

# Package ‘mastif’

January 27, 2023

**Type** Package

**Title** Mast Inference and Forecasting

**Version** 2.0

**Date** 2023-1-20

**Author** James S. Clark

**Maintainer** James S. Clark <jimclark@duke.edu>

**Description** Analyzes production and dispersal of seeds dispersed from trees and recovered in seed traps. Motivated by long-term inventory plots where seed collections are used to infer seed production by each individual plant.

**License** GPL (>= 2)

**Imports** Rcpp (>= 0.11.5), RANN, corrplot, xtable, repmis, robustbase, stringi, stringr

**Depends** R (>= 2.10)

**LinkingTo** Rcpp, RcppArmadillo

**Suggests** knitr, rmarkdown

**VignetteBuilder** knitr

**NeedsCompilation** yes

**Repository** CRAN

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mastif-package	<i>Mast Inference and Forecasting</i>
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## Description

Seed production is estimated from censuses of trees and seed collections from traps. From locations of known trees and seed traps, infers source strength, coefficients for predictor variables, and parameters for a dispersal kernel. Fecundity is a state-space model allowing for random individual (tree) effects, random year effects and random AR(p) lag effects. Estimates unknown redistribution of seed types to known species identities of trees. Functions begin with 'mast' to avoid conflicts with other packages.

## Details

Package: mastif  
Type: Package  
Version: 2.0  
Date: 2023-1-20  
License: GPL (>= 2)  
URL: <http://sites.nicholas.duke.edu/clarklab/code/>

The package mastif estimates fecundity of trees and dispersion of seed observed at seed traps, using information on locations of sources and detectors, and covariates that could explain source strength. Data sets of this type are common and used to understand a range of processes related to seed dispersal, masting, environmental controls on reproduction, sex ratio, and allocation.

Posterior simulation is done by Gibbs sampling. Analysis is done by these functions:

[mastif](#) fits model with Gibbs sampling.

[mastSim](#) simulates data for analysis by [mastif](#).

[mastFillCensus](#) aligns sample years in tree census data with seed trap data.

[mastClimate](#) annotates tree data with covariates for fecundity modeling.

[mastPlot](#) generates plots of the output from [mastif](#).

## Author(s)

Author: James S Clark, <[jimclark@duke.edu](mailto:jimclark@duke.edu)>

## References

Clark, JS, C Nunes, and B Tomasek. 2019. Masting as an unreliable resource: spatio-temporal host diversity merged with consumer movement, storage, and diet. *Ecological Monographs*, e01381.

**See Also**

[mastif](#), [mastSim](#)

A more detailed vignette is can be obtained with:

```
browseVignettes('mastif')
```

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mastClimate	<i>Covariates for mast data</i>
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**Description**

Annotates treeData for mastif to include covariates.

**Usage**

```
mastClimate( file, plots, years, months = 1:12, FUN = 'mean',
             vname = '', normYr = c( 1990:2020 ), lastYear = 2021 )
```

**Arguments**

file	character giving the path and file name where covariate is stored in mastif covariate format (see Details).
plots	character vector from treeData\$plot.
years	numeric vector from treeData\$year.
months	numeric vector of months to include in annual values.
FUN	character specifies how monthly covariate values are combined into annual values.
vname	name to use for a variable in the model that comes from file.
normYr	years for climate norm for calculating anomalies.
lastYear	last data year to include.

**Details**

The version of treeData used in mastif can have additional tree years included when there are seed trap years that were not censused or when AR(p) effects extend observations to impute the p years before and after a tree was observed. The function mastFillCensus makes this version of treeData available to the user. The function mastClimate provides a quick way to add plot-year covariates to treeData.

A covariate like minimum monthly temperature is stored in a plot by year\_month format, where rownames of file are plot names matching treeData\$plot, and colnames of file could be 2012\_1, 2012\_2, ... for the 12 months in the year. The numeric vector months holds the months to be included in the annual values, e.g., c(3, 4) for minimum winter temperatures during the period from March through April. To find the minimum for this period, set FUN to 'min'.

More detailed vignettes can be obtained with: browseVignettes('mastif')

**Value**

A numeric vector equal in length to the number of rows in `treeData` that can be added as a column and included in `formulaFec`.

**Author(s)**

James S Clark, <jimclark@duke.edu>

**References**

Clark, J.S., C. Nunes, and B. Tomasek. 2019. Foodwebs based on unreliable foundations: spatio-temporal masting merged with consumer movement, storage, and diet. *Ecological Monographs*, e01381.

