Package ‘coda’

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as.ts.mcmc

Coerce mcmc object to time series

Description

the as.ts method for mcmc objects coerces an mcmc object to a time series.

Usage

## S3 method for class 'mcmc'
as.ts(x, ...)

Arguments

x an mcmc object

... unused arguments for compatibility with generic as.ts

Author(s)

Martyn Plummer

See Also

as.ts

autocorr

Autocorrelation function for Markov chains

Description

autocorr calculates the autocorrelation function for the Markov chain mcmc.obj at the lags given by lags. The lag values are taken to be relative to the thinning interval if relative=TRUE.

High autocorrelations within chains indicate slow mixing and, usually, slow convergence. It may be useful to thin out a chain with high autocorrelations before calculating summary statistics: a thinned chain may contain most of the information, but take up less space in memory. Re-running the MCMC sampler with a different parameterization may help to reduce autocorrelation.

Usage

autocorr(x, lags = c(0, 1, 5, 10, 50), relative=TRUE)
Arguments

\(x\) \hspace{1cm} \text{an mcmc object}

\(\text{lags}\) \hspace{1cm} \text{a vector of lags at which to calculate the autocorrelation}

\(\text{relative}\) \hspace{1cm} \text{a logical flag. TRUE if lags are relative to the thinning interval of the chain, or FALSE if they are absolute difference in iteration numbers}

Value

A vector or array containing the autocorrelations.

Author(s)

Martyn Plummer

See Also

\(\text{acf, autocorr.plot.}\)

\autocorr.diag \hspace{1cm} \text{Autocorrelation function for Markov chains}

Description

\autocorr.diag calculates the autocorrelation function for the Markov chain \text{mcmc.obj} at the lags given by \text{lags}. The lag values are taken to be relative to the thinning interval if \text{relative}=\text{TRUE}. Unlike \autocorr, if \text{mcmc.obj} has many parameters it only computes the autocorrelations with itself and not the cross correlations. In cases where \autocorr would return a matrix, this function returns the diagonal of the matrix. Hence it is more useful for chains with many parameters, but may not be as helpful at spotting parameters.

If \text{mcmc.obj} is of class \text{mcmc.list} then the returned vector is the average autocorrelation across all chains.

Usage

\autocorr.diag(mcmc.obj, ...)

Arguments

\text{mcmc.obj} \hspace{1cm} \text{an object of class mcmc or mcmc.list}

\(...\) \hspace{1cm} \text{optional arguments to be passed to autocorr}

Value

A vector containing the autocorrelations.
**autocorr.plot**

**Author(s)**
Russell Almond

**See Also**
autocorr.acf, autocorr.plot.

---

**autocorr.plot**

Plot autocorrelations for Markov Chains

**Description**
Plots the autocorrelation function for each variable in each chain in x.

**Usage**
autocorr.plot(x, lag.max, auto.layout = TRUE, ask, ...)

**Arguments**
x A Markov Chain
lag.max Maximum value at which to calculate acf
auto.layout If TRUE then, set up own layout for plots, otherwise use existing one.
ask If TRUE then prompt user before displaying each page of plots. Default is dev.interactive() in R and interactive() in S-PLUS.
...
graphical parameters

**See Also**
autocorr.

---

**batchSE**

Batch Standard Error

**Description**
Effective standard deviation of population to produce the correct standard errors.

**Usage**
batchSE(x, batchSize=100)
Arguments

- `x`: An `mcmc` or `mcmc.list` object.
- `batchSize`: Number of observations to include in each batch.

Details

Because of the autocorrelation, the usual method of taking \( \text{var}(x)/n \) overstates the precision of the estimate. This method works around the problem by looking at the means of batches of the parameter. If the batch size is large enough, the batch means should be approximately uncorrelated and the normal formula for computing the standard error should work.

The batch standard error procedure is usually thought to be not as accurate as the time series methods used in `summary` and `effectiveSize`. It is included here for completeness.

Value

A vector giving the standard error for each column of `x`.

Author(s)

Russell Almond

References


See Also

`spectrumPNar`, `effectiveSize`, `summary.mcmc`

Description

`bugs2jags` converts a WinBUGS data in the format called "S-Plus" (i.e. the format created by the `dput` function) and writes it in dump format used by JAGS.

NB WinBUGS stores its arrays in row order. This is different from R and JAGS which both store arrays in column order. This difference is taken into account by `bugs2jags` which will automatically reorder the data in arrays, without changing the dimension.

Not yet available in S-PLUS.

Usage

`bugs2jags(infile, outfile)`
coda.options

Arguments

  infile     name of the input file
  outfile    name of the output file

Note

If the input file is saved from WinBUGS, it must be saved in plain text format. The default format for files saved from WinBUGS is a binary compound document format (with extension odc) that cannot be read by bugs2jags.

Author(s)

Martyn Plummer

References


See Also

dput, dump.

coda.options  Options settings for the codamenu driver

Description

coda.options is a utility function that queries and sets options for the codamenu() function. These settings affect the behaviour of the functions in the coda library only when they are called via the codamenu() interface.

The coda.options() function behaves just like the options() function in the base library, with the additional feature that coda.options(default=TRUE) will reset all options to the default values.

Options can be pretty-printed using the display.coda.options() function, which groups the options into sections.

Available options are

  bandwidth  Bandwidth function used when smoothing samples to produce density estimates. Defaults to Silverman’s “Rule of thumb”.
  combine.corr Logical option that determines whether to combine multiple chains when calculating cross-correlations.
  combine.plots Logical option that determines whether to combine multiple chains when plotting.
  combine.plots Logical option that determines whether to combine multiple chains when calculating summary statistics.
  data.saved  For internal use only.
**densplot**  Logical option that determines whether to plot a density plot when plot methods are called for mcmc objects.

**digits**  Number of significant digits to use when printing.

**frac1**  For Geweke diagnostic, fraction to use from start of chain. Defaults to 0.1

**frac2**  For Geweke diagnostic, fraction to use from end of chain. Default to 0.5.

**gr.bin**  For Geweke-Brooks plot, number of iterations to use per bin.

**gr.max**  For Geweke-Brooks plot, maximum number of bins to use. This option overrides gr.bin.

**halfwidth**  For Heidelberger and Welch diagnostic, the target value for the ratio of half width to sample mean.

**lowess**  Logical option that controls whether to plot a smooth line through a trace plot when plotting MCMC output.

**q**  For Raftery and Lewis diagnostic, the target quantile to be estimated

**r**  For Raftery and Lewis diagnostic, the required precision.

**s**  For Raftery and Lewis diagnostic, the probability of obtaining an estimate in the interval (q-r, \(q+r\)).

**quantiles**  Vector of quantiles to print when calculating summary statistics for MCMC output.

**trace**  Logical option that determines whether to plot a trace of the sampled output when plotting MCMC output.

**user.layout**  Logical option that determines whether current value of par("mfrow") should be used for plots (TRUE) or whether the optimal layout should be calculated (FALSE).

**Usage**

```r
coda.options(...)
display.coda.options(stats = FALSE, plots = FALSE, diags = FALSE)
.Coda.Options
.Coda.Options.Default
```

**Arguments**

- **stats**  logical flag: show summary statistic options?
- **plots**  logical flag: show plotting options?
- **diags**  logical flag: show plotting options?
- **...**  list of options

**See Also**

- `options`
codamenu

---

**Main menu driver for the coda package**

---

**Description**

codamenu presents a simple menu-based interface to the functions in the coda package. It is designed for users who know nothing about the R/S language.

**Usage**

codamenu()

**Author(s)**

Kate Cowles, Nicky Best, Karen Vines, Martyn Plummer

---

**Cramer**

*The Cramer-von Mises Distribution*

---

**Description**

Distribution function of the Cramer-von Mises distribution.

**Usage**

pcramer(q, eps)

**Arguments**

- q vector of quantiles.
- eps accuracy required

**Value**

pcramer gives the distribution function.

**References**


crosscorr

Cross correlations for MCMC output

Description

crosscorr calculates cross-correlations between variables in Markov Chain Monte Carlo output. If x is an mcmc.list then all chains in x are combined before calculating the correlation.

Usage

crosscorr(x)

Arguments

x an mcmc or mcmc.list object.

Value

A matrix or 3-d array containing the correlations.

See Also

crosscorr.plot, autocorr.

crosscorr.plot

Plot image of correlation matrix

Description

crosscorr.plot provides an image of the correlation matrix for x. If x is an mcmc.list object, then all chains are combined. The range [-1,1] is divided into a number of equal-length categories given by the length of col and assigned the corresponding color. By default, topographic colours are used as this makes it easier to distinguish positive and negative correlations.

Usage

crosscorr.plot (x, col = topo.colors(10), ...)

Arguments

x an mcmc or mcmc.list object
col color palette to use
... graphical parameters
cumuplot

Cumulative quantile plot

Description
Plots the evolution of the sample quantiles as a function of the number of iterations.

Usage
cumuplot(x, probs=c(0.025, 0.5, 0.975), ylab="", lty=c(2,1), lwd=c(1,2), type="l", ask, auto.layout=TRUE, col=1, ...)

Arguments
x
probs
ylab, lty, lwd, type, col
auto.layout
ask
...

Author(s)
Arni Magnusson

densplot
Probability density function estimate from MCMC output

Description
Displays a plot of the density estimate for each variable in x, calculated by the density function. For discrete-valued variables, a histogram is produced.

Usage
densplot(x, show.obs = TRUE, bwf,
          ylim, xlab, ylab = "", type="l", main, right=TRUE, ...)

See Also
crosscorr, image, topo.colors.
Arguments

- **x**: An `mcmc` or `mcmc.list` object
- **show.obs**: Function for calculating the bandwidth. If omitted, the bandwidth is calculated as 1.06 times the minimum of the standard deviation and the interquartile range divided by 1.34 times the sample size to the negative one fifth power
- **bwf**: Limits on y axis. See `plot.window`
- **xlim**: X-axis label. By default this will show the sample size and the bandwidth used for smoothing. See `plot`
- **ylab**: Y-axis label. By default, this is blank. See `plot`
- **type**: Plot type. See `plot`
- **main**: An overall title for the plot. See `title`
- **right**: Logical flag for discrete-valued distributions passed to the `hist` function. See `Details`
- **...**: Further graphical parameters

Details

For discrete-valued distributions, a histogram is produced and values are aggregated using the `pretty()` function. By default, tick marks appear to the right of the corresponding bar in the histogram and give the inclusive upper limit of the hist (`right=True`). This can be modified by specifying `right=False`. In this case tick marks appear on the left and specify the inclusive lower limit of the bar.

For continuous distributions, if a variable is bounded below at 0, or bounded in the interval [0,1], then the data are reflected at the boundary before being passed to the `density()` function. This allows correct estimation of a non-zero density at the boundary.

Note

You can call this function directly, but it is more usually called by the `plot.mcmc` function.

See Also

density, hist, plot.mcmc.

effectiveSize

**effectiveSize**

*Effective sample size for estimating the mean*

Description

Sample size adjusted for autocorrelation.

Usage

effectiveSize(x)
Arguments

- **x**: An mcmc or mcmc.list object.

Details

For a time series \( x \) of length \( N \), the standard error of the mean is the square root of \( \text{var}(x)/n \) where \( n \) is the effective sample size. \( n = N \) only when there is no autocorrelation.

Estimation of the effective sample size requires estimating the spectral density at frequency zero. This is done by the function `spectrum0.ar`.

For a mcmc.list object, the effective sizes are summed across chains. To get the size for each chain individually use `lapply(x, effectiveSize)`.

Value

A vector giving the effective sample size for each column of \( x \).

See Also

- `spectrum0.ar`.

Description

The ‘potential scale reduction factor’ is calculated for each variable in \( x \), together with upper and lower confidence limits. Approximate convergence is diagnosed when the upper limit is close to 1. For multivariate chains, a multivariate value is calculated that bounds above the potential scale reduction factor for any linear combination of the (possibly transformed) variables.

The confidence limits are based on the assumption that the stationary distribution of the variable under examination is normal. Hence the ‘transform’ parameter may be used to improve the normal approximation.

Usage

```r
gelman.diag(x, confidence = 0.95, transform=FALSE, autoburnin=TRUE,
            multivariate=TRUE)
```

Arguments

- **x**: An mcmc.list object with more than one chain, and with starting values that are overdispersed with respect to the posterior distribution.
- **confidence**: the coverage probability of the confidence interval for the potential scale reduction factor.
transform a logical flag indicating whether variables in \( x \) should be transformed to improve the normality of the distribution. If set to TRUE, a log transform or logit transform, as appropriate, will be applied.

autoburnin a logical flag indicating whether only the second half of the series should be used in the computation. If set to TRUE (default) and \( \text{start}(x) \) is less than \( \text{end}(x)/2 \) then start of series will be adjusted so that only second half of series is used.

multivariate a logical flag indicating whether the multivariate potential scale reduction factor should be calculated for multivariate chains

Value

An object of class `gelman.diag`. This is a list with the following elements:

- **psrf**: A list containing the point estimates of the potential scale reduction factor (labelled `Point est.`) and their upper confidence limits (labelled `Upper C.I.`).
- **mpsrf**: The point estimate of the multivariate potential scale reduction factor. This is NULL if there is only one variable in \( x \).

The `gelman.diag` class has its own `print` method.

Theory

Gelman and Rubin (1992) propose a general approach to monitoring convergence of MCMC output in which \( m > 1 \) parallel chains are run with starting values that are overdispersed relative to the posterior distribution. Convergence is diagnosed when the chains have ‘forgotten’ their initial values, and the output from all chains is indistinguishable. The `gelman.diag` diagnostic is applied to a single variable from the chain. It is based a comparison of within-chain and between-chain variances, and is similar to a classical analysis of variance.

There are two ways to estimate the variance of the stationary distribution: the mean of the empirical variance within each chain, \( W \), and the empirical variance from all chains combined, which can be expressed as

\[
\hat{\sigma}^2 = \frac{(n-1)W}{n} + \frac{B}{n}
\]

where \( n \) is the number of iterations and \( B/n \) is the empirical between-chain variance.

If the chains have converged, then both estimates are unbiased. Otherwise the first method will underestimate the variance, since the individual chains have not had time to range all over the stationary distribution, and the second method will overestimate the variance, since the starting points were chosen to be overdispersed.

The convergence diagnostic is based on the assumption that the target distribution is normal. A Bayesian credible interval can be constructed using a \( t \)-distribution with mean

\[
\hat{\mu} = \text{Sample mean of all chains combined}
\]

and variance

\[
\hat{V} = \hat{\sigma}^2 + \frac{B}{mn}
\]
and degrees of freedom estimated by the method of moments

\[ d = \frac{2 \times \hat{V}^2}{\text{Var}(\hat{V})} \]

Use of the t-distribution accounts for the fact that the mean and variance of the posterior distribution are estimated.

The convergence diagnostic itself is

\[ R = \sqrt{\frac{(d + 3)\hat{V}}{(d + 1)W}} \]

Values substantially above 1 indicate lack of convergence. If the chains have not converged, Bayesian credible intervals based on the t-distribution are too wide, and have the potential to shrink by this factor if the MCMC run is continued.

**Note**

The multivariate a version of Gelman and Rubin’s diagnostic was proposed by Brooks and Gelman (1998). Unlike the univariate proportional scale reduction factor, the multivariate version does not include an adjustment for the estimated number of degrees of freedom.

**References**


**See Also**

gelman.plot.
Arguments

- **x**: an mcmc object
- **bin.width**: Number of observations per segment, excluding the first segment which always has at least 50 iterations.
- **max.bins**: Maximum number of bins, excluding the last one.
- **confidence**: Coverage probability of confidence interval.
- **transform**: Automatic variable transformation (see gelman.diag)
- **autoburnin**: Remove first half of sequence (see gelman.diag)
- **auto.layout**: If TRUE then, set up own layout for plots, otherwise use existing one.
- **ask**: Prompt user before displaying each page of plots. Default is dev.interactive() in R and interactive() in S-PLUS.
- **col**: graphical parameter (see par)
- **lty**: graphical parameter (see par)
- **xlab**: graphical parameter (see par)
- **ylab**: graphical parameter (see par)
- **type**: graphical parameter (see par)
- **...**: further graphical parameters.

Details

The Markov chain is divided into bins according to the arguments `bin.width` and `max.bins`. Then the Gelman-Rubin shrink factor is repeatedly calculated. The first shrink factor is calculated with observations 1:50, the second with observations 1 : (50 + `bin.width`), the third contains samples 1 : (50 + 2 * `bin.width`) and so on. If the chain has less than 50 + `bin.width` iterations then `gelman.diag` will exit with an error.

Theory

A potential problem with `gelman.diag` is that it may mis-diagnose convergence if the shrink factor happens to be close to 1 by chance. By calculating the shrink factor at several points in time, `gelman.plot` shows if the shrink factor has really converged, or whether it is still fluctuating.

References


See Also

`gelman.diag`
Geweke's convergence diagnostic

**Description**

Geweke (1992) proposed a convergence diagnostic for Markov chains based on a test for equality of the means of the first and last part of a Markov chain (by default the first 10% and the last 50%). If the samples are drawn from the stationary distribution of the chain, the two means are equal and Geweke's statistic has an asymptotically standard normal distribution.

The test statistic is a standard Z-score: the difference between the two sample means divided by its estimated standard error. The standard error is estimated from the spectral density at zero and so takes into account any autocorrelation.

The Z-score is calculated under the assumption that the two parts of the chain are asymptotically independent, which requires that the sum of \( \frac{1}{2} \) and \( \frac{R}{N} \) be strictly less than 1.

**Usage**

\[
geweke.diag(x, frac1=0.1, frac2=0.5)\]

**Arguments**

- **x**: an mcmc object
- **frac1**: fraction to use from beginning of chain
- **frac2**: fraction to use from end of chain

**Value**

Z-scores for a test of equality of means between the first and last parts of the chain. A separate statistic is calculated for each variable in each chain.

**References**


**See Also**

- `geweke.plot`
Description

If `geweke.diag` indicates that the first and last part of a sample from a Markov chain are not drawn from the same distribution, it may be useful to discard the first few iterations to see if the rest of the chain has "converged". This plot shows what happens to Geweke’s Z-score when successively larger numbers of iterations are discarded from the beginning of the chain. To preserve the asymptotic conditions required for Geweke’s diagnostic, the plot never discards more than half the chain.

The first half of the Markov chain is divided into \( \text{nbins} - 1 \) segments, then Geweke’s Z-score is repeatedly calculated. The first Z-score is calculated with all iterations in the chain, the second after discarding the first segment, the third after discarding the first two segments, and so on. The last Z-score is calculated using only the samples in the second half of the chain.

Usage

```r
geweke.plot(x, frac1 = 0.1, frac2 = 0.5, nbins = 20,
             pvalue = 0.05, auto.layout = TRUE, ask, ...)
```

Arguments

- `x`: an `mcmc` object
- `frac1`: fraction to use from beginning of chain.
- `frac2`: fraction to use from end of chain.
- `nbins`: Number of segments.
- `pvalue`: p-value used to plot confidence limits for the null hypothesis.
- `auto.layout`: If `TRUE` then, set up own layout for plots, otherwise use existing one.
- `ask`: If `TRUE` then prompt user before displaying each page of plots. Default is `dev.interactive()` in R and `interactive()` in S-PLUS.
- `...`: Graphical parameters.

Note

The graphical implementation of Geweke’s diagnostic was suggested by Steve Brooks.

See Also

`geweke.diag`
Description

heidel.diag is a run length control diagnostic based on a criterion of relative accuracy for the estimate of the mean. The default setting corresponds to a relative accuracy of two significant digits.

heidel.diag also implements a convergence diagnostic, and removes up to half the chain in order to ensure that the means are estimated from a chain that has converged.

Usage

heidel.diag(x, eps=0.1, pvalue=0.05)

Arguments

x        An mcmc object
eps      Target value for ratio of halfwidth to sample mean
pvalue   significance level to use

Details

The convergence test uses the Cramer-von-Mises statistic to test the null hypothesis that the sampled values come from a stationary distribution. The test is successively applied, firstly to the whole chain, then after discarding the first 10%, 20%, ..., of the chain until either the null hypothesis is accepted, or 50% of the chain has been discarded. The latter outcome constitutes ‘failure’ of the stationarity test and indicates that a longer MCMC run is needed. If the stationarity test is passed, the number of iterations to keep and the number to discard are reported.

The half-width test calculates a 95% confidence interval for the mean, using the portion of the chain which passed the stationarity test. Half the width of this interval is compared with the estimate of the mean. If the ratio between the half-width and the mean is lower than eps, the halfwidth test is passed. Otherwise the length of the sample is deemed not long enough to estimate the mean with sufficient accuracy.

Theory

The heidel.diag diagnostic is based on the work of Heidelberger and Welch (1983), who combined their earlier work on simulation run length control (Heidelberger and Welch, 1981) with the work of Schruben (1982) on detecting initial transients using Brownian bridge theory.

Note

If the half-width test fails then the run should be extended. In order to avoid problems caused by sequential testing, the test should not be repeated too frequently. Heidelberger and Welch (1981) suggest increasing the run length by a factor 1 > 1.5, each time, so that estimate has the same, reasonably large, proportion of new data.
HPDinterval

References


HPDinterval

Highest Posterior Density intervals

Description

Create Highest Posterior Density (HPD) intervals for the parameters in an MCMC sample.

Usage

HPDinterval(obj, prob = 0.95, ...)

## S3 method for class 'mcmc'
HPDinterval(obj, prob = 0.95, ...)

## S3 method for class 'mcmc.list'
HPDinterval(obj, prob = 0.95, ...)

Arguments

obj The object containing the MCMC sample - usually of class "mcmc" or "mcmc.list".

prob A numeric scalar in the interval (0,1) giving the target probability content of the intervals. The nominal probability content of the intervals is the multiple of 1/ncrow(obj) nearest to prob.

... Optional additional arguments for methods. None are used at present.

Details

For each parameter the interval is constructed from the empirical cdf of the sample as the shortest interval for which the difference in the ecdf values of the endpoints is the nominal probability. Assuming that the distribution is not severely multimodal, this is the HPD interval.

Value

For an "mcmc" object, a matrix with columns "lower" and "upper" and rows corresponding to the parameters. The attribute "Probability" is the nominal probability content of the intervals. A list of such matrices is returned for an "mcmc.list" object.

Author(s)

Douglas Bates
Examples

data(line)
HPDinterval(line)

Simple linear regression example

Description

Sample MCMC output from a simple linear regression model given in the BUGS manual.

Usage

data(line)

Format

An mcmc object

Source


Markov Chain Monte Carlo Objects

Description

The function mcmc is used to create a Markov Chain Monte Carlo object. The input data are taken to be a vector, or a matrix with one column per variable.

If the optional arguments start, end, and thin are omitted then the chain is assumed to start with iteration 1 and have thinning interval 1. If data represents a chain that starts at a later iteration, the first iteration in the chain should be given as the start argument. Likewise, if data represents a chain that has already been thinned, the thinning interval should be given as the thin argument.

An mcmc object may be summarized by the summary function and visualized with the plot function.

MCMC objects resemble time series (ts) objects and have methods for the generic functions time, start, end, frequency and window.

Usage

mcmc(data= NA, start = 1, end = numeric(0), thin = 1)
as.mcmc(x, ...)
is.mcmc(x)
Arguments

- **data**: a vector or matrix of MCMC output
- **start**: the iteration number of the first observation
- **end**: the iteration number of the last observation
- **thin**: the thinning interval between consecutive observations
- **x**: An object that may be coerced to an mcmc object
- **...**: Further arguments to be passed to specific methods

Note

The format of the mcmc class has changed between coda version 0.3 and 0.4. Older mcmc objects will now cause `is.mcmc` to fail with an appropriate warning message. Obsolete mcmc objects can be upgraded with the `mcmcupgrade` function.

Author(s)

Martyn Plummer

See Also

`mcmc.list`, `mcmcupgrade`, `thin`, `window.mcmc`, `summary.mcmc`, `plot.mcmc`.

mcmc.convert

Conversions of MCMC objects

Description

These are methods for the generic functions `as.matrix()`, `as.array()` and `as.mcmc()`.

- `as.matrix()` strips the MCMC attributes from an mcmc object and returns a matrix. If `iters = TRUE` then a column is added with the iteration number. For mcmc.list objects, the rows of multiple chains are concatenated and, if `chains = TRUE` a column is added with the chain number.

- mcmc.list objects can be coerced to 3-dimensional arrays with the `as.array()` function.

An mcmc.list object with a single chain can be coerced to an mcmc object with `as.mcmc()`. If the argument has multiple chains, this causes an error.

Usage

```r
## S3 method for class 'mcmc'
as.matrix(x, iters = FALSE, ...)
## S3 method for class 'mcmc.list'
as.matrix(x, iters = FALSE, chains = FALSE, ...)
## S3 method for class 'mcmc.list'
as.array(x, drop, ...)
```
mcmc.list

Arguments

x An mcmc or mcmc.list object
iters logical flag: add column for iteration number?
chains logical flag: add column for chain number? (if mcmc.list)
drop logical flag: if TRUE the result is coerced to the lowest possible dimension
... optional arguments to the various methods

See Also

as.matrix, as.array, as.mcmc

Description

The function 'mcmc.list' is used to represent parallel runs of the same chain, with different starting values and random seeds. The list must be balanced: each chain in the list must have the same iterations and the same variables.

Diagnostic functions which act on mcmc objects may also be applied to mcmc.list objects. In general, the chains will be combined, if this makes sense, otherwise the diagnostic function will be applied separately to each chain in the list.

Since all the chains in the list have the same iterations, a single time dimension can be ascribed to the list. Hence there are time series methods time, window, start, end, frequency and thin for mcmc.list objects.

An mcmc.list can be indexed as if it were a single mcmc object using the [ operator (see examples below). The [[ operator selects a single mcmc object from the list.

Usage

mcmc.list(...) as.mcmc.list(x, ...) is.mcmc.list(x)

Arguments

... a list of mcmc objects
x an object that may be coerced to mcmc.list

Author(s)

Martyn Plummer
mcmc.subset

See Also

mcmc.

Examples

data(line)
x1 <- line[1]  # Select first chain
x2 <- line[,1, drop=FALSE]  # Select first var from all chains
varnames(x2) == varnames(line)[1]  # TRUE

Description

These are methods for subsetting mcmc objects. You can select iterations using the first dimension and variables using the second dimension. Selecting iterations will return a vector or matrix, not an mcmc object. If you want to do row-subsetting of an mcmc object and preserve its dimensions, use the window function.

Subsetting applied to an mcmc.list object will simultaneously affect all the parallel chains in the object.

Usage

## S3 method for class 'mcmc'
x[i,j, drop=missing(i)]
## S3 method for class 'mcmc.list'
x[i,j, drop=TRUE]

Arguments

x  An mcmc object
i  Row to extract
j  Column to extract
drop  if TRUE, the redundant dimensions are dropped

See Also

[, window.mcmc

mcmc.subset
**mcmcUpgrade**  
*Upgrade mcmc objects in obsolete format*

**Description**

In previous releases of CODA, an mcmc object could be a single or multiple chains. A new class mcmc.list has now been introduced to deal with multiple chains and mcmc objects can only have data from a single chain.

Objects stored in the old format are now obsolete and must be upgraded.

**Usage**

```r
mcmcUpgrade(x)
```

**Arguments**

- `x` an obsolete mcmc object.

**Author(s)**

Martyn Plummer

**See Also**

`mcmc`

---

**mcpar**  
*Mcpar attribute of MCMC objects*

**Description**

The `mcpar` attribute of an MCMC object gives the start iteration the end iteration and the thinning interval of the chain.

It resembles the ‘tsp’ attribute of time series (ts) objects.

**Usage**

```r
mcpar(x)
```

**Arguments**

- `x` An mcmcm or mcmc.list object

**See Also**

`ts, mcmc, mcmc.list`
multi.menu

Choose multiple options from a menu

Description

multi.menu presents the user with a menu of choices labelled from 1 to the number of choices. The user may choose one or more options by entering a comma separated list. A range of values may also be specified using the ":" operator. Mixed expressions such as "1,3:5, 6" are permitted. If allow.zero is set to TRUE, one can select ‘0’ to exit without choosing an item.

Usage

multi.menu(choices, title, header, allow.zero = TRUE)

Arguments

choices Character vector of labels for choices
title Title printed before menu
header Character vector of length 2 giving column titles
allow.zero Permit 0 as an acceptable response

Value

Numeric vector giving the numbers of the options selected, or 0 if no selection is made.

Author(s)

Martyn Plummer

See Also

menu.

nchain

Dimensions of MCMC objects

Description

These functions give the dimensions of an MCMC object

niter(x) returns the number of iterations.
nvar(x) returns the number of variables.
nchain(x) returns the number of parallel chains.
Usage

niter(x)
nvar(x)
nchain(x)

Arguments

x An mcmc or mcmc.list object

Value

A numeric vector of length 1:

See Also

mcmc, mcmc.list,

---

plot.mcmc

Summary plots of mcmc objects

Description

plot.mcmc summarizes an mcmc or mcmc.list object with a trace of the sampled output and a density estimate for each variable in the chain.

Usage

## S3 method for class 'mcmc'
plot(x, trace = TRUE, density = TRUE, smooth = FALSE, bwf, auto.layout = TRUE, ask = dev.interactive(), ...)

Arguments

x an object of class mcmc or mcmc.list
trace Plot trace of each variable
density Plot density estimate of each variable
smooth Draw a smooth line through trace plots
bwf Bandwidth function for density plots
auto.layout Automatically generate output format
ask Prompt user before each page of plots
... Further arguments

Author(s)

Martyn Plummer
See Also
densplot, traceplot.

raftery.diag  Raftery and Lewis's diagnostic

Description
raftery.diag is a run length control diagnostic based on a criterion of accuracy of estimation of the quantile q. It is intended for use on a short pilot run of a Markov chain. The number of iterations required to estimate the quantile q to within an accuracy of +/- r with probability p is calculated. Separate calculations are performed for each variable within each chain.

If the number of iterations in data is too small, an error message is printed indicating the minimum length of pilot run. The minimum length is the required sample size for a chain with no correlation between consecutive samples. Positive autocorrelation will increase the required sample size above this minimum value. An estimate I (the 'dependence factor') of the extent to which autocorrelation inflates the required sample size is also provided. Values of I larger than 5 indicate strong autocorrelation which may be due to a poor choice of starting value, high posterior correlations or 'stickiness' of the MCMC algorithm.

The number of 'burn in' iterations to be discarded at the beginning of the chain is also calculated.

Usage
raftery.diag(data, q=0.025, r=0.005, s=0.95, converge.eps=0.001)

Arguments
  data  an mcmc object
  q     the quantile to be estimated.
  r     the desired margin of error of the estimate.
  s     the probability of obtaining an estimate in the interval (q-r,q+r).
  converge.eps  Precision required for estimate of time to convergence.

Value
A list with class raftery.diag. A print method is available for objects of this class. the contents of the list are

   tspar  The time series parameters of data
   params A vector containing the parameters r, s and q
   Niters The number of iterations in data
   resmatrix A 3-d array containing the results: M the length of "burn in", N the required sample size, Nmin the minimum sample size based on zero autocorrelation and I = (M + N)/Nmin the "dependence factor"
Theory

The estimated sample size for variable $U$ is based on the process $Z_t = d(U_t <= u)$ where $d$ is the indicator function and $u$ is the $q$th quantile of $U$. The process $Z_t$ is derived from the Markov chain data by marginalization and truncation, but is not itself a Markov chain. However, $Z_t$ may behave as a Markov chain if it is sufficiently thinned out. 

Note

raftery.diag calculates the smallest value of thinning interval $k$ which makes the thinned chain $Z_t^k$ behave as a Markov chain. The required sample size is calculated from this thinned sequence. Since some data is ‘thrown away’ the sample size estimates are conservative.

The criterion for the number of ‘burn in’ iterations $m$ to be discarded, is that the conditional distribution of $Z_m^k$ given $Z_0$ should be within converge.eps of the equilibrium distribution of the chain $Z_t^k$.

Note

raftery.diag is based on the FORTRAN program ‘gibbsit’ written by Steven Lewis, and available from the Statlib archive.

References


read.and.check

Read data interactively and check that it satisfies conditions

Description

Input is read interactively and checked against conditions specified by the arguments what, lower, upper and answer.in. If the input does not satisfy all the conditions, an appropriate error message is produced and the user is prompted to provide input. This process is repeated until a valid input value is entered.

Usage

read.and.check(message = "", what = numeric(), lower, upper, answer.in, default)

Arguments

message message displayed before prompting for user input.
what the type of what gives the type of data to be read.
lower lower limit of input, for numeric input only.
upper
upper limit of input, for numeric input only.
answer.in
the input must correspond to one of the elements of the vector answer.in, if supplied.
default
value assumed if user enters a blank line.

Value
The value of the valid input. When the default argument is specified, a blank line is accepted as valid input and in this case read.coda.check returns the value of default.

Note
Since the function does not return a value until it receives valid input, it extensively checks the conditions for consistency before prompting the user for input. Inconsistent conditions will cause an error.

Author(s)
Martyn Plummer

read.coda  Read output files in CODA format

Description
read.coda reads Markov Chain Monte Carlo output in the CODA format produced by OpenBUGS and JAGS. By default, all of the data in the file is read, but the arguments start, end and thin may be used to read a subset of the data. If the arguments given to start, end or thin are incompatible with the data, they are ignored.

Usage
read.coda(output.file, index.file, start, end, thin, quiet=FALSE)
read.jags(file = "jags.out", start, end, thin, quiet=FALSE)

Arguments
output.file  The name of the file containing the monitored output
index.file  The name of the file containing the index, showing which rows of the output file correspond to which variables
file  For JAGS output, the name of the output file. The extension ".out" may be omitted. There must be a corresponding ".ind" file with the same file stem.
start  First iteration of chain
end  Last iteration of chain
thin  Thinning interval for chain
quiet  Logical flag. If true, a progress summary will be printed
read.coda.interactive

Value
An object of class mcmc containing a representation of the data in the file.

Author(s)
Karen Vines, Martyn Plummer

References
Gibbs Sampling, Version 0.50. MRC Biostatistics Unit, Cambridge.

See Also
mcmc, read.coda.interactive, read.openbugs.

Description
read.coda.interactive reads Markov Chain Monte Carlo output in the format produced by the
classic BUGS program. No arguments are required. Instead, the user is prompted for the required
information.

Usage
read.coda.interactive()

Value
An object of class mcmc.list containing a representation of the data in one or more BUGS output
files.

Note
This function is normally called by the codamenu function, but can also be used on a stand-alone
basis.

Author(s)
Nicky Best, Martyn Plummer

References
Gibbs Sampling, Version 0.50. MRC Biostatistics Unit, Cambridge.
read.openbugs

Read CODA output files produced by OpenBUGS

Description

read.openbugs reads Markov Chain Monte Carlo output in the CODA format produced by OpenBUGS.

This is a convenience wrapper around the function read.coda which allows you to read all the data output by OpenBUGS by specifying only the file stem.

Usage

read.openbugs(stem="", start, end, thin, quiet=FALSE)

Arguments

stem Character string giving the stem for the output files. OpenBUGS produces files with names "<stem>CODAindex.txt", "<stem>CODAchain1.txt", "<stem>CODAchain2.txt", ...
start First iteration of chain
end Last iteration of chain
thin Thinning interval for chain
quiet Logical flag. If true, a progress summary will be printed

Value

An object of class mcmc.list containing output from all chains.

Author(s)

Martyn Plummer

References


See Also

read.coda.
rejectionRate

Description

rejectionRate calculates the fraction of time that a Metropolis–Hastings type chain rejected a proposed move. The rejection rate is calculated separately for each variable in the mcmc.obj argument, irregardless of whether the variables were drawn separately or in a block. In the latter case, the values returned should be the same.

Usage

rejectionRate(x)

Arguments

x An mcmc or mcmc.list object.

Details

For the purposes of this function, a "rejection" has occurred if the value of the time series is the same at two successive time points. This test is done naively using == and may produce problems due to rounding error.

Value

A vector containing the rejection rates, one for each variable.

Author(s)

Russell Almond

spectrum0

Describe spectral density at zero

Description

The spectral density at frequency zero is estimated by fitting a glm to the low-frequency end of the periodogram. spectrum0(x)/length(x) estimates the variance of mean(x).

Usage

spectrum0(x, max.freq = 0.5, order = 1, max.length = 200)
Arguments

- **x**: A time series.
- **max.freq**: The glm is fitted on the frequency range \((0, \text{max.freq}]\)
- **order**: Order of the polynomial to fit to the periodogram.
- **max.length**: The data \(x\) is aggregated if necessary by taking batch means so that the length of the series is less than \(\text{max.length}\). If this is set to \text{NULL} no aggregation occurs.

Details

The raw periodogram is calculated for the series \(x\) and a generalized linear model with family Gamma and log link is fitted to the periodogram.

The linear predictor is a polynomial in terms of the frequency. The degree of the polynomial is determined by the parameter \(\text{order}\).

Value

A list with the following values

- **spec**: The predicted value of the spectral density at frequency zero.

Theory

Heidelberger and Welch (1991) observed that the usual non-parametric estimator of the spectral density, obtained by smoothing the periodogram, is not appropriate for frequency zero. They proposed an alternative parametric method which consisted of fitting a linear model to the log periodogram of the batched time series. Some technical problems with model fitting in their original proposal can be overcome by using a generalized linear model.

Batching of the data, originally proposed in order to save space, has the side effect of flattening the spectral density and making a polynomial fit more reasonable. Fitting a polynomial of degree zero is equivalent to using the ‘batched means’ method.

Note

The definition of the spectral density used here differs from that used by \text{specogram}. We consider the frequency range to be between 0 and 0.5, not between 0 and \(\text{frequency}(x)/2\).

The model fitting may fail on chains with very high autocorrelation.

References


See Also

- \text{spectrum, spectrum0.ar, glm.}
Description

The spectral density at frequency zero is estimated by fitting an autoregressive model. \texttt{spectrum0(x)/length(x)} estimates the variance of mean(x).

Usage

\texttt{spectrum0.ar(x)}

Arguments

\indent \texttt{x} \hspace{1cm} A time series.

Details

The \texttt{ar()} function to fit an autoregressive model to the time series \texttt{x}. For multivariate time series, separate models are fitted for each column. The value of the spectral density at zero is then given by a well-known formula.

Value

A list with the following values

\begin{itemize}
\item \texttt{spec} \hspace{1cm} The predicted value of the spectral density at frequency zero.
\item \texttt{order} \hspace{1cm} The order of the fitted model
\end{itemize}

Note

The definition of the spectral density used here differs from that used by \texttt{spec.pgram}. We consider the frequency range to be between 0 and 0.5, not between 0 and \texttt{frequency(x)/2}.

See Also

\texttt{spectrum, spectrum0, glm}.
**summary.mcmc**  
*Summary statistics for Markov Chain Monte Carlo chains*

**Description**

`summary.mcmc` produces two sets of summary statistics for each variable:
- Mean, standard deviation, naive standard error of the mean (ignoring autocorrelation of the chain) and time-series standard error based on an estimate of the spectral density at 0.
- Quantiles of the sample distribution using the `quantiles` argument.

**Usage**

```r
## S3 method for class 'mcmc'
summary(object, quantiles = c(0.025, 0.25, 0.5, 0.75, 0.975), ...)
```

**Arguments**

- `object` an object of class `mcmc` or `mcmc.list`
- `quantiles` a vector of quantiles to evaluate for each variable
- `...` a list of further arguments

**Author(s)**

Martyn Plummer

**See Also**

`mcmc, mcmc.list`.

---

**thin**  
*Thinning interval*

**Description**

`thin` returns the interval between successive values of a time series. `thin(x)` is equivalent to `1/frequency(x)`. This is a generic function. Methods have been implemented for `mcmc` objects.

**Usage**

```r
thin(x, ...)
```
Arguments

x  a regular time series
... a list of arguments

Author(s)

Martyn Plummer

See Also

time.

time.mcmc          Time attributes for mcmc objects

Description

These are methods for mcmc objects for the generic time series functions.

Usage

## S3 method for class 'mcmc'
time(x, ...)
## S3 method for class 'mcmc'
start(x, ...)
## S3 method for class 'mcmc'
end(x, ...)
## S3 method for class 'mcmc'
thin(x, ...)

Arguments

x  an mcmc or mcmc.list object
... extra arguments for future methods

See Also

time, start, frequency, thin.
traceplot  

*Trace plot of MCMC output*

**Description**

Displays a plot of iterations vs. sampled values for each variable in the chain, with a separate plot per variable.

**Usage**

```
traceplot(x, smooth = FALSE, 
          col = 1:6, type = "l", xlab = "Iterations", ylab = "", ...)
```

**Arguments**

- `x`: An `mcmc` or `mcmc.list` object
- `smooth`: draw smooth line through trace plot
- `col`: graphical parameter (see `par`)
- `type`: graphical parameter (see `plot`)
- `xlab`: graphical parameter (see `plot`)
- `ylab`: graphical parameter (see `plot`)
- `...`: further graphical parameters

**Note**

You can call this function directly, but it is more usually called by the `plot.mcmc` function.

**See Also**

- `densplot`, `plot.mcmc`

---

trellisplots  

*Trellis plots for mcmc objects*

**Description**

These methods use the Trellis framework as implemented in the `lattice` package to produce space-conserving diagnostic plots from "mcmc" and "mcmc.list" objects. The `xyplot` methods produce trace plots. The `densityplot` methods and `qqmath` methods produce empirical density and probability plots. The `levelplot` method depicts the correlation of the series. The `acfplot` methods plot the auto-correlation in the series.

Not yet available in S-PLUS.
Usage

```r
## S3 method for class 'mcmc'
densityplot(x, data,
    outer, aspect = "xy",
    default.scales = list(relation = "free"),
    start = 1, thin = 1,
    main = attr(x, "title"),
    xlab = "",
    plot.points = "rug",
    ...
)
## S3 method for class 'mcmc.list'
densityplot(x, data,
    outer = FALSE, groups = !outer,
    aspect = "xy",
    default.scales = list(relation = "free"),
    start = 1, thin = 1,
    main = attr(x, "title"),
    xlab = "",
    plot.points = "rug",
    ...
)
```

```r
## S3 method for class 'mcmc'
levelplot(x, data, main = attr(x, "title"),
    start = 1, thin = 1,
    ...
    xlab = "", ylab = "",
    cuts = 10, at,
    col.regions = topo.colors(100),
    subset)
## S3 method for class 'mcmc.list'
levelplot(x, data,
    outer = FALSE, groups = !outer,
    aspect = "xy",
    default.scales = list(y = list(relation = "free")),
    prepanel = prepanel.qqmathline,
    start = 1, thin = 1,
    main = attr(x, "title"),
    ylab = "",
    ...
)
```

```r
## S3 method for class 'mcmc'
qqmath(x, data,
    outer, aspect = "xy",
    default.scales = list(y = list(relation = "free")),
    prepanel = prepanel.qqmathline,
    start = 1, thin = 1,
    main = attr(x, "title"),
    ...
)
## S3 method for class 'mcmc.list'
qqmath(x, data,
    outer = FALSE, groups = !outer,
    aspect = "xy",
    default.scales = list(y = list(relation = "free")),
    prepanel = prepanel.qqmathline,
    start = 1, thin = 1,
    main = attr(x, "title"),
```
```r
xyplot(x, data,
   outer, layout = c(1, nvar(x)),
   default.scales = list(y = list(relation = "free")),
   type = 'l',
   start = 1, thin = 1,
   xlab = "Iteration number",
   ylab = "",
   main = attr(x, "title"),
   ...
)
```

```r
acfplot(x, data, ...
```

```r
xyplot(x, data, outer = FALSE, groups = !outer,
   aspect = "xy", layout = c(1, nvar(x)),
   default.scales = list(y = list(relation = "free")),
   type = 'l',
   start = 1, thin = 1,
   xlab = "Iteration number",
   ylab = "",
   main = attr(x, "title"),
   ...
)
```

```r
acfplot(x, data, ...
```

```r
xyplot(x, data, outer, prepanel, panel,
   type = 'h',
   aspect = "xy",
   start = 1, thin = 1,
   lag.max = NULL,
   ylab = "Autocorrelation",
   xlab = "Lag",
   main = attr(x, "title"),
   ...
)
```

```r
acfplot(x, data, outer = FALSE, groups = !outer,
   prepanel, panel,
   type = if (groups) 'b' else 'h',
   aspect = "xy",
   start = 1, thin = 1,
   lag.max = NULL,
   ylab = "Autocorrelation",
   xlab = "Lag",
   main = attr(x, "title"),
```
Arguments

x an "mcmc" or "mcmc.list" object.
data ignored, present for consistency with generic.
outer for the "mcmc.list" methods, a logical flag to control whether multiple runs of a series are displayed in the same panel (they are if FALSE, not if TRUE). If specified in the "mcmc" methods, this argument is ignored with a warning.
groups for the "mcmc.list" methods, a logical flag to control whether the underlying lattice call will be supplied a groups argument indicating which run a data point originated from. The panel function is responsible for handling such an argument, and will usually differentiate runs within a panel by using different graphical parameters. When outer=FALSE, the default of groups is TRUE if the corresponding default panel function is able to make use of such information. When outer=FALSE, groups=TRUE will be ignored with a warning.
aspect controls the physical aspect ratio of the panel. See xyplot for details. The default for these methods is chosen carefully - check what the default plot looks like before changing this parameter.
default.scales this parameter provides a reasonable default value of the scales parameter for the method. It is unlikely that a user will wish to change this parameter. Pass a value for scales (see xyplot) instead, which will override values specified here.
type a character vector that determines if lines, points, etc. are drawn on the panel. The default values for the methods are carefully chosen. See panel.xyplot for possible values.
thin an optional thinning interval that is applied before the plot is drawn.
start an optional value for the starting point within the series. Values before the starting point are considered part of the "burn-in" of the series and dropped.
plot.points character argument giving the style in which points are added to the plot. See panel.densityplot for details.
layout a method-specific default for the layout argument to the lattice functions.
xlab,ylab,main Used to provide default axis annotations and plot labels.
cuts, at defines number and location of values where colors change
col.regions color palette used
lag.max maximum lag for which autocorrelation is computed. By default, the value chosen by acf is used
prepanel,panel suitable prepanel and panel functions for acfplot. The prepanel function omits the lag-0 auto-correlation (which is always 1) from the range calculations.
... other arguments, passed to the lattice function. Documentation of the corresponding generics in the lattice package should be consulted for possible arguments.
subset indices of the subset of the series to plot. The default is constructed from the start and thin arguments.
Value

An object of class "trellis". The relevant update method can be used to update components of the object and the print method (usually called by default) will plot it on an appropriate plotting device.

Author(s)

Deepayan Sarkar <Deepayan.Sarkar@R-project.org>

See Also

Lattice for a brief introduction to lattice displays and links to further documentation.

Examples

data(line)

## Not run:
xyplot(line)
xyplot(line[[1]], start = 10)
densityplot(line, start = 10)
qqmath(line, start = 10)
levelplot(line[[2]])
acfplot(line, outer = TRUE)

## End(Not run)

---

`varnames` Named dimensions of MCMC objects

Description

`varnames()` returns the variable names and `chanames` returns the chain names, or NULL if these are not set.

If `allow.null = FALSE` then NULL values will be replaced with canonical names.

Usage

`varnames(x, allow.null=TRUE)`
`chanames(x, allow.null=TRUE)`
`varnames(x) <- value`
`chanames(x) <- value`

Arguments

- `x` an mcmc or mcmc.list object
- `allow.null` Logical argument that determines whether the function may return NULL
- `value` A character vector, or NULL
window.mcmc

Value

A character vector, or NULL.

See Also

mcmc, mcmc.list.

Description

window.mcmc is a method for mcmc objects which is normally called by the generic function window. In addition to the generic parameters, start and end the additional parameter thin may be used to thin out the Markov chain. Setting thin=k selects every kth iteration starting with the first. Note that the value of thin is absolute not relative. The value supplied given to the parameter thin must be a multiple of thin(x).

Values of start, end and thin which are inconsistent with x are ignored, but a warning message is issued.

Usage

## S3 method for class 'mcmc'
window(x, start, end, thin, ...)

Arguments

x  an mcmc object
start  the first iteration of interest
end  the last iteration of interest
thin  the required interval between successive samples
...  further arguments for future methods

See Also

window, thin.
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