

# Package ‘mice’

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**Title** Multivariate Imputation by Chained Equations

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**Depends** methods, R (>= 2.10.0)

**Imports** lattice, grDevices, graphics, MASS, nnet, rpart, splines,  
stats, survival, utils, Rcpp

**Suggests** AGD, CALIBERrfimpute, gamlss, lme4, mitools, nlme, pan,  
randomForest, Zelig, BSDA, knitr, rmarkdown

**LinkingTo** Rcpp

**Description** Multiple imputation using Fully Conditional Specification (FCS) implemented by the MICE algorithm as described in Van Buuren and Groothuis-Oudshoorn (2011) <doi:10.18637/jss.v045.i03>. Each variable has its own imputation model. Built-in imputation models are provided for continuous data (predictive mean matching, normal), binary data (logistic regression), unordered categorical data (polytomous logistic regression) and ordered categorical data (proportional odds). MICE can also impute continuous two-level data (normal model, pan, second-level variables). Passive imputation can be used to maintain consistency between variables. Various diagnostic plots are available to inspect the quality of the imputations.

**License** GPL-2 | GPL-3

**LazyLoad** yes

**LazyData** yes

**VignetteBuilder** knitr

**URL** <http://www.stefvanbuuren.nl> , <http://www.multiple-imputation.com>

**BugReports** <https://github.com/stefvanbuuren/mice/issues>

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**NeedsCompilation** yes

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## R topics documented:

.pmm.match . . . . .	4
ampute . . . . .	5
ampute.continuous . . . . .	9
ampute.default.freq . . . . .	10
ampute.default.odds . . . . .	11
ampute.default.patterns . . . . .	11
ampute.default.type . . . . .	12
ampute.default.weights . . . . .	13
ampute.discrete . . . . .	13
ampute.mcar . . . . .	14
appendbreak . . . . .	15
as.mids . . . . .	16
as.mira . . . . .	17
boys . . . . .	18
bwplot.mads . . . . .	19
bwplot.mids . . . . .	21
cbind.mids . . . . .	24
cc . . . . .	26
cci . . . . .	26
complete . . . . .	27
densityplot.mids . . . . .	29
extractBS . . . . .	32
fdd . . . . .	33
fdgs . . . . .	35
fico . . . . .	36
flux . . . . .	37
fluxplot . . . . .	38
getfit . . . . .	40
glm.mids . . . . .	41
ibind . . . . .	42
ic . . . . .	43
ici . . . . .	44
is.mads . . . . .	44

is.mids . . . . .	45
is.mipo . . . . .	45
is.mira . . . . .	46
leiden85 . . . . .	46
lm.mids . . . . .	47
mads-class . . . . .	48
mammalsleep . . . . .	49
md.pairs . . . . .	50
md.pattern . . . . .	52
mdc . . . . .	53
mice . . . . .	54
mice.impute.2l.norm . . . . .	59
mice.impute.2l.pan . . . . .	60
mice.impute.2lonly.mean . . . . .	62
mice.impute.2lonly.norm . . . . .	63
mice.impute.2lonly.pmm . . . . .	65
mice.impute.cart . . . . .	67
mice.impute.fastpmm . . . . .	68
mice.impute.lda . . . . .	70
mice.impute.logreg . . . . .	71
mice.impute.logreg.boot . . . . .	72
mice.impute.mean . . . . .	73
mice.impute.midastouch . . . . .	74
mice.impute.norm . . . . .	76
mice.impute.norm.boot . . . . .	77
mice.impute.norm.nob . . . . .	78
mice.impute.norm.predict . . . . .	79
mice.impute.passive . . . . .	80
mice.impute.pmm . . . . .	81
mice.impute.polr . . . . .	83
mice.impute.polyreg . . . . .	84
mice.impute.quadratic . . . . .	86
mice.impute.rf . . . . .	87
mice.impute.ri . . . . .	89
mice.impute.sample . . . . .	90
mice.mids . . . . .	91
mice.theme . . . . .	92
mids-class . . . . .	93
mids2mplus . . . . .	94
mids2spss . . . . .	95
mipo-class . . . . .	97
mira-class . . . . .	98
ncc . . . . .	99
nelsonaalen . . . . .	99
nhanes . . . . .	101
nhanes2 . . . . .	102
nic . . . . .	103
norm.draw . . . . .	103

pattern . . . . .	104
plot.mids . . . . .	105
pool . . . . .	106
pool.compare . . . . .	108
pool.r.squared . . . . .	110
pool.scalar . . . . .	112
popmis . . . . .	113
pops . . . . .	114
potthoffroy . . . . .	115
print.mads . . . . .	116
print.mids . . . . .	117
quickpred . . . . .	118
rbind.mids . . . . .	120
selfreport . . . . .	121
squeeze . . . . .	123
stripplot.mids . . . . .	124
summary.mira . . . . .	127
supports.transparent . . . . .	128
tbc . . . . .	129
version . . . . .	130
walking . . . . .	131
windspeed . . . . .	132
with.mids . . . . .	133
xyplot.mads . . . . .	134
xyplot.mids . . . . .	135

**Index****139**


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<code>.pmm.match</code>	<i>Finds an imputed value from matches in the predictive metric</i>
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**Description**

This function finds matches among the observed data in the predictive mean metric. It selects the donor's closest matches, randomly samples one of the donors, and returns the observed value of the match.

**Usage**

```
.pmm.match(z, yhat = yhat, y = y, donors = 5, ...)
```

**Arguments**

<code>z</code>	A scalar containing the predicted value for the current case to be imputed.
<code>yhat</code>	A vector containing the predicted values for all cases with an observed outcome.
<code>y</code>	A vector of <code>length(yhat)</code> elements containing the observed outcome

donors	The size of the donor pool among which a draw is made. The default is donors = 5. Setting donors = 1 always selects the closest match. Values between 3 and 10 provide the best results. Note: This setting was changed from 3 to 5 in version 2.19, based on simulation work by Tim Morris (UCL).
...	Other parameters (not used).

### Details

Not used after mice 2.21. The `mice.impute.pmm()` function now calls the much faster C function `matcher` instead of `.pmm.match()`. Use `mice(..., version = "2.21")` to call `.pmm.match()`

### Value

A scalar containing the observed value of the selected donor.

### Author(s)

Stef van Buuren

### References

Schenker N & Taylor JMG (1996) Partially parametric techniques for multiple imputation. *Computational Statistics and Data Analysis*, 22, 425-446.

Little RJA (1988) Missing-data adjustments in large surveys (with discussion). *Journal of Business Economics and Statistics*, 6, 287-301.

---

ampute

*Generate Missing Data for Simulation Purposes*

---

### Description

This function generates multivariate missing data in a MCAR, MAR or MNAR manner. Imputation of data sets containing missing values can be performed with [mice](#).