**See Also**

[mastFillCensus](#) to fill tree census

[mastif](#) for analysis

A more detailed vignette is can be obtained with:

`browseVignettes('mastif')`

website '<http://sites.nicholas.duke.edu/clarklab/code/>'.

**Examples**

```
d <- "https://github.com/jimclarkatduke/mast/blob/master/liriodendronExample.rData?raw=True"
repmis::source_data(d)

inputs <- list( specNames = specNames, seedNames = seedNames,
               treeData = treeData, seedData = seedData,
               xytree = xytree, xytrap = xytrap)

# interpolate census, add years for AR(p) model
inputs <- mastFillCensus(inputs, p = 3)
treeData <- inputs$treeData #now includes additional years

# include minimum spring temperature of previous year
cfile <- tempfile(fileext = '.csv')
d <- "https://github.com/jimclarkatduke/mast/blob/master/tmin.csv?raw=True"
download.file(d, destfile=cfile)

tyears <- treeData$year - 1
tplots <- treeData$plot

tmp <- mastClimate( file = cfile, plots = tplots,
                  years = tyears, months = 1:4, FUN = 'min')
treeData$tminSprAnomaly <- tmp$x[,3]
inputs$treeData <- treeData

formulaRep <- as.formula( ~ diam )
```

```

formulaFec <- as.formula( ~ diam + tminSprAnomaly )

inputs$yearEffect <- list(groups = 'species', p = 3) # AR(3) model, species are lag groups

output <- mastif(inputs = inputs, formulaFec, formulaRep, ng = 1000, burnin = 400)

```

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mastFillCensus	<i>Interpolate census data for seed trap years</i>
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### Description

Provides interpolated census data to include years when seed data are available. This is used when tree sampling is at a lower frequency than seed-trap collections.

### Usage

```

mastFillCensus(inputs, beforeFirst = 15, afterLast = 15, p = 0, verbose = FALSE)

```

### Arguments

inputs	list used as input to <code>mastif</code> that must include two character vectors, ( <code>specNames</code> , <code>seedNames</code> ) and four <code>data.frames</code> ( <code>treeData</code> , <code>seedData</code> , <code>xytree</code> , <code>xytrap</code> ). See details.
beforeFirst	number of years before a tree is first observed in a census that it should be considered as potentially present.
afterLast	number of years after a tree is last observed in a census that it should be considered as potentially present.
p	if AR(p) model is used (in <code>yearEffect</code> ), the number of lag terms.
verbose	if <code>verbose = TRUE</code> information is provided on filling progress.

### Details

Masting data sets contain tree census data, held in `treeData`, and seed trap data, held in `seedData`. Most studies monitor seed rain frequently (e.g., annual), while tree censuses occur at intervals of 2 to 5 years. `mastFillCensus` 'fills in' the tree census so that the annual seed data can be used. It is made available to the user so covariates can be added, e.g., with `mastClimate`.

`mastFillCensus` accepts the list of inputs used in `mastif`. The missing years are inserted for each tree with interpolated diameters. `inputs` is returned with objects updated to include the missing census years and modified slightly for analysis by `mastif`.

The function `mastFillCensus` is made accessible to the user, because covariates may be needed for the missing census years. For example, models often include climate variables that change annually. The version of `treeData` returned by `mastFillCensus` can be annotated with additional

columns that can then be included in the model, as specified in `formulaFec`, `formulaRep`, and/or `randomEffect$formulaRan`.

`beforeFirst` and `afterLast` allow the user to control the assumptions about treatment of trees between (and before and after) tree census years. Seed trap data may begin before the first tree census or after the last tree census. Trees may appear in the middle of the study due to ingrowth. They may be lost to mortality. In other words, census data can be left-, right-, and interval-censored.

For the AR(p) model, values are imputed for p years before a tree is first observed and p years after the tree is last observed (`mastif`). These years are inserted by `mastFillCensus`, such that they too can then be annotated with covariate data.

More detailed vignettes can be obtained with:

```
browseVignettes('mastif')
```

## Value

`inputs` list includes all inputs to the model, cleaned for `mastif`, with years inserted to `treeData`.

## Author(s)

James S Clark, <jimclark@duke.edu>

## References

Clark, J.S., C. Nunes, and B. Tomasek. 2019. Foodwebs based on unreliable foundations: spatio-temporal masting merged with consumer movement, storage, and diet. *Ecological Monographs*, e01381.

## See Also

`mastSim` simulates data

A more detailed vignette can be obtained with:

```
browseVignettes('mastif')
```

website '<http://sites.nicholas.duke.edu/clarklab/code/>'.

