

# Package ‘beast’

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**Type** Package

**Title** Bayesian Estimation of Change-Points in the Slope of Multivariate Time-Series

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**Description** Assume that a temporal process is composed of contiguous segments with differing slopes and replicated noise-corrupted time series measurements are observed. The unknown mean of the data generating process is modelled as a piecewise linear function of time with an unknown number of change-points. The package infers the joint posterior distribution of the number and position of change-points as well as the unknown mean parameters per time-series by MCMC sampling. A-priori, the proposed model uses an overfitting number of mean parameters but, conditionally on a set of change-points, only a subset of them influences the likelihood. An exponentially decreasing prior distribution on the number of change-points gives rise to a posterior distribution concentrating on sparse representations of the underlying sequence, but also available is the Poisson distribution. See Papastamoulis et al (2017) <[arXiv:1709.06111](#)> for a detailed presentation of the method.

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**Index****21****beast-package***Bayesian Estimation of Change-Points in the Slope of Multivariate Time-Series***Description**

Assume that a temporal process is composed of contiguous segments with differing slopes and replicated noise-corrupted time series measurements are observed. The unknown mean of the data generating process is modelled as a piecewise linear function of time with an unknown number of change-points. The package infers the joint posterior distribution of the number and position of change-points as well as the unknown mean parameters per time-series by MCMC sampling. A-priori, the proposed model uses an overfitting number of mean parameters but, conditionally on a set of change-points, only a subset of them influences the likelihood. An exponentially decreasing prior distribution on the number of change-points gives rise to a posterior distribution concentrating on sparse representations of the underlying sequence, but also available is the Poisson distribution. See Papastamoulis et al (2017) <arXiv:1709.06111> for a detailed presentation of the method.

**Details**

The beast package deals with Bayesian estimation of change-points in the slope of multivariate time-series, introduced by Papastamoulis et al (2017). For a given period  $t = 1, \dots, T$  we observe multiple time series which are assumed independent, each one consisting of multiple measurements (replicates). Each time series is assumed to have its own segmentation, which is common among its replicates. Thus, different time series have distinct mean parameters in the underlying normal distribution. The variance, which is assumed known, can be either shared between different time series or not and in practice it is estimated at a pre-processing stage.

Our model infers the joint posterior distribution of the number and location of change-points by MCMC sampling. For this purpose a Metropolis-Hastings MCMC sampler is used. The main function of the package is **beast**.

Assume that the observed data consists of  $N$  time-series, each one consisting of  $R$  variables (or replicates) measured at  $T$  consecutive time-points. The input of the main function **beast** should be given in the form of a list `myDataList` with the following attributes:

- `length(myDataList)` should be equal to  $R$ , that is, the number of variables (or replicates)
- `dim(myDataList)[[1]], ..., dim(myDataList)[[R]]` should be all  $T \times N$ , that is, rows and columns should correspond to time-points and different series, respectively.

Then, a basic usage of the package consists of the following commands:

- `beastRun <- beast( myDataList = myDataList )`
- `print(beastRun)`
- `plot(beastRun)`

which correspond to running the MCMC sampler, printing and plotting output summaries, respectively.

## Author(s)

Panagiotis Papastamoulis

Maintainer: Panagiotis Papastamoulis <papapast@yahoo.gr>

## References

Papastamoulis P., Furukawa T., van Rhijn N., Bromley M., Bignell E. and Rattray M. (2017). Bayesian detection of piecewise linear trends in replicated time-series with application to growth data modelling. arXiv:1709.06111 [stat.AP]

