

Package ‘EMC’

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R topics documented:

evolMonteCarlo	2
findMaxTemper	6
MetropolisHastings	11
parallelTempering	13
placeTempers	17
print	21
randomWalkMetropolis	22
utilsForExamples	25
Index	26

evolMonteCarlo *evolutionary Monte Carlo algorithm*

Description

Given a multi-modal and multi-dimensional target density function, a (possibly asymmetric) proposal distribution and a temperature ladder, this function produces samples from the target using the evolutionary Monte Carlo algorithm.

Below `sampDim` refers to the dimension of the sample space, `temperLadderLen` refers to the length of the temperature ladder, and `levelsSaveSampForLen` refers to the length of the `levelsSaveSampFor`.

Usage

```
evolMonteCarlo(nIters,
               temperLadder,
               startingVals,
               logTarDensFunc,
               MHPropNewFunc,
               logMHPropDensFunc = NULL,
               MHBlocks           = NULL,
               MHBlockNTimes     = NULL,
               moveProbsList     = NULL,
               moveNTimesList    = NULL,
               SCRWMNTimes       = NULL,
               SCRWMPropSD       = NULL,
               levelsSaveSampFor = NULL,
               nThin             = 1,
               saveFitness       = FALSE,
               verboseLevel      = 0,
               ...)
```

Arguments

<code>nIters</code>	integer > 0.
<code>temperLadder</code>	double vector with all <i>positive</i> entries, in <i>decreasing</i> order.
<code>startingVals</code>	double matrix of dimension <code>temperLadderLen</code> × <code>sampDim</code> or vector of length <code>sampDim</code> , in which case the same starting values are used for every temperature level.
<code>logTarDensFunc</code>	function of two arguments (<code>draw</code> , ...) that returns the target density evaluated in the log scale.
<code>MHPropNewFunc</code>	function of four arguments (<code>temperature</code> , <code>block</code> , <code>currentDraw</code> , ...) that returns new Metropolis-Hastings proposals. <i>See details below on the argument block.</i>

logMHPropDensFunc	function of five arguments (temperature, block, currentDraw, proposalDraw, ...) that returns the proposal density evaluated in the log scale. <i>See details below on the argument block.</i>
MHBLOCKS	list of integer vectors giving dimensions to be blocked together for sampling. It defaults to <code>as.list(1:sampDim)</code> , i.e., each dimension is treated as a block on its own. <i>See details below for an example.</i>
MHBlockNTimes	integer vector of number of times each block given by MHBLOCKS should be sampled in each iteration. It defaults to <code>rep(1, length(MHBLOCKS))</code> . <i>See details below for an example.</i>
moveProbsList	named list of probabilities adding upto 1.
moveNTimesList	named list of integers ≥ 0 .
SCRWMNTimes	integer > 0 .
SCRWMPPropSD	double > 0 .
levelsSaveSampFor	integer vector with <i>positive</i> entries.
nThin	integer ≥ 1 . Every nThin draw is saved.
saveFitness	logical.
verboseLevel	integer, a value ≥ 2 produces a lot of output.
...	optional arguments to be passed to logTarDensFunc, MHPropNewFunc and logMHPropDensFunc.

Details

MHPropNewFunc and logMHPropDensFunc The MHPropNewFunc and the logMHPropDensFunc are called multiple times by varying the block argument over `1:length(MHBLOCKS)`, so these functions should know how to generate a proposal from the currentDraw or to evaluate the proposal density depending on which block was passed as the argument. *See the example section for sample code.*

MHBLOCKS and MHBlockNTimes Blocking is an important and useful tool in MCMC that helps speed up sampling and hence mixing. Example: Let `sampDim = 6`. Let we want to sample dimensions 1, 2, 4 as one block, dimensions 3 and 5 as another and treat dimension 6 as the third block. Suppose we want to sample the three blocks mentioned above 1, 5 and 10 times in each iteration, respectively. Then we could set `MHBLOCKS = list(c(1, 2, 4), c(3, 5), 6)` and `MHBlockNTimes = c(1, 5, 10)`.

The EMC and the TOEMC algorithm The evolutionary Monte Carlo (EMC; Liang and Wong, 2001) algorithm is composed of the following moves:

MH	Metropolis-Hastings or mutation
RC	real crossover
SC	snooker crossover
RE	(random) exchange

The target oriented EMC (TOEMC; Goswami and Liu, 2007) algorithm has the following additional moves on top of EMC:

BCE best chromosome exchange
 BIRE best importance ratio exchange
 BSE best swap exchange
 CE cyclic exchange

The current function could be used to run both EMC and TOEMC algorithms by specifying what moves to employ using the following variables.

`moveProbsList` **and** `moveNTimesList` The allowed names for components of `moveProbsList` and `moveNTimesList` come from the abbreviated names of the moves above. For example, the following specifications are valid:

```
moveProbsList = list(MH = 0.4,
                    RC = 0.3,
                    SC = 0.3)

moveNTimesList = list(MH = 1,
                    RC = floor(temperLadderLen / 2),
                    SC = floor(temperLadderLen / 2),
                    RE = temperLadderLen)
```

`SCRWMNTimes` **and** `SCRWMPPropSD` The conditional sampling step of the snooker crossover (SC) move is done using random walk Metropolis (RWM) with normal proposals; `SCRWMNTimes` and `SCRWMPPropSD` are the number of RWM draws and the proposal standard deviation for the RWM step, respectively. Note these variables are only required if the SC move has positive probability in `moveProbsList` or a positive number of times in `moveNTimesList`.

`levelsSaveSampFor` By default, samples are saved and returned for temperature level `temperLadderLen`. The `levelsSaveSampFor` could be used to save samples from other temperature levels as well (e.g., `levelsSaveSampFor = 1:temperLadderLen` saves samples from all levels).