## Examples

```
d <- "https://github.com/jimclarkatduke/mast/blob/master/liriodendronExample.rData?raw=True"
repmis::source_data(d)
```

```
inputs <- list( specNames = specNames, seedNames = seedNames,
               treeData = treeData, seedData = seedData,
               xytree = xytree, xytrap = xytrap)
inputs <- mastFillCensus(inputs)
```

```
formulaFec <- formulaRep <- as.formula(~ diam)
```

```
output <- mastif(inputs = inputs, formulaFec, formulaRep, ng = 1000,
```

```
burnin = 400)
```

---

```
mastif Gibbs sampler for mast data
```

---

## Description

Estimates productivity and dispersion of seeds observed at seed traps, using information on locations, and covariates that could explain source strength. Data can be simulated with [mastSim](#).

## Usage

```
mastif( inputs, formulaFec = NULL, formulaRep = NULL,
        ng = NULL, burnin = NULL )

## S3 method for class 'mastif'
print(x, ...)

## S3 method for class 'mastif'
summary(object, verbose = TRUE, latex = FALSE, ...)
```

## Arguments

<code>inputs</code>	list that must include two character vectors, ( <code>specNames</code> , <code>seedNames</code> ) and four <code>data.frames</code> ( <code>treeData</code> , <code>seedData</code> , <code>xytree</code> , <code>xytrap</code> ). See details.
<code>formulaFec</code>	R formula for fecundity model, e.g., <code>~ diam + x1</code> .
<code>formulaRep</code>	R formula for maturation model, e.g., <code>~ diam</code> .
<code>ng</code>	numeric number of Gibbs steps.
<code>burnin</code>	numeric number of burnin steps, before predictions are saved.
<code>object</code>	currently, also an object of class <code>mastif</code> .
<code>verbose</code>	if TRUE summary sent to screen.
<code>latex</code>	if TRUE tables written to screen in latex format.
<code>x</code>	object of class <code>mastif</code> .
<code>...</code>	further arguments not used here.

## Details

`inputs` includes the following:

`specNames` is a character vector containing names of species, `specNames`, that appear in the `treeData$species` column.

`seedNames` is a character vector of seed types that appear as column names in `seedData`.

`treeData` is a `data.frame` holding tree information, including predictors and tree-year identification. Required columns are `plot`, `tree`, `species`, `year`, `diam`, and any other predictors for fecundity or maturation.

`seedData` is a `data.frame` holding seed counts with seed trap and year identification. Required columns are `plot`, `trap`, `year`, and `seedNames`, the latter holding seed counts.

`xytree` is a `data.frame` holding tree locations. Required columns are `plot`, `tree`, `x`, and `y`.

`xytrap` is `data.frame` holding seed trap locations. Required columns are `plot`, `trap`, `x`, and `y`.

`formulaFec` and `formulaRep` specify the models for plant fecundity and maturation. Variables listed in formulas appear as column headings in `treeData`. Note that `formulaFec` and `formulaRep` begin with  $\sim$ , not  $y \sim$ . The response matrix is constructed from seed types in `seedData`.

The `treeData$tree` column has values that are unique for a tree within a plot. These reference the same unique identifiers in `xytree$tree`. In addition to these identifiers, the `data.frame` `xytree` holds columns `x` and `y` for map locations.

The character vector `seedNames` holds the names of columns in `seedData` for seed counts. The elements of `seedNames` are seed types produced by one or more of the species in `specNames`. `seedData` must also include columns for `trap`, `plot`, and `year`, which link with columns in `xytrap`, which additionally includes columns `x` and `y`.

`predList` includes the names of plots and years to be predicted. It can include a numeric value `mapMeters` for the distance between lattice points in the prediction grid. See examples.

`yearEffect` is a list indicating the column names in `treeData` for random groups in year effects or AR(p) models. See examples.

`randomEffect` is a list indicating the column names in `treeData` for random groups in fecundity estimates, the character `randGroups` and the `formulaRan` for random effects. The `formulaRan` must be a subset of predictors from `formulaFec`. See examples.

`modelYears` is a numeric vector of years to include in the analysis.

`ng` is the number of Gibbs steps. `burnin` is the number of initial steps, must be less than `ng`.

Additional arguments to inputs can include prior parameters; default values are:

`priorDist = 10` is a prior mean dispersal distance in meters.

`priorVDist = 1` is the prior variance on mean dispersal distance in meters.

`minDist = 2` and `maxDist = 60` are the minimum and maximum values for the mean dispersal kernel in meters.

`minDiam = 2` is the minimum diameter that a tree could be reproductively mature, in cm.

