

# Package ‘HelpersMG’

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**Type** Package

**Title** Tools for Environmental Analyses, Ecotoxicology and Various R Functions

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**Author** Marc Girondot

**Maintainer** Marc Girondot <[marc.girondot@gmail.com](mailto:marc.girondot@gmail.com)>

**Depends** R (>= 4.1), MASS, ggplot2, rlang, coda, Matrix

**Suggests** lme4, RNetCDF, ncdf4, maps, fields, shiny, ppcor, pbmcapply, pbapply, parallel, visNetwork, igraph, shinyWidgets, cranlogs

**Description** Contains miscellaneous functions useful for managing 'NetCDF' files (see <<https://en.wikipedia.org/wiki/NetCDF>>), get moon phase and time for sun rise and fall, tide level, analyse and reconstruct periodic time series of temperature with irregular sinusoidal pattern, show scales and wind rose in plot with change of color of text, Metropolis-Hastings algorithm for Bayesian MCMC analysis, plot graphs or boxplot with error bars, search files in disk by there names or their content, read the contents of all files from a folder at one time.

**License** GPL-2

**LazyLoad** yes

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**RoxygenNote** 7.2.0

**NeedsCompilation** no

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## R topics documented:

HelpersMG-package . . . . .	4
addS3Class . . . . .	7
as.mcmc.mcmcComposite . . . . .	8
as.parameters . . . . .	9

as.quantiles . . . . .	12
asc . . . . .	13
barplot_errbar . . . . .	14
cArrows . . . . .	15
ChangeCoordinate . . . . .	17
char . . . . .	18
compare . . . . .	19
compare_AIC . . . . .	20
compare_AICc . . . . .	21
compare_BIC . . . . .	23
contingencyTable.compare . . . . .	24
convert.tz . . . . .	27
cutter . . . . .	28
d . . . . .	36
dbeta_new . . . . .	37
dcutter . . . . .	38
dggamma . . . . .	40
DIx . . . . .	42
dnbinom_new . . . . .	43
dSnbnom . . . . .	45
duplicated_packages . . . . .	47
ellipse . . . . .	48
ExtractAIC.glm . . . . .	50
fitdistrquantiles . . . . .	52
flexit . . . . .	53
FormatCompareAIC . . . . .	55
format_ncdf . . . . .	56
iCutter . . . . .	58
IC_clean_data . . . . .	59
IC_correlation_simplify . . . . .	61
IC_threshold_matrix . . . . .	62
index.periodic . . . . .	65
ind_long_lat . . . . .	66
inside . . . . .	68
invlogit . . . . .	69
LD50 . . . . .	70
LD50_MHmcmc . . . . .	72
LD50_MHmcmc_p . . . . .	75
list.packages . . . . .	76
local.search . . . . .	77
logit . . . . .	78
logLik.compareAIC . . . . .	79
logLik.cutter . . . . .	80
logLik.LD50 . . . . .	81
merge.mcmcComposite . . . . .	82
MHalgoGen . . . . .	84
minmax.periodic . . . . .	88
modeled.hist . . . . .	90

modifyVector . . . . .	91
moon.info . . . . .	92
MovingWindow . . . . .	93
NagelkerkeScaledR2 . . . . .	94
newcompassRose . . . . .	95
newmap.scale . . . . .	96
openwd . . . . .	97
plot.cutter . . . . .	98
plot.IconoCorel . . . . .	102
plot.LD50 . . . . .	104
plot.mcmcComposite . . . . .	106
plot.PriorsmcmcComposite . . . . .	111
plot_add . . . . .	112
plot_errbar . . . . .	113
predict.LD50 . . . . .	115
print.cutter . . . . .	117
pSnbinom . . . . .	120
qSnbinom . . . . .	122
qvlmer . . . . .	123
r2norm . . . . .	124
RandomFromHessianOrMCMC . . . . .	125
rcutter . . . . .	128
read_folder . . . . .	130
RectangleRegression . . . . .	131
rmnorm . . . . .	132
RM_add . . . . .	134
RM_delete . . . . .	135
RM_duplicate . . . . .	136
RM_get . . . . .	138
RM_list . . . . .	139
rnbnom_new . . . . .	140
rSnbinom . . . . .	142
ScalePreviousPlot . . . . .	143
SEfromHessian . . . . .	144
series.compare . . . . .	146
setPriors . . . . .	148
show_name . . . . .	150
similar . . . . .	151
specify_decimal . . . . .	152
summary.mcmcComposite . . . . .	153
sun.info . . . . .	154
symbol.Female . . . . .	156
symbol.Male . . . . .	157
symmetricize . . . . .	158
tide.info . . . . .	159
tnirp . . . . .	162
universalmclapply . . . . .	163
wget . . . . .	165

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HelpersMG-package      *Tools for Environmental Analyses, Ecotoxicology and Various R Functions*

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**Description**

Contains miscellaneous functions useful for managing  
'NetCDF' files (see <http://en.wikipedia.org/wiki/NetCDF>),  
get tide levels on any point of the globe,  
get moon phase and time for sun rise and fall,  
analyse and reconstruct daily time series of temperature  
with irregular sinusoidal pattern,  
show scales and wind rose in plot with change of color of text,  
Metropolis-Hastings algorithm for Bayesian MCMC analysis,  
plot graphs or boxplot with error bars,  
search files in disk by their names or their content,  
read the contents of all files from a folder at one time,  
calculate IC50 for ecotoxicological studies,  
calculate the probability mass function of the sum of negative binomial  
distributions, calculate distribution of unobserved values in censored or truncated distributions.  
The latest version of this package can always be installed using:  
`install.packages("https://hebergement.universite-paris-saclay.fr/marcgirondot/CRAN/HelpersMG.tar.gz",  
repos=NULL, type="source")`



## Details

Helpers functions for several packages

Package:	HelpersMG
Type:	Package
Version:	5.5 build 1268
Date:	2022-07-14
License:	GPL (>= 2)
LazyLoad:	yes

## Author(s)

Marc Girondot <[marc.girondot@gmail.com](mailto:marc.girondot@gmail.com)>

## Examples

```
## Not run:  
library.HelpersMG
```

```

print('-----')
print('Examples for mcmcComposite objects')
print('-----')
require(coda)
x <- rnorm(30, 10, 2)
dnormx <- function(x, par) return(-sum(dnorm(x, mean=par['mean'], sd=par['sd'], log=TRUE)))
parameters_mcmc <- data.frame(Density=c('dnorm', 'dlnorm'),
Prior1=c(10, 0.5), Prior2=c(2, 0.5), SDProp=c(0.35, 0.2),
Min=c(-3, 0), Max=c(100, 10), Init=c(10, 2), stringsAsFactors = FALSE,
row.names=c('mean', 'sd'))
mcmc_run <- MHalgoGen(n.iter=100000, parameters=parameters_mcmc, data=x,
likelihood=dnormx, n.chains=1, n.adapt=100, thin=1, trace=1)
plot(mcmc_run, xlim=c(0, 20))
plot(mcmc_run, xlim=c(0, 10), parameters="sd")
mcmcforcoda <- as.mcmc(mcmc_run)
# Optimal rejection rate should be 0.234
rejectionRate(mcmcforcoda)
heidel.diag(mcmcforcoda)
raftery.diag(mcmcforcoda)
autocorr.diag(mcmcforcoda)
acf(mcmcforcoda[[1]][,"mean"], lag.max=20, bty="n", las=1)
acf(mcmcforcoda[[1]][,"sd"], lag.max=20, bty="n", las=1)
batchSE(mcmcforcoda, batchSize=100)
# The batch standard error procedure is usually thought to
# be not as accurate as the time series methods used in summary
summary(mcmcforcoda)$statistics[, "Time-series SE"]
summary(mcmc_run)
as.parameters(mcmc_run)
lastp <- as.parameters(mcmc_run, index="last")
parameters_mcmc[,"Init"] <- lastp
# The n.adapt set to 1 is used to not record the first set of parameters
# then it is not duplicated (as it is also the last one for
# the object mcmc_run)
mcmc_run2 <- MHalgoGen(n.iter=10000, parameters=parameters_mcmc, data=x,
likelihood=dnormx, n.chains=1, n.adapt=1, thin=1, trace=1)
mcmc_run3 <- merge(mcmc_run, mcmc_run2)
##### no adaptation, n.adapt must be 0
parameters_mcmc[,"Init"] <- c(mean(x), sd(x))
mcmc_run3 <- MHalgoGen(n.iter=10000, parameters=parameters_mcmc, data=x,
likelihood=dnormx, n.chains=1, n.adapt=0, thin=1, trace=1)
print('-----')
print('Examples for Daily patterns of temperature')
print('-----')
# Generate a timeserie of time
time.obs <- NULL
for (i in 0:9) time.obs <- c(time.obs, c(0, 6, 12, 18)+i*24)
# For these time, generate a timeseries of temperatures
temp.obs <- rep(NA, length(time.obs))
temp.obs[3+(0:9)*4] <- rnorm(10, 25, 3)
temp.obs[1+(0:9)*4] <- rnorm(10, 10, 3)
for (i in 1:(length(time.obs)-1))
if (is.na(temp.obs[i]))
temp.obs[i] <- mean(c(temp.obs[i-1], temp.obs[i+1]))

```

```

if (is.na(temp.obs[length(time.obs)]))
  temp.obs[length(time.obs)] <- temp.obs[length(time.obs)-1]/2
observed <- data.frame(time=time.obs, temperature=temp.obs)
# Search for the minimum and maximum values
r <- minmax.periodic(time.minmax.daily=c(Min=2, Max=15),
observed=observed, period=24)

# Estimate all the temperatures for these values
t <- temperature.periodic(minmax=r)

plot_errbar(x=t[, "time"], y=t[, "temperature"],
errbar.y=ifelse(is.na(t[, "sd"]), 0, 2*t[, "sd"]),
type="l", las=1, bty="n", errbar.y.polygon = TRUE,
xlab="hours", ylab="Temperatures", ylim=c(0, 35),
errbar.y.polygon.list = list(col="grey"))

plot_add(x=t[, "time"], y=t[, "temperature"], type="l")

# How many times this package has been download
library(cranlogs)
HelpersMG <- cran_downloads("HelpersMG", from = "2015-04-07",
                             to = Sys.Date() - 1)
sum(HelpersMG$count)
plot(HelpersMG$date, HelpersMG$count, type="l", bty="n")

## End(Not run)

```

addS3Class

*Add a S3 class to an object.***Description**

Add a S3 class as first class to an object.

**Usage**

```
addS3Class(x, class = NULL)
```

**Arguments**

- |       |                          |
|-------|--------------------------|
| x     | The object to add class. |
| class | The class to add.        |

**Details**

addS3Class add a S3 class to an object

**Value**

The same object with the new class as first class

**Author(s)**

Marc Girondot <marc.girondot@gmail.com>

**Examples**

```
print.CF <- function(x) {cat("print.CF ", x)}
result <- "Je suis donc je pense"
result <- addS3Class(result, class="CF")
class(result)
print(result)
result <- addS3Class(result, class=c("ECF", "OCF"))
class(result)
print(result)
```

**as.mcmc.mcmcComposite** *Extract mcmc object from a mcmcComposite object*

**Description**

Take a mcmcComposite object and create a mcmc.list object to be used with coda package.

**Usage**

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

**Arguments**

x	A mcmcComposite obtained as a result of MHalgoGen() function
...	Not used

**Details**

as.mcmc Extract mcmc object from the result of phenology\_MHmcmc to be used with coda package

**Value**

A mcmc.list object

**Author(s)**

Marc Girondot <marc.girondot@gmail.com>

**See Also**

Other mcmcComposite functions: [MHalgoGen\(\)](#), [as.parameters\(\)](#), [as.quantiles\(\)](#), [merge.mcmcComposite\(\)](#), [plot.PriorsmcmcComposite\(\)](#), [plot.mcmcComposite\(\)](#), [setPriors\(\)](#), [summary.mcmcComposite\(\)](#)

## Examples

```

## Not run:
library(HelpersMG)
require(coda)
x <- rnorm(30, 10, 2)
dnormx <- function(data, x) {
  data <- unlist(data)
  return(-sum(dnorm(data, mean=x['mean'], sd=x['sd'], log=TRUE)))
}
parameters_mcmc <- data.frame(Density=c('dnorm', 'dlnorm'),
Prior1=c(10, 0.5), Prior2=c(2, 0.5), SDProp=c(1, 1),
Min=c(-3, 0), Max=c(100, 10), Init=c(10, 2), stringsAsFactors = FALSE,
row.names=c('mean', 'sd'))
mcmc_run <- MHalgoGen(n.iter=1000, parameters=parameters_mcmc, data=x,
likelihood=dnormx, n.chains=1, n.adapt=100, thin=1, trace=1)
plot(mcmc_run, xlim=c(0, 20))
plot(mcmc_run, xlim=c(0, 10), parameters="sd")
mcmcforcoda <- as.mcmc(mcmc_run)
#' heidel.diag(mcmcforcoda)
raftery.diag(mcmcforcoda)
autocorr.diag(mcmcforcoda)
acf(mcmcforcoda[[1]][,"mean"], lag.max=20, bty="n", las=1)
acf(mcmcforcoda[[1]][,"sd"], lag.max=20, bty="n", las=1)
batchSE(mcmcforcoda, batchSize=100)
# The batch standard error procedure is usually thought to
# be not as accurate as the time series methods used in summary
summary(mcmcforcoda)$statistics[,"Time-series SE"]
summary(mcmc_run)
as.parameters(mcmc_run)
lastp <- as.parameters(mcmc_run, index="last")
parameters_mcmc[,"Init"] <- lastp
# The n.adapt set to 1 is used to not record the first set of parameters
# then it is not duplicated (as it is also the last one for
# the object mcmc_run)
mcmc_run2 <- MHalgoGen(n.iter=1000, parameters=parameters_mcmc, data=x,
likelihood=dnormx, n.chains=1, n.adapt=1, thin=1, trace=1)
mcmc_run3 <- merge(mcmc_run, mcmc_run2)
##### no adaptation, n.adapt must be 0
parameters_mcmc[,"Init"] <- c(mean(x), sd(x))
mcmc_run3 <- MHalgoGen(n.iter=1000, parameters=parameters_mcmc, data=x,
likelihood=dnormx, n.chains=1, n.adapt=0, thin=1, trace=1)

## End(Not run)

```

as.parameters

*Extract parameters from mcmcComposite object*

## Description

Take a mcmcComposite object and create a vector object with parameter value at specified iteration. If index="best", the function will return the parameters for the highest likelihood. It also indicates

at which iteration the maximum likelihood has been observed.  
 If `index="last"`, the function will return the parameters for the last likelihood.  
 If `index="median"`, the function will return the median value of the parameter.  
 if `index="quantile"`, the function will return the probs defined by quantiles parameter.  
 If `index="mode"`, the function will return the mode value of the parameter based on Asselin de Beauville (1978) method.  
`index` can also be a numeric value.  
 This function uses the complete iterations available except the adaptation part, even if `thin` parameter is not equal to 1.

## Usage

```
as.parameters(x, index = "best", chain = 1, probs = 0.025)
```

## Arguments

<code>x</code>	A mcmcComposite obtained as a result of <code>MHalgoGen()</code> function
<code>index</code>	At which iteration the parameters must be taken, see description
<code>chain</code>	The number of the chain in which to get parameters
<code>probs</code>	Quantiles to be returned, see description

## Value

A vector with parameters at maximum likelihood or index position

## Author(s)

Marc Girondot <[marc.girondot@gmail.com](mailto:marc.girondot@gmail.com)>

## References

Asselin de Beauville J.-P. (1978). Estimation non paramétrique de la densité et du mode, exemple de la distribution Gamma. *Revue de Statistique Appliquée*, 26(3):47-70.

## See Also

Other mcmcComposite functions: [MHalgoGen\(\)](#), [as.mcmc.mcmcComposite\(\)](#), [as.quantiles\(\)](#), [merge.mcmcComposite\(\)](#), [plot.PriorsmcmcComposite\(\)](#), [plot.mcmcComposite\(\)](#), [setPriors\(\)](#), [summary.mcmcComposite\(\)](#)

## Examples

```
## Not run:
library(HelpersMG)
require(coda)
x <- rnorm(30, 10, 2)
dnormx <- function(data, x) {
  data <- unlist(data)
  return(-sum(dnorm(data, mean=x['mean'], sd=x['sd'], log=TRUE)))
}
```

```

parameters_mcmc <- data.frame(Density=c('dnorm', 'dlnorm'),
                                Prior1=c(10, 0.5),
                                Prior2=c(2, 0.5),
                                SDProp=c(1, 1),
                                Min=c(-3, 0),
                                Max=c(100, 10),
                                Init=c(10, 2),
                                stringsAsFactors = FALSE,
                                row.names=c('mean', 'sd'))
mcmc_run <- MHalgoGen(n.iter=1000, parameters=parameters_mcmc, data=x,
                       likelihood=dnormx, n.chains=1, n.adapt=100,
                       thin=1, trace=1)
plot(mcmc_run, xlim=c(0, 20))
plot(mcmc_run, xlim=c(0, 10), parameters="sd")
mcmcforcoda <- as.mcmc(mcmc_run)
#' heidel.diag(mcmcforcoda)
raftery.diag(mcmcforcoda)
autocorr.diag(mcmcforcoda)
acf(mcmcforcoda[[1]][,"mean"], lag.max=20, bty="n", las=1)
acf(mcmcforcoda[[1]][,"sd"], lag.max=20, bty="n", las=1)
batchSE(mcmcforcoda, batchSize=100)

# The batch standard error procedure is usually thought to
# be not as accurate as the time series methods used in summary
summary(mcmcforcoda)$statistics[, "Time-series SE"]
summary(mcmc_run)
as.parameters(mcmc_run)
lastp <- as.parameters(mcmc_run, index="last")
parameters_mcmc[,"Init"] <- lastp

# The n.adapt set to 1 is used to not record the first set of parameters
# then it is not duplicated (as it is also the last one for
# the object mcmc_run)
mcmc_run2 <- MHalgoGen(n.iter=1000, parameters=parameters_mcmc, data=x,
                        likelihood=dnormx, n.chains=1, n.adapt=1,
                        thin=1, trace=1)
mcmc_run3 <- merge(mcmc_run, mcmc_run2)

##### no adaptation, n.adapt must be 0
parameters_mcmc[,"Init"] <- c(mean(x), sd(x))
mcmc_run3 <- MHalgoGen(n.iter=1000, parameters=parameters_mcmc, data=x,
                        likelihood=dnormx, n.chains=1, n.adapt=0,
                        thin=1, trace=1)
# With index being median, it returns the median value for each parameter
as.parameters(mcmc_run3, index="median")
as.parameters(mcmc_run3, index="mode")
as.parameters(mcmc_run3, index="best")
as.parameters(mcmc_run3, index="quantile", probs=0.025)
as.parameters(mcmc_run3, index="quantile", probs=0.975)
as.parameters(mcmc_run3, index="quantile", probs=c(0.025, 0.975))

## End(Not run)

```

**as.quantiles***Extract quantile distribution from mcmcComposite object***Description**

Extract quantile distribution from mcmcComposite object

**Usage**

```
as.quantiles(
  x,
  chain = 1,
  fun = function(...) return(as.numeric(list(...))),
  probs = c(0.025, 0.975),
  xlim = NULL,
  nameparxlim = NULL,
  namepar = NULL
)
```

**Arguments**

<code>x</code>	A mcmcComposite obtained as a result of <code>MHalgoGen()</code> function
<code>chain</code>	The number of the chain in which to get parameters
<code>fun</code>	The function to apply the parameters
<code>probs</code>	The probability to get quantiles
<code>xlim</code>	The values to apply in <code>fun</code>
<code>nameparxlim</code>	The name of the parameter for <code>xlim</code>
<code>namepar</code>	The name of parameters from mcmc object to be used in <code>fun</code>

**Value**

A data.frame with quantiles

**Author(s)**

Marc Girondot <[marc.girondot@gmail.com](mailto:marc.girondot@gmail.com)>

**See Also**

Other mcmcComposite functions: `MHalgoGen()`, `as.mcmc.mcmcComposite()`, `as.parameters()`, `merge.mcmcComposite()`, `plot.PriorsmcmcComposite()`, `plot.mcmcComposite()`, `setPriors()`, `summary.mcmcComposite()`

## Examples

```
## Not run:
library(HelperMG)
require(coda)
x <- rnorm(30, 10, 2)
dnormx <- function(data, x) {
  data <- unlist(data)
  return(-sum(dnorm(data, mean=x['mean'], sd=x['sd'], log=TRUE)))
}
parameters_mcmc <- data.frame(Density=c('dnorm', 'dlnorm'),
Prior1=c(10, 0.5), Prior2=c(2, 0.5), SDProp=c(1, 1),
Min=c(-3, 0), Max=c(100, 10), Init=c(10, 2), stringsAsFactors = FALSE,
row.names=c('mean', 'sd'))
mcmc_run <- MHalgoGen(n.iter=10000, parameters=parameters_mcmc, data=x,
likelihood=dnormx, n.chains=1, n.adapt=100, thin=1, trace=1)
k <- as.quantiles(x=mcmc_run, namepar="mean")
k <- as.quantiles(x=mcmc_run, namepar="mean",
                  xlim=c(1:5), nameparxlim="sd",
                  fun=function(...) return(sum(as.numeric(list(...)))))

## End(Not run)
```

asc

*Return the codes (in UTF-8) of a string*

## Description

Return the codes (in UTF-8) of a string.

## Usage

```
asc(x)
```

## Arguments

x	The string to be analyzed
---	---------------------------

## Details

asc returns the codes (in UTF-8) of a string

## Value

A vector with ITF-8 codes of a string

## Author(s)

Based on this blog: <http://datadebri.blogs.com/2011/03/ascii-code-table-in-r.html>

**See Also**

Other Characters: [char\(\)](#), [d\(\)](#), [tnirp\(\)](#)

**Examples**

```
asc("abcd")
asc("ABCD")
```

<b>barplot_errbar</b>	<i>Plot a barplot graph with error bar on y</i>
-----------------------	---

**Description**

To plot data, just use it as a normal barplot but add the errbar.y values or errbar.y.minus, errbar.y.plus if bars for y axis are asymmetric. Use y.plus and y\_MINUS to set absolut limits for error bars. Note that y.plus and y\_MINUS have priority over errbar.y, errbar.y\_MINUS and errbar.y\_PLUS.

**Usage**

```
barplot_errbar(
  ...,
  errbar.y = NULL,
  errbar.y.plus = NULL,
  errbar.y_MINUS = NULL,
  y.plus = NULL,
  y_MINUS = NULL,
  errbar.tick = 1/50,
  errbar.lwd = par("lwd"),
  errbar.lty = par("lty"),
  errbar.col = par("fg"),
  add = FALSE
)
```

**Arguments**

...	Parameters for barplot() such as main= or ylim=
errbar.y	The length of error bars for y. Recycled if necessary.
errbar.y.plus	The length of positive error bars for y. Recycled if necessary.
errbar.y_MINUS	The length of negative error bars for y. Recycled if necessary.
y.plus	The absolut position of the positive error bar for y. Recycled if necessary.
y_MINUS	The absolut position of the nagative error bar for y. Recycled if necessary.
errbar.tick	Size of small ticks at the end of error bars defined as a proportion of total width or height graph size.
errbar.lwd	Error bar line width, see par("lwd")
errbar.lty	Error bar line type, see par("lwd")
errbar.col	Error bar line color, see par("col")
add	If true, add the graph to the previous one.

**Details**

`barplot_errbar` plot a barplot with error bar on y

**Value**

A numeric vector (or matrix, when `beside = TRUE`), say `mp`, giving the coordinates of all the bar midpoints drawn, useful for adding to the graph.

If `beside` is true, use `colMeans(mp)` for the midpoints of each group of bars, see example.

**Author(s)**

Marc Girondot <[marc.girondot@gmail.com](mailto:marc.girondot@gmail.com)>

**See Also**

`plot_errorbar`

Other plot and barplot functions: `ScalePreviousPlot()`, `plot_add()`, `plot_errbar()`, `show_name()`

**Examples**

```
## Not run:
barplot_errbar(rnorm(10, 10, 3),
  xlab="axe x", ylab="axe y", bty="n",
  errbar.y.plus=rnorm(10, 1, 0.1), col=rainbow(10),
  names.arg=paste("Group",1:10), cex.names=0.6)
y <- rnorm(10, 10, 3)
barplot_errbar(y,
  xlab="axe x", ylab="axe y", bty="n",
  y.plus=y+2)

## End(Not run)
```

**cArrows**

*Draw curved lines with arrowhead*

**Description**

Draw a curved line with arrowhead.

**Usage**

```
cArrows(
  x1,
  y1,
  x2,
  y2,
  code = 2,
  size = 1,
```

```

width = 1.2/4/cin,
open = TRUE,
sh.adj = 0.1,
sh.lwd = 1,
sh.col = if (is.R()) par("fg") else 1,
sh.lty = 1,
h.col = sh.col,
h.col.bo = sh.col,
h.lwd = sh.lwd,
h.lty = sh.lty,
curved = FALSE,
beautiful.arrow = 2/3
)

```

### Arguments

x1	coordinates of points from which to draw.
y1	coordinates of points from which to draw.
x2	coordinates of points to which to draw.
y2	coordinates of points to which to draw.
code	integer code (1, 2, or 3), determining kind of arrows to be drawn.
size	size of the arrowhead.
width	width of the arrowhead.
open	shape of the arrowhead.
sh.adj	Shift the beginning of the line.
sh.lwd	width of the line.
sh.col	color of the line.
sh.lty	type of line.
h.col	color of the arrowhead.
h.col.bo	color of the arrowhead border.
h.lwd	width of the arrowhead.
h.lty	type of line for the arrowhead.
curved	0 is a straigth line, positive of negative value make the line curved.
beautiful.arrow	if open is false, make the arrowhead more beautiful.

### Details

cArrows draws curved lines with arrowhead

### Value

A list wit lab.x and lab.y being the position where to draw label

**Author(s)**

Modified from iGraph

**Examples**

```
plot(c(1, 10), c(1, 10), type="n", bty="n")
cArrows(x1=2, y1=2, x2=6, y2=6, curved=1)
cArrows(x1=2, y1=2, x2=6, y2=6, curved=0)
cArrows(x1=2, y1=2, x2=6, y2=6, curved=1, sh.adj=1)
cArrows(x1=2, y1=2, x2=6, y2=6, curved=-1, open=FALSE)
cArrows(x1=9, y1=2, x2=6, y2=6, curved=-1, open=FALSE, sh.col="red")
cArrows(x1=9, y1=9, x2=6, y2=6, curved=-1, open=FALSE, h.col="red")
cArrows(x1=2, y1=9, x2=6, y2=6, curved=1, open=FALSE, h.col="red", h.col.bo="red")
```

---

ChangeCoordinate      *Return a value in a changed coordinate*

---

**Description**

Return a value in a changed coordinate system.

**Usage**

```
ChangeCoordinate(
  x = stop("At least one value to convert must be provided"),
  initial = stop("Set of two values must be provided as references"),
  transformed = stop("Set of two transformed values must be provided")
)
```

**Arguments**

x	value to convert
initial	Set of two values in the original system
transformed	Set of the two values in the converted system

**Details**

ChangeCoordinate returns a value in a changed coordinate

**Value**

A value in the new system

**Author(s)**

Marc Girondot <marc.girondot@gmail.com>

## Examples

```
ChangeCoordinate(x=c(10, 20), initial=c(1, 100), transformed=c(0, 1))
```

**char**

*Return the characters defined by the codes*

## Description

Return a string with characters defined by the codes.

## Usage

```
char(n)
```

## Arguments

n	The code to be used to return a character
---	---

## Details

char returns the characters defined by the codes

## Value

A string with characters defined by the codes

## Author(s)

Based on this blog: <http://datadebrief.blogspot.com/2011/03/ascii-code-table-in-r.html>

## See Also

Other Characters: [asc\(\)](#), [d\(\)](#), [tnirp\(\)](#)

## Examples

```
char(65:75)
char(unlist(tapply(144:175, 144:175, function(x) {c(208, x)})))
```

---

compare

*Run a shiny application for basic functions of comparison*

---

## Description

Run a shiny application for basic functions of comparison.

## Usage

```
compare()
```

## Details

compare runs a shiny application for basic functions of comparison

## Value

Nothing

## Author(s)

Marc Girondot <marc.girondot@gmail.com>

## References

Girondot, M., Guillon, J.-M., 2018. The w-value: An alternative to t- and X<sup>2</sup> tests. Journal of Biostatistics & Biometrics 1, 1-3.

## See Also

Other w-value functions: [contingencyTable.compare\(\)](#), [series.compare\(\)](#)

## Examples

```
## Not run:  
library(HelpersMG)  
compare()  
  
## End(Not run)
```

---

<code>compare_AIC</code>	<i>Compares the AIC of several outputs</i>
--------------------------	--

---

## Description

This function is used to compare the AIC of several outputs obtained with the same data but with different set of parameters.

The parameters must be lists with \$aic or \$AIC or \$value and \$par elements or if AIC(element) is defined.

if \$value and \$par are present in the object, the AIC is calculated as  $2 * \text{factor.value} * \text{value} + 2 * \text{length}(\text{par})$ .

If \$value is -log(likelihood), then factor.value must be 1 and if \$value is log(likelihood), then factor.value must be -1.

If several objects are within the same list, their AIC are summed.

For example, `compare_AIC(g1=list(group), g2=list(separe1, separe2))` can be used to compare a single model onto two different sets of data against each set of data fitted with its own set of parameters.

Take a look at `ICtab` in package `bbmle` which is similar.

## Usage

```
compare_AIC(
  ...,
  factor.value = 1,
  silent = FALSE,
  FUN = function(x) specify_decimal(x, decimals = 2)
)
```

## Arguments

...	Successive results to be compared as lists.
<code>factor.value</code>	The \$value of the list object is multiplied by factor.value to calculate AIC.
<code>silent</code>	If TRUE, nothing is displayed.
<code>FUN</code>	Function used to show values

## Details

`compare_AIC` compares the AIC of several outputs obtained with the same data.

## Value

A list with DeltaAIC and Akaike weight for the models.

## Author(s)

Marc Girondot <[marc.girondot@gmail.com](mailto:marc.girondot@gmail.com)>

**See Also**

Other AIC: [ExtractAIC.glm\(\)](#), [FormatCompareAIC\(\)](#), [compare\\_AICc\(\)](#), [compare\\_BIC\(\)](#)

**Examples**

```
## Not run:
library("HelpersMG")
# Here two different models are fitted
x <- 1:30
y <- rnorm(30, 10, 2)+log(x)
plot(x, y)
d <- data.frame(x=x, y=y)
m1 <- lm(y ~ x, data=d)
m2 <- lm(y ~ log(x), data=d)
compare_AIC(linear=m1, log=m2)
# Here test if two datasets can be modeled with a single model
x2 <- 1:30
y2 <- rnorm(30, 15, 2)+log(x2)
plot(x, y, ylim=c(5, 25))
plot_add(x2, y2, col="red")
d2 <- data.frame(x=x2, y=y2)
m1_2 <- lm(y ~ x, data=d2)
x_grouped <- c(x, x2)
y_grouped <- c(y, y2)
d_grouped <- data.frame(x=x_grouped, y=y_grouped)
m1_grouped <- lm(y ~ x, data=d_grouped)
compare_AIC(separate=list(m1, m1_2), grouped=m1_grouped)

## End(Not run)
```

**compare\_AICc**

*Compares the AICc of several outputs*

**Description**

This function is used to compare the AICc of several outputs obtained with the same data but with different set of parameters.

Each object must have associated `logLik()` method with `df` and `nobs` attributes.

AICc for object `x` will be calculated as `2*factor.value*logLik(x)+(2*attributes(logLik(x))$df*(attributes(logL`

**Usage**

```
compare_AICc(
  ...,
  factor.value = -1,
  silent = FALSE,
  FUN = function(x) specify_decimal(x, decimals = 2)
)
```

## Arguments

...	Successive results to be compared as lists.
factor.value	The \$value of the list object is multiplied by factor.value to calculate BIC.
silent	If TRUE, nothing is displayed.
FUN	Function used to show values

## Details

compare\_AICc compares the AICc of several outputs obtained with the same data.

## Value

A list with DeltaAICc and Akaike weight for the models.

## Author(s)

Marc Girondot <marc.girondot@gmail.com>

## See Also

Other AIC: [ExtractAIC.glm\(\)](#), [FormatCompareAIC\(\)](#), [compare\\_AIC\(\)](#), [compare\\_BIC\(\)](#)

## Examples

```
## Not run:
library("HelpersMG")
# Here two different models are fitted
x <- 1:30
y <- rnorm(30, 10, 2)+log(x)
plot(x, y)
d <- data.frame(x=x, y=y)
m1 <- lm(y ~ x, data=d)
m2 <- lm(y ~ log(x), data=d)
compare_BIC(linear=m1, log=m2, factor.value=-1)
# Here test if two datasets can be modeled with a single model
x2 <- 1:30
y2 <- rnorm(30, 15, 2)+log(x2)
plot(x, y, ylim=c(5, 25))
plot_add(x2, y2, col="red")
d2 <- data.frame(x=x2, y=y2)
m1_2 <- lm(y ~ x, data=d2)
x_grouped <- c(x, x2)
y_grouped <- c(y, y2)
d_grouped <- data.frame(x=x_grouped, y=y_grouped)
m1_grouped <- lm(y ~ x, data=d_grouped)
compare_AICc(separate=list(m1, m1_2), grouped=m1_grouped, factor.value=-1)
# Or simply
compare_AICc(m1=list(AICc=100), m2=list(AICc=102))

## End(Not run)
```

---

compare_BIC	<i>Compares the BIC of several outputs</i>
-------------	--

---

## Description

This function is used to compare the BIC of several outputs obtained with the same data but with different set of parameters.

Each object must have associated `logLik()` method with `df` and `nobs` attributes.

BIC for object `x` will be calculated as  $2 * \text{factor.value} * \sum(\text{logLik}(x)) + \sum(\text{attributes}(\text{logLik}(x))\$df) * \log(\text{attributes}(\text{logLik}(x))\$nobs)$

When several data (`i..n`) are included, the global BIC is calculated as:

$2 * \text{factor.value} * \sum(\text{logLik}(x)) \text{ for } i..n + \sum(\text{attributes}(\text{logLik}(x))\$df) \text{ for } i..n * \log(\text{attributes}(\text{logLik}(x))\$nobs) \text{ for } i..n$

## Usage

```
compare_BIC(  
  ...,  
  factor.value = -1,  
  silent = FALSE,  
  FUN = function(x) specify_decimal(x, decimals = 2)  
)
```

## Arguments

- `...` Successive results to be compared as lists.
- `factor.value` The \$value of the list object is multiplied by `factor.value` to calculate BIC.
- `silent` If TRUE, nothing is displayed.
- `FUN` Function used to show values

## Details

`compare_BIC` compares the BIC of several outputs obtained with the same data.

## Value

A list with DeltaBIC and Akaike weight for the models.

## Author(s)

Marc Girondot <[marc.girondot@gmail.com](mailto:marc.girondot@gmail.com)>

## See Also

Other AIC: [ExtractAIC.glm\(\)](#), [FormatCompareAIC\(\)](#), [compare\\_AICc\(\)](#), [compare\\_AIC\(\)](#)

## Examples

```
## Not run:
library("HelpersMG")
# Here two different models are fitted
x <- 1:30
y <- rnorm(30, 10, 2)+log(x)
plot(x, y)
d <- data.frame(x=x, y=y)
m1 <- lm(y ~ x, data=d)
m2 <- lm(y ~ log(x), data=d)
compare_BIC(linear=m1, log=m2, factor.value=-1)
# Here test if two datasets can be modeled with a single model
x2 <- 1:30
y2 <- rnorm(30, 15, 2)+log(x2)
plot(x, y, ylim=c(5, 25))
plot_add(x2, y2, col="red")
d2 <- data.frame(x=x2, y=y2)
m1_2 <- lm(y ~ x, data=d2)
x_grouped <- c(x, x2)
y_grouped <- c(y, y2)
d_grouped <- data.frame(x=x_grouped, y=y_grouped)
m1_grouped <- lm(y ~ x, data=d_grouped)
compare_BIC(separate=list(m1, m1_2), grouped=m1_grouped, factor.value=-1)

## End(Not run)
```

## contingencyTable.compare

*Contingency table comparison using Akaike weight*

## Description

This function is used as a replacement of chisq.test() to not use p-value.

## Usage

```
contingencyTable.compare(
  table,
  criterion = c("AIC", "AICc", "BIC"),
  probs = NULL
)
```

## Arguments

table	A matrix or a data.frame with series in rows and number of each category in column
criterion	Which criterion is used for model selection
probs	Series of probabilities used for conformity comparison

## Details

`contingencyTable.compare` compares contingency table using Akaike weight.

## Value

The probability that a single proportion model is sufficient to explain the data

## Author(s)

Marc Girondot <[marc.girondot@gmail.com](mailto:marc.girondot@gmail.com)>

## References

Girondot, M., Guillon, J.-M., 2018. The w-value: An alternative to t- and X<sup>2</sup> tests. *Journal of Biostatistics & Biometrics* 1, 1-4.

## See Also

Other w-value functions: `compare()`, `series.compare()`

## Examples

```
## Not run:
library("HelpersMG")

# Symmetry of Lepidochelys olivacea scutes
table <- t(data.frame(SriLanka=c(200, 157), AfricaAtl=c(19, 12),
                      Guyana=c(8, 6), Suriname=c(162, 88),
                      MexicoPac1984=c(42, 34), MexicoPac2014Dead=c(8, 9),
                      MexicoPac2014Alive=c(13, 12),
                      row.names =c("Symmetric", "Asymmetric")))
table
contingencyTable.compare(table)

table <- t(data.frame(SriLanka=c(200, 157), AfricaAtl=c(19, 12), Guyana=c(8, 6),
                      Suriname=c(162, 88), MexicoPac1984=c(42, 34),
                      MexicoPac2014Dead=c(8, 9),
                      MexicoPac2014Alive=c(13, 12), Lepidochelys.kempii=c(99, 1),
                      row.names =c("Symmetric", "Asymmetric")))
table
contingencyTable.compare(table)

# Conformity to a model
table <- matrix(c(33, 12, 25, 75), ncol = 2, byrow = TRUE)
probs <- c(0.5, 0.5)
contingencyTable.compare(table, probs=probs)

# Conformity to a model
table <- matrix(c(33, 12), ncol = 2, byrow = TRUE)
probs <- c(0.5, 0.5)
contingencyTable.compare(table, probs=probs)
```

```

# Conformity to a model
table <- matrix(c(33, 12, 8, 25, 75, 9), ncol = 3, byrow = TRUE)
probs <- c(0.8, 0.1, 0.1)
contingencyTable.compare(table, probs=probs)

# Comparison of chisq.test() and this function
table <- matrix(c(NA, NA, 25, 75), ncol = 2, byrow = TRUE)

pv <- NULL
aw <- NULL
par(new=FALSE)
n <- 100

for (GroupA in 0:n) {
  table[1, 1] <- GroupA
  table[1, 2] <- n-GroupA
  pv <- c(pv, chisq.test(table)$p.value)
  aw <- c(aw, contingencyTable.compare(table, criterion="BIC")[1])
}

x <- 0:n
y <- pv
y2 <- aw
plot(x=x, y=y, type="l", bty="n", las=1, xlab="Number of type P in Group B", ylab="Probability",
     main="", lwd=2)
lines(x=x, y=y2, type="l", col="red", lwd=2)

# w-value
(l1 <- x[which(aw>0.05)[1]])
(l2 <- rev(x)[which(rev(aw)>0.05)[1]])

aw[l1]
pv[l1]

aw[l2+2]
pv[l2+2]

# p-value
l1 <- which(pv>0.05)[1]
l2 <- max(which(pv>0.05))

aw[l1]
pv[l1]

aw[l2]
pv[l2]

y[which(y2>0.05)[1]]
y[which(rev(y2)>0.05)[1]]

par(xpd=TRUE)
text(x=25, y=1.15, labels="Group A: 25 type P / 100", pos=1)

```

```
segments(x0=25, y0=0, x1=25, y1=1, lty=3)

# plot(1, 1)

v1 <- c(expression(italic("p")*"-value"), expression("after *\chi^2*-test"))
v2 <- c(expression(italic("w")*"-value for A"), expression("and B identical models"))
legend("topright", legend=c(v1, v2),
       y.intersp = 1,
       col=c("black", "black", "red", "red"), bty="n", lty=c(1, 0, 1, 0))

segments(x0=0, x1=n, y0=0.05, y1=0.05, lty=2)
text(x=101, y=0.05, labels = "0.05", pos=4)

## End(Not run)
```

---

**convert.tz***Convert one Date-Time from one timezone to another*

---

**Description**

Convert one Date-Time from one timezone to another.  
Available timezones can be shown using OlsonNames().

**Usage**

```
convert.tz(x, tz = Sys.timezone())
```

**Arguments**

x	The date-time in POSIXlt or POSIXct format
tz	The timezone

**Details**

convert.tz Convert one Date-Time from one timezone to another

**Value**

A POSIXlt or POSIXct date converted

**Author(s)**

Marc Girondot <marc.girondot@gmail.com>

**See Also**

Function `with_tz()` from `lubridate` package does the same. I keep it here only for compatibility with old scripts.

## Examples

```
d <- as.POSIXlt("2010-01-01 17:34:20", tz="UTC")
convert.tz(d, tz="America/Guatemala")
```

cutter

*Distribution of the fitted distribution without cut.*

## Description

If observations is a data.frame, it can have 4 columns:

A column for the measurements;

A column for the lower detection limit;

A column for the upper detection limit;

A column for the truncated or censored nature of the data.

The names of the different columns are in the observations.colname, lower\_detection\_limit.colname, upper\_detection\_limit.colname and cut\_method.colname.

If lower\_detection\_limit.colname is NULL or if the column does not exist, the data are supposed to not be left-cut and if upper\_detection\_limit.colname is NULL or if the column does not exist, the data are supposed to not be right-cut.

If observations is a vector, then the parameters lower\_detection\_limit and/or upper\_detection\_limit must be given. Then cut\_method must be also provided.

In observations, -Inf must be used to indicate a value below the lower detection limit and +Inf must be used for a value above the upper detection limit.

Be careful: NA is used to represent a missing data and not a value below or above the detection limit.

If lower\_detection\_limit, upper\_detection\_limit or cut\_method are only one value, they are supposed to be used for all the observations.

Definitions for censored or truncated distribution vary, and the two terms are sometimes used interchangeably. Let the following data set:

1 1.25 2 4 5

Censoring: some observations will be censored, meaning that we only know that they are below (or above) some bound. This can for instance occur if we measure the concentration of a chemical in a water sample. If the concentration is too low, the laboratory equipment cannot detect the presence of the chemical. It may still be present though, so we only know that the concentration is below the laboratory's detection limit.

If the detection limit is 1.5, so that observations that fall below this limit is censored, our example data set would become:

<1.5 <1.5 2 4 5; that is, we don't know the actual values of the first two observations, but only that they are smaller than 1.5.

Truncation: the process generating the data is such that it only is possible to observe outcomes above (or below) the truncation limit. This can for instance occur if measurements are taken using a detector which only is activated if the signals it detects are above a certain limit. There may be lots of weak incoming signals, but we can never tell using this detector.

If the truncation limit is 1.5, our example data set would become: 2 4 5; and we would not know that there in fact were two signals which were not recorded.

If `n.iter` is `NULL`, no Bayesian MCMC is performed but credible interval will not be available.

## Usage

```
cutter(
  observations = stop("Observations must be provided"),
  observations.colname = "Observations",
  lower_detection_limit.colname = "LDL",
  upper_detection_limit.colname = "UDL",
  cut_method.colname = "Cut",
  par = NULL,
  lower_detection_limit = NULL,
  upper_detection_limit = NULL,
  cut_method = "censored",
  distribution = "gamma",
  n.mixture = 1,
  n.iter = 5000,
  debug = FALSE,
  progress.bar = TRUE,
  session = NULL
)
```

## Arguments

<code>observations</code>	The observations; see description
<code>observations.colname</code>	If <code>observations</code> is a <code>data.frame</code> , the name of column with <code>observations</code>
<code>lower_detection_limit.colname</code>	If <code>observations</code> is a <code>data.frame</code> , the name of column with lower detection limit
<code>upper_detection_limit.colname</code>	If <code>observations</code> is a <code>data.frame</code> , the name of column with upper detection limit
<code>cut_method.colname</code>	If <code>observations</code> is a <code>data.frame</code> , the name of column with cut method, being " <code>censored</code> " or " <code>truncated</code> "
<code>par</code>	Initial values for parameters of distribution
<code>lower_detection_limit</code>	Value for lower detection limit
<code>upper_detection_limit</code>	Value for upper detection limit
<code>cut_method</code>	Value for cut method, being " <code>censored</code> " or " <code>truncated</code> "
<code>distribution</code>	Can be <code>gamma</code> , <code>normal</code> , <code>weibull</code> , <code>lognormal</code> , or <code>generalized.gamma</code>
<code>n.mixture</code>	Number of distributions
<code>n.iter</code>	Number of iteration for Bayesian MCMC and to estimate the goodness-of-fit

debug	If TRUE, show some information
progress.bar	If TRUE, show a progress bar for MCMC
session	The session of a shiny process

**Details**

*cutter* returns the fitted distribution without cut

**Value**

The parameters of distribution of values below or above the detection limit.

**Author(s)**

Marc Girondot <marc.girondot@gmail.com>

**See Also**

Other Distributions: [dbeta\\_new\(\)](#), [dcutter\(\)](#), [dggamma\(\)](#), [logLik.cutter\(\)](#), [pSnbinoM\(\)](#), [plot.cutter\(\)](#), [print.cutter\(\)](#), [qSnbinoM\(\)](#), [r2norm\(\)](#), [rcutter\(\)](#), [rmnrm\(\)](#), [rnbinom\\_new\(\)](#)

**Examples**

```
## Not run:
library(HelpersMG)
#
# -----
# right censored distribution with gamma distribution
# -----
# Detection limit
DL <- 100
# Generate 100 random data from a gamma distribution
obc <- rgamma(100, scale=20, shape=2)
# remove the data below the detection limit
obc[obc>DL] <- +Inf
# search for the parameters the best fit these censored data
result <- cutter(observations=obc, upper_detection_limit=DL,
                  cut_method="censored")
result
plot(result, xlim=c(0, 150), breaks=seq(from=0, to=150, by=10), col.mcmc=NULL)
plot(result, xlim=c(0, 150), breaks=seq(from=0, to=150, by=10))
#
# -----
# The same data seen as truncated data with gamma distribution
#
# -----
obc <- obc[is.finite(obc)]
# search for the parameters the best fit these truncated data
result <- cutter(observations=obc, upper_detection_limit=DL,
                  cut_method="truncated")
result
plot(result, xlim=c(0, 150), breaks=seq(from=0, to=150, by=10))
#
# -----
# left censored distribution with gamma distribution
```

```

# -----
# Detection limit
DL <- 10
# Generate 100 random data from a gamma distribution
obc <- rgamma(100, scale=20, shape=2)
# remove the data below the detection limit
obc[obc<DL] <- -Inf
# search for the parameters the best fit these truncated data
result <- cutter(observations=obc, lower_detection_limit=DL,
                  cut_method="censored")
result
plot(result, xlim=c(0, 200), breaks=seq(from=0, to=300, by=10))
# -----
# left censored distribution with mixture of gamma distribution
# -----
#' # Detection limit
library(HelpersMG)
# Generate 200 random data from a gamma distribution
set.seed(1234)
obc <- c(rgamma(100, scale=10, shape=5), rgamma(100, scale=20, shape=10))
LDL <- 20
l <- seq(from=0, to=LDL, length.out=1001)
p <- pgamma(l, scale=10, shape=5)*0.5+pgamma(l, scale=20, shape=10)
deltal <- l[2]-l[1]
expected_LDL <- sum((l[-1]-deltal/2)*(p[-1]-p[-length(p)]))/sum((p[-1]-p[-length(p)]))
# remove the data below the detection limit
obc[obc<LDL] <- -Inf

UDL <- 300
l <- seq(from=UDL, to=1000, length.out=1001)
p <- pgamma(l, scale=10, shape=5)*0.5+pgamma(l, scale=20, shape=10)
deltal <- l[2]-l[1]
expected_UDL <- sum((l[-1]-deltal/2)*(p[-1]-p[-length(p)]))/sum((p[-1]-p[-length(p)]))
obc[obc>UDL] <- +Inf

# search for the parameters the best fit these truncated data
result1_gamma <- cutter(observations=obc, lower_detection_limit=LDL,
                        upper_detection_limit = UDL,
                        distribution="gamma",
                        cut_method="censored", n.iter=5000, debug=0)
result1_normal <- cutter(observations=obc, lower_detection_limit=LDL,
                         upper_detection_limit = UDL,
                         distribution="normal",
                         cut_method="censored", n.iter=5000)
result1_lognormal <- cutter(observations=obc, lower_detection_limit=LDL,
                            upper_detection_limit = UDL,
                            distribution="lognormal",
                            cut_method="censored", n.iter=5000)
result1_Weibull <- cutter(observations=obc, lower_detection_limit=LDL,
                           upper_detection_limit = UDL,
                           distribution="Weibull",
                           cut_method="censored", n.iter=5000)
result1_generalized.gamma <- cutter(observations=obc, lower_detection_limit=LDL,

```

```

upper_detection_limit = UDL,
distribution="generalized.gamma",
cut_method="censored", n.iter=5000)
result2_gamma <- cutter(observations=obc, lower_detection_limit=LDL,
                        upper_detection_limit = UDL,
                        distribution="gamma",
                        n.mixture=2,
                        cut_method="censored", n.iter=5000)
result2_normal <- cutter(observations=obc, lower_detection_limit=LDL,
                         upper_detection_limit = UDL,
                         distribution="normal",
                         n.mixture=2,
                         cut_method="censored", n.iter=5000)
result2_lognormal <- cutter(observations=obc, lower_detection_limit=LDL,
                            upper_detection_limit = UDL,
                            distribution="lognormal",
                            n.mixture=2,
                            cut_method="censored", n.iter=5000)
result2_Weibull <- cutter(observations=obc, lower_detection_limit=LDL,
                           upper_detection_limit = UDL,
                           distribution="Weibull",
                           n.mixture=2,
                           cut_method="censored", n.iter=5000)
result2_generalized.gamma <- cutter(observations=obc, lower_detection_limit=LDL,
                                      upper_detection_limit = UDL,
                                      distribution="generalized.gamma",
                                      n.mixture=2,
                                      cut_method="censored", n.iter=5000)

compare_AIC(nomixture.gamma=result1_gamma,
            nomixture.normal=result1_normal,
            nomixture.lognormal=result1_lognormal,
            nomixture.Weibull=result1_Weibull,
            nomixture.generalized.gamma=result1_generalized.gamma,
            mixture.gamma=result2_gamma,
            mixture.normal=result2_normal,
            mixture.lognormal=result2_lognormal,
            mixture.Weibull=result2_Weibull,
            mixture.generalized.gamma=result2_generalized.gamma)

plot(result2_gamma, xlim=c(0, 600), breaks=seq(from=0, to=600, by=10))
plot(result2_generalized.gamma, xlim=c(0, 600), breaks=seq(from=0, to=600, by=10))

# -----
# left and right censored distribution
# -----
# Generate 100 random data from a gamma distribution
obc <- rgamma(100, scale=20, shape=2)
# Detection limit
LDL <- 10
# remove the data below the detection limit
obc[obc<LDL] <- -Inf
# Detection limit

```

```

UDL <- 100
# remove the data below the detection limit
obc[obc>UDL] <- +Inf
# search for the parameters the best fit these censored data
result <- cutter(observations=obc, lower_detection_limit=LDL,
                  upper_detection_limit=UDL,
                  cut_method="censored")
result
plot(result, xlim=c(0, 150), col.DL=c("black", "grey"),
      col.unobserved=c("green", "blue"),
      breaks=seq(from=0, to=150, by=10))
#
# -----
# Example with two values for lower detection limits
# corresponding at two different methods of detection for example
# with gamma distribution
#
# -----
obc <- rgamma(50, scale=20, shape=2)
# Detection limit for sample 1 to 50
LDL1 <- 10
# remove the data below the detection limit
obc[obc<LDL1] <- -Inf
obc2 <- rgamma(50, scale=20, shape=2)
# Detection limit for sample 1 to 50
LDL2 <- 20
# remove the data below the detection limit
obc2[obc2<LDL2] <- -Inf
obc <- c(obc, obc2)
# search for the parameters the best fit these censored data
result <- cutter(observations=obc,
                  lower_detection_limit=c(rep(LDL1, 50), rep(LDL2, 50)),
                  cut_method="censored")
result
# It is difficult to choose the best set of colors
plot(result, xlim=c(0, 150), col.dist="red",
      col.unobserved=c(rgb(red=1, green=0, blue=0, alpha=0.1),
                        rgb(red=1, green=0, blue=0, alpha=0.2)),
      col.DL=c(rgb(red=0, green=0, blue=1, alpha=0.5),
                rgb(red=0, green=0, blue=1, alpha=0.9)),
      breaks=seq(from=0, to=200, by=10))

#
# -----
# left censored distribution comparison of normal, lognormal,
# weibull, generalized gamma, and gamma without Bayesian MCMC
# Comparison with Akaike Information Criterion
#
# -----
# Detection limit
DL <- 10
# Generate 100 random data from a gamma distribution
obc <- rgamma(100, scale=20, shape=2)
# remove the data below the detection limit
obc[obc<DL] <- -Inf

result_gamma <- cutter(observations=obc, lower_detection_limit=DL,

```

```

            cut_method="censored", distribution="gamma",
            n.iter=NULL)
plot(result_gamma)

result_lognormal <- cutter(observations=obc, lower_detection_limit=DL,
                           cut_method="censored", distribution="lognormal",
                           n.iter=NULL)
plot(result_lognormal)

result_weibull <- cutter(observations=obc, lower_detection_limit=DL,
                           cut_method="censored", distribution="weibull",
                           n.iter=NULL)
plot(result_weibull)

result_normal <- cutter(observations=obc, lower_detection_limit=DL,
                        cut_method="censored", distribution="normal",
                        n.iter=NULL)
plot(result_normal)

result_generalized.gamma <- cutter(observations=obc, lower_detection_limit=DL,
                                      cut_method="censored", distribution="generalized.gamma",
                                      n.iter=NULL)
plot(result_generalized.gamma)

compare_AIC(gamma=result_gamma,
            lognormal=result_lognormal,
            normal=result_normal,
            Weibull=result_weibull,
            Generalized.gamma=result_generalized.gamma)

# -----
# left censored distribution comparison of normal, lognormal,
# weibull, generalized gamma, and gamma
# -----
# Detection limit
DL <- 10
# Generate 100 random data from a gamma distribution
obc <- rgamma(100, scale=20, shape=2)
# remove the data below the detection limit
obc[obc<DL] <- -Inf
# search for the parameters the best fit these truncated data
result_gamma <- cutter(observations=obc, lower_detection_limit=DL,
                       cut_method="censored", distribution="gamma")
result_gamma
plot(result_gamma, xlim=c(0, 250), breaks=seq(from=0, to=250, by=10))

result_lognormal <- cutter(observations=obc, lower_detection_limit=DL,
                           cut_method="censored", distribution="lognormal")
result_lognormal
plot(result_lognormal, xlim=c(0, 250), breaks=seq(from=0, to=250, by=10))

result_weibull <- cutter(observations=obc, lower_detection_limit=DL,
                         cut_method="censored", distribution="weibull")

```

```
result_weibull
plot(result_weibull, xlim=c(0, 250), breaks=seq(from=0, to=250, by=10))

result_normal <- cutter(observations=obc, lower_detection_limit=DL,
                        cut_method="censored", distribution="normal")
result_normal
plot(result_normal, xlim=c(0, 250), breaks=seq(from=0, to=250, by=10))

result_generalized.gamma <- cutter(observations=obc, lower_detection_limit=DL,
                                      cut_method="censored", distribution="generalized.gamma")
result_generalized.gamma
plot(result_generalized.gamma, xlim=c(0, 250), breaks=seq(from=0, to=250, by=10))

# -----
# Test for similarity in gamma left censored distribution between two
# datasets
# -----
obc1 <- rgamma(100, scale=20, shape=2)
# Detection limit for sample 1 to 50
LDL <- 10
# remove the data below the detection limit
obc1[obc1<LDL] <- -Inf
obc2 <- rgamma(100, scale=10, shape=2)
# remove the data below the detection limit
obc2[obc2<LDL] <- -Inf
# search for the parameters the best fit these censored data
result1 <- cutter(observations=obc1,
                   distribution="gamma",
                   lower_detection_limit=LDL,
                   cut_method="censored", n.iter=NULL)
logLik(result1)
plot(result1, xlim=c(0, 200),
      breaks=seq(from=0, to=200, by=10))
result2 <- cutter(observations=obc2,
                   distribution="gamma",
                   lower_detection_limit=LDL,
                   cut_method="censored", n.iter=NULL)
logLik(result2)
plot(result2, xlim=c(0, 200),
      breaks=seq(from=0, to=200, by=10))
result_totl <- cutter(observations=c(obc1, obc2),
                      distribution="gamma",
                      lower_detection_limit=LDL,
                      cut_method="censored", n.iter=NULL)
logLik(result_totl)
plot(result_totl, xlim=c(0, 200),
      breaks=seq(from=0, to=200, by=10))

compare_AIC(Separate=list(result1, result2),
             Common=result_totl, factor.value=1)
compare_BIC(Separate=list(result1, result2),
             Common=result_totl, factor.value=1)
```

---

```
## End(Not run)
```

---

**d***Write an ASCII Representation of a vector object***Description**

Writes an ASCII text representation of an R object.

It can be used as a replacement of dput() for named vectors.

The controls "keepNA", "keepInteger" and "showAttributes" are utilized for named vectors.

**Usage**

```
d(
  x,
  file = "",
  control = c("keepNA", "keepInteger", "showAttributes"),
  collapse = ", \n"
)
```

**Arguments**

<code>x</code>	A named vector object
<code>file</code>	either a character string naming a file or a connection. "" indicates output to the console.
<code>control</code>	character vector indicating deparsing options. See .deparseOpts for their description.
<code>collapse</code>	Characters used to separate values.

**Details**

`d` Write an ASCII Representation of a vector object

**Value**

A string

**Author(s)**

Marc Girondot <marc.girondot@gmail.com>

**See Also**

Other Characters: [asc\(\)](#), [char\(\)](#), [tnirp\(\)](#)

**Examples**

```
d(c(A=10, B=20))
dput(c(A=10, B=20))
```

---

<code>dbeta_new</code>	<i>Density for the Beta distributions.</i>
------------------------	--

---

## Description

Density for the Beta distribution with parameters mu and v or shape1 and shape2 (and optional non-centrality parameter ncp).

The returned object has three attributes:

shape1, shape2, and ncp

Note that if x has attributes, they are preserved.

## Usage

```
dbeta_new(
  x,
  mu = NULL,
  v = NULL,
  shape1,
  shape2,
  ncp = 0,
  log = FALSE,
  silent = FALSE
)
```

## Arguments

<code>x</code>	vector of quantiles.
<code>mu</code>	mean of the Beta distribution.
<code>v</code>	variance of the Beta distribution.
<code>shape1</code>	non-negative parameters of the Beta distribution.
<code>shape2</code>	non-negative parameters of the Beta distribution.
<code>ncp</code>	non-centrality parameter.
<code>log</code>	logical; if TRUE, probabilities p are given as log(p).
<code>silent</code>	If FALSE, show the shape1 and shape 2 values.

## Details

`dbeta_new` returns the density for the Beta distributions

The Beta distribution with parameters shape1 = a and shape2 = b has density

$$\text{gamma}(a+b)/(\text{gamma}(a)\text{gamma}(b))x^{(a-1)}(1-x)^{(b-1)}$$

for  $a > 0, b > 0$  and  $0 \leq x \leq 1$  where the boundary values at  $x=0$  or  $x=1$  are defined as by continuity (as limits).

The mean is  $a/(a+b)$  and the variance is  $ab/((a+b)^2 (a+b+1))$ . These moments and all distributional properties can be defined as limits.

**Value**

`dbeta_new` gives the density for the Beta distributions

**Author(s)**

Marc Girondot <[marc.girondot@gmail.com](mailto:marc.girondot@gmail.com)>

**See Also**

Other Distributions: [cutter\(\)](#), [dcutter\(\)](#), [dggamma\(\)](#), [logLik.cutter\(\)](#), [pSnbino\(\)](#), [plot.cutter\(\)](#), [print.cutter\(\)](#), [qSnbino\(\)](#), [r2norm\(\)](#), [rcutter\(\)](#), [rmnrm\(\)](#), [rnbinom\\_new\(\)](#)

**Examples**

```
pi <- rbeta(100, shape1=0.48, shape2=0.12)
hist(pi, freq=FALSE, breaks=seq(from=0, to=1, by=0.1), ylim=c(0, 8), las=1)
library("HelpersMG")
mx <- ScalePreviousPlot()$ylim["end"]/
      max(dbeta_new(seq(from=0.01, to=0.99, by=0.01), mu = 0.8, v=0.1))
curve(dbeta_new(x, mu = 0.8, v=0.1)*mx, add=TRUE, col="red")
out <- dbeta_new(0.1, mu = 0.8, v=0.1)
out
attributes(out)$shape1; attributes(out)$shape2; attributes(out)$ncp
dbeta(0.1, shape1=attributes(out)$shape1, shape2=attributes(out)$shape2,
      ncp=attributes(out)$ncp)

# It can be used to generate random numbers using mu and v
out <- dbeta_new(0.1, mu = 0.8, v=0.1, silent=TRUE)
pi <- rbeta(100, shape1=attributes(out)$shape1, shape2=attributes(out)$shape2,
           ncp=attributes(out)$ncp)
hist(pi, freq=FALSE, breaks=seq(from=0, to=1, by=0.1), ylim=c(0, 8), las=1)
```

**dcutter**

*Distribution of the fitted distribution without cut.*

**Description**

If observations must be a data.frame with 4 columns:  
 observations: A column for the measurements;  
 LDL: A column for the lower detection limit;  
 UDL: A column for the upper detection limit;  
 Cut: A column for the truncated or censored nature of the data.

**Usage**

```
dcutter(
  par,
  observations = NULL,
  distribution = "gamma",
  n.mixture = NULL,
  debug = FALSE,
  limits.lower = NULL,
  limits.upper = NULL,
  log = TRUE
)
```

**Arguments**

<code>par</code>	Values for parameters of distribution
<code>observations</code>	The observations; see description.
<code>distribution</code>	Can be gamma, normal, weibull, lognormal, or generalized.gamma.
<code>n.mixture</code>	Number of distributions
<code>debug</code>	If TRUE, show some information. If 2, show more information.
<code>limits.lower</code>	Value for lower detection limit
<code>limits.upper</code>	Value for upper detection limit
<code>log</code>	If TRUE, return the log likelihood

**Details**

`dcutter` returns the density of the cutter function

**Value**

The density of the cutter function according to observations.

**Author(s)**

Marc Girondot <[marc.girondot@gmail.com](mailto:marc.girondot@gmail.com)>

**See Also**

Other Distributions: [cutter\(\)](#), [dbeta\\_new\(\)](#), [dggamma\(\)](#), [logLik.cutter\(\)](#), [pSnbinoM\(\)](#), [plot.cutter\(\)](#), [print.cutter\(\)](#), [qSnbinoM\(\)](#), [r2norm\(\)](#), [rcutter\(\)](#), [rmnrm\(\)](#), [rnbinom\\_new\(\)](#)

**Examples**

```
## Not run:
library(HelperMG)
par <- c('shape1' = 0.42265849507444225,
       'scale1' = 14.139457094879594,
       'shape2' = 1.667131542489706,
       'scale2' = 0.10763344388223803,
```

```

'p1' = 0.12283307526788023)
obs <- data.frame(Observations=c(0.755, 1.013, 2.098, 6.265, 4.708, 0.078, 2.169, 0.403, 1.251,
0.008, 1.419, 1.078, 2.744, 81.534, 1.426, 13.486, 7.813, 0.165,
0.118, 0.864, 0.369, 7.159, 2.605, 1.579, 1.646, 0.484, 4.492,
0.139, 0.28, 0.154, 0.106, 0.104, 4.185, 0.735, 0.149, 0.183,
0.062, 8.246, 0.165, 0.121, 0.109, 0.092, 0.162, 0.108, 0.139,
0.141, 0.124, 0.124, 0.151, 0.141, 0.364, 0.295, 0.09, 0.135,
0.154, 0.218, 0.167, -Inf, 0.203, 0.228, 0.107, 0.162, 0.194,
0.322, 0.351, 0.17, 0.236, 0.176, 0.107, 0.12, 0.095, 0.27, 0.194,
0.125, 0.123, 0.085, 0.164, 0.106, 0.079, 0.162),
LDL=0.001, UDL=NA, Cut="censored")
dcutter(par=par, observations=obs, distribution="gamma",
n.mixture=NULL, debug=FALSE, limits.lower=NULL,
limits.upper=NULL, log=FALSE)
dcutter(par=par, observations=obs, distribution="gamma",
n.mixture=NULL, debug=FALSE, limits.lower=NULL,
limits.upper=NULL, log=TRUE)

## End(Not run)

```

**dggamma***Generalized gamma distribution.***Description**

Generalized gamma distribution

**Usage**

```

dggamma(x, theta, kappa, delta, log = FALSE)

pggamma(q, theta, kappa, delta, lower.tail = TRUE, log.p = FALSE)

qggamma(p, theta, kappa, delta, lower.tail = TRUE, log.p = FALSE)

rggamma(n, theta, kappa, delta)

```

**Arguments**

<code>x, q</code>	vector of quantiles.
<code>theta</code>	scale parameter.
<code>kappa</code>	shape parameter.
<code>delta</code>	shape parameter.
<code>log, log.p</code>	logical; if TRUE, probabilities $p$ are given as $\log(p)$ .
<code>lower.tail</code>	logical; if TRUE (default), probabilities are $P[X \leq x]$ otherwise, $P[X > x]$ .
<code>p</code>	vector of probabilities.
<code>n</code>	number of observations.

## Details

`pgamma`, `qgamma`, `dgamma`, and `rgamma` are used to model the generalized gamma distribution.

The code is modified from <https://rpubs.com/FJRUBIO/GG>.

## Value

`dggamma` gives the density, `pgamma` gives the distribution function, `qgamma` gives the quantile function, and `rgamma` generates random deviates.

## Functions

- `dggamma`: Density of the generalized gamma.
- `pgamma`: Distribution function of the generalized gamma.
- `qgamma`: Quantile of the generalized gamma.
- `rgamma`: Random of the generalized gamma.

## More details here

The generalized gamma is described here [https://en.wikipedia.org/wiki/Generalized\\_gamma\\_distribution](https://en.wikipedia.org/wiki/Generalized_gamma_distribution).

With  $a$  being theta,  $b$  being kappa, and  $p$  being delta.  
 $\theta$ ,  $\kappa$  and  $\delta$  must be all  $> 0$ .

## Author(s)

Marc Girondot <[marc.girondot@gmail.com](mailto:marc.girondot@gmail.com)>

## See Also

Other Distributions: `cutter()`, `dbeta_new()`, `dcutter()`, `logLik.cutter()`, `pSnbnom()`, `plot.cutter()`, `print.cutter()`, `qSnbnom()`, `r2norm()`, `rcutter()`, `rmnrm()`, `rnbnom_new()`

## Examples

```
# To reproduce the wikipedia page graphic
x <- seq(from=0, to=8, by=0.1)
plot(x, dgamma(x, theta=2, kappa=0.5, delta=0.5), lty=1, col="blue",
     type="l", lwd=2, xlab="x", ylab="PDF")
lines(x, dgamma(x, theta=1, kappa=1, delta=0.5), lty=1, col="green", lwd=2)
lines(x, dgamma(x, theta=2, kappa=1, delta=2), lty=1, col="red", lwd=2)
lines(x, dgamma(x, theta=5, kappa=1, delta=5), lty=1, col="yellow", lwd=2)
lines(x, dgamma(x, theta=7, kappa=1, delta=7), lty=1, col="grey", lwd=2)
legend("topright", legend=c("a=2, d=0.5, p=0.5",
                            "a=2, d=1, p=2", "a=5, d=1, p=5", "a=7, d=1, p=7"),
       col=c("blue", "green", "red", "yellow", "grey"),
       lty=1, lwd=2, bty="n")
```

---

DIx	<i>Return an index of quantitative asymmetry and complexity named Developmental Instability Index (DIx)</i>
-----	---

---

## Description

Return an index of quantitative asymmetry and complexity.

Higher is the value, higher is the complexity (number of objects) and diversity (difference between them).

The indice is based on the product of the average angular distance of Edwards (1971) for all permutations of measures for both sides with the geometric mean of the inverse of Shannon entropy H for both sides. Let p1 and p2 two vectors of relative measures of objects with sum(p1) = 1 and sum(p2)=1 and n1 being the number of objects in p1 and n2 being the number of objects in p2.

Edwards distance for all permutations of p1 and p2 objects are computed and the average value E is calculated.

The maximun possible Shannon index for identical n1 is max1 = sum((1/n1) \* log(1/n1)).

Shannon index is v1 = sum(p1 \* log(p1)).

If version == 2, the complementary of Shannon index for these n1 objects is used: c1 = 2 \* max1 - v1

If version == 1, the Shannon index is used directly.

The geometry mean between both sides defined the measure of diversity within each side: S=sqrt(c1 \* c2)

The Developmental Instability Index is then S \* E

## Usage

```
DIx(l1, l2, details = FALSE, version = 1)
```

## Arguments

l1	Set of measures at one side of an organism
l2	Set of measures at the other side of an organism
details	If TRUE, will show the details of computing
version	Can be 1 or 2; see description

## Details

DIx returns an index of quantitative asymmetry and complexity

## Value

A numeric value

## Author(s)

Marc Girondot <[marc.girondot@gmail.com](mailto:marc.girondot@gmail.com)>

## References

- Edwards, A.W.F., 1971. Distances between populations on the basis of gene frequencies. *Biometrics* 27, 873–881.  
 Shannon C.E. 1948 A mathematical theory of communication. *Bell System Technical Journal* 27(3), 379-423.

## Examples

```
## Not run:
11 <- c(0.1, 0.1, 0.05, 0.2, 0.3, 0.25)
12 <- c(0.2, 0.3, 0.5)
DIx(11, 12)

11 <- c(0.1, 0.1, 0.05, 0.2, 0.3, 0.25)
12 <- c(0.1, 0.1, 0.05, 0.2, 0.3, 0.25)
DIx(11, 12)

11 <- c(0.2, 0.3, 0.5)
12 <- c(0.2, 0.3, 0.5)
DIx(11, 12)

11 <- c(0.2, 0.2, 0.2, 0.2, 0.2)
12 <- c(0.2, 0.3, 0.5)
DIx(11, 12)

11 <- c(0.2, 0.2, 0.2, 0.2, 0.2)
12 <- c(0.3333, 0.3333, 0.3333)
DIx(11, 12)

11 <- c(0.2, 0.2, 0.2, 0.2, 0.2)
12 <- c(0.2, 0.2, 0.2, 0.2, 0.2)
DIx(11, 12)

11 <- c(0.3333, 0.3333, 0.3333)
12 <- c(0.3333, 0.3333, 0.3333)
DIx(11, 12)

## End(Not run)
```

## Description

Density for the negative binomial distribution with parameters mu, sd, var, size or prob. See dnbinom.

**Usage**

```
dnbinom_new(
  x,
  size = NULL,
  prob = NULL,
  mu = NULL,
  sd = NULL,
  var = NULL,
  log = FALSE
)
```

**Arguments**

<code>x</code>	vector of (non-negative integer) quantiles.
<code>size</code>	target for number of successful trials, or dispersion parameter (the shape parameter of the gamma mixing distribution). Must be strictly positive, need not be integer.
<code>prob</code>	probability of success in each trial. $0 < \text{prob} \leq 1$ .
<code>mu</code>	alternative parametrization via mean.
<code>sd</code>	alternative parametrization via standard deviation.
<code>var</code>	alternative parametrization via variance.
<code>log</code>	logical; if TRUE, probabilities p are given as log(p).

**Details**

`dnbinom_new` returns density for the negative binomial distribution

**Value**

Random numbers for the negative binomial distribution

**Author(s)**

Marc Girondot <[marc.girondot@gmail.com](mailto:marc.girondot@gmail.com)>

**Examples**

```
## Not run:
library("HelpersMG")
set.seed(1)
x <- rnbinom_new(n=100, mu=2, sd=3)
LnL <- NULL
df <- data.frame(mu=seq(from=0.1, to=8, by=0.1), "-LnL"=NA)
for (mu in df[, "mu"]){
  LnL <- c(LnL, -sum(dnbinom_new(x=x, mu=mu, sd=3, log=TRUE)))
}
df[, "-LnL"] <- LnL
ggplot(data = df, aes(x = .data[["mu"]], y = .data[["-LnL"]])) + geom_line()
# Examples of wrong parametrization
```

```
dnbinom_new(x=x, mu=c(1, 2), sd=3, log=TRUE)
## End(Not run)
```

dSnbnom

*Density for the sum of random variable with negative binomial distributions.*

## Description

Density for the sum of random variable with negative binomial distributions.  
If all prob values are the same, infinite is automatically set to 0.

## Usage

```
dSnbnom(
  x = stop("You must provide a x value"),
  size = NULL,
  prob = NULL,
  mu = NULL,
  log = FALSE,
  tol = 1e-06,
  infinite = 1000
)
```

## Arguments

x	vector of (non-negative integer) quantiles.
size	target for number of successful trials, or dispersion parameter (the shape parameter of the gamma mixing distribution). Must be strictly positive, need not be integer.
prob	probability of success in each trial. $0 < \text{prob} \leq 1$ .
mu	alternative parametrization via mean.
log	logical; if TRUE, probabilities p are given as log(p).
tol	Tolerance for recurrence
infinite	Maximum level of recursion

## Details

dSnbnom returns the density for the sum of random variable with negative binomial distributions

## Value

dSnbnom gives the density

**Author(s)**

Marc Girondot <marc.girondot@gmail.com>

**References**

Furman, E., 2007. On the convolution of the negative binomial random variables. Statistics & Probability Letters 77, 169-172.

**See Also**

Other Distribution of sum of random variable with negative binomial distributions: [rSnbinom\(\)](#)

**Examples**

```
## Not run:
library(HelpersMG)
alpha <- c(1, 2, 5, 1, 2)
p <- c(0.1, 0.12, 0.13, 0.14, 0.14)
dSnbinom(20, size=alpha, prob=p)
dSnbinom(20, size=alpha, prob=p, log=TRUE)
dSnbinom(20, size=2, mu=c(0.01, 0.02, 0.03))
dSnbinom(20, size=2, mu=c(0.01, 0.02, 0.03), log=TRUE)
# Test with a single distribution
dSnbinom(20, size=1, mu=20)
# when only one distribution is available, it is the same as dnbinom()
dnbinom(20, size=1, mu=20)
# If a parameter is supplied as only one value, it is supposed to be constant
dSnbinom(20, size=1, mu=c(14, 15, 10))
# The function is vectorized:
plot(0:200, dSnbinom(0:200, size=alpha, prob=p), bty="n", type="h", xlab="x", ylab="Density")
# Comparison with simulated distribution using rep replicates
alpha <- c(2.1, 2.05, 2)
mu <- c(10, 30, 20)
rep <- 10000
distEmpirique <- rSnbinom(rep, size=alpha, mu=mu)
tabledistEmpirique <- rep(0, 301)
names(tabledistEmpirique) <- as.character(0:300)
tabledistEmpirique[names(table(distEmpirique))] <- table(distEmpirique)/rep

plot(0:300, dSnbinom(0:300, size=alpha, mu=mu), type="h", bty="n",
     xlab="x", ylab="Density", ylim=c(0,0.02))
plot_add(0:300, tabledistEmpirique, type="l", col="red")
legend(x=200, y=0.02, legend=c("Empirical", "Theoretical"),
       text.col=c("red", "black"), bty="n")

# Example with the approximation mu=mean(mu)
plot(0:300, dSnbinom(0:300, size=alpha, mu=mu), type="h", bty="n",
     xlab="x", ylab="Density", ylim=c(0,0.02))
plot_add(0:300, tabledistEmpirique, type="l", col="red")
legend(x=200, y=0.02, legend=c("Empirical", "Theoretical"),
       text.col=c("red", "black"), bty="n")
```

```

# example to fit the distribution
data <- rnbnom(1000, size=1, mu=10)
hist(data)
ag <- rep(1:100, 10)
r <- aggregate(data, by=list(ag), FUN=sum)
hist(r[,2])

parx <- c(size=1, mu=10)

dSnbomx <- function(x, par) {
  -sum(dSnbom(x=x[,2], mu=rep(par["mu"], 10), size=par["size"], log=TRUE))
}

fit_mu_size <- optim(par = parx, fn=dSnbomx, x=r, method="BFGS", control=c(trace=TRUE))
fit_mu_size$par

## End(Not run)

```

duplicated\_packages    *List the duplicated packages with their locations*

## Description

A data.frame with the duplicated packages and their locations and version.  
The columns Lib1 and Version1 should have the oldest version of the packages. Then you can try:  
li <- duplicated\_packages()  
if (nrow(li) != 0)  
for (i in 1:nrow(li))  
remove.packages(rownames(li)[i], lib=li[i, "Lib1"])

## Usage

```
duplicated_packages()
```

## Details

duplicated\_packages lists the duplicated packages with their locations

## Value

A data.frame with 4 elements for each duplicated packages:  
- versions: the version of the packages  
- libraries: the locations

## Author(s)

Marc Girondot <marc.girondot@gmail.com>

## Examples

```
## Not run:
library(HelpersMG)
duplicated_packages()

## End(Not run)
```

**ellipse**

*Plot an ellipse*

## Description

Plot a ellipse dined by the center and the radius. The options for binomial confidence are:

- alpha is 1 - confidence interval
- method must be one of these "wilson", "exact", "asymptotic"
- col parameter can be a list of colors. See examples

## Usage

```
ellipse(
  center.x = 0,
  center.y = 0,
  radius.x = 1,
  radius.y = 1,
  radius.x.lower = NULL,
  radius.x.upper = NULL,
  radius.y.lower = NULL,
  radius.y.upper = NULL,
  alpha = 0,
  binconf.x = NULL,
  binconf.y = NULL,
  control.binconf = list(alpha = 0.05, method = "wilson"),
  length = 100,
  ...
)
```

## Arguments

center.x	Center of the ellipse on x axis
center.y	Center of the ellipse on y axis
radius.x	Radius along the x axis
radius.y	Radius along the y axis
radius.x.lower	Radius along the x axis, at left of center
radius.x.upper	Radius along the x axis, at right of center
radius.y.lower	Radius along the y axis, at bottom of center

radius.y.upper	Radius along the y axis, at top of center
alpha	Rotation in radians
binconf.x	A data.frame or a matrix with two columns, x and n or with three columns, PointEst, Lower, and Upper
binconf.y	A data.frame or a matrix with two columns, x and n or with three columns, PointEst, Lower, and Upper
control.binconf	A list with options for binomial confidence
length	Number of points to draw the ellipse
...	Graphical parameters

## Details

ellipse plots an ellipse

## Value

Nothing

## Author(s)

Marc Girondot <marc.girondot@gmail.com>

## Examples

```
plot(0:1, 0:1, xlim=c(0, 1), ylim=c(0,1), lty=2, type="l", las=1, bty="n",
     xlab="Variable x", ylab="variable y")

ellipse(center.x = c(0.2, 0.3, 0.25), center.y = c(0.7, 0.6, 0.55),
        radius.x = c(0.1, 0.1, 0.1), radius.y = c(0.15, 0.2, 0.4),
        border=NA, col=rgb(red = 0.1, green = 0.1, blue = 0.1, alpha = 0.1))

ellipse(center.x = 0.5, center.y = 0.5,
        radius.x.lower = 0.1, radius.x.upper = 0.3,
        radius.y = 0.2,
        border=NA, col=rgb(red = 0.1, green = 0.1, blue = 0.1, alpha = 0.1))

ellipse(center.x = 0.6, center.y = 0.3,
        radius.x.lower = 0.3, radius.x.upper = 0.3,
        radius.y.lower = 0.2, radius.y.upper = 0.4,
        border=NA, col=rgb(red = 0.1, green = 0.1, blue = 0.1, alpha = 0.1))

plot(0:1, 0:1, xlim=c(0, 1), ylim=c(0,1), lty=2, type="l", bty="n", asp=1,
     xlab="Variable x", ylab="variable y", axes=FALSE)
axis(1, at=c(0, 0.25, 0.5, 0.75, 1))
axis(2, at=c(0, 0.25, 0.5, 0.75, 1), las=1)

ellipse(center.x = 0.5, center.y = 0.5, radius.x = 0.2, radius.y = 0.4,
        border=NA, col=rgb(red = 0.1, green = 0.1, blue = 0.1, alpha = 0.1))
ellipse(center.x = 0.5, center.y = 0.5, radius.x = 0.2, radius.y = 0.4,
```

```

border=NA, col=rgb(red = 0.1, green = 0.1, blue = 0.1, alpha = 0.1), alpha = pi/4)

plot(0:1, 0:1, xlim=c(0, 1), ylim=c(0,1), lty=2, type="l", las=1, bty="n",
     xlab="Variable x", ylab="variable y")

for (k in 0:8)
  ellipse(center.x=0.5, center.y=0.5, radius.x=0.1, radius.y=0.4,
          alpha=seq(from=0, to=pi/4, length=9)[k],
          border=rainbow(9)[k])

# Exemple with confidence of proportions
males <- c(10, 25, 3, 4)
N <- c(12, 52, 17, 10)

males2 <- c(12, 20, 3, 6)
N2 <- c(15, 50, 20, 12)

plot(0:1, 0:1, xlim=c(0, 1), ylim=c(0,1), lty=2, type="l", las=1, bty="n",
     xlab="Variable x", ylab="variable y")

ellipse(binconf.x = data.frame(x=males, n=N), binconf.y = data.frame(x=males2, n=N2),
        border=NA, col=rgb(red = 0.1, green = 0.5, blue = 0.1, alpha = 0.1))

plot(0:1, 0:1, xlim=c(0, 1), ylim=c(0,1), lty=2, type="l", las=1, bty="n",
     xlab="Variable x", ylab="variable y")

ellipse(binconf.x = data.frame(x=males, n=N),
        binconf.y = data.frame(PointEst=c(0.1, 0.2, 0.3, 0.5),
                               Lower=c(0.02, 0.12, 0.25, 0.30),
                               Upper=c(0.18, 0.29, 0.35, 0.67)),
        border=NA, col=rgb(red = 0.1, green = 0.5, blue = 0.1, alpha = 0.1))

# Examples with a gradient
plot(0:1, 0:1, xlim=c(0, 1), ylim=c(0,1), lty=2, type="l", las=1, bty="n",
      xlab="Variable x", ylab="variable y")
ellipse(center.x = 0.6, center.y = 0.3,
        radius.x.lower = 0.3, radius.x.upper = 0.3,
        radius.y.lower = 0.2, radius.y.upper = 0.4,
        border=NA, col=grey.colors(100, alpha = 0.1))

plot(0:1, 0:1, xlim=c(0, 1), ylim=c(0,1), lty=2, type="l", las=1, bty="n",
      xlab="Variable x", ylab="variable y")
ellipse(binconf.x = data.frame(x=males, n=N), binconf.y = data.frame(x=males2, n=N2),
        border=NA, col=grey.colors(100, alpha = 0.1))

```

## Description

For `glm` fits the family's `aic()` function is used to compute the AIC.

The choice between different criteria is done by setting a global option `AIC`. It can be checked using `show.option=TRUE`. Indeed, it is not possible to use the `...` parameter due to a bug in some functions of MASS package. If you want to use this function as a replacement for `setpAIC()`, do `extractAIC.glm <- ExtractAIC.glm` before.

## Usage

```
ExtractAIC.glm(fit, scale = 0, k = 2, ...)
```

## Arguments

<code>fit</code>	fitted model, the result of a fitter <code>glm</code> .
<code>scale</code>	unused for <code>glm</code> .
<code>k</code>	numeric specifying the ‘weight’ of the equivalent degrees of freedom ( <code>=: edf</code> ) part in the AIC formula.
<code>...</code>	further arguments (currently unused because <code>addterm.glm</code> and <code>dropterm.glm</code> using this function do not transmit them).

## Details

`ExtractAIC.glm` returns AIC, AICc or BIC from a `glm` object

## Value

A numeric named vector of length 2, with first and second elements giving  
`edf` the ‘equivalent degrees of freedom’ for the fitted model fit.  
`x` the Information Criterion for fit.

## Author(s)

Modified from `stats:::extract.AIC.glm`

## See Also

Other AIC: [FormatCompareAIC\(\)](#), [compare\\_AICc\(\)](#), [compare\\_AIC\(\)](#), [compare\\_BIC\(\)](#)

## Examples

```
extractAIC.glm <- ExtractAIC.glm
n <- 100
x <- rnorm(n, 20, 2)
A <- rnorm(n, 20, 5)
g <- glm(x ~ A)
extractAIC(g, show.option=TRUE)
options(AIC="AIC")
extractAIC(g)
options(AIC="BIC")
```

```
extractAIC(g)
options(AIC="AICc")
extractAIC(g)
```

**fitdistrquantiles**      *Parameters of beta, normal or gamma distribution based on quantiles.*

## Description

Return the parameters of beta or gamm that fits the best the quantiles. The vector of probabilities can be obtained from names of quantiles.

## Usage

```
fitdistrquantiles(
  quantiles = stop("At least two quantiles must be provided"),
  probs = NULL,
  scaled = FALSE,
  distribution = "beta"
)
```

## Arguments

quantiles	Vector of quantiles.
probs	Numeric vector of probabilities with values in [0,1].
scaled	Used scaled least-square.
distribution	Distribution to be fitted: beta, normal, or gamma.

## Details

fitdistrquantiles returns the parameters of beta, normal or gamma distribution

## Value

Parameters of beta, normal or gamma distribution based on quantiles.

## Author(s)

Marc Girondot <marc.girondot@gmail.com>

## Examples

```

rd <- rbeta(100000, shape1 = 0.7, shape2 = 6.2, ncp=0)
(q <- quantile(rd, probs=c(0.025, 0.5, 0.975)))

(best <- fitdistrquantiles(quantiles = q, probs = c(0.025, 0.5, 0.975),
                           scaled=FALSE, distribution = "beta"))
rd10000 <- rbeta(10000, shape1 = best["shape1"], shape2 = best["shape2"], ncp=best["ncp"])
quantile(rd10000, probs=c(0.025, 0.5, 0.975))

# Here the probabilities are obtained from names of quantiles
(best <- fitdistrquantiles(quantiles = q, scaled=FALSE, distribution = "beta"))
rd10000 <- rbeta(10000, shape1 = best["shape1"], shape2 = best["shape2"], ncp=best["ncp"])
quantile(rd10000, probs=c(0.025, 0.5, 0.975))

# If only two quantiles are provided, ncp cannot be fitted
(q2 <- quantile(rd, probs=c(0.025, 0.975)))
(best <- fitdistrquantiles(quantiles = q2, scaled=FALSE, distribution = "beta"))
rd10000 <- rbeta(10000, shape1 = best["shape1"], shape2 = best["shape2"])
quantile(rd10000, probs=c(0.025, 0.975))
x <- seq(from=0.00, to=1, by=0.001)
plot(x=x, y=pbeta(x, shape1 = best["shape1"], shape2 = best["shape2"]),
      las=1, bty="n", type="l", ylim=c(0, 1))
segments(x0=q2[1], x1=q2[1], y0=0, y1=1, lty=2)
segments(x0=q2[2], x1=q2[2], y0=0, y1=1, lty=2)

(best <- fitdistrquantiles(quantiles = q, probs = c(0.025, 0.5, 0.975),
                           scaled=FALSE, distribution = "gamma"))
rd10000 <- rgamma(10000, shape = best["shape"], scale = best["scale"])
quantile(rd10000, probs=c(0.025, 0.5, 0.975))

(best <- fitdistrquantiles(quantiles = c(10, 20, 30), probs = c(0.025, 0.5, 0.975),
                           scaled=FALSE, distribution = "normal"))
rd10000 <- rnorm(10000, mean = best["mean"], sd = best["sd"])
quantile(rd10000, probs=c(0.025, 0.5, 0.975))

```

flexit

*Return the flexit*

## Description

Return a vector with the probabilities. The flexit equation is published in:

Abreu-Grobois, F.A., Morales-Mérida, B.A., Hart, C.E., Guillón, J.-M., Godfrey, M.H., Navarro, E. & Girondot, M. (2020) Recent advances on the estimation of the thermal reaction norm for sex ratios. PeerJ, 8, e8451.

If dose < P then  $(1 + (2^K 1 - 1) * \exp(4 * S1 * (P - x)))^{(-1/K1)}$

If dose > P then  $1 - ((1 + (2^K 2 - 1) * \exp(4 * S2 * (x - P)))^{(-1/K2)})$

with:

$$S1 = (2^K 1 - 1) * S * K1 / (2^K 1 - 1)$$

$$S2 = (2^{(K2 - 1)} * S * K2) / (2^K 2 - 1)$$

New in version 4.7-3 and larger:

If  $2^K 1$  is too large to be estimated, the approximation  $S1 = S * K1/2$  is used.

Demonstration:

$$S1 = (2^{(K1 - 1)} * S * K1) / (2^K 1 - 1)$$

$$S1 = \exp(\log((2^{(K1 - 1)} * S * K1) / (2^K 1 - 1)))$$

$$S1 = \exp(\log(2^{(K1 - 1)}) + \log(S * K1) - \log(2^K 1 - 1))$$

When  $K1$  is very large,  $2^K 1 - 1 = 2^K 1$  then

$$S1 = \exp((K1 - 1) * \log(2) + \log(S * K1) - K1 * \log(2))$$

$$S1 = \exp((K1 * \log(2) - \log(2) + \log(S * K1) - K1 * \log(2))$$

$$S1 = \exp(\log(S * K1) - \log(2))$$

$$S1 = S * K1/2$$

If  $2^K 2$  is too large to be estimated, the approximation  $S2 = S * K2/2$  is used.

If  $(1 + (2^K 1 - 1) * \exp(4 * S1 * (P - x)))^{-1/K1}$  is not finite, the following approximation is used:

$$\exp((-1/K1) * (K1 * \log(2) + (4 * S1 * (P - x))))$$

If  $1 - ((1 + (2^K 2 - 1) * \exp(4 * S2 * (x - P)))^{-1/K2})$  is not finite, the following approximation is used:

$$1 - \exp((-1/K2) * (K2 * \log(2) + (4 * S2 * (x - P))))$$

## Usage

```
flexit(
  x,
  par = NULL,
  P = NULL,
  S = NULL,
  K1 = NULL,
  K2 = NULL,
  zero = 1e-09,
  error0 = 0,
  error1 = 1
)
```

**Arguments**

x	The values at which the flexit model must be calculated
par	The vector with P, S, K1, and K2 values
P	P value
S	S value
K1	K1 value
K2	K2 value
zero	Value to replace zero
error0	Value to return if an error is observed toward 0
error1	Value to return if an error is observed toward 1

**Details**

Return the flexit value

**Value**

A vector with the probabilities

**Author(s)**

Marc Girondot <marc.girondot@gmail.com>

**See Also**

Other logit: [invlogit\(\)](#), [logit\(\)](#)

**Examples**

```
n <- flexit(x=1:100, par=c(P=50, S=0.001, K1=0.01, K2=0.02))
n <- flexit(x=1:100, P=50, S=0.001, K1=0.01, K2=0.02)

1/(1+exp(0.01*4*(50-1:100)))
flexit(1:100, P=50, S=0.01, K1=1, K2=1)
```

**Description**

Format data to be used with `compare_AIC()`, `compare_AICC()` and `compare_BIC()`. Note that `logLik` is supposed to not be `-logLik`.

**Usage**

```
FormatCompareAIC(logLik, nobs, df)
```

**Arguments**

<code>logLik</code>	The log likelihood
<code>nobs</code>	Number of observations
<code>df</code>	Number of parameters

**Details**

`FormatCompareAIC` formats data to be used with `compare_AIC()`

**Value**

An object to be used with `compare_AIC()`

**Author(s)**

Marc Girondot <[marc.girondot@gmail.com](mailto:marc.girondot@gmail.com)>

**See Also**

Other AIC: [ExtractAIC.glm\(\)](#), [compare\\_AICc\(\)](#), [compare\\_AIC\(\)](#), [compare\\_BIC\(\)](#)

**Examples**

```
## Not run:
ED <- FormatCompareAIC(logLik=-140, nobs=100, df=3)
L <- FormatCompareAIC(logLik=-145, nobs=100, df=4)
compare_AIC(L=L, ED=ED)
compare_AICc(L=L, ED=ED)
compare_BIC(L=L, ED=ED)

## End(Not run)
```

**format\_ncdf** *Return an array with ncdf data*

**Description**

Return a list with two elements: data is an array and time is the POSIX.lt time.  
 Or if `label.time` is `NULL` or if `bathy` is `TRUE`, a bathy object.  
 If `varid` is `NULL`, it shows the available variable and dimensions of the file.  
 Bathymetry data can be download here:  
[https://www.gebco.net/data\\_and\\_products/gridded\\_bathymetry\\_data/#global](https://www.gebco.net/data_and_products/gridded_bathymetry_data/#global)

**Usage**

```
format_ncdf(  
  ncdf,  
  label.latitude = "latitude",  
  label.longitude = "longitude",  
  label.time = "time",  
  varid = NULL,  
  longitude1 = NA,  
  latitude1 = NA,  
  longitude2 = NA,  
  latitude2 = NA,  
  package = "ncdf4",  
  bathy = TRUE  
)
```

**Arguments**

ncdf	An object read from package ncdf4 or a file name of ncdf file
label.latitude	Label of latitude
label.longitude	Label of longitude
label.time	Label of time
varid	Name of variable to extract
longitude1	Longitude for first corner
latitude1	latitude for first corner
longitude2	Longitude for second corner
latitude2	latitude for second corner
package	If ncdf is a file, give the package to use to open the file
bathy	If TRUE, return a bathy object

**Details**

format\_ncdf is used extract information from ncdf file

**Value**

A list with two element: data is an array and time is the POSIX.lt time

**Author(s)**

Marc Girondot <marc.girondot@gmail.com>

**See Also**

Other ncdf: [ind\\_long\\_lat\(\)](#)

## Examples

```
## Not run:
url <- "https://downloads.psl.noaa.gov/Datasets/noaa.oisst.v2.highres/"
url <- paste0(url, "sst.day.mean.2012.v2.nc")
dest <- paste(Sys.getenv("HOME"), "/sst.day.mean.2012.v2.nc", sep="")
download.file(url, dest)
format_ncdf(dest)

## End(Not run)
```

**iCutter**

*Run a shiny application to fit bone section*

## Description

Run a shiny application to fit bone section

## Usage

```
iCutter()
```

## Details

BP runs a shiny application to fit bone section

## Value

Nothing

## Author(s)

Marc Girondot <marc.girondot@gmail.com>

## Examples

```
## Not run:
# Not run:
library(HelpersMG)
iCutter()

## End(Not run)
```

---

IC\_clean\_data*Clean the dataframe before to be used with IC\_threshold\_matrix*

---

**Description**

This function must be used if missing values are present in the dataset.

It ensures that all correlations and partial correlations can be calculated. The columns of the dataframe are removed one per one until all can be calculated without error. It is possible to say that one or more columns must be retained because they are of particular importance in the analysis. The use and method parameters are used by cor() function. The function uses by default a parallel computing in Unix or Mac OSX systems. If progress is TRUE and the package pbmcapply is present, a progress bar is displayed. If debug is TRUE, some informations are shown during the process. [https://fr.wikipedia.org/wiki/Iconographie\\_des\\_corrélations](https://fr.wikipedia.org/wiki/Iconographie_des_corrélations)

**Usage**

```
IC_clean_data(
  data = stop("A dataframe object is required"),
  use = c("pairwise.complete.obs", "everything", "all.obs", "complete.obs",
         "na.or.complete"),
  method = c("pearson", "kendall", "spearman"),
  variable.retain = NULL,
  test.partial.correlation = TRUE,
  progress = TRUE,
  debug = FALSE
)
```

**Arguments**

data	The data.frame to be cleaned
use	an optional character string giving a method for computing covariances in the presence of missing values. This must be (an abbreviation of) one of the strings "everything", "all.obs", "complete.obs", "na.or.complete", or "pairwise.complete.obs".
method	a character string indicating which correlation coefficient (or covariance) is to be computed. One of "pearson" (default), "kendall", or "spearman": can be abbreviated.
variable.retain	a vector with the name of columns to keep
test.partial.correlation	should the partial correlations be tested ?
progress	Show a progress bar
debug	if TRUE, information about progression of cleaning are shown

**Details**

IC\_clean\_data checks and corrects the dataframe to be used with IC\_threshold\_matrix

**Value**

A dataframe

**Author(s)**

Marc Girondot <marc.girondot@gmail.com>

**References**

Lesty, M., 1999. Une nouvelle approche dans le choix des régresseurs de la régression multiple en présence d'interactions et de colinéarités. Revue de Modulad 22, 41-77.

**See Also**

Other Iconography of correlations: [IC\\_correlation\\_simplify\(\)](#), [IC\\_threshold\\_matrix\(\)](#), [plot.IconoCorel\(\)](#)

**Examples**

```
## Not run:
library("HelpersMG")
# based on https://fr.wikipedia.org/wiki/Iconographie_des_corrélations
es <- structure(list(Student = c("e1", "e2", "e3", "e4", "e5", "e6", "e7", "e8"),
                      Mass = c(52, 59, 55, 58, 66, 62, 63, 69),
                      Age = c(12, 12.5, 13, 14.5, 15.5, 16, 17, 18),
                      Assiduity = c(12, 9, 15, 5, 11, 15, 12, 9),
                      Note = c(5, 5, 9, 5, 13.5, 18, 18, 18)),
                     row.names = c(NA, -8L), class = "data.frame")
es

df_clean <- IC_clean_data(es, debug = TRUE)
cor_matrix <- IC_threshold_matrix(data=df_clean, threshold = NULL, progress=FALSE)
cor_threshold <- IC_threshold_matrix(data=df_clean, threshold = 0.3)
plot(cor_threshold, show.legend.strength=FALSE, show.legend.direction = FALSE)
cor_threshold_Note <- IC_correlation_simplify(matrix=cor_threshold, variable="Note")
plot(cor_threshold_Note, show.legend.strength=FALSE, show.legend.direction = FALSE)

cor_threshold <- IC_threshold_matrix(data=df_clean, threshold = 0.6)
plot(cor_threshold,
      layout=matrix(data=c(53, 53, 55, 55,
                          55, 53, 55, 53), ncol=2, byrow=FALSE),
      show.legend.direction = FALSE,
      show.legend.strength = FALSE, xlim=c(-2, 2), ylim=c(-2, 2))

## End(Not run)
```

---

**IC\_correlation\_simplify**

*Simplify the correlation matrix*

---

**Description**

This function can be used to simplify the network of correlations.

If no vector of variables is given, the variables not linked to any other variable are removed. If a vector of variables is given, only link to these variables are retained. [https://fr.wikipedia.org/wiki/Iconographie\\_des\\_c](https://fr.wikipedia.org/wiki/Iconographie_des_c)

**Usage**

```
IC_correlation_simplify(matrix, variable = NULL)
```

**Arguments**

matrix	The correlation matrix to simplify
variable	a vector with the name of columns to keep

**Details**

IC\_correlation\_simplify simplifies the correlation matrix

**Value**

A list

**Author(s)**

Marc Girondot <[marc.girondot@gmail.com](mailto:marc.girondot@gmail.com)>

**References**

Lesty, M., 1999. Une nouvelle approche dans le choix des régresseurs de la régression multiple en présence d'interactions et de colinéarités. Revue de Modulad 22, 41-77.

**See Also**

Other Iconography of correlations: [IC\\_clean\\_data\(\)](#), [IC\\_threshold\\_matrix\(\)](#), [plot.IconoCorel\(\)](#)

**Examples**

```
## Not run:  
library("HelpersMG")  
es <- structure(list(Student = c("e1", "e2", "e3", "e4", "e5", "e6", "e7", "e8"),  
    Mass = c(52, 59, 55, 58, 66, 62, 63, 69),  
    Age = c(12, 12.5, 13, 14.5, 15.5, 16, 17, 18),  
    Assiduity = c(12, 9, 15, 5, 11, 15, 12, 9),
```

```

Note = c(5, 5, 9, 5, 13.5, 18, 18, 18)),
row.names = c(NA, -8L), class = "data.frame")

es

df <- IC_clean_data(es, debug = TRUE)
cor_matrix <- IC_threshold_matrix(data=df, threshold = NULL, progress=FALSE)
cor_threshold <- IC_threshold_matrix(data=df, threshold = 0.3)
par(mar=c(1,1,1,1))
set.seed(4)
plot(cor_threshold)
cor_threshold_Note <- IC_correlation_simplify(matrix=cor_threshold, variable="Note")
plot(cor_threshold_Note)

## End(Not run)

```

### **IC\_threshold\_matrix      Calculate correlation matrix**

#### **Description**

This function calculates the matrix of correlations thresholded using partial correlation.

If the threshold is not given, the object that is produced can be used later for thresholding.

For model OAT: The link between A and B is “remarkable” if and only if the total correlation between them is higher than a given threshold and if the partial correlation between A and B in respect to any other variable C is also higher in absolute values than this threshold and with the same sign as the total correlation. For model AAT: A correlation is retained if it is higher than the threshold and the partial correlation is lower than the threshold. In this case, no missing value is accepted.

The use and method parameters are used by cor() function. The function uses by default a parallel computing in Unix or Mac OSX systems. If progress is TRUE and the package pbmcapply is present, a progress bar is displayed. If debug is TRUE, some informations are shown during the process but parallel computing is not used.

[https://fr.wikipedia.org/wiki/Iconographie\\_des\\_corrélations](https://fr.wikipedia.org/wiki/Iconographie_des_corrélations)

#### **Usage**

```

IC_threshold_matrix(
  data = stop("A dataframe or an IconoCorel object is required"),
  threshold = NULL,
  use = c("pairwise.complete.obs", "everything", "all.obs", "complete.obs",
         "na.or.complete"),
  method = c("pearson", "kendall", "spearman"),
  model = c("OAT", "ATT"),
  significance.level = FALSE,
  correction.multiple.comparisons = "fdr",
  progress = TRUE,
  debug = FALSE
)

```

## Arguments

data	A datafram or an IconoCorel object from a previous run of IC_threshold_matrix
threshold	threshold for partial and full correlations
use	an optional character string giving a method for computing covariances in the presence of missing values. This must be (an abbreviation of) one of the strings "everything", "all.obs", "complete.obs", "na.or.complete", or "pairwise.complete.obs".
method	a character string indicating which correlation coefficient (or covariance) is to be computed. One of "pearson" (default), "kendall", or "spearman": can be abbreviated.
model	a character string indicating if linear model uses all variables at a time (AAT) or one at a time (OAT).
significance.level	if FALSE, does not use significance level; or use this significance level.
correction.multiple.comparisons	"holm", "hochberg", "hommel", "bonferroni", "BH", "BY", "fdr", or "none".
progress	show a progress bar
debug	display information about progression of computing

## Details

IC\_threshold\_matrix calculates correlation matrix thresholded by partial correlation

## Value

A list

## Author(s)

Marc Girondot <[marc.girondot@gmail.com](mailto:marc.girondot@gmail.com)>

## References

Lesty, M., 1999. Une nouvelle approche dans le choix des régresseurs de la régression multiple en présence d'interactions et de colinéarités. Revue de Modulad 22, 41-77.

## See Also

Other Iconography of correlations: [IC\\_clean\\_data\(\)](#), [IC\\_correlation\\_simplify\(\)](#), [plot.IconoCorel\(\)](#)

## Examples

```
## Not run:
library("HelpersMG")
es <- structure(list(Student = c("e1", "e2", "e3", "e4", "e5", "e6", "e7", "e8"),
                      Mass = c(52, 59, 55, 58, 66, 62, 63, 69),
                      Age = c(12, 12.5, 13, 14.5, 15.5, 16, 17, 18),
                      Assiduity = c(12, 9, 15, 5, 11, 15, 12, 9),
                      Note = c(5, 5, 9, 5, 13.5, 18, 18, 18)),
```

```

row.names = c(NA, -8L), class = "data.frame")

es

df_clean <- IC_clean_data(es, debug = TRUE)
cor_matrix <- IC_threshold_matrix(data=df_clean, threshold = NULL, progress=FALSE)
cor_threshold <- IC_threshold_matrix(data=df_clean, threshold = 0.3)
plot(cor_threshold, show.legend.strength=FALSE, show.legend.direction = FALSE)
cor_threshold_Note <- IC_correlation_simplify(matrix=cor_threshold, variable="Note")
plot(cor_threshold_Note)

cor_threshold <- IC_threshold_matrix(data=df_clean, threshold = 0.8, progress=FALSE)
gr <- plot(cor_threshold, plot=FALSE)
ly <- getFromNamespace("layout_nicely", ns="igraph")(gr)
plot(cor_threshold,
      layout=matrix(data=c(53, 53, 55, 55,
                         55, 53, 55, 53), ncol=2, byrow=FALSE),
      show.legend.direction = FALSE,
      show.legend.strength = FALSE, xlim=c(-2, 2), ylim=c(-2, 2))

# Using significance level

cor_threshold <- IC_threshold_matrix(data=df_clean, threshold = 0.3,
                                       significance.level=0.05)
plot(cor_threshold, show.legend.strength=FALSE, show.legend.direction = FALSE)
cor_threshold_Note <- IC_correlation_simplify(matrix=cor_threshold, variable="Note")
plot(cor_threshold_Note)

# Using the model All at a time

cor_threshold_AAT <- IC_threshold_matrix(data=df_clean, threshold = 0.3, model="AAT")
par(mar=c(1,1,1,1))
set.seed(4)
plot(cor_threshold_AAT, show.legend.strength="bottomleft")

#####
dta <- structure(list(Student = c("e1", "e2", "e3", "e4", "e5", "e6", "e7", "e8"),
                        Mass = c(52, 59, 55, 58, 66, 62, 63, 69),
                        Age = c(12, 12.5, 13, 14.5, 15.5, 16, 17, 18),
                        Assiduity = c(12, 9, 15, 5, 11, 15, 12, 9),
                        Note = c(5, 5, 9, 5, 13.5, 18, 18, 18)),
                        row.names = c(NA, -8L), class = "data.frame")

dta0 <- dta[, 2:ncol(dta)]
ic0 <- IC_threshold_matrix(data = dta0)
cor_threshold <- IC_threshold_matrix(data=ic0, threshold = 0.3)
par(mar=c(1,1,1,1))
set.seed(4)
library("igraph")

plot(cor_threshold, vertex.color="red", show.legend.strength = FALSE)

```

```
plot(IC_correlation_simplify(matrix=cor_threshold),  
     show.legend.strength = FALSE, show.legend.direction = FALSE)  
  
## End(Not run)
```

---

index.periodic	<i>Estimate indices in periodic timeseries based on anchored minimum and maximum</i>
----------------	--

---

## Description

Estimate indices in periodic timeseries based on anchored minimum and maximum. The data.frame minmax can be generated manually. It should have three columns (time, index, SD), with all the successive minimum and maximum indices. It can be used with sun.info() to get the time of minimum and maximum air temperature or with getTide() to reconstruct the sea level.

## Usage

```
index.periodic(minmax, time = NULL, replicates = 100, progressbar = FALSE)
```

## Arguments

minmax	A data.frame returned by minmax.periodic
time	The time at which produced the estimate
replicates	Number of replicates to estimate SD
progressbar	Does a progression bar must be shown

## Details

index.periodic estimate indices in periodic timeseries based on anchored minimum and maximum

## Value

A data.frame with a column time and a column index

## Author(s)

Marc Girondot <marc.girondot@gmail.com>

## See Also

Other Periodic patterns of indices: [minmax.periodic\(\)](#), [moon.info\(\)](#), [sun.info\(\)](#), [tide.info\(\)](#)

## Examples

```

## Not run:
# Generate a timeserie of time
time.obs <- NULL
for (i in 0:9) time.obs <- c(time.obs, c(0, 6, 12, 18)+i*24)
# For these time, generate a timeseries of temperatures
temp.obs <- rep(NA, length(time.obs))
temp.obs[3+(0:9)*4] <- rnorm(10, 25, 3)
temp.obs[1+(0:9)*4] <- rnorm(10, 10, 3)
for (i in 1:(length(time.obs)-1))
  if (is.na(temp.obs[i]))
    temp.obs[i] <- mean(c(temp.obs[i-1], temp.obs[i+1]))
  if (is.na(temp.obs[length(time.obs)]))
    temp.obs[length(time.obs)] <- temp.obs[length(time.obs)-1]/2
observed <- data.frame(time=time.obs, temperature=temp.obs)
# Search for the minimum and maximum values
r <- minmax.periodic(time.minmax.daily=c(Min=2, Max=15),
observed=observed, period=24, colname.index="temperature")

# Estimate all the temperatures for these values
t <- index.periodic(minmax=r)

plot_errbar(x=t[, "time"], y=t[, "index"],
errbar.y=ifelse(is.na(t[, "sd"]), 0, 2*t[, "sd"]),
type="l", las=1, bty="n", errbar.y.polygon = TRUE,
xlab="hours", ylab="Temperatures", ylim=c(0, 35),
errbar.y.polygon.list = list(col="grey"))

plot_add(x=t[, "time"], y=t[, "index"], type="l")
plot_add(observed$time, observed$temperature, pch=19, cex=0.5)

## End(Not run)

```

*ind\_long\_lat*

*Return or the index in ncdf object from lat/longitude or inverse*

## Description

Return or the index in ncdf object from lat/longitude or reverse.

## Usage

```

ind_long_lat(
  ncdf = stop("The ncdf data must be supplied"),
  long = NULL,
  lat = NULL,
  indice.long = NULL,
  indice.lat = NULL,

```

```

    label.longitude = "lon",
    label.latitude = "lat"
)

```

## Arguments

ncdf	An object read from package ncdf4, ncdf or RNetCDF
long	Longitude in decimal format
lat	Latitude in decimal format
indice.long	Index of longitude
indice.lat	Index of latitude
label.longitude	Name of argument for longitude, default is lon
label.latitude	Name of argument for latitude, default is lat

## Details

ind\_long\_lat is used to manage ncdf information

## Value

Or the index in ncdf object from lat/longitude or inverse

## Author(s)

Marc Girondot <marc.girondot@gmail.com>

## See Also

Other ncdf: [format\\_ncdf\(\)](#)

## Examples

```

## Not run:
url <- "https://downloads.psl.noaa.gov/Datasets/noaa.oisst.v2.highres/"
url <- paste0(url, "sst.day.mean.2012.v2.nc")
dest <- paste(Sys.getenv("HOME"), "/sst.day.mean.2012.v2.nc", sep="")
download.file(url, dest)
library("ncdf4")
dta2012 <- nc_open(dest)
indices <- ind_long_lat(ncdf=dta2012, lat=5.89, long=-20.56)
coordinates <- ind_long_lat(ncdf=dta2012, indice.lat=20, indice.long=30)
# library("RNetCDF")
# dta2012 <- open.nc(dest)
# indices <- ind_long_lat(ncdf=dta2012, lat=5.89, long=-20.56)
# coordinates <- ind_long_lat(ncdf=dta2012, indice.lat=20, indice.long=30)
# ncdf library is depreciated in CRAN
# library("ncdf")
# dta2012 <- open.ncdf(dest)
# indices <- ind_long_lat(ncdf=dta2012, lat=5.89, long=-20.56)

```

```
# coordinates <- ind_long_lat(ncdf=dta2012, indice.lat=20, indice.long=30)
## End(Not run)
```

**inside***Search a string within files of a folder***Description**

Search for a string inside the files of a folder and return where the string is found.  
The pattern for files that must be included uses regex for filtering.

**Usage**

```
inside(
  text = stop("A text to be searched for is necessary"),
  path = ".",
  pattern = "*\\\\.R$",
  showallfilenames = FALSE,
  ...,
  fixed = TRUE,
  ignore.case = FALSE
)
```

**Arguments**

<code>text</code>	Text to search in files
<code>path</code>	Path of the folder to search in
<code>pattern</code>	Pattern for file names to search in
<code>showallfilenames</code>	logical. Show all the filenames search for in
<code>...</code>	Options for readLines(), example warn = FALSE
<code>fixed</code>	logical. If TRUE, pattern is a string to be matched as is. Overrides all conflicting arguments (see gsub)
<code>ignore.case</code>	logical. if FALSE, the pattern matching for text is case sensitive and if TRUE, case is ignored during matching.

**Details**

`inside` Search a string within files of a folder

**Value**

Return an invisible vector with filenames in which the pattern occurs

**Author(s)**

Marc Girondot <marc.girondot@gmail.com>

**Examples**

```
## Not run:  
library(HelpersMG)  
# Search for files in path with names based on pattern that have the string search inside.  
inside("embryogrowth", path=". ", pattern="*\\".R$")  
  
## End(Not run)
```

---

invlogit

*Return the inverse logit*

---

**Description**

Return the inverse logit.

**Usage**

```
invlogit(n)
```

**Arguments**

n               The value to inverse to get the probability

**Details**

invlogit returns the inverse logit

**Value**

A value

**Author(s)**

Marc Girondot <marc.girondot@gmail.com>

**See Also**

Other logit: [flexit\(\)](#), [logit\(\)](#)

**Examples**

```
n <- logit(0.5)  
invlogit(n)
```

---

LD50*Estimate the parameters that best describe LD50*

---

**Description**

Estimate the parameters that best describe LD50

Logistic and logit models are the same but with different parametrization:

logistic =  $1/(1+\exp((1/S)*(P-d)))$

logit =  $1/(1+\exp(P+d*S))$

See these publications for the description of equations:

Girondot, M. 1999. Statistical description of temperature-dependent sex determination using maximum likelihood. Evolutionary Ecology Research, 1, 479-486.

Godfrey, M.H., Delmas, V., Girondot, M., 2003. Assessment of patterns of temperature-dependent sex determination using maximum likelihood model selection. Ecoscience 10, 265-272.

Hulin, V., Delmas, V., Girondot, M., Godfrey, M.H., Guillon, J.-M., 2009. Temperature-dependent sex determination and global change: are some species at greater risk? Oecologia 160, 493-506.

The flexit equation is not still published :

$$\text{if } dose < P \text{ then } (1 + (2^K 1 - 1) * \exp(4 * S1 * (P - x)))^{(1/K1)} - 1/K1 \\ \text{if } dose > P \text{ then } 1 - ((1 + (2^K 2 - 1) * \exp(4 * S2 * (x - P)))^{(1/K2)} - 1/K2)$$

with:

$$S1 = S / ((4/K1) * (2^{(1/K1)} - 1) * (2^K 1 - 1)) \\ S2 = S / ((4/K2) * (2^{(1/K2)} - 1) * (2^K 2 - 1))$$

**Usage**

```
LD50(
  df = NULL,
  alive = NULL,
  dead = NULL,
  N = NULL,
  doses = NULL,
  l = 0.05,
  parameters.initial = NULL,
  fixed.parameters = NULL,
  SE = NULL,
  equation = "logistic",
  replicates = 1000,
  range.CI = 0.95,
  limit.low.TRD.minimum = 5,
  limit.high.TRD.maximum = 1000,
  print = TRUE,
  doses.plot = seq(from = 0, to = 1000, by = 0.1)
)
```

## Arguments

df	A dataframe with at least two columns named alive, dead or N and doses columns
alive	A vector with alive individuals at the end of experiment
dead	A vector with dead individuals at the end of experiment
N	A vector with total numbers of tested individuals
doses	The doses
l	The limit to define TRD (see Girondot, 1999)
parameters.initial	Initial values for P, S or K search as a vector, ex. c(P=29, S=-0.3)
fixed.parameters	Parameters that will not be changed during fit
SE	Standard errors for parameters
equation	Could be "logistic", "logit", "probit", "Hill", "Richards", "Hulin", "flexit" or "Double-Richards"
replicates	Number of replicates to estimate confidence intervals
range.CI	The range of confidence interval for estimation, default=0.95
limit.low.TRD.minimum	Minimum lower limit for TRD
limit.high.TRD.maximum	Maximum higher limit for TRD
print	Do the results must be printed at screen? TRUE (default) or FALSE
doses.plot	Sequences of doses that will be used for plotting. If NULL, does not estimate them

## Details

LD50 estimates the parameters that best describe LD50

## Value

A list with the LD50, Transitional Range of Doses and their SE

## Author(s)

Marc Girondot <marc.girondot@gmail.com>

## See Also

Other LD50 functions: [LD50\\_MHmcmc\\_p\(\)](#), [LD50\\_MHmcmc\(\)](#), [logLik.LD50\(\)](#), [plot.LD50\(\)](#), [predict.LD50\(\)](#)

## Examples

```

## Not run:
library("HelpersMG")
data <- data.frame(Doses=c(80, 120, 150, 150, 180, 200),
Alive=c(10, 12, 8, 6, 2, 1),
Dead=c(0, 1, 5, 6, 9, 15))
LD50_logistic <- LD50(data, equation="logistic")
predict(LD50_logistic, doses=c(140, 170))
plot(LD50_logistic, xlim=c(0, 300), at=seq(from=0, to=300, by=50))
LD50_probit <- LD50(data, equation="probit")
predict(LD50_probit, doses=c(140, 170))
plot(LD50_probit)
LD50_logit <- LD50(data, equation="logit")
predict(LD50_logit, doses=c(140, 170))
plot(LD50_logit)
LD50_hill <- LD50(data, equation="hill")
predict(LD50_hill, doses=c(140, 170))
plot(LD50_hill)
LD50_Richards <- LD50(data, equation="Richards")
predict(LD50_Richards, doses=c(140, 170))
plot(LD50_Richards)
LD50_Hulin <- LD50(data, equation="Hulin")
predict(LD50_Hulin, doses=c(140, 170))
plot(LD50_Hulin)
LD50_DoubleRichards <- LD50(data, equation="Double-Richards")
predict(LD50_DoubleRichards, doses=c(140, 170))
plot(LD50_DoubleRichards)
LD50_flexit <- LD50(data, equation="flexit")
predict(LD50_flexit, doses=c(140, 170))
plot(LD50_flexit)

## End(Not run)

```

## Description

Run the Metropolis-Hastings algorithm for tsd.

Deeply modified from a MCMC script by Olivier Martin (INRA, Paris-Grignon).

The number of iterations is n.iter+n.adapt+1 because the initial likelihood is also displayed.

I recommend that thin=1 because the method to estimate SE uses resampling.

If initial point is maximum likelihood, n.adapt = 0 is a good solution.

To get the SE from result\_mcmc <- tsd\_MHmcmc(result=try), use:

result\_mcmc\$BatchSE or result\_mcmc\$TimeSeriesSE

The batch standard error procedure is usually thought to be not as accurate as the time series methods.

Based on Jones, Haran, Caffo and Neath (2005), the batch size should be equal to sqrt(n.iter).

Jones, G.L., Haran, M., Caffo, B.S. and Neath, R. (2006) Fixed Width Output Analysis for Markov

chain Monte Carlo , Journal of the American Statistical Association, 101:1537-1547.  
 coda package is necessary for this function.  
 The parameters intermediate and filename are used to save intermediate results every 'intermediate' iterations (for example 1000). Results are saved in a file of name filename.  
 The parameter previous is used to indicate the list that has been save using the parameters intermediate and filename. It permits to continue a mcmc search.  
 These options are used to prevent the consequences of computer crash or if the run is very very long and processes at time limited.

## Usage

```
LD50_MHmcmc(
  result = stop("A result of LD50() fit must be provided"),
  n.iter = 10000,
  parametersMCMC = NULL,
  n.chains = 1,
  n.adapt = 0,
  thin = 1,
  trace = FALSE,
  batchSize = sqrt(n.iter),
  adaptive = FALSE,
  adaptive.lag = 500,
  adaptive.fun = function(x) {
    ifelse(x > 0.234, 1.3, 0.7)
  },
  intermediate = NULL,
  filename = "intermediate.Rdata",
  previous = NULL
)
```

## Arguments

result	An object obtained after a SearchR fit
n.iter	Number of iterations for each step
parametersMCMC	A set of parameters used as initial point for searching with information on priors
n.chains	Number of replicates
n.adapt	Number of iterations before to store outputs
thin	Number of iterations between each stored output
trace	True or False, shows progress
batchSize	Number of observations to include in each batch fo SE estimation
adaptive	Should an adaptive process for SDProp be used
adaptive.lag	Lag to analyze the SDProp value in an adaptive content
adaptive.fun	Function used to change the SDProp
intermediate	Period for saving intermediate result, NULL for no save

filename	If intermediate is not NULL, save intermediate result in this file
previous	Previous result to be continued. Can be the filename in which intermediate results are saved.

## Details

LD50\_MHmcmc runs the Metropolis-Hastings algorithm for LD50 (Bayesian MCMC)

## Value

A list with resultMCMC being mcmc.list object, resultLnL being likelihoods and parametersMCMC being the parameters used

## Author(s)

Marc Girondot <marc.girondot@gmail.com>

## See Also

Other LD50 functions: [LD50\\_MHmcmc\\_p\(\)](#), [LD50\(\)](#), [logLik.LD50\(\)](#), [plot.LD50\(\)](#), [predict.LD50\(\)](#)

## Examples

```
## Not run:
library("HelpersMG")
data <- data.frame(Doses=c(80, 120, 150, 150, 180, 200),
Alive=c(10, 12, 8, 6, 2, 1),
Dead=c(0, 1, 5, 6, 9, 15))
LD50_logistic <- LD50(data, equation="logistic")
pMCMC <- LD50_MHmcmc_p(LD50_logistic, accept=TRUE)
# Take care, it can be very long
result_mcmc_LD50 <- LD50_MHmcmc(result=LD50_logistic,
parametersMCMC=pMCMC, n.iter=10000, n.chains = 1,
n.adapt = 0, thin=1, trace=1000, adaptive=TRUE, )
# summary() permits to get rapidly the standard errors for parameters
summary(result_mcmc_LD50)
plot(x=result_mcmc_LD50, parameters="S", scale.prior=TRUE, las=1)
plot(result_mcmc_LD50, parameters="S", scale.prior=TRUE, las=1, xlim=c(-20, 20))
plot(result_mcmc_LD50, parameters="P", scale.prior=TRUE, las=1)
1-rejectionRate(as.mcmc(result_mcmc_LD50))
raftery.diag(as.mcmc(result_mcmc_LD50))
heidel.diag(as.mcmc(result_mcmc_LD50))

#### Example with Uniforms priors

pMCMC <- structure(list(Density = c("dunif", "dunif"),
Prior1 = c(77.6216005852911, -31.0438095277258),
Prior2 = c(310.486402341165, 31.0438095277258),
SDProp = c(2, 0.5),
Min = c(77.6216005852911, -31.0438095277258),
Max = c(310.486402341165, 31.0438095277258),
Init = c(155.243201170582, -15.5219047638629)),
```

```

row.names = c("P", "S"), class = "data.frame")
result_mcmc_LD50 <- LD50_MHmcmc(result=LD50_logistic,
parametersMCMC=pMCMC, n.iter=10000, n.chains = 1,
n.adapt = 0, thin=1, trace=1000, adaptive=TRUE, )
# summary() permits to get rapidly the standard errors for parameters
summary(result_mcmc_LD50)
plot(x=result_mcmc_LD50, parameters="S", scale.prior=TRUE, las=1)
plot(result_mcmc_LD50, parameters="S", scale.prior=TRUE, las=1, xlim=c(-40, 40))
plot(result_mcmc_LD50, parameters="P", scale.prior=TRUE, las=1)
1-rejectionRate(as.mcmc(result_mcmc_LD50))
raftery.diag(as.mcmc(result_mcmc_LD50))
heidel.diag(as.mcmc(result_mcmc_LD50))

## End(Not run)

```

LD50\_MHmcmc\_p

*Generates set of parameters to be used with LD50\_MHmcmc()*

## Description

Interactive script used to generate set of parameters to be used with LD50\_MHmcmc().

## Usage

```

LD50_MHmcmc_p(
  result = stop("An output from LD50() must be provided"),
  accept = FALSE
)

```

## Arguments

- |        |  |
|--------|--|
| result | An object obtained after a LD50 fit                |
| accept | If TRUE, the script does not wait user information |

## Details

LD50\_MHmcmc\_p generates set of parameters to be used with LD50\_MHmcmc()

## Value

A matrix with the parameters

## Author(s)

Marc Girondot <marc.girondot@gmail.com>

## See Also

Other LD50 functions: [LD50\\_MHmcmc\(\)](#), [LD50\(\)](#), [logLik.LD50\(\)](#), [plot.LD50\(\)](#), [predict.LD50\(\)](#)

## Examples

```
## Not run:
library("HelpersMG")
data <- data.frame(Doses=c(80, 120, 150, 150, 180, 200),
Alive=c(10, 12, 8, 6, 2, 1),
Dead=c(0, 1, 5, 6, 9, 15))
LD50_logistic <- LD50(data, equation="logistic")
pmcmc <- LD50_MHmcmc_p(LD50_logistic, accept=TRUE)

## End(Not run)
```

**list.packages**

*List the installed packages with their locations*

## Description

List the installed packages with their locations and version.

## Usage

```
list.packages()
```

## Details

`list.packages` lists the installed packages with their locations

## Value

A list with the installed packages and their version.

## Author(s)

Marc Girondot

## Examples

```
## Not run:
library(HelpersMG)
list.packages()

## End(Not run)
```

---

**local.search***Return path of file searched for in local disk based on its file name*

---

## Description

Return path of file searched for in local disk based on its file name.

It has been tested only with Windows XP and MacOSX. In MacOSX, you must have created the locate database first. Use OnyX utilities for this purpose.

## Usage

```
local.search(  
  pattern,  
  directory = "",  
  folder = "$HOME",  
  intern = TRUE,  
  ignore.stdout = FALSE,  
  ignore.stderr = TRUE  
)
```

## Arguments

pattern	The name of file to be searched for. Can use wildcards *
directory	The path of directory to be explored in for Windows
folder	The path of folder to be explored in for Unix based systems
intern	A logical (not NA) which indicates whether to capture the output of the command as an R character vector (see system()).
ignore.stdout	a logical (not NA) indicating whether messages written to 'stdout' should be ignored (see system()).
ignore.stderr	a logical (not NA) indicating whether messages written to 'stderr' should be ignored (see system()).

## Details

local.search() returns path of file serached in local disk based on its file name

## Value

A vector with paths

## Author(s)

Marc Girondot

**Examples**

```
## Not run:  
RnwFiles <- local.search("*.Rnw")  
nc.files <- local.search("*.nc", folder=paste0("'",getwd(),"'))  
  
## End(Not run)
```

---

logit

*Return the logit*

---

**Description**

Return the logit.

**Usage**

```
logit(p)
```

**Arguments**

p                   The probability

**Details**

logit returns the logit

**Value**

A value

**Author(s)**

Marc Girondot <marc.girondot@gmail.com>

**See Also**

Other logit: [flexit\(\)](#), [invlogit\(\)](#)

**Examples**

```
n <- logit(0.5)  
invlogit(n)
```

---

logLik.compareAIC      *Return Log Likelihood generated by FormatCompareAIC*

---

## Description

Return Log Likelihood generated by FormatCompareAIC

## Usage

```
## S3 method for class 'compareAIC'  
logLik(object, ...)
```

## Arguments

object	A result generated by FormatCompareAIC
...	Not used

## Details

logLik.compareAIC Return Log Likelihood of a fit

## Value

The Log Likelihood value for the fitted model with data

## Author(s)

Marc Girondot <marc.girondot@gmail.com>

## Examples

```
## Not run:  
ED <- FormatCompareAIC(logLik=-140, nobs=100, df=3)  
logLik(ED)  
  
## End(Not run)
```

**logLik.cutter**      *Return log likelihood of a cutter fitted model*

## Description

Return log likelihood of a cutter fitted model.

## Usage

```
## S3 method for class 'cutter'
logLik(object, ...)
```

## Arguments

object	A result file generated by cutter
...	Not used

## Details

`logLik.cutter` return log likelihood of a cutter fitted model

## Value

Nothing

## Author(s)

Marc Girondot <[marc.girondot@gmail.com](mailto:marc.girondot@gmail.com)>

## See Also

Other Distributions: [cutter\(\)](#), [dbeta\\_new\(\)](#), [dcutter\(\)](#), [dggamma\(\)](#), [pSnbino\(\)](#), [plot.cutter\(\)](#), [print.cutter\(\)](#), [qSnbino\(\)](#), [r2norm\(\)](#), [rcutter\(\)](#), [rmnrm\(\)](#), [rnbinom\\_new\(\)](#)

## Examples

```
## Not run:
#
# -----
# Test for similarity in gamma left censored distribution between two
# datasets
#
# -----
obc1 <- rgamma(100, scale=20, shape=2)
# Detection limit for sample 1 to 50
LDL <- 10
# remove the data below the detection limit
obc1[obc1<LDL] <- -Inf
obc2 <- rgamma(100, scale=10, shape=2)
# remove the data below the detection limit
obc2[obc2<LDL] <- -Inf
```

```

# search for the parameters the best fit these censored data
result1 <- cutter(observations=obc1,
                   lower_detection_limit=LDL,
                   cut_method="censored")
logLik(result1)
result2 <- cutter(observations=obc2,
                   lower_detection_limit=LDL,
                   cut_method="censored")
logLik(result2)
result_totl <- cutter(observations=c(obc1, obc2),
                      lower_detection_limit=LDL,
                      cut_method="censored")
logLik(result_totl)
compare_AICc(Separate=list(result1, result2),
              Common=result_totl, factor.value=1)
compare_BIC(Separate=list(result1, result2),
             Common=result_totl, factor.value=1)

## End(Not run)

```

**logLik.LD50***Return Log Likelihood of a fit generated by LD50***Description**

Return Log Likelihood of a fit generated by LD50

**Usage**

```
## S3 method for class 'LD50'
logLik(object, ...)
```

**Arguments**

object	A result file generated by fitRMU
...	Not used

**Details**

**logLik.LD50** Return Log Likelihood of a fit for LD50

**Value**

The Log Likelihood value for the fitted model with data

**Author(s)**

Marc Girondot <marc.girondot@gmail.com>

**See Also**

Other LD50 functions: [LD50\\_MHmcmc\\_p\(\)](#), [LD50\\_MHmcmc\(\)](#), [LD50\(\)](#), [plot.LD50\(\)](#), [predict.LD50\(\)](#)

**Examples**

```
## Not run:
data <- data.frame(Doses=c(80, 120, 150, 150, 180, 200),
Alive=c(10, 12, 8, 6, 2, 1),
Dead=c(0, 1, 5, 6, 9, 15))
LD50_logistic <- LD50(data, equation="logistic")
logLik(LD50_logistic)
AIC(LD50_logistic)

## End(Not run)
```

**merge.mcmcComposite**    *Merge two mcmcComposite results*

**Description**

Merge two mcmcComposite results and produced a new one mcmcComposite object.  
Note that the initial value for the second run must use the last value of the first one as shown in example.

**Usage**

```
## S3 method for class 'mcmcComposite'
merge(x, y, ...)
```

**Arguments**

x	A mcmcComposite obtained as a result of <code>MHalgoGen()</code> function
y	A mcmcComposite obtained as a result of <code>MHalgoGen()</code> function
...	not used

**Details**

`merge.mcmcComposite` Merge two mcmcComposite results

**Value**

A mcmcComposite result

**Author(s)**

Marc Girondot <[marc.girondot@gmail.com](mailto:marc.girondot@gmail.com)>

**See Also**

Other mcmcComposite functions: `MHalgoGen()`, `as.mcmc.mcmcComposite()`, `as.parameters()`, `as.quantiles()`, `plot.PriorsmcmcComposite()`, `plot.mcmcComposite()`, `setPriors()`, `summary.mcmcComposite()`

**Examples**

```
## Not run:
library(HelpersMG)
require(coda)
x <- rnorm(30, 10, 2)
dnormx <- function(data, x) {
  data <- unlist(data)
  return(-sum(dnorm(data, mean=x['mean'], sd=x['sd'], log=TRUE)))
}
parameters_mcmc <- data.frame(Density=c('dnorm', 'dlnorm'),
Prior1=c(10, 0.5), Prior2=c(2, 0.5), SDProp=c(1, 1),
Min=c(-3, 0), Max=c(100, 10), Init=c(10, 2), stringsAsFactors = FALSE,
row.names=c('mean', 'sd'))
mcmc_run <- MHalgoGen(n.iter=1000, parameters=parameters_mcmc, data=x,
likelihood=dnormx, n.chains=1, n.adapt=100, thin=1, trace=1)
plot(mcmc_run, xlim=c(0, 20))
plot(mcmc_run, xlim=c(0, 10), parameters="sd")
mcmcforcoda <- as.mcmc(mcmc_run)
#' heidel.diag(mcmcforcoda)
raftery.diag(mcmcforcoda)
autocorr.diag(mcmcforcoda)
acf(mcmcforcoda[[1]][,"mean"], lag.max=20, bty="n", las=1)
acf(mcmcforcoda[[1]][,"sd"], lag.max=20, bty="n", las=1)
batchSE(mcmcforcoda, batchSize=100)
# The batch standard error procedure is usually thought to
# be not as accurate as the time series methods used in summary
summary(mcmcforcoda)$statistics[,"Time-series SE"]
summary(mcmc_run)
as.parameters(mcmc_run)
lastp <- as.parameters(mcmc_run, index="last")
parameters_mcmc[,"Init"] <- lastp
# The n.adapt set to 1 is used to not record the first set of parameters
# then it is not duplicated (as it is also the last one for
# the object mcmc_run)
mcmc_run2 <- MHalgoGen(n.iter=1000, parameters=parameters_mcmc, data=x,
likelihood=dnormx, n.chains=1, n.adapt=1, thin=1, trace=1)
mcmc_run3 <- merge(mcmc_run, mcmc_run2)
##### no adaptation, n.adapt must be 0
parameters_mcmc[,"Init"] <- c(mean(x), sd(x))
mcmc_run3 <- MHalgoGen(n.iter=1000, parameters=parameters_mcmc, data=x,
likelihood=dnormx, n.chains=1, n.adapt=0, thin=1, trace=1)

## End(Not run)
```

## Description

The parameters must be stored in a data.frame with named rows for each parameter with the following columns:

- Density. The density function name, example `dnorm`, `dlnorm`, `dunif`, `dbeta`
- Prior1. The first parameter to send to the Density function
- Prior2. The second parameter to send to the Density function
- SDProp. The standard error from new proposition value of this parameter
- Min. The minimum value for this parameter
- Max. The maximum value for this parameter
- Init. The initial value for this parameter

This script has been deeply modified from a MCMC script provided by Olivier Martin (INRA, Paris-Grignon).

The likelihood function must use a parameter named `parameters_name` for the nammed parameters. For adaptive mcmc, see:

Rosenthal, J. S. 2011. Optimal Proposal Distributions and Adaptive MCMC. Pages 93-112 in S. Brooks, A. Gelman, G. Jones, and X.-L. Meng, editors. MCMC Handbook. Chapman and Hall/CRC.

## Usage

```
MHalgoGen(
  likelihood = stop("A likelihood function must be supplied"),
  parameters_name = "x",
  parameters = stop("Priors must be supplied"),
  ...,
  n.iter = 10000,
  n.chains = 1,
  n.adapt = 100,
  thin = 30,
  trace = FALSE,
  traceML = FALSE,
  progress.bar.ini = NULL,
  progress.bar = NULL,
  adaptive = FALSE,
  adaptive.lag = 500,
  adaptive.fun = function(x) {
    ifelse(x > 0.234, 1.3, 0.7)
  },
  ...)
```

```

    intermediate = NULL,
    filename = "intermediate.Rdata",
    previous = NULL,
    session = NULL
)

```

## Arguments

<code>likelihood</code>	The function that returns -ln likelihood using data and parameters
<code>parameters_name</code>	The name of the parameters in the likelihood function, default is "x"
<code>parameters</code>	A data.frame with priors; see description and examples
<code>...</code>	Parameters to be transmitted to likelihood function
<code>n.iter</code>	Number of iterations for each chain
<code>n.chains</code>	Number of chains
<code>n.adapt</code>	Number of iteration to stabilize likelihood
<code>thin</code>	Interval for thinning likelihoods
<code>trace</code>	Or FALSE or period to show progress
<code>traceML</code>	TRUE or FALSE to show ML
<code>progress.bar.ini</code>	The command to initialize progress bar
<code>progress.bar</code>	The command to run the progress bar
<code>adaptive</code>	Should an adaptive process for SDProp be used
<code>adaptive.lag</code>	Lag to analyze the SDProp value in an adaptive context
<code>adaptive.fun</code>	Function used to change the SDProp
<code>intermediate</code>	Or NULL of period to save intermediate result
<code>filename</code>	Name of file in which intermediate results are saved
<code>previous</code>	The content of the file in which intermediate results are saved
<code>session</code>	The shiny session

## Details

MHalgoGen is a function to use mcmc with Metropolis-Hastings algorithm

## Value

A mcmcComposite object with all characteristics of the model and mcmc run

## Author(s)

Marc Girondot <[marc.girondot@gmail.com](mailto:marc.girondot@gmail.com)>

## See Also

Other mcmcComposite functions: [as.mcmc.mcmcComposite\(\)](#), [as.parameters\(\)](#), [as.quantiles\(\)](#), [merge.mcmcComposite\(\)](#), [plot.PriorsmcmcComposite\(\)](#), [plot.mcmcComposite\(\)](#), [setPriors\(\)](#), [summary.mcmcComposite\(\)](#)

## Examples

```
## Not run:
library(HelpersMG)
require(coda)
val <- rnorm(30, 10, 2)
dnormx <- function(data, x) {
  data <- unlist(data)
  return(-sum(dnorm(data, mean=x['mean'], sd=x['sd'], log=TRUE)))
}
parameters_mcmc <- data.frame(Density=c('dnorm', 'dlnorm'),
Prior1=c(10, 0.5), Prior2=c(2, 0.5), SDProp=c(0.35, 0.2),
Min=c(-3, 0), Max=c(100, 10), Init=c(10, 2), stringsAsFactors = FALSE,
row.names=c('mean', 'sd'))
# Use of trace and traceML parameters
# trace=1 : Only one likelihood is printed
mcmc_run <- MHalgoGen(n.iter=50000, parameters=parameters_mcmc, data=val,
likelihood=dnormx, n.chains=1, n.adapt=100, thin=1, trace=1)
# trace=10 : 10 likelihoods are printed
mcmc_run <- MHalgoGen(n.iter=50000, parameters=parameters_mcmc, data=val,
likelihood=dnormx, n.chains=1, n.adapt=100, thin=1, trace=10)
# trace=TRUE : all likelihoods are printed
mcmc_run <- MHalgoGen(n.iter=50000, parameters=parameters_mcmc, data=val,
likelihood=dnormx, n.chains=1, n.adapt=100, thin=1, trace=TRUE)
# trace=FALSE : No likelihood is printed
mcmc_run <- MHalgoGen(n.iter=50000, parameters=parameters_mcmc, data=val,
likelihood=dnormx, n.chains=1, n.adapt=100, thin=1, trace=FALSE)
# traceML=TRUE : values when likelihood is better are shown
mcmc_run <- MHalgoGen(n.iter=100, parameters=parameters_mcmc, data=val,
likelihood=dnormx, n.chains=1, n.adapt=100, thin=1, trace=TRUE, traceML=TRUE)
mcmc_run <- MHalgoGen(n.iter=100, parameters=parameters_mcmc, data=val,
likelihood=dnormx, n.chains=1, n.adapt=100, thin=1, trace=FALSE, traceML=TRUE)

plot(mcmc_run, xlim=c(0, 20))
plot(mcmc_run, xlim=c(0, 10), parameters="sd")
library(graphics)
library(fields)
# show a scatter plot of the result
x <- mcmc_run$resultMCMC[[1]][, 1]
y <- mcmc_run$resultMCMC[[1]][, 2]
marpre <- par(mar=c(4, 4, 2, 6)+0.4)
smoothScatter(x, y)
# show a scale
n <- matrix(0, ncol=128, nrow=128)
xrange <- range(x)
yrange <- range(y)
for (i in 1:length(x)) {
```

```

posx <- 1+floor(127*(x[i]-xrange[1])/(xrange[2]-xrange[1]))
posy <- 1+floor(127*(y[i]-yrange[1])/(yrange[2]-yrange[1]))
n[posx, posy] <- n[posx, posy]+1
}
image.plot(legend.only=TRUE, zlim= c(0, max(n)), nlevel=128,
  col=colorRampPalette(c("white", "blues9))(128))
# Compare with a heatmap
x <- seq(from=8, to=12, by=0.2)
y <- seq(from=1, to=4, by=0.2)
df <- expand.grid(mean=x, sd=y)
df <- cbind(df, L=rep(0, length(nrow(df))))
for (i in 1:nrow(df)) df[i, "L"] <- -sum(dnorm(val, df[i, 1], df[i, 2], log = TRUE))
hm <- matrix(df[, "L"], nrow=length(x))
par(mar = marpre)
image.plot(x=x, y=y, z=hm, las=1)
# Diagnostic function from coda library
mcmcforcoda <- as.mcmc(mcmc_run)
#' heidel.diag(mcmcforcoda)
raftery.diag(mcmcforcoda)
autocorr.diag(mcmcforcoda)
acf(mcmcforcoda[[1]][,"mean"], lag.max=20, bty="n", las=1)
acf(mcmcforcoda[[1]][,"sd"], lag.max=20, bty="n", las=1)
batchSE(mcmcforcoda, batchSize=100)
# The batch standard error procedure is usually thought to
# be not as accurate as the time series methods used in summary
summary(mcmcforcoda)$statistics[, "Time-series SE"]
summary(mcmc_run)
as.parameters(mcmc_run)
lastp <- as.parameters(mcmc_run, index="last")
parameters_mcmc[,"Init"] <- lastp
# The n.adapt set to 1 is used to not record the first set of parameters
# then it is not duplicated (as it is also the last one for
# the object mcmc_run)
mcmc_run2 <- MHalgoGen(n.iter=1000, parameters=parameters_mcmc, x=x, data=val,
likelihood=dnormx, n.chains=1, n.adapt=1, thin=1, trace=1)
mcmc_run3 <- merge(mcmc_run, mcmc_run2)
##### no adaptation, n.adapt must be 0
parameters_mcmc[,"Init"] <- c(mean(x), sd(x))
mcmc_run3 <- MHalgoGen(n.iter=1000, parameters=parameters_mcmc, x=x, data=val,
likelihood=dnormx, n.chains=1, n.adapt=0, thin=1, trace=1)
# Here is how to use adaptive mcmc
mcmc_run <- MHalgoGen(n.iter=50000, parameters=parameters_mcmc, data=val, adaptive = FALSE,
likelihood=dnormx, n.chains=1, n.adapt=100, thin=1, trace=1)
1-rejectionRate(as.mcmc(mcmc_run))
mcmc_run <- MHalgoGen(n.iter=50000, parameters=parameters_mcmc, data=val, adaptive = TRUE,
likelihood=dnormx, n.chains=1, n.adapt=100, thin=1, trace=1)
1-rejectionRate(as.mcmc(mcmc_run))
# To see the dynamics :
var <- "mean"
par(mar=c(4, 4, 1, 1)+0.4)
plot(1:nrow(mcmc_run$resultMCMC[[1]]), mcmc_run$resultMCMC[[1]][, var], type="l",
  xlab="Iterations", ylab=var, bty="n", las=1)
# Exemple with a progress bar

```

```

val <- rnorm(30, 10, 2)
dnormx <- function(data, x) {
  data <- unlist(data)
  return(-sum(dnorm(data, mean=x['mean'], sd=x['sd'], log=TRUE)))
}
parameters_mcmc <- data.frame(Density=c('dnorm', 'dlnorm'),
Prior1=c(10, 0.5), Prior2=c(2, 0.5), SDProp=c(0.35, 0.2),
Min=c(-3, 0), Max=c(100, 10), Init=c(10, 2), stringsAsFactors = FALSE,
row.names=c('mean', 'sd'))
# Set up the progress bar
mcmc_run <- MHalgoGen(n.iter=50000, parameters=parameters_mcmc, data=val,
                      likelihood=dnormx, n.chains=1, n.adapt=100, thin=1, trace=FALSE,
                      progress.bar.ini=function(n.iter) {
                        assign("pb", txtProgressBar(min=0, max=n.iter, style=3),
                           env = parent.frame())},
                      progress.bar=function(iter) {setTxtProgressBar(get("pb", envir = parent.frame()), iter)})}

## End(Not run)

```

**minmax.periodic***Search for minimum and maximum indices in periodic timeseries***Description**

Search for minimum and maximum for periodic timeseries when only intermediate values are known.

For each couple of value with an increasing or decreasing segment of the sinusoid function, it is possible to estimate a minimum and maximum values using analytical algebra.

Then the average and standard deviations of all minima and maxima are evaluated.

It should be noted that any extremum can be estimated at least twice, one by increasing segment and one by decreasing segment. Both are used here to produce SD.

`time.minmax.daily` should be used when the time at which maximum and minimum indices are regular and `time.minmax` permits to define this time day by day.

**Usage**

```

minmax.periodic(
  time.minmax.daily = NULL,
  time.minmax = NULL,
  progressbar = FALSE,
  observed = stop("data.frame with observed indices"),
  period = 24,
  colname.time = "time",
  colname.index = "index",
  colname.SD = "SD",
  plot = FALSE
)

```

## Arguments

<code>time.minmax.daily</code>	A named vector with Min and Max being the time in the day with minimum and maximum indices (temperature or level)
<code>time.minmax</code>	A named vector daily with time in the day at which minimum and maximum indices are observed
<code>progressbar</code>	Tell if a progression bar must be shown
<code>observed</code>	A datafram with at least two columns: time and temperatures. A third column SD can indicate the know error in index
<code>period</code>	The unit of day period (24 for hours, 24*60 for minutes)
<code>colname.time</code>	The name of the column for time in observed
<code>colname.index</code>	The name of the column for indices in observed
<code>colname.SD</code>	The name of the column for SD in observed
<code>plot</code>	If TRUE, show a plot with the different estimates

## Details

`minmax.periodic` search for minimum and maximum indices (temperatures or levels) in periodic timeseries

## Value

A data.frame with a column time, a column index and a column SD

## Author(s)

Marc Girondot <[marc.girondot@gmail.com](mailto:marc.girondot@gmail.com)>

## See Also

Other Periodic patterns of indices: [index.periodic\(\)](#), [moon.info\(\)](#), [sun.info\(\)](#), [tide.info\(\)](#)

## Examples

```
## Not run:
library("HelpersMG")
# Generate a timeserie of time
time.obs <- NULL
for (i in 0:9) time.obs <- c(time.obs, c(0, 6, 12, 18)+i*24)
# For these time, generate a timeseries of temperatures
temp.obs <- rep(NA, length(time.obs))
temp.obs[3+(0:9)*4] <- rnorm(10, 25, 3)
temp.obs[1+(0:9)*4] <- rnorm(10, 10, 3)
for (i in 1:(length(time.obs)-1))
  if (is.na(temp.obs[i]))
    temp.obs[i] <- mean(c(temp.obs[i-1], temp.obs[i+1]))
  if (is.na(temp.obs[length(time.obs)]))
    temp.obs[length(time.obs)] <- temp.obs[length(time.obs)-1]/2
```

```

observed <- data.frame(time=time.obs, temperature=temp.obs)
# Search for the minimum and maximum values
r <- minmax.periodic(time.minmax.daily=c(Min=2, Max=15),
observed=observed, period=24, colname.index="temperature")

# Estimate all the temperatures for these values
t <- index.periodic(minmax=r)

plot_errbar(x=t[, "time"], y=t[, "index"],
errbar.y=ifelse(is.na(t[, "sd"]), 0, 2*t[, "sd"]),
type="l", las=1, bty="n", errbar.y.polygon = TRUE,
xlab="hours", ylab="Temperatures", ylim=c(0, 35),
errbar.y.polygon.list = list(col="grey"))

plot_add(x=t[, "time"], y=t[, "index"], type="l")

plot_add(observed$time, observed$temperature, pch=19, cex=0.5)

## End(Not run)

```

**modeled.hist***Return the theoretical value for the histogram bar***Description**

Return the theoretical value for the histogram bar based on a model of distribution.

**Usage**

```
modeled.hist(breaks, FUN, ..., sum = 1)
```

**Arguments**

<code>breaks</code>	Vector with the breaks; it can be obtained directly from <code>hist()</code>
<code>FUN</code>	Function to be used to integrate the density, ex. <code>pnorm</code>
<code>...</code>	Parameters to be used by <code>FUN</code>
<code>sum</code>	Total numbers in the histogram; 1 for emperical frequencies

**Details**

`modeled.hist` returns the theoretical value for the histogram bar based on a model of distribution.

**Value**

A list with x (the center of the bar) and y components

**Author(s)**

Marc Girondot <[marc.girondot@gmail.com](mailto:marc.girondot@gmail.com)>

## Examples

```
## Not run:  
n <- rnorm(100, mean=10, sd=2)  
breaks <- 0:20  
hist(n, breaks=breaks)  
  
s <- modeled.hist(breaks=breaks, FUN=pnorm, mean=10, sd=2, sum=100)  
  
points(s$x, s$y, pch=19)  
lines(s$x, s$y)  
  
n <- rlnorm(100, meanlog=2, sdlog=0.4)  
b <- hist(n, ylim=c(0, 70))  
  
s <- modeled.hist(breaks=b$breaks, FUN=plnorm, meanlog=2, sdlog=0.4, sum=100)  
  
points(s$x, s$y, pch=19)  
lines(s$x, s$y)  
  
## End(Not run)
```

---

modifyVector	<i>Modifies Elements of a Vector</i>
--------------	--------------------------------------

---

## Description

Modifies a vector by changing a subset of elements to match a second vector.

## Usage

```
modifyVector(x, val, add = TRUE)
```

## Arguments

- |     |  |
|-----|--|
| x   | A named vector.  |
| val | A named vector with components to replace corresponding components in x. |
| add | If FALSE, only existing elements of x are returned.                      |

## Details

modifyVector modifies elements of a vector

## Value

A modified version of x, with the elements of val replacing the elements of x

## Author(s)

Marc Girondot

## Examples

```
library("HelpersMG")
e <- c(M=10, L=20, J=30)
modifyVector(e, c(U=10, M=30))
modifyVector(e, c(U=10, M=30), add=FALSE)
```

**moon.info**

*Moon phase based on a date*

## Description

The script gives an index (base 100) that represents moon phase.

If the return value (from 0 to 100) is between:

0 and 1.6931595 or 98.3068405 and 100, it is full moon,  
23.3068405 and 26.6931595, last quarter,  
48.3068405 and 51.6931595, new moon,  
73.3068405 and 76.6931595, first quarter

When phase is set to TRUE, a character representing the moon phase is returned.

## Usage

```
moon.info(date = Sys.Date(), phase = FALSE)
```

## Arguments

- |       |   |
|-------|---|
| date  | A date in class Date. By default, it will use today date            |
| phase | If TRUE, a vector of characters with NM, FQ, FL LQ will be returned |

## Details

moon.info calculates the moon phase based on a date.

## Value

Return a value describing the moon phase:

0 and 100 are full moon, 50 is new moon, 25 last quarter and 75 first quarter

## Author(s)

Marc Girondot <marc.girondot@gmail.com>

## See Also

Other Periodic patterns of indices: [index.periodic\(\)](#), [minmax.periodic\(\)](#), [sun.info\(\)](#), [tide.info\(\)](#)

## Examples

```
## Not run:
library("HelpersMG")
moon.info(as.Date("2001-12-31"))
moon.info(as.Date("14/04/2010", "%d/%m/%Y"))
moon.info(as.Date("22/06/07", "%d/%m/%y"))
moon.info(seq(from=as.Date("2012-03-01"),
to=as.Date("2012-04-15"), by="days"))
moon.info(seq(from=as.Date("2012-03-01"),
to=as.Date("2012-04-15"), by="days"), phase=TRUE)

## End(Not run)
```

MovingWindow

*Return a moving average of a vector.*

## Description

Return a moving average of a vector./cr hole parameter can be none, bothL, bothR, both, begin, end.

## Usage

```
MovingWindow(x, window, hole = "begin", fill = TRUE, FUN = mean)
```

## Arguments

x	The vector to analyze
window	The window size
hole	Should the returned vector have the same length than x
fill	TRUE or FALSE, should the vector return NA
FUN	Function to apply to the window

## Details

MovingWindow returns a moving average of a vector.

## Value

A vector

## Author(s)

Marc Girondot <marc.girondot@gmail.com>

## Examples

```
MovingWindow(1:10, window = 4, fill = TRUE, hole="bothL")
MovingWindow(1:10, window = 4, fill = TRUE, hole="bothR")
MovingWindow(1:10, window = 4, fill = TRUE, hole="both")
MovingWindow(1:10, window = 4, fill = TRUE, hole="none")
MovingWindow(1:10, window = 4, fill = TRUE, hole="begin")
MovingWindow(1:10, window = 4, fill = TRUE, hole="end")
MovingWindow(1:10, window = 4, fill = TRUE, hole="end", FUN=sd)
```

NagelkerkeScaledR2      *Return the scaled R2 defined by Nagelkerke (1991)*

## Description

Return the scaled R2 of a binomial model based on:

Nagelkerke NJD (1991) A note on a general definition of the coefficient of determination. Biometrika 78:691-192.

This definition of scaled R2 by Nagelkerke (1991) has the following properties:

- (i) It is consistent with classical R2, that is the general definition applied to e.g. linear regression yields the classical R2.
- (ii) It is consistent with maximum likelihood as an estimation method, i.e. the maximum likelihood estimates of the model parameters maximize R2.
- (iii) It is asymptotically independent of the sample size n.
- (iv) 1-R2 has the interpretation of the proportion of unexplained 'variation'.
- (v) It is dimensionless, i.e. it does not depend on the units used.

The reported value is similar to the value estimated with nagelkerke() function from rcompanion package but not from the NagelkerkeR2() function from fmsb package. I don't know why.

## Usage

```
NagelkerkeScaledR2(x, size, prediction, scaled = TRUE)
```

## Arguments

x	The number of observations
size	Number of trials
prediction	Prediction of x/size
scaled	If TRUE, return the scaled R2

## Details

NagelkerkeScaledR2 returns the scaled R2 defined by Nagelkerke (1991)

## Value

The scaled R2 value

**Author(s)**

Marc Girondot <marc.girondot@gmail.com>

**Examples**

```
x <- c(10, 9, 6, 4, 3, 1, 0)
size <- c(10, 10, 10, 10, 10, 10, 10)
prediction <- c(0.9, 0.8, 0.7, 0.5, 0.4, 0.3, 0.2)
NagelkerkeScaledR2(x, size, prediction)

# Using the example in fmsb::NagelkerkeR2
res <- glm(cbind(ncases,ncontrols) ~ agegp+alcgp+tobgp, data=esoph, family=binomial())
NagelkerkeScaledR2(x=esoph$ncases, size = esoph$ncases+esoph$ncontrols,
                    prediction = res$fitted.values)
```

newcompassRose

*Display a compass rose***Description**

Displays a basic compass rose, usually to orient a map.

`newcompassRose` displays a conventional compass rose at the position requested.

The size of the compass rose is determined by the character expansion, as the central "rose" is calculated relative to the character size.

Rotation is in degrees counterclockwise.

**Usage**

```
newcompassRose(
  x,
  y,
  rot = 0,
  cex = 1,
  col = "black",
  col.arrows.light = "white",
  col.arrows.dark = "black"
)
```

**Arguments**

<code>x</code>	The position of the center of the compass rose in user units.
<code>y</code>	The position of the center of the compass rose in user units.
<code>rot</code>	Rotation for the compass rose in degrees. See Details.
<code>cex</code>	The character expansion to use in the display.
<code>col</code>	The color of text
<code>col.arrows.light</code>	The color of lighter lines

```
col.arrows.dark
The color of darker lines
```

## Details

`newcompassRose` Display a compass rose

## Value

`none`

## Author(s)

modified from Jim Lemon; See `compassRose` sp

## Examples

```
## Not run:
library(HelpersMG)
require("maps")
map("world", "China")
newcompassRose(x=110, y=35, col.arrows.light="grey")

## End(Not run)
```

`newmap.scale`

*Add Scale to Existing Unprojected Map*

## Description

Adds a scale to an existing map, both as a ratio and a distance gauge. If `x` or `y` are not specified, this will be taken to be near the lower left corner of the map.

## Usage

```
newmap.scale(
  x,
  y,
  relwidth = 0.15,
  metric = TRUE,
  ratio = TRUE,
  col.line = "black",
  ...
)
```

**Arguments**

x	Location of left end of distance gauge.
y	Location of left end of distance gauge.
relwidth	Proportion of width of display to be used for the scale. The default is 0.15.
metric	If TRUE, the distance gauge will be in km, otherwise miles.
ratio	If FALSE, the scale ratio of the map is not displayed.
col.line	The color of lines for the gauge.
...	Further plotting parameters may be specified as for the command text().

**Details**

`newmap.scale` Add Scale to Existing Unprojected Map

**Value**

The exact calculated scale is returned.

**Author(s)**

See `map.scale` maps

**Examples**

```
## Not run:  
library("maps")  
library("HelpersMG")  
map("world", "China")  
newmap.scale(col.line = "red", col="blue")  
  
## End(Not run)
```

---

openwd

*Open a finder window with current working directory in MacOS X and windows*

---

**Description**

This function opens a finder window with directory files in MacOS X. It has not been fully tested in Windows. In linux, it just returns the list of files in directory.  
By default, it uses the current working directory.

**Usage**

```
openwd(directory = getwd())
```

**Arguments**

`directory`      The directory you want to open

**Details**

`openwd` will open a finder window with current working directory

**Value**

A vector with the list of files.

**Author(s)**

Marc Girondot <[marc.girondot@gmail.com](mailto:marc.girondot@gmail.com)>

**Examples**

```
## Not run:  
openwd()  
  
## End(Not run)
```

`plot.cutter`

*Plot results of cutter that best describe distribution*

**Description**

Plot the estimates of cut distribution.

**Usage**

```
## S3 method for class 'cutter'  
plot(  
  x,  
  col.hist = "grey",  
  col.DL = "blue",  
  col.dist = "black",  
  col.unobserved = "green",  
  col.mcmc = rgb(red = 0.6, green = 0, blue = 0, alpha = 0.01),  
  legend = TRUE,  
  ...  
)
```

## Arguments

x	A result file generated by cutter
col.hist	The color of histogram
col.DL	The color of below of above samples
col.dist	The color of distribution
col.unobserved	The color of unobserved states
col.mcmc	The color of mcmc outputs
legend	If TRUE, a legend is shown
...	Parameters for plot

## Details

plot.cutter plot result of cutter

## Value

Nothing

## Author(s)

Marc Girondot <marc.girondot@gmail.com>

## See Also

Other Distributions: [cutter\(\)](#), [dbeta\\_new\(\)](#), [dcutter\(\)](#), [dggamma\(\)](#), [logLik.cutter\(\)](#), [pSnbino](#)(), [print.cutter\(\)](#), [qSnbino](#)(), [r2norm\(\)](#), [rcutter\(\)](#), [rmnrm\(\)](#), [rnbinom\\_new\(\)](#)

## Examples

```
## Not run:
library(HelperMG)
# -----
# right censored distribution with gamma distribution
# -----
# Detection limit
DL <- 100
# Generate 100 random data from a gamma distribution
obc <- rgamma(100, scale=20, shape=2)
# remove the data below the detection limit
obc[obc>DL] <- +Inf
# search for the parameters the best fit these censored data
result <- cutter(observations=obc, upper_detection_limit=DL,
                  cut_method="censored")
result
plot(result, xlim=c(0, 150), breaks=seq(from=0, to=150, by=10))
# -----
# The same data seen as truncated data with gamma distribution
# -----
```

```

obc <- obc[is.finite(obc)]
# search for the parameters the best fit these truncated data
result <- cutter(observations=obc, upper_detection_limit=DL,
                  cut_method="truncated")
result
plot(result, xlim=c(0, 150), breaks=seq(from=0, to=150, by=10))
#
# -----
# left censored distribution with gamma distribution
# -----
# Detection limit
DL <- 10
# Generate 100 random data from a gamma distribution
obc <- rgamma(100, scale=20, shape=2)
# remove the data below the detection limit
obc[obc<DL] <- -Inf
# search for the parameters the best fit these truncated data
result <- cutter(observations=obc, lower_detection_limit=DL,
                  cut_method="censored")
result
plot(result)
plot(result, xlim=c(0, 200), breaks=seq(from=0, to=200, by=10))
#
# -----
# left and right censored distribution
# -----
# Generate 100 random data from a gamma distribution
obc <- rgamma(100, scale=20, shape=2)
# Detection limit
LDL <- 10
# remove the data below the detection limit
obc[obc<LDL] <- -Inf
# Detection limit
UDL <- 100
# remove the data below the detection limit
obc[obc>UDL] <- +Inf
# search for the parameters the best fit these censored data
result <- cutter(observations=obc, lower_detection_limit=LDL,
                  upper_detection_limit=UDL,
                  cut_method="censored")
result
plot(result, xlim=c(0, 150), col.DL=c("black", "grey"),
      col.unobserved=c("green", "blue"),
      breaks=seq(from=0, to=150, by=10))
#
# -----
# Example with two values for lower detection limits
# corresponding at two different methods of detection for example
# with gamma distribution
#
obc <- rgamma(50, scale=20, shape=2)
# Detection limit for sample 1 to 50
LDL1 <- 10
# remove the data below the detection limit
obc[obc<LDL1] <- -Inf
obc2 <- rgamma(50, scale=20, shape=2)

```

```
# Detection limit for sample 1 to 50
LDL2 <- 20
# remove the data below the detection limit
obc2[obc2<LDL2] <- -Inf
obc <- c(obc, obc2)
# search for the parameters the best fit these censored data
result <- cutter(observations=obc,
                  lower_detection_limit=c(rep(LDL1, 50), rep(LDL2, 50)),
                  cut_method="censored")
result
# It is difficult to choose the best set of colors
plot(result, xlim=c(0, 150), col.dist="red",
      col.unobserved=c(rgb(red=1, green=0, blue=0, alpha=0.1),
                        rgb(red=1, green=0, blue=0, alpha=0.2)),
      col.DL=c(rgb(red=0, green=0, blue=1, alpha=0.5),
                rgb(red=0, green=0, blue=1, alpha=0.9)),
      breaks=seq(from=0, to=200, by=10))
#
# -----
# left censored distribution comparison of normal, lognormal and gamma
#
# Detection limit
DL <- 10
# Generate 100 random data from a gamma distribution
obc <- rgamma(100, scale=20, shape=2)
# remove the data below the detection limit
obc[obc<DL] <- -Inf
# search for the parameters the best fit these truncated data
result_gamma <- cutter(observations=obc, lower_detection_limit=DL,
                       cut_method="censored", distribution="gamma")
result_gamma
plot(result_gamma, xlim=c(0, 250), breaks=seq(from=0, to=250, by=10))

result_lognormal <- cutter(observations=obc, lower_detection_limit=DL,
                           cut_method="censored", distribution="lognormal")
result_lognormal
plot(result_lognormal, xlim=c(0, 250), breaks=seq(from=0, to=250, by=10))

result_normal <- cutter(observations=obc, lower_detection_limit=DL,
                        cut_method="censored", distribution="normal")
result_normal
plot(result_normal, xlim=c(0, 250), breaks=seq(from=0, to=250, by=10))

compare_AICc(gamma=result_gamma,
              lognormal=result_lognormal,
              normal=result_normal)
#
# -----
# Test for similarity in gamma left censored distribution between two
# datasets
#
# -----
obc1 <- rgamma(100, scale=20, shape=2)
# Detection limit for sample 1 to 50
LDL <- 10
# remove the data below the detection limit
```

```

obc1[obc1<LDL] <- -Inf
obc2 <- rgamma(100, scale=10, shape=2)
# remove the data below the detection limit
obc2[obc2<LDL] <- -Inf
# search for the parameters the best fit these censored data
result1 <- cutter(observations=obc1,
                   distribution="gamma",
                   lower_detection_limit=LDL,
                   cut_method="censored")
logLik(result1)
plot(result1, xlim=c(0, 200),
      breaks=seq(from=0, to=200, by=10))
result2 <- cutter(observations=obc2,
                   distribution="gamma",
                   lower_detection_limit=LDL,
                   cut_method="censored")
logLik(result2)
plot(result2, xlim=c(0, 200),
      breaks=seq(from=0, to=200, by=10))
result_tot1 <- cutter(observations=c(obc1, obc2),
                      distribution="gamma",
                      lower_detection_limit=LDL,
                      cut_method="censored")
logLik(result_tot1)
plot(result_tot1, xlim=c(0, 200),
      breaks=seq(from=0, to=200, by=10))

compare_AICc(Separate=list(result1, result2),
              Common=result_tot1, factor.value=1)
compare_BIC(Separate=list(result1, result2),
             Common=result_tot1, factor.value=1)

## End(Not run)

```

**plot.IconoCorel***Clean the dataframe before to be used with IC\_threshold\_matrix*

## Description

This function plots the data as a network. It returns an invisible object that can be used with visIgraph from package visNetwork. [https://fr.wikipedia.org/wiki/Iconographie\\_des\\_corrélations](https://fr.wikipedia.org/wiki/Iconographie_des_corrélations)

## Usage

```

## S3 method for class 'IconoCorel'
plot(
  x,
  ...,
  show.legend.direction = "bottomright",
  show.legend.strength = "topleft",

```

```

title = "Correlation iconography",
vertex.label.color = "black",
vertex.label = NULL,
vertex.color = "white",
vertex.label.cex = 1,
plot = TRUE
)

```

## Arguments

x	The correlation matrix to show
...	other options of plot.igraph()
show.legend.direction	the position of the legend of direction; FALSE to not show it
show.legend.strength	the position of the legend with intensity of correlation; FALSE to not show it
title	the title of the plot
vertex.label.color	a vector with the colors of labels
vertex.label	a vector with the labels
vertex.color	a vector of colors
vertex.label.cex	a vector of cex
plot	if TRUE, the plot is shown

## Details

plot.IconoCorel checks and corrects the dataframe to be used with IC\_threshold\_matrix

## Value

A igraph object

## Author(s)

Marc Girondot <[marc.girondot@gmail.com](mailto:marc.girondot@gmail.com)>

## References

Lesty, M., 1999. Une nouvelle approche dans le choix des régresseurs de la régression multiple en présence d'interactions et de colinéarités. Revue de Modulad 22, 41-77.

## See Also

Other Iconography of correlations: [IC\\_clean\\_data\(\)](#), [IC\\_correlation\\_simplify\(\)](#), [IC\\_threshold\\_matrix\(\)](#)

## Examples

```

## Not run:
library("HelpersMG")
es <- structure(list(Student = c("e1", "e2", "e3", "e4", "e5", "e6", "e7", "e8"),
                      Mass = c(52, 59, 55, 58, 66, 62, 63, 69),
                      Age = c(12, 12.5, 13, 14.5, 15.5, 16, 17, 18),
                      Assiduity = c(12, 9, 15, 5, 11, 15, 12, 9),
                      Note = c(5, 5, 9, 5, 13.5, 18, 18, 18)),
                      row.names = c(NA, -8L), class = "data.frame")

es

df <- IC_clean_data(es, debug = TRUE)
cor_matrix <- IC_threshold_matrix(data=df, threshold = NULL, progress=FALSE)
cor_threshold <- IC_threshold_matrix(data=df, threshold = 0.3)
par(mar=c(1,1,1,1))
set.seed(4)
library("igraph")
library("visNetwork")
kk <- plot(cor_threshold, vertex.color="red")
# it can be shown also with the visNetwork package
visIgraph(kk)
cor_threshold_Note <- IC_correlation_simplify(matrix=cor_threshold, variable="Note")
plot(cor_threshold_Note)

# You can record the position of elements and use them later
ly <- layout_nicely(kk)
plot(cor_threshold, vertex.color="red", layout=ly)

## End(Not run)

```

plot.LD50

*Plot results of LD50() that best describe LD50*

## Description

Plot the estimates that best describe lethality of doses.

## Usage

```

## S3 method for class 'LD50'
plot(
  x,
  ...,
  las.x = 1,
  las.y = 1,
  lab.PT = "LD50",
  at = NULL,
  lab.TRD = paste0("Transitional range of doses l=", l * 100, "%"),

```

```

col.TRD = "gray",
col.TRD.CI = rgb(0.8, 0.8, 0.8, 0.5),
col.PT.CI = rgb(0.8, 0.8, 0.8, 0.5),
show.CI = TRUE
)

```

**Arguments**

x	A result file generated by IC50()
...	Parameters for plot()
las.x	las parameter for x axis
las.y	las parameter for y axis
lab.PT	Label to describe pivotal dose
at	Position of ticks in x-axis
lab.TRD	Label to describe transitional range of dose
col.TRD	The color of TRD
col.TRD.CI	The color of CI of TRD based on range.CI
col.PT.CI	The color of CI of PT based on range.CI
show.CI	Do the CI for the curve should be shown

**Details**

plot.LD50 plot result of IC50() that best describe IC50

**Value**

Nothing

**Author(s)**

Marc Girondot <marc.girondot@gmail.com>

**See Also**

Other LD50 functions: [LD50\\_MHmcmc\\_p\(\)](#), [LD50\\_MHmcmc\(\)](#), [LD50\(\)](#), [logLik.LD50\(\)](#), [predict.LD50\(\)](#)

**Examples**

```

## Not run:
data <- data.frame(Doses=c(80, 120, 150, 150, 180, 200),
Alive=c(10, 12, 8, 6, 2, 1),
Dead=c(0, 1, 5, 6, 9, 15))
LD50_logistic <- LD50(data, equation="logistic")
predict(LD50_logistic, doses=c(140, 170))
plot(LD50_logistic, xlim=c(0, 300))

## End(Not run)

```

---

**plot.mcmcComposite**      *Plot the result of a mcmcComposite object*

---

### Description

Plot the results within a mcmcComposite object.  
 If scale.prior is TRUE, another scale is shown at right.  
 legend can take these values:  
 FALSE, TRUE, topleft, topright, bottomleft, bottomright, c(x=, y=)

### Usage

```
## S3 method for class 'mcmcComposite'
plot(
  x,
  ...,
  chain = 1,
  parameters = 1,
  transform = NULL,
  scale.prior = TRUE,
  legend = "topright",
  ylab = "Posterior density",
  las = 1,
  show.prior = TRUE,
  col.prior = "red",
  lty.prior = 1,
  lwd.prior = 1,
  col.posterior = "white",
  lty.posterior = 1,
  lwd.posterior = 1,
  ylab.prior = "Prior density"
)
```

### Arguments

<b>x</b>	A mcmcComposite object
<b>...</b>	Graphical parameters to be sent to hist()
<b>chain</b>	The chain to use
<b>parameters</b>	Name of parameters or "all"
<b>transform</b>	Function to be used to transform the variable
<b>scale.prior</b>	If TRUE, the prior is scaled at the same size as posterior
<b>legend</b>	If FALSE, the legend is not shown; see description
<b>ylab</b>	y-label for posterior
<b>las</b>	las parameter (orientation of y-axis graduation)

show.prior	whether the prior be shown?
col.prior	Color for prior curve
lty.prior	Type of line for prior curve
lwd.prior	Width of line for prior curve
col.posterior	Color for posterior histogram
lty.posterior	Type of line for posterior histogram
lwd.posterior	Width of line for posterior histogram
ylab.prior	y-label for prior

**Details**

plot.mcmcComposite plots the result of a MCMC search

**Value**

None

**Author(s)**

Marc Girondot <marc.girondot@gmail.com>

**See Also**

Other mcmcComposite functions: [MHalgoGen\(\)](#), [as.mcmc.mcmcComposite\(\)](#), [as.parameters\(\)](#), [as.quantiles\(\)](#), [merge.mcmcComposite\(\)](#), [plot.PriorsmcmcComposite\(\)](#), [setPriors\(\)](#), [summary.mcmcComposite\(\)](#)

**Examples**

```
## Not run:
library(HelpersMG)
require(coda)
x <- rnorm(30, 10, 2)
dnormx <- function(data, x) {
  data <- unlist(data)
  return(-sum(dnorm(data, mean=x['mean'], sd=x['sd'], log=TRUE)))
}
parameters_mcmc <- data.frame(Density=c('dnorm', 'dlnorm'),
Prior1=c(10, 0.5), Prior2=c(2, 0.5), SDProp=c(1, 1),
Min=c(-3, 0), Max=c(100, 10), Init=c(10, 2), stringsAsFactors = FALSE,
row.names=c('mean', 'sd'))
mcmc_run <- MHalgoGen(n.ITER=50000, parameters=parameters_mcmc, data=x,
adaptive = TRUE,
likelihood=dnormx, n.chains=1, n.adapt=100, thin=1, trace=1)
plot(mcmc_run, xlim=c(0, 20))
plot(mcmc_run, xlim=c(0, 10), parameters="sd")
mcmcforcoda <- as.mcmc(mcmc_run)
#' heidel.diag(mcmcforcoda)
raftery.diag(mcmcforcoda)
autocorr.diag(mcmcforcoda)
```

```

acf(mcmcforcoda[[1]][,"mean"], lag.max=20, bty="n", las=1)
acf(mcmcforcoda[[1]][,"sd"], lag.max=20, bty="n", las=1)
batchSE(mcmcforcoda, batchSize=100)
# The batch standard error procedure is usually thought to
# be not as accurate as the time series methods used in summary
summary(mcmcforcoda)$statistics[, "Time-series SE"]
summary(mcmc_run)
as.parameters(mcmc_run)
lastp <- as.parameters(mcmc_run, index="last")
parameters_mcmc[,"Init"] <- lastp
# The n.adapt set to 1 is used to not record the first set of parameters
# then it is not duplicated (as it is also the last one for
# the object mcmc_run)
mcmc_run2 <- MHalgoGen(n.iter=50000, parameters=parameters_mcmc, data=x,
adaptive = TRUE,
likelihood=dnormx, n.chains=1, n.adapt=1, thin=1, trace=1)
mcmc_run3 <- merge(mcmc_run, mcmc_run2)
##### no adaptation, n.adapt must be 0
parameters_mcmc[,"Init"] <- c(mean(x), sd(x))
mcmc_run3 <- MHalgoGen(n.iter=1000, parameters=parameters_mcmc, data=x,
adaptive = TRUE,
likelihood=dnormx, n.chains=1, n.adapt=0, thin=1, trace=1)

#####
## Example with transform
#####

x.1<-rnorm(6000, 2.4, 0.6)
x.2<-rlnorm(10000, 1.3, 0.1)

X<-c(x.1, x.2)
hist(X,100,freq=FALSE, ylim=c(0,1.5))
Lnormlnorm <- function(par, val) {
  p <- invlogit(par["p"])
  return(-sum(log(p*dnorm(val, par["m1"], abs(par["s1"])), log = FALSE) +
    (1-p)*dlnorm(val, par["m2"], abs(par["s2"])), log = FALSE)))
}
# Mean 1
m1=2.3; s1=0.5
# Mean 2
m2=1.3; s2=0.1
# proportion of category 1 - logit transform
p=0

par<-c(m1=m1, s1=s1, m2=m2, s2=s2, p=p)

result2<-optim(par, Lnormlnorm, method="BFGS", val=X,
                hessian=FALSE, control=list(trace=1))

lines(seq(from=0, to=5, length=100),
      dnorm(seq(from=0, to=5, length=100),
            result2$par["m1"], abs(result2$par["s1"])), col="red")

```

```

lines(seq(from=0, to=5, length=100),
      dlnorm(seq(from=0, to=5, length=100),
             result2$par["m2"], abs(result2$par["s2"])), col="green")

p <- invlogit(result2$par["p"])

paste("Proportion of Gaussian data", p)

lines(seq(from=0, to=5, length=100),
      p*dnorm(seq(from=0, to=5, length=100),
               result2$par["m1"], result2$par["s1"])+
      (1-p)*dlnorm(seq(from=0, to=5, length=100),
                    result2$par["m2"], result2$par["s2"])), col="blue")

parameters_mcmc <- data.frame(Density=c('dunif', 'dunif', 'dunif', 'dunif', 'dunif'),
                                 Prior1=c(0, 0.001, 0, 0.001, -3),
                                 Prior2=c(10, 10, 10, 10, 3),
                                 SDProp=c(1, 1, 1, 1, 1),
                                 Min=c(0, 0.001, 0, 0.001, -3),
                                 Max=c(10, 10, 10, 10, 3),
                                 Init=result2$par, stringsAsFactors = FALSE,
                                 row.names=c('m1', 's1', 'm2', 's2', 'p'))

mcmc_run <- MHalgoGen(n.iter=50000, parameters=parameters_mcmc, val=X,
                       parameters_name = "par",
                       adaptive = TRUE,
                       likelihood=LnormLnorm, n.chains=1,
                       n.adapt=100, thin=1, trace=100)
plot(mcmc_run, parameters="m1", breaks=seq(from=0, to =10, by=0.1),
      legend=c(x=6, y=0.10))
plot(mcmc_run, parameters="p", transform=invlogit, xlim=c(0,1),
      breaks=seq(from=0, to=1, by=0.01), legend=c(x=0.6, y=0.10))
plot(mcmc_run, parameters="p", xlim=c(-3,3),
      breaks=seq(from=-3, to =3, by=0.05), legend=c(x=1, y= 0.10))

parameters_mcmc <- data.frame(Density=c('dunif', 'dunif', 'dunif', 'dunif', 'dnorm'),
                                 Prior1=c(0, 0.001, 0, 0.001, 0.5),
                                 Prior2=c(10, 10, 10, 10, 1),
                                 SDProp=c(1, 1, 1, 1, 1),
                                 Min=c(0, 0.001, 0, 0.001, -3),
                                 Max=c(10, 10, 10, 10, 3),
                                 Init=result2$par, stringsAsFactors = FALSE,
                                 row.names=c('m1', 's1', 'm2', 's2', 'p'))

mcmc_run_pnorm <- MHalgoGen(n.iter=50000, parameters=parameters_mcmc, val=X,
                             parameters_name = "par",
                             adaptive = TRUE,
                             likelihood=LnormLnorm, n.chains=1,
                             n.adapt=100, thin=1, trace=100)
plot(mcmc_run_pnorm, parameters="m1", breaks=seq(from=0, to =10, by=0.1),
      legend=c(x=6, y=0.10))
plot(mcmc_run_pnorm, parameters="p", transform=invlogit, xlim=c(0,1),
      breaks=seq(from=0, to=1, by=0.01), legend=c(x=0.6, y=0.10))

```

```

plot(x=mcmc_run_pnorm, parameters="p", xlim=c(-3,3),
      breaks=seq(from=-3, to =3, by=0.05), legend=c(x=1, y= 0.10))

# Note that it is more logic to use beta distribution for p as a
# proportion. However p value must be checked to be used in optim
# The use of logit transform can be a problem because it can stuck
# the p value to 1 or 0 during fit.

Lnormlnorm <- function(par, val) {
  p <- par["p"]
  return(-sum(log(p*dnorm(val, par["m1"]), abs(par["s1"])), log = FALSE) +
         (1-p)*dlnorm(val, par["m2"], abs(par["s2"])), log = FALSE)))
}

# Example of beta distribution

# Mean is alpha/(alpha+beta)
# Variance is (alpha*beta)/((alpha+beta)^2*(alpha+beta+1))
alpha = 5
beta = 9
plot(x = seq(0.0001, 1, by = .0001),
      y = dbeta(seq(0.0001, 1, by = .0001), alpha, beta),
      type = "l", ylab="Density", xlab="p", bty="n")
points(x=alpha/(alpha+beta), y=0, pch=4)
segments(x0=alpha/(alpha+beta)-sqrt((alpha*beta)/((alpha+beta)^2*(alpha+beta+1))),
          x1=alpha/(alpha+beta)+sqrt((alpha*beta)/((alpha+beta)^2*(alpha+beta+1))), y0=0, y1=0)

# Use of optim with L-BFGS-B to limit p between 0 and 1 and s > 0

# Mean 1
m1=2.3; s1=0.5
# Mean 2
m2=1.3; s2=0.1
# proportion of category 1 - logit transform
p=0.5

par <- c(m1=m1, s1=s1, m2=m2, s2=s2, p=p)

result2 <- optim(par, Lnormlnorm, method="L-BFGS-B", val=X,
                  lower = c(-Inf, 0, -Inf, 0, 0),
                  upper = c(Inf, Inf, Inf, Inf, 1),
                  hessian=FALSE, control=list(trace=1))

parameters_mcmc <- data.frame(Density=c('dunif', 'dunif', 'dunif', 'dunif', 'dbeta'),
                                 Prior1=c(0, 0.001, 0, 0.001, 5),
                                 Prior2=c(10, 10, 10, 10, 9),
                                 SDProp=c(1, 1, 1, 1, 1),
                                 Min=c(0, 0.001, 0, 0.001, 0),
                                 Max=c(10, 10, 10, 10, 1),
                                 Init=c('m1' = 2.4,
                                       's1' = 0.6,

```

```

'm2' = 1.3,
's2' = 0.1,
'p' = 0.5), stringsAsFactors = FALSE,
row.names=c('m1', 's1', 'm2', 's2', 'p'))

mcmc_run_pbeta <- MHalgoGen(n.iter=50000, parameters=parameters_mcmc, val=X,
                             parameters_name = "par",
                             adaptive = TRUE,
                             likelihood=Lnormlnorm, n.chains=1,
                             n.adapt=100, thin=1, trace=100)
plot(mcmc_run_pbeta, parameters="m1", breaks=seq(from=0, to =10, by=0.1),
      legend=c(x=6, y=0.10))
plot(mcmc_run_pbeta, parameters="p", xlim=c(0,1),
      breaks=seq(from=0, to=1, by=0.01), legend=c(x=0.6, y=2))

## End(Not run)

```

**plot.PriorsmcmcComposite***Plot a prior defined with setPriors function***Description**

Create a ggplot graph with prior.  
The function makes minimal effort to decorate the plot.

**Usage**

```
## S3 method for class 'PriorsmcmcComposite'
plot(x, parameter = 1, ...)
```

**Arguments**

- x The priors to show
- parameter The name or rank of prior to show
- ... Not used

**Details**

`plot.PriorsmcmcComposite` plot a prior

**Value**

A ggplot object

**Author(s)**

Marc Girondot <marc.girondot@gmail.com>

**See Also**

Other mcmcComposite functions: [MHalgoGen\(\)](#), [as.mcmc.mcmcComposite\(\)](#), [as.parameters\(\)](#), [as.quantiles\(\)](#), [merge.mcmcComposite\(\)](#), [plot.mcmcComposite\(\)](#), [setPriors\(\)](#), [summary.mcmcComposite\(\)](#)

**Examples**

```
## Not run:
library("HelpersMG")
par <- c(a0=10, a1=2, b2=20, b1=-1)
rules <- rbind(data.frame(Name="^a", Min=0, Max="x*2"),
                 data.frame(Name="^b", Min=0, Max=100))
p <- setPriors(par=par, se=NULL, density="dgamma", rules=rules)
plot(p, parameter="a0")
q <- plot(p, parameter="b1")
q + geom_line(color = "red") + theme_bw() +
  theme(plot.margin=unit(c(2,1,1,1), 'cm'),
        panel.border = element_blank(),
        axis.line.x.bottom = element_line(colour = "black"),
        axis.line.y.left = element_line(colour = "black")) +
  labs(title="Parameter: b1") + theme(plot.title = element_text(hjust = 0.5))

## End(Not run)
```

**plot\_add**

*Add a plot to a previous one*

**Description**

To plot data, just add use it as a normal plot. It will plot the new data without axes, or labels for axes.

This function is complementary to matlines() and matpoints() from package graphics.

**Usage**

```
plot_add(...)
```

**Arguments**

...	Parameters for plot()
-----	-----------------------

**Details**

`plot_add` adds a plot to a previous one

**Value**

Nothing

**Author(s)**

Marc Girondot <marc.girondot@gmail.com>

**See Also**

Other plot and barplot functions: [ScalePreviousPlot\(\)](#), [barplot\\_errbar\(\)](#), [plot\\_errbar\(\)](#), [show\\_name\(\)](#)

**Examples**

```
## Not run:  
plot(x=1:100, y=sin(1:100), type="l", bty="n", xlim=c(1,200), xlab="x", ylab="y")  
plot_add(x=1:200, y=cos(1:200), type="l", bty="n", col="red")  
  
## End(Not run)
```

---

plot\_errbar

*Plot a xy graph with error bar on x and/or y*

---

**Description**

To plot data, just use it as a normal plot but add the errbar.x and errbar.y values or errbar.x.minus, errbar.x.plus if bars for x axis are asymmetric and errbar.y.minus, errbar.y.plus if bars for y axis are asymmetric. Use x.plus, x.minus, y.plus and y.minus to set absolute limits for error bars. Note that x.plus and x.minus have priority over errbar.x, errbar.x.minus and errbar.x.plus and that y.plus and y.minus have priority over errbar.y, errbar.y.minus and errbar.y.plus.

The parameter errbar.y.polygon=TRUE permits to define error as an envelope for y axis.

**Usage**

```
plot_errbar(  
  ...,  
  errbar.x = NULL,  
  errbar.y = NULL,  
  errbar.x.plus = NULL,  
  errbar.x.minus = NULL,  
  errbar.y.plus = NULL,  
  errbar.y.minus = NULL,  
  x.plus = NULL,  
  x.minus = NULL,  
  y.plus = NULL,  
  y.minus = NULL,  
  errbar.tick = 1/50,
```

```

errbar.lwd = par("lwd"),
errbar.lty = par("lty"),
errbar.col = par("fg"),
errbar.y.polygon = FALSE,
errbar.y.polygon.list = list(NULL),
names = NULL,
add = FALSE
)

```

### Arguments

...	Parameters for plot() such as main= or ylim=
errbar.x	The length of error bars for x. Recycled if necessary.
errbar.y	The length of error bars for y. Recycled if necessary.
errbar.x.plus	The length of positive error bars for x. Recycled if necessary.
errbar.x_MINUS	The length of negative error bars for x. Recycled if necessary.
errbar.y.plus	The length of positive error bars for y. Recycled if necessary.
errbar.y_MINUS	The length of negative error bars for y. Recycled if necessary.
x.plus	The absolut position of the positive error bar for x. Recycled if necessary.
x_MINUS	The absolut position of the negative error bar for x. Recycled if necessary.
y.plus	The absolut position of the positive error bar for y. Recycled if necessary.
y_MINUS	The absolut position of the nagative error bar for y. Recycled if necessary.
errbar.tick	Size of small ticks at the end of error bars defined as a proportion of total width or height graph size.
errbar.lwd	Error bar line width, see par("lwd")
errbar.lty	Error bar line type, see par("lwd")
errbar.col	Error bar line color, see par("col")
errbar.y.polygon	If true, the errors are shown as a filed polygon.
errbar.y.polygon.list	List of parameters to be used for polygon.
names	The names of the points to be used with show_name().
add	If true, add the graph to the previous one.

### Details

*plot\_errbar* plot a xy graph with error bar on x and/or y

### Value

A list with x, y and names for points

### Author(s)

Marc Girondot <marc.girondot@gmail.com>

**See Also**

[barplot\\_errorbar](#)

Other plot and barplot functions: [ScalePreviousPlot\(\)](#), [barplot\\_errbar\(\)](#), [plot\\_add\(\)](#), [show\\_name\(\)](#)

**Examples**

```
## Not run:
plot_errbar(1:100, rnorm(100, 1, 2),
            xlab="axe x", ylab="axe y", bty="n", xlim=c(1,100),
            errbar.x=2, errbar.y=rnorm(100, 1, 0.1))
x <- 1:100
plot_errbar(x=1:100, rnorm(100, 1, 2),
            xlab="axe x", ylab="axe y", bty="n", xlim=c(1,100),
            x.minus=x-2, x.plus=x+2)
x <- 1:100
plot_errbar(x=1:100, rnorm(100, 1, 2),
            xlab="axe x", ylab="axe y", bty="n",
            pch=21, bg="white",
            x.minus=x-10, x.plus=x+10)
x <- (1:200)/10
y <- sin(x)
plot_errbar(x=x, y=y, xlab="axe x", ylab="axe y", bty="n", xlim=c(1,20),
            y.minus=y-1, y.plus=y+1, ylim=c(-3, 3), type="l",
            errbar.y.polygon=TRUE,
            errbar.y.polygon.list=list(border=NA, col=rgb(0, 0, 0.5)))

## End(Not run)
```

**predict.LD50**

*Estimate survival according to doses*

**Description**

Estimate survival according to doses.

The returned data.frame has the following components:

doses, SE, survival, CI.minus.sexratio, CI.plus.sexratio, range.CI

**Usage**

```
## S3 method for class 'LD50'
predict(
  object,
  doses = NULL,
  SE = NULL,
  range.CI = 0.95,
  replicates = 1000,
  progressbar = FALSE,
```

```
  ...
)
```

### Arguments

object	A result file generated by LD50
doses	A vector of temperatures
SE	The standard error for doses, optional
range.CI	The range of confidence interval for estimation, default=0.95
replicates	Number of replicates to estimate CI
progressbar	Logical. Does a progression bar must be shown
...	Not used

### Details

*predict.LD50* Estimate survival according to doses

### Value

A data.frame with informations about survival

### Author(s)

Marc Girondot <marc.girondot@gmail.com>

### See Also

Other LD50 functions: [LD50\\_MHmcmc\\_p\(\)](#), [LD50\\_MHmcmc\(\)](#), [LD50\(\)](#), [logLik.LD50\(\)](#), [plot.LD50\(\)](#)

### Examples

```
## Not run:
#' data <- data.frame(Doses=c(80, 120, 150, 150, 180, 200),
#Alive=c(10, 12, 8, 6, 2, 1),
#Dead=c(0, 1, 5, 6, 9, 15))
LD50_logistic <- LD50(data, equation="logistic")
predict(LD50_logistic, doses=c(140, 170))
plot(LD50_logistic

## End(Not run)
```

---

**print.cutter***Print results of cutter that best describe distribution*

---

## Description

Print the estimates of cut distribution.

## Usage

```
## S3 method for class 'cutter'  
print(x, silent = FALSE, ...)
```

## Arguments

x	A result file generated by cutter
silent	If TRUE does not show the output
...	Not used

## Details

print.cutter plot result of cutter

## Value

Nothing

## Author(s)

Marc Girondot <marc.girondot@gmail.com>

## See Also

Other Distributions: [cutter\(\)](#), [dbeta\\_new\(\)](#), [dcutter\(\)](#), [dggamma\(\)](#), [logLik.cutter\(\)](#), [pSnbnom\(\)](#), [plot.cutter\(\)](#), [qSnbnom\(\)](#), [r2norm\(\)](#), [rcutter\(\)](#), [rmnrm\(\)](#), [rnbinom\\_new\(\)](#)

## Examples

```
## Not run:  
library(HelpersMG)  
# -----  
# right censored distribution with gamma distribution  
# -----  
# Detection limit  
DL <- 100  
# Generate 100 random data from a gamma distribution  
obc <- rgamma(100, scale=20, shape=2)  
# remove the data below the detection limit  
obc[obc>DL] <- +Inf
```

```

# search for the parameters the best fit these censored data
result <- cutter(observations=obc, upper_detection_limit=DL,
                  cut_method="censored")
result
plot(result, xlim=c(0, 150), breaks=seq(from=0, to=150, by=10))
#
# -----
# The same data seen as truncated data with gamma distribution
#
# -----
obc <- obc[is.finite(obc)]
# search for the parameters the best fit these truncated data
result <- cutter(observations=obc, upper_detection_limit=DL,
                  cut_method="truncated")
result
plot(result, xlim=c(0, 150), breaks=seq(from=0, to=150, by=10))
#
# -----
# left censored distribution with gamma distribution
#
# -----
# Detection limit
DL <- 10
# Generate 100 random data from a gamma distribution
obc <- rgamma(100, scale=20, shape=2)
# remove the data below the detection limit
obc[obc<DL] <- -Inf
# search for the parameters the best fit these truncated data
result <- cutter(observations=obc, lower_detection_limit=DL,
                  cut_method="censored")
result
plot(result, xlim=c(0, 200), breaks=seq(from=0, to=200, by=10))
#
# -----
# left and right censored distribution
#
# -----
# Generate 100 random data from a gamma distribution
obc <- rgamma(100, scale=20, shape=2)
# Detection limit
LDL <- 10
# remove the data below the detection limit
obc[obc<LDL] <- -Inf
# Detection limit
UDL <- 100
# remove the data below the detection limit
obc[obc>UDL] <- +Inf
# search for the parameters the best fit these censored data
result <- cutter(observations=obc, lower_detection_limit=LDL,
                  upper_detection_limit=UDL,
                  cut_method="censored")
result
plot(result, xlim=c(0, 150), col.DL=c("black", "grey"),
                  col.unobserved=c("green", "blue"),
                  breaks=seq(from=0, to=150, by=10))
#
# -----
# Example with two values for lower detection limits
# corresponding at two different methods of detection for example
# with gamma distribution

```

```
# -----
# obc <- rgamma(50, scale=20, shape=2)
# Detection limit for sample 1 to 50
LDL1 <- 10
# remove the data below the detection limit
obc[obc<LDL1] <- -Inf
obc2 <- rgamma(50, scale=20, shape=2)
# Detection limit for sample 1 to 50
LDL2 <- 20
# remove the data below the detection limit
obc2[obc2<LDL2] <- -Inf
obc <- c(obc, obc2)
# search for the parameters the best fit these censored data
result <- cutter(observations=obc,
                  lower_detection_limit=c(rep(LDL1, 50), rep(LDL2, 50)),
                  cut_method="censored")
result
# It is difficult to choose the best set of colors
plot(result, xlim=c(0, 150), col.dist="red",
      col.unobserved=c(rgb(red=1, green=0, blue=0, alpha=0.1),
                        rgb(red=1, green=0, blue=0, alpha=0.2)),
      col.DL=c(rgb(red=0, green=0, blue=1, alpha=0.5),
                rgb(red=0, green=0, blue=1, alpha=0.9)),
      breaks=seq(from=0, to=200, by=10))
# -----
# left censored distribution comparison of normal, lognormal and gamma
# -----
# Detection limit
DL <- 10
# Generate 100 random data from a gamma distribution
obc <- rgamma(100, scale=20, shape=2)
# remove the data below the detection limit
obc[obc<DL] <- -Inf
# search for the parameters the best fit these truncated data
result_gamma <- cutter(observations=obc, lower_detection_limit=DL,
                      cut_method="censored", distribution="gamma")
result_gamma
plot(result_gamma, xlim=c(0, 250), breaks=seq(from=0, to=250, by=10))

result_lognormal <- cutter(observations=obc, lower_detection_limit=DL,
                           cut_method="censored", distribution="lognormal")
result_lognormal
plot(result_lognormal, xlim=c(0, 250), breaks=seq(from=0, to=250, by=10))

result_normal <- cutter(observations=obc, lower_detection_limit=DL,
                        cut_method="censored", distribution="normal")
result_normal
plot(result_normal, xlim=c(0, 250), breaks=seq(from=0, to=250, by=10))

compare_AICc(gamma=result_gamma,
              lognormal=result_lognormal,
              normal=result_normal)
# -----
```

```

# Test for similarity in gamma left censored distribution between two
# datasets
#
# -----
obc1 <- rgamma(100, scale=20, shape=2)
# Detection limit for sample 1 to 50
LDL <- 10
# remove the data below the detection limit
obc1[obc1<LDL] <- -Inf
obc2 <- rgamma(100, scale=10, shape=2)
# remove the data below the detection limit
obc2[obc2<LDL] <- -Inf
# search for the parameters the best fit these censored data
result1 <- cutter(observations=obc1,
                    distribution="gamma",
                    lower_detection_limit=LDL,
                    cut_method="censored")
logLik(result1)
plot(result1, xlim=c(0, 200),
      breaks=seq(from=0, to=200, by=10))
result2 <- cutter(observations=obc2,
                    distribution="gamma",
                    lower_detection_limit=LDL,
                    cut_method="censored")
logLik(result2)
plot(result2, xlim=c(0, 200),
      breaks=seq(from=0, to=200, by=10))
result_totl <- cutter(observations=c(obc1, obc2),
                      distribution="gamma",
                      lower_detection_limit=LDL,
                      cut_method="censored")
logLik(result_totl)
plot(result_totl, xlim=c(0, 200),
      breaks=seq(from=0, to=200, by=10))

compare_AICc(Separate=list(result1, result2),
              Common=result_totl, factor.value=1)
compare_BIC(Separate=list(result1, result2),
             Common=result_totl, factor.value=1)

## End(Not run)

```

## Description

Distribution function for the sum of random variable with negative binomial distributions.

**Usage**

```
pSnbinom(
  q = stop("At least one quantile must be provided"),
  size = NULL,
  prob = NULL,
  mu = NULL,
  lower.tail = TRUE,
  log.p = FALSE,
  tol = 1e-06
)
```

**Arguments**

<code>q</code>	vector of quantiles.
<code>size</code>	target for number of successful trials, or dispersion parameter (the shape parameter of the gamma mixing distribution). Must be strictly positive, need not be integer.
<code>prob</code>	probability of success in each trial. $0 < \text{prob} \leq 1$ .
<code>mu</code>	alternative parametrization via mean.
<code>lower.tail</code>	logical; if TRUE (default), probabilities are $P[X \leq x]$ , otherwise, $P[X > x]$ .
<code>log.p</code>	logical; if TRUE, probabilities p are given as log(p).
<code>tol</code>	Tolerance for recurrence

**Details**

`pSnbinom` returns the distribution function for the sum of random variable with negative binomial distributions

**Value**

`pSnbinom` returns distribution function

**Author(s)**

Marc Girondot <[marc.girondot@gmail.com](mailto:marc.girondot@gmail.com)>

**See Also**

Other Distributions: [cutter\(\)](#), [dbeta\\_new\(\)](#), [dcutter\(\)](#), [dggamma\(\)](#), [logLik.cutter\(\)](#), [plot.cutter\(\)](#), [print.cutter\(\)](#), [qSnbinom\(\)](#), [r2norm\(\)](#), [rcutter\(\)](#), [rmnorm\(\)](#), [rnbinom\\_new\(\)](#)

**Examples**

```
## Not run:
alpha <- c(2.1, 2.05, 2)
mu <- c(10, 30, 20)
p <- pSnbinom(q=10, size=alpha, mu=mu, lower.tail = TRUE)

## End(Not run)
```

**qSnb***n**om**Quantile function for the sum of random variable with negative binomial distributions.***Description**

Quantile function for the sum of random variable with negative binomial distributions.

**Usage**

```
qSnb(
  p = stop("At least one probability must be provided"),
  size = stop("size parameter is mandatory"),
  prob = NULL,
  mu = NULL,
  lower.tail = TRUE,
  log.p = FALSE,
  tol = 1e-06
)
```

**Arguments**

<b>p</b>	vector of probabilities.
<b>size</b>	target for number of successful trials, or dispersion parameter (the shape parameter of the gamma mixing distribution). Must be strictly positive, need not be integer.
<b>prob</b>	probability of success in each trial. $0 < \text{prob} \leq 1$ .
<b>mu</b>	alternative parametrization via mean.
<b>lower.tail</b>	logical; if TRUE (default), probabilities are $P[X \leq x]$ , otherwise, $P[X > x]$ .
<b>log.p</b>	logical; if TRUE, probabilities p are given as log(p).
<b>tol</b>	Tolerance for recurrence

**Details**

**qSnb** returns the quantile function for the sum of random variable with negative binomial distributions

**Value**

**qSnb** returns quantile function

**Author(s)**

Marc Girondot <[marc.girondot@gmail.com](mailto:marc.girondot@gmail.com)>

**See Also**

Other Distributions: [cutter\(\)](#), [dbeta\\_new\(\)](#), [dcutter\(\)](#), [dggamma\(\)](#), [logLik.cutter\(\)](#), [pSnbino\(m\)](#), [plot.cutter\(\)](#), [print.cutter\(\)](#), [r2norm\(\)](#), [rcutter\(\)](#), [rmnrm\(\)](#), [rnbinom\\_new\(\)](#)

**Examples**

```
## Not run:
alpha <- c(2.1, 2.05, 2)
mu <- c(10, 30, 20)
q <- qSnbino(m(p=0.1, size=alpha, mu=mu, lower.tail = TRUE))

## End(Not run)
```

qvlmer

*Quasi Variances for lmer Model Coefficients***Description**

Computes a set of quasi variances (and corresponding quasi standard errors) for estimated model coefficients relating to the levels of a categorical (i.e., factor) explanatory variable. For details of the method see Firth (2000), Firth (2003) or Firth and de Menezes (2004). Quasi variances generalize and improve the accuracy of “floating absolute risk” (Easton et al., 1991). This device for economical model summary was first suggested by Ridout (1989).

Modified from qvcalc.lm() of packages qvcalc by David Firth, d.firth@warwick.ac.uk

**Usage**

```
qvlmer(object, factorname = NULL, coef.indices = NULL, dispersion = NULL, ...)
```

**Arguments**

<code>object</code>	A object obtained using lmer from package lme4
<code>factorname</code>	Either NULL, or a character vector of length 1
<code>coef.indices</code>	Either NULL, or a numeric vector of length at least 3
<code>dispersion</code>	An optional scalar multiplier for the covariance matrix, to cope with overdispersion for example
<code>...</code>	Other arguments to pass to qvcalc.default

**Details**

qvlmer is Quasi Variances for lmer Model Coefficients

**Value**

A list of class qv.

**Author(s)**

Marc Girondot <marc.girondot@gmail.com>

**References**

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- Firth, D. (2000) Quasi-variances in Xlisp-Stat and on the web. *Journal of Statistical Software* 5.4, 1–13. At <http://www.jstatsoft.org>
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- Menezes, R. X. de (1999) More useful standard errors for group and factor effects in generalized linear models. D.Phil. Thesis, Department of Statistics, University of Oxford.
- Ridout, M.S. (1989). Summarizing the results of fitting generalized linear models to data from designed experiments. In: Statistical Modelling: Proceedings of GLIM89 and the 4th International Workshop on Statistical Modelling held in Trento, Italy, July 17–21, 1989 (A. Decarli et al., eds.), pp 262–269. New York: Springer.

**Examples**

```
## Not run:
x <- rnorm(100)
y <- rnorm(100)
G <- as.factor(sample(c("A", "B", "C", "D"), 100, replace = TRUE))
R <- as.factor(rep(1:25, 4))
library(lme4)
m <- lmer(y ~ x + G + (1 | R))
qvlmer(m, factorname="G")

## End(Not run)
```

**r2norm**

*Random generation for Gaussian distributions different at left and right*

**Description**

Random generation for Gaussian distributions different at left and right

**Usage**

```
r2norm(n, mean = 0, sd_low = 1, sd_high = 1)
```

**Arguments**

<code>n</code>	number of observations.
<code>mean</code>	vector of means
<code>sd_low</code>	vector of standard deviations below the mean.
<code>sd_high</code>	vector of standard deviations above the mean.

**Details**

`r2norm` returns random numbers for Gaussian distributions different at left and right

**Value**

`r2norm` returns random numbers

**Author(s)**

Marc Girondot <[marc.girondot@gmail.com](mailto:marc.girondot@gmail.com)>

**See Also**

Other Distributions: `cutter()`, `dbeta_new()`, `dcutter()`, `dggamma()`, `logLik.cutter()`, `pSnbino`(`m`), `plot.cutter()`, `print.cutter()`, `qSnbino`(`m`), `rcutter()`, `rnnorm()`, `rnbino`\_new()

**Examples**

```
## Not run:
n <- r2norm(1000, mean=25, sd_low=2, sd_high=10)

hist(n)

## End(Not run)
```

**Description**

If it is very long, use silent parameter to check if something goes wrong.  
 If replicates is null or is 0, or if method is NULL, parameters are just copied into data.frame.

**Usage**

```
RandomFromHessianOrMCMC(
  se = NULL,
  Hessian = NULL,
  mcmc = NULL,
  chain = 1,
  regularThin = TRUE,
  MinMax = NULL,
  fitted.parameters = NULL,
  fixed.parameters = NULL,
  method = NULL,
  probs = c(0.025, 0.5, 0.975),
  replicates = 10000,
  fn = NULL,
  silent = FALSE,
  ParTofn = "par",
  ...
)
```

**Arguments**

<code>se</code>	A named vector with SE of parameters
<code>Hessian</code>	An Hessian matrix
<code>mcmc</code>	A result from <code>MHalgogen()</code>
<code>chain</code>	MCMC chain to be used
<code>regularThin</code>	If TRUE, use regular thin for MCMC
<code>MinMax</code>	A data.frame with at least two columns: Min and Max and rownames being the variable names
<code>fitted.parameters</code>	The fitted parameters
<code>fixed.parameters</code>	The fixed parameters
<code>method</code>	Can be <code>NULL</code> , "SE", "Hessian", "MCMC", or "PseudoHessianFromMCMC"
<code>probs</code>	Probability for quantiles
<code>replicates</code>	Number of replicates to generate the randoms
<code>fn</code>	The function to apply to each replicate
<code>silent</code>	Should the function display some information
<code>ParTofn</code>	Name of the parameter to send random values to <code>fn</code>
<code>...</code>	Parameters send to <code>fn</code> function

**Details**

`RandomFromHessianOrMCMC` returns random numbers based on Hessian matrix or MCMC

**Value**

Returns a list with three data.frames named random, fn, and quantiles

**Author(s)**

Marc Girondot <marc.girondot@gmail.com>

**Examples**

```
## Not run:
library(HelpersMG)
val <- rnorm(100, mean=20, sd=5)+(1:100)/10
# Return -ln L of values in val in Gaussian distribution with mean and sd in par
fitnorm <- function(par, data) {
  -sum(dnorm(data, par["mean"], abs(par["sd"])), log = TRUE)}
}
# Initial values for search
p<-c(mean=20, sd=5)
# fit the model
result <- optim(par=p, fn=fitnorm, data=val, method="BFGS", hessian=TRUE)
# Using Hessian
df <- RandomFromHessianOrMCMC(Hessian=result$hessian,
                                 fitted.parameters=result$par,
                                 method="Hessian")$random
hist(df[, 1], main="mean")
hist(df[, 2], main="sd")
plot(df[, 1], df[, 2], xlab="mean", ylab="sd", las=1, bty="n")

# Using MCMC
parameters_mcmc <- data.frame(Density=c('dnorm', 'dlnorm'),
Prior1=c(10, 0.5), Prior2=c(2, 0.5), SDProp=c(0.35, 0.2),
Min=c(-3, 0), Max=c(100, 10), Init=c(10, 2), stringsAsFactors = FALSE,
row.names=c('mean', 'sd'))
# Use of trace and traceML parameters
# trace=1 : Only one likelihood is printed
mcmc_run <- MHalgoGen(n.iter=50000, parameters=parameters_mcmc, data=val,
parameters_name = "par",
likelihood=fitnorm, n.chains=1, n.adapt=100, thin=1, trace=1)
df <- RandomFromHessianOrMCMC(mcmc=mcmc_run, fitted.parameters=NULL,
method="MCMC")$random
hist(df[, 1], main="mean")
hist(df[, 2], main="sd")
plot(df[, 1], df[, 2], xlab="mean", ylab="sd", las=1, bty="n")

# Using a function fn
fitnorm <- function(par, data, x) {
  y=par["a"]*(x)+par["b"]
  -sum(dnorm(data, y, abs(par["sd"])), log = TRUE)}
}
p<-c(a=0.1, b=20, sd=5)
# fit the model
x <- 1:100
```

```

result <- optim(par=p, fn=fitnorm, data=val, x=x, method="BFGS", hessian=TRUE)
# Using Hessian
df <- RandomFromHessianOrMCMC(Hessian=result$hessian, fitted.parameters=result$par,
                                    method="Hessian",
                                    fn=function(par) (par["a"]*(x)+par["b"]))
plot(1:100, val)
lines(1:100, df$quantiles["50%", ])
lines(1:100, df$quantiles["2.5%", ], lty=2)
lines(1:100, df$quantiles["97.5%", ], lty=2)

## End(Not run)

```

rcutter

*Random values of unobserved values of cut distribution.*

## Description

Return n random numbers.

It can be used to get the posterior predictive distribution; see example.

If random\_method is "ML", the parameter values obtained using maximum likelihood are used.

If random\_method is "medianMCMC", the parameter values obtained using median of posterior distribution are used.

If random\_method is "MCMC", the parameter values are one sample of the MCMC posterior distribution.

if observed\_detection\_limit is set to TRUE, the number of random number is equal to the number of observations; n is not used.

rcutter is the abbreviation for random-cutter.

## Usage

```

rcutter(
  cutter = stop("A result of cutter() must be provided"),
  n = 1,
  lower_detection_limit = NULL,
  upper_detection_limit = NULL,
  method_cut = c("censored", "truncated"),
  observed_detection_limit = FALSE,
  random_method = c("medianMCMC", "MCMC", "ML"),
  index_mcmc = NULL
)

```

## Arguments

- |                       |   |
|-----------------------|---|
| cutter                | The fitted model obtained with cutter() |
| n                     | number of random numbers                |
| lower_detection_limit | The lower detection limit               |

```

upper_detection_limit
  The upper detection limit
method_cut      What method is used to cut the distribution: "censored", "truncated"?
observed_detection_limit
  If TRUE, will use the pattern of detection limit as in observations
random_method   How to get parameters; it can be "ML", "medianMCMC", or "MCMC"
index_mcmc      For MCMC random_method, the index of data to be used.

```

## Details

rcutter returns random values based on fitted distribution with cut.

## Value

A vector with the random numbers.

## Author(s)

Marc Girondot <[marc.girondot@gmail.com](mailto:marc.girondot@gmail.com)>

## See Also

Other Distributions: [cutter\(\)](#), [dbeta\\_new\(\)](#), [dcutter\(\)](#), [dggamma\(\)](#), [logLik.cutter\(\)](#), [pSnbino](#)[m\(\)](#), [plot.cutter\(\)](#), [print.cutter\(\)](#), [qSnbino](#)[m\(\)](#), [r2norm\(\)](#), [rmnorm\(\)](#), [rnbinom\\_new\(\)](#)

## Examples

```

## Not run:
library(HelpersMG)
# -----
# right censored distribution with gamma distribution
# -----
# Detection limit
DL <- 100
# Generate 100 random data from a gamma distribution
obc <- rgamma(100, scale=20, shape=2)
# remove the data below the detection limit
obc[obc>DL] <- +Inf
# search for the parameters the best fit these censored data
result <- cutter(observations=obc, upper_detection_limit=DL,
                  cut_method="censored")
result
# Posterior predictive distribution
r <- rcutter(cutter=result, upper_detection_limit=DL, n=100)
hist(r)
# -----
# left censored distribution with gamma distribution
# -----
# Detection limit
DL <- 10
# Generate 100 random data from a gamma distribution

```

```

obc <- rgamma(100, scale=20, shape=2)
# remove the data below the detection limit
obc[obc<DL] <- -Inf
# search for the parameters the best fit these truncated data
result <- cutter(observations=obc, lower_detection_limit=DL,
                  cut_method="censored")
result
plot(result, breaks=seq(from=0, to=200, by=10))
r <- rcutter(cutter=result, n=100)
hist(r, breaks=seq(from=0, to=200, by=10))
r <- rcutter(cutter=result, lower_detection_limit=DL, n=100)
hist(r, breaks=seq(from=0, to=250, by=10))
# With censored method, some values are replaced with +Inf or -Inf
any(is.infinite(r))
r <- rcutter(cutter=result, upper_detection_limit=DL, n=100,
              method_cut="truncated")
# With truncated method, the values below LDL or upper UDL are not present
any(is.infinite(r))
hist(r, breaks=seq(from=0, to=10, by=0.25))
r <- rcutter(cutter=result, observed_detection_limit=TRUE)
hist(r, breaks=seq(from=0, to=300, by=10))

## End(Not run)

```

**read\_folder**

*Read files present in a folder and creates a list with the content of these files*

**Description**

To create a list, the syntax is:

```
datalist <- read_folder(folder=".", read=read.delim, header=FALSE)
```

It returns an error if the folder does not exist.

The names of the elements of the list are the filenames.

The parameter file can be used to predefined a list of file. If file is NULL, all the files of the folder/directory are used.

**Usage**

```

read_folder(
  folder = try(file.choose(), silent = TRUE),
  file = NULL,
  wildcard = "*.*",
  read = read.delim,
  ...
)

```

**Arguments**

folder	Where to search for files; can be or a file path or a folder path
file	list of files
wildcard	Define which files are to be read (examples: "*.*", "*.xls", "essai*.txt"). It can be also a vector with all filenames.
read	Function used to read file. Ex: read.delim or read.xls from gdata package
...	Parameters send to the read function

**Details**

read\_folder reads all files present in a folder

**Value**

Return a list with the data in the files of the folder (directory for windows users)

**Author(s)**

Marc Girondot <marc.girondot@gmail.com>

**Examples**

```
## Not run:
library(HelpersMG)
# Read all the .csv files from the current folder/directory
contentaslist <- read_folder(folder=".", wildcard=".csv", read=read.csv2)
# Read all the files from the current folder/directory
contentaslist <- read_folder(folder=".", wildcard="*.*", read=read.csv2)
# Read two files from the current folder/directory
files <- c("filename1.csv", "filename2.csv")
contentaslist <- read_folder(folder=".", wildcard=files, read=read.csv2)

## End(Not run)
```

RectangleRegression    *Return parameters of rectangle regression*

**Description**

Fit a line using least rectangle method.

**Usage**

```
RectangleRegression(
  x1,
  x2,
  replicate = 1000,
  x1new = seq(from = min(x1), to = max(x1), length.out = 100)
)
```

**Arguments**

x1	The first series of data
x2	The second series of data
replicate	Number of replicates for bootstrap
x1new	Values for x1 to generate x2

**Details**

RectangleRegression performs rectangle regression

**Value**

A list with parameters of rectangle regression

**Author(s)**

Marc Girondot <marc.girondot@gmail.com>

**Examples**

```

x1 <- runif(100, min=10, max=20)
x2 <- runif(100, min=10, max=20)+x1

rectreg <- RectangleRegression(x1, x2)

plot(x=x1, y=x2, bty="n", las=1, xlim=c(10, 20), ylim=c(20, 40))
abline(a=rectreg$par["Intercept"], b=rectreg$par["Slope"], lwd=2)
par(xpd=FALSE)
lines(rectreg$x2new["x1new", ], rectreg$x2new["50%", ])
lines(rectreg$x2new["x1new", ], rectreg$x2new["2.5%", ], lty=2)
lines(rectreg$x2new["x1new", ], rectreg$x2new["97.5%", ], lty=2)

abline(a=rectreg$Intercept[1], b=rectreg$Slope[3], col="red")
abline(a=rectreg$Intercept[3], b=rectreg$Slope[1], col="red")

```

**Description**

*rmnorm* generate random numbers from a multivariate normal distribution.

**Usage**

```
rmnorm(n = 1, mean = rep(0, d), varcov)
```

## Arguments

n	the number of random vectors to be generated.
mean	a vector with means of length d.
varcov	a variance-covariance matrix with dimentions d * d.

## Details

rmnrm Generate random numbers from the multivariate normal distribution

## Value

For n > 1 rmnrm returns a matrix of n rows of random vectors, while for n = 1 rmnrm returns a named random vector.

## Author(s)

Based on lmf package

## See Also

Other Distributions: [cutter\(\)](#), [dbeta\\_new\(\)](#), [dcutter\(\)](#), [dggamma\(\)](#), [logLik.cutter\(\)](#), [pSnbino](#)(), [plot.cutter\(\)](#), [print.cutter\(\)](#), [qSnbino](#)(), [r2norm\(\)](#), [rcutter\(\)](#), [rnbinom\\_new\(\)](#)

## Examples

```
## Not run:
#Variance-covariance matrix
varcov <- matrix(c(2.047737e-03, 3.540039e-05, 0.0075178920, 3.540039e-05,
6.122832e-07, 0.0001299661, 7.517892e-03, 1.299661e-04, 0.0276005740), ncol = 3)
#Set names
nam <- c("a", "b", "c")
dimnames(varcov) <- list(nam, nam)
#Check positive definiteness (all positive eigenvalues = positive definite)
eigen(varcov) $values
#Mean
mean <- c(1, 0.3, 0.5)
#Generate n = 1 random vector
rmnrm(n = 1, mean = mean, varcov = varcov)
#Generate n = 10 random vectors
rmnrm(n = 10, mean = mean, varcov = varcov)
#Generate n = 1 random vectors when varcov is non-positive definite
#Non-positive definite varcov matrix
varcov2 <- matrix(c(2.04e-03, 3.54e-05, 7.52e-03, 3.54e-05, 6.15e-07,
1.30e-04, 7.52e-03, 1.30e-04, 2.76e-02), ncol = 3)
dimnames(varcov2) <- dimnames(varcov)
eigen(varcov2)
#Random vector
rmnrm(n = 1, mean = mean, varcov = varcov2)

## End(Not run)
```

---

RM_add	<i>Create a results managment or add a value in a results managment to an object</i>
--------	--

---

## Description

Return original object with a new value or a new results managment.

## Usage

```
RM_add(
  x = stop("An object with results managment must be provided"),
  RM = "RM",
  RMname = stop("A results managment name must be provided"),
  valuename = NULL,
  value = NULL
)
```

## Arguments

x	The object to add a results managment or a result in a results managment
RM	The name of results managment stored
RMname	The name of the results managment to be modified or created
valuename	The name of the new value to be added
value	The value to be added

## Details

RM\_add adds a results managment or a value in results managment to an object

## Value

The original object with a new value in a results managment object or a new results managment

## Author(s)

Marc Girondot <marc.girondot@gmail.com>

## See Also

Other Results Management: [RM\\_delete\(\)](#), [RM\\_duplicate\(\)](#), [RM\\_get\(\)](#), [RM\\_list\(\)](#)

## Examples

```
## Not run:
library("HelpersMG")
# Let an object of class objclass being created
obj <- list(A=100, name="My object")
class(obj) <- "objclass"
# And now I create a RM to this object
obj <- RM_add(x=obj, RMname="NewAnalysis1")
RM_list(obj)
obj <- RM_add(x=obj, RMname="NewAnalysis2")
RM_list(obj)
obj <- RM_add(x=obj, RMname="NewAnalysis2", valuename="V1", value=100)
RM_get(x=obj, RMname="NewAnalysis2", valuename="V1")
obj <- RM_add(x=obj, RMname="NewAnalysis2", valuename="V1", value=200)
RM_get(x=obj, RMname="NewAnalysis2", valuename="V1")
obj <- RM_add(x=obj, RMname="NewAnalysis2", valuename="V2", value=300)
RM_get(x=obj, RMname="NewAnalysis2", valuename="V2")
RM_list(obj)

## End(Not run)
```

**RM\_delete**

*Delete a results managment or a result within a results managment from an object*

## Description

Return the original object with the deleted results managment or result.

## Usage

```
RM_delete(
  x = stop("An object with results managment must be provided"),
  RM = "RM",
  RMname = stop("A name must be provided"),
  valuename = NULL
)
```

## Arguments

x	The object to delete a results managment
RM	The name of results managment stored
RMname	The name of the result that will be deleted or its rank
valuename	The name of the result that will be deleted

## Details

RM\_delete deletes a results managment or a result within a results managment from an object

**Value**

The original object with the deleted results management

**Author(s)**

Marc Girondot <marc.girondot@gmail.com>

**See Also**

Other Results Management: [RM\\_add\(\)](#), [RM\\_duplicate\(\)](#), [RM\\_get\(\)](#), [RM\\_list\(\)](#)

**Examples**

```
## Not run:
library("HelpersMG")
# Let an object of class objclass being created
obj <- list(A=100, name="My object")
class(obj) <- "objclass"
# And now I create a RM to this object
obj <- RM_add(x=obj, RMname="NewAnalysis1")
obj <- RM_add(x=obj, RMname="NewAnalysis2")
RM_list(obj)
obj <- RM_delete(x=obj, RMname="NewAnalysis1")
RM_list(obj)
obj <- RM_delete(x=obj, RMname=1)
RM_list(obj)
obj <- RM_add(x=obj, RMname="NewAnalysis1", valuename="V1", value=100)
RM_list(obj)
RM_get(x=obj, RMname="NewAnalysis1", valuename="V1")
obj <- RM_add(x=obj, RMname="NewAnalysis1", valuename="V2", value=200)
RM_get(x=obj, RMname="NewAnalysis1", valuename="V2")
obj <- RM_delete(x=obj, RMname="NewAnalysis1", valuename="V1")
RM_get(x=obj, RMname="NewAnalysis1", valuename="V1")
RM_get(x=obj, RMname="NewAnalysis1", valuename="V2")

## End(Not run)
```

**RM\_duplicate**

*Duplicate a results management within an object.*

**Description**

**RM\_duplicate** duplicates a results management within an object.

**Usage**

```
RM_duplicate(  
  x = stop("An object with results management must be provided"),  
  RM = "RM",  
  RMnamefrom = 1,  
  RMnameto = 2  
)
```

**Arguments**

x	The object to duplicate a results management
RM	The name of results management stored
RMnamefrom	The name of the results management to be duplicated
RMnameto	The new name of the results management

**Details**

RM\_duplicate duplicates a results management within an object

**Value**

The original object with a duplicated results management.

**Author(s)**

Marc Girondot <marc.girondot@gmail.com>

**See Also**

Other Results Management: [RM\\_add\(\)](#), [RM\\_delete\(\)](#), [RM\\_get\(\)](#), [RM\\_list\(\)](#)

**Examples**

```
## Not run:  
library("HelpersMG")  
# Let an object of class objclass being created  
obj <- list(A=100, name="My object")  
class(obj) <- "objclass"  
# And now I create a RM to this object  
obj <- RM_add(x=obj, RMname="NewAnalysis1")  
RM_list(obj)  
obj <- RM_duplicate(x=obj, RMnamefrom="NewAnalysis1", RMnameto="NewAnalysis2")  
RM_list(obj)  
  
## End(Not run)
```

RM\_get

*Get a value in a results managment to an object***Description**

Return the value valuename of the results managment RMname.

**Usage**

```
RM_get(
  x = stop("An object with results managment must be provided"),
  RM = "RM",
  RMname = stop("A results managment name must be provided"),
  valuename = NULL
)
```

**Arguments**

x	The object in which to get a result in a results managment
RM	The name of results managment stored
RMname	The name of the results managment to be read
valuename	The name of the value to be read

**Details**

RM\_get gets a value in results managment to an object

**Value**

Return a value in a results managment object

**Author(s)**

Marc Girondot <marc.girondot@gmail.com>

**See Also**

Other Results Managment: [RM\\_add\(\)](#), [RM\\_delete\(\)](#), [RM\\_duplicate\(\)](#), [RM\\_list\(\)](#)

**Examples**

```
## Not run:
library("HelpersMG")
# Let an object of class objclass being created
obj <- list(A=100, name="My object")
class(obj) <- "objclass"
# And now I create a RM to this object
obj <- RM_add(x=obj, RMname="NewAnalysis1")
```

```
RM_list(obj)
obj <- RM_add(x=obj, RMname="NewAnalysis2")
RM_list(obj)
obj <- RM_add(x=obj, RMname="NewAnalysis2", valuename="V1", value=100)
RM_get(x=obj, RMname="NewAnalysis2", valuename="V1")

## End(Not run)
```

---

**RM\_list**

*Return the list of results managment of an object.*

---

**Description**

RM\_list returns the list of results managment of an object.

**Usage**

```
RM_list(
  x = stop("An object with results managment must be provided"),
  RM = "RM",
  silent = FALSE,
  max.level = FALSE
)
```

**Arguments**

x	The object to add a results managment
RM	The name of results managment stored
silent	Should the results be shown ?
max.level	If TRUE, will return all list element of the objects

**Details**

RM\_list returns the list of results managment of an object

**Value**

A list with the names of results stored in an object

**Author(s)**

Marc Girondot <marc.girondot@gmail.com>

**See Also**

Other Results Management: [RM\\_add\(\)](#), [RM\\_delete\(\)](#), [RM\\_duplicate\(\)](#), [RM\\_get\(\)](#)

## Examples

```
## Not run:
library("HelpersMG")
# Let an object of class objclass being created
obj <- list(A=100, name="My object")
class(obj) <- "objclass"
# And now I create a RM to this object
obj <- RM_add(x=obj, RMname="NewAnalysis1")
RM_list(obj)
obj <- RM_add(x=obj, RMname="NewAnalysis2")
RM_list(obj)
obj <- RM_add(x=obj, RMname="NewAnalysis2", valuename="V1", value=100)
RM_get(x=obj, RMname="NewAnalysis2", valuename="V1")
obj <- RM_add(x=obj, RMname="NewAnalysis2", valuename="V1", value=200)
RM_get(x=obj, RMname="NewAnalysis2", valuename="V1")
obj <- RM_add(x=obj, RMname="NewAnalysis2", valuename="V2", value=300)
RM_get(x=obj, RMname="NewAnalysis2", valuename="V2")
RM_list(obj)
rmlist <- RM_list(obj, max.level=TRUE)
rmlist

## End(Not run)
```

**rnbinom\_new**

*Random numbers for the negative binomial distribution.*

## Description

See `rnbinom`.

## Usage

```
rnbinom_new(n, size = NULL, prob = NULL, mu = NULL, sd = NULL, var = NULL)
```

## Arguments

<code>n</code>	number of observations.
<code>size</code>	target for number of successful trials, or dispersion parameter (the shape parameter of the gamma mixing distribution). Must be strictly positive, need not be integer.
<code>prob</code>	probability of success in each trial. $0 < \text{prob} \leq 1$ .
<code>mu</code>	alternative parametrization via mean.
<code>sd</code>	alternative parametrization via standard deviation.
<code>var</code>	alternative parametrization via variance.

## Details

`rnbinom_new` returns random numbers for the negative binomial distribution

**Value**

Random numbers for the negative binomial distribution

**Author(s)**

Marc Girondot <marc.girondot@gmail.com>

**See Also**

Other Distributions: [cutter\(\)](#), [dbeta\\_new\(\)](#), [dcutter\(\)](#), [dggamma\(\)](#), [logLik.cutter\(\)](#), [pSnbnom\(\)](#), [plot.cutter\(\)](#), [print.cutter\(\)](#), [qSnbnom\(\)](#), [r2norm\(\)](#), [rcutter\(\)](#), [rmnorm\(\)](#)

**Examples**

```
## Not run:  
library("HelpersMG")  
set.seed(1)  
x <- rnbinom_new(n=1000, prob=6.25/(5+6.25), size=6.25)  
mean(x)  
sd(x)  
set.seed(1)  
x <- rnbinom_new(n=1000, mu=5, sd=3)  
mean(x)  
sd(x)  
set.seed(1)  
x <- rnbinom_new(n=1000, mu=5, var=3^2)  
mean(x)  
sd(x)  
set.seed(1)  
x <- rnbinom_new(n=1000, mu=5, size=6.25)  
mean(x)  
sd(x)  
set.seed(1)  
x <- rnbinom_new(n=1000, size=6.25, var=3^2)  
mean(x)  
sd(x)  
set.seed(1)  
x <- rnbinom_new(n=1000, prob=6.25/(5+6.25), var=3^2)  
mean(x)  
sd(x)  
# Example of wrong parametrization  
set.seed(1)  
x <- rnbinom_new(n=1000, sd=3, var=3^2)  
set.seed(1)  
x <- rnbinom_new(n=1000, mu=10, var=3^2)  
  
## End(Not run)
```

**rSnb***n**om**Random generation for the sum of random variable with negative binomial distributions.***Description**

Random numbers for the sum of random variable with negative binomial distributions.

**Usage**

```
rSnb(n = 1, size = NULL, prob = NULL, mu = NULL)
```

**Arguments**

<code>n</code>	number of observations.
<code>size</code>	target for number of successful trials, or dispersion parameter (the shape parameter of the gamma mixing distribution). Must be strictly positive, need not be integer.
<code>prob</code>	probability of success in each trial. $0 < \text{prob} \leq 1$ .
<code>mu</code>	alternative parametrization via mean.

**Details**

`rSnb` returns random numbers for the sum of random variable with negative binomial distributions

**Value**

`rSnb` returns random number

**Author(s)**

Marc Girondot <[marc.girondot@gmail.com](mailto:marc.girondot@gmail.com)>

**See Also**

Other Distribution of sum of random variable with negative binomial distributions: [dSnb](#)*n**om*()

**Examples**

```
## Not run:
alpha <- c(2.1, 2.05, 2)
mu <- c(10, 30, 20)
rep <- 100000
distEmpirique <- rSnb(n=rep, size=alpha, mu=mu)
tabledistEmpirique <- rep(0, 301)
names(tabledistEmpirique) <- as.character(0:300)
tabledistEmpirique[names(table(distEmpirique))] <- table(distEmpirique)/rep
```

```
plot(0:300, dSnbinom(0:300, size=alpha, mu=mu), type="h", bty="n",
     xlab="x", ylab="Density", ylim=c(0,0.02))
plot_add(0:300, tabledistEmpirique, type="l", col="red")
legend(x=200, y=0.02, legend=c("Empirical", "Theoretical"),
       text.col=c("red", "black"), bty="n")

## End(Not run)
```

---

ScalePreviousPlot      *Return the scale of the previous plot*

---

## Description

Return a list with the limits of the previous plot, the center, the range, and the position of label on this axe.

## Usage

```
ScalePreviousPlot(x = NULL, y = NULL)
```

## Arguments

- |   |  |
|---|--|
| x | The position in x as relative position |
| y | The position in y as relative position |

## Details

ScalePreviousPlot returns the scale of the previous plot

## Value

A list with xlim and ylim

## Author(s)

Marc Girondot <marc.girondot@gmail.com>

## See Also

Other plot and barplot functions: [barplot\\_errbar\(\)](#), [plot\\_add\(\)](#), [plot\\_errbar\(\)](#), [show\\_name\(\)](#)

## Examples

```

## Not run:
par(xaxs="i", yaxs="i")
plot(x=1:100, y=sin(1:100), type="l", bty="n", xlim=c(1,200), xlab="x", ylab="y")
xlim= ScalePreviousPlot()$xlim[1:2]
ylim= ScalePreviousPlot()$ylim[1:2]
par(xaxs="r", yaxs="i")
plot(x=1:100, y=sin(1:100), type="l", bty="n", xlim=c(1,200), xlab="x", ylab="y")
xlim= ScalePreviousPlot()$xlim[1:2]
ylim= ScalePreviousPlot()$ylim[1:2]
# Here is an example of the use of the label output
plot(x=1:100, y=sin(1:100), type="l", bty="n", xlim=c(1,200), xlab="", ylab="")
text(x=ScalePreviousPlot()$xlim["label"], y=ScalePreviousPlot()$ylim["center"],
     xpd=TRUE, "Legend for Y axes", pos=3, srt=90)
text(x=ScalePreviousPlot()$xlim["center"], y=ScalePreviousPlot()$ylim["label"],
     xpd=TRUE, "Legend for X axes", pos=1)
Example to plot legend always in the same place
layout(1:2)
plot(x=1:100, y=sin(1:100), type="l", bty="n", xlim=c(1,200), xlab="", ylab="")
text(x=ScalePreviousPlot(x=0.95, y=0.05)$x,
     y=ScalePreviousPlot(x=0.95, y=0.05)$y,
     labels="A", cex=2)
plot(x=0:1, y=0:1, type="p", bty="n")
text(x=ScalePreviousPlot(x=0.95, y=0.05)$x,
     y=ScalePreviousPlot(x=0.95, y=0.05)$y,
     labels="B", cex=2)

## End(Not run)

```

## Description

Standard error of parameters based on Hessian matrix.

The strategy is as follow:

First it tries to inverse the Hessian matrix. If it fails, it uses the near positive definite matrix of the Hessian.

So now the inverse of the Hessian matrix can be computed.

The diagonal of the inverse of the Hessian matrix is calculated. If all values are positive, the SEs are the square root of the inverse of the Hessian.

If not all values are positive, it will estimate the pseudo-variance matrix based on Gill & King (2004). It necessitates a Cholesky matrix.

If from some reason it fails (for example all SE are 0 in output), then the strategy of Rebonato and Jackel (2000) will be used to generate the Cholesky matrix.

## Usage

```
SEfromHessian(a, hessian = FALSE, silent = FALSE)
```

### Arguments

a	An Hessian matrix
hessian	If TRUE, return a list with the hessian and SE
silent	If TRUE, report some problems

### Details

SEfromHessian returns standard error of parameters based on Hessian matrix

### Value

SEfromHessian returns a vector with standard errors

### Author(s)

Marc Girondot <marc.girondot@gmail.com>

### References

Gill J. and G. King 2004. What to do when your Hessian is not invertible: Alternatives to model respecification in nonlinear estimation. *Sociological Methods & Research* 33: 54-87.

Rebonato and Jackel, “The most general methodology for creating a valid correlation matrix for risk management and option pricing purposes”, *Journal of Risk*, Vol 2, No 2, 2000.

### Examples

```
## Not run:
val=rnorm(100, mean=20, sd=5)
# Return -ln L of values in val in Gaussian distribution with mean and sd in par
fitnorm<-function(par, val) {
  -sum(dnorm(val, par["mean"], par["sd"], log = TRUE))
}
# Initial values for search
p<-c(mean=20, sd=5)
# fit the model
result <- optim(par=p, fn=fitnorm, val=val, method="BFGS", hessian=TRUE)
SE <- SEfromHessian(result$hessian)
library(MASS)
fitdistr(val, densfun = "normal")

## End(Not run)
```

`series.compare`      *Data series comparison using Akaike weight*

## Description

This function is used as a replacement of `t.test()` to not use p-value.

## Usage

```
series.compare(..., criterion = c("BIC", "AIC", "AICc"), var.equal = TRUE)
```

## Arguments

...	Series of data (at least two or data are in a table with series in different rows)
criterion	Which criterion is used for model selection. can be AIC, AICc or BIC
var.equal	Should the variances of all series being equal? Default TRUE

## Details

`series.compare` compares series of data using Akaike weight.

## Value

The probability that a single proportion model is sufficient to explain the data

## Author(s)

Marc Girondot <[marc.girondot@gmail.com](mailto:marc.girondot@gmail.com)>

## References

Girondot, M., Guillon, J.-M., 2018. The w-value: An alternative to t- and X<sup>2</sup> tests. Journal of Biostatistics & Biometrics 1, 1-4.

## See Also

Other w-value functions: [compare\(\)](#), [contingencyTable.compare\(\)](#)

## Examples

```
## Not run:
library("HelpersMG")
A <- rnorm(100, 10, 2)
B <- rnorm(100, 11.1, 2)
series.compare(A, B, criterion = "BIC", var.equal=TRUE)
B <- B[1:10]
series.compare(A, B, criterion = "BIC", var.equal=TRUE)
A <- rnorm(100, 10, 2)
```

```
B <- rnorm(100, 10.1, 2)
C <- rnorm(100, 10.5, 2)
series.compare(A, B, C, criterion = "BIC", var.equal=TRUE)
B <- B[1:10]
series.compare(A, B, criterion = "BIC", var.equal=TRUE)
t.test(A, B, var.equal=TRUE)
# Example with a data.frame
series.compare(t(data.frame(A=c(10, 27, 19, 20, NA), B=c(10, 20, NA, NA, NA))))
# Test in the context of big data
A <- rnorm(10000, 10, 2)
B <- rnorm(10000, 10.1, 2)
series.compare(A, B, criterion = "BIC", var.equal=TRUE)
t.test(A, B, var.equal=TRUE)
#####
w <- NULL
p <- NULL

for (i in 1:1000) {

  A <- rnorm(50000, 10, 2)
  B <- rnorm(50000, 10.01, 2)
  w <- c(w, unname(series.compare(A, B, criterion = "BIC", var.equal=TRUE)[1]))
  p <- c(p, t.test(A, B, var.equal=TRUE)$p.value)

}

layout(mat = 1:2)
par(mar=c(4, 4, 1, 1)+0.4)
hist(p, main="", xlim=c(0, 1), las=1, breaks = (0:20)/20,
     freq=FALSE, xlab = expression(italic("p")*"-value"))
hist(w, main="", xlim=c(0, 1), las=1, breaks = (0:20)/20,
     freq=FALSE, xlab = expression(italic("w")*"-value"))
#####

x <- seq(from=8, to=13, by=0.1)

pv <- NULL
aw <- NULL
A <- rnorm(100, mean=10, sd=2)
B <- A-2

for (meanB in x) {
  pv <- c(pv, t.test(A, B, var.equal = FALSE)$p.value)
  aw <- c(aw, series.compare(A, B, criterion="BIC", var.equal = FALSE)[1])
  B <- B + 0.1
}

par(mar=c(4, 4, 2, 1)+0.4)
y <- pv
plot(x=x, y=y, type="l", lwd=2,
      bty="n", las=1, xlab="Mean B value (SD = 4)", ylab="Probability", ylim=c(0,1),
      main="")
y2 <- aw
```

```

lines(x=x, y=y2, type="l", col="red", lwd=2)

l1 <- which(aw>0.05)[1]
l2 <- max(which(aw>0.05))

aw[l1]
pv[l1]

aw[l2]
pv[l2]

l1 <- which(pv>0.05)[1]
l2 <- max(which(pv>0.05))

aw[l1]
pv[l1]

aw[l2]
pv[l2]

par(xpd=TRUE)
segments(x0=10-1.96*2/10, x1=10+1.96*2/10, y0=1.1, y1=1.1, lwd=2)
segments(x0=10, x1=10, y0=1.15, y1=1.05, lwd=2)
par(xpd=TRUE)
text(x=10.5, y=1.1, labels = "Mean A = 10, SD = 2", pos=4)

v1 <- c(expression(italic("p")*"-value"), expression("based on "*italic("t")*"-test"))
v2 <- c(expression(italic("w")*"-value for A"), expression("and B identical models"))
legend("topright", legend=c(v1, v2),
      y.intersp = 1,
      col=c("black", "black", "red", "red"), bty="n", lty=c(1, 0, 1, 0))

segments(x0=min(x), x1=max(x), y0=0.05, y1=0.05, lty=2)
par(xpd = TRUE)
text(x=13.05, y=0.05, labels = "0.05", pos=4)

## End(Not run)

```

**setPriors***Set priors for MHalgoGen()***Description**

Set priors for MHalgoGen()

**Usage**

```
setPriors(
  par = stop("A vector with init values is necessary."),
```

```

    se = NULL,
    density = "dunif",
    rules = NULL,
    silent = FALSE
)

```

## Arguments

par	Named vector with init value of parameters
se	Named vector with standard error of parameters
density	Named vector with density or single value
rules	List of rules to define priors
silent	If TRUE, do not show warning.

## Details

setPriors is a general function to set priors for MHalgoGen()

## Value

Return a data.frame with priors

## Author(s)

Marc Girondot <marc.girondot@gmail.com>

## See Also

Other mcmcComposite functions: [MHalgoGen\(\)](#), [as.mcmc.mcmcComposite\(\)](#), [as.parameters\(\)](#), [as.quantiles\(\)](#), [merge.mcmcComposite\(\)](#), [plot.PriorsmcmcComposite\(\)](#), [plot.mcmcComposite\(\)](#), [summary.mcmcComposite\(\)](#)

## Examples

```

## Not run:
library(HelpersMG)
rules <- rbind(data.frame(Name="^a", Min=0, Max="x*2"),
                 data.frame(Name="^b", Min=0, Max=100))
par <- c(a0=10, a1=2, b2=20)
(p <- setPriors(par=par, se=NULL, density="dgamma", rules=rules))
(p <- setPriors(par=par, se=NULL, density="dnorm", rules=rules))
(p <- setPriors(par=par, se=NULL, density="dunif", rules=rules))
par <- c(a0=10, a1=2, b2=20, b1=-1)
(p <- setPriors(par=par, se=NULL, density="dgamma", rules=rules))

## End(Not run)

```

`show_name`      *Show the name of a point*

## Description

Click on a point in plot region and it will tell you what is the point.

## Usage

```
show_name(
  points = NULL,
  x = NULL,
  y = NULL,
  names = NULL,
  col = "red",
  silent = FALSE
)
```

## Arguments

<code>points</code>	A list with x, y and names elements
<code>x</code>	The x coordinates
<code>y</code>	The y coordinates.
<code>names</code>	The names of the points
<code>col</code>	Color of the legend.
<code>silent</code>	TRUE or FALSE

## Details

Show the name of a point

## Value

Name of the point

## Author(s)

Marc Girondot <[marc.girondot@gmail.com](mailto:marc.girondot@gmail.com)>

## See Also

[plot\\_errorbar](#)

Other plot and barplot functions: [ScalePreviousPlot\(\)](#), [barplot\\_errbar\(\)](#), [plot\\_add\(\)](#), [plot\\_errbar\(\)](#)

## Examples

```
## Not run:
k <- plot_errbar(1:100, rnorm(100, 1, 2),
xlab="axe x", ylab="axe y", bty="n", xlim=c(1,100),
errbar.x=2, errbar.y=rnorm(100, 1, 0.1))
show_name(k)
k <- plot_errbar(1:10, rnorm(10, 1, 2),
xlab="axe x", ylab="axe y", bty="n", xlim=c(1,10),
errbar.x=2, errbar.y=rnorm(10, 1, 0.1),
names=LETTERS[1:10])
show_name(k)
k <- plot_errbar(1:10, rnorm(10, 1, 2),
xlab="axe x", ylab="axe y", bty="n", xlim=c(1,10),
errbar.x=2, errbar.y=rnorm(10, 1, 0.1))
show_name(k, names=LETTERS[1:10])

## End(Not run)
```

**similar**

*Test if two vectors contains the same elements independently of their order*

## Description

Return TRUE only if all elements of x are present and only once in y.

## Usage

```
similar(x, y, test.names = FALSE)
```

## Arguments

x	A vector with numeric or character elements
y	A vector with numeric or character elements
test.names	Logical. If TRUE, the names of the vector elements must be also identical and unique

## Value

A logical TRUE or FALSE

## Author(s)

Marc Girondot <marc.girondot@gmail.com>

## Examples

```
## Not run:
A <- c("A", "B", "C", "D")
B <- c("A", "B", "C", "D")
similar(A, B)
similar(B, A)
A <- c(x="A", y="B", z="C", k="D")
B <- c(x="A", y="B", z="C", l="D")
similar(B, A)
similar(A, B, test.names=TRUE)
A <- c(x="A", y="B", z="C", k="D")
B <- c(x="A", z="C", k="D", y="B")
similar(B, A)
similar(A, B, test.names=TRUE)

## End(Not run)
```

### **specify\_decimal**

*Return a number as character with specified number of decimals*

## Description

Return a number as character with specified number of decimals. If *a* is a matrix, it will return a matrix of the same size and the same attributes.

## Usage

```
specify_decimal(x, decimals = NULL, decimal.point = ".")
```

## Arguments

- x*              The numbers to be formated
- decimals*       Number of decimals to print
- decimal.point* Character to be used as decimal point

## Details

*specify\_decimals* format a number with specified number of decimals

## Value

A character

## Author(s)

Marc Girondot <marc.girondot@gmail.com>

## Examples

```
specify_decimal(x=pi, decimal.point=".")  
specify_decimal(x=pi, decimals=4, decimal.point=".")  
specify_decimal(x=c(pi, exp(1)), decimals=3, decimal.point=",")  
specify_decimal(x=c(pi, exp(1)), decimal.point=",")  
specify_decimal(x=c(pi*10, pi, pi/10, pi/100, pi/1000))  
specify_decimal(x=c(pi=pi), decimal.point=".")  
specify_decimal(x=matrix(pi*1:4, ncol=2), decimal.point=".")  
m <- matrix(pi*1:4, ncol=2)  
rownames(m) <- c("A", "B")  
colnames(m) <- c("C", "D")  
specify_decimal(x=m, decimal.point=".")
```

**summary.mcmcComposite** *Summarize the result of a mcmcComposite object*

## Description

Summary for the result of a mcmcComposite object.

## Usage

```
## S3 method for class 'mcmcComposite'  
summary(object, chain = NULL, ...)
```

## Arguments

object	A mcmcComposite object
chain	The chain to use
...	Not used

## Details

**summary.mcmcComposite** get info on the result of a mcmcComposite object

## Value

A summary of the result

## Author(s)

Marc Girondot <marc.girondot@gmail.com>

## See Also

Other mcmcComposite functions: [MHalgoGen\(\)](#), [as.mcmc.mcmcComposite\(\)](#), [as.parameters\(\)](#), [as.quantiles\(\)](#), [merge.mcmcComposite\(\)](#), [plot.PriorsmcmcComposite\(\)](#), [plot.mcmcComposite\(\)](#), [setPriors\(\)](#)

## Examples

```

## Not run:
library(HelpersMG)
require(coda)
x <- rnorm(30, 10, 2)
dnormx <- function(data, x) {
  data <- unlist(data)
  return(-sum(dnorm(data, mean=x['mean'], sd=x['sd'], log=TRUE)))
}
parameters_mcmc <- data.frame(Density=c('dnorm', 'dlnorm'),
Prior1=c(10, 0.5), Prior2=c(2, 0.5), SDProp=c(1, 1),
Min=c(-3, 0), Max=c(100, 10), Init=c(10, 2), stringsAsFactors = FALSE,
row.names=c('mean', 'sd'))
mcmc_run <- MHalgoGen(n.iter=1000, parameters=parameters_mcmc, data=x,
likelihood=dnormx, n.chains=1, n.adapt=100, thin=1, trace=1)
plot(mcmc_run, xlim=c(0, 20))
plot(mcmc_run, xlim=c(0, 10), parameters="sd")
mcmcforcoda <- as.mcmc(mcmc_run)
#' heidel.diag(mcmcforcoda)
raftery.diag(mcmcforcoda)
autocorr.diag(mcmcforcoda)
acf(mcmcforcoda[[1]][,"mean"], lag.max=20, bty="n", las=1)
acf(mcmcforcoda[[1]][,"sd"], lag.max=20, bty="n", las=1)
batchSE(mcmcforcoda, batchSize=100)
# The batch standard error procedure is usually thought to
# be not as accurate as the time series methods used in summary
summary(mcmcforcoda)$statistics[,"Time-series SE"]
summary(mcmc_run)
as.parameters(mcmc_run)
lastp <- as.parameters(mcmc_run, index="last")
parameters_mcmc[,"Init"] <- lastp
# The n.adapt set to 1 is used to not record the first set of parameters
# then it is not duplicated (as it is also the last one for
# the object mcmc_run)
mcmc_run2 <- MHalgoGen(n.iter=1000, parameters=parameters_mcmc, data=x,
likelihood=dnormx, n.chains=1, n.adapt=1, thin=1, trace=1)
mcmc_run3 <- merge(mcmc_run, mcmc_run2)
##### no adaptation, n.adapt must be 0
parameters_mcmc[,"Init"] <- c(mean(x), sd(x))
mcmc_run3 <- MHalgoGen(n.iter=1000, parameters=parameters_mcmc, data=x,
likelihood=dnormx, n.chains=1, n.adapt=0, thin=1, trace=1)

## End(Not run)

```

## Description

Estimate the sun fates according to latitude and date.

Can be compared with the function `sunrise.set()` of package `StreamMetabolism`.

## Usage

```
sun.info(date, latitude, longitude)
```

## Arguments

<code>date</code>	A vector with the time at which sun fates are needed
<code>latitude</code>	The latitude at which estimate the sun fates
<code>longitude</code>	The longitude at which estimate the sun fates

## Details

`sun.info` estimate the time of sunrise and sunset according to longitude, latitude and date

## Value

A data.frame with information about daily sun

## Author(s)

Marc Girondot <[marc.girondot@gmail.com](mailto:marc.girondot@gmail.com)>

## References

Teets, D.A. 2003. Predicting sunrise and sunset times. *The College Mathematics Journal* 34(4):317-321.

## See Also

Other Periodic patterns of indices: `index.periodic()`, `minmax.periodic()`, `moon.info()`, `tide.info()`

## Examples

```
## Not run:
# Generate a timeserie of time
date <- seq(from=as.Date("2000-01-01"), to=as.Date("2000-12-31"), by="1 day")
plot(date, sun.info(date, latitude=23, longitude=0)$day.length, bty="n",
  las=1, type="l", xlab="Ordinal days", ylab="Day length in hours")
plot(date, sun.info(date, latitude=23, longitude=0)$sunrise, bty="n",
  las=1, type="l", xlab="Ordinal days", ylab="Sun rise in hours")

## End(Not run)
```

`symbol.Female`      *Plot a female symbol in the plotting region*

## Description

Plot a female symbol in the plotting region.

## Usage

```
symbol.Female(centerx, centery, rayonx, lwd = 2, col = "black")
```

## Arguments

<code>centerx</code>	The x position of the center of the circle
<code>centery</code>	The y position of the center of the circle
<code>rayonx</code>	The size of the rayon in the scale of the x axis
<code>lwd</code>	The width of the line of the symbol
<code>col</code>	The color of the symbol

## Details

`symbol.Female` plot a female symbol in the plotting region

## Value

Nothing

## Author(s)

Marc Girondot <[marc.girondot@gmail.com](mailto:marc.girondot@gmail.com)>

## See Also

Other Symbol: [symbol.Male\(\)](#)

## Examples

```
## Not run:
plot(x=1:2, y=c(10,20), type="n", bty="n", xlab="", ylab="")

rayonx <- 0.01
centerx <- 1.2
centery <- 15

symbol.Male(centerx=centerx, centery = centery, rayonx=rayonx)
symbol.Female(centerx=centerx+0.5, centery = centery, rayonx=rayonx)

rayonx <- 0.03
```

```
centerx <- 1.2
centery <- 18

symbol.Male(centerx=centerx, centery = centery, rayonx=rayonx, lwd=3)
symbol.Female(centerx=centerx+0.5, centery = centery, rayonx=rayonx, lwd=3, col="red")

rayonx <- 0.05
centerx <- 1.4
centery <- 13

symbol.Male(centerx=centerx, centery = centery, rayonx=rayonx, lwd=4, col="blue")
symbol.Female(centerx=centerx+0.5, centery = centery, rayonx=rayonx, lwd=4, col="red")

## End(Not run)
```

---

**symbol.Male**

*Plot a male symbol in the plotting region*

---

## Description

Plot a male symbol in the plotting region.

## Usage

```
symbol.Male(centerx, centery, rayonx, lwd = 2, col = "black")
```

## Arguments

centerx	The x position of the center of the circle
centery	The y position of the center of the circle
rayonx	The size of the rayon in the scale of the x axis
lwd	The width of the line of the symbol
col	The color of the symbol

## Details

`symbol.Male` plot a male symbol in the plotting region

## Value

Nothing

## Author(s)

Marc Girondot <[marc.girondot@gmail.com](mailto:marc.girondot@gmail.com)>

## See Also

Other Symbol: [symbol.Female\(\)](#)

## Examples

```
## Not run:
plot(x=1:2, y=c(10,20), type="n", bty="n", xlab="", ylab="")

rayonx <- 0.01
centerx <- 1.2
centery <- 15

symbol.Male(centerx=centerx, centery = centery, rayonx=rayonx)
symbol.Female(centerx=centerx+0.5, centery = centery, rayonx=rayonx)

rayonx <- 0.03
centerx <- 1.2
centery <- 18

symbol.Male(centerx=centerx, centery = centery, rayonx=rayonx, lwd=3)
symbol.Female(centerx=centerx+0.5, centery = centery, rayonx=rayonx, lwd=3, col="red")

rayonx <- 0.05
centerx <- 1.4
centery <- 13

symbol.Male(centerx=centerx, centery = centery, rayonx=rayonx, lwd=4, col="blue")
symbol.Female(centerx=centerx+0.5, centery = centery, rayonx=rayonx, lwd=4, col="red")

## End(Not run)
```

**symmetricize**

*Make a matrix symmetric*

## Description

This function was part of the package ENA. This package is no more available and it cannot be installed from archive because some dependencies are no more available.

## Usage

```
symmetricize(
  matrix,
  method = c("max", "min", "avg", "ld", "ud"),
  adjacencyList = FALSE
)
```

## Arguments

<b>matrix</b>	The matrix to make symmetric
<b>method</b>	The method to use to make the matrix symmetric. Default is to take the maximum.

- "max" For each position,  $m_{i,j}$ , use the maximum of  $(m_{i,j}, m_{j,i})$
- "min" For each position,  $m_{i,j}$ , use the minimum of  $(m_{i,j}, m_{j,i})$
- "avg" For each position,  $m_{i,j}$ , use the mean:  $(m_{i,j} + m_{j,i})/2$
- "ld" Copy the lower triangular portion of the matrix to the upper triangular portion.
- "ud" Copy the upper triangular portion of the matrix to the lower triangular portion.

**adjacencyList** Logical. If false, returns the symmetric matrix (the same format as the input). If true, returns an adjacency list representing the upper triangular portion of the adjacency matrix with addressing based on the row.names of the matrix provided.

## Details

Make the matrix symmetric by making all "mirrored" positions consistent. A variety of methods are provided to make the matrix symmetrical.

## Value

The symmetric matrix

## Author(s)

Jeffrey D. Allen <Jeffrey.Allen@UTSouthwestern.edu>

## Examples

```
#Create a sample 3x3 matrix
mat <- matrix(1:9, ncol=3)

#Copy the upper diagonal portion to the lower
symmetricize(mat, "ud")

#Take the average of each symmetric location
symmetricize(mat, "avg")
```

## Description

Annual tide information.

The columns are: Location, Longitude, Latitude, Phase, DateTime.local, DateTime.UTC, Tide.meter  
This function uses an API linking xtide software (<https://flaterco.com/xtide/>) with tide.info() function.

You must have a working internet connection for this function.

**Usage**

```
tide.info(
  location = NULL,
  year = 2021,
  longitude = NULL,
  latitude = NULL,
  force.tide.height = TRUE
)
```

**Arguments**

<code>location</code>	Textual information about location name
<code>year</code>	Year to get the calendar
<code>longitude</code>	Longitude to search for
<code>latitude</code>	Latitude to search for
<code>force.tide.height</code>	If FALSE, can return a current speed rather than tide height

**Details**

`tide.info` gets the annual tide calendar for one particular location.

**Value**

Return a `data.frame` with annual tide calendar.

**Author(s)**

Marc Girondot <[marc.girondot@gmail.com](mailto:marc.girondot@gmail.com)>

**See Also**

Other Periodic patterns of indices: `index.periodic()`, `minmax.periodic()`, `moon.info()`, `sun.info()`

**Examples**

```
## Not run:
library("HelpersMG")
Location <- "Les Hattes"
Year <- 2010
tide <- tide.info(Location, Year)
plot(tide[, "DateTime.local"], tide[, "Tide.meter"],
  type="l", bty="n", las=1,
  main=tide[1, "Location"],
  xlab=as.character(Year), ylab="Tide level in meter")

Location <- "Hawaii"
Year <- 2010
tide <- tide.info(Location, Year)
```

```

Location <- "Hanamaulu Bay, Kauai Island, Hawaii"
Year <- 2010
tide <- tide.info(Location, Year)
plot(tide[, "DateTime.local"], tide[, "Tide.meter"],
     type="l", bty="n", las=1,
     main=tide[1, "Location"],
     xlab=as.character(Year), ylab="Tide level in meter")

tide <- tide.info(year=2010, longitude=-32, latitude=-4)
library(maps)
map(database = "world", regions = "Brazil", asp=1,
     xlim=c(-80, -30), ylim=c(-33, 5))
points(tide[1, "Longitude"], tide[1, "Latitude"], col="red", pch=19)
points(-32, -4, col="blue", pch=19)
axis(1)
axis(2, las=1)

# Show the locations with data
library(maps)
map(xlim=c(-180, 180), ylim=c(-90, 90))
title("Locations with harmonics data")
axis(1, at=seq(from=-180, to=180, by=45))
axis(2, las=1, at=seq(from=-90, to=90, by=15))
points(getFromNamespace(x="tide_location", ns="HelpersMG")[, c("longitude")],
       getFromNamespace(x="tide_location", ns="HelpersMG")[, c("latitude")],
       pch=". ", col="red", cex=2)
# Another example
tikei_lon <- (-144.5465183)
tikei_lat <- -14.9505897
Year <- 2021
tikei_tide <- tide.info(year=Year, longitude=tikei_lon, latitude=tikei_lat)
plot(tikei_tide[, "DateTime.local"], tikei_tide[, "Tide.meter"],
     type="l", bty="n", las=1,
     main=tikei_tide[1, "Location"],
     xlab=as.character(Year), ylab="Tide level in meter")
## Another one
tikei_lon <- (-75.56861111)
tikei_lat <- 39.50083333
Year <- 2012
tikei_tide <- tide.info(year=Year, longitude=tikei_lon, latitude=tikei_lat)

library(mapdata)
map('worldHires', xlim=c(-77, -74), ylim=c(37, 40))
points(x=tikei_lon, y=tikei_lat, pch=19, col="red", cex=1)
points(x=tikei_tide$Longitude[1], y=tikei_tide$Latitude[2],
       pch=19, col="blue", cex=1)

par(mar=c(4, 4, 2, 2))
plot(tikei_tide$DateTime.local, tikei_tide$Tide.meter, type="l")

## End(Not run)

```

**tnirp***Read an ASCII text representation of a named or not vector object***Description**

Read an ASCII text representation of a named or not vector object.  
 Note that paste0(rev(c("p", "r", "i", "n", "t"))), collapse="") = "tnirp"

**Usage**

```
tnirp(x, named = TRUE)
```

**Arguments**

x	A string or a vector of strings with value and possibly names.
named	TRUE if names are included.

**Details**

tnirp reads an ASCII text representation of a named or not vector object

**Value**

A vector

**Author(s)**

Marc Girondot <marc.girondot@gmail.com>

**See Also**

Other Characters: [asc\(\)](#), [char\(\)](#), [d\(\)](#)

**Examples**

```
A <- structure(runif(26), .Names=letters)
text <- capture.output(A)
tnirp(text)

tnirp("      mu    mu_season        OTN      p1.09      p1.10      p1.11
      4.63215947 10.78627511  0.36108497  0.08292101 -0.52558196 -0.76430859
      p1.12      p1.13      p1.14      p1.15      p1.16      p1.17
      -0.75186542 -0.57632291 -0.58017174 -0.57048696 -0.56234135 -0.80645122
      p1.18      p1.19      p1.20      p1.21      p1.22      p1.23
      -0.77752524 -0.80909494 -0.56920540 -0.55317302  0.45757298 -0.64155368
      p1.24      p1.25      p1.26      p1.27      p1.28      p1.29
      -0.59119637 -0.66006794 -0.66582399 -0.66772684 -0.67351412 -0.66941992
      p1.30      p1.31      p1.32      p1.33      p1.34      p1.35
      -0.67038245 -0.68938726 -0.68889078 -0.68779016 -0.68604629 -0.68361820
```

```

p1.36      p1.37      p2.09      p2.10      p2.11      p2.12
-0.67045238 -0.66115613 2.55403149 2.31060620 2.31348160 2.20958757
p2.13      p2.14      p2.15      p2.16      p2.17      p2.18
2.14304918 2.19699719 2.30705457 2.18740019 2.32305811 2.31668302
p2.19      p2.20      p2.21      p2.22      p2.23      p2.24
1.99424288 2.06613445 2.38092301 2.40551276 2.31987342 2.30344402
p2.25      p2.26      p2.27      p2.28      p2.29      p2.30
2.26869058 2.25008836 2.23385204 2.22768782 2.25341904 1.77043360
p2.31      p2.32      p2.33      p2.34      p2.35      p2.36
2.21606813 2.21581431 2.21153872 2.21118013 2.21375660 2.21182196
p2.37
1.86137833 ")
tnirp(" 27.89 289.99
90.56", named=FALSE)

```

**universalmcapply***Run the function FUN on X using parallel computing***Description**

Return the results of the function FUN applied to X. It uses forking in unix system and not in windows system.

By default, it will send all the content of environment.

**Usage**

```

universalmcapply(
  X,
  FUN,
  ...,
  mc.cores = getOption("mc.cores", parallel::detectCores()),
  mc.preschedule = TRUE,
  clusterExport = list(),
  clusterEvalQ = list(),
  forking = ifelse(.Platform$OS.type == "windows", FALSE, TRUE),
  progressbar = FALSE
)

```

**Arguments**

- |          |   |
|----------|---|
| X        | A vector (atomic or list) or an expressions vector. Other objects (including classed objects) will be coerced by as.list. |
| FUN      | The function to be applied to each element of X   |
| ...      | Optional arguments to FUN   |
| mc.cores | The number of cores to use, i.e. at most how many child processes will be run simultaneously.                             |

**mc.preschedule** if set to TRUE then the computation is first divided to (at most) as many jobs are there are cores and then the jobs are started, each job possibly covering more than one value. If set to FALSE then one job is forked for each value of X. The former is better for short computations or large number of values in X, the latter is better for jobs that have high variance of completion time and not too many values of X compared to mc.cores.

**clusterExport** List of clusterExport parameters as list

**clusterEvalQ** List of clusterEvalQ parameters as list

**forking** If TRUE will use forking

**progressbar** If pbapply package is installed, show a progressbar

## Details

*universalmcapply* runs the function FUN on X using parallel computing

## Value

The results of the function FUN applied to X

## Author(s)

Marc Girondot <marc.girondot@gmail.com>

## Examples

```
## Not run:
library.HelpersMG)
x <- 1:1000
funx <- function(y) {
  mint <- rep(NA, length(y))
  for (i in seq_along(y)) {
    k <- rnorm(runif(n = 1, 50, 50), mean=10, sd=2)
    mint[i] <- mean(k)
  }
  mint
}
# Note that parallel computing is not always the best solution !
(tp <- system.time({
  m <- lapply(X=x, FUN=funx)
}))
(tp <- system.time({
  m <- universalmcapply(X=x, FUN=funx, forking=FALSE)
}))
(tp <- system.time({
  m <- universalmcapply(X=x, FUN=funx, forking=TRUE)
}))

### An example using clusterExport
# Here no error is generated because environment was exported
# However forking is not possible in windows and non parallel code is ran
```

```
pp <- runif(100)
x <- 1:100
funx1 <- function(y) {pp[y]*10}
u <- universalmclapply(x, FUN=funx1, forking=TRUE)

# Here an error is generated because environment was not exported when parLapplyLB is used
pp <- runif(100)
x <- 1:100
u <- universalmclapply(x, FUN=funx1, forking=FALSE)
u <- universalmclapply(x, FUN=funx1, forking=FALSE,
                        clusterExport=list())

# here no error is generated because the variable pp is exported
pp <- runif(100)
x <- 1:100
u <- universalmclapply(x, FUN=funx1, forking=FALSE,
                        clusterExport=list(varlist=c("pp"), envir=environment()))

# here no error is generated because all the environment is exported
pp <- runif(100)
x <- 1:100
u <- universalmclapply(x, FUN=funx1, forking=FALSE,
                        clusterExport=list(varlist=c(ls()), envir=environment()))

#### An example using clusterEvalQ
asc("a") # asc() is a function from packages HelpersMG
funx2 <- function(y) {asc("a")*10}
# In unix, the loaded packages are visible from all cores
x <- 1:100
u <- universalmclapply(x, FUN=funx2, forking=TRUE)
# In windows, the loaded packages are not visible from all cores
x <- 1:100
u <- universalmclapply(x, FUN=funx2, forking=FALSE)
# In windows, the loaded packages are not visible from all cores
x <- 1:100
u <- universalmclapply(x, FUN=funx2, forking=FALSE,
                        clusterEvalQ=list(expr=expression(library(HelpersMG))))
)

#### If package pbapply is available, progress bar can be shown
m <- universalmclapply(X=x, FUN=funx, forking=FALSE, progressbar=TRUE)
m <- universalmclapply(X=x, FUN=funx, forking=TRUE, progressbar=TRUE)

## End(Not run)
```

### Description

Download a file from internet and save it locally. This function is a wrapper for download.files() that keep the name identical and can get several files at once. It was written to simplify downloading of file. It does not use the true wget function (<https://www.gnu.org/software/wget/>) which is much more complex but also powerful.

### Usage

```
wget(url = stop("At least one internet address is required"), ...)
```

### Arguments

url	The url where to download file
...	The parameters send to download.file()

### Details

wget download a file from internet and save it locally

### Value

Nothing

### Author(s)

Marc Girondot

### Examples

```
## Not run:  
library.HelpersMG  
# Save locally the files send in the parameter url  
wget(c("https://cran.r-project.org/web/packages/HelpersMG/HelpersMG.pdf",  
      "https://cran.r-project.org/web/packages/embryogrowth/embryogrowth.pdf"))  
  
## End(Not run)
```

# Index

- \* **AIC functions**
  - logLik.compareAIC, 79
- \* **AIC**
  - compare\_AIC, 20
  - compare\_AICc, 21
  - compare\_BIC, 23
  - ExtractAIC.glm, 50
  - FormatCompareAIC, 55
- \* **Characters**
  - asc, 13
  - char, 18
  - d, 36
  - tnirp, 162
- \* **Distribution of sum of random variable with negative binomial distributions**
  - dSnbinom, 45
  - rSnbinom, 142
- \* **Distributions**
  - cutter, 28
  - dbeta\_new, 37
  - dcutter, 38
  - ddgamma, 40
  - logLik.cutter, 80
  - plot.cutter, 98
  - print.cutter, 117
  - pSnbinom, 120
  - qSnbinom, 122
  - r2norm, 124
  - rcutter, 128
  - rmnorm, 132
  - rnbinom\_new, 140
- \* **Iconography of correlations**
  - IC\_clean\_data, 59
  - IC\_correlation\_simplify, 61
  - IC\_threshold\_matrix, 62
  - plot.IconoCorel, 102
- \* **LD50 functions**
  - LD50, 70
- \* **Lunar**
  - moon.info, 92
- \* **Lune**
  - moon.info, 92
- \* **Moon**
  - moon.info, 92
- \* **Periodic patterns of indices**
  - index.periodic, 65
  - minmax.periodic, 88
  - moon.info, 92
  - sun.info, 154
  - tide.info, 159
- \* **Rectangle Regression**
  - RectangleRegression, 131
- \* **Results Management**
  - RM\_add, 134
  - RM\_delete, 135
  - RM\_duplicate, 136
  - RM\_get, 138
  - RM\_list, 139
- \* **Symbol**
  - symbol.Female, 156
  - symbol.Male, 157
- \* **Tide**
  - tide.info, 159
- \* **logit**
  - flexit, 53
  - invlogit, 69
  - logit, 78
- \* **mcmcComposite functions**
  - as.mcmc.mcmcComposite, 8
  - as.parameters, 9
  - as.quantiles, 12
  - merge.mcmcComposite, 82

MHalgoGen, 84  
 plot.mcmcComposite, 106  
 plot.PriorsmcmcComposite, 111  
 setPriors, 148  
 summary.mcmcComposite, 153  
**\* ncdf**  
 format\_ncdf, 56  
 ind\_long\_lat, 66  
**\* plot and barplot functions**  
 barplot\_errbar, 14  
 plot\_add, 112  
 plot\_errbar, 113  
 ScalePreviousPlot, 143  
 show\_name, 150  
**\* w-value functions**  
 compare, 19  
 contingencyTable.compare, 24  
 series.compare, 146  
  
 addS3Class, 7  
 as.mcmc.mcmcComposite, 8, 10, 12, 83, 86, 107, 112, 149, 153  
 as.parameters, 8, 9, 12, 83, 86, 107, 112, 149, 153  
 as.quantiles, 8, 10, 12, 83, 86, 107, 112, 149, 153  
 asc, 13, 18, 36, 162  
  
 barplot\_errbar, 14, 113, 115, 143, 150  
  
 cArrows, 15  
 ChangeCoordinate, 17  
 char, 14, 18, 36, 162  
 compare, 19, 25, 146  
 compare\_AIC, 20, 22, 23, 51, 56  
 compare\_AICC, 21, 21, 23, 51, 56  
 compare\_BIC, 21, 22, 23, 51, 56  
 contingencyTable.compare, 19, 24, 146  
 convert.tz, 27  
 cutter, 28, 38, 39, 41, 80, 99, 117, 121, 123, 125, 129, 133, 141  
  
 d, 14, 18, 36, 162  
 dbeta\_new, 30, 37, 39, 41, 80, 99, 117, 121, 123, 125, 129, 133, 141  
 dcutter, 30, 38, 38, 41, 80, 99, 117, 121, 123, 125, 129, 133, 141  
 dggamma, 30, 38, 39, 40, 80, 99, 117, 121, 123, 125, 129, 133, 141  
  
 DIx, 42  
 dnbinom\_new, 43  
 dSnbnom, 45, 142  
 duplicated\_packages, 47  
  
 ellipse, 48  
 ExtractAIC.glm, 21–23, 50, 56  
  
 fitdistrquantiles, 52  
 flexit, 53, 69, 78  
 format\_ncdf, 56, 67  
 FormatCompareAIC, 21–23, 51, 55  
  
 HelpersMG-package, 4  
  
 IC\_clean\_data, 59, 61, 63, 103  
 IC\_correlation\_simplify, 60, 61, 63, 103  
 IC\_threshold\_matrix, 60, 61, 62, 103  
 iCutter, 58  
 ind\_long\_lat, 57, 66  
 index.periodic, 65, 89, 92, 155, 160  
 inside, 68  
 invlogit, 55, 69, 78  
  
 LD50, 70, 74, 75, 82, 105, 116  
 LD50\_MHmcmc, 71, 72, 75, 82, 105, 116  
 LD50\_MHmcmc\_p, 71, 74, 75, 82, 105, 116  
 list.packages, 76  
 local.search, 77  
 logit, 55, 69, 78  
 logLik.compareAIC, 79  
 logLik.cutter, 30, 38, 39, 41, 80, 99, 117, 121, 123, 125, 129, 133, 141  
 logLik.LD50, 71, 74, 75, 81, 105, 116  
  
 merge.mcmcComposite, 8, 10, 12, 82, 86, 107, 112, 149, 153  
 MHalgoGen, 8, 10, 12, 83, 84, 107, 112, 149, 153  
 minmax.periodic, 65, 88, 92, 155, 160  
 modeled.hist, 90  
 modifyVector, 91  
 moon.info, 65, 89, 92, 155, 160  
 MovingWindow, 93  
  
 NagelkerkeScaledR2, 94  
 newcompassRose, 95  
 newmap.scale, 96  
  
 openwd, 97

pgamma (dggamma), 40  
 plot.cutter, 30, 38, 39, 41, 80, 98, 117, 121, 123, 125, 129, 133, 141  
 plot.IconoCorel, 60, 61, 63, 102  
 plot.LD50, 71, 74, 75, 82, 104, 116  
 plot.mcmcComposite, 8, 10, 12, 83, 86, 106, 112, 149, 153  
 plot.PriorsmcmcComposite, 8, 10, 12, 83, 86, 107, 111, 149, 153  
 plot\_add, 15, 112, 115, 143, 150  
 plot\_errbar, 15, 113, 113, 143, 150  
 predict.LD50, 71, 74, 75, 82, 105, 115  
 print.cutter, 30, 38, 39, 41, 80, 99, 117, 121, 123, 125, 129, 133, 141  
 pSnbnom, 30, 38, 39, 41, 80, 99, 117, 120, 123, 125, 129, 133, 141  
  
 qggamma (dggamma), 40  
 qSnbnom, 30, 38, 39, 41, 80, 99, 117, 121, 122, 125, 129, 133, 141  
 qvlmer, 123  
  
 r2norm, 30, 38, 39, 41, 80, 99, 117, 121, 123, 124, 129, 133, 141  
 RandomFromHessianOrMCMC, 125  
 rcutter, 30, 38, 39, 41, 80, 99, 117, 121, 123, 125, 128, 133, 141  
 read\_folder, 130  
 RectangleRegression, 131  
 rgamma (dggamma), 40  
 RM\_add, 134, 136–139  
 RM\_delete, 134, 135, 137–139  
 RM\_duplicate, 134, 136, 136, 138, 139  
 RM\_get, 134, 136, 137, 138, 139  
 RM\_list, 134, 136–138, 139  
 rmnorm, 30, 38, 39, 41, 80, 99, 117, 121, 123, 125, 129, 132, 141  
 rnbinom\_new, 30, 38, 39, 41, 80, 99, 117, 121, 123, 125, 129, 133, 140  
 rSnbnom, 46, 142  
  
 ScalePreviousPlot, 15, 113, 115, 143, 150  
 SEfromHessian, 144  
 series.compare, 19, 25, 146  
 setPriors, 8, 10, 12, 83, 86, 107, 112, 148, 153  
 show\_name, 15, 113, 115, 143, 150  
 similar, 151  
 specify\_decimal, 152  
  
 summary.mcmcComposite, 8, 10, 12, 83, 86, 107, 112, 149, 153  
 sun.info, 65, 89, 92, 154, 160  
 symbol.Female, 156, 157  
 symbol.Male, 156, 157  
 symmetricize, 158  
  
 tide.info, 65, 89, 92, 155, 159  
 tnirp, 14, 18, 36, 162  
  
 universalmcapply, 163  
  
 wget, 165