Package 'synlik'

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Type Package Title Synthetic Likelihood Methods for Intractable Likelihoods Version 0.1.2 Date 2018-05-22 Author Matteo Fasiolo and Simon Wood Maintainer Matteo Fasiolo <matteo.fasiolo@gmail.com> **Description** Framework to perform synthetic likelihood inference for models where the likelihood function is unavailable or intractable. URL http://mfasiolo.github.io/synlik_release License GPL (>= 2) **Depends** R (>= 2.10), Rcpp (>= 0.12.0) Imports methods, graphics, Matrix, compiler, stats, parallel Suggests knitr, stabledist LinkingTo Rcpp, RcppArmadillo VignetteBuilder knitr RoxygenNote 5.0.1 NeedsCompilation yes **Repository** CRAN

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Synthetic Likelihood Methods for Intractable Likelihoods

Description

Package that provides Synthetic Likelihood methods for intractable likelihoods. The package is meant to be as general purpose as possible: as long as you are able to simulate data from your model you should be able to fit it.

Details

Package:	synlik
Type:	Package
Version:	0.1.2
Date:	2018-05-22
License:	GPL (>=2)

The package allows users to create objects of class synlik (S4), which are essentially constituted of a simulator function and a function (summaries) that transforms the data into summary statistics. The simulator can output any kind of data (vector, list, etc) and this will be passed directly to the summaries function. This allow the package to fit a large variety of models.

Once the model of interest has been set up as a synlik object, it is possible several methods on it. The function most useful function is slik, which can be used to evaluate the synthetic likelihood. The slice.synlik function allows to obtain and plot slices of the synthetic likelihood with respect

synlik-package

to model parameters. It is possible to simulate data or statistics from the model using the generic simulate, and to check the normality of the statistics using the checkNorm function. Unknow parameters can be estimated by MCMC, through the smcmc function. This function will return an object of class smcmc (S4), which contains all the inputs and results of the MCMC procedure.

Many functions in the package support parallel simulation on multiple cores.

Author(s)

Matteo Fasiolo and Simon N. Wood

Maintainer: Matteo Fasiolo <matteo.fasiolo@gmail.com>

References

Simon N Wood. Statistical inference for noisy nonlinear ecological dynamic systems. Nature, 466(7310):1102–1104, 2010.

See Also

For some examples see the Vignettes (type vignette("synlik")).

```
## Not run:
#### Here I put a simple example,
#### if you want to see more type: vignette("synlik")
## End(Not run)
#### Create synlik object
ricker_sl <- synlik(simulator = rickerSimul,
                    summaries = rickerStats,
                    param = c(logR = 3.8, logSigma = log(0.3), logPhi = log(10)),
                    extraArgs = list("nObs" = 50, "nBurn" = 50),
                    plotFun = function(input, ...){
                           plot(drop(input), type = 'l', ylab = "Pop", xlab = "Time", ...)
                             }
)
#### Simulate from the object
ricker_sl@data <- simulate(ricker_sl)</pre>
ricker_sl@extraArgs$obsData <- ricker_sl@data</pre>
#### Simulate statistics (each row is a vector of statistics)
simulate(ricker_sl, seed = 523, nsim = 10, stats = TRUE)
#### Plotting the data
plot(ricker_sl)
#### Checking multivariate normality of the statistics
```

```
checkNorm(ricker_sl)
#### Evaluate the likelihood
set.seed(4234)
slik(ricker_sl,
     param = c(logR = 3.8, logSigma = log(0.3), logPhi = log(10)),
           = 1e3)
     nsim
#### Plotting a slice of the log-Likelihood possibly using multiple cores
slice(object = ricker_sl,
      ranges = list("logR" = seq(3.5, 3.9, by = 0.02),
                    "logPhi" = seq(2, 2.6, by = 0.02),
                    "logSigma" = seq(-2, -0.5, by = 0.05)),
      param = c(logR = 3.8, logSigma = log(0.3), logPhi = log(10)),
      nsim = 500, multicore = FALSE)
#### MCMC estimation possibly using multiple cores
set.seed(4235)
ricker_sl <- smcmc(ricker_sl,</pre>
                   initPar = c(3.2, -1, 2.6),
                   niter = 50,
                   burn = 3,
                   priorFun = function(input, ...) 0,
                   propCov = diag(c(0.1, 0.1, 0.1))^2,
                   nsim = 1e3,
                   multicore = FALSE)
# Continue with additional 50 iterations
ricker_sl <- continue(ricker_sl, niter = 50)</pre>
# Plotting results on transformed scale (exponential)
trans <- rep("exp", 3)</pre>
names(trans) <- names(ricker_sl@param)</pre>
plot(ricker_sl)
```

ANYOrNULL-class Dummy class

Description

Class unions for internal use only

Description

Data from figures 3 and 4 of Nicholson, 1954.