`saveFitness` The term *fitness* refers to the function $H(x)$, where the target density of interest is given by:

$$g(x) \propto \exp[-H(x)/\tau_{min}]$$

$H(x)$ is also known as the *energy* function. By default, the fitness values are not saved, but one can do so by setting `saveFitness = TRUE`.

Value

Below `nSave` refers to `ceil(nIters / nThin)`. This function returns a list with the following components:

`draws` array of dimension `nSave × sampDim × levelsSaveSampForLen`, if `saveFitness = FALSE`. If `saveFitness = TRUE`, then the returned array is of dimension `nSave × (sampDim + 1) × levelsSaveSampForLen`; i.e., each of the `levelsSaveSampForLen` matrices contain the fitness values in their last column.

`acceptRatios` matrix of the acceptance rates for various moves used.

detailedAcceptRatios	list of matrices with detailed summary of the acceptance rates for various moves used.
nIters	the nIters argument.
nThin	the nThin argument.
nSave	as defined above.
temperLadder	the temperLadder argument.
startingVals	the startingVals argument.
moveProbsList	the moveProbsList argument.
moveNTimesList	the moveNTimesList argument.
levelsSaveSampFor	the levelsSaveSampFor argument.
time	the time taken by the run.

Note

The effect of leaving the default value NULL for some of the arguments above are as follows:

logMHPPropDensFunc	the proposal density MHPPropNewFunc is deemed symmetric.
MHBlocks	as.list(1:sampDim).
MHBlockNTimes	rep(1, length(MHBlocks)).
moveProbsList	list(MH = 0.4, RC = 0.3, SC = 0.3).
moveNTimesList	list(MH = 1, RC = mm, SC = mm, RE = nn), where mm <- floor(nn / 2) and nn <- temperLadderLen.
SCRWMNTimes	1, if SC is used.
SCRWMPPropSD	needs to be provided by the user, if SC is used.
levelsSaveSampFor	temperLadderLen.

Author(s)

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References

- Gopi Goswami and Jun S. Liu (2007). On learning strategies for evolutionary Monte Carlo. Statistics and Computing 17:1:23-38.*
- Faming Liang and Wing H. Wong (2001). Real-Parameter Evolutionary Monte Carlo with Applications to Bayesian Mixture Models. Journal of the American Statistical Association 96:653-666.*

See Also

[parallelTempering](#)

Examples

```
## Not run:
samplerObj <-
```

```

with(VShapedFuncGenerator(-13579),
{
  allMovesNTimesList <- list(MH = 1, RC = 2, SC = 2, RE = 4,
                             BIRE = 2, BCE = 2, BSE = 2, CE = 2)
  evolMonteCarlo(nIters      = 2000,
                 temperLadder = c(15, 6, 2, 1),
                 startingVals = c(0, 0),
                 logTarDensFunc = logTarDensFunc,
                 MHPropNewFunc = MHPropNewFunc,
                 moveNTimesList = allMovesNTimesList,
                 SCRWMNTimes = 1,
                 SCRWMPropSD = 3.0,
                 levelsSaveSampFor = seq_len(4),
                 verboseLevel = 1)
})
print(samplerObj)
print(names(samplerObj))
with(samplerObj,
{
  print(detailedAcceptRatios)
  print(dim(draws))
  par(mfcol = c(2, 2))
  for (ii in seq_along(levelsSaveSampFor)) {
    main <- paste('temper:', round(temperLadder[levelsSaveSampFor[ii]], 3))
    plot(draws[ , , ii],
         xlim = c(-5, 20),
         ylim = c(-8, 8),
         pch = '.',
         ask = FALSE,
         main = as.expression(main),
         xlab = as.expression(substitute(x[xii], list(xii = 1))),
         ylab = as.expression(substitute(x[xii], list(xii = 2))))
  }
})
## End(Not run)

```

findMaxTemper

Find the maximum temperature for parallel MCMC chains

Description

Multiple MCMC chains based algorithms (e.g., parallel tempering, evolutionary Monte Carlo) need a temperature ladder. This function finds the maximum temperature for constructing the ladder.

Below `sampDim` refers to the dimension of the sample space, `temperLadderLen` refers to the length of the temperature ladder, and `levelsSaveSampForLen` refers to the length of `levelsSaveSampFor`. Note, this function calls `evolMonteCarlo`, so some of the arguments below have the same name and meaning as the corresponding ones for `evolMonteCarlo`. See details below for explanation on the arguments.

Usage

```

findMaxTemper(nIters,
              statsFuncList,
              startingVals,
              logTarDensFunc,
              MHPropNewFunc,
              logMHPropDensFunc = NULL,
              temperLadder      = NULL,
              temperLimits     = NULL,
              ladderLen        = 10,
              scheme            = 'exponential',
              schemeParam      = 0.5,
              cutoffDStats     = 1.96,
              cutoffESS        = 50,
              guideMe           = TRUE,
              levelsSaveSampFor = NULL,
              saveFitness       = FALSE,
              doFullAnal       = TRUE,
              verboseLevel     = 0,
              ...)

```

Arguments

nIters	integer > 0.
statsFuncList	list of functions of one argument each, which return the value of the statistic evaluated at one MCMC sample or draw.
startingVals	double matrix of dimension temperLadderLen × sampDim or vector of length sampDim, in which case the same starting values are used for every temperature level.
logTarDensFunc	function of two arguments (draw, ...) that returns the target density evaluated in the log scale.
MHPropNewFunc	function of four arguments (temperature, block, currentDraw, ...) that returns new Metropolis-Hastings proposals. <i>See details below on the argument block.</i>
logMHPropDensFunc	function of five arguments (temperature, block, currentDraw, proposalDraw, ...) that returns the proposal density evaluated in the log scale. <i>See details below on the argument block.</i>
temperLadder	double vector with all <i>positive</i> entries, in <i>decreasing</i> order.
temperLimits	double vector with <i>two positive</i> entries.
ladderLen	integer > 0.
scheme	character.
schemeParam	double > 0.
cutoffDStats	double > 0.
cutoffESS	double > 0.

guideMe logical.
 levelsSaveSampFor integer vector with *positive* entries.
 saveFitness logical.
 doFullAnal logical.
 verboseLevel integer, a value ≥ 2 produces a lot of output.
 ... optional arguments to be passed to logTarDensFunc, MHPPropNewFunc and logMHPPropDensFunc.

