

# Package ‘Boom’

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**Title** Bayesian Object Oriented Modeling

**Description** A C++ library for Bayesian modeling, with an emphasis on Markov chain Monte Carlo. Although boom contains a few R utilities (mainly plotting functions), its primary purpose is to install the BOOM C++ library on your system so that other packages can link against it.

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**Depends** R(>= 4.5.0)

**Imports** MASS

**Suggests** testthat

**Encoding** UTF-8

**SystemRequirements** GNU Make

**NeedsCompilation** yes

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## Contents

Boom-package	3
add.segments	4
ar1.coefficient.prior	5
beta.prior	6
boxplot.mcmc.matrix	6
boxplot.true	8
check	9
check.data	10
circles	11
compare.den	12

compare.dynamic.distributions . . . . .	13
compare.many.densities . . . . .	14
compare.many.ts . . . . .	16
compare.vector.distribution . . . . .	18
diff.double.model . . . . .	18
dirichlet-distribution . . . . .	19
dirichlet.prior . . . . .	20
discrete-uniform-prior . . . . .	21
dmvn . . . . .	22
double.model . . . . .	22
external.legend . . . . .	23
gamma.prior . . . . .	25
GenerateFactorData . . . . .	26
histabunch . . . . .	27
inverse-wishart . . . . .	28
invgamma . . . . .	29
is.even . . . . .	30
lmgamma . . . . .	30
log.integrated.gaussian.likelihood . . . . .	31
lognormal.prior . . . . .	32
markov.prior . . . . .	33
match_data_frame . . . . .	34
mscan . . . . .	35
mvn.diagonal.prior . . . . .	36
mvn.independent.sigma.prior . . . . .	36
mvn.prior . . . . .	37
MvnGivenSigmaMatrixPrior . . . . .	38
normal.inverse.gamma.prior . . . . .	38
normal.inverse.wishart.prior . . . . .	39
normal.prior . . . . .	40
pairs.density . . . . .	41
plot.density.contours . . . . .	42
plot.dynamic.distribution . . . . .	43
plot.macf . . . . .	45
plot.many.ts . . . . .	46
regression.coefficient.conjugate.prior . . . . .	47
replist . . . . .	48
rmvn . . . . .	49
rvectorfunction . . . . .	50
scaled.matrix.normal.prior . . . . .	50
sd.prior . . . . .	51
sufstat.Rd . . . . .	52
suggest.burn.log.likelihood . . . . .	53
thin . . . . .	54
thin.matrix . . . . .	55
TimeSeriesBoxplot . . . . .	56
ToString . . . . .	57
traceproduct . . . . .	57

<i>Boom-package</i>	3
uniform.prior . . . . .	58
wishart . . . . .	59
<b>Index</b>	<b>61</b>

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Boom-package	<i>Boom</i>
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## Description

The Boom package provides access to the C++ BOOM library for Bayesian computation.

## Details

**Installation note for Linux users:** If you are installing Boom using [install.packages](#) on a Linux machine (and thus compiling yourself) you will almost certainly want to set the `Ncpus` argument to a large number. Windows and Mac users can ignore this advice.

The main purpose of the Boom package is not to be used directly, but to provide the BOOM C++ library for other packages to link against. The Boom package provides additional utility code for C++ authors to use when writing R packages with C++ internals. These are described in `.../inst/include/r_interface/boom_r_tools.hpp` among the package's include files.

Boom provides a collection of R functions and objects to help users format data in the manner expected by the underlying C++ code. Standard distributions that are commonly used as Bayesian priors can be specified using [BetaPrior](#), [GammaPrior](#), etc.

Boom provides a set of utilities helpful when writing unit tests for Bayesian models. See [CheckMcmcMatrix](#) and [CheckMcmcVector](#) for MCMC output, and functions like [check.probability.distribution](#) for checking function inputs

Boom provides a collection of useful plots (using base R graphics) that have proven useful for summarizing MCMC output. See [PlotDynamicDistribution](#), [PlotManyTs](#), [BoxplotTrue](#), and other code in the index with `Plot` in the title.

## See Also

Please see the following packages

- [bsts](#)
- [CausalImpact](#)

---

`add.segments`*Function to add horizontal line segments to an existing plot*

---

**Description**

Adds horizontal line segments to an existing plot. The segments are centered at  $x$  with height  $y$ . The  $x$  values are assumed to be equally spaced, so that  $\text{diff}(x)$  is a constant 'dx'. The line segments go from  $x \pm \text{half.width.factor} * dx$ , so if  $\text{half.width.factor} = .5$  there will be no gaps between segments. The default is to leave a small gap.

This function was originally used to add reference lines to side-by-side boxplots.

**Usage**

```
AddSegments(x, y, half.width.factor = 0.45, ...)
```

**Arguments**

<code>x</code>	A numeric vector giving the midpoints of the line segments.
<code>y</code>	A numeric vector of the same length as <code>x</code> giving the vertical position of the line segments
<code>half.width.factor</code>	See 'description' above.
<code>...</code>	graphical parameters controlling the type of lines used in the line segments

**Value**

Called for its side effect.

**Author(s)**

Steven L. Scott

**See Also**

[boxplot.true](#)

**Examples**

```
x <- rnorm(100)
y <- rnorm(100, 1)
boxplot(list(x=x,y=y))
AddSegments(1:2, c(0, 1)) ## add segments to the boxplot
```

---

ar1.coefficient.prior *Normal prior for an AR1 coefficient*

---

### Description

A (possibly truncated) Gaussian prior on the autoregression coefficient in an AR1 model.

### Usage

```
Ar1CoefficientPrior(mu = 0, sigma = 1, force.stationary = TRUE,  
  force.positive = FALSE, initial.value = mu)
```

### Arguments

mu	The mean of the prior distribution.
sigma	The standard deviation of the prior distribution.
force.stationary	Logical. If TRUE then the prior support for the AR1 coefficient will be truncated to (-1, 1).
force.positive	Logical. If TRUE then the prior for the AR1 coefficient will be truncated so that zero support is given to values less than zero.
initial.value	The initial value of the parameter being modeled in the MCMC algorithm.

### Details

The `Ar1CoefficientPrior()` syntax is preferred, as it more closely matches R's syntax for other constructors.

### Author(s)

Steven L. Scott <steve.the.bayesian@gmail.com>

### References

Gelman, Carlin, Stern, Rubin (2003), "Bayesian Data Analysis", Chapman and Hall.

---

beta.prior	<i>Beta prior for a binomial proportion</i>
------------	---

---

**Description**

Specifies beta prior distribution for a binomial probability parameter.

**Usage**

```
BetaPrior(a = 1, b = 1, mean = NULL, sample.size = NULL,
          initial.value = NULL)
```

**Arguments**

a	A positive real number interpretable as a prior success count.
b	A positive real number interpretable as a prior failure count.
mean	A positive real number representing $a/(a+b)$ .
sample.size	A positive real number representing $a+b$ .
initial.value	An initial value to be used for the variable being modeled. If NULL then the mean of the distribution will be used instead.

**Details**

The distribution should be specified either with a and b, or with mean and sample.size.

**Author(s)**

Steven L. Scott <steve.the.bayesian@gmail.com>

**References**

Gelman, Carlin, Stern, Rubin (2003), "Bayesian Data Analysis", Chapman and Hall.

---

boxplot.mcmc.matrix	<i>Plot the distribution of a matrix</i>
---------------------	--

---

**Description**

Plot the marginal distribution of each element in the Monte Carlo distribution of a matrix (e.g. a variance matrix or transition probability matrix). Rows and columns in the boxplots correspond to rows and columns in the matrix being plotted.

**Usage**

```
BoxplotMcmcMatrix(X, ylim = range(X), col.names,
                  row.names, truth, colors = NULL,
                  las = 0, ...)
```

**Arguments**

<code>X</code>	3 dimensional array. The first dimension is the Monte Carlo index (e.g. MCMC iteration). The second and third dimensions are the row and column of the matrix being plotted. E.g. $X[i, j, k]$ is Monte Carlo draw $i$ of matrix element $j, k$ .
<code>ylim</code>	2-vector giving the lower and upper limits of the vertical axis.
<code>col.names</code>	(optional) character vector giving the names of matrix columns (third dimension of $X$ ).
<code>row.names</code>	(optional) character vector giving the names of matrix rows (second dimension of $X$ ).
<code>truth</code>	(optional) scalar or matrix giving the values of reference lines to be plotted on each boxplot. If a scalar then the same value will be used for each boxplot. If a matrix then the rows and columns of the matrix correspond to the second and third dimension of $X$ .
<code>colors</code>	A vector of colors to use for the boxplots. Each row uses the same color scheme.
<code>las</code>	Controls the orientation of axis labels. See the <code>las</code> section in the help page for <a href="#">par</a> .
<code>...</code>	Extra arguments passed to <a href="#">boxplot</a>

**Value**

Called for its side effect, which is to draw a set of side-by-side boxplots on the current graphics device.

**Author(s)**

Steven L. Scott

**See Also**

[boxplot.true](#), [boxplot](#)

**Examples**

```
X <- array(rnorm(1000 * 3 * 4), dim=c(1000, 3, 4))
dimnames(X)[[2]] <- paste("row", 1:3)
dimnames(X)[[3]] <- paste("col", 1:4)
BoxplotMcmcMatrix(X)

truth <- 0
BoxplotMcmcMatrix(X, truth=truth)

truth <- matrix(rnorm(12), ncol=4)
BoxplotMcmcMatrix(X, truth=truth)
```

---

boxplot.true                      *Compare Boxplots to True Values*

---

### Description

Plots side-by-side boxplots of the columns of the matrix `x`. Each boxplot can have its own reference line (`truth`) and standard error lines `se.truth`, if desired. This function was originally written to display MCMC output, where the reference lines were true values used to test an MCMC simulation.

### Usage

```
BoxplotTrue(x, truth = NULL, vnames = NULL, center = FALSE,
            se.truth = NULL, color = "white", truth.color = "black",
            ylim = NULL, ...)
```

### Arguments

<code>x</code>	The matrix whose columns are to be plotted.
<code>truth</code>	(optional) A vector of reference values with length equal to <code>ncol(x)</code> .
<code>vnames</code>	(optional) character vector giving the column names of <code>x</code> .
<code>center</code>	(optional) logical. If <code>truth</code> is supplied then <code>center=TRUE</code> will center each column of <code>x</code> around <code>truth</code> to show the variation around the reference line.
<code>se.truth</code>	(optional) numeric vector of length <code>ncol(x)</code> . If <code>truth</code> is supplied then additional reference lines will be drawn at <code>truth +/- 2*se.truth</code> .
<code>color</code>	(optional) vector of colors for each boxplot.
<code>truth.color</code>	A color (or vector of colors) to use for the segments representing true values.
<code>ylim</code>	Limits for the vertical axis. If <code>NULL</code> then the axis will be scaled to fit <code>x</code> , <code>truth</code> , and <code>truth + c(2, -2) * se.truth</code>
<code>...</code>	additional arguments to <a href="#">boxplot</a> .

### Value

called for its side effect

### Author(s)

Steven L. Scott

### See Also

[boxplot.matrix](#), [boxplot](#),

### Examples

```
x <- t(matrix(rnorm(5000, 1:5, 1:5), nrow=5))
BoxplotTrue(x, truth=1:5, se.truth=1:5, col=rainbow(5), vnames =
  c("EJ", "TK", "JT", "OtherEJ", "TJ") )
```



---

check	<i>Check MCMC Output</i>
-------	--------------------------

---

### Description

Verify that MCMC output covers expected values.

### Usage

```
CheckMcmcMatrix(draws, truth, confidence = .95,
                 control.multiple.comparisons = TRUE,
                 burn = 0)
```

```
CheckMcmcVector(draws, truth, confidence = .95, burn = 0)
```

```
McmcMatrixReport(draws, truth, confidence = .95, burn = 0)
```

### Arguments

draws	The array of MCMC draws to check. This must be a matrix for <code>CheckMcmcMatrix</code> and a vector for <code>CheckMcmcVector</code> .
truth	The vector of true values that must be covered by draws in order for the check to succeed.
confidence	Specifies the probability width of the intervals used to determine whether draws covers truth. Central intervals are used, not HPD intervals.
control.multiple.comparisons	If <code>FALSE</code> then every interval must cover its corresponding true value. Otherwise a fraction of intervals (given by confidence) must cover.
burn	The number of MCMC iterations to discard as burn-in.

### Details

`CheckMcmcVector` checks a vector of draws corresponding to a scalar random variable. `CheckMcmcMatrix` checks a matrix of draws corresponding to a vector of random variables. In either case the check is made by constructing a central confidence interval (obtained by removing half of  $1 - \text{confidence}$  from the upper and lower tails of the distribution).

If a single variable is being checked with `CheckMcmcVector` then the check passes if and only if the interval covers the true value.

If multiple values are being checked with `CheckMcmcMatrix` then the user has control over how strict to make the check. If `control.multiple.comparisons` is `FALSE` then the check passes if and only if all intervals cover true values. Otherwise a fraction of intervals must cover. The fraction is the lower bound of the binomial confidence interval for the coverage rate under the hypothesis that the true coverage rate is confidence.

**Value**

CheckMcmcVector and CheckMcmcMatrix return TRUE if the check passes, and FALSE if it does not.

McmcMatrixReport returns a string that can be put in the info field of an `expect_true` expression, to give useful information about a failed test case. The return value is a textual representation of a three column matrix. Each row matches a variable in draws, and gives the lower and upper bounds for the credible interval used to check the values. The final column lists the true values that are supposed to be inside the credible intervals. The value is returned as a character string that is expected to be fed to `cat()` or `print()` so that it will render correctly in R CMD CHECK output.

**Author(s)**

Steven L. Scott

**Examples**

```
ndraws <- 100
draws <- rnorm(ndraws, 0, 1)
CheckMcmcVector(draws, 0) ## Returns TRUE
CheckMcmcVector(draws, 17) ## Returns FALSE

draws <- matrix(nrow = ndraws, ncol = 5)
for (i in 1:5) {
  draws[, i] <- rnorm(ndraws, i, 1)
}
CheckMcmcMatrix(draws, truth = 1:5) ## Returns TRUE
CheckMcmcMatrix(draws, truth = 5:1) ## Returns FALSE
```

---

check.data

*Checking data formats*

---

**Description**

Checks that data matches a concept

**Usage**

```
check.scalar.probability(x)
check.positive.scalar(x)
check.nonnegative.scalar(x)
check.probability.distribution(x)
check.scalar.integer(x)
check.scalar.boolean(x)
```

**Arguments**

x                    An object to be checked.

**Details**

If the object does not match the concept being checked, [stop](#) is called. Otherwise TRUE is returned.

**Author(s)**

Steven L. Scott <steve.the.bayesian@gmail.com>

---

circles

*Draw Circles*

---

**Description**

Draw circles on the current graphics device.

**Usage**

```
circles(center, radius, ...)
```

**Arguments**

center	A two-column matrix giving the coordinates of the circle center. If a single circle is to be drawn then a 2-element vector can be passed instead.
radius	The radii of the circles. A scalar value will be repeated if center is a matrix with more than one row.
...	Extra arguments passed to 'segments'. See <a href="#">par</a> for options controlling line type, line width, color, etc.

**Details**

Draws circles on the current graphics device. This is a low-level plotting function similar to [points](#), [lines](#), [segments](#), etc.

**Value**

Returns invisible NULL.

**Author(s)**

Steven L. Scott <steve.the.bayesian@gmail.com>

**Examples**

```
plot(1:10, type = "n")
circles(cbind(c(2, 3, 4), c(4, 5, 6)), radius = c(.3, .4, .5))
```

---

 compare.den

---

*Compare several density estimates.*


---

### Description

Produces multiple density plots on a single axis, to compare the columns of a matrix or the elements of a list.

### Usage

```
CompareDensities(x,
                 legend.text = NULL,
                 legend.location = "topright",
                 legend.title = NULL,
                 xlim = NULL,
                 ylim = NULL,
                 xlab = "parameter",
                 ylab = "density",
                 main = "",
                 lty = NULL,
                 col = "black",
                 axes = TRUE,
                 na.rm = TRUE,
                 ...)
```

### Arguments

x	matrix or list of numeric vectors. A density plot is produced for each column of the matrix or element of the list.
legend.text	(optional) character vector giving names of each density plot.
legend.location	Entry that can be passed to <a href="#">legend</a> .
legend.title	The legend title.
xlim	(optional) horizontal range of the plotting region. If omitted the region will be sized to fit all the observations in x.
ylim	(optional) vertical range of the plotting region. If omitted the region will be sized to fit all empirical density plots.
xlab	label to be placed on the horizontal axis
ylab	label to be placed on the vertical axis
main	main title for the plot
lty	The line types to use for the different densities. See <a href="#">par</a> . If NULL then a different line type will be used for each density.
col	vector of colors for the densities to be plotted.
axes	Logical. Should axes and a box be drawn around the figure?

na.rm            Logical value indicating whether NA's should be removed.  
...              Other graphical parameters passed to [plot.density](#), and [lines](#).

**Value**

Called for its side effect, which is to produce multiple density plots on the current graphics device.

**Author(s)**

Steven L. Scott

**See Also**

[density](#)

**Examples**

```
x <- t(matrix(rnorm(5000, 1:5, 1:5), nrow=5))
CompareDensities(x, legend.text=c("EJ", "TK", "JT", "OtherEJ", "TJ"),
                 col=rainbow(5), lwd=2)
```

---

compare.dynamic.distributions

*Compare Dynamic Distributions*

---

**Description**

Produce a plot showing several stacked dynamic distributions over the same horizontal axis.

**Usage**

```
CompareDynamicDistributions(
  list.of.curves,
  timestamps,
  style = c("dynamic", "boxplot"),
  xlab = "Time",
  ylab = "",
  frame.labels = rep("", length(list.of.curves)),
  main = "",
  actuals = NULL,
  col.actuals = "blue",
  pch.actuals = 1,
  cex.actuals = 1,
  vertical.cuts = NULL,
  ...)
```

**Arguments**

<code>list.of.curves</code>	A list of matrices, all having the same number of columns. Each matrix represents a distribution of curves, with rows corresponding to individual curves, and columns to time points.
<code>timestamps</code>	A vector of time stamps, with length matching the number of columns in each element of <code>list.of.curves</code> .
<code>style</code>	Should the curves be represented using a dynamic distribution plot, or boxplots. Boxplots are better for small numbers of time points. Dynamic distribution plots are better for large numbers of time points.
<code>xlab</code>	Label for the horizontal axis.
<code>ylab</code>	Label for the (outer) vertical axis.
<code>frame.labels</code>	Labels for the vertical axis of each subplot. The length must match the number of plot.
<code>main</code>	Main title for the plot.
<code>actuals</code>	If non-NULL, actuals should be a numeric vector giving the actual "true" value at each time point.
<code>col.actuals</code>	Color to use for the actuals. See <a href="#">par</a> .
<code>pch.actuals</code>	Plotting character(s) to use for the actuals. See <a href="#">par</a> .
<code>cex.actuals</code>	Scale factor for actuals. See <a href="#">par</a> .
<code>vertical.cuts</code>	If non-NULL then this must be a vector of the same type as <code>timestamps</code> with length matching the number of plots. A vertical line will be drawn at this location for each plot. Entries with the value NA signal that no vertical line should be drawn for that entry.
<code>...</code>	Extra arguments passed to <a href="#">PlotDynamicDistribution</a> or <a href="#">TimeSeriesBoxplot</a> .

**Author(s)**

Steven L. Scott

---

`compare.many.densities`

*Compare several density estimates.*

---

**Description**

Produce a plot that compares the kernel density estimates for each element in a series of Monte Carlo draws of a vector or matrix.

**Usage**

```
CompareManyDensities(list.of.arrays,
                     style = c("density", "box"),
                     main = "",
                     color = NULL,
                     gap = 0,
                     burn = 0,
                     suppress.labels = FALSE,
                     x.same.scale = TRUE,
                     y.same.scale = FALSE,
                     xlim = NULL,
                     ylim = NULL,
                     legend.location = c("top", "right"),
                     legend.cex = 1,
                     reflines = NULL,
                     ...)
```

**Arguments**

<code>list.of.arrays</code>	A list of arrays representing the MCMC draws of the vector or matrix in question. Each list element represents a different group. The first index in each list element represents the Monte Carlo draw number (or iteration). The remaining indices represent the variables to be plotted. If the first list element has variable names assigned to its indices, these will be used to label the plots.
<code>style</code>	The style of plot to use for comparing distributions.
<code>main</code>	The main title of the plot.
<code>color</code>	A vector of colors to be used for representing the groups.
<code>gap</code>	The gap (in lines) between plots.
<code>burn</code>	The number of MCMC iterations to be discarded as burn-in.
<code>suppress.labels</code>	Logical. If FALSE then the dimnames (if any) of the first element in <code>list.of.arrays</code> will be used to annotate the plot. If TRUE then no labels will be used.
<code>x.same.scale</code>	Logical indicating whether the same horizontal scale should be used for all the plots.
<code>y.same.scale</code>	Logical indicating whether the same vertical scale should be used for all the plots. This argument is ignored if <code>style == "box"</code> .
<code>xlim</code>	Either NULL, or a pair of numbers giving limits for the horizontal axis. If <code>xlim</code> is set then the same <code>xlim</code> values will be used for all plots and the <code>x.same.scale</code> argument will be ignored.
<code>ylim</code>	Either NULL, or a pair of numbers giving limits for the vertical axis. If <code>ylim</code> is set then the same <code>ylim</code> values will be used for all plots and the <code>y.same.scale</code> argument will be ignored. This argument is ignored if <code>style == "box"</code> .
<code>legend.location</code>	The location of the legend, either on top or at the right. It can also be NULL in which case no legend will appear. The legend names will be taken from <code>names(list.of.arrays)</code> . If it does not have names, then no legend will be produced.

legend.cex	The relative scale factor to use for the legend text.
reflines	This can be NULL, in which case no reference lines are drawn, it can be a single real number in which case a reference line will be drawn at that value in each panel, or it can be a vector with length equal to the number of panels, in which case a reference line will be drawn at each panel-specific value.
...	Extra arguments passed to <a href="#">CompareDen</a> .

**Author(s)**

Steven L. Scott

**See Also**[density](#), [CompareManyTs](#)**Examples**

```
x <- array(rnorm(9000), dim = c(1000, 3, 3))
dimnames(x) <- list(NULL, c("Larry", "Moe", "Curly"), c("Larry", "Eric", "Sergey"))
y <- array(rnorm(9000), dim = c(1000, 3, 3))
z <- array(rnorm(9000), dim = c(1000, 3, 3))
data <- list(x = x, y = y, z = z)
CompareManyDensities(data, color = c("red", "blue", "green"))
CompareManyDensities(data, style = "box")

x <- matrix(rnorm(5000), nrow = 1000)
colnames(x) <- c("Larry", "Moe", "Curly", "Shemp", "???)")
y <- matrix(rnorm(5000), nrow = 1000)
z <- matrix(rnorm(5000), nrow = 1000)
data <- list(x = x, y = y, z = z)
CompareManyDensities(data, color = c("red", "blue", "green"))
CompareManyDensities(data, style = "box")
```

---

`compare.many.ts`*Compares several density estimates.*

---

**Description**

Produce a plot that compares the kernel density estimates for each element in a series of Monte Carlo draws of a vector or matrix.

**Usage**

```
CompareManyTs(list.of.ts, burn = 0, type = "l", gap = 0,
              boxes = TRUE, thin = 1, labels = NULL,
              same.scale = TRUE, ylim = NULL, reline = NULL,
              color = NULL, ...)
```



**Arguments**

<code>list.of.ts</code>	A list of time series matrices, data.frames or 3-dimensional arrays, all of the same size. The list elements correspond to groups. The first index of the array in each list element corresponds to time. The subsequent indices correspond to variables to be plotted.
<code>burn</code>	The number of initial observations to be discarded as burn-in (when plotting MCMC output).
<code>type</code>	The plotting type to use when plotting the time series. See <a href="#">plot</a> .
<code>gap</code>	The amount of space to put between plots.
<code>boxes</code>	Logical. Should boxes be drawn around the plots?
<code>thin</code>	Plot every thin'th observation. This can reduce the amount of time it takes to make the plot if there are many long time series.
<code>labels</code>	A character vector to use as labels for individual plots.
<code>same.scale</code>	Logical. If TRUE then all plots are shown on the same vertical scale, and vertical axes are drawn. If FALSE then each plot gets its own scale.
<code>ylim</code>	The scale of the vertical axis. If non-NULL then same.scale will be set to TRUE.
<code>refline</code>	The scalar value at which a thin dotted horizontal line should be plotted in each panel. This is useful for highlighting zero, for example.
<code>color</code>	A vector of colors to use for the plots.
<code>...</code>	Extra arguments passed to 'plot' and 'axis'.

**Author(s)**

Steven L. Scott

**See Also**

[PlotManyTs](#), [CompareManyDensities](#)

**Examples**

```
x <- array(rnorm(9000), dim = c(1000, 3, 3))
dimnames(x) <- list(NULL, c("Larry", "Moe", "Curly"), c("Larry", "Eric", "Sergey"))
y <- array(rnorm(9000), dim = c(1000, 3, 3))
z <- array(rnorm(9000), dim = c(1000, 3, 3))
data <- list(x = x, y = y, z = z)
CompareManyTs(data, color = c("red", "blue", "green"))

x <- matrix(rnorm(5000), nrow = 1000)
colnames(x) <- c("Larry", "Moe", "Curly", "Shemp", "???" )
y <- matrix(rnorm(5000), nrow = 1000)
z <- matrix(rnorm(5000), nrow = 1000)
data <- list(x = x, y = y, z = z)
CompareManyTs(data, color = c("red", "blue", "green"))
```

---

compare.vector.distribution

*Boxplots to compare distributions of vectors*

---

### Description

Uses boxplots to compare distributions of vectors.

### Usage

```
CompareVectorBoxplots(draws, main = NULL, colors = NULL, burn = 0, ...)
```

### Arguments

draws	A list of MCMC draws. Each list element is a matrix with rows corresponding to MCMC iterations and columns to variables. The matrices can have different numbers of rows, but should have the same numbers of columns.
main	Main title of the plot.
colors	Colors to use for the boxplots. The length must match the number entries in draws.
burn	The number of initial MCMC iterations to discard before making the plot.
...	Extra arguments passed to <a href="#">boxplot</a> .

### Details

Creates side-by-side boxplots with the dimensions of each vector grouped together.

### Examples

```
x <- matrix(rnorm(300, mean = 1:3, sd = .4), ncol = 3, byrow = TRUE)
y <- matrix(rnorm(600, mean = 3:1, sd = .2), ncol = 3, byrow = TRUE)
CompareVectorBoxplots(list(x = x, y = y), colors = c("red", "blue"))
```

**Description**

A 'DiffDoubleModel' is tag given to a probability distribution that measures a real-valued scalar outcome, and whose log density is twice differentiable with respect to the random variable. The tag is a signal to underlying C++ code that the object being passed is one of a subset of understood distributions. Presently that subset includes the following distributions.

- [SdPrior](#)
- [NormalPrior](#)
- [BetaPrior](#)
- [UniformPrior](#)
- [GammaPrior](#)
- [TruncatedGammaPrior](#)
- [LognormalPrior](#)

Clearly this list is non-exhaustive, and other distributions may be added in the future.

**Author(s)**

Steven L. Scott <steve.the.bayesian@gmail.com>

---

dirichlet-distribution

*The Dirichlet Distribution*

---

**Description**

Density and random generation for the Dirichlet distribution.

**Usage**

```
ddirichlet(probabilities, nu, logscale = FALSE)
rdirichlet(n, nu)
```

**Arguments**

probabilities	A vector representing a discrete probability distribution, or a matrix where each row is a discrete probability distribution. Zero probabilities are not allowed.
nu	The parameters of the Dirichlet distribution. This can be a vector of positive numbers, interpretable as prior counts, of length matching the dimension of probabilities. If probabilities is a matrix (or if $n > 1$ ) then nu can also be a matrix of the same dimension, in which case each row of nu is used to evaluate the corresponding row of probabilities.
logscale	Logical. If TRUE then return the density on the log scale. Otherwise return the density on the raw scale.
n	The number of desired draws.

**Details**

The Dirichlet distribution is a generalization of the beta distribution. Whereas beta distribution is a model for probabilities, the Dirichlet distribution is a model for discrete distributions with several possible outcome values.

Let  $\pi$  denote a discrete probability distribution (a vector of positive numbers summing to 1), and let  $\nu$  be a vector of positive numbers (the parameters of the Dirichlet distribution), which can be thought of as prior counts. Then the density of the Dirichlet distribution can be written

$$f(\pi) = \frac{\Gamma(\sum_i \nu_i)}{\prod_i \Gamma(\nu_i)} \prod_i \pi_i^{\nu_i-1}.$$

**Value**

`ddirichlet` returns a vector of density values, with one entry per row in probabilities. `rdirichlet` returns a matrix (if  $n > 1$ ) or a vector (if  $n=1$ ) containing the draws from the Dirichlet distribution with the specified parameters.

**Author(s)**

Steven L. Scott <steve.the.bayesian@gmail.com>

**References**

Gelman, Carlin, Stern, Rubin (2003), "Bayesian Data Analysis", Chapman and Hall.

---

`dirichlet.prior`      *Dirichlet prior for a multinomial distribution*

---

**Description**

Specifies Dirichlet prior for a discrete probability distribution.

**Usage**

```
DirichletPrior(prior.counts, initial.value = NULL)
```

**Arguments**

`prior.counts`      A vector of positive numbers representing prior counts.  
`initial.value`      The initial value in the MCMC algorithm of the distribution being modeled.

**Author(s)**

Steven L. Scott <steve.the.bayesian@gmail.com>

**References**

Gelman, Carlin, Stern, Rubin (2003), "Bayesian Data Analysis", Chapman and Hall.

---

`discrete-uniform-prior`*Discrete prior distributions*

---

**Description**

Prior distributions over a discrete quantities.

**Usage**

```
PointMassPrior(location)
PoissonPrior(mean, lower.limit = 0, upper.limit = Inf)
DiscreteUniformPrior(lower.limit, upper.limit)
```

**Arguments**

<code>location</code>	The location of the point mass.
<code>mean</code>	The mean of the Poisson distribution.
<code>lower.limit</code>	The smallest value within the support of the distribution. The prior probability for numbers less than <code>lower.limit</code> is zero.
<code>upper.limit</code>	The largest value within the support of the distribution. The prior probability for numbers greater than <code>upper.limit</code> is zero.

**Value**

Each function returns a prior object whose class is the same as the function name. All of these inherit from "DiscreteUniformPrior" and from "Prior".

The `PoissonPrior` assumes a potentially truncated Poisson distribution with the given mean.

**Author(s)**

Steven L. Scott <steve.the.bayesian@gmail.com>

**Examples**

```
## Specify an exact number of trees in a Bart model (see the BoomBart
## package).

ntrees <- PointMassPrior(200)

## Uniform prior between 50 and 100 trees, including the endpoints.
ntrees <- DiscreteUniformPrior(50, 100)

## Truncated Poisson prior, with a mean of 20, a lower endpoint of 1,
## and an upper endpoint of 50.
ntrees <- PoissonPrior(20, 1, 50)
```

---

dmvn *Multivariate Normal Density*

---

### Description

Evaluate the multivariate normal density.

### Usage

```
dmvn(y, mu, sigma, siginv = NULL, ldsi = NULL, logscale = FALSE)
```

### Arguments

y	A numeric vector or matrix containing the data whose density is desired.
mu	The mean of the distribution. A vector.
sigma	The variance matrix of the distribution. A matrix.
siginv	The inverse of sigma, or NULL. If siginv is non-NULL then sigma will not be used.
ldsi	The log determinant of siginv or NULL.
logscale	Logical. If TRUE then the density is returned on the log scale. Otherwise the density is returned on the density scale.

### Value

A vector containing the density of each row of y.

### Author(s)

Steven L. Scott <steve.the.bayesian@gmail.com>

---

double.model *Prior distributions for a real valued scalar*

---

### Description

A DoubleModel is a class of prior distributions for real valued scalar parameters. A DoubleModel is sometimes used by a probability model that does not have a conjugate prior.

### Author(s)

Steven L. Scott <steve.the.bayesian@gmail.com>

### See Also

[BetaPrior](#), [GammaPrior](#), [LognormalPrior](#), [NormalPrior](#), [SdPrior](#), [TruncatedGammaPrior](#), and [UniformPrior](#).

---

external.legend	<i>Add an external legend to an array of plots.</i>
-----------------	---

---

### Description

ExternalLegendLayout sets up a plotting region to plot a regular grid of points, with an optional legend on the top or right of the grid.

AddExternalLegend adds a legend to a grid of plots that was set up using ExternalLegendLayout.

### Usage

```
ExternalLegendLayout(nrow,
                    ncol,
                    legend.labels,
                    legend.location = c("top", "right"),
                    outer.margin.lines = rep(4, 4),
                    gap.between.plots = rep(0, 4),
                    legend.cex = 1,
                    x.axis = TRUE,
                    y.axis = TRUE)
```

```
AddExternalLegend(legend.labels,
                  legend.location = c("top", "right"),
                  legend.cex = 1,
                  bty = "n",
                  ...)
```

### Arguments

nrow	The number of rows in the array of plots.
ncol	The number of columns in the array of plots.
legend.labels	The labels to be used in the legend.
legend.location	Specifies whether the legend should appear on the top or the right hand side. It can also be NULL, indicating that no legend is desired.
outer.margin.lines	A vector of length four giving the number of lines of text desired for the outer margins of the plot. See the oma argument of <a href="#">par</a> . This can also be specified as a single number, to be repeated 4 times.
gap.between.plots	A vector of length 4 giving the number of lines of text to leave between grid panels. See the mar argument of <a href="#">par</a> . This can also be specified as a single number, to be repeated 4 times.
legend.cex	The scale factor that will be used for legend text. This must match the scale factor used in add.external.legend.

x.axis	Will any plots have a horizontal axis?
y.axis	Will any plots have a vertical axis?
bty	Type of box to draw around the legend. Can be "n" (for no box) or "o" for a box. See <a href="#">legend</a> .
...	Extra arguments passed to <a href="#">legend</a> .

**Value**

ExternalLegendLayout returns the original graphical parameters, intended for use with [on.exit](#).  
AddExternalLegend returns invisible NULL.

**Author(s)**

Steven L. Scott

**See Also**

[legend layout](#)

**Examples**

```
example.plot <- function() {
  x <- rnorm(100)
  y <- rnorm(100)
  scale <- range(x, y)
  opar <- ExternalLegendLayout(nrow = 2,
                               ncol = 2,
                               legend.labels = c("foo", "bar"))
  on.exit({par(opar); layout(1)})
  hist(x, xlim = scale, axes = FALSE, main = "")
  mtext("X", side = 3, line = 1)
  box()
  plot(x, y, xlim = scale, ylim = scale, axes = FALSE)
  box()
  axis(3)
  axis(4)
  plot(y, x, xlim = scale, ylim = scale, axes = FALSE, pch = 2, col = 2)
  box()
  axis(1)
  axis(2)
  hist(y, xlim = scale, axes = FALSE, main = "")
  mtext("Y", side = 1, line = 1)
  box()
  AddExternalLegend(legend.labels = c("foo", "bar"),
                    pch = 1:2,
                    col = 1:2,
                    legend.cex = 1.5)
}

## Now call example.plot().
example.plot()
```



```
## Because of the call to on.exit(), in example.plot,
## the original plot layout is restored.
hist(1:10)
```

---

gamma.prior

*Gamma prior distribution*


---

## Description

Specifies gamma prior distribution.

## Usage

```
GammaPrior(a = NULL, b = NULL, prior.mean = NULL, initial.value = NULL)
TruncatedGammaPrior(a = NULL, b = NULL, prior.mean = NULL,
  initial.value = NULL,
  lower.truncation.point = 0,
  upper.truncation.point = Inf)
```

## Arguments

a	The shape parameter in the Gamma(a, b) distribution.
b	The scale parameter in the Gamma(a, b) distribution.
prior.mean	The mean of the Gamma(a, b) distribution, which is a/b.
initial.value	The initial value in the MCMC algorithm of the variable being modeled.
lower.truncation.point	The lower limit of support for the truncated gamma distribution.
upper.truncation.point	The upper limit of support for the truncated gamma distribution.

## Details

The mean of the Gamma(a, b) distribution is a/b and the variance is a/b<sup>2</sup>. If prior.mean is not NULL, then one of either a or b must be non-NULL as well.

GammaPrior is the conjugate prior for a Poisson mean or an exponential rate. For a Poisson mean a corresponds to a prior sum of observations and b to a prior number of observations. For an exponential rate the roles are reversed a represents a number of observations and b the sum of the observed durations. The gamma distribution is a generally useful for parameters that must be positive.

The gamma distribution is the conjugate prior for the reciprocal of a Gaussian variance, but [SdPrior](#) should usually be used in that case.

A TruncatedGammaPrior is a GammaPrior with support truncated to the interval (lower.truncation.point, upper.truncation.point).

If an object specifically needs a GammaPrior you typically cannot pass a TruncatedGammaPrior.

**Author(s)**

Steven L. Scott <steve.the.bayesian@gmail.com>

**References**

Gelman, Carlin, Stern, Rubin (2003), "Bayesian Data Analysis", Chapman and Hall.

---

GenerateFactorData      *Generate a data frame of all factor data*

---

**Description**

This function is mainly intended for example code and unit testing. It generates a `data.frame` containing all factor data.

**Usage**

```
GenerateFactorData(factor.levels.list, sample.size)
```

**Arguments**

`factor.levels.list`

A list of character vectors giving factor level names. The names attribute of this list becomes the set of variables names for the return data frame.

`sample.size`

The desired number of rows in the returned data frame.

**Author(s)**

Steven L. Scott <steve.the.bayesian@gmail.com>

**Examples**

```
foo <- GenerateFactorData(list(a = c("foo", "bar", "baz"),
                              b = c("larry", "moe", "curly", "shemp")),
                          50)
```

```
head(foo)
#   a   b
# 1 bar curly
# 2 foo curly
# 3 bar  moe
# 4 bar  moe
# 5 baz curly
# 6 bar curly
```

---

`histabunch`*A Bunch of Histograms*

---

### Description

Plot a bunch of histograms describing the marginal distributions the columns in a data frame.

### Usage

```
histabunch(x, gap = 1, same.scale = FALSE, boxes = FALSE,  
           min.continuous = 12, max.factor = 40,  
           vertical.axes = FALSE, ...)
```

### Arguments

<code>x</code>	A matrix or data frame containing the variables to be plotted.
<code>gap</code>	The gap between the plots, measured in lines of text.
<code>same.scale</code>	Logical. Indicates whether the histograms should all be plotted on the same scale.
<code>boxes</code>	Logical. Indicates whether boxes should be drawn around the histograms.
<code>min.continuous</code>	Numeric variables with more than <code>min.continuous</code> unique values will be plotted as continuous. Otherwise they will be treated as factors.
<code>max.factor</code>	Factors with more than <code>max.factor</code> levels will be beautified (ha!) by combining their remaining levels into an "other" category.
<code>vertical.axes</code>	Logical value indicating whether the histograms should be given vertical "Y" axes.
<code>...</code>	Extra arguments passed to <code>hist</code> (for numeric variables) or <code>barplot</code> (for factors).

### Value

Called for its side effect, which is to produce multiple histograms on the current graphics device.

### Author(s)

Steven L. Scott

### See Also

[hist barplot](#)

### Examples

```
data(airquality)  
histabunch(airquality)
```

---

inverse-wishart	<i>Inverse Wishart Distribution</i>
-----------------	-------------------------------------

---

**Description**

Density for the inverse Wishart distribution.

**Usage**

```
dInverseWishart(Sigma, sum.of.squares, nu, logscale = FALSE,
                log.det.sumsq = log(det(sum.of.squares)))
```

```
InverseWishartPrior(variance.guess, variance.guess.weight)
```

**Arguments**

Sigma	Argument (random variable) for the inverse Wishart distribution. A positive definite matrix.
nu	The "degrees of freedom" parameter of the inverse Wishart distribution. The distribution is only defined for $\text{nu} \geq \text{nrow}(\text{Sigma}) - 1$ .
sum.of.squares	A positive definite matrix. Typically this is the sum of squares that is the sufficient statistic for the inverse Wishart distribution.
logscale	Logical. If TRUE then the density is returned on the log scale. Otherwise the density is returned on the density scale.
log.det.sumsq	The log determinant of sum.of.squares. If this function is to be called many times then precomputing the log determinant can save considerable compute time.
variance.guess	A prior guess at the value of the variance matrix the prior is modeling.
variance.guess.weight	A positive scalar indicating the number of observations worth of weight to place on variance.guess.

**Details**

The inverse Wishart distribution has density function

$$\frac{|\text{Sigma}|^{-\frac{\nu+p+1}{2}} \exp(-\text{tr}(\Sigma^{-1}S)/2)}{2^{\frac{\nu p}{2}} |\Sigma|^{\frac{\nu}{2}} \Gamma_p(\nu/2)}$$

**Value**

`dInverseWishart` returns the scalar density (or log density) at the specified value. This function is not vectorized, so only one random variable (matrix) can be evaluated at a time.

`InverseWishartPrior` returns a list that encodes the parameters of the distribution in a format expected by underlying C++ code.

**Author(s)**

Steven L. Scott <steve.the.bayesian@gmail.com>

**See Also**

[dWishart](#), [rWishart](#), [NormalInverseWishartPrior](#)

---

 invgamma

*Inverse Gamma Distribution*


---

**Description**

Density, distribution function, quantile function, and random draws from the inverse gamma distribution.

**Usage**

```
dinvgamma(x, shape, rate, logscale = FALSE)
pinvgamma(x, shape, rate, lower.tail = TRUE, logscale = FALSE)
qinvgamma(p, shape, rate, lower.tail = TRUE, logscale = FALSE)
rinvgamma(n, shape, rate)
```

**Arguments**

x	A vector of deviates where the density or distribution function is to be evaluated.
p	A vector of probabilities representing CDF values (if <code>lower.tail == TRUE</code> ) or survivor function values (if <code>lower.tail == FALSE</code> ) from the inverse gamma distribution.
n	The desired number of draws from the inverse gamma distribution.
shape	The shape parameter.
rate	The 'rate' parameter. NOTE: The term 'rate' is used to match the corresponding parameter in <a href="#">rgamma</a> . Much of the rest of the world calls this the 'scale' parameter.
logscale	Logical. If TRUE then probabilities or density values are interpreted on the log scale. Otherwise the scale is the probability or probability density scale.
lower.tail	Logical. If TRUE then cumulative probabilities are measured from zero, as in a CDF. If FALSE then cumulative are measured from infinity, as in a survivor function.

**Value**

- `rinvgamma` returns draws from the distribution.
- `dinvgamma` returns the density function.
- `pinvgamma` returns the cumulative distribution function (or survivor function, if `lower.tail == FALSE`).
- `qinvgamma` returns quantiles from the distribution. `qinvgamma` and `pinvgamma` are inverse functions.

**Author(s)**

Steven L. Scott <steve.the.bayesian@gmail.com>

---

is.even

*Check whether a number is even or odd.*

---

**Description**

Check whether a number is even or odd.

**Usage**

```
IsEven(x)  
IsOdd(x)
```

**Arguments**

x                    An integer or vector of integers.

**Value**

Logical indicating whether the argument is even (or odd).

**Author(s)**

Steven L. Scott

**Examples**

```
IsEven(2) ## TRUE  
IsOdd(2)  ## FALSE
```

---

lmgamma

*Log Multivariate Gamma Function*

---

**Description**

Returns the log of the multivariate gamma function.

**Usage**

```
lmgamma(y, dimension)
```

**Arguments**

y	The function argument, which must be a positive scalar.
dimension	The dimension of the multivariate gamma function, which must be an integer $\geq 1$ .

**Details**

The multivariate gamma function is

$$\Gamma_p(y) = \pi^{p(p-1)/4} \prod_{j=1}^p \Gamma(y + (1 - j)/2).$$

The multivariate gamma function shows up as part of the normalizing constant for the Wishart and inverse Wishart distributions.

**Value**

Returns the log of the multivariate gamma function. Note that this function is not vectorized. Both y and dimension must be scalars, and the return value is a scalar.

**Author(s)**

Steven L. Scott <steve.the.bayesian@gmail.com>

---

log.integrated.gaussian.likelihood

*Log Integrated Gaussian Likelihood*

---

**Description**

Compute the log of the integrated Gaussian likelihood, where the model parameters are integrated out with respect to a normal-inverse gamma prior.

**Usage**

```
LogIntegratedGaussianLikelihood(suf, prior)
```

**Arguments**

suf	A <a href="#">GaussianSuf</a> object describing the data.
prior	A <a href="#">NormalInverseGammaPrior</a> describing the prior distribution.

**Value**

Returns a scalar giving the log integrated likelihood.

**Author(s)**

Steven L. Scott <steve.the.bayesian@gmail.com>

**Examples**

```
prior <- NormalInverseGammaPrior(10, 2, sqrt(2), 1)
y <- c(7.8949, 9.17438, 8.38808, 5.52521)
suf <- GaussianSuf(y)
loglike <- LogIntegratedGaussianLikelihood(suf, prior)
# -9.73975
```

---

lognormal.prior      *Lognormal Prior Distribution*

---

**Description**

Specifies a lognormal prior distribution.

**Usage**

```
LognormalPrior(mu = 0.0, sigma = 1.0, initial.value = NULL)
```

**Arguments**

mu	mean of the corresponding normal distribution.
sigma	standard deviation of the corresponding normal distribution. <b>WARNING:</b> If something looks strange in your program, look out for SD != Variance errors.
initial.value	Initial value of the variable to be modeled (e.g. in an MCMC algorithm). If NULL then the prior mean will be used.

**Details**

A lognormal distribution, where  $\log(y) \sim N(\mu, \sigma)$ . The mean of this distribution is  $\exp(\mu + 0.5 * \sigma^2)$ , so don't only focus on the mean parameter here.

**Author(s)**

Steven L. Scott <steve.the.bayesian@gmail.com>

**References**

Gelman, Carlin, Stern, Rubin (2003), "Bayesian Data Analysis", Chapman and Hall.  
[https://en.wikipedia.org/wiki/Log-normal\\_distribution](https://en.wikipedia.org/wiki/Log-normal_distribution)



---

markov.prior	<i>Prior for a Markov chain</i>
--------------	---------------------------------

---

### Description

The conjugate prior distribution for the parameters of a homogeneous Markov chain. The rows in the transition probability matrix modeled with independent Dirichlet priors. The distribution of the initial state is modeled with its own independent Dirichlet prior.

### Usage

```
MarkovPrior(prior.transition.counts = NULL,  
            prior.initial.state.counts = NULL,  
            state.space.size = NULL,  
            uniform.prior.value = 1)
```

### Arguments

`prior.transition.counts`

A matrix of the same dimension as the transition probability matrix being modeled. Entry (i, j) represents the "prior count" of transitions from state i to state j.

`prior.initial.state.counts`

A vector of positive numbers representing prior counts of initial states.

`state.space.size`

If both `prior.transition.counts` and `prior.initial.state.counts` are missing, then they will be filled with an object of dimension `state.space.size` where all entries are set to `uniform.prior.value`.

`uniform.prior.value`

The default value to use for entries of `prior.transition.counts` and `prior.initial.state.counts`, when they are not supplied by the user.

### Author(s)

Steven L. Scott <steve.the.bayesian@gmail.com>

### References

Gelman, Carlin, Stern, Rubin (2003), "Bayesian Data Analysis", Chapman and Hall.

---

match_data_frame	<i>MatchDataFrame</i>
------------------	-----------------------

---

### Description

Given two data frames with the same data, but with rows and columns in potentially different orders, produce a pair of permutations such that `data2[row.permutation, column.permutation]` matches `data1`.

### Usage

```
MatchDataFrame(data.to.match, data.to.permute)
```

### Arguments

`data.to.match` The data frame to be matched.  
`data.to.permute`  
The data frame to be permuted.

### Value

Returns a list with two elements.

`column.permutation`

A vector of indices such that the columns of `data2[, column.permutation]` match the columns of `data1`. The matching is based on column names.

`row.permutation`

A vector of indices such that the rows of `data2[row.permutation, column.permutation]` match the rows of `data1`. The matching is done by converting rows to strings, and matching the strings.

### Author(s)

Steven L. Scott <steve.the.bayesian@gmail.com>

### Examples

```
x1 <- data.frame(larry = rnorm(10), moe = 1:10, curly = rpois(10, 2))
x2 <- x1[c(1:5, 10:6), c(3, 1, 2)]

m <- MatchDataFrame(x1, x2)
x2[m$row.permutation, m$column.permutation] == x1 ## all TRUE
```

---

mscan	<i>Scan a Matrix</i>
-------	----------------------

---

### Description

Quickly scan a matrix from a file.

### Usage

```
mscan(fname, nc = 0, header = FALSE, burn = 0, thin = 0, nlines = 0L,  
      sep = "", ...)
```

### Arguments

fname	The name of the file from which to scan the data.
nc	The number of columns in the matrix to be read. If zero then the number of columns will be determined by the number of columns in the first line of the file.
header	logical indicating whether the file contains a header row.
burn	An integer giving the number of initial lines of the matrix to discard.
thin	An integer. If thin > 1 then keep every thin`th line. This is useful for reading in very large files of MCMC output, for example.
nlines	If positive, the number of data lines to scan from the data file (e.g. for an MCMC algorithm that is only partway done). Otherwise the entire file will be read.
sep	Field separator in the data file.
...	Extra arguments passed to 'scan'.

### Details

This function is similar to [read.table](#), but scanning a matrix of homogeneous data is much faster because there is much less format deduction.

### Value

The matrix stored in the data file.

### Author(s)

Steven L. Scott <steve.the.bayesian@gmail.com>

**Examples**

```
filename <- file.path(tempdir(), "example.data")
cat("foo bar baz", "1 2 3", "4 5 6", file = filename, sep = "\n")
m <- mscan(filename, header = TRUE)
m
##      foo bar baz
## [1,] 1  2  3
## [2,] 4  5  6
```

---

mvn.diagonal.prior      *diagonal MVN prior*

---

**Description**

A multivariate normal prior distribution formed by the product of independent normal margins.

**Usage**

```
MvnDiagonalPrior(mean.vector, sd.vector)
```

**Arguments**

mean.vector      A vector giving the mean of the prior distribution.  
sd.vector      The standard deviations of the components in the distribution. I.e. the square root of the diagonal of the variance matrix.

**Author(s)**

Steven L. Scott <steve.the.bayesian@gmail.com>

**References**

Gelman, Carlin, Stern, Rubin (2003), "Bayesian Data Analysis", Chapman and Hall.

---

mvn.independent.sigma.prior  
*Independence prior for the MVN*

---

**Description**

A prior for the parameters of the multivariate normal distribution that assumes Sigma to be a diagonal matrix with elements modeled by independent inverse Gamma priors.

**Usage**

```
MvnIndependentSigmaPrior(mvn.prior, sd.prior.list)
```

**Arguments**

- mvn.prior      An object of class `MvnPrior` that is the prior distribution for the multivariate normal mean parameter.
- sd.prior.list      A list of `SdPrior` objects modeling the diagonal elements of the multivariate normal variance matrix. The off-diagonal elements are assumed to be zero.

**Author(s)**

Steven L. Scott <steve.the.bayesian@gmail.com>

**References**

Gelman, Carlin, Stern, Rubin (2003), "Bayesian Data Analysis", Chapman and Hall.

---

mvn.prior      *Multivariate normal prior*

---

**Description**

A multivariate normal prior distribution.

**Usage**

```
MvnPrior(mean, variance)
```

**Arguments**

- mean      A vector giving the mean of the prior distribution.
- variance      A symmetric positive definite matrix giving the variance of the prior distribution.

**Author(s)**

Steven L. Scott <steve.the.bayesian@gmail.com>

**References**

Gelman, Carlin, Stern, Rubin (2003), "Bayesian Data Analysis", Chapman and Hall.

---

MvnGivenSigmaMatrixPrior

*Conditional Multivariate Normal Prior Given Variance*


---

### Description

A multivariate normal prior distribution, typically used as the prior distribution for the mean of multivariate normal data. The variance of this distribution is proportional to another parameter "Sigma" that exists elsewhere. Usually "Sigma" is the variance of the data.

This distribution is the "normal" part of a "normal-inverse Wishart" distribution.

### Usage

```
MvnGivenSigmaMatrixPrior(mean, sample.size)
```

### Arguments

mean            A vector giving the mean of the prior distribution.  
sample.size    A positive scalar. The variance of the distribution is Sigma / sample.size.

### Author(s)

Steven L. Scott <steve.the.bayesian@gmail.com>

### References

Gelman, Carlin, Stern, Rubin (2003), "Bayesian Data Analysis", Chapman and Hall.

---

normal.inverse.gamma.prior

*Normal inverse gamma prior*


---

### Description

The NormalInverseGammaPrior is the conjugate prior for the mean and variance of the scalar normal distribution. The model says that

$$\frac{1}{\sigma^2} \sim \text{Gamma}(df/2, ss/2) \mid \mu \mid \sigma \sim N(\mu_0, \sigma^2/\kappa)$$

### Usage

```
NormalInverseGammaPrior(mu.guess, mu.guess.weight = .01,  
sigma.guess, sigma.guess.weight = 1, ...)
```

**Arguments**

mu.guess	The mean of the prior distribution. This is $\mu_0$ in the description above.
mu.guess.weight	The number of observations worth of weight assigned to mu.guess. This is $\kappa$ in the description above.
sigma.guess	A prior estimate at the value of sigma. This is $\sqrt{ss/df}$ .
sigma.guess.weight	The number of observations worth of weight assigned to sigma.guess. This is $df$ .
...	blah

**Author(s)**

Steven L. Scott <steve.the.bayesian@gmail.com>

**References**

Gelman, Carlin, Stern, Rubin (2003), "Bayesian Data Analysis", Chapman and Hall.

---

normal.inverse.wishart.prior  
*Normal inverse Wishart prior*

---

**Description**

The NormalInverseWishartPrior is the conjugate prior for the mean and variance of the multivariate normal distribution. The model says that

$$\Sigma^{-1} \sim Wishart(\nu, S) | \sigma \sim N(\mu_0, \Sigma/\kappa)$$

The  $Wishart(S, \nu)$  distribution is parameterized by S, the *inverse* of the sum of squares matrix, and the scalar degrees of freedom parameter nu.

The distribution is improper if  $\nu < dim(S)$ .

**Usage**

```
NormalInverseWishartPrior(mean.guess,
                           mean.guess.weight = .01,
                           variance.guess,
                           variance.guess.weight = nrow(variance.guess) + 1)
```

**Arguments**

mean.guess	The mean of the prior distribution. This is $\mu_0$ in the description above.
mean.guess.weight	The number of observations worth of weight assigned to mean.guess. This is $\kappa$ in the description above.
variance.guess	A prior estimate at the value of $\Sigma$ . This is $S^{-1}/\nu$ in the notation above.
variance.guess.weight	The number of observations worth of weight assigned to variance.guess. This is $df$ .

**Author(s)**

Steven L. Scott <steve.the.bayesian@gmail.com>

**References**

Gelman, Carlin, Stern, Rubin (2003), "Bayesian Data Analysis", Chapman and Hall.

---

normal.prior	<i>Normal (scalar Gaussian) prior distribution</i>
--------------	--

---

**Description**

Specifies a scalar Gaussian prior distribution.

**Usage**

```
NormalPrior(mu, sigma, initial.value = mu, fixed = FALSE)
```

**Arguments**

mu	The mean of the prior distribution.
sigma	The standard deviation of the prior distribution.
initial.value	The initial value of the parameter being modeled in the MCMC algorithm.
fixed	Should the deviate modeled by this distribution be fixed at its initial value? (Used for debugging in some code. Not universally respected.)

**Author(s)**

Steven L. Scott <steve.the.bayesian@gmail.com>

**References**

Gelman, Carlin, Stern, Rubin (2003), "Bayesian Data Analysis", Chapman and Hall.



---

pairs.density                      *Pairs plot for posterior distributions.*

---

### Description

A pairs plot showing the posterior distribution of the given list of Monte Carlo draws. Plots above the diagonal show the posterior distribution on a scale just wide enough to fit the plots. The diagonal shows a marginal density plot, and the subdiagonal shows the distribution with all plots on a common scale.

### Usage

```
PairsDensity(draws,
              nlevels = 20,
              lty = NULL,
              color = NULL,
              subset = NULL,
              labels,
              legend.location = "top",
              legend.cex = 1,
              label.cex = 1,
              ...)
```

### Arguments

draws	Either a matrix or a list of matrices. If a list is provided then each list element is plotted as a separate set of contours, and all matrices must have the same number of columns (though the number of rows can differ).
nlevels	The number of contour levels to plot.
lty	The line types to use for the different elements in draws.
color	The color to use for different elements in draws.
subset	If draws is a list, then this can be a numerical vector. If draws has names, then subset can be a character vector naming which elements to include. If NULL then all elements of draws are plotted.
labels	If labels is missing and the first element of draws has non-NULL colnames then these will be used to label the pairs plot. If a character vector of length <code>ncol(draws[[1]])</code> then this character vector will be used in place of the colnames. If NULL then no labels will be used.
legend.location	Either "top", or "right" specifying the location for the legend, or NULL, indicating that no legend is desired. if draws is a matrix or a singleton list then no legend is produced.
legend.cex	Scale factor to use for the legend labels.
label.cex	Scale factor to use for the row and column labels.
...	Extra arguments (graphical parameters), passed to <a href="#">plot</a> , <a href="#">PlotDensityContours</a> , <a href="#">axis</a> , and <a href="#">AddExternalLegend</a> .

**Author(s)**

Steven L. Scott

**See Also**[pairs](#), [CompareDensities](#), [CompareManyDensities](#)**Examples**

```
## You can see the pairs plot for a single set of draws.  
y <- matrix(rnorm(5000, mean = 1:5), ncol = 5, byrow = TRUE)  
PairsDensity(y)
```

```
## You can also compare two or more sets of draws.  
z <- matrix(rnorm(2500, mean = 2:6), ncol = 5, byrow = TRUE)  
PairsDensity(list("first set" = y, "second set" = z))
```

---

plot.density.contours *Contour plot of a bivariate density.*

---

**Description**

Contour plot of one or more bivariate densities. This function was originally created to implement `PairsDensity`, but might be useful on its own.

**Usage**

```
PlotDensityContours(draws,  
                    x.index = 1,  
                    y.index = 2,  
                    xlim = NULL,  
                    ylim = NULL,  
                    nlevels = 20,  
                    subset = NULL,  
                    color = NULL,  
                    lty = NULL,  
                    axes = TRUE,  
                    ...)
```

**Arguments**

draws	Either a matrix or a list of matrices. If a list is provided then each list element is plotted as a separate set of contours, and all matrices must have the same number of columns (though the number of rows can differ).
x.index	The index of the parameter to plot on the horizontal axis.

y.index	The index of the beta coefficient to plot on the vertical axis.
xlim	Limits on the horizontal axis. If NULL then the plot is just wide enough to fit the contours.
ylim	Limits on the vertical axis. If NULL then the plot is just tall enough to fit the contours.
nlevels	The number of contour levels to plot.
subset	If draws is a list, then this can be a numerical vector. If draws has names, then subset can be a character vector naming which elements to include. If NULL then all elements of draws are plotted.
color	The color to use for different elements in draws.
lty	The line types to use for the different elements in draws.
axes	Logical. Should x and y axes be drawn?
...	Extra arguments passed to <a href="#">contour</a> .

**Author(s)**

Steven L. Scott

**See Also**[contour](#), [kde2d](#)**Examples**

```
## You can see the pairs plot for a single set of draws.
y <- matrix(rnorm(5000, mean = 1:5), ncol = 5, byrow = TRUE)
PlotDensityContours(y, 3, 1)

## You can also compare two or more sets of draws.
z <- matrix(rnorm(2500, mean = 2:6), ncol = 5, byrow = TRUE)
PlotDensityContours(list("first set" = y, "second set" = z), 3, 1)
```

---

plot.dynamic.distribution

*Plots the pointwise evolution of a distribution over an index set.*


---

**Description**

Produces an dynamic distribution plot where gray scale shading is used to show the evolution of a distribution over an index set. This function is particularly useful when the index set is too large to do side-by-side boxplots.

**Usage**

```
PlotDynamicDistribution(curves,
                       timestamps = NULL,
                       quantile.step=.01,
                       xlim = NULL,
                       xlab = "Time",
                       ylim = range(curves, na.rm = TRUE),
                       ylab = "distribution",
                       add = FALSE,
                       axes = TRUE,
                       ...)
```

**Arguments**

curves	A matrix where each row represents a curve (e.g. a simulation of a time series from a posterior distribution) and columns represent different points in the index set. For example, a long time series would be a wide matrix.
timestamps	An optional vector of "time stamps" that curves will be plotted against. The length of timestamps must match the number of columns in curves. If timestamps is NULL then the function attempts to extract time stamps from the colnames(curves). If no appropriate time stamps can be found then the positive integers will be used as time stamps.
quantile.step	Each color step in the plot corresponds to this difference in quantiles. Smaller values make prettier plots, but the plots take longer to produce.
xlim	The x limits (x1, x2) of the plot. Note that x1 > x2 is allowed and leads to a "reversed axis".
xlab	Label for the horizontal axis.
ylim	The y limits (y1, y2) of the plot. Note that y1 > y2 is allowed and leads to a "reversed axis".
ylab	Label for the vertical axis.
add	Logical. If true then add the plot to the current plot. Otherwise a fresh plot will be created.
axes	Logical. Should axes be added to the plot?
...	Extra arguments to pass on to <a href="#">plot</a>

**Details**

The function works by passing many calls to [polygon](#). Each polygon is associated with a quantile level, with darker shading near the median.

**Value**

This function is called for its side effect, which is to produce a plot on the current graphics device.

**Author(s)**

Steven L. Scott <steve.the.bayesian@gmail.com>

**Examples**

```
x <- t(matrix(rnorm(1000 * 100, 1:100, 1:100), nrow=100))
## x has 1000 rows, and 100 columns. Column i is N(i, i^2) noise.

PlotDynamicDistribution(x)
time <- as.Date("2010-01-01", format = "%Y-%m-%d") + (0:99 - 50)*7
PlotDynamicDistribution(x, time)
```

plot.macf

*Plots individual autocorrelation functions for many-valued time series***Description**

Produces individual autocorrelation functions for many-valued time series such as those produced by highly multivariate MCMC output. Cross-correlations such as those produced by [acf](#) are not shown.

**Usage**

```
PlotMacf(x, lag.max = 40, gap = 0.5, main = NULL, boxes = TRUE,
         xlab = "lag", ylab = "ACF", type = "h")
```

**Arguments**

x	matrix or 3-way array of MCMC output (or other time series). The first dimension represents discrete time.
lag.max	maximum lag to use when computing ACF's.
gap	non-negative scalar. gap between plots
main	character. main title for the plot
boxes	logical. Should boxes be drawn around the plots
xlab	character label for horizontal axis.
ylab	character label for vertical axis.
type	type of line plot to show. Defaults to "h". See <a href="#">plot.default</a> for other options.

**Value**

Called for its side effect

**Author(s)**

Steven L. Scott

**See Also**

[acf](#), [plot.many.ts](#).

**Examples**

```
x <- matrix(rnorm(1000), ncol=10)
PlotMacf(x)
```

---

plot.many.ts

*Multiple time series plots*


---

**Description**

Plots many time series plots on the same graphical device. Each plot gets its own frame. Scales can be adjusted to see variation in each plot (each plot gets its own scale), or variation between plots (common scale).

**Usage**

```
PlotManyTs(x, type = "l", gap = 0, boxes = TRUE, truth = NULL,
           thin = 1, labs, same.scale = TRUE, ylim = NULL,
           refline = NULL, color = NULL, ...)
```

**Arguments**

x	Matrix, data frame, list, or 3-dimensional array to be plotted.
type	type of line plots to produce. See <a href="#">plot.default</a> for other options.
gap	Number of lines of space to put between plots.
boxes	Logical indicating whether boxes should be drawn around each plot.
truth	A vector or matrix of reference values to be added to each plot as a horizontal line. The dimension should match <code>dim(x)[-1]</code>
thin	Frequency of observations to plot. E.g. <code>thin=10</code> means plot every 10 <sup>th</sup> observation. Thinning can speed things up when plotting large amounts MCMC output.
labs	Optional character vector giving the title (e.g. variable name) for each plot. If <code>labs</code> is missing then column names or <code>dimnames</code> will be used to label the plots. If <code>labs</code> is <code>NULL</code> then no labels will be used.
same.scale	Logical indicating whether plots should be drawn with a common vertical axis, which is displayed on alternating rows of the plot. If <code>FALSE</code> then the vertical axis of each plot is scaled to the range of data in that plot, but no tick marks are displayed.
ylim	Scale of the vertical axis. If non- <code>NULL</code> then <code>same.scale</code> is set to <code>TRUE</code> and the supplied scale is used for all plots.
refline	a vector or scalar value to use as a reference line. This is a supplement to the <code>truth</code> argument. It can be useful when comparing true values (used in a simulation), estimated values (e.g. point estimates of parameters) and MCMC output.
color	Vector of colors to use in the plots.
...	Extra arguments passed to <a href="#">plot</a> and <a href="#">axis</a> .

**Author(s)**

Steven L. Scott

**See Also**[plot.ts](#) (for plotting a small number of time series) [plot.macf](#)**Examples**

```
x <- matrix(rnorm(1000), ncol = 10)
PlotManyTs(x)
PlotManyTs(x, same = FALSE)
```

---

 regression.coefficient.conjugate.prior

*Regression Coefficient Conjugate Prior*


---

**Description**

A conjugate prior for regression coefficients, conditional on residual variance, and sample size.

**Usage**

```
RegressionCoefficientConjugatePrior(
  mean,
  sample.size,
  additional.prior.precision = numeric(0),
  diagonal.weight = 0)
```

**Arguments**

<code>mean</code>	The mean of the prior distribution, denoted 'b' below. See Details.
<code>sample.size</code>	The value denoted $\kappa$ below. This can be interpreted as a number of observations worth of weight to be assigned to mean in the posterior distribution.
<code>additional.prior.precision</code>	A vector of non-negative numbers representing the diagonal matrix $\Lambda^{-1}$ below. Positive values for <code>additional.prior.precision</code> will ensure the distribution is proper even if the regression model has no data. If all columns of the design matrix have positive variance then <code>additional.prior.precision</code> can safely be set to zero. A zero-length numeric vector is a slightly more efficient equivalent to a vector of all zeros.
<code>diagonal.weight</code>	The weight given to the diagonal when $X^T X$ is averaged with its diagonal. The purpose of <code>diagonal.weight</code> is to keep the prior distribution proper even if $X$ is less than full rank. If the design matrix is full rank then <code>diagonal.weight</code> can be set to zero.

**Details**

A conditional prior for the coefficients (beta) in a linear regression model. The prior is conditional on the residual variance  $\sigma^2$ , the sample size n, and the design matrix X. The prior is

$$\beta|\sigma^2, X \sim N(b, \sigma^2(\Lambda^{-1} + V$$

where

$$V^{-1} = \frac{\kappa}{n}((1-w)X^T X + w \text{diag}(X^T X)).$$

The prior distribution also depends on the cross product matrix  $X^T X$  and the sample size n, which are not arguments to this function. It is expected that the underlying C++ code will get those quantities elsewhere (presumably from the regression modeled by this prior).

**Author(s)**

Steven L. Scott <steve.the.bayesian@gmail.com>

**References**

Gelman, Carlin, Stern, Rubin (2003), "Bayesian Data Analysis", Chapman and Hall.

---

replist

*Repeated Lists of Objects*

---

**Description**

Produces repeated copies of an object.

**Usage**

```
ReplList(object, times)
```

**Arguments**

object	The object to be replicated.
times	The desired number of replications.

**Value**

Returns a list containing times copies of object.

**Author(s)**

Steven L. Scott

**Examples**

```
alist <- list(x = "foo", y = 12, z = c(1:3))
three.copies <- ReplList(alist, 3)
```



---

rmvn	<i>Multivariate Normal Simulation</i>
------	---------------------------------------

---

**Description**

Simulate draws from the multivariate normal distribution.

**Usage**

```
rmvn(n = 1, mu, sigma = diag(rep(1., length(mu))))
```

**Arguments**

n	The desired number of draws.
mu	The mean of the distribution. A vector.
sigma	The variance matrix of the distribution. A matrix.

**Details**

Note that `mu` and `sigma` are the same for all `n` draws. This function cannot handle separate parameters for each draw the way `rnorm` and similar functions for scalar random variables can.

**Value**

If `n == 1` the return value is a vector. Otherwise it is a matrix with `n` rows and `length(mu)` columns.

**Author(s)**

Steven L. Scott <steve.the.bayesian@gmail.com>

**Examples**

```
y1 <- rnorm(1, 1:3)
## y1 is a vector
y2 <- rnorm(10, 1:3)
## y2 is a matrix
```

rvectorfunction      *RVectorFunction*

---

**Description**

A wrapper for passing R functions to C++ code.

**Usage**

```
RVectorFunction(f, ...)
```

**Arguments**

f	A scalar-valued function of a vector-valued argument. The function can depend on other arguments as long as the vector valued argument appears in the first position.
...	Optional, named, extra arguments to be passed to f. These arguments are fixed at the time this object is created. For the purpose of evaluating f, these arguments do not update.

**Details**

The Boom library can handle the output of this function as a C++ function object. Note that evaluating R functions in C is no faster than evaluating them in R, but a wrapper like this is useful for interacting with C and C++ libraries that expect to operate on C and C++ functions.

**Value**

A list containing the information needed to evaluate the function f in C++ code.

**Author(s)**

Steven L. Scott <steve.the.bayesian@gmail.com>

---

scaled.matrix.normal.prior  
*Scaled Matrix-Normal Prior*

---

**Description**

A matrix-normal prior distribution, intended as the conjugate prior for the regression coefficients in a multivariate linear regression.

**Usage**

```
ScaledMatrixNormalPrior(mean, nu)
```

**Arguments**

mean	A matrix giving the mean of the distributions
nu	A scale factor affecting the variance.

**Details**

The matrix normal distribution is a 3-parameter distribution  $MN(\mu, \Omega, V)$ , where  $\mu$  is the mean. A deviate from the distribution is a matrix  $B$ , where  $\text{Cov}(B[i, j], B[k, m]) = \Omega[i, k] * \text{Sigma}[j, m]$ . If  $b = \text{Vec}(B)$  is the vector obtained by stacking columns of  $B$ , then  $b$  is multivariate normal with mean  $\text{Vec}(\mu)$  and covariance matrix

$$\Sigma \otimes \Omega$$

(the kronecker product).

This prior distribution assumes the underlying C++ code knows where to find the predictor ( $X$ ) matrix in the regression, and the residual variance matrix  $\text{Sigma}$ . The assumed prior distribution is  $B \sim MN(\mu, X'X / \nu, \text{Sigma})$ .

Like most other priors in Boom, this function merely encodes information expected by the underlying C++ code, ensuring correct names and formatting.

**Author(s)**

Steven L. Scott <steve.the.bayesian@gmail.com>

---

sd.prior

*Prior for a standard deviation or variance*

---

**Description**

Specifies an inverse Gamma prior for a variance parameter, but inputs are defined in terms of a standard deviation.

**Usage**

```
SdPrior(sigma.guess, sample.size = .01, initial.value = sigma.guess,
        fixed = FALSE, upper.limit = Inf)
```

**Arguments**

sigma.guess	A prior guess at the value of the standard deviation.
sample.size	The weight given to sigma.guess. Interpretable as a prior observation count.
initial.value	The initial value of the parameter in the MCMC algorithm.
fixed	Logical. Some algorithms allow you to fix sigma at a particular value. If TRUE then sigma will remain fixed at initial.value, if supported.
upper.limit	If positive, this is the upper limit on possible values of the standard deviation parameter. Otherwise the upper limit is assumed infinite. Not supported by all MCMC algorithms.

**Author(s)**

Steven L. Scott <steve.the.bayesian@gmail.com>

**References**

Gelman, Carlin, Stern, Rubin (2003), "Bayesian Data Analysis", Chapman and Hall.

---

sufstat.Rd

*Sufficient Statistics*

---

**Description**

Sufficient statistics for various models.

**Usage**

```
RegressionSuf(X = NULL,
              y = NULL,
              xtx = crossprod(X),
              xty = crossprod(X, y),
              yty = sum(y^2),
              n = length(y),
              xbar = colMeans(X),
              ybar = mean(y))
```

```
GaussianSuf(y)
```

**Arguments**

x	The predictor matrix for a regression problem.
y	The data, or the regression response variable.
xtx	The cross product of the design matrix. "X transpose X."
xty	The cross product of the design matrix with the response vector. "X transpose y."
yty	The sum of the squares of the response vector. "y transpose y."
n	The sample size.
xbar	A vector giving the average of each column in the predictor matrix.
ybar	The (scalar) mean of the response variable y.

**Value**

The returned value is a function containing the sufficient statistics for a regression model. Arguments are checked to ensure they have legal values. List names match the names expected by underlying C++ code.

**Author(s)**

Steven L. Scott <steve.the.bayesian@gmail.com>

**Examples**

```
X <- cbind(1, matrix(rnorm(3 * 100), ncol = 3))
y <- rnorm(100)

## Sufficient statistics can be computed from raw data, if it is
## available.
suf1 <- RegressionSuf(X, y)

## The individual components can also be computed elsewhere, and
## provided as arguments. If n is very large, this can be a
## substantial computational savings.
suf2 <- RegressionSuf(xtx = crossprod(X),
                     xty = crossprod(X, y),
                     yty = sum(y^2),
                     n = 100,
                     xbar = colMeans(X))
```

---

suggest.burn.log.likelihood

*Suggest MCMC Burn-in from Log Likelihood*

---

**Description**

Suggests a burn-in period for an MCMC chain based on the log likelihood values simulated in the final few iterations of the chain.

**Usage**

```
SuggestBurnLogLikelihood(log.likelihood, fraction = .10, quantile = .9)
```

**Arguments**

log.likelihood	A numeric vector giving the log likelihood values for each MCMC iteration.
fraction	The fraction of the chain that should be used to determine the log likelihood lower bound. The default setting looks in the final 25% of the MCMC run. Must be an number less than 1. If fraction <= 0 then a 0 burn-in is returned.
quantile	The quantile of the values in the final fraction that must be exceeded before the burn-in period is declared over.

**Details**

Looks at the last fraction of the log.likelihood sequence and finds a specified quantile to use as a threshold. Returns the first iteration where log.likelihood exceeds this threshold.

**Value**

Returns a suggested number of iterations to discard. This can be 0 if `fraction == 0`, which is viewed as a signal that no burn-in is desired.

**Author(s)**

Steven L. Scott <steve.the.bayesian@gmail.com>

---

thin

*Thin the rows of a matrix*

---

**Description**

Systematic sampling of every `thin`'th row of a matrix or vector. Useful for culling MCMC output or denoising a plot.

**Usage**

```
thin(x, thin)
```

**Arguments**

<code>x</code>	The array to be thinned. The first dimension is the one sampled over.
<code>thin</code>	The frequency of observations to keep. With <code>thin=10</code> you will keep every 10th observation.

**Value**

The thinned vector, matrix, or array is returned.

**Author(s)**

Steven L. Scott

**Examples**

```
x <- rnorm(100)
thin(x, 10)
# returns a 10 vector

y <- matrix(rnorm(200), ncol=2)
thin(y, 10)
# returns a 10 by 2 matrix
```

---

thin.matrix	<i>Thin a Matrix</i>
-------------	----------------------

---

**Description**

Return discard all but every k'th row of a matrix.

**Usage**

```
ThinMatrix(mat, thin)
```

**Arguments**

mat	The matrix to be thinned.
thin	The distance between kept lines from mat. The larger the number the fewer lines are kept.

**Details**

The bigger the value of thin the more thinning that gets done. For example, thin = 10 will keep every 10 lines from mat.

**Value**

The matrix mat, after discarding all but every thin lines.

**Author(s)**

Steven L. Scott <steve.the.bayesian@gmail.com>

**Examples**

```
m <- matrix(1:100, ncol = 2)
ThinMatrix(m, thin = 10)
##      [,1] [,2]
## [1,]  10  60
## [2,]  20  70
## [3,]  30  80
## [4,]  40  90
## [5,]  50 100
```

---

TimeSeriesBoxplot      *Time Series Boxplots*

---

## Description

Creates a series of boxplots showing the evolution of a distribution over time.

## Usage

```
TimeSeriesBoxplot(x, time, ylim = NULL, add = FALSE, ...)
```

## Arguments

x	A matrix where each row represents a curve (e.g. a simulation of a time series from a posterior distribution) and columns represent time. A long time series would be a wide matrix.
time	A vector of class <code>Date</code> with length matching the number of columns in x.
ylim	limits for the y axis.
add	logical, if TRUE then add boxplots to current plot.
...	Extra arguments to pass on to <code>boxplot</code>

## Value

Called for its side effect, which is to produce a plot on the current graphics device.

## Author(s)

Steven L. Scott <steve.the.bayesian@gmail.com>

## Examples

```
x <- t(matrix(rnorm(1000 * 100, 1:100, 1:100), nrow=100))
## x has 1000 rows, and 100 columns. Column i is N(i, i^2) noise.
time <- as.Date("2010-01-01", format = "%Y-%m-%d") + (0:99 - 50)*7
TimeSeriesBoxplot(x, time)
```



---

ToString	<i>Convert to Character String</i>
----------	------------------------------------

---

**Description**

Convert an object to a character string, suitable for including in error messages.

**Usage**

```
ToString(object, ...)
```

**Arguments**

object	An object to be printed to a string.
...	Extra arguments passed to <code>print</code> .

**Value**

A string (a character vector of length 1) containing the printed value of the object.

**Author(s)**

Steven L. Scott <steve.the.bayesian@gmail.com>

**Examples**

```
m <- matrix(1:6, ncol = 2)
printed.matrix <- ToString(m)

y <- c(1, 2, 3, 3, 3, 3, 3, 3)
tab <- table(y)
printed.table <- ToString(tab)
```

---

traceproduct	<i>Trace of the Product of Two Matrices</i>
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**Description**

Returns the trace of the product of two matrices.

**Usage**

```
TraceProduct(A, B, b.is.symmetric = FALSE)
```

**Arguments**

- A                    The first matrix in the product.
- B                    The second matrix in the product.
- b.is.symmetric   Logical. A TRUE value indicates that B is a symmetric matrix. A slight computational savings is possible if B is symmetric.

**Value**

Returns a number equivalent to `sum(diag(A %% B))`.

**Author(s)**

Steven L. Scott <steve.the.bayesian@gmail.com>

---

uniform.prior                    *Uniform prior distribution*

---

**Description**

Specifies a uniform prior distribution on a real-valued scalar parameter.

**Usage**

```
UniformPrior(lo = 0, hi = 1, initial.value = NULL)
```

**Arguments**

- lo                    The lower limit of support.
- hi                    The upper limit of support.
- initial.value        The initial value of the parameter in question to use in the MCMC algorithm. If NULL then the mean  $(lo + hi)/2$  is used.

**Author(s)**

Steven L. Scott <steve.the.bayesian@gmail.com>

**References**

Gelman, Carlin, Stern, Rubin (2003), "Bayesian Data Analysis", Chapman and Hall.

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wishart	<i>Wishart Distribution</i>
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---

### Description

Density and random generation for the Wishart distribution.

### Usage

```
dWishart(W, Sigma, nu, logscale = FALSE)
rWishart(nu, scale.matrix, inverse = FALSE)
```

### Arguments

W	Argument (random variable) for the Wishart density. A symmetric positive definite matrix.
Sigma	Scale or "variance" parameter of the Wishart distribution. See the "details" section below.
nu	The "degrees of freedom" parameter of the Wishart distribution. The distribution is only defined for $\text{nu} \geq \text{nrow}(W) - 1$ .
logscale	Logical. If TRUE then the density is returned on the log scale. Otherwise the density is returned on the density scale.
scale.matrix	For the Wishart distribution the <code>scale.matrix</code> parameter means the same thing as the <code>Sigma</code> parameter in <code>dWishart</code> . It is the variance parameter of the generating multivariate normal distribution. If simulating from the inverse Wishart, <code>scale.matrix</code> is the INVERSE of the "sum of squares" matrix portion of the multivariate normal sufficient statistics.
inverse	Logical. If TRUE then simulate from the inverse Wishart distribution. If FALSE then simulate from the Wishart distribution.

### Details

If  $\text{nu}$  is an integer then a  $W(\Sigma, \nu)$  random variable can be thought of as the sum of  $\text{nu}$  outer products:  $y_i y_i^T$ , where  $y_i$  is a zero-mean multivariate normal with variance matrix `Sigma`.

The Wishart distribution is

$$\frac{|W|^{\frac{\nu-p-1}{2}} \exp(-\text{tr}(\Sigma^{-1}W)/2)}{2^{\frac{\nu p}{2}} |\Sigma|^{\frac{\nu}{2}} \Gamma_p(\nu/2)}$$

where  $p == \text{nrow}(W)$  and  $\Gamma_p(\nu)$  is the multivariate gamma function (see [lmgamma](#)).

**Value**

`dWishart` returns the density of the Wishart distribution. It is not vectorized, so only one random variable (matrix) can be evaluated at a time.

`rWishart` returns one or more draws from the Wishart or inverse Wishart distributions. If  $n > 0$  the result is a 3-way array. Unlike the `rWishart` function from the `stats` package, the first index corresponds to draws. This is in keeping with the convention of other models from the `Boom` package.

**Author(s)**

Steven L. Scott <steve.the.bayesian@gmail.com>

# Index

- \* **aplot**
  - add.segments, 4
- \* **dplot**
  - thin, 54
- \* **hplot**
  - boxplot.mcmc.matrix, 6
  - boxplot.true, 8
  - compare.den, 12
  - compare.dynamic.distributions, 13
  - compare.many.densities, 14
  - compare.many.ts, 16
  - external.legend, 23
  - histabunch, 27
  - pairs.density, 41
  - plot.density.contours, 42
  - plot.dynamic.distribution, 43
  - plot.macf, 45
  - plot.many.ts, 46
  - TimeSeriesBoxplot, 56
- \* **package**
  - Boom-package, 3
- acf, 45
- add.segments, 4
- AddExternalLegend, 41
- AddExternalLegend (external.legend), 23
- AddSegments (add.segments), 4
- ar1.coefficient.prior, 5
- Ar1CoefficientPrior
  - (ar1.coefficient.prior), 5
- axis, 41, 46
- barplot, 27
- beta.prior, 6
- BetaPrior, 3, 19, 22
- BetaPrior (beta.prior), 6
- Boom (Boom-package), 3
- Boom-package, 3
- boxplot, 7, 8, 18, 56
- boxplot.matrix, 8
- boxplot.mcmc.matrix, 6
- boxplot.true, 4, 7, 8
- BoxplotMcmcMatrix
  - (boxplot.mcmc.matrix), 6
- BoxplotTrue, 3
- BoxplotTrue (boxplot.true), 8
- check, 9
- check.data, 10
- check.nonnegative.scalar (check.data), 10
- check.positive.scalar (check.data), 10
- check.probability.distribution, 3
- check.probability.distribution (check.data), 10
- check.scalar.boolean (check.data), 10
- check.scalar.integer (check.data), 10
- check.scalar.probability (check.data), 10
- CheckMcmcMatrix, 3
- CheckMcmcMatrix (check), 9
- CheckMcmcVector, 3
- CheckMcmcVector (check), 9
- circles, 11
- compare.den, 12
- compare.densities (compare.den), 12
- compare.dynamic.distributions, 13
- compare.many.densities, 14
- compare.many.ts, 16
- compare.vector.boxplots (compare.vector.distribution), 18
- compare.vector.distribution, 18
- CompareDen, 16
- CompareDen (compare.den), 12
- CompareDensities, 42
- CompareDensities (compare.den), 12
- CompareDynamicDistributions (compare.dynamic.distributions), 13

- CompareManyDensities, [17, 42](#)
- CompareManyDensities
  - (compare.many.densities), [14](#)
- CompareManyTs, [16](#)
- CompareManyTs (compare.many.ts), [16](#)
- CompareVectorBoxplots
  - (compare.vector.distribution), [18](#)
- contour, [43](#)
- Date, [56](#)
- ddirichlet (dirichlet-distribution), [19](#)
- density, [13, 16](#)
- diff.double.model, [18](#)
- DiffDoubleModel (diff.double.model), [18](#)
- dInverseWishart (inverse-wishart), [28](#)
- dinvgamma (invgamma), [29](#)
- dirichlet-distribution, [19](#)
- dirichlet.prior, [20](#)
- DirichletPrior (dirichlet.prior), [20](#)
- discrete-uniform-prior, [21](#)
- DiscreteUniformPrior
  - (discrete-uniform-prior), [21](#)
- dmvn, [22](#)
- double.model, [22](#)
- DoubleModel (double.model), [22](#)
- dWishart, [29](#)
- dWishart (wishart), [59](#)
- expect\_true, [10](#)
- external.legend, [23](#)
- ExternalLegendLayout (external.legend), [23](#)
- gamma.prior, [25](#)
- GammaPrior, [3, 19, 22](#)
- GammaPrior (gamma.prior), [25](#)
- GaussianSuf, [31](#)
- GaussianSuf (sufstat.Rd), [52](#)
- GenerateFactorData, [26](#)
- hist, [27](#)
- histabunch, [27](#)
- install.packages, [3](#)
- inverse-wishart, [28](#)
- InverseWishartPrior (inverse-wishart), [28](#)
- invgamma, [29](#)
- is.even, [30](#)
- is.odd (is.even), [30](#)
- IsEven (is.even), [30](#)
- IsOdd (is.even), [30](#)
- kde2d, [43](#)
- layout, [24](#)
- legend, [12, 24](#)
- lines, [11, 13](#)
- lmgamma, [30, 59](#)
- log.integrated.gaussian.likelihood, [31](#)
- LogIntegratedGaussianLikelihood
  - (log.integrated.gaussian.likelihood), [31](#)
- lognormal.prior, [32](#)
- LognormalPrior, [19, 22](#)
- LognormalPrior (lognormal.prior), [32](#)
- markov.prior, [33](#)
- MarkovPrior (markov.prior), [33](#)
- match\_data\_frame, [34](#)
- MatchDataFrame (match\_data\_frame), [34](#)
- McmcMatrixReport (check), [9](#)
- mscan, [35](#)
- mvn.diagonal.prior, [36](#)
- mvn.independent.sigma.prior, [36](#)
- mvn.prior, [37](#)
- MvnDiagonalPrior (mvn.diagonal.prior), [36](#)
- MvnGivenSigmaMatrixPrior, [38](#)
- MvnIndependentSigmaPrior
  - (mvn.independent.sigma.prior), [36](#)
- MvnPrior, [37](#)
- MvnPrior (mvn.prior), [37](#)
- normal.inverse.gamma.prior, [38](#)
- normal.inverse.wishart.prior, [39](#)
- normal.prior, [40](#)
- NormalInverseGammaPrior, [31](#)
- NormalInverseGammaPrior
  - (normal.inverse.gamma.prior), [38](#)
- NormalInverseWishartPrior, [29](#)
- NormalInverseWishartPrior
  - (normal.inverse.wishart.prior), [39](#)
- NormalPrior, [19, 22](#)

- NormalPrior (normal.prior), 40
- on.exit, 24
- pairs, 42
- pairs.density, 41
- PairsDensity (pairs.density), 41
- par, 7, 11, 12, 14, 23
- pinvgamma (invgamma), 29
- plot, 17, 41, 44, 46
- plot.default, 45, 46
- plot.density, 13
- plot.density.contours, 42
- plot.dynamic.distribution, 43
- plot.macf, 45, 47
- plot.many.ts, 45, 46
- plot.ts, 47
- PlotDensityContours, 41
- PlotDensityContours
  - (plot.density.contours), 42
- PlotDynamicDistribution, 3, 14
- PlotDynamicDistribution
  - (plot.dynamic.distribution), 43
- PlotMacf (plot.macf), 45
- PlotManyTs, 3, 17
- PlotManyTs (plot.many.ts), 46
- PointMassPrior
  - (discrete-uniform-prior), 21
- points, 11
- PoissonPrior (discrete-uniform-prior), 21
- polygon, 44
- print, 57
- qinvgamma (invgamma), 29
- rdirichlet (dirichlet-distribution), 19
- read.table, 35
- regression.coefficient.conjugate.prior, 47
- RegressionCoefficientConjugatePrior
  - (regression.coefficient.conjugate.prior), 47
- RegressionSuf (sufstat.Rd), 52
- Replist (replist), 48
- replist, 48
- rgamma, 29
- rinvgamma (invgamma), 29
- rmvn, 49
- rnorm, 49
- RVectorFunction (rvectorfunction), 50
- rvectorfunction, 50
- rWishart, 29, 60
- rWishart (wishart), 59
- scaled.matrix.normal.prior, 50
- ScaledMatrixNormalPrior
  - (scaled.matrix.normal.prior), 50
- sd.prior, 51
- SdPrior, 19, 22, 25, 37
- SdPrior (sd.prior), 51
- segments, 11
- stop, 11
- sufstat.Rd, 52
- suggest.burn.log.likelihood, 53
- SuggestBurnLogLikelihood
  - (suggest.burn.log.likelihood), 53
- thin, 54
- thin.matrix, 55
- ThinMatrix (thin.matrix), 55
- TimeSeriesBoxplot, 14, 56
- ToString, 57
- TraceProduct (traceproduct), 57
- traceproduct, 57
- TruncatedGammaPrior, 19, 22
- TruncatedGammaPrior (gamma.prior), 25
- uniform.prior, 58
- UniformPrior, 19, 22
- UniformPrior (uniform.prior), 58
- wishart, 59