Usage

data(bf1)

Arguments

bf1 the dataset name

Details

bf1 is Nisbet and Gurney's run 1, and Nicholson's (1954) figure 3 (adult food limitation). The data are actually from the global population dynamics database at Silwood. They are daily: Nicholson's figure 3 plots data every other day, but the text says that measurements were taken daily. However elsewhere they are reported every other day. Probably best to assume that they have been interpolated to daily.

bf2 and bf3 are digitized from Nicholson's (1954) figure 4. bf2 is the upper series: larval food limitation, with 50g per day of larval food provided. bf3 is the lower series: same set up, half as much food. These are Nisbet and Gurney's runs 2 and 3, respectively.

Value

matrix of replicate data series

Author(s)

Simon N. Wood, maintainer Matteo Fasiolo <matteo.fasiolo@@gmail.com>

References

Alexander J Nicholson. An outline of the dynamics of animal populations. Australian Journal of Zoology, 2(1):9–65, 1954.

See Also

blowfly

bf

Examples

```
library(synlik)
data(bf1)
data(bf2)
data(bf3)
par(mfrow=c(3,1),mar=c(4,4,1,1))
with(bf1,plot(day,pop,type="1"))
with(bf1,points(day,pop,pch=20,cex=.8))
abline(mean(bf1$pop),0,col=2); abline(median(bf1$pop),0,col=3);
with(bf2,plot(day,pop,type="1"))
with(bf2,points(day,pop,pch=20,cex=.8))
abline(mean(bf2$pop),0,col=2); abline(median(bf2$pop),0,col=3);
with(bf3,plot(day,pop,type="1"))
with(bf3,plot(day,pop,type="1"))
with(bf3,points(day,pop,pch=20,cex=.8))
abline(mean(bf3$pop),0,col=2); abline(median(bf3$pop),0,col=3);
```

blowSimul

Simulates from the blowfly model

Description

Simulator for the blowfly model proposed by Wood (2010).

Usage

```
blowSimul(param, nsim, extraArgs, ...)
```

Arguments

param	vector of log-parameters: delta, P, N0, var.p, tau and var.d. The interpretation of these parameters is described in Wood (2010).
nsim	Number of simulations from the model.
extraArgs	A named list of additional arguments:
	• n0bs = Length of each simulated time series.
	• nBurn = Number of initial steps to be discarded before saving the following nObs steps.
	• steps = Positive integer. If steps == n the observations correspond to n time steps.
	Need for compatibility with synlik, but not used.

Value

A matrix nsim by n0bs, where each row is a simulated path.

Author(s)

Simon Wood and Matteo Fasiolo <matteo.fasiolo@gmail.com>.

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blow_sl

References

Simon N Wood. Statistical inference for noisy nonlinear ecological dynamic systems. Nature, 466(7310):1102–1104, 2010.

Brillinger, D. R., J. Guckenheimer, P. Guttorp, and G. Oster. 1980. Empirical modelling of population time series data: the case of age and density dependent vital rates. Lectures on Mathematics in the Life Sciences13:65-90.

Nicholson, A. J. 1957. The self-adjustment of populations to change. Cold Spring Harbor Symposia on Quantitative Biology22:153-173.

See Also

blow_sl

Examples

blow_sl

Blowfly model

Description

synlik object containing the blowfly model proposed by Wood (2010). The main components are the simulator blowSimul and the statistics blowStats, described in the same reference.

Author(s)

Simon Wood and Matteo Fasiolo <matteo.fasiolo@gmail.com>.

References

Simon N Wood. Statistical inference for noisy nonlinear ecological dynamic systems. Nature, 466(7310):1102–1104, 2010.