### Usage

```
ampute(data, prop = 0.5, patterns = NULL, freq = NULL, mech = "MAR",
        weights = NULL, cont = TRUE, type = NULL, odds = NULL,
        bycases = TRUE, run = TRUE)
```

### Arguments

data	A complete data matrix or dataframe. Values should be numeric. Categorical variables should have been transformed into dummies.
prop	A scalar specifying the proportion of missingness. Should be a value between 0 and 1. Default is a missingness proportion of 0.5.

patterns	A matrix or data frame of size #patterns by #variables where 0 indicates a variable should have missing values and 1 indicates a variable should remain complete. The user may specify as many patterns as desired. One pattern (a vector) or double patterns are possible as well. Default is a square matrix of size #variables where each pattern has missingness on one variable only (created with <code>ampute.default.patterns</code> ). After the amputation procedure, <code>md.pattern</code> can be used to investigate the missing data patterns in the data.
freq	A vector of length #patterns containing the relative frequency with which the patterns should occur. For example, for three missing data patterns, the vector could be <code>c(0.4, 0.4, 0.2)</code> , meaning that of all cases with missing values, 40 percent should have pattern 1, 40 percent pattern 2 and 20 percent pattern 3. The vector should sum to 1. Default is an equal probability for each pattern, created with <code>ampute.default.freq</code> .
mech	A string specifying the missingness mechanism, either MCAR (Missing Complete At Random), MAR (Missing At Random) or MNAR (Missing Not At Random). Default is a MAR missingness mechanism.
weights	A matrix or data frame of size #patterns by #variables. The matrix contains the weights that will be used to calculate the weighted sum scores. For a MAR mechanism, weights of the variables that will be made incomplete, should be zero. For a MNAR mechanism, these weights might have any possible value. Furthermore, the weights may differ between patterns and between variables. They may be negative as well. Within each pattern, the relative size of the values are of importance. The default weights matrix is made with <code>ampute.default.weights</code> and returns a matrix with equal weights for all variables. In case of MAR, variables that will be amputed will be weighted with 0. If it is MNAR, variables that will be observed will be weighted with 0. If mechanism is MCAR, the weights matrix will not be used.
cont	Logical. Whether the probabilities should be based on a continuous or discrete distribution. If TRUE, the probabilities of being missing are based on a continuous logit distribution. <code>ampute.continuous</code> will be used to calculate and assign the probabilities. These will be based on argument type. If FALSE, the probabilities of being missing are based on a discrete distribution ( <code>ampute.discrete</code> ) based on the odds argument. Default is TRUE.
type	A vector of strings containing the type of missingness for each pattern. Either "LEFT", "MID", "TAIL" or "RIGHT". If a single missingness type is entered, all patterns will be created by the same type. If missingness types should differ over patterns, a vector of missingness types should be entered. Default is RIGHT for all patterns and is the result of <code>ampute.default.type</code> .
odds	A matrix where #patterns defines the #rows. Each row should contain the odds of being missing for the corresponding pattern. The amount of odds values defines in how many quantiles the sum scores will be divided. The values are relative probabilities: a quantile with odds value 4 will have a probability of being missing that is four times higher than a quantile with odds 1. The #quantiles may differ between the patterns, specify NA for cells remaining empty. Default is 4 quantiles with odds values 1, 2, 3 and 4, the result of <code>ampute.default.odds</code> .
bycases	Logical. If TRUE, the proportion of missingness is defined in terms of cases. If FALSE, the proportion of missingness is defined in terms of cells. Default is

TRUE.

run Logical. If TRUE, the amputations are implemented. If FALSE, the return object will contain everything but the amputed data set.

## Details

When new multiple imputation techniques are tested, missing values need to be generated in simulated data sets. The generation of missing values is what we call: amputation. The function `ampute` is developed to perform any kind of amputation desired by the researcher. An extensive example and more explanation of the function can be found in the vignette *Multivariate Amputation using Ampute*, available in `mice` as well. For imputation, the function `mice` is advised.

Until recently, univariate amputation procedures were used to generate missing data in complete, simulated data sets. With this approach, variables are made incomplete one variable at a time. When several variables need to be amputed, the procedure is repeated multiple times.

With this univariate approach, it is difficult to relate the missingness on one variable to the missingness on another variable. A multivariate amputation procedure solves this issue and moreover, it does justice to the multivariate nature of data sets. Hence, `ampute` is developed to perform the amputation according the researcher's desires.

The idea behind the function is the specification of several missingness patterns. Each pattern is a combination of variables with and without missing values (denoted by 0 and 1 respectively). For example, one might want to create two missingness patterns on a data set with four variables. The patterns could be something like: 0, 0, 1, 1 and 1, 0, 1, 0. Each combination of zeros and ones may occur.

Furthermore, the researcher specifies the proportion of missingness, either the proportion of missing cases or the proportion of missing cells, and the relative frequency each pattern occurs. Consequently, the data is divided over the patterns with these probabilities. Now, each case is candidate for a certain missingness pattern, but whether the case will have missing values eventually, depends on other specifications.

The first of these specifications is the missing mechanism. There are three possible mechanisms: the missingness depends completely on chance (MCAR), the missingness depends on the values of the observed variables (i.e. the variables that remain complete) (MAR) or on the values of the variables that will be made incomplete (MNAR). For a more thorough explanation of these definitions, I refer to Van Buuren (2012).

When the user sets the missingness mechanism to "MCAR", the candidates have an equal probability of having missing values. No other specifications have to be made. For a "MAR" or "MNAR" mechanism, weighted sum scores are calculated. These scores are a linear combination of the variables.

In order to calculate the weighted sum scores, the data is standardized. That is the reason the data has to be numeric. Second, for each case, the values in the data set are multiplied with the weights, specified by argument `weights`. These weighted scores will be summed, resulting in a weighted sum score for each case.

The weights may differ between patterns and they may be negative or zero as well. Naturally, in case of a MAR mechanism, the weights corresponding to the variables that will be made incomplete, have a 0. Note that this might be different for each pattern. In case of MNAR missingness, especially the weights of the variables that will be made incomplete are of importance. However, the other variables might be weighted as well.

It is the relative difference between the weights that will result in an effect in the sum scores. For example, for the first missing data pattern mentioned above, the weights for the third and fourth variables might be set to 2 and 4. However, weight values of 0.2 and 0.4 will have the exact same effect on the weighted sum score: the fourth variable is weighted twice as much as variable 3.

Based on the weighted sum scores, either a discrete or continuous distribution of probabilities is used to calculate whether a candidate will have missing values.

For a discrete distribution of probabilities, the weighted sum scores are divided into subgroups of equal size (quantiles). Thereafter, the user specifies for each subgroup the odds of being missing. Both the number of subgroups and the odds values are important for the generation of missing data. For example, for a RIGHT-like mechanism, scoring in one of the higher quantiles should have high missingness odds, whereas for a MID-like mechanism, the central groups should have higher odds. Again, not the size of the odds values are of importance, but the relative distance between the values.

The continuous distributions of probabilities are based on the logit function, as described by Van Buuren (2012). The user can specify the type of missingness, which, again, may differ between patterns.

For an extensive example of the working of the function, I gladly refer to the vignette *Multivariate Amputation using Ampute*.

### Value

Returns an S3 object of class `mads-class` (multivariate amputed data set)

### Author(s)

Rianne Schouten [aut, cre], Gerko Vink [aut], Peter Lugtig [ctb], 2016

### References

Brand, J.P.L. (1999). *Development, implementation and evaluation of multiple imputation strategies for the statistical analysis of incomplete data sets* (pp. 110-113). Dissertation. Rotterdam: Erasmus University.

Van Buuren, S., Brand, J.P.L., Groothuis-Oudshoorn, C.G.M., Rubin, D.B. (2006). Fully conditional specification in multivariate imputation. *Journal of Statistical Computation and Simulation*, 76(12), Appendix B.

Van Buuren, S. (2012). *Flexible imputation of missing data*. Boca Raton, FL.: Chapman & Hall/CRC Press.

Vink, G. (2016). Towards a standardized evaluation of multiple imputation routines.

