## from 2 chains after discarding the first 500 MCMC steps
m = sampleMcmc(TD$m, samples=1000, transient=500, thin=2, nChains=2, nParallel=1)

## End(Not run)
```

## Description
Samples the parameter vector from prior

## Usage
```r
samplePrior(hM, dataParList = NULL)
```

## Arguments
- `hM`: a fitted Hmsc model object
- `dataParList`: list of data parameters (see `computeDataParameters`)

## Value
A named list containing the Hmsc model parameters

## Description
Sets or resets priors to objects

## Usage
```r
setPriors(...)```

## Arguments
- `...`: Hmsc or HmseRandolLevel object and other arguments.
Value

Object of same type as first input

See Also

setPriors.Hmsc, setPriors.HmscRandomLevel

Examples

# Set priors for random level so that there is minimum of 2 latent factors and maximum of 3
rL1 = HmscRandomLevel(units=TD$studyDesign$plot)
rL1 = setPriors(rL1, nfMax=3, nfMin=2)

# Set shrinkage parameters for priors of random level
rL1 = HmscRandomLevel(units=TD$studyDesign$plot)
rL1 = setPriors(rL1, a1=10, a2=10, b1=1, b2=1)

Description

Sets or resets priors to the Hmsc object

Usage

## S3 method for class 'Hmsc'
setPriors(
  hM,
  V0 = NULL,
  f0 = NULL,
  mGamma = NULL,
  UGamma = NULL,
  aSigma = NULL,
  bSigma = NULL,
  nuRRR = NULL,
  a1RRR = NULL,
  b1RRR = NULL,
  a2RRR = NULL,
  b2RRR = NULL,
  rhopw = NULL,
  setDefault = FALSE,
  ...
)
Arguments

- **hM**: a fitted Hmsc model object
- **V0**: scale matrix in the Wishart prior distribution for the V matrix
- **f0**: number of degrees of freedom in the Wishart prior distribution for the V matrix
- **mGamma**: mean for the prior multivariate Gaussian distribution for Gamma parameters
- **UGamma**: covariance matrix for the prior multivariate Gaussian distribution for Gamma parameters
- **aSigma**: shape parameter for the prior gamma distribution for the variance parameter, only for normal & lognormal Poisson models
- **bSigma**: rate parameter for the prior gamma distribution for the variance parameter, only for normal & lognormal Poisson models
- **nuRRR, a1RRR, b1RRR, a2RRR, b2RRR**: parameters of the multiplicative gamma process shrinking prior for reduced rank regression
- **alphapw**: discrete grid prior for phylogenetic signal, should be a matrix of 2 columns
- **setDefault**: logical indicating whether default priors should be used
- **...**: other parameters passed to the function.

Value

Modified Hmsc object

---

Description

Sets or resets priors to the Hmsc object

Usage

```r
## S3 method for class 'HmscRandomLevel'
setPriors(rl,
  nu = NULL,
  a1 = NULL,
  a2 = NULL,
  b1 = NULL,
  b2 = NULL,
  alphapw = NULL,
  nfMax = NULL,
  nfMin = NULL,
  setDefault = FALSE,
  ...
)
```
Arguments

- rL: a fitted HmscRandomLevel model object
- nu, a1, b1, a2, b2: parameters of the multiplicative gamma process shrinking prior
- alphapw: discrete grid prior for spatial scale parameter
- nfMax: maximum number of latent factors to be sampled
- nfMin: minimum number of latent factors to be sampled
- setDefault: logical indicating whether default priors should be used
- ...: other arguments (ignored)

Value

Modified HmscRandomLevel object

Simulated data and a fitted Hmsc model for a small species community.

Description

This dataset contains simulated occurrence data for 4 species in 50 sampling units. The data is based on a hierarchical study design consisting of 50 sampling units in 10 georeferenced plots. Occurrences of 4 species were simulated using one continuous environmental variable (x1) and spatial autocorrelation between the plots. Response of species to the environment are related to one species trait which is fully phylogenetically structured. This dataset is used for the examples and package testing. The variables are as follows:

Usage

TD

Format

A list of 12 objects

- ns: Number of species in the dataset
- units: Number of sampling units
- plots: Number of plots
- X: A 3 by 50 environmental matrix consisting of one continuous and one categorical variable. Also includes intercept column
- phy: A list containing the simulated phylogenetic tree for 4 species
- C: A 4 by 4 phylogenetic variance covariance matrix
- Tr: A 4 by 3 trait matrix with one phylogenetically phylogenetically structured continuous trait, one categorical trait and an intercept
xycoords simulated 2 dimensional coordinates

studyDesign Sampling unit and plot IDs

Y Simulated species occurrences

m A fitted Hmsc object with 100 posterior samples
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