See Also

blowSimul

Examples

```
data(blow_sl)
plot(blow_sl)
simulate(blow_sl, stats = TRUE)
slik(blow_sl,
     param = log( c( "delta" = 0.16, "P" = 6.5, "N0" = 400,
                      "var.p" = 0.1, "tau" = 14, "var.d" = 0.1) ),
     nsim
             = 1e3)
# Using Nicholson's data
data(bf1)
plot(blow_sl)
blow_sl@data <- bf1$pop</pre>
blow_sl@extraArgs$obsData <- bf1$pop #Important: blow_sl@blowStats uses the observed data</pre>
slik(blow_sl,
     param = log( c( "delta" = 0.16, "P" = 6.5, "N0" = 400,
                      "var.p" = 0.1, "tau" = 14, "var.d" = 0.1) ),
     nsim
             = 1e3)
```

checkNorm

Checking the multivariate normal approximation.

Description

Given an object of class synlik this routine provides a graphical check of whether the distribution of the random summary statistics is multivariate normal.

Usage

```
checkNorm(object, param = object@param, nsim = 1000, observed = NULL,
    cex.axis = 1, cex.lab = 1, ...)
```

Arguments

object	An object of class synlik or a matrix where each row is a random vector.
param	A vector of model's parameters at which the summary statistics will be simulated.
nsim	number of summary statistics to be simulated if object is of class synlik, otherwise it is not used.
observed	A vector of observed summary statistics. By default NULL, so object@data will be used as observed statistics. It will be looked at only if object is a matrix.
cex.axis	Axis scale expansion factor.

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checkNorm

cex.lab	Axis label expansion factor.
	additional arguments to be passed to object@simulator and object@summaries In general I would avoid using it and including in object@extraArgs every- thing they need.

Details

The method is from section 7.5 of Krzanowski (1988). The replicate vectors of summary statistic S are transformed to variables which should be univariate chi squared r.v.s with DoF given by the number of rows of S. An appropriate QQ-plot is produced, and the proportion of the data differing substantially from the ideal line is reported. Deviations at the right hand end of the plot indicate that the tail behaviour of the Normal approximation is poor: in the context of synthetic likelihood this is of little consequence. Secondly, s is transformed to a vector which should be i.i.d. N(0,1) under multivariate normality, and a QQ plot is produced. Unfortunately this approach is not very useful unless the dimension of s is rather large. In simulations, perfectly MVN data produce highly variable results, so that the approach lacks any real power.

Value

Mainly produces plots and prints output. Also an array indicating proportion of simulated statistics smaller than observed.

Author(s)

Simon N. Wood, maintained by Matteo Fasiolo <matteo.fasiolo@gmail.com>.

References

Krzanowski, W.J. (1988) Principles of Multivariate Analysis. Oxford.

```
#### Create Object
data(ricker_sl)
```

```
#### Simulate from the object
ricker_sl@data <- simulate(ricker_sl)
ricker_sl@extraArgs$obsData <- ricker_sl@data</pre>
```

```
#### Checking multivariate normality
checkNorm(ricker_sl)
```

```
# With matrix input
checkNorm(matrix(rnorm(200), 100, 2))
```

continue

Description

Generic function, that given the results of an estimation procedure (ex. MCMC or maximum likelihood optimization) continues the procedure for some more iterations.

Usage

```
continue(object, ...)
## S4 method for signature 'smcmc'
continue(object, niter = object@niter, nsim = object@nsim,
    propCov = object@propCov, targetRate = object@targetRate,
    recompute = object@recompute, multicore = object@multicore,
    ncores = object@ncores, cluster = NULL, control = object@control, ...)
```

Arguments

object	An object representing the results of an estimation procedure which we wish to continue. For example it might represents an MCMC chain.
	additional arguments to be passed to slik function, see slik.
niter	see smcmc-class.
nsim	see smcmc-class.
propCov	see smcmc-class.
targetRate	see smcmc-class.
recompute	see smcmc-class.
multicore	see smcmc-class.
ncores	see smcmc-class.
cluster	an object of class c("SOCKcluster", "cluster"). This allowes the user to pass her own cluster, which will be used if multicore == TRUE. The user has to remember to stop the cluster.
control	see smcmc-class.