## See Also

`beast`, `print.beast.object`, `plot.beast.object`

## Examples

```
# toy-example (MCMC iterations not enough)
library('beast') # load package
data("FungalGrowthDataset") # load dataset
myIndex <- c(392, 62, 3, 117) # run the sampler only for the
#                                     specific subset of time-series
set.seed(1)
# Run MCMC sampler with very small number of iterations (nIter):
run_mcmc <- beast(myDataList = FungalGrowthDataset, subsetIndex = myIndex,
zeroNormalization = TRUE, nIter = 40, burn = 20)
# Print output:
print(run_mcmc)
# Plot output to file: "beast_plot.pdf"
plot(run_mcmc, fileName = "beast_plot_toy.pdf", timeScale=1/6, xlab = "hours", ylab = "growth")
```

```
# Run the following commands to obtain convergence:

## Not run:
# This example illustrates the package using a subset of four
#   time-series of the fungal dataset.
library('beast') # load package
data("FungalGrowthDataset") # load dataset
myIndex <- c(392, 62, 3, 117) # run the sampler only for the
#                           specific subset of time-series
set.seed(1) # optional
# Run MCMC sampler with the default number of iterations (nIter =70000):
run_mcmc <- beast(myDataList = FungalGrowthDataset, subsetIndex = myIndex,
zeroNormalization = TRUE)
# Print output:
print(run_mcmc)
# Plot output to file: "beast_plot.pdf"
plot(run_mcmc, fileName = "beast_plot.pdf", timeScale=1/6, xlab = "hours", ylab = "growth")
# NOTE 1: for a complete analysis remove the `subsetIndex = myIndex` argument.
# NOTE 2: `zeroNormalization = TRUE` is an optional argument that forces all
#   time-series to start from zero. It is not supposed to be used
#   for other applications.

## End(Not run)
```

**beast***Main function***Description**

This is the main function of the package, implementing the MCMC sampler described in Papastamoulis et al (2017).

**Usage**

```
beast(myDataList, burn, nIter, mhPropRange, mhSinglePropRange, startPoint,
      timeScale, savePlots, zeroNormalization, LRange, tau,
      gammaParameter, nu0, alpha0, beta0, subsetIndex, saveTheta, sameVariance,
      Prior
    )
```

**Arguments**

<code>myDataList</code>	Observed data in the form of a list with length $R$ , denoting the dimensionality of the multivariate time-series data. For each $r = 1, \dots, R$ , <code>myDataList[[r]]</code> should correspond to $T \times N$ array, with <code>myDataList[[r]][t, n]</code> corresponding to the observed data for time-series $n = 1, \dots, N$ and time-point $t = 1, \dots, T$ .
<code>burn</code>	Number of iterations that will be discarded as burn-in period. Default value: <code>burn = 20000</code> .

<code>nIter</code>	Number of MCMC iterations. Default value: <code>nIter = 70000</code> .
<code>mhPropRange</code>	Positive integer corresponding to the parameter $d_1$ of MCMC Move 3.a of Papastamoulis et al (2017). Default value: <code>mhPropRange = 1</code> .
<code>mhSinglePropRange</code>	Positive integer denoting the parameter $d_2$ of Papastamoulis et al (2017). Default value: <code>mhPropRange = 40</code> .
<code>startPoint</code>	An (optional) positive integer pointing at the minimum time-point where changes are allowed to occur. Default value: <code>startPoint = 2</code> (all possible values are taken into account).
<code>timeScale</code>	Null.
<code>savePlots</code>	Character denoting the name of the folder where various plots will be saved to.
<code>zeroNormalization</code>	Logical value denoting whether to normalize to zero all time time-series for $t = 1$ . Default: <code>zeroNormalization = FALSE</code> .
<code>LRange</code>	Range of possible values for the number of change-points. Default value: <code>LRange = 0:30</code> .
<code>tau</code>	Positive real number corresponding to parameter $c$ in Move 2 of Papastamoulis et al (2017). Default value: <code>tau = 0.05</code> .
<code>gammaParameter</code>	Positive real number corresponding to parameter $\alpha$ of the exponential prior distribution. Default value: <code>gammaParameter = 2</code> .
<code>nu0</code>	Positive real number corresponding to prior parameter $\nu_0$ in Papastamoulis et al (2017). Default value: <code>nu0 = 0.1</code> .
<code>alpha0</code>	Positive real number corresponding to prior parameter $\alpha_0$ in Papastamoulis et al (2017). Default value: <code>alpha0 = 1</code> .
<code>beta0</code>	Positive real number corresponding to prior parameter $\beta_0$ in Papastamoulis et al (2017). Default value: <code>beta0 = 1</code> .
<code>subsetIndex</code>	Optional subset of integers corresponding to time-series indexes. If not null, the sampler will be applied only to the specified subset.
<code>saveTheta</code>	Logical value indicating whether to save the generated values of the mean per time-point across the MCMC trace. Default: <code>FALSE</code> .
<code>sameVariance</code>	Logical value indicating whether to assume the same variance per time-point across time-series. Default value: <code>sameVariance = TRUE</code> .
<code>Prior</code>	Character string specifying the prior distribution of the number of change-points. Allowed values: <code>Prior = "complexity"</code> (default) or <code>Prior = "Poisson"</code> (not suggested).