`sigmaMu = .5` and `sigmaWt = nrow(inputs$treeData)` are the prior mean and the prior weight on log fecundity variance.

`maxF = 1e+8`, maximum fecundity, helps stabilize analysis of especially noisy data.

More detailed vignettes can be obtained with:

```
browseVignettes('mastif')
```

## Value

Returns an object of class `"mast"`, which is a list containing the following components:



inputs	list includes all inputs to the model, including transformations not resulting from posterior simulation.
chains	list of MCMC matrices, each with <code>ng</code> rows, includes: <code>bfec</code> : fecundity coefficients <code>brep</code> : maturation coefficients <code>ugibbs</code> : dispersal parameters <code>sgibbs</code> : residual variance and, after burnin, <code>rmspe</code> (root mean square prediction error) <code>rgibbs</code> : if multiple seed types, then rows are the <code>specNames</code> to <code>seedNames</code> matrix <code>R</code> . <code>bygibbsF</code> , <code>bygibbsR</code> : if <code>yearEffects</code> are included in the model, these are fixed year/lag and random year/lag effects by random group. <code>agibbs</code> : if <code>randomEffects</code> on individuals, rows are covariance matrix.
parameters	list of parameter estimates summarized from chains. <code>acfMat</code> : autocorrelation on fecundity by random group. <code>betaFec</code> : fecundity regression coefficients (log scale). <code>betaRep</code> : maturation regression coefficients (probit scale). <code>dpars</code> : dispersal kernel coefficient, by random group, on meter scale. <code>omegalist</code> : fecundity covariance between trees for the same plot-year. <code>pacfMat</code> : partial autocorrelation on fecundity. <code>pacfSe</code> : standard errors for <code>pacfMat</code> . <code>pacsMat</code> : partial autocorrelation on seed counts. <code>sigma</code> : estimate of residual log fecundity variance. <code>sigmaList</code> : fecundity covariance over years. <code>upars</code> : dispersal kernel coefficient, by random group, on meter <sup>2</sup> scale. <code>rMu</code> , <code>rSe</code> : if more than one <code>seedName</code> , posterior mean and standard error on <code>R</code> matrix If <code>yearEffects</code> , then the following will be included in <code>\$parameters</code> : <code>betaYrMu</code> , <code>betaYrSe</code> : posterior mean and standard errors on fixed year effects. <code>betaYrRand</code> , <code>betaYrRandSE</code> : posterior mean and standard errors on random year effects. <code>alphaMu</code> , <code>alphaSe</code> : posterior mean and standard errors on random tree effects. <code>aMu</code> , <code>aSe</code> : posterior mean and standard errors on random effects covariance.
prediction	list of latent variable estimates and prediction: <code>fecPred</code> includes <code>matrEst</code> , <code>fecEstMu</code> , <code>fecEstSe</code> maturation and fecundity estimates and <code>matrPred</code> , <code>fecPred</code> , <code>fecEst</code> maturation and fecundity predictions. <code>seedPred</code> includes seed predictions by species, from estimates of latent fecundity and maturation ( <code>_estMean</code> , <code>_estSe</code> ) and from the fully generative model ( <code>_predMean</code> , <code>_predSe</code> ). If <code>predictList</code> is passed to <code>mast</code> , then predictions are returned as seeds per m <sup>2</sup> (not per trap) for the regular prediction grid spaced <code>mapMeters</code> apart in the <code>data.frame</code> <code>seedPredGrid</code> . Tree predictions are returned in <code>treePredGrid</code> are included. If the AR(p) model is used ( <code>yearEffect\$p</code> is supplied), then both <code>data.frames</code> include p-yr hind casts and p-yr forecasts.

**Author(s)**

James S Clark, <jimclark@duke.edu>

**References**

Clark, J.S., C. Nunes, and B. Tomasek. 2019. Foodwebs based on unreliable foundations: spatio-temporal masting merged with consumer movement, storage, and diet. *Ecological Monographs*, e01381.

**See Also**

[mastSim](#) simulates data

A more detailed vignette is can be obtained with:

```
browseVignettes('mastif')
```

website 'http://sites.nicholas.duke.edu/clarklab/code/.'