Details

This function is based on the method to find the temperature range introduced in section 4.1 of Goswami and Liu (2007).

statsFuncList The user specifies this list of functions, each of which is known to be sensitive to the presence of modes. For example, if both dimension 1 and 3 are sensitive to presence of modes, then one could use:

```

coord1           <- function (xx) { xx[1] }

coord3           <- function (xx) { xx[3] }

statsFuncList <- list(coord1, coord3)
  
```

temperLadder This is the temperature ladder needed for the first stage preliminary run. One can either specify a temperature ladder via `temperLadder` or specify `temperLimits`, `ladderLen`, `scheme` and `schemeParam`. For details on the later set of parameters, see below. Note, `temperLadder` overrides `temperLimits`, `ladderLen`, `scheme` and `schemeParam`.

temperLimits `temperLimits = c(lowerLimit, upperLimit)` is a two-tuple of positive numbers, where the `lowerLimit` is usually 1 and `upperLimit` is a number in [100, 1000]. If stochastic optimization (via sampling) is the goal, then `lowerLimit` is taken to be in [0, 1].

ladderLen, **scheme** **and** **schemeParam** These three parameters are required (along with `temperLimits`) if `temperLadder` *is not* provided. We recommend taking `ladderLen` in [15, 30]. The allowed choices for `scheme` and `schemeParam` are:

scheme	schemeParam
=====	=====
linear	NA
log	NA
geometric	NA
mult-power	NA
add-power	≥ 0
reciprocal	NA
exponential	≥ 0
tangent	≥ 0

We recommended using `scheme = 'exponential'` and `schemeParam` in `[0.3, 0.5]`.

`cutoffDStats` This cutoff comes from $Normal_1(0, 1)$, the standard normal distribution (Goswami and Liu, 2007); the default value 1.96 is a conservative cutoff. Note if you have more than one statistic in `statsFuncList`, which is usually the case, using this cutoff may result in different suggested maximum temperatures (as can be seen by calling the `print` function on the result of `findMaxTemper`). A conservative recommendation is that you choose the maximum of the suggested temperatures as the final maximum temperature for use in `placeTemperers` and later in `parallelTempering` or `evolMonteCarlo`.

`cutoffESS` a cutoff for the effective sample size (ESS) of the underlying Markov chain ergodic estimator and the importance sampling estimators.

`guideMe` If `guideMe = TRUE`, then the function suggests different modifications to alter the setting towards a re-run, in case there are problems with the underlying MCMC run.

`doFullAnal` If `doFullAnal = TRUE`, then the search for the maximum temperature is conducted among *all* the levels of the `temperLadder`. In case this switch is turned off, the search for maximum temperature is done in a greedy (and faster) manner, namely, search is stopped as soon as all the statistic(s) in the `statsFuncList` find some maximum temperature(s). Note, the greedy search may result in much higher maximum temperature (and hence sub-optimal) than needed, so it is not recommended.

`levelsSaveSampFor` This is passed to `evolMonteCarlo` for the underlying MCMC run.

Value

This function returns a list with the following components:

<code>temperLadder</code>	the temperature ladder used for the underlying MCMC run.
<code>DStats</code>	the D -statistic (Goswami and Liu, 2007) values used to find the maximum temperature.
<code>cutoffDStats</code>	the <code>cutoffDStats</code> argument.
<code>nIters</code>	the post burn-in <code>nIters</code> .
<code>levelsSaveSampFor</code>	the <code>levelsSaveSampFor</code> argument.
<code>draws</code>	array of dimension <code>nIters × sampDim × levelsSaveSampForLen</code> , if <code>saveFitness = FALSE</code> . If <code>saveFitness = TRUE</code> , then the returned array is of dimension <code>nIters × (sampDim + 1) × levelsSaveSampForLen</code> ; i.e., each of the <code>levelsSaveSampForLen</code> matrices contain the fitness values in their last column.
<code>startingVals</code>	the <code>startingVals</code> argument.
<code>intermediate statistics</code>	a bunch of intermediate statistics used in the computation of <code>DStats</code> , namely, <code>MCEsts</code> , <code>MCVarEsts</code> , <code>MCESS</code> , <code>ISEsts</code> , <code>ISVarEsts</code> , <code>ISESS</code> , each being computed for all the statistics provided by <code>statsFuncList</code> argument.
<code>time</code>	the time taken by the run.

Note

The effect of leaving the default value `NULL` for some of the arguments above are as follows:

logMHPropDensFunc	the proposal density MHPropNewFunc is deemed symmetric.
temperLadder	valid temperLimits, ladderLen, scheme and schemeParam are provided, which are used to construct the temperLadder.
temperLimits	a valid temperLadder is provided.
levelsSaveSampFor	temperLadderLen.

Author(s)

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References

Gopi Goswami and Jun S. Liu (2007). On learning strategies for evolutionary Monte Carlo. Statistics and Computing 17:1:23-38.