## Value

The output of the sampler is returned as a list, with the following features:

### `Cutpoint_posterior_median`

The estimated medians per change-point, conditionally on the most probable number of change-points per time-series.

Cutpoint_posterior_variance	The estimated variances per change-points, conditionally on the most probable number of change-points per time-series.
NumberOfCutPoints_posterior_distribution	Posterior distributions of number of change-points per time-series.
NumberOfCutPoints_MAP	The most probable number of change-points per time-series.
Metropolis-Hastings_acceptance_rate	Acceptance of the MCMC move-types.
subject_ID	the identifier of individual time-series.
Cutpoint_mcmc_trace_map	The sampled values of each change-point per time series, conditionally on the MAP values.
theta	The sampled values of the means per time-series, conditionally on the MAP values.
nCutPointsTrace	The sampled values of the number of change-points, per time-series.

### Note

The complexity prior distribution with parameter `gammaParameter = 2` is the default prior assumption imposed on the number of change-points. Smaller (larger) values of `gammaParameter` will a-priori support larger (respectively: smaller) number of change-points.

For completeness purposes, the Poisson distribution is also allowed in the `Prior`. In this latter case, the `gammaParameter` denotes the rate parameter of the Poisson distribution. Note that in this case the interpretation of `gammaParameter` is reversed: Smaller (larger) values of `gammaParameter` will a-priori support smaller (respectively: larger) number of change-points.

### Author(s)

Panagiotis Papastamoulis

### References

Papastamoulis P., Furukawa T., van Rhijn N., Bromley M., Bignell E. and Rattray M. (2017). Bayesian detection of piecewise linear trends in replicated time-series with application to growth data modelling. arXiv:1709.06111 [stat.AP]

### Examples

```
# toy-example (MCMC iterations not enough)
library('beast') # load package
data("FungalGrowthDataset") # load dataset
myIndex <- c(392, 62, 3, 117) # run the sampler only for the
# specific subset of time-series
set.seed(1)
# Run MCMC sampler with very small number of iterations (nIter):
run_mcmc <- beast(myDataList = FungalGrowthDataset, subsetIndex = myIndex,
```

```

zeroNormalization = TRUE, nIter = 40, burn = 20)
# Print output:
print(run_mcmc)
# Plot output to file: "beast_plot.pdf"
plot(run_mcmc, fileName = "beast_plot_toy.pdf", timeScale=1/6, xlab = "hours", ylab = "growth")
# Run the following commands to obtain convergence:

## Not run:
# This example illustrates the package using a subset of four
#      time-series of the fungal dataset.
library('beast') # load package
data("FungalGrowthDataset") # load dataset
myIndex <- c(392, 62, 3, 117) # run the sampler only for the
#                           specific subset of time-series
set.seed(1) # optional
# Run MCMC sampler with the default number of iterations (nIter =70000):
run_mcmc <- beast(myDataList = FungalGrowthDataset, subsetIndex = myIndex,
zeroNormalization = TRUE)
# Print output:
print(run_mcmc)
# Plot output to file: "beast_plot.pdf"
plot(run_mcmc, fileName = "beast_plot.pdf", timeScale=1/6, xlab = "hours", ylab = "growth")
# NOTE 1: for a complete analysis remove the `subsetIndex = myIndex` argument.
# NOTE 2: `zeroNormalization = TRUE` is an optional argument that forces all
#      time-series to start from zero. It is not supposed to be used
#      for other applications.

## End(Not run)

```

**birthProbs***Birth Probabilities***Description**

This function defines the probability of proposing an addition of a change-point.

