Package ‘rjags’

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Description

The rjags package provides an interface from R to the JAGS library for Bayesian data analysis. JAGS uses Markov Chain Monte Carlo (MCMC) to generate a sequence of dependent samples from the posterior distribution of the parameters.

Details

JAGS is a clone of BUGS (Bayesian analysis Using Gibbs Sampling). See Lunn et al (2009) for a history of the BUGS project. Note that the rjags package does not include a copy of the JAGS library: you must install this separately. For instructions on downloading JAGS, see the home page at http://mcmc-jags.sourceforge.net.

To fully understand how JAGS works, you need to read the JAGS User Manual. The manual explains the basics of modelling with JAGS and shows the functions and distributions available in the dialect of the BUGS language used by JAGS. It also describes the command line interface. The rjags package does not use the command line interface but provides equivalent functionality using R functions.

Analysis using the rjags package proceeds in steps:

1. Define the model using the BUGS language in a separate file.
2. Read in the model file using the jags.model function. This creates an object of class “jags”.
3. Update the model using the update method for “jags” objects. This constitutes a ‘burn-in’ period.
4. Extract samples from the model object using the coda.samples function. This creates an object of class “mcmc.list” which can be used to summarize the posterior distribution. The coda package also provides convergence diagnostics to check that the output is valid for analysis (see Plummer et al 2006).

Author(s)

Martyn Plummer
References


### adapt

*Adaptive phase for JAGS models*

**Description**

Update the model in adaptive mode.

**Usage**

```r
adapt(object, n.iter, end.adaptation=FALSE, ...)
```

**Arguments**

<table>
<thead>
<tr>
<th>Argument</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>object</td>
<td>a jags model object</td>
</tr>
<tr>
<td>n.iter</td>
<td>length of the adaptive phase</td>
</tr>
<tr>
<td>end.adaptation</td>
<td>logical flag. If TRUE then adaptive mode will be turned off on exit.</td>
</tr>
<tr>
<td>...</td>
<td>additional arguments to the update method</td>
</tr>
</tbody>
</table>

**Details**

This function is not normally called by the user. It is called by the `jags.model` function when the model object is created.

When a JAGS model is compiled, it may require an initial sampling phase during which the samplers adapt their behaviour to maximize their efficiency (e.g. a Metropolis-Hastings random walk algorithm may change its step size). The sequence of samples generated during this adaptive phase is not a Markov chain, and therefore may not be used for posterior inference on the model.

The `adapt` function updates the model for `n.iter` iterations in adaptive mode. Then each sampler reports whether it has achieved optimal performance (e.g. whether the rejection rate of a Metropolis-Hasting sampler is close to the theoretical optimum). If any sampler reports failure of this test then `adapt` returns `FALSE`.

If `end.adaptation = TRUE`, then adaptive mode is turned off on exit, and further calls to `adapt()` do nothing. The model may be maintained in adaptive mode with the default option `end.adaptation = FALSE` so that successive calls to `adapt()` may be made until adaptation is satisfactory.

**Value**

Returns `TRUE` if all the samplers in the model have successfully adapted their behaviour to optimum performance and `FALSE` otherwise.
coda.samples  

Author(s)

Martyn Plummer


coda.samples  Generate posterior samples in mcmc.list format

Description

This is a wrapper function for jags.samples which sets a trace monitor for all requested nodes, updates the model, and coerces the output to a single mcmc.list object.

Usage

coda.samples(model, variable.names, n.iter, thin = 1, na.rm=TRUE, ...)

Arguments

- **model**: a jags model object
- **variable.names**: a character vector giving the names of variables to be monitored
- **n.iter**: number of iterations to monitor
- **thin**: thinning interval for monitors
- **na.rm**: logical flag that indicates whether variables containing missing values should be omitted. See details.
- **...**: optional arguments that are passed to the update method for jags model objects

Details

If na.rm=TRUE (the default) then elements of a variable that are missing (NA) for any iteration in at least one chain will be dropped.

This argument was added to handle incompletely defined variables. From JAGS version 4.0.0, users may monitor variables that are not completely defined in the BUGS language description of the model, e.g. if y[i] is defined in a for loop starting from i=3 then y[1],y[2] are not defined. The user may still monitor variable y and the monitored values corresponding to y[1],y[2] will have value NA for all iterations in all chains. Most of the functions in the coda package cannot handle missing values so these variables are dropped by default.

Value

An mcmc.list object.

Author(s)

Martyn Plummer
See Also

jags.samples

Examples

```r
data(LINE)
LINE$recompile()
LINE.out <- coda.samples(LINE, c("alpha","beta","sigma"), n.iter=1000)
summary(LINE.out)
```

control

Advanced control over JAGS

Description

JAGS modules contain factory objects for samplers, monitors, and random number generators for a JAGS model. These functions allow fine-grained control over which factories are active.

Usage

`list.factories(type)`

`set.factory(name, type, state)`

Arguments

- **name**: name of the factory to set
- **type**: type of factory to query or set. Possible values are "sampler", "monitor", or "rng"
- **state**: a logical. If TRUE then the factory will be active, otherwise the factory will become inactive.

Value

`list.factories` returns a data frame with two columns, the first column shows the names of the factory objects in the currently loaded modules, and the second column is a logical vector indicating whether the corresponding factory is active or not.

`set.factory` is called to change the future behaviour of factory objects. If a factory is set to inactive then it will be skipped.

Note

When a module is loaded, all of its factory objects are active. This is also true if a module is unloaded and then reloaded.

Author(s)

Martyn Plummer
Examples

```r
list.factories("sampler")
list.factories("monitor")
list.factories("rng")
set.factory("base::Slice", "sampler", FALSE)
list.factories("sampler")
set.factory("base::Slice", "sampler", TRUE)
```

dic.samples  Generate penalized deviance samples

Description

Function to extract random samples of the penalized deviance from a jags model.

Usage

```r
dic.samples(model, n.iter, thin = 1, type, ...)
```

Arguments

- `model`: a jags model object
- `n.iter`: number of iterations to monitor
- `thin`: thinning interval for monitors
- `type`: type of penalty to use
- `...`: optional arguments passed to the update method for jags model objects

Details

The `dic.samples` function generates penalized deviance statistics for use in model comparison. The two alternative penalized deviance statistics generated by `dic.samples` are the deviance information criterion (DIC) and the penalized expected deviance. These are chosen by giving the values "pD" and "popt" respectively as the `type` argument.

DIC (Spiegelhalter et al 2002) is calculated by adding the “effective number of parameters” (pD) to the expected deviance. The definition of pD used by `dic.samples` is the one proposed by Plummer (2002) and requires two or more parallel chains in the model.

DIC is an approximation to the penalized plug-in deviance, which is used when only a point estimate of the parameters is of interest. The DIC approximation only holds asymptotically when the effective number of parameters is much smaller than the sample size, and the model parameters have a normal posterior distribution.

The penalized expected deviance (Plummer 2008) is calculated by adding the optimism (popt) to the expected deviance. The popt penalty is at least twice the size of the pD penalty, and penalizes complex models more severely.
Value
An object of class “dic”. This is a list containing the following elements:

- deviance: A numeric vector, with one element for each observed stochastic node, containing the mean deviance for that node
- penalty: A numeric vector, with one element for each observed stochastic node, containing an estimate of the contribution towards the penalty
- type: A string identifying the type of penalty: “pD” or “popt”

Note
The popt penalty is estimated by importance weighting, and may be numerically unstable.

Author(s)
Martyn Plummer

References


See Also
diffdic

diffdic  Differences in penalized deviance

Description
Compare two models by the difference of two dic objects.

Usage
dic1 - dic2
diffdic(dic1, dic2)

Arguments
dic1, dic2  Objects inheriting from class “dic”
A `diffdic` object represents the difference in penalized deviance between two models. A negative value indicates that `dic1` is preferred and vice versa.

An object of class “`diffdic`”. This is a numeric vector with an element for each observed stochastic node in the model.

The `diffdic` class has its own print method, which will display the sum of the differences, and its sample standard deviation.

The problem of determining what is a noteworthy difference in DIC (or other penalized deviance) between two models is currently unsolved. Following the results of Ripley (1996) on the Akaike Information Criterion, Plummer (2008) argues that there is no absolute scale for comparison of two penalized deviance statistics, and proposes that the difference should be calibrated with respect to the sample standard deviation of the individual contributions from each observed stochastic node.

**Author(s)**

Martyn Plummer

**References**


See Also

dic

---

**Description**

`jags.model` is used to create an object representing a Bayesian graphical model, specified with a BUGS-language description of the prior distribution, and a set of data.

**Usage**

```r
jags.model(file, data, inits,
            n.chains = 1, n.adapt=1000, quiet=FALSE)
```
Arguments

- **file**: the name of the file containing a description of the model in the JAGS dialect of the BUGS language. Alternatively, `file` can be a readable text-mode connection, or a complete URL.
- **data**: a list or environment containing the data. Any numeric objects in `data` corresponding to node arrays used in `file` are taken to represent the values of observed nodes in the model.
- **inits**: optional specification of initial values in the form of a list or a function (see initialization below). If omitted, initial values will be generated automatically. It is an error to supply an initial value for an observed node.
- **n.chains**: the number of parallel chains for the model.
- **n.adapt**: the number of iterations for adaptation. See `adapt` for details. If `n.adapt = 0` then no adaptation takes place.
- **quiet**: if `TRUE` then messages generated during compilation will be suppressed, as well as the progress bar during adaptation.

Value

- `jags.model` returns an object inheriting from class `jags` which can be used to generate dependent samples from the posterior distribution of the parameters.

An object of class `jags` is a list of functions that share a common environment. This environment encapsulates the state of the model, and the functions can be used to query or modify the model state.

- **ptr()**: Returns an external pointer to an object created by the JAGS library.
- **data()**: Returns a list containing the data that define the observed nodes in the model.
- **model()**: Returns a character vector containing the BUGS-language representation of the model.
- **state(internal=FALSE)**: Returns a list of length equal to the number of parallel chains in the model. Each element of the list is itself a list containing the current parameter values in that chain. If `internal=TRUE` then the returned lists also include the RNG names (`.RNG.name`) and states (`.RNG.state`). This is not the user-level interface: use the `coef.jags` method instead.
- **update(n.iter)**: Updates the model by `n.iter` iterations. This is not the user-level interface: use the `update.jags` method instead.

Initialization

There are various ways to specify initial values for a JAGS model. If no initial values are supplied, then they will be generated automatically by JAGS. See the JAGS User Manual for details. Otherwise, the options are as follows:

1. A list of numeric values. Initial values for a single chain may be supplied as a named list of numeric values. If there are multiple parallel chains then the same list is re-used for each chain.
2. A list of lists. Distinct initial values for each chain may be given as a list of lists. In this case, the list should have the same length as the number of chains in the model.

3. A function. A function may be supplied that returns a list of initial values. The function is called repeatedly to generate initial values for each chain. Normally this function should call some random number generating functions so that it returns different values every time it is called. The function should either have no arguments, or have a single argument named chain. In the latter case, the supplied function is called with the chain number as argument. In this way, initial values may be generated that depend systematically on the chain number.

**Random number generators**

Each chain in a model has its own random number generator (RNG). RNGs and their initial seed values are assigned automatically when the model is created. The automatic seeds are calculated from the current time.

If you wish to make the output from the model reproducible, you may specify the RNGs to be used for each chain, and their starting seeds as part of the inits argument (see Initialization above). This is done by supplementing the list of initial parameter values for a given chain with two additional elements named “.RNG.name”, and “.RNG.seed”:

- **.RNG.name** a character vector of length 1. The names of the RNGs supplied in the base module are:
  - “base::Wichmann-Hill”
  - “base::Marsaglia-Multicarry”
  - “base::Super-Duper”
  - “base::Mersenne-Twister”

- **.RNG.seed** a numeric vector of length 1 containing an integer value.

If the lecuyer module is loaded, it provides “lecuyer::RngStream”

Note that it is also possible to specify “.RNG.state” rather than “.RNG.seed” - see for example the output of `parallel.seeds`

**Author(s)**

Martyn Plummer

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**Description**

A JAGS module is a dynamically loaded library that extends the functionality of JAGS. These functions load and unload JAGS modules and show the names of the currently loaded modules.

**Usage**

```
load.module(name, path, quiet=FALSE)
unload.module(name, quiet=FALSE)
list.modules()
```
jags.object

Arguments

name name of the load module to be loaded
path file path to the location of the DLL. If omitted, the option jags.moddir is used to locate the modules
quiet a logical. If TRUE, no message will be printed about loading the package

Author(s)

Martyn Plummer

Examples

list.modules()
load.module("glm")
list.modules()
unload.module("glm")
list.modules()

---

table

| jags.object | Functions for manipulating jags model objects |

Description

A jags object represents a Bayesian graphical model described using the BUGS language.

Usage

```r
## S3 method for class 'jags'
coef(object, chain=1, ...)
## S3 method for class 'jags'
variable.names(object, ...)
list.samplers(object)
```

Arguments

object a jags model object
chain chain number to query
... additional arguments to the call (ignored)

Value

The coef function returns a list with an entry for each Node array that contains an unobserved Node. Elements corresponding to observed Nodes or deterministic Nodes are given missing values.

The variable.names function returns a character vector of names of node arrays used in the model.

The list.samplers function returns a named list with an entry for each Sampler used by the model. Each list element is a character vector containing the names of stochastic Nodes that are updated.
The names of the list elements indicate the sampling methods that are used to update each block. Stochastic nodes that are updated by forward sampling from the prior are not listed.

**Author(s)**

Martyn Plummer

**Examples**

```r
data(LINE)
LINE$recompile()
coef(LINE)
variable.names(LINE)
list.samplers(LINE)
```

---

**jags.samples**  
*Generate posterior samples*

**Description**

Function to extract random samples from the posterior distribution of the parameters of a jags model.

**Usage**

```r
jags.samples(model, variable.names, n.iter, thin = 1,
        type="trace", force.list=FALSE, ...)
```

**Arguments**

- **model**: a jags model object
- **variable.names**: a character vector giving the names of variables to be monitored
- **n.iter**: number of iterations to monitor
- **thin**: thinning interval for monitors
- **type**: type of monitor (can be vectorised)
- **force.list**: option to consistently return a named list of monitor types even if a single monitor type is requested
- **...**: optional arguments passed to the update method for jags model objects

**Details**

The `jags.samples` function creates monitors for the given variables, runs the model for `n.iter` iterations and returns the monitored samples.
jags.version

Description
Get the version of the JAGS library which is currently linked to this R session

Usage
jags.version()

Value
The version of JAGS formatted as a package version string (see package_version)
Description
The LINE model is a trivial linear regression model with only 5 observations. It's main use is to allow automated checks of the \texttt{rjags} package.

Format
A \texttt{jags.model} object, which must be recompiled before use.

\begin{verbatim}
mcarray.object  Objects for representing MCMC output
\end{verbatim}

Description
An \texttt{mcarray} object is used by the \texttt{jags.samples} function to represent MCMC output from a JAGS model. It is an array with named dimensions, for which the dimensions "iteration" and "chain" have a special status.

Usage
\begin{verbatim}
## S3 method for class 'mcarray'
summary(object, FUN, ...)
## S3 method for class 'mcarray'
print(x, ...)
## S3 method for class 'mcarray'
as.mcmc.list(x, ...)
\end{verbatim}

Arguments
\begin{itemize}
  \item \texttt{object,x} an \texttt{mcarray} object
  \item \texttt{FUN} a function to be used to generate summary statistics
  \item \ldots additional arguments to the call
\end{itemize}

Details
The \texttt{coda} package defines \texttt{mcmc} objects for representing output from an MCMC sampler, and \texttt{mcmc.list} for representing output from multiple parallel chains. These objects emphasize the time-series aspect of the MCMC output, but lose the original array structure of the variables they represent. The \texttt{mcarray} class attempts to rectify this by preserving the dimensions of the original node array defined in the JAGS model.
Value

The summary method for mcarray objects applies the given function to the array, marginalizing the "chain" and "iteration" dimensions.
The print method applies the summary function with FUN=mean.
The as.mcmc.list method coerces an mcarray to an mcmc.list object so that the diagnostics provided by the coda package can be applied to the MCMC output it represents.

Author(s)

Martyn Plummer

parallel

Get initial values for parallel RNGs

Description

On a multi-processor system, you may wish to run parallel chains using multiple jags.model objects, each running a single chain on a separate processor. This function returns a list of values that may be used to initialize the random number generator of each chain.

Usage

parallel.seeds(factory, nchain)

Arguments

factory Name of the RNG factory to use.
nchain Number of chains for which to initialize RNGs.

Value

parallel.seeds returns a list of RNG states. Each element is a list of length 2 with the following elements:

>RNG.name The name of the RNG
>RNG.state An integer vector giving the state of the RNG.

Note

It is not yet possible to make the results of parallel.seeds reproducible. This will be fixed in a future version of JAGS.

Author(s)

Martyn Plummer
See Also

`jags.model`, section “Random number generators”, for further details on RNG initialization; `list.factories` to find the names of available RNG factories.

Examples

```r
## The BaseRNG factory generates up to four distinct types of RNG. If
## more than 4 chains are requested, it will recycle the RNG types, but
## use different initial values
parallel.seeds("base::BaseRNG", 3)

## The lecuyer module provides the RngStream factory, which allows large
## numbers of independent parallel RNGs to be generated.
load.module("lecuyer")
list.factories(type="rng")
parallel.seeds("lecuyer::RngStream", 5);
```

### Description

Read data for a JAGS model from a file.

#### Usage

```r
read.jagsdata(file)
read.bugsdata(file)
```

#### Arguments

- `file` name of a file containing a text representation of the data for a jags model

#### Details

The command line interface for JAGS reads data and initial values from a text file. The data format used for jags data files is the same as the R `dump` function. Thus the data values can be read into an R session using the `source` function, but this will create objects in the global environment. The `read.jagsdata` function is a simple wrapper that reads the data into a list instead.

OpenBUGS also reads data and initial values from a text file. The format of these files is described as "S-PLUS" format by the OpenBUGS authors. It superficially resembles the format used by the `dput` function (and in fact can be parsed by the `dget` function). However, in BUGS "S-PLUS" format, arrays are stored in row-major order instead of the column-major order used by R. The `read.bugsdata` function reads OpenBUGS "S-PLUS" format files and permutes the elements of arrays so that they appear in the correct order.

Either function returns a list which can be used as the `data` or `inits` argument of `jags.model`. 
Value

A named list of numeric vectors or arrays.

Note

Earlier versions of the rjags package had a read.data function which read data in either format, but the function name was ambiguous (There are many data file format in R) so this is now deprecated.

Author(s)

Martyn Plummer

---

**rjags-deprecated**

*Deprecated Functions in the rjags package*

---

Description

These functions are provided for compatibility with older versions of the rjags package and will soon be defunct.

Usage

```r
read.data(file, format=c("jags","bugs"))
```

Arguments

- `file`: name of a file containing a text representation of the data for a jags model
- `format`: format of the data

Details

read.data with format="jags" is a deprecated synonym for `read.jagsdata` and with format="bugs" is a deprecated synonym for `read.bugsdata`. 
update  

Update jags models

Description

Update the Markov chain associated with the model.

Usage

```r
## S3 method for class 'jags'
update(object, n.iter=1, by, progress.bar, ...)
```

Arguments

- `object`: a jags model object
- `n.iter`: number of iterations of the Markov chain to run
- `by`: refresh frequency for progress bar. See Details
- `progress.bar`: type of progress bar. Possible values are "text", "gui", and "none". See Details.
- `...`: additional arguments to the update method (ignored)

Details

Since MCMC calculations are typically long, a progress bar is displayed during the call to `update`. The type of progress bar is determined by the `progress.bar` argument. Type "text" is displayed on the R console. Type "gui" is a graphical progress bar in a new window. The progress bar is suppressed if `progress.bar` is "none" or `NULL`, if the update is less than 100 iterations, or if R is not running interactively.

The default progress bar type is taken from the option `jags.pb`.

The progress bar is refreshed every `by` iterations. The update can only be interrupted when the progress bar is refreshed. Therefore it is advisable not to set `by` to a very large value. By default `by` is either `n.iter/50` or 100, whichever is smaller.

Value

The `update` method for jags model objects modifies the original object and returns `NULL`.

Author(s)

Martyn Plummer
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