### See Also

`mads-class`, `bwplot`, `xyplot`, `ampute.mcar`, `ampute.continuous`, `ampute.discrete`, `mice`

### Examples

```
# Simulate data set with \code{mvrnorm} from package \code{\pkg{MASS}}.
require(MASS)
sigma <- matrix(data = c(1, 0.2, 0.2, 0.2, 1, 0.2, 0.2, 0.2, 1), nrow = 3)
complete.data <- mvrnorm(n = 100, mu = c(5, 5, 5), Sigma = sigma)
```



```

# Perform quick amputation
result1 <- ampute(data = complete.data)
# Change default matrices as desired
patterns <- result1$patterns
patterns[1:3, 2] <- 0
odds <- result1$odds
odds[2,3:4] <- c(2, 4)
odds[3,] <- c(3, 1, NA, NA)
# Rerun amputation
result3 <- ampute(data = complete.data, patterns = patterns, freq =
c(0.3, 0.3, 0.4), cont = FALSE, odds = odds)
# Run an amputation procedure with continuous probabilities
result4 <- ampute(data = complete.data, type = c("RIGHT", "TAIL", "LEFT"))

```

---

ampute.continuous

*Multivariate Amputation Based On Continuous Probability Functions*


---

## Description

This function creates a missing data indicator for each pattern. The continuous probability distributions (Van Buuren, 2012, pp. 63, 64) will be induced on the weighted sum scores, calculated earlier in the multivariate amputation function [ampute](#).

## Usage

```
ampute.continuous(P, scores, prop, type)
```

## Arguments

P	A vector containing the pattern numbers of the cases's candidacies. For each case, a value between 1 and #patterns is given. For example, a case with value 2 is candidate for missing data pattern 2.
scores	A list containing vectors with the candidates's weighted sum scores, the result of an underlying function in <a href="#">ampute</a> .
prop	A scalar specifying the proportion of missingness. Should be a value between 0 and 1. Default is a missingness proportion of 0.5.
type	A vector of strings containing the type of missingness for each pattern. Either "LEFT", "MID", "TAIL" or "RIGHT". If a single missingness type is entered, all patterns will be created by the same type. If missingness types should differ over patterns, a vector of missingness types should be entered. Default is RIGHT for all patterns and is the result of <a href="#">ampute.default.type</a> .

## Value

A list containing vectors with 0 if a case should be made missing and 1 if a case should remain complete. The first vector refers to the first pattern, the second vector to the second pattern, etcetera.

**Author(s)**

Rianne Schouten [aut, cre], Gerko Vink [aut], Peter Lugtig [ctb], 2016

**References**

Van Buuren, S. (2012). *Flexible imputation of missing data*. Boca Raton, FL.: Chapman & Hall/CRC Press.

**See Also**

[ampute](#), [ampute.default.type](#)

---

`ampute.default.freq`    *Default freq in ampute*

---

**Description**

Defines the default relative frequency vector for the multivariate amputation function `ampute`.

**Usage**

```
ampute.default.freq(patterns)
```

**Arguments**

`patterns`    A matrix of size `#patterns` by `#variables` where 0 indicates a variable should have missing values and 1 indicates a variable should remain complete. Could be the result of [ampute.default.patterns](#).

**Value**

A vector of length `#patterns` containing the relative frequencies with which the patterns should occur. An equal probability is given to each pattern.

**Author(s)**

Rianne Schouten, 2016

**See Also**

[ampute](#), [ampute.default.patterns](#)

---

ampute.default.odds     *Default odds in ampute()*

---

**Description**

Defines the default odds matrix for the multivariate amputation function `ampute`.

**Usage**

```
ampute.default.odds(patterns)
```

**Arguments**

`patterns`     A matrix of size `#patterns` by `#variables` where 0 indicates a variable should have missing values and 1 indicates a variable should remain complete. Could be the result of `ampute.default.patterns`.

**Value**

A matrix where `#rows` equals `#patterns`. Default is 4 quantiles with odds values 1, 2, 3 and 4, for each pattern, imitating a RIGHT type of missingness.

**Author(s)**

Rianne Schouten, 2016

**See Also**

`ampute`, `ampute.default.patterns`

---

ampute.default.patterns  
                          *Default patterns in ampute*

---

**Description**

This function creates a default pattern matrix for the multivariate amputation function `ampute()`.

**Usage**

```
ampute.default.patterns(n)
```

**Arguments**

`n`     A scalar specifying the `#variables` in the data.

**Value**

A square matrix of size #variables where 0 indicates a variable

**Author(s)**

Rianne Schouten, 2016

**See Also**

[ampute](#), [md.pattern](#)

---

`ampute.default.type`    *Default type in ampute()*

---

**Description**

Defines the default type vector for the multivariate amputation function `ampute`.

**Usage**

```
ampute.default.type(patterns)
```

**Arguments**

`patterns`    A matrix of size #patterns by #variables where 0 indicates a variable should have missing values and 1 indicates a variable should remain complete. Could be the result of [ampute.default.patterns](#).

**Value**

A string vector of length #patterns containing the missingness types. Each pattern will be amputated with a "RIGHT" missingness.

**Author(s)**

Rianne Schouten, 2016

**See Also**

[ampute](#), [ampute.default.patterns](#)

---

ampute.default.weights  
*Default weights in ampute*

---

**Description**

Defines the default weights matrix for the multivariate amputation function `ampute`.

**Usage**

```
ampute.default.weights(patterns, mech)
```

**Arguments**

patterns	A matrix of size #patterns by #variables where 0 indicates a variable should have missing values and 1 indicates a variable should remain complete. Could be the result of <a href="#">ampute.default.patterns</a> .
mech	A string specifying the missingness mechanism.

**Value**

A matrix of size #patterns by #variables containing the weights that will be used to calculate the weighted sum scores. Equal weights are given to all variables. When mechanism is MAR, variables that will be amputed will be weighted with 0. If it is MNAR, variables that will be observed will be weighted with 0. If mechanism is MCAR, the weights matrix will not be used. A default MAR matrix will be returned.

**Author(s)**

Rianne Schouten, 2016

**See Also**

[ampute](#), [ampute.default.patterns](#)

---

ampute.discrete      *Multivariate Amputation Based On Discrete Probability Functions*

---

**Description**

This function creates a missing data indicator for each pattern. Odds probabilities (Brand, 1999, pp. 110-113) will be induced on the weighted sum scores, calculated earlier in the multivariate amputation function `ampute`.

## Usage

```
ampute.discrete(P, scores, prop, odds)
```

## Arguments

P	A vector containing the pattern numbers of the cases's candidacies. For each case, a value between 1 and #patterns is given. For example, a case with value 2 is candidate for missing data pattern 2.
scores	A list containing vectors with the candidates's weighted sum scores, the result of an underlying function in <a href="#">ampute</a> .
prop	A scalar specifying the proportion of missingness. Should be a value between 0 and 1. Default is a missingness proportion of 0.5.
odds	A matrix where #patterns defines the #rows. Each row should contain the odds of being missing for the corresponding pattern. The amount of odds values defines in how many quantiles the sum scores will be divided. The values are relative probabilities: a quantile with odds value 4 will have a probability of being missing that is four times higher than a quantile with odds 1. The #quantiles may differ between the patterns, specify NA for cells remaining empty. Default is 4 quantiles with odds values 1, 2, 3 and 4, the result of <a href="#">ampute.default.odds</a> .

## Value

A list containing vectors with 0 if a case should be made missing and 1 if a case should remain complete. The first vector refers to the first pattern, the second vector to the second pattern, etcetera.

## Author(s)

Rianne Schouten, 2016

## References

Brand, J.P.L. (1999). *Development, implementation and evaluation of multiple imputation strategies for the statistical analysis of incomplete data sets*. Dissertation. Rotterdam: Erasmus University.