**Usage**

```
birthProbs(LRange)
```

**Arguments**

LRange	The range of possible values for the number of change-points.
--------	---

**Value**

probs	Vector of probabilities
-------	-------------------------

**Author(s)**

Panagiotis Papastamoulis

<code>complexityPrior</code>	<i>Complexity prior distribution</i>
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### Description

This function computes the complexity prior distribution on the number of change-points, defined as  $f(\ell) = P(\ell_n = \ell) \propto e^{-\alpha\ell \log(bT/\ell)}$ ,  $a, b > 0$ ;  $\ell = 0, 1, 2, \dots$ . Note that this distribution has exponential decrease (Castillo and van der Vaart, 2012) when  $b > 1 + e$ , so we set  $b = 3.72$ .

### Usage

```
complexityPrior(Lmax = 20, gammaParameter, nTime)
```

### Arguments

<code>Lmax</code>	maximum number of change-points (default = 20).
<code>gammaParameter</code>	positive real number, corresponding to $\alpha$ .
<code>nTime</code>	positive integer denoting the total number of time-points.

### Value

<code>logPrior</code>	Prior distribution values in the log-scale.
-----------------------	---

### Author(s)

Panagiotis Papastamoulis

### References

Castillo I. and van der Vaart A (2012). Needles and Straw in a Haystack: Posterior concentration for possibly sparse sequences. *The Annals of Statistics*, 40(4), 2069–2101.

<code>computeEmpiricalPriorParameters</code>	<i>Compute the empirical mean.</i>
--	------------------------------------

### Description

This function computes the empirical mean of the time-series.

### Usage

```
computeEmpiricalPriorParameters(myDataList, nu0 = 1, alpha0 = 1, beta0 = 1)
```

**Arguments**

myDataList	Observed multivariate time-series.
nu0	positive real number.
alpha0	positive real number.
beta0	positive real number.

**Value**

mu0	Empirical mean
-----	----------------

**Author(s)**

Panagiotis Papastamoulis

---

computePosteriorParameters  
*Compute empirical posterior parameters*

---

**Description**

Compute empirical posterior parameters

**Usage**

```
computePosteriorParameters(myDataList, priorParameters)
```

**Arguments**

myDataList	Observed data.
priorParameters	Prior parameters.

**Value**

list of posterior parameters

**Author(s)**

Panagiotis Papastamoulis

**computePosteriorParametersFree**  
*Posterior parameters*

### Description

Posterior parameters

### Usage

```
computePosteriorParametersFree(myDataList, priorParameters)
```

### Arguments

myDataList	observed data.
priorParameters	list of prior parameters.

### Value

list of posterior parameters.

### Author(s)

Panagiotis Papastamoulis

*FungalGrowthDataset*      *Fungal Growth Dataset*

### Description

Time-series dataset with  $N \times R \times T$  growth levels for  $R = 3$  replicates of  $N = 411$  objects (mutants) measured every 10 minutes for  $T = 289$  time-points. See Papastamoulis et al (2017) for a detailed description.

### Usage

*FungalGrowthDataset*

### Format

Time-series data.

### References

Papastamoulis P., Furukawa T., van Rhijn N., Bromley M., Bignell E. and Rattray M. (2017). Bayesian detection of piecewise linear trends in replicated time-series with application to growth data modelling. arXiv:[1709.06111](https://arxiv.org/abs/1709.06111) [stat.AP]

---

localProposal            *Move 3.b*

---

**Description**

Implements Move 3.b of the Metropolis-Hastings MCMC sampler.

**Usage**

```
localProposal(cutPoints, nTime, mhPropRange, startPoint)
```

**Arguments**

cutPoints	Current configuration of change-points.
nTime	Total number of time-points.
mhPropRange	Parameter $d_2$ .
startPoint	Integer, at least equal to 2.

**Value**

newState	Candidate state of the chain.
propRatio	Proposal ratio.