Details

When is("smcmc", object) == TRUE continues MCMC estimation of an object of class smcmc. All input parameters are defaulted to the corresponding slots in the input object, with the exception of cluster. The chain restarts were it ended, burn-in is set to zero, the same prior (if any) is used.

Value

An object of the same class as object, where the results of the estimation have been updated.

extractCorr

See Also

For examples, see smcmc-class.

extractCorr

Extracting correlations from a covariance matrix

Description

Extracting correlations from a covariance matrix

Usage

extractCorr(mat)

Arguments

mat A covariance matrix.

Value

The correlation matrix embedded in mat.

Examples

```
# 2 dimensional case
d <- 2
tmp <- matrix(rnorm(d^2), d, d)
mcov <- tcrossprod(tmp, tmp)
# Covariance matrix
mcov
# Correlation matrix
extractCorr(mcov)
```

functionOrNULL-class Dummy class

Description

Class unions for internal use only

internal_C

Description

This functions are for internal use only.

Author(s)

Simon Wood and Matteo Fasiolo <matteo.fasiolo@gmail.com>.

nlar

Estimate non-linear autoregressive coefficients

Description

Function that, give time series data, transforms them into summary statistics using polynomial autoregression.

Usage

nlar(x, lag, power)

Arguments

х	a matrix. Each column contains a replicate series.
lag	vector of lags, for rhs terms.
power	vector of powers, for rhs terms.

Value

a matrix where each column contains the coefficients for a different replicate.

Author(s)

Simon N. Wood, maintainer Matteo Fasiolo <matteo.fasiolo@gmail.com>.

Examples

```
library(synlik)
set.seed(10)
x <- matrix(runif(200),100,2)</pre>
beta <- nlar(x,lag=c(1,1),power=c(1,2))</pre>
y <- x[,1]
y <- y - mean(y)
z <- y[1:99];y <- y[2:100]
lm(y^{z+I}(z^{2})-1)
beta
## NA testing
x[5,1] <- x[45,2] <- NA
beta <- nlar(x,lag=c(1,1),power=c(1,2))</pre>
y <- x[,1]
y <- y - mean(y,na.rm=TRUE)</pre>
z <- y[1:99];y <- y[2:100]
lm(y~z+I(z^2)-1)
beta
## higher order...
set.seed(10)
x <- matrix(runif(100),100,2)</pre>
beta <- nlar(x,lag=c(6,6,6,1,1),power=c(1,2,3,1,2))</pre>
k <- 2
y <- x[,k]
y <- y - mean(y)
ind <- (1+6):100
y6 <- y[ind-6];y1 <- y[ind-1];y <- y[ind]</pre>
beta0 <- coef(lm(y~y6+I(y6^2)+I(y6^3)+y1+I(y1^2)-1))</pre>
as.numeric(beta[,k]);beta0;beta0-as.numeric(beta[,k])
```

numericOrNULL-class Dummy class

Description

Class unions for internal use only

orderDist

Summarize marginal distribution of (differenced) series.

Description

Summarizes (difference) distribution of replicate series, by regressing ordered differenced series on a reference series (which might correspond to observed data).

Usage

orderDist(x, z, np = 3, diff = 1)

Arguments

х	a matrix. Each column contains a replicate series.
Z	vector of lags, for rhs terms.
np	maximum power on rhs of regression.
diff	order of differencing (zero for none).

Value

a matrix where each column contains the coefficients for a different replicate.

Author(s)

Simon N. Wood, maintainer Matteo Fasiolo <matteo.fasiolo@gmail.com>.

Examples

```
library(synlik)
set.seed(10)
n <- 100;nr <- 3
x <- matrix(runif(n*nr),n,nr)
z <- runif(n)
beta <- orderDist(x,z,np=3,diff=1)
zd <- z;xd <- x[,3]
zd <- diff(zd,1);xd <- diff(xd,1)
zd <- sort(zd);zd <- zd - mean(zd)
xd <- sort(xd);xd <- xd - mean(xd)
lm(xd~zd+I(zd^2)+I(zd^3)-1)</pre>
```

plot-smcmc

Plotting objects of class smcmc.

Description

Method for plotting an object of class smcmc.

Usage

```
## S4 method for signature 'smcmc,missing'
plot(x, trans = NULL, addPlot1 = NULL,
    addPlot2 = NULL, ...)
```

plot-synlik

Arguments

х	An object of class smcmc.
trans	Name list or vector containing names of transforms for some parameters (ex: list("par1" = "exp", "par2" = "log")). The transformations will be applied before plotting.
addPlot1	Name of additional plotting function that will be call after the MCMC chain have been plotted. It has to have prototype fun(nam,) where nam will be the parameter name. See "examples".
addPlot2	Name of additional plotting function that will be call after the histograms have been plotted. It has to have prototype fun(nam,) where nam will be the parameter name. See "examples".
	additional arguments to be passed to the plotting functions.

See Also

smcmc-class, plot.

Examples

plot-synlik

Method for plotting an object of class synlik.

Description

It basically calls the slot dject@plotFun with input dject@data, if it has been provided by the user. Otherwise it tries to use the plot(x = object@data, y, ...) generic.

Usage

```
## S4 method for signature 'synlik,missing'
plot(x, y, ...)
```

Arguments

х	An object of class synlik.
У	Useless argument, only here for compatibility reasons.
	additional arguments to be passed to object@plotFun.

Author(s)

Matteo Fasiolo <matteo.fasiolo@gmail.com>

See Also

synlik-class, plot.

Examples

```
data(ricker_sl)
```

Using ricker_sl@plotFun
plot(ricker_sl)

```
# Using generic plot, doesn't work well because object@data is a matrix.
ricker_sl@plotFun <- NULL
plot(ricker_sl)
```

rickerSimul

Simulates from the ricker model

Description

Simulator for the stochastic Ricker model, as described by Wood (2010). The observations are $Y_t \sim Pois(Phi * N_t)$, and the dynamics of the hidden state are given by $N_t = r * N_{t-1} * exp(-N_{t-1} + e_t)$, where $e_t \sim N(0, Sigma^2)$.

Usage

```
rickerSimul(param, nsim, extraArgs, ...)
```

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rickerSimul

Arguments

param	vector of log-parameters: logR, logSigma, logPhi. Alternatively a matrix nsim by 3 were each row is a different parameter vector.	
nsim	Number of simulations from the model.	
extraArgs	A named list of additional arguments:	
	 nObs = Length of each simulated time series. nBurn = Number of initial steps to be discarded before saving the following nObs steps. randInit = if TRUE (default) the initial state NO is runif(0, 1), otherwise it is equal to extraArgs\$initVal. initVal = initial value NO, used only if extraArgs\$randInit == TRUE. 	
	Need for compatibility with synlik, but not used.	

Value

A matrix nsim by n0bs, where each row is a simulated path.

Author(s)

Simon Wood and Matteo Fasiolo <matteo.fasiolo@gmail.com>.

References

Simon N Wood. Statistical inference for noisy nonlinear ecological dynamic systems. Nature, 466(7310):1102–1104, 2010.

See Also

ricker_sl

ricker_sl

Description

ricker_sl is synlik object containing the stochastic Ricker model, ricker_smcmc is a smcmc object which also contains the results of some MCMC iterations. The model is described rickerSimul and in Wood (2010). The main components of the object are the simulator rickerSimul and the statistics rickerStats, described in the same reference.

Author(s)

Simon Wood and Matteo Fasiolo <matteo.fasiolo@gmail.com>.

References

Simon N Wood. Statistical inference for noisy nonlinear ecological dynamic systems. Nature, 466(7310):1102–1104, 2010.

See Also

rickerSimul

```
data(ricker_sl)
plot(ricker_sl)
simulate(ricker_sl, stats = TRUE)
slik(ricker_sl,
    param = c( logR = 3.8, logSigma = log(0.3), logPhi = log(10) ),
    nsim = 1e3)
# Using Nicholson's data
data(ricker_smcmc)
plot(ricker_smcmc)
```

robCov

Description

Obtains a robust estimate of the covariance matrix of a sample of multivariate data, using Campbell's (1980) method as described on p231-235 of Krzanowski (1988).

Usage

robCov(sY, alpha = 2, beta = 1.25)

Arguments

sY	A matrix, where each column is a replicate observation on a multivariate r.v.
alpha	tuning parameter, see details.
beta	tuning parameter, see details.