**Examples**

```
# simulate data (see \link{\code{mastSim}})
seedNames <- specNames <- 'acerRubr'
sim <- list(nyr=10, ntree=20, nplot=5, ntrap=40,
           specNames = specNames, seedNames = seedNames)

inputs <- mastSim(sim)      # simulate data
inputs$predList <- list( mapMeters = 3, plots = inputs$plots[1],
                       years = inputs$years )
output <- mastif( inputs = inputs, ng = 3000, burnin = 2000 )
# mastPlot(output)

# for Liriodendron
d <- "https://github.com/jimclarkatduke/mast/blob/master/liriodendronExample.rData?raw=True"
repmis::source_data(d)

formulaFec <- as.formula( ~ diam ) # fecundity model
formulaRep <- as.formula( ~ diam ) # maturation model

yearEffect <- list(groups = 'species')
randomEffect <- list(randGroups = 'treeID',
                    formulaRan = as.formula( ~ 1 ) )
inputs <- list( specNames = specNames, seedNames = seedNames,
              treeData = treeData, seedData = seedData,
              xytree = xytree, xytrap = xytrap,
              yearEffect = yearEffect, randomEffect = randomEffect )
output <- mastif(inputs = inputs, formulaFec, formulaRep, ng = 1000,
                burnin = 400 )

summary(output)
# plot output:
# mastPlot(output)
```

---

mastMap	<i>Map data and predictions for mastif model</i>
---------	--

---

### Description

Maps dispersal data (trees and seed traps) with predictions.

### Usage

```
mastMap(mapList)
```

### Arguments

mapList	<p>list that includes data.frames described in the mastif help page: treeData, seedData, xytree, xytrap. Alternatively, mapList can be an object of class mastif simulated by mastSim or fitted by mast. If a class mastif object is used, it must have appended mapPlot (name of plot to map) and mapYears (years to map). These and additional objects in mapList are summarized here:</p> <p>mapPlot: character for the plot name, which appears in treeData\$plot and seedData\$plot.</p> <p>mapYears: numeric vector containing years to plot, which appear in treeData\$year and seedData\$year.</p> <p>treeSymbol: character indicating a column name in treeData holding numeric values that scale the sizes of trees in the map, e.g., diam for diameter.</p> <p>PREDICT: if TRUE, then predicted seed will be mapped. Must be predicted in mastif.</p> <p>xlim: vector with upper and lower plot limits.</p> <p>ylim: vector with upper and lower plot limits.</p> <p>MAPTRAPS: include seed traps in map.</p> <p>MAPTREES: include trees in map.</p> <p>seedMax: numeric value can be used to set maximum seed symbol/prediction contour.</p> <p>fecMax: numeric value can be used to set maximum fecundity symbol size.</p> <p>mfrow: numeric vector of length-2 to set arrangement of maps on plot, as in pars().</p> <p>LEGEND: add legend with species names to map.</p> <p>plotScale: some level of control on size of plot is possible with values &gt; 1.</p> <p>treeScale: relative scale for tree symbols, default value = 1.</p> <p>trapScale: relative scale for trap symbols, default value = 1.</p> <p>SCALEBAR: add a scale bar with distance to bottom of map.</p> <p>scaleValue: numeric value in meters used if SCALEBAR = TRUE.</p> <p>mfrow: numeric vector of length-2, giving numbers of rows and columns for plots, see help(pars).</p> <p>COLORSCALE: if TRUE, then a color scale is added for predicted seed.</p>
---------	--

**Details**

Generates of map of seed traps and trees, with symbols scaled to the sizes relative to seed counts in `sdata$seedNames` and `treeSymbol`. Sizes are adjusted with `scaleTree` and `scaleTrap`.

If `PREDICT = TRUE`, then predictions come in the object fitted in `mastif` with `predictList` used to specify prediction plots and years. See the help page for `mastif`.

More detailed vignettes can be obtained with:

```
browseVignettes('mastif')
```

**Value**

Only graphical outputs.

**Author(s)**

James S Clark, <jimclark@duke.edu>

**References**

Clark, J.S., C. Nunes, and B. Tomasek. 2019. Foodwebs based on unreliable foundations: spatio-temporal masting merged with consumer movement, storage, and diet. *Ecological Monographs*, e01381.

**See Also**

[mastSim](#) simulates data

A more detailed vignette is can be obtained with:

```
browseVignettes('mastif')
```

website `'http://sites.nicholas.duke.edu/clarklab/code/'`.