**Author(s)**

Panagiotis Papastamoulis

---

logLikelihoodFullModel  
Log-likelihood function.

---

**Description**

Log-likelihood function.

**Usage**

```
logLikelihoodFullModel(myData, cutPoints, theta, startPoint)
```

**Arguments**

myData	data
cutPoints	change-points.
theta	means.
startPoint	optional integer at least equal to 2.

**Value**

log-likelihood value.

**Author(s)**

Panagiotis Papastamoulis

---

logPrior

*Log-prior.*

---

**Description**

Logarithm of the prior distribution on the number of change-points.

**Usage**

```
logPrior(cutPoints, nTime, startPoint)
```

**Arguments**

cutPoints	change-points.
nTime	number of time-points.
startPoint	optional integer, at least equal to 2.

**Value**

logarithm of the prior distribution.

**Author(s)**

Panagiotis Papastamoulis

---

mcmcSampler

*MCMC sampler*

---

**Description**

This function implements the Metropolis-Hastings MCMC sampler for individual time-series.

**Usage**

```
mcmcSampler(myData, nIter, finalIterationPdf, modelVariance, mhPropRange,
mhSinglePropRange, movesRange, startPoint, postPar, dName, timeScale,
burn, iterPerPlotPrefix, priorParameters, L = 3, LRange, tau,
gammaParameter, saveTheta, Prior = "complexity")
```

**Arguments**

<code>myData</code>	observed data.
<code>nIter</code>	number of mcmc iterations
<code>finalIterationPdf</code>	output folder
<code>modelVariance</code>	null
<code>mhPropRange</code>	positive integer
<code>mhSinglePropRange</code>	positive integer
<code>movesRange</code>	null
<code>startPoint</code>	positive integer
<code>postPar</code>	list of empirically estimated parameters
<code>dName</code>	subject ID
<code>timeScale</code>	null
<code>burn</code>	burn-in period.
<code>iterPerPlotPrefix</code>	null
<code>priorParameters</code>	prior parameters.
<code>L</code>	null
<code>LRange</code>	range of possible values of the number of change-points.
<code>tau</code>	real.
<code>gammaParameter</code>	real.
<code>saveTheta</code>	TRUE.
<code>Prior</code>	character.

**Value**

MCMC output.

**Author(s)**

Panagiotis Papastamoulis

---

myUnicodeCharacters     *Printing*

---

**Description**

Printing various unicode symbols.

**Usage**

```
myUnicodeCharacters()
```

**Value**

printed symbol

---

normalizeTime0        *Zero normalization*

---

**Description**

Zero normalization at 1st time-point

**Usage**

```
normalizeTime0(myDataList)
```

**Arguments**

myDataList     data

**Value**

null

**Author(s)**

Panagiotis Papastamoulis

---

**plot.beast.object** *Plot function*

---

**Description**

This function plots objects returned by the `beast` function. All output is diverted to a pdf file provided in the `fileName` argument.

**Usage**

```
## S3 method for class 'beast.object'  
plot(x, fileName, width, height, pointsize, ylab, xlab, timeScale, myPal, boxwex, ...)
```

**Arguments**

<code>x</code>	An object of class <code>beast.object</code> , which is returned by the <code>beast</code> function.
<code>fileName</code>	Name of the output pdf file.
<code>width</code>	Width of pdf file. Default: 9
<code>height</code>	Height pdf file. Default: 6
<code>pointsize</code>	Pointsize. Default: 12
<code>ylab</code>	<i>y</i> -axis label. Default: <code>x</code> .
<code>xlab</code>	<i>x</i> -axis label. Default: <code>t</code> .
<code>timeScale</code>	A multiplicative-factor which will be used to scale the <i>x</i> -axis labels. For example, if time-points correspond to 10-minute periods, then setting <code>timeScale</code> = 1/6 will make the <i>x</i> -axis labels correspond to hours. Default: <code>timeScale</code> = 1.
<code>myPal</code>	Vector of colors that will be used to produce the plot with all time-series overlaid. If the distinct values of the inferred numbers of change-points is less than 10, the Set1 palette of the <code>RColorBrewer</code> library is used. Otherwise, the user has to manually define the colors.
<code>boxwex</code>	A scale factor to be applied to all boxes. The appearance of the plot can be improved by making the boxes narrower or wider. Default: 0.2.
...	ignored.

**Details**

The function will produce a plot with all time-series coloured according to the corresponding number of change-points. In addition, it will generate individual plots per time-series displaying the observed data with boxplots which summarize the posterior distribution of change-points locations, conditionally on the most probable number of change-points.

**Author(s)**

Panagiotis Papastamoulis

`print.beast.object`      *Print function*

### Description

This function prints a summary of objects returned by the `beast` function.

### Usage

```
## S3 method for class 'beast.object'
print(x, ...)
```

### Arguments

<code>x</code>	An object of class <code>beast.object</code> , which is returned by the <code>beast</code> function.
<code>...</code>	ignored.

### Details

The function prints a summary of the most probable number (MAP) of change-points per time-series in the form of a table, as well as a list containing the MAP number of change-points and the corresponding locations (posterior medians) per time-series.

### Author(s)

Panagiotis Papastamoulis

`proposeTheta`      *Move 2*

### Description

Proposes an update of  $\theta$  according to Metropolis-Hastings move 2.

### Usage

```
proposeTheta(thetaOld, tau, alpha, beta)
```

### Arguments

<code>thetaOld</code>	Current values
<code>tau</code>	Parameter $c$ .
<code>alpha</code>	null
<code>beta</code>	null

**Value**

`mean` proposed values.

**Author(s)**

Panagiotis Papastamoulis

---

`simMultiIndNormInvGamma`

*Prior random numbers*

---

**Description**

Generation of mean values according to the prior

**Usage**

```
simMultiIndNormInvGamma(mu, nu, alpha, beta)
```

**Arguments**

<code>mu</code>	means
<code>nu</code>	precision parameter
<code>alpha</code>	prior parameter
<code>beta</code>	prior parameter

**Value**

`null`

**Author(s)**

Panagiotis Papastamoulis

`simulateFromPrior`      *Generate change-points according to the prior*

### Description

Generate change-points according to the prior distribution conditionally on a given number of change-points.

### Usage

```
simulateFromPrior(nTime, startPoint, L = 3)
```

### Arguments

<code>nTime</code>	Number of time-points
<code>startPoint</code>	At least equal to 2.
<code>L</code>	null

### Value

<code>cutPoints</code>	Change-point locations
------------------------	------------------------

### Author(s)

Panagiotis Papastamoulis

`singleLocalProposal`      *Move 3.b*

### Description

Implement Metropolis-Hastings Move 3.b.

### Usage

```
singleLocalProposal(cutPoints, nTime, mhSinglePropRange, startPoint)
```

### Arguments

<code>cutPoints</code>	Current state
<code>nTime</code>	Number of time-points
<code>mhSinglePropRange</code>	Prior parameter.
<code>startPoint</code>	Optional.

**Value**

<code>newState</code>	Candidate state
<code>propRatio</code>	Proposal ratio

**Author(s)**

Panagiotis Papastamoulis

`truncatedPoisson`      *Truncated Poisson pdf*

**Description**

Probability density function of the truncated Poisson distribution.

**Usage**

```
truncatedPoisson(Lmax = 20, gammaParameter = 1)
```

**Arguments**

<code>Lmax</code>	Max number
<code>gammaParameter</code>	Location parameter.

**Value**

<code>logPrior</code>	Log-pdf values
-----------------------	----------------

**Author(s)**

Panagiotis Papastamoulis

`updateNumberOfCutpoints`  
    *Move 1*

**Description**

Update the number of change-points according to Metropolis-Hastings move 1.

**Usage**

```
updateNumberOfCutpoints(cutPoints, nTime, startPoint, LRange, birthProbs)
```

**Arguments**

cutPoints	Current configuration
nTime	Number of time-points
startPoint	Optional integer
LRange	Range of possible values
birthProbs	Birth probabilities

**Value**

newState	Candidate state
propRatio	Proposal ratio

**Author(s)**

Panagiotis Papastamoulis

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