Details

Campbell (1980) suggests an estimator of the covariance matrix which downweights observations at more than some Mahalanobis distance d.0 from the mean. d.0 is sqrt(nrow(sY))+alpha/sqrt(2). Weights are one for observations with Mahalanobis distance, d, less than d.0. Otherwise weights are d.0*exp(-.5*(d-d.0)^2/beta)/d. The defaults are as recommended by Campbell. This routine also uses pre-conditioning to ensure good scaling and stable numerical calculations.

Value

A list where:

- Ea square root of the inverse covariance matrix. i.e. the inverse cov matrix is t(E)%*%E;
- half.ldet.VHalf the log of the determinant of the covariance matrix;
- mYThe estimated mean;
- sdThe estimated standard deviations of each variable.

Author(s)

Simon N. Wood, maintained by Matteo Fasiolo <matteo.fasiolo@gmail.com>.

References

Krzanowski, W.J. (1988) Principles of Multivariate Analysis. Oxford. Campbell, N.A. (1980) Robust procedures in multivariate analysis I: robust covariance estimation. JRSSC 29, 231-237.

```
p <- 5;n <- 100
Y <- matrix(runif(p*n),p,n)
robCov(Y)</pre>
```

simulate,synlik-method

Simulate data or statistics from an object of class synlik.

Description

Simulate data or statistics from an object of class synlik.

Usage

```
## S4 method for signature 'synlik'
simulate(object, nsim, seed = NULL, param = object@param,
   stats = FALSE, clean = TRUE, verbose = TRUE, ...)
```

Arguments

object	An object of class synlik.
nsim	Number of simulations from the model.
seed	Random seed to be used. It is not passed to the simulator, but simply passed to set.seed() from within simulate.synlik.
param	Vector of parameters passed to object@simulator.
stats	If TRUE the function trasforms the simulated data into statistics using object@summaries.
clean	If TRUE the function tries to clean the statistics from NaNs or non-finite values. Given that object@summaries has to returns a numeric vector or a matrix where each row is a simulation, rows containing non-finite values will be discarded.
verbose	If TRUE the function will complain if, for instance, the simulations contain lots of non-finite values.
	additional arguments to be passed to object@simulator and object@summaries. In general I would avoid using it and including object@extraArgs everything they need.

Value

If stats == FALSE the output will that of object@simulator, which depends on the simulator used by the user. If stats == TRUE the output will be a matrix where each row is vector of simulated summary statistics.

Author(s)

Matteo Fasiolo <matteo.fasiolo@gmail.com>

See Also

synlik-class, simulate.

slAcf

Examples

```
data(ricker_sl)
# Simulate data
simulate(ricker_sl, nsim = 2)
# Simulate statistics
simulate(ricker_sl, nsim = 2, stats = TRUE)
```

slAcf

Estimate auto-covariances for multiple datasets.

Description

Function that, give time series data, transforms them into auto-covariances with different lags.

Usage

slAcf(x, max.lag = 10)

Arguments

х	a matrix. Each column contains a replicate series.
max.lag	How many lags to use.

Value

a matrix where each column contains the coefficients for a different replicate. The first coefficient corresponds to lag == 0, hence it is the variance, the second is the covariance one step ahead and so on.

Author(s)

Simon N. Wood, maintainer Matteo Fasiolo <matteo.fasiolo@gmail.com>.

```
library(synlik)
set.seed(10)
x <- matrix(runif(1000),100,10)
acf <- slAcf(x)</pre>
```

Description

Plot slices of the synthetic log-likelihood.

Usage

```
slice(object, ranges, nsim, param = object@param, pairs = FALSE,
draw = TRUE, trans = NULL, multicore = FALSE, ncores = detectCores() -
1, cluster = NULL, ...)
```

Arguments

object	synlik object.
ranges	ranges of values along which we want the slices. If length(parName) == 1 than range has a vector, while if length(parName) == 2 it have to be a named list of 2 vectors (ex: list("alpha" = 1:10, "beta" = 10:1)).
nsim	Number of simulations used to evaluate the synthetic likelihood at each location.
param	Named vector containing the value of the ALL parameters (including the sliced one). Parameters that are not in parName will be fixed to the values in param.
pairs	if TRUE the function will produce a 2D slice for every pair of parameters in ranges. FALSE by default.
draw	If TRUE the slice will be plotted.
trans	Named vector or list of transformations to be applied to the parameters in parName before plotting ex: trans = c(s = "exp", d = "exp")/
multicore	If TRUE the object@simulator and object@summaries functions will be exe- cuted in parallel. That is the nsim simulations will be divided in multiple cores.
ncores	Number of cores to use if multicore == TRUE.
cluster	An object of class c("SOCKcluster", "cluster"). This allowes the user to pass her own cluster, which will be used if multicore == TRUE. The user has to remember to stop the cluster.
	additional arguments to be passed to slik(), see slik.