## See Also

[ampute](#), [ampute.default.odds](#)

## Description

This function creates a missing data indicator for each pattern, based on a MCAR missingness mechanism. The function is used in the multivariate amputation function [ampute](#).

**Usage**

```
ampute.mcar(P, patterns, prop)
```

**Arguments**

P	A vector containing the pattern numbers of the cases's candidacies. For each case, a value between 1 and #patterns is given. For example, a case with value 2 is candidate for missing data pattern 2.
patterns	A matrix of size #patterns by #variables where 0 indicates a variable should have missing values and 1 indicates a variable should remain complete. The user may specify as many patterns as desired. One pattern (a vector) is also possible. Could be the result of <code>ampute.default.patterns</code> , default will be a square matrix of size #variables where each pattern has missingness on one variable only.
prop	A scalar specifying the proportion of missingness. Should be a value between 0 and 1. Default is a missingness proportion of 0.5.

**Value**

A list containing vectors with 0 if a case should be made missing and 1 if a case should remain complete. The first vector refers to the first pattern, the second vector to the second pattern, etcetera.

**Author(s)**

Rianne Schouten, 2016

**See Also**

[ampute](#)

---

appendbreak

*Appends specified break to the data*

---

**Description**

A custom function to insert rows in long data with new pseudo-observations that are being done on the specified break ages. There should be a column called `first` in data with logical data that codes whether the current row is the first for subject `id`. Furthermore, the function assumes that columns `age`, `occ`, `hgt.z`, `wgt.z` and `bmi.z` are available. This function is used on the `tbc` data in FIMD chapter 9. Check that out to see it in action.

**Usage**

```
appendbreak(data, brk, warp.model = warp.model, id = NULL, typ = "pred")
```

**Arguments**

data	A data frame in the long long format
brk	A vector of break ages
warp.model	A time warping model
id	The subject identifier
typ	Label to signal that this is a newly added observation

**Value**

A long data frame with additional rows for the break ages

---

as.mids	<i>Converts an multiply imputed dataset (long format) into a mids object</i>
---------	--

---

**Description**

This function converts imputed data stored in long format into an object of class `mids`. The original incomplete data set needs to be available so that we know where the missing data are. The function is useful to convert back operations applied to the imputed data back in a `mids` object. It may also be used to store multiply imputed data sets from other software into the format used by `mice`.

**Usage**

```
as.mids(data, .imp=1, .id=2)
```

**Arguments**

data	A multiply imputed data set in long format
.imp	Mandatory column indicator for the multiple imputation stream, where 0 indicates the incomplete data and 1 through $m$ indicate the $m$ multiple imputation streams. Default is 1.
.id	Optional column indicator for the row numbers. Default is 2.

**Details**

If `.id` is specified, row names from the original data (if supplied) will be copied to the `mids` object.

**Value**

An object of class `mids`

**Author(s)**

Gerko Vink



**Examples**

```

# impute the nhanes dataset
imp <- mice(nhanes, print = FALSE)
# extract the data in long format
X <- complete(imp, action = "long", include = TRUE)
# create dataset with .imp variable that is not a factor
X2 <- X
X2$.imp <- as.numeric(levels(X$.imp))[X$.imp]
# nhanes example without .id
test1 <- as.mids(X[, -2], .id = NULL)
is.mids(test1)
all(complete(test1, action = "long", include = TRUE) == X, na.rm = TRUE)
# nhanes example without .id where .imp is not a factor
test2 <- as.mids(X2[, -2], .id = NULL)
is.mids(test2)
all(complete(test1, action = "long", include = TRUE) == X, na.rm = TRUE)
# nhanes example with .id
test3 <- as.mids(X, .id = 2)
is.mids(test3)
all(complete(test1, action = "long", include = TRUE) == X, na.rm = TRUE)
# nhanes example with .id where .imp is not a factor
test4 <- as.mids(X2, .id = 2)
is.mids(test4)
all(complete(test1, action = "long", include = TRUE) == X, na.rm = TRUE)

```

---

as.mira

*Create a mira object from repeated analyses*


---

**Description**

The `as.mira()` function takes the results of repeated complete-data analysis stored as a list, and turns it into a `mira` object that can be pooled. Pooling requires that `coef()` and `vcov()` methods are available for fitted object.

**Usage**

```
as.mira(fitlist)
```

**Arguments**

`fitlist`            A list containing `$m$` fitted analysis objects

**Value**

An S3 object of class `mira`.

**Author(s)**

Stef van Buuren, 2011

**See Also**[mira](#)

---

**boys***Growth of Dutch boys*

---

**Description**

Height, weight, head circumference and puberty of 748 Dutch boys.

**Format**

A data frame with 748 rows on the following 9 variables:

**age** Decimal age (0-21 years)

**hgt** Height (cm)

**wgt** Weight (kg)

**bmi** Body mass index

**hc** Head circumference (cm)

**gen** Genital Tanner stage (G1-G5)

**phb** Pubic hair (Tanner P1-P6)

**tv** Testicular volume (ml)

**reg** Region (north, east, west, south, city)

**Details**

Random sample of 10% from the cross-sectional data used to construct the Dutch growth references 1997. Variables *gen* and *phb* are ordered factors. *reg* is a factor.

**Source**

Fredriks, A.M., van Buuren, S., Burgmeijer, R.J., Meulmeester JF, Beuker, R.J., Brugman, E., Roede, M.J., Verloove-Vanhorick, S.P., Wit, J.M. (2000) Continuing positive secular growth change in The Netherlands 1955-1997. *Pediatric Research*, **47**, 316-323.

Fredriks, A.M., van Buuren, S., Wit, J.M., Verloove-Vanhorick, S.P. (2000). Body index measurements in 1996-7 compared with 1980. *Archives of Disease in Childhood*, **82**, 107-112.

**Examples**

```

# create two imputed data sets
imp <- mice(boys, m=1, maxit=2)
z <- complete(imp, 1)

# create imputations for age <8yrs
plot(z$age, z$gen, col=mdc(1:2)[1+is.na(boys$gen)],
     xlab = "Age (years)", ylab = "Tanner Stage Genital")

# figure to show that the default imputation method does not impute BMI
# consistently
plot(z$bmi, z$wgt/(z$hgt/100)^2, col=mdc(1:2)[1+is.na(boys$bmi)],
     xlab = "Imputed BMI", ylab="Calculated BMI")

# also, BMI distributions are somewhat different
require(MASS)
oldpar <- par(mfrow=c(1,2))
truehist(z$bmi[!is.na(boys$bmi)], h=1, xlim=c(10,30), ymax=0.25,
         col=mdc(1), xlab="BMI observed")
truehist(z$bmi[is.na(boys$bmi)], h=1, xlim=c(10,30), ymax=0.25,
         col=mdc(2), xlab="BMI imputed")
par(oldpar)

# repair the inconsistency problem by passive imputation
meth <- imp$meth
meth["bmi"] <- "~I(wgt/(hgt/100)^2)"
pred <- imp$predictorMatrix
pred["hgt", "bmi"] <- 0
pred["wgt", "bmi"] <- 0
imp2 <- mice(boys, m=1, maxit=2, meth=meth, pred=pred)
z2 <- complete(imp2, 1)

# show that new imputations are consistent
plot(z2$bmi, z2$wgt/(z2$hgt/100)^2, col=mdc(1:2)[1+is.na(boys$bmi)],
     ylab="Calculated BMI")

# and compare distributions
oldpar <- par(mfrow=c(1,2))
truehist(z2$bmi[!is.na(boys$bmi)], h=1, xlim=c(10,30), ymax=0.25, col=mdc(1),
         xlab="BMI observed")
truehist(z2$bmi[is.na(boys$bmi)], h=1, xlim=c(10,30), ymax=0.25, col=mdc(2),
         xlab="BMI imputed")
par(oldpar)

```

## Description

Plotting method to investigate the result of function [ampute](#). Based on [lattice](#). bwplot produces box-and-whisker plots of the relation between the data variables and the amputed data. The function does not show which data is amputed. It does show how the amputed values are related to the variable values.