See Also

[placeTempers](#), [parallelTempering](#), [evolMonteCarlo](#)

Examples

```
## Not run:
coord1      <- function (xx) { xx[1] }
ss          <- function (xx) { sum(xx) }
pp          <- function (xx) { prod(xx) }
statsFuncList <- list(coord1, ss, pp)
maxTemperObj <-
  with(VShapedFuncGenerator(-13579),
    findMaxTemper(nIters      = 15000,
                  statsFuncList = statsFuncList,
                  temperLadder = c(20, 15, 10, 5, 1),
                  startingVals  = c(0, 0),
                  logTarDensFunc = logTarDensFunc,
                  MHPropNewFunc = MHPropNewFunc,
                  levelsSaveSampFor = seq_len(5),
                  doFullAnal     = TRUE,
                  verboseLevel   = 1))

print(maxTemperObj)
print(names(maxTemperObj))
with(maxTemperObj,
  {
    par(mfcol = c(3, 3))
    for (ii in seq_along(levelsSaveSampFor)) {
      main <- paste('temper:', round(temperLadder[levelsSaveSampFor[ii]], 3))
      plot(draws[ , , ii],
           xlim = c(-10, 25),
           ylim = c(-10, 10),
           pch = '.',
           ask = FALSE,
           main = as.expression(main),
           xlab = as.expression(substitute(x[xii], list(xii = 1))),
           ylab = as.expression(substitute(x[xii], list(xii = 2))))
    }
  }
}
```

```

    }
  })

  ## End(Not run)

```

MetropolisHastings *The Metropolis-Hastings algorithm*

Description

Given a target density function and an asymmetric proposal distribution, this function produces samples from the target using the Metropolis Hastings algorithm.

Below sampDim refers to the dimension of the sample space.

Usage

```

MetropolisHastings(nIters,
                   startingVal,
                   logTarDensFunc,
                   propNewFunc,
                   logPropDensFunc,
                   MHBlocks = NULL,
                   MHBlockNTimes = NULL,
                   nThin = 1,
                   saveFitness = FALSE,
                   verboseLevel = 0,
                   ...)

```

Arguments

nIters	integer > 0.
startingVal	double vector of length sampDim.
logTarDensFunc	function of two arguments (draw, ...) that returns the target density evaluated in the log scale.
propNewFunc	function of three arguments (block, currentDraw, ...) that returns new Metropolis-Hastings proposals. <i>See details below on the argument block.</i>
logPropDensFunc	function of four arguments (block, currentDraw, proposalDraw, ...) that returns the proposal density evaluated in the log scale. <i>See details below on the argument block.</i>
MHBlocks	list of integer vectors giving dimensions to be blocked together for sampling. It defaults to <code>as.list(1:sampDim)</code> , i.e., each dimension is treated as a block on its own. <i>See details below for an example.</i>
MHBlockNTimes	integer vector of number of times each block given by MHBlocks should be sampled in each iteration. It defaults to <code>rep(1, length(MHBlocks))</code> . <i>See details below for an example.</i>

nThin	integer ≥ 1 . Every nThin draw is saved.
saveFitness	logical indicating whether fitness values should be saved. <i>See details below.</i>
verboseLevel	integer, a value ≥ 2 produces a lot of output.
...	optional arguments to be passed to logTarDensFunc, propNewFunc and logPropDensFunc.

Details

propNewFunc **and** logPropDensFunc The propNewFunc and the logPropDensFunc are called multiple times by varying the block argument over $1:\text{length}(\text{MHBlocks})$, so these functions should know how to generate a proposal from the currentDraw or to evaluate the proposal density depending on which block was passed as the argument. *See the example section for sample code.*

MHBlocks **and** MHBlockNTimes Blocking is an important and useful tool in MCMC that helps speed up sampling and hence mixing. Example: Let sampDim = 6. Let we want to sample dimensions 1, 2, 4 as one block, dimensions 3 and 5 as another and treat dimension 6 as the third block. Suppose we want to sample the three blocks mentioned above 1, 5 and 10 times in each iteration, respectively. Then we could set MHBlocks = list(c(1, 2, 4), c(3, 5), 6) and MHBlockNTimes = c(1, 5, 10)

saveFitness The term *fitness* refers to the negative of the logTarDensFunc values. By default, the fitness values are not saved, but one can do so by setting saveFitness = TRUE.

Value

Below nSave refers to $\text{ceil}(\text{nIters} / \text{nThin})$. This function returns a list with the following components:

draws	matrix of dimension nSave \times sampDim, if saveFitness = FALSE. If saveFitness = TRUE, then the returned matrix is of dimension nSave \times (sampDim + 1), where the fitness values appear in its last column.
acceptRatios	matrix of the acceptance rates.
detailedAcceptRatios	matrix with detailed summary of the acceptance rates.
nIters	the nIters argument.
nThin	the nThin argument.
nSave	as defined above.
startingVal	the startingVal argument.
time	the time taken by the run.

Note

The effect of leaving the default value NULL for some of the arguments above are as follows:

```
MHBlocks  as.list(1:sampDim).
MHBlockNTimes rep(1, length(MHBlocks)).
```

Author(s)

Gopi Goswami <goswami@stat.harvard.edu>

References

Jun S. Liu (2001). Monte Carlo strategies for scientific computing. *Springer*.

See Also

[randomWalkMetropolis](#), [parallelTempering](#), [evolMonteCarlo](#)

Examples

```
## Not run:
samplerObj <-
  with(CigarShapedFuncGenerator2(-13579),
    MetropolisHastings(nIters      = 5000,
                      startingVal  = c(0, 0),
                      logTarDensFunc = logTarDensFunc,
                      propNewFunc   = propNewFunc,
                      logPropDensFunc = logPropDensFunc,
                      verboseLevel  = 2))

print(samplerObj)
print(names(samplerObj))
with(samplerObj,
  {
    print(detailedAcceptRatios)
    print(dim(draws))
    plot(draws,
         xlim = c(-3, 5),
         ylim = c(-3, 4),
         pch = '.',
         ask = FALSE,
         main = as.expression(paste('# draws:', nIters)),
         xlab = as.expression(substitute(x[xii], list(xii = 1))),
         ylab = as.expression(substitute(x[xii], list(xii = 2))))
  })

## End(Not run)
```

parallelTempering *The parallel Tempering algorithm*

Description

Given a multi-modal and multi-dimensional target density function, a (possibly asymmetric) proposal distribution and a temperature ladder, this function produces samples from the target using the parallel tempering algorithm.

Below sampDim refers to the dimension of the sample space, temperLadderLen refers to the length of the temperature ladder, and levelsSaveSampForLen refers to the length of the levelsSaveSampFor.

Usage

```
parallelTempering(nIters,
                  temperLadder,
                  startingVals,
                  logTarDensFunc,
                  MHPropNewFunc,
                  logMHPropDensFunc = NULL,
                  MHBlocks          = NULL,
                  MHBlockNTimes    = NULL,
                  moveProbsList     = NULL,
                  moveNTimesList    = NULL,
                  levelsSaveSampFor = NULL,
                  nThin             = 1,
                  saveFitness       = FALSE,
                  verboseLevel      = 0,
                  ...)
```

Arguments

nIters integer > 0.

temperLadder double vector with all *positive* entries, in *decreasing* order.

startingVals double matrix of dimension `temperLadderLen × sampDim` or vector of length `sampDim`, in which case the same starting values are used for every temperature level.

logTarDensFunc function of two arguments (`draw`, ...) that returns the target density evaluated in the log scale.

MHPropNewFunc function of four arguments (`temperature`, `block`, `currentDraw`, ...) that returns new Metropolis-Hastings proposals. *See details below on the argument block.*

logMHPropDensFunc function of five arguments (`temperature`, `block`, `currentDraw`, `proposalDraw`, ...) that returns the proposal density evaluated in the log scale. *See details below on the argument block.*

MHBlocks list of integer vectors giving dimensions to be blocked together for sampling. It defaults to `as.list(1:sampDim)`, i.e., each dimension is treated as a block on its own. *See details below for an example.*

MHBlockNTimes integer vector of number of times each block given by `MHBlocks` should be sampled in each iteration. It defaults to `rep(1, length(MHBlocks))`. *See details below for an example.*

moveProbsList named list of probabilities adding upto 1.

moveNTimesList named list of integers ≥ 0 .

levelsSaveSampFor integer vector with *positive* entries.

nThin integer ≥ 1 . Every `nThin` draw is saved.

saveFitness logical.

verboseLevel integer, a value ≥ 2 produces a lot of output.
 ... optional arguments to be passed to logTarDensFunc, MHPropNewFunc and logMHPropDensFunc.

Details

MHPropNewFunc **and** logMHPropDensFunc The MHPropNewFunc and the logMHPropDensFunc are called multiple times by varying the block argument over $1:\text{length}(\text{MHBlocks})$, so these functions should know how to generate a proposal from the currentDraw or to evaluate the proposal density depending on which block was passed as the argument. *See the example section for sample code.*

MHBlocks **and** MHBlockNTimes Blocking is an important and useful tool in MCMC that helps speed up sampling and hence mixing. Example: Let `sampDim = 6`. Let we want to sample dimensions 1, 2, 4 as one block, dimensions 3 and 5 as another and treat dimension 6 as the third block. Suppose we want to sample the three blocks mentioned above 1, 5 and 10 times in each iteration, respectively. Then we could set `MHBlocks = list(c(1, 2, 4), c(3, 5), 6)` and `MHBlockNTimes = c(1, 5, 10)`.

The parallel tempering algorithm The parallel tempering (PT; Liang and Wong, 2001) algorithm is composed of the following moves:

MH Metropolis-Hastings or mutation
 RE (random) exchange

The current function could be used to run the PT algorithm by specifying what moves to employ using the following variables.

moveProbsList **and** moveNTimesList The allowed names for components of moveProbsList and moveNTimesList come from the abbreviated names of the moves above. For example, the following specifications are valid:

```
moveProbsList = list(MH = 0.4,
                     RE = 0.6)
```

```
moveNTimesList = list(MH = 1,
                     RE = temperLadderLen)
```

levelsSaveSampFor By default, samples are saved and returned for temperature level temperLadderLen.

The levelsSaveSampFor could be used to save samples from other temperature levels as well (e.g., levelsSaveSampFor = $1:\text{temperLadderLen}$ saves samples from all levels).

saveFitness The term *fitness* refers to the function $H(x)$, where the target density of interest is given by:

$$g(x) \propto \exp[-H(x)/\tau_{min}]$$

$H(x)$ is also known as the *energy* function. By default, the fitness values are not saved, but one can do so by setting saveFitness = TRUE.

Value

Below nSave refers to $\text{ceil}(\text{nIters} / \text{nThin})$. This function returns a list with the following components:

draws	array of dimension $nSave \times sampDim \times levelsSaveSampForLen$, if <code>saveFitness = FALSE</code> . If <code>saveFitness = TRUE</code> , then the returned array is of dimension $nSave \times (sampDim + 1) \times levelsSaveSampForLen$; i.e., each of the <code>levelsSaveSampForLen</code> matrices contain the fitness values in their last column.
acceptRatios	matrix of the acceptance rates for various moves used.
detailedAcceptRatios	list of matrices with detailed summary of the acceptance rates for various moves used.
nIters	the <code>nIters</code> argument.
nThin	the <code>nThin</code> argument.
nSave	as defined above.
temperLadder	the <code>temperLadder</code> argument.
startingVals	the <code>startingVals</code> argument.
moveProbsList	the <code>moveProbsList</code> argument.
moveNTimesList	the <code>moveNTimesList</code> argument.
levelsSaveSampFor	the <code>levelsSaveSampFor</code> argument.
time	the time taken by the run.

Note

The effect of leaving the default value NULL for some of the arguments above are as follows:

logMHPropDensFunc	the proposal density <code>MHPropNewFunc</code> is deemed symmetric.
MHBlocks	<code>as.list(1:sampDim)</code> .
MHBlockNTimes	<code>rep(1, length(MHBlocks))</code> .
moveProbsList	<code>list(MH = 0.4, RC = 0.3, SC = 0.3)</code> .
moveNTimesList	<code>list(MH = 1, RC = mm, SC = mm, RE = nn)</code> , where <code>mm <- floor(nn / 2)</code> and <code>nn <- temperLadderLen</code> .
levelsSaveSampFor	<code>temperLadderLen</code> .

Author(s)

Gopi Goswami <goswami@stat.harvard.edu>

References

Faming Liang and Wing H. Wong (2001). Real-Parameter Evolutionary Monte Carlo with Applications to Bayesian Mixture Models. Journal of the American Statistical Association 96:653-666.

See Also

[evolMonteCarlo](#)

Examples

```
## Not run:
```

```

samplerObj <-
  with(VShapedFuncGenerator(-13579),
    parallelTempering(nIters           = 2000,
                      temperLadder    = c(15, 6, 2, 1),
                      startingVals     = c(0, 0),
                      logTarDensFunc   = logTarDensFunc,
                      MHPropNewFunc    = MHPropNewFunc,
                      levelsSaveSampFor = seq_len(4),
                      verboseLevel     = 1))

print(samplerObj)
print(names(samplerObj))
with(samplerObj,
  {
    print(detailedAcceptRatios)
    print(dim(draws))
    par(mfcol = c(2, 2))
    for (ii in seq_along(levelsSaveSampFor)) {
      main <- paste('temper:', round(temperLadder[levelsSaveSampFor[ii]], 3))
      plot(draws[ , , ii],
           xlim = c(-5, 20),
           ylim = c(-8, 8),
           pch = '.',
           ask = FALSE,
           main = as.expression(main),
           xlab = as.expression(substitute(x[xii], list(xii = 1))),
           ylab = as.expression(substitute(x[xii], list(xii = 2))))
    }
  })

## End(Not run)

```

placeTempers

Place the intermediate temperatures between the temperature limits

Description

Multiple MCMC chains based algorithms (e.g., parallel tempering, evolutionary Monte Carlo) need a temperature ladder. This function places the intermediate temperatures between the minimum and the maximum temperature for the ladder.

Below `sampDim` refers to the dimension of the sample space, `temperLadderLen` refers to the length of the temperature ladder, and `levelsSaveSampForLen` refers to the length of `levelsSaveSampFor`. Note, this function calls `evolMonteCarlo`, so some of the arguments below have the same name and meaning as the corresponding ones for `evolMonteCarlo`. See details below for explanation on the arguments.

Usage

```

placeTempers(nIters,
             acceptRatLiimits,

```

```

ladderLenMax,
startingVals,
logTarDensFunc,
MHPropNewFunc,
logMHPropDensFunc = NULL,
temperLadder      = NULL,
temperLimits      = NULL,
ladderLen         = 15,
scheme            = 'exponential',
schemeParam       = 1.5,
guideMe           = TRUE,
levelsSaveSampFor = NULL,
saveFitness       = FALSE,
verboseLevel      = 0,
...)
```

Arguments

nIters	integer > 0.
acceptRatioLimits	double vector of <i>two probabilities</i> .
ladderLenMax	integer > 0.
startingVals	double matrix of dimension $\text{temperLadderLen} \times \text{sampDim}$ or vector of length sampDim , in which case the same starting values are used for every temperature level.
logTarDensFunc	function of two arguments (<code>draw</code> , ...) that returns the target density evaluated in the log scale.
MHPropNewFunc	function of four arguments (<code>temperature</code> , <code>block</code> , <code>currentDraw</code> , ...) that returns new Metropolis-Hastings proposals. <i>See details below on the argument block.</i>
logMHPropDensFunc	function of five arguments (<code>temperature</code> , <code>block</code> , <code>currentDraw</code> , <code>proposalDraw</code> , ...) that returns the proposal density evaluated in the log scale. <i>See details below on the argument block.</i>
temperLadder	double vector with all <i>positive entries</i> , in <i>decreasing</i> order.
temperLimits	double vector with <i>two positive entries</i> .
ladderLen	integer > 0.
scheme	character.
schemeParam	double > 0.
guideMe	logical.
levelsSaveSampFor	integer vector with <i>positive entries</i> .
saveFitness	logical.
verboseLevel	integer, a value ≥ 2 produces a lot of output.
...	optional arguments to be passed to <code>logTarDensFunc</code> , <code>MHPropNewFunc</code> and <code>logMHPropDensFunc</code> .

Details

This function is based on the temperature placement method introduced in section 4.2 of Goswami and Liu (2007).

`acceptRatioLimits` This is a range for the estimated acceptance ratios for the random exchange move for the consecutive temperature levels of the final ladder. It is recommended that specified range is between 0.3 and 0.6.

`ladderLenMax` It is preferred that one specifies `acceptRatioLimits` for constructing the final temperature ladder. However, If one has some computational limitations then one could also specify `ladderLenMax` which will limit the length of the final temperature ladder produced. This also serves as an upper bound on the number of temperature levels while placing the intermediate temperatures using the `acceptRatioLimits`.

`temperLadder` This is the temperature ladder needed for the second stage preliminary run. One can either specify a temperature ladder via `temperLadder` or specify `temperLimits`, `ladderLen`, `scheme` and `schemeParam`. For details on the later set of parameters, see below. Note, `temperLadder` overrides `temperLimits`, `ladderLen`, `scheme` and `schemeParam`.

`temperLimits` `temperLimits = c(lowerLimit, upperLimit)` is a two-tuple of positive numbers, where the `lowerLimit` is usually 1 and `upperLimit` is a number in [100, 1000]. If stochastic optimization (via sampling) is the goal, then `lowerLimit` is taken to be in [0, 1]. Often the `upperLimit` is the maximum temperature as suggested by `findMaxTemper`.

`ladderLen`, `scheme` **and** `schemeParam` These three parameters are required (along with `temperLimits`) if `temperLadder` *is not* provided. We recommend taking `ladderLen` in [15, 30]. The allowed choices for `scheme` and `schemeParam` are:

scheme	schemeParam
=====	=====
linear	NA
log	NA
geometric	NA
mult-power	NA
add-power	≥ 0
reciprocal	NA
exponential	≥ 0
tangent	≥ 0

We recommended using `scheme = 'exponential'` and `schemeParam` in [1.5, 2].

`guideMe` If `guideMe = TRUE`, then the function suggests different modifications to alter the setting towards a re-run, in case there are problems with the underlying MCMC run.

`levelsSaveSampFor` This is passed to `evolMonteCarlo` for the underlying MCMC run.

Value

This function returns a list with the following components:

`finalLadder` the final temperature ladder found by placing the intermediate temperatures to be used in `parallelTempering` or `evolMonteCarlo`.

temperLadder	the temperature ladder used for the underlying MCMC run.
acceptRatiosEst	the estimated acceptance ratios for the random exchange move for the consecutive temperature levels of temperLadder.
CVSqWeights	this is the square of the coefficient of variation of the weights of the importance sampling estimators used to estimate the acceptance ratios, namely, estAcceptRatios.
temperLimits	the sorted temperLimits argument.
acceptRatioLimits	the sorted acceptRatioLimits argument.
nIters	the post burn-in nIters.
levelsSaveSampFor	the levelsSaveSampFor argument.
draws	array of dimension $nIters \times sampDim \times levelsSaveSampForLen$, if saveFitness = FALSE. If saveFitness = TRUE, then the returned array is of dimension $nIters \times (sampDim + 1) \times levelsSaveSampForLen$; i.e., each of the levelsSaveSampForLen matrices contain the fitness values in their last column.
startingVals	the startingVals argument.
time	the time taken by the run.

Note

The effect of leaving the default value NULL for some of the arguments above are as follows:

logMHPropDensFunc	the proposal density MHPropNewFunc is deemed symmetric.
temperLadder	valid temperLimits, ladderLen, scheme and schemeParam are provided, which are used to construct the temperLadder.
temperLimits	a valid temperLadder is provided.
levelsSaveSampFor	temperLadderLen.

Author(s)

Gopi Goswami <goswami@stat.harvard.edu>

References

Gopi Goswami and Jun S. Liu (2007). On learning strategies for evolutionary Monte Carlo. Statistics and Computing 17:1:23-38.

See Also

[findMaxTemper](#), [parallelTempering](#), [evolMonteCarlo](#)

Examples

```
## Not run:
placeTempersObj <-
  with(VShapedFuncGenerator(-13579),
```

```

        placeTempers(nIters          = 10000,
                    acceptRatioLimits = c(0.5, 0.6),
                    ladderLenMax      = 50,
                    startingVals      = c(0, 0),
                    logTarDensFunc     = logTarDensFunc,
                    MHPropNewFunc      = MHPropNewFunc,
                    temperLimits      = c(1, 5),
                    ladderLen          = 10,
                    levelsSaveSampFor = seq_len(10),
                    verboseLevel       = 1))
print(placeTempersObj)
print(names(placeTempersObj))
with(placeTempersObj,
  {
    par(mfcol = c(3, 3))
    for (ii in seq_along(levelsSaveSampFor)) {
      main <- paste('temper:', round(temperLadder[levelsSaveSampFor[ii]], 3))
      plot(draws[ , , ii],
           xlim = c(-4, 20),
           ylim = c(-8, 8),
           pch = '.',
           ask = FALSE,
           main = as.expression(main),
           xlab = as.expression(substitute(x[xii], list(xii = 1))),
           ylab = as.expression(substitute(x[xii], list(xii = 2))))
    }
  })
## End(Not run)

```

print

The printing family of functions

Description

The printing family of functions for this package.

Usage

```

## S3 method for class 'EMC'
print(x, ...)
## S3 method for class 'EMCMaxTemper'
print(x, ...)
## S3 method for class 'EMCPlaceTempers'
print(x, ...)

```

Arguments

x an object inheriting from class EMC (generated by functions randomWalkMetropolis, MetropolisHastings, parallelTempering and evolMonteCarlo), EMCMaxTemper

(generated by function `findMaxTemper`) or `EMCPlaceTempers` (generated by function `placeTempers`).

... optional arguments passed to `print.default`; see its documentation.

Author(s)

Gopi Goswami <goswami@stat.harvard.edu>

See Also

[randomWalkMetropolis](#), [MetropolisHastings](#), [parallelTempering](#), [evolMonteCarlo](#), [findMaxTemper](#), [placeTempers](#)

randomWalkMetropolis *The Random Walk Metropolis algorithm*

Description

Given a target density function and a symmetric proposal generating function, this function produces samples from the target using the random walk Metropolis algorithm.

Below `sampDim` refers to the dimension of the sample space.

Usage

```
randomWalkMetropolis(nIters,
                     startingVal,
                     logTarDensFunc,
                     propNewFunc,
                     MHBlocks = NULL,
                     MHBlockNTimes = NULL,
                     nThin = 1,
                     saveFitness = FALSE,
                     verboseLevel = 0,
                     ...)
```

Arguments

<code>nIters</code>	integer > 0.
<code>startingVal</code>	double vector of length <code>sampDim</code> .
<code>logTarDensFunc</code>	function of two arguments (<code>draw</code> , ...) that returns the target density evaluated in the log scale.
<code>propNewFunc</code>	function of three arguments (<code>block</code> , <code>currentDraw</code> , ...) that returns new Metropolis-Hastings proposals. <i>See details below on the argument block.</i>
<code>MHBlocks</code>	list of integer vectors giving dimensions to be blocked together for sampling. It defaults to <code>as.list(1:sampDim)</code> , i.e., each dimension is treated as a block on its own. <i>See details below for an example.</i>

MHBlockNTimes	integer vector of number of times each block given by MHBlocks should be sampled in each iteration. It defaults to <code>rep(1, length(MHBlocks))</code> . See <i>details below for an example</i> .
nThin	integer ≥ 1 . Every nThin draw is saved.
saveFitness	logical indicating whether fitness values should be saved. See <i>details below</i> .
verboseLevel	integer, a value ≥ 2 produces a lot of output.
...	optional arguments to be passed to logTarDensFunc and propNewFunc.

Details

propNewFunc The propNewFunc is called multiple times by varying the block argument over `1:length(MHBlocks)`, so this function should know how to generate a proposal from the currentDraw depending on which block was passed as the argument. See *the example section for sample code*.

MHBlocks **and** MHBlockNTimes Blocking is an important and useful tool in MCMC that helps speed up sampling and hence mixing. Example: Let `sampDim = 6`. Let we want to sample dimensions 1, 2, 4 as one block, dimensions 3 and 5 as another and treat dimension 6 as the third block. Suppose we want to sample the three blocks mentioned above 1, 5 and 10 times in each iteration, respectively. Then we could set `MHBlocks = list(c(1, 2, 4), c(3, 5), 6)` and `MHBlockNTimes = c(1, 5, 10)`

saveFitness The term *fitness* refers to the negative of the logTarDensFunc values. By default, the fitness values are not saved, but one can do so by setting `saveFitness = TRUE`.

Value

Below nSave refers to `ceil(nIters / nThin)`. This function returns a list with the following components:

draws	matrix of dimension <code>nSave × sampDim</code> , if <code>saveFitness = FALSE</code> . If <code>saveFitness = TRUE</code> , then the returned matrix is of dimension <code>nSave × (sampDim + 1)</code> , where the fitness values appear in its last column.
acceptRatios	matrix of the acceptance rates.
detailedAcceptRatios	matrix with detailed summary of the acceptance rates.
nIters	the nIters argument.
nThin	the nThin argument.
nSave	as defined above.
startingVal	the startingVal argument.
time	the time taken by the run.

Note

The effect of leaving the default value NULL for some of the arguments above are as follows:

```
MHBlocks    as.list(1:sampDim).
MHBlockNTimes rep(1, length(MHBlocks)).
```

Author(s)

Gopi Goswami <goswami@stat.harvard.edu>

References

Jun S. Liu (2001). Monte Carlo strategies for scientific computing. Springer.

See Also

[MetropolisHastings](#), [parallelTempering](#), [evolMonteCarlo](#)

Examples

```
## Not run:
samplerObj <-
  with(CigarShapedFuncGenerator1(-13579),
    randomWalkMetropolis(nIters      = 5000,
                        startingVal  = c(0, 0),
                        logTarDensFunc = logTarDensFunc,
                        propNewFunc   = propNewFunc,
                        verboseLevel  = 1))

print(samplerObj)
print(names(samplerObj))
with(samplerObj,
  {
    print(detailedAcceptRatios)
    print(dim(draws))
    plot(draws,
         xlim = c(-3, 5),
         ylim = c(-3, 4),
         pch = '.',
         ask = FALSE,
         main = as.expression(paste('# draws:', nIters)),
         xlab = as.expression(substitute(x[xii], list(xii = 1))),
         ylab = as.expression(substitute(x[xii], list(xii = 2))))
  })

samplerObj <-
  with(threeDimNormalFuncGenerator(-13579),
    {
      randomWalkMetropolis(nIters      = 5000,
                          startingVal  = c(0, 0, 0),
                          logTarDensFunc = logTarDensFunc,
                          propNewFunc   = propNewFunc,
                          MHBlocks     = list(c(1, 2), 3),
                          verboseLevel  = 1)
    })
print(samplerObj)
print(names(samplerObj))
with(samplerObj,
  {
```

```

print(detailedAcceptRatios)
print(dim(draws))
pairs(draws,
      pch = '.',
      ask = FALSE,
      main = as.expression(paste('# draws:', nIters)),
      labels = c(as.expression(substitute(x[xii], list(xii = 1))),
                 as.expression(substitute(x[xii], list(xii = 2))),
                 as.expression(substitute(x[xii], list(xii = 3)))))
})

## End(Not run)

```

utilsForExamples

The utility function(s) for examples

Description

The utility function(s) that are used in the example sections of the exported functions in this package.

Usage

```

CigarShapedFuncGenerator1(seed)
CigarShapedFuncGenerator2(seed)
VShapedFuncGenerator(seed)
WShapedFuncGenerator(seed)
uniModeFuncGenerator(seed)
twentyModeFuncGenerator(seed)
threeDimNormalFuncGenerator(seed)

```

Arguments

`seed` the seed for random number generation.

Value

A list containing the objects to be used as arguments to the exported functions in the respective example sections of this package.

Author(s)

Gopi Goswami <goswami@stat.harvard.edu>

See Also

[randomWalkMetropolis](#), [MetropolisHastings](#), [parallelTempering](#), [evolMonteCarlo](#), [findMaxTemper](#), [placeTempers](#)

Index

- *Topic **datagen**
 - utilsForExamples, 25
- *Topic **methods**
 - evolMonteCarlo, 2
 - findMaxTemper, 6
 - MetropolisHastings, 11
 - parallelTempering, 13
 - placeTempers, 17
 - randomWalkMetropolis, 22
- *Topic **print**
 - print, 21
- CigarShapedFuncGenerator1
 - (utilsForExamples), 25
- CigarShapedFuncGenerator2
 - (utilsForExamples), 25
- evolMonteCarlo, 2, 6, 10, 13, 16, 17, 20, 22, 24, 25
- findMaxTemper, 6, 19, 20, 22, 25
- MetropolisHastings, 11, 22, 24, 25
- parallelTempering, 5, 10, 13, 13, 20, 22, 24, 25
- placeTempers, 10, 17, 22, 25
- print, 21
- randomWalkMetropolis, 13, 22, 22, 25
- threeDimNormalFuncGenerator
 - (utilsForExamples), 25
- twentyModeFuncGenerator
 - (utilsForExamples), 25
- uniModeFuncGenerator
 - (utilsForExamples), 25
- utilsForExamples, 25
- VShapedFuncGenerator
 - (utilsForExamples), 25
- WShapedFuncGenerator
 - (utilsForExamples), 25