**Examples**

```
# simulate data (see \link{\code{mastSim}})
seedNames <- specNames <- 'acerRubr'
sim <- list(nyr=10, ntree=30, nplot=5,
           specNames = specNames, seedNames = seedNames)
inputs <- mastSim(sim)

inputs$mapPlot <- 'p1'
inputs$mapYears = inputs$years[1]

mastMap( inputs )

# for Pinus
d <- "https://github.com/jimclarkatduke/mast/blob/master/pinusExample.rdata?raw=True"
repmis::source_data(d)

specNames <- c("pinuEchi", "pinuRigi", "pinuStro", "pinuTaed", "pinuVirg")
seedNames <- c(specNames, "pinuUNKN")
```

```
mapList <- list( treeData = treeData, seedData = seedData,
  specNames = specNames, seedNames = seedNames,
  xytree = xytree, xytrap = xytrap, mapPlot = 'DUKE_BW',
  mapYears = c(2004:2007), treeScale = .5, trapScale=1.2,
  plotScale = 1.2, LEGEND=TRUE)
mastMap(mapList)
```

---

mastPlot

*Plot mast model*


---

### Description

Plots data fitted with `mastif` in package `mastif`.

### Usage

```
mastPlot(output, plotPars = NULL)
```

### Arguments

<code>output</code>	list of class <code>mast</code> .
<code>plotPars</code>	list of inputs for plotting can include: <code>outFolder</code> : if <code>SAVEPLOTS = TRUE</code> path to saved files. <code>SAVEPLOTS = FALSE</code> : if <code>TRUE</code> writes plots to <code>.pdf</code> files. <code>trueValues</code> : if passed from <code>mastSim</code> output then <code>trueValues</code> will be included in plots. <code>RMD</code> : if set to <code>"pdf"</code> or <code>"html"</code> writes output to R markdown file in working directory.

### Details

If `SAVEPLOTS = TRUE` plots are saved to files in `outFolder`. If `RMD = "pdf"`, output is written to a R markdown file that can be edited and knitted. Maps are not included in this option. Otherwise, plots are rendered to the screen.

More detailed vignettes can be obtained with:

```
browseVignettes('mastif')
```

### Value

Currently, there are graphical outputs.

### Author(s)

James S Clark, <jimclark@duke.edu>

## References

Clark, J.S., C. Nunes, and B. Tomasek. 2019. Foodwebs based on unreliable foundations: spatio-temporal masting merged with consumer movement, storage, and diet. *Ecological Monographs*, e01381.

## See Also

`mastSim` simulates data

A more detailed vignette is can be obtained with:

```
browseVignettes('mast')
```

website '<http://sites.nicholas.duke.edu/clarklab/code/>'.

## Examples

```
# simulate data
seedNames <- specNames <- 'acerRubr'
sim <- list(nyr=10, ntree=30, nplot=5, specNames = specNames, seedNames = seedNames)
inputs <- mastSim(sim)
output <- mastif( inputs = inputs, ng = 4000, burnin = 2000 )

# plot output
# mastPlot( output, plotPars = list(trueValues = inputs>trueValues) )
```

---

mastPriors

*Obtain prior parameter values for mastif from file*

---

## Description

Prior parameter values may be saved in a file by species or by genus. `mastPriors` looks for a species-level prior first. If not found, it can substitute a genus-level prior.

## Usage

```
mastPriors(file, specNames, code, genus = 'NULL')
```

## Arguments

<code>file</code>	character giving the path and file name where parameters are stored (see Details).
<code>specNames</code>	character vector if names in <code>treeData\$species</code> .
<code>code</code>	character name of column holding <code>specNames</code> in parameter file.
<code>genus</code>	character name of genus held in "genus" column of parameter file.

## Details

The file includes rows with genera, given in column "genus", or "species". Species rows also have an entry for genus, with the species code given in the column named code. Additional columns are names of prior parameters, including:

priorDist: mean parameter for dispersal kernel (m), related to kernel parameter  $u$  as  $d \leftarrow \pi * \sqrt{u} / 2$ .

The estimated values for these parameters are found in `output$parameters$upars` and `output$parameters$dparams`, where `output` is an object fitted by `mastif`.

minDist: the lower bound for the mean parameter  $d$  of the dispersal kernel (m).

maxDist: the upper bound for the mean parameter  $d$  of the dispersal kernel (m).

priorVDist: variance on the mean parameter for dispersal kernel ( $m^2$ ). For large values, the prior distribution of  $d$  (and by variable change,  $u$ ) becomes `dunif(d, minDist, maxDist)`.

minDiam: below this diameter trees of unknown status are assumed immature (cm).

maxDiam: above this diameter trees of unknown status are assumed mature (cm).

maxFec: maximum seeds per tree per year

More detailed vignettes can be obtained with: `browseVignettes('mastif')`

## Value

A data.frame with a row for each `specNames` and columns for prior parameter values. Where file contains species-level parameter values, they will be used. If a separate row in file holds genus-level parameters, with the entry for `code == 'NA'`, then genus-level parameters will be substituted. In other words, these genus rows are default values.

## Author(s)

James S Clark, <jimclark@duke.edu>

## References

Clark, J.S., C. Nunes, and B. Tomasek. 2019. Foodwebs based on unreliable foundations: spatio-temporal masting merged with consumer movement, storage, and diet. *Ecological Monographs*, e01381.

## See Also

[mastFillCensus](#) to fill tree census

[mastif](#) for analysis

A more detailed vignette is can be obtained with:

`browseVignettes('mastif')`

website '<http://sites.nicholas.duke.edu/clarklab/code/>'.

## Examples

```
d <- "https://github.com/jimclarkatduke/mast/blob/master/pinusExample.rdata?raw=True"
repmis::source_data(d)

# prior parameter values
pfile <- tempfile(fileext = '.txt')
d <- "https://github.com/jimclarkatduke/mast/blob/master/priorParameters.txt?raw=True"
download.file(d, destfile = pfile)

specNames <- c("pinuEchi", "pinuRigi", "pinuStro", "pinuTaed", "pinuVirg")
seedNames <- c(specNames, "pinuUNKN")

priorTable <- mastPriors(file = pfile, specNames,
                        code = 'code4', genus = 'pinus')

inputs <- list( specNames = specNames, seedNames = seedNames,
                treeData = treeData, seedData = seedData,
                xytree = xytree, xytrap = xytrap,
                priorTable = priorTable, seedTraits = seedTraits)

formulaRep <- as.formula( ~ diam )
formulaFec <- as.formula( ~ diam )

output <- mastif(inputs = inputs, formulaFec, formulaRep,
                ng = 1000, burnin = 400)
```

---

 mastSim

*Data simulation for mast model*


---

## Description

Simulates data for analysis by `mastif` in package `mastif`.

## Usage

```
mastSim(sim)
```

## Arguments

`sim` list of inputs for simulation, see Details.

## Details

The list `sim` contains the following:

`specNames`: character vector of species names.

`seedNames`: character vector of seed names.



nyr = 5: average number of years for a plot  
 ntree = 10: average number of trees in specNames on a plot  
 plotWide = 100: diameter of plot  
 nplot = 3: number of plots  
 ntrap = 20: average number of seed traps on a plot  
 meanDist = 25: mean dispersal (meters)

## Value

Returns an object of `class` "mastif", a list containing the following components:

distall	matrix of distances between all seed traps (rows) and trees (columns). Block diagonal structure by plot, padded with the value 10000 for tree/trap combinations on different plots.
distall	seed trap by tree matrix of distances.
formulaFec	formula for the fecundity model.
formulaRep	formula for the maturation model.
plots	character vector of plot names.
R	species to seed type matrix.
seedData	data.frame of seed data needed for mastif.
seedNames	character vector of seed types.
sim	inputs to mastSim.
specNames	character vector of species.
treeData	data.frame of tree data needed for mastif.
trueValues	list holding true parameter values includes: fec: vector of log fecundity states. repr: vector of maturation states. betaFec: fecundity parameter values. betaRep: maturation parameter values. upar: dispersal parameter, $m^2$ .
xytrap	data.frame of seed locations needed for mastif.
xytree	data.frame of tree locations needed for mastif.
years	numeric vector years.

## Author(s)

James S Clark, <jimclark@duke.edu>

## References

Clark, J.S., C. Nunes, and B. Tomasek. 2019. Foodwebs based on unreliable foundations: spatio-temporal masting merged with consumer movement, storage, and diet. *Ecological Monographs*, in press.

**See Also**

`mastSim` simulates data

A more detailed vignette is can be obtained with:

```
browseVignettes('mastif')
```

website `'http://sites.nicholas.duke.edu/clarklab/code/'`.

**Examples**

```
# simulate data
seedNames <- specNames <- 'acerRubr'
sim <- list(nyr = 10, ntree = 30, nplot = 5, ntrap = 40,
           specNames = specNames, seedNames = seedNames)
inputs <- mastSim(sim)
output <- mastif( inputs = inputs, ng = 500, burnin = 200 )

# increase iterations, then plot:
# output <- mastif( inputs = output, ng = 2000, burnin = 1000 )
# plot output
# mastPlot(output, plotPars = list(trueValues = inputs>trueValues) )
```

---

`mastSpectralDensity`    *Volatility and period for mast data*

---

**Description**

Extracts time series attributes for tree or population fecundity.

**Usage**

```
mastSpectralDensity( x, maxPeriod = length(x)/2, PLOT = FALSE, ylim = NULL )
```

**Arguments**

<code>x</code>	numeric vector of sequential fecundity values.
<code>maxPeriod</code>	the number of frequencies/periods to include.
<code>PLOT</code>	logical to plot spectrum.
<code>ylim</code>	if PLOT, values for lower and upper bounds.

**Details**

Returns attributes of volatility and period for a sequence of fecundity values for a single tree (or population) in `x`, which may often be on a log scale.

More detailed examples can be obtained with: `browseVignettes('mastif')`

**Value**

Returns a list that includes `spect`, a matrix of power values ordered by frequency (1/period). To permit comparisons between series that differ in length, `totVar` (total variance) and `volatility` (period-weighted variance) are divided by the length of the series. Mean and standard deviation for the weighted period are `periodMu` and `periodSd`.

**Author(s)**

James S Clark, <jimclark@duke.edu>

**References**

Qiu, T, ..., and J.S. Clark. 2023. Mutualist dispersers and the global distribution of masting: mediation by climate and fertility. in review.

**See Also**

[mastif](#) for analysis

A more detailed vignette is can be obtained with:

```
browseVignettes('mastif')
website 'http://sites.nicholas.duke.edu/clarklab/code/'.
```

**Examples**

```
d <- "https://github.com/jimclarkatduke/mast/blob/master/outputAbies.rdata?raw=True"
repmis::source_data( d )

# single Abies tree from fitted output$prediction$fecPred:

wt <- which( fecPred$treeID == "BAMT1-1" )
s <- mastSpectralDensity( log(fecPred$fecEstMu[ wt ]), PLOT = TRUE )

# population year effects (log scale) for an ecoRegion_species in output$parameters$betaYrRand:

x <- betaYrRand['3_abiesAmabilis', ]
x <- x[ x != 0 ] # ecoRegion_species vary in observation years
s <- mastSpectralDensity( x, PLOT = TRUE )
```

---

mastVolatility

*Volatility and period for mast data, combining trees in populations*

---

**Description**

Synthesis of volatility and period at the population scale.

**Usage**

```
mastVolatility( treeID, year, fec, minLength = 6, minFrequency = 1/20 )
```

**Arguments**

treeID	vector of labels for individual trees
year	numeric vector for year
fec	numeric vector for fecundity values
minLength	determines the minimum number of years to a tree to be included in population estimates
minFrequency	lowest frequency to include in volatility, period evaluation

**Details**

The three vectors `treeID`, `year`, `fec` are aligned by tree and year and, thus, of the same length. Tree fecundity values in the numeric vector `fec` can differ in number of years due to maturation times, deaths, and observation years. Trees having fewer than `minLength` observations are omitted from the analysis. `minFrequency` is high enough to omit low frequencies that are missing in the shortest series to be compared.

More detailed examples can be obtained with: `browseVignettes('mastif')`

**Value**

Returns a list that includes `stats`, which holds the period- and fecundity-weighted estimates of volatility and period at the population scale. The matrix `statsDensity` holds the means and standard deviations by period (1/frequency). The matrix `mastMatrix` holds for each tree the number of years, mean log fecundity, variance, volatility, and period mean and standard deviation. Returned as tree by frequency are density and frequency.

**Author(s)**

James S Clark, <jimclark@duke.edu>

**References**

Qiu, T, ..., and J.S. Clark. 2023. Mutualist dispersers and the global distribution of masting: mediation by climate and fertility. in review.

**See Also**

[mastif](#) for analysis

A more detailed vignette is can be obtained with:

```
browseVignettes('mastif')
```

website `'http://sites.nicholas.duke.edu/clarklab/code/'`.

**Examples**

```
d <- "https://github.com/jimclarkatduke/mast/blob/master/outputAbies.rdata?raw=True"
repmis::source_data( d )

# all trees in a plot:

wi <- which( fecPred$plotSpec == 'BERK28 abiesGrandis' ) # tree-years in group
tmp <- mastVolatility( treeID = fecPred$treeID[wi], year = fecPred$year[wi],
                     fec = fecPred$fecEstMu[wi], minLength = 10 )
period <- 1/tmp$frequency
density <- tmp$density

plot( NA, xlim = range( period, na.rm = TRUE ), ylim = range( density, na.rm = TRUE ),
      xlab = 'Period (yr)', ylab = 'Density', log = 'xy' )

for( i in 1:nrow(density) )lines( period[i,], density[i, ], col = 'grey' )
lines( tmp$statsDensity['Period', ], tmp$statsDensity['Mean', ], lwd = 2 )
```

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