Value

Either a vector or matrix of log-synthetic likelihood estimates, depending on whether length(parNames) == 1 or 2. These are returned invisibly.

Author(s)

Matteo Fasiolo <matteo.fasiolo@gmail.com>

slice

Examples

slik

```
data(ricker_sl)
# Plotting slices of the logLikelihood
slice(object = ricker_sl,
     ranges = list("logR" = seq(3.5, 3.9, by = 0.01),
                   "logPhi" = seq(2, 2.6, by = 0.01),
                    "logSigma" = seq(-2, -0.5, by = 0.01)),
     param = c(logR = 3.8, logSigma = log(0.3), logPhi = log(10)),
     nsim = 500)
## Not run:
# Plotting a contour of the logLikelihood
slice(object = ricker_sl,
     ranges = list("logR" = seq(3.5, 3.9, by = 0.01),
                   "logPhi" = seq(2, 2.6, by = 0.01),
                   "logSigma" = seq(-2, -0.5, by = 0.04)),
     pairs = TRUE,
    param = c(logR = 3.8, logSigma = log(0.3), logPhi = log(10)),
    nsim = 500, multicore = TRUE)
```

End(Not run)

slik

Evaluates the synthetic log-likelihood.

Description

Evaluates the synthetic log-likelihood.

Usage

```
slik(object, param, nsim, multicore = FALSE, ncores = detectCores() - 1,
cluster = NULL, ...)
```

Arguments

object	An object of class synlik.
param	Vector of parameters at which the synthetic likelihood will be evaluated.
nsim	Number of simulation from the model.
multicore	(logical) if TRUE the object@simulator and object@summaries functions will be executed in parallel. That is the nsim simulations will be divided in multiple cores.
ncores	(integer) number of cores to use if multicore == TRUE.
cluster	an object of class c("SOCKcluster", "cluster"). This allowes the user to pass her own cluster, which will be used if multicore == TRUE. The user has to remember to stop the cluster.

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smcmc

additional arguments to be passed to object@simulator and object@summaries. In general I would avoid using it and including object@extraArgs everything they need.

Value

. . .

The estimated value of the synthetic log-likelihood at param.

Author(s)

Matteo Fasiolo <matteo.fasiolo@gmail.com>

References

Simon N Wood. Statistical inference for noisy nonlinear ecological dynamic systems. Nature, 466(7310):1102–1104, 2010.

Examples

```
data(ricker_sl)
set.seed(643)
slik(ricker_sl, param = c(3.8, -1.2, 2.3), nsim = 500)
```

SMCMC

MCMC parameter estimation for objects of class synlik.

Description

MCMC parameter estimation for objects of class synlik.

Usage

```
smcmc(object, initPar, niter, nsim, propCov, burn = 0,
priorFun = function(param, ...) 0, targetRate = NULL, recompute = FALSE,
multicore = !is.null(cluster), cluster = NULL, ncores = detectCores() -
1, control = list(), ...)
```

Arguments

object	An object of class synlik.
initPar	see smcmc-class.
niter	see smcmc-class.
nsim	see smcmc-class.
propCov	see smcmc-class.
burn	see smcmc-class.
priorFun	see smcmc-class.

smcmc-class

targetRate	see smcmc-class.
recompute	see smcmc-class.
multicore	see smcmc-class.
cluster	an object of class c("SOCKcluster", "cluster"). This allowes the user to pass her own cluster, which will be used if multicore == TRUE. The user has to remember to stop the cluster.
ncores	see smcmc-class.
control	see smcmc-class.
	additional arguments to be passed to slik function, see slik.

Value

An object of class smcmc.

Author(s)

Matteo Fasiolo <matteo.fasiolo@gmail.com>, code for adaptive step from the adaptMCMC package.

References

Vihola, M. (2011) Robust adaptive Metropolis algorithm with coerced acceptance rate. Statistics and Computing.

smcmc-class smcmc-class

Description

Object representing the results of MCMC estimation on an object of class synlik, from which it inherits.

Slots

initPar Vector of initial parameters where the MCMC chain will start (numeric).

niter Number of MCMC iterations (integer).

- nsim Number of simulations from the simulator at each step of the MCMC algorithm (integer).
- burn Number of initial MCMC iterations that are discarded (integer).
- **priorFun** Function that takes a vector of parameters as input and the log-density of the prior as output. If the output is not finite the proposed point will be discarded. (function). The function needs to have signature fun(x, ...), where x represents the input parameters (function).
- **propCov** Matrix representing the covariance matrix to be used to perturb the parameters at each step of the MCMC chain (matrix).

- **targetRate** Target rate for the adaptive MCMC sampler. Should be in (0, 1), default is NULL (no adaptation). The adaptation uses the approach of Vihola (2011). (numeric)
- **recompute** If TRUE the synthetic likelihood will be evaluated at the current and proposed positions in the parameter space (thus doubling the computational effort). If FALSE the likelihood of the current position won't be re-estimated (logical).
- **multicore** If TRUE the object@simulator and object@summaries functions will be executed in parallel. That is the nsim simulations will be divided in multiple cores (logical).

ncores Number of cores to use if multicore == TRUE (integer).

accRate Acceptance rate of the MCMC chain, between 0 and 1 (numeric).

- **chains** Matrix of size niter by length(initPar) where the i-th row contains the position of the MCMC algorithm in the parameter space at the i-th (matrix).
- **llkChain** Vector of niter elements where the i-th element is contains the estimate of the synthetic likelihood at the i-th iteration (numeric).

control Control parameters used by the MCMC sampler:

- theta = controls the speed of adaption. Should be between 0.5 and 1. A lower gamma leads to faster adaption.
- adaptStart = iteration where the adaption starts. Default 0.
- adaptStop = iteration where the adaption stops. Default burn + niter
- saveFile = path to the file where the intermediate results will be stored (ex: "~/Res.RData").
- saveFreq = frequency with which the intermediate results will be saved on saveFile. Default 100.
- verbose = if TRUE intermediate posterior means will be printer.
- verbFreq = frequency with which the intermediate posterior means will be printer. Default 500.

Author(s)

Matteo Fasiolo <matteo.fasiolo@gmail.com>

References

Vihola, M. (2011) Robust adaptive Metropolis algorithm with coerced acceptance rate. Statistics and Computing.

synlik-class

```
priorFun = function(input, ...) 1,
propCov = diag( c(0.1, 0.1, 0.1) )^2,
nsim = 200,
multicore = FALSE)
# Continue with additional 50 iterations
ricker_sl <- continue(ricker_sl, niter = 50)</pre>
```

```
plot(ricker_sl)
```

synlik-class synlik-class

Description

Basic class for simulation-based approximate inference using Synthetic Likelihood methods.

Usage

synlik(...)

Arguments

... See section "Slots".

Slots

param Named vector of parameters used by object@simulator (numeric).

- simulator Function that simulates from the model (function). It has to have prototype fun(param, nsim, extraArgs, ... If summaries() is not specified the simulator() has output a matrix with nsim rows, where each row is a vector of simulated statistics. Otherwise it can output any kind of object, and this output will be passed to summaries().
- **summaries** Function that transforms simulated data into summary statistics (function). It has to have prototype fun(x, extraArgs, ...) and it has to output a matrix with nsim rows, where each row is a vector of simulated statistics. Parameter x contains the data.
- data Object containing the observed data or statistics (ANY).
- **extraArgs** List containing all the extra arguments to be passed to object@simulator and object@summaries (list).
- plotFun Function that will be used to plot object@data. Prototype should be fun(x, ...)
 (function).

Author(s)

Matteo Fasiolo <matteo.fasiolo@gmail.com>

References

Simon N Wood. Statistical inference for noisy nonlinear ecological dynamic systems. Nature, 466(7310):1102–1104, 2010.

```
plot(ricker_sl)
```

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