## Usage

```
## S3 method for class 'mads'  
bwplot(x, yvar = NULL, which.pat = NULL,  
        standardized = TRUE, descriptives = TRUE, layout = NULL)
```

## Arguments

x	A mads ( <a href="#">mads-class</a> ) object, typically created by <a href="#">ampute</a> .
yvar	A string or vector of variable names that needs to be plotted. As a default, all variables will be plotted.
which.pat	A scalar or vector indicating which patterns need to be plotted. As a default, all patterns are plotted.
standardized	Logical. Whether the box-and-whisker plots need to be created from standardized data or not. Default is TRUE.
descriptives	Logical. Whether the mean, variance and n of the variables need to be printed. This is useful to examine the effect of the amputation. Default is TRUE.
layout	A vector of two values indicating how the boxplots of one pattern should be divided over the plot. For example, <code>c(2, 3)</code> indicates that the boxplots of six variables need to be placed on 3 rows and 2 columns. Default is 1 row and an amount of columns equal to #variables. Note that for more than 6 variables, multiple plots will be created automatically.

## Value

A list containing the box-and-whisker plots. Note that a new pattern will always be shown in a new plot.

## Note

The mads object contains all the information you need to make any desired plots. Check [mads-class](#) or the vignette *Multivariate Amputation using Ampute* to understand the contents of class object mads.

## Author(s)

Rianne Schouten, 2016

## See Also

[ampute](#), [bwplot](#), [Lattice](#) for an overview of the package, [mads-class](#)

bwplot.mids

*Box-and-whisker plot of observed and imputed data***Description**

Plotting methods for imputed data using **lattice**. `bwplot` produces box-and-whisker plots. The function automatically separates the observed and imputed data. The functions extend the usual features of **lattice**.

**Usage**

```
## S3 method for class 'mids'
bwplot(x, data, na.groups = NULL, groups = NULL,
       as.table = TRUE, theme = mice.theme(), mayreplicate = TRUE,
       allow.multiple = TRUE, outer = TRUE,
       drop.unused.levels = lattice::lattice.getOption("drop.unused.levels"), ...,
       subscripts = TRUE, subset = TRUE)
```

**Arguments**

<code>x</code>	A <code>mids</code> object, typically created by <code>mice()</code> or <code>mice.mids()</code> .
<code>data</code>	<p>Formula that selects the data to be plotted. This argument follows the <b>lattice</b> rules for <i>formulas</i>, describing the primary variables (used for the per-panel display) and the optional conditioning variables (which define the subsets plotted in different panels) to be used in the plot.</p> <p>The formula is evaluated on the complete data set in the long form. Legal variable names for the formula include <code>names(x\$data)</code> plus the two administrative factors <code>.imp</code> and <code>.id</code>.</p> <p><b>Extended formula interface:</b> The primary variable terms (both the LHS <code>y</code> and RHS <code>x</code>) may consist of multiple terms separated by a '+' sign, e.g., <code>y1 + y2 ~ x   a * b</code>. This formula would be taken to mean that the user wants to plot both <code>y1 ~ x   a * b</code> and <code>y2 ~ x   a * b</code>, but with the <code>y1 ~ x</code> and <code>y2 ~ x</code> in <i>separate panels</i>. This behavior differs from standard <b>lattice</b>. <i>Only combine terms of the same type</i>, i.e. only factors or only numerical variables. Mixing numerical and categorical data occasionally produces odds labeling of vertical axis.</p> <p>For convenience, in <code>stripplot()</code> and <code>bwplot</code> the formula <code>y~.imp</code> may be abbreviated as <code>y</code>. This applies only to a single <code>y</code>, and does not (yet) work for <code>y1+y2~.imp</code>.</p>
<code>na.groups</code>	<p>An expression evaluating to a logical vector indicating which two groups are distinguished (e.g. using different colors) in the display. The environment in which this expression is evaluated is the response indicator <code>is.na(x\$data)</code>.</p> <p>The default <code>na.group = NULL</code> contrasts the observed and missing data in the LHS <code>y</code> variable of the display, i.e. groups created by <code>is.na(y)</code>. The expression <code>y</code> creates the groups according to <code>is.na(y)</code>. The expression <code>y1 &amp; y2</code> creates groups by <code>is.na(y1) &amp; is.na(y2)</code>, and <code>y1   y2</code> creates groups as <code>is.na(y1)   is.na(y2)</code>, and so on.</p>

groups	This is the usual groups arguments in <b>lattice</b> . It differs from <code>na.groups</code> because it evaluates in the completed data <code>data.frame(complete(x, "long", inc=TRUE))</code> (as usual), whereas <code>na.groups</code> evaluates in the response indicator. See <a href="#">xyplot</a> for more details. When both <code>na.groups</code> and <code>groups</code> are specified, <code>na.groups</code> takes precedence, and <code>groups</code> is ignored.
as.table	See <a href="#">xyplot</a> .
theme	A named list containing the graphical parameters. The default function <code>mice.theme</code> produces a short list of default colors, line width, and so on. The extensive list may be obtained from <code>trellis.par.get()</code> . Global graphical parameters like <code>col</code> or <code>cex</code> in high-level calls are still honored, so first experiment with the global parameters. Many setting consists of a pair. For example, <code>mice.theme</code> defines two symbol colors. The first is for the observed data, the second for the imputed data. The theme settings only exist during the call, and do not affect the trellis graphical parameters.
mayreplicate	A logical indicating whether color, line widths, and so on, may be replicated. The graphical functions attempt to choose "intelligent" graphical parameters. For example, the same color can be replicated for different element, e.g. use all reds for the imputed data. Replication may be switched off by setting the flag to <code>FALSE</code> , in order to allow the user to gain full control.
allow.multiple	See <a href="#">xyplot</a> .
outer	See <a href="#">xyplot</a> .
drop.unused.levels	See <a href="#">xyplot</a> .
subscripts	See <a href="#">xyplot</a> .
subset	See <a href="#">xyplot</a> .
...	Further arguments, usually not directly processed by the high-level functions documented here, but instead passed on to other functions.

## Details

The argument `na.groups` may be used to specify (combinations of) missingness in any of the variables. The argument `groups` can be used to specify groups based on the variable values themselves. Only one of both may be active at the same time. When both are specified, `na.groups` takes precedence over `groups`.

Use the `subset` and `na.groups` together to plots parts of the data. For example, select the first imputed data set by `subset=.imp==1`.

Graphical parameters like `col`, `pch` and `cex` can be specified in the arguments list to alter the plotting symbols. If `length(col)==2`, the color specification to define the observed and missing groups. `col[1]` is the color of the 'observed' data, `col[2]` is the color of the missing or imputed data. A convenient color choice is `col=mdc(1:2)`, a transparent blue color for the observed data, and a transparent red color for the imputed data. A good choice is `col=mdc(1:2)`, `pch=20`, `cex=1.5`. These choices can be set for the duration of the session by running `mice.theme()`.

**Value**

The high-level functions documented here, as well as other high-level Lattice functions, return an object of class "trellis". The `update` method can be used to subsequently update components of the object, and the `print` method (usually called by default) will plot it on an appropriate plotting device.

**Note**

The first two arguments (`x` and `data`) are reversed compared to the standard Trellis syntax implemented in **lattice**. This reversal was necessary in order to benefit from automatic method dispatch.

In **mice** the argument `x` is always a `mids` object, whereas in **lattice** the argument `x` is always a formula.

In **mice** the argument `data` is always a formula object, whereas in **lattice** the argument `data` is usually a data frame.

All other arguments have identical interpretation.

**Author(s)**

Stef van Buuren

**References**

Sarkar, Deepayan (2008) *Lattice: Multivariate Data Visualization with R*, Springer. <http://lmdvr.r-forge.r-project.org/>

van Buuren S and Groothuis-Oudshoorn K (2011). `mice`: Multivariate Imputation by Chained Equations in R. *Journal of Statistical Software*, **45**(3), 1-67. <http://www.jstatsoft.org/v45/i03/>

**See Also**

[mice](#), [xyplot](#), [densityplot](#), [stripplot](#), [Lattice](#) for an overview of the package, as well as [bwplot](#), [panel.bwplot](#), [print.trellis](#), [trellis.par.set](#)

**Examples**

```
require(lattice)

imp <- mice(boys, maxit=1)

### box-and-whisker plot per imputation of all numerical variables
bwplot(imp)

### tv (testicular volume), conditional on region
bwplot(imp, tv~.imp|reg)

### same data, organized in a different way
bwplot(imp, tv~reg|.imp, theme=list())
```

---

 cbind.mids

*Columnwise combination of a mids object.*


---

### Description

This function combines two `mids` objects columnwise into a single object of class `mids`, or combines a `mids` object with a vector, matrix, factor or `data.frame` columnwise into an object of class `mids`. The number of rows in the (incomplete) data `x$data` and `y` (or `y$data` if `y` is a `mids` object) should be equal. If `y` is a `mids` object then the number of imputations in `x` and `y` should be equal. Note: If `y` is a vector or factor its original name is lost and it will be denoted with `y` in the `mids` object.

### Usage

```
cbind.mids(x, y, ...)
```

### Arguments

<code>x</code>	A <code>mids</code> object.
<code>y</code>	A <code>mids</code> object or a <code>data.frame</code> , matrix, factor or vector.
<code>...</code>	Additional <code>data.frame</code> , matrix, vector or factor. These can be given as named arguments.

### Value

An S3 object of class `mids`

### Note

Component `call` is a vector, with first argument the `mice()` statement that created `x` and second argument the call to `cbind.mids()`. Component `data` is the `codebind` of the (incomplete) data in `x$data` and `y$data`. Component `m` is the number of imputations. Component `nmis` is an array containing the number of missing observations per column. Component `imp` is a list of `nvar` components with the generated multiple imputations. Each part of the list is a `nmis[j]` by `m` matrix of imputed values for variable `j`. The original data of `y` will be copied into this list, including the missing values of `y` then `y` is not imputed. Component `method` is a vector of strings of length(`nvar`) specifying the elementary imputation method per column. If `y` is a `mids` object this vector is a combination of `x$method` and `y$method`, otherwise this vector is `x$method` and for the columns of `y` the method is set to `''`. Component `predictorMatrix` is a square matrix of size `ncol(data)` containing integer data specifying the predictor set. If `x` and `y` are `mids` objects then the predictor matrices of `x` and `y` are combined with zero matrices on the off-diagonal blocks. Otherwise the variables in `y` are included in the predictor matrix of `x` such that `y` is not used as predictor(s) and not imputed as well. Component `visitSequence` is the sequence in which columns are visited. The same as `x$visitSequence`. Component `seed` is the seed value of the solution, `x$seed`. Component `iteration` is the last Gibbs sampling iteration number, `x$iteration`. Component `lastSeedValue`



is the most recent seed value, `x$lastSeedValue` Component `chainMean` is the combination of `x$chainMean` and `y$chainMean`. If `y$chainMean` does not exist this element equals `x$chainMean`. Component `chainVar` is the combination of `x$chainVar` and `y$chainVar`. If `y$chainVar` does not exist this element equals `x$chainVar`. Component `pad` is a list containing various settings of the padded imputation model, i.e. the imputation model after creating dummy variables. This list is defined by combining `x$pad` and `y$pad` if `y` is a `mids` object. Otherwise, it is defined by the settings of `x` and the combination of the data `x$data` and `y`. Component `loggedEvents` is set to `x$loggedEvents`. If a column of `y` is categorical this is ignored in the padded model since that column is not used as predictor for another column.

### Author(s)

Karin Groothuis-Oudshoorn, Stef van Buuren, 2009

### See Also

[rbind.mids](#), [ibind](#), [mids](#)

### Examples

```
# append 'forgotten' variable bmi to imp
temp <- boys[,c(1:3,5:9)]
imp <- mice(temp,maxit=1,m=2)
imp2 <- cbind.mids(imp, data.frame(bmi=boys$bmi))

# append maturation score to imp (numerical)
mat <- (as.integer(temp$gen) + as.integer(temp$phb)
+ as.integer(cut(temp$tv,breaks=c(0,3,6,10,15,20,25))))
imp2 <- cbind.mids(imp, as.data.frame(mat))

# append maturation score to imp (factor)
# known issue: new column name is 'y', not 'mat'
mat <- as.factor(mat)
imp2 <- cbind.mids(imp, mat)

# append data frame with two columns to imp
temp2 <- data.frame(bmi=boys$bmi,mat=as.factor(mat))
imp2 <- cbind.mids(imp, temp2)

# combine two mids objects
impa <- mice(temp, maxit=1, m=2)
impb <- mice(temp2, maxit=2, m=2)

# first a then b
impab <- cbind.mids(impa, impb)

# first b then a
impba <- cbind.mids(impb, impa)
```

---

cc *Select complete cases*

---

**Description**

Extracts the complete cases, also known as *listwise deletion*. `cc(x)` is similar to `na.omit(x)`, but returns an object of the same class as the input data. Dimensions are not dropped. For extracting incomplete cases, use [ici](#).

**Usage**

```
cc(x)
```

**Arguments**

x An R object. Methods are available for classes `mids`, `data.frame` and `matrix`. Also, x could be a vector.

**Value**

A vector, matrix or data.frame containing the data of the complete cases.

**Author(s)**

Stef van Buuren, 2017.

**See Also**

[na.omit](#), [cci](#), [ici](#)

**Examples**

```
# cc(nhanes) # get the 13 complete cases
# cc(nhanes$bmi) # extract complete bmi
```

---

cci *Complete case indicator*

---

**Description**

The complete case indicator is useful for extracting the subset of complete cases. The function `cci(x)` calls `complete.cases(x)`. The companion function `ici()` selects the incomplete cases.

**Usage**

```
cci(x)
```

**Arguments**

`x` An R object. Currently supported are methods for the following classes: `mids`.

**Value**

Logical vector indicating the complete cases.

**Author(s)**

Stef van Buuren, 2017.

**See Also**

[complete.cases](#), [ici](#), [cc](#)

**Examples**

```
cci(nhanes) # indicator for 13 complete cases
cci(mice(nhanes, maxit = 0))
f <- cci(nhanes[,c("bmi", "hyp")]) # complete data for bmi and hyp
nhanes[f,] # obtain all data from those with complete bmi and hyp
```

---

complete

*Creates imputed data sets from a mids object*

---

**Description**

Takes an object of class `mids`, fills in the missing data, and returns the completed data in a specified format.

**Usage**

```
complete(x, action = 1, include = FALSE)
```

**Arguments**

`x` An object of class `mids` as created by the function `mice()`.

`action` If `action` is a scalar between 1 and `x$m`, the function returns the data with imputation number `action` filled in. Thus, `action=1` returns the first completed data set, `action=2` returns the second completed data set, and so on. The value of `action` can also be one of the following strings: `'long'`, `'broad'`, `'repeated'`. See 'Details' for the interpretation.

`include` Flag to indicate whether the original data with the missing values should be included. This requires that `action` is specified as `'long'`, `'broad'` or `'repeated'`.

## Details

The argument `action` can also be a string, which is partially matched as follows:

**`list('\long')`** produces a long data frame of vertically stacked imputed data sets with `nrow(x$data) * x$m` rows and `ncol(x$data)+2` columns. The two additional columns are labeled `.id` containing the row names of `x$data`, and `.imp` containing the imputation number. If `include=TRUE` then `nrow(x$data)` additional rows with the original data are appended with `.imp` set equal to `0`.

**`list('\broad')`** produces a broad data frame with `nrow(x$data)` rows and `ncol(x$data) * x$m` columns. Columns are ordered such that the first `ncol(x$data)` columns corresponds to the first imputed data matrix. The imputation number is appended to each column name. If `include=TRUE` then `ncol(x$data)` additional columns with the original data are appended. The number `.0` is appended to the column names.

**`list('\repeated')`** produces a broad data frame with `nrow(x$data)` rows and `ncol(x$data) * x$m` columns. Columns are ordered such that the first `x$m` columns correspond to the `x$m` imputed versions of the first column in `x$data`. The imputation number is appended to each column name. If `include=TRUE` then `ncol(x$data)` additional columns with the original data are appended. The number `.0` is appended to the column names.

## Value

A data frame with the imputed values filled in. Optionally, the original data are appended.

## Author(s)

Stef van Buuren, Karin Groothuis-Oudshoorn, 2009

## See Also

[mice](#), [mids](#)

## Examples

```
# do default multiple imputation on a numeric matrix
imp <- mice(nhanes)

# obtain first imputed matrix
mat <- complete(imp)

# fill in the third imputation
mat <- complete(imp, 3)

# long matrix with stacked complete data
mat <- complete(imp, 'long')

# long matrix with stacked complete data, including the original data
mat <- complete(imp, 'long', inc=TRUE)
```

```
# repeated matrix with complete data
mat <- complete(imp, 'r')

# for numeric data, produces a blocked correlation matrix, where
# each block contains of the same variable pair over different
# multiple imputations.
cor(mat)
```

---

densityplot.mids      *Density plot of observed and imputed data*

---

### Description

Plotting methods for imputed data using **lattice**. `densityplot` produces plots of the densities. The function automatically separates the observed and imputed data. The functions extend the usual features of **lattice**.

### Usage

```
## S3 method for class 'mids'
densityplot(x, data, na.groups = NULL, groups = NULL,
  as.table = TRUE, plot.points = FALSE, theme = mice.theme(),
  mayreplicate = TRUE, thicker = 2.5, allow.multiple = TRUE,
  outer = TRUE,
  drop.unused.levels = lattice::lattice.getOption("drop.unused.levels"),
  panel = lattice::lattice.getOption("panel.densityplot"),
  default.prepanel = lattice::lattice.getOption("prepanel.default.densityplot"),
  ..., subscripts = TRUE, subset = TRUE)
```

### Arguments

<code>x</code>	A <code>mids</code> object, typically created by <code>mice()</code> or <code>mice.mids()</code> .
<code>data</code>	<p>Formula that selects the data to be plotted. This argument follows the <b>lattice</b> rules for <i>formulas</i>, describing the primary variables (used for the per-panel display) and the optional conditioning variables (which define the subsets plotted in different panels) to be used in the plot.</p> <p>The formula is evaluated on the complete data set in the long form. Legal variable names for the formula include <code>names(x\$data)</code> plus the two administrative factors <code>.imp</code> and <code>.id</code>.</p> <p><b>Extended formula interface:</b> The primary variable terms (both the LHS <code>y</code> and RHS <code>x</code>) may consist of multiple terms separated by a '+' sign, e.g., <code>y1 + y2 ~ x   a * b</code>. This formula would be taken to mean that the user wants to plot both <code>y1 ~ x   a * b</code> and <code>y2 ~ x   a * b</code>, but with the <code>y1 ~ x</code> and <code>y2 ~ x</code> in <i>separate panels</i>. This behavior differs from standard <b>lattice</b>. <i>Only combine terms of the same type</i>, i.e. only factors or only numerical variables. Mixing numerical and categorical data occasionally produces odds labeling of vertical axis.</p>

	The function <code>densityplot</code> does not use the <code>y</code> terms in the formula. Density plots for <code>x1</code> and <code>x2</code> are requested as <code>~ x1 + x2</code> .
<code>na.groups</code>	An expression evaluating to a logical vector indicating which two groups are distinguished (e.g. using different colors) in the display. The environment in which this expression is evaluated in the response indicator is <code>is.na(x\$data)</code> . The default <code>na.group = NULL</code> contrasts the observed and missing data in the LHS <code>y</code> variable of the display, i.e. groups created by <code>is.na(y)</code> . The expression <code>y</code> creates the groups according to <code>is.na(y)</code> . The expression <code>y1 &amp; y2</code> creates groups by <code>is.na(y1) &amp; is.na(y2)</code> , and <code>y1   y2</code> creates groups as <code>is.na(y1)   is.na(y2)</code> , and so on.
<code>groups</code>	This is the usual <code>groups</code> arguments in <b>lattice</b> . It differs from <code>na.groups</code> because it evaluates in the completed data <code>data.frame(complete(x, "long", inc=TRUE))</code> (as usual), whereas <code>na.groups</code> evaluates in the response indicator. See <a href="#">xyplot</a> for more details. When both <code>na.groups</code> and <code>groups</code> are specified, <code>na.groups</code> takes precedence, and <code>groups</code> is ignored.
<code>as.table</code>	See <a href="#">xyplot</a> .
<code>plot.points</code>	A logical used in <code>densityplot</code> that signals whether the points should be plotted.
<code>theme</code>	A named list containing the graphical parameters. The default function <code>mice.theme</code> produces a short list of default colors, line width, and so on. The extensive list may be obtained from <code>trellis.par.get()</code> . Global graphical parameters like <code>col</code> or <code>cex</code> in high-level calls are still honored, so first experiment with the global parameters. Many setting consists of a pair. For example, <code>mice.theme</code> defines two symbol colors. The first is for the observed data, the second for the imputed data. The theme settings only exist during the call, and do not affect the <code>trellis</code> graphical parameters.
<code>mayreplicate</code>	A logical indicating whether color, line widths, and so on, may be replicated. The graphical functions attempt to choose "intelligent" graphical parameters. For example, the same color can be replicated for different element, e.g. use all reds for the imputed data. Replication may be switched off by setting the flag to <code>FALSE</code> , in order to allow the user to gain full control.
<code>thicker</code>	Used in <code>densityplot</code> . Multiplication factor of the line width of the observed density. <code>thicker=1</code> uses the same thickness for the observed and imputed data.
<code>allow.multiple</code>	See <a href="#">xyplot</a> .
<code>outer</code>	See <a href="#">xyplot</a> .
<code>drop.unused.levels</code>	See <a href="#">xyplot</a> .
<code>panel</code>	See <a href="#">xyplot</a> .
<code>default.prepanel</code>	See <a href="#">xyplot</a> .
<code>subscripts</code>	See <a href="#">xyplot</a> .
<code>subset</code>	See <a href="#">xyplot</a> .
<code>...</code>	Further arguments, usually not directly processed by the high-level functions documented here, but instead passed on to other functions.

## Details

The argument `na.groups` may be used to specify (combinations of) missingness in any of the variables. The argument `groups` can be used to specify groups based on the variable values themselves. Only one of both may be active at the same time. When both are specified, `na.groups` takes precedence over `groups`.

Use the `subset` and `na.groups` together to plots parts of the data. For example, select the first imputed data set by `subset=.imp==1`.

Graphical parameters like `col`, `pch` and `cex` can be specified in the arguments list to alter the plotting symbols. If `length(col)==2`, the color specification to define the observed and missing groups. `col[1]` is the color of the 'observed' data, `col[2]` is the color of the missing or imputed data. A convenient color choice is `col=mdc(1:2)`, a transparent blue color for the observed data, and a transparent red color for the imputed data. A good choice is `col=mdc(1:2)`, `pch=20`, `cex=1.5`. These choices can be set for the duration of the session by running `mice.theme()`.

## Value

The high-level functions documented here, as well as other high-level Lattice functions, return an object of class "trellis". The `update` method can be used to subsequently update components of the object, and the `print` method (usually called by default) will plot it on an appropriate plotting device.

## Note

The first two arguments (`x` and `data`) are reversed compared to the standard Trellis syntax implemented in **lattice**. This reversal was necessary in order to benefit from automatic method dispatch.

In **mice** the argument `x` is always a `mids` object, whereas in **lattice** the argument `x` is always a formula.

In **mice** the argument `data` is always a formula object, whereas in **lattice** the argument `data` is usually a data frame.

All other arguments have identical interpretation.

`densityplot` errs on empty groups, which occurs if all observations in the subgroup contain NA. The relevant error message is: `Error in density.default: ... need at least 2 points to select a bandwidth automatically`. There is yet no workaround for this problem. Use the more robust `bwplot` or `stripplot` as a replacement.

## Author(s)

Stef van Buuren

## References

Sarkar, Deepayan (2008) *Lattice: Multivariate Data Visualization with R*, Springer. <http://lmdvr.r-forge.r-project.org/>

van Buuren S and Groothuis-Oudshoorn K (2011). `mice`: Multivariate Imputation by Chained Equations in R. *Journal of Statistical Software*, **45**(3), 1-67. <http://www.jstatsoft.org/v45/i03/>

**See Also**

[mice](#), [xyplot](#), [stripplot](#), [bwplot](#), [Lattice](#) for an overview of the package, as well as [densityplot](#), [panel.densityplot](#), [print.trellis](#), [trellis.par.set](#)

**Examples**

```
require(lattice)

imp <- mice(boys, maxit=1)

### density plot of head circumference per imputation
### blue is observed, red is imputed
densityplot(imp, ~hc|.imp)

### All combined in one panel.
densityplot(imp, ~hc)
```

---

extractBS

*Extract broken stick estimates from a lmer object*

---

**Description**

Extract broken stick estimates from a lmer object

**Usage**

```
extractBS(fit)
```

**Arguments**

`fit` An object of class lmer

**Value**

A matrix containing broken stick estimates

**Author(s)**

Stef van Buuren, 2012



fdd

*SE Fireworks disaster data***Description**

Multiple outcomes of a randomized study to reduce post-traumatic stress.

**Format**

fdd is a data frame with 52 rows and 65 columns:

**id** Client number  
**trt** Treatment (E=EMDR, C=CBT)  
**pp** Per protocol (Y/N)  
**trtp** Number of parental treatments  
**sex** Sex: M/F  
**etn** Ethnicity: NL/OTHER  
**age** Age (years)  
**trauma** Trauma count (1-5)  
**prop1** PROPS total score T1  
**prop2** PROPS total score T2  
**prop3** PROPS total score T3  
**crop1** CROPS total score T1  
**crop2** CROPS total score T2  
**crop3** CROPS total score T3  
**masc1** MASC score T1  
**masc2** MASC score T2  
**masc3** MASC score T3  
**cbcl1** CBCL T1  
**cbcl3** CBCL T3  
**prs1** PRS total score T1  
**prs2** PRS total score T2  
**prs3** PRS total score T3  
**ypa1** PTSD-RI B intrusive recollection parent T1  
**ypb1** PTSD-RI C avoidant/numbing parent T1  
**ypc1** PTSD-RI D hyper-arousal parent T1  
**yp1** PTSD-RI B+C+D parent T1  
**ypa2** PTSD-RI B intrusive recollection parent T2  
**ypb2** PTSD-RI C avoidant/numbing parent T2

**ypc2** PTSD-RI D hyper-arousal parent T2  
**yp2** PTSD-RI B+C+D parent T1  
**ypa3** PTSD-RI B intrusive recollection parent T3  
**ypb3** PTSD-RI C avoidant/numbing parent T3  
**ypc3** PTSD-RI D hyper-arousal parent T3  
**yp3** PTSD-RI B+C+D parent T3  
**yca1** PTSD-RI B intrusive recollection child T1  
**ycb1** PTSD-RI C avoidant/numbing child T1  
**ycc1** PTSD-RI D hyper-arousal child T1  
**yc1** PTSD-RI B+C+D child T1  
**yca2** PTSD-RI B intrusive recollection child T2  
**ycb2** PTSD-RI C avoidant/numbing child T2  
**ycc2** PTSD-RI D hyper-arousal child T2  
**yc2** PTSD-RI B+C+D child T2  
**yca3** PTSD-RI B intrusive recollection child T3  
**ycb3** PTSD-RI C avoidant/numbing child T3  
**ycc3** PTSD-RI D hyper-arousal child T3  
**yc3** PTSD-RI B+C+D child T3  
**ypf1** PTSD-RI parent full T1  
**ypf2** PTSD-RI parent full T2  
**ypf3** PTSD-RI parent full T3  
**ypp1** PTSD parent partial T1  
**ypp2** PTSD parent partial T2  
**ypp3** PTSD parent partial T3  
**ycf1** PTSD child full T1  
**ycf2** PTSD child full T2  
**ycf3** PTSD child full T3  
**ycp1** PTSD child partial T1  
**ycp2** PTSD child partial T2  
**ycp3** PTSD child partial T3  
**cbin1** CBCL Internalizing T1  
**cbin3** CBCL Internalizing T3  
**cbex1** CBCL Externalizing T1  
**cbex3** CBCL Externalizing T3  
**bir1** Birlison T1  
**bir2** Birlison T2  
**bir3** Birlison T3

fdd.pred is the 65 by 65 binary predictor matrix used to impute fdd.

## Details

Data from a randomized experiment to reduce post-traumatic stress by two treatments: Eye Movement Desensitization and Reprocessing (EMDR) (experimental treatment), and cognitive behavioral therapy (CBT) (control treatment). 52 children were randomized to one of these two treatments. Outcomes were measured at three time points: at baseline (pre-treatment, T1), post-treatment (T2, 4-8 weeks), and at follow-up (T3, 3 months). For more details, see de Roos et al (2011). Some person covariates were reshuffled. The imputation methodology is explained in Chapter 9 of van Buuren (2012).

## Source

de Roos, C., Greenwald, R., den Hollander-Gijsman, M., Noorthoorn, E., van Buuren, S., de Jong, A. (2011). A Randomised Comparison of Cognitive Behavioral Therapy (CBT) and Eye Movement Desensitisation and Reprocessing (EMDR) in disaster-exposed children. *European Journal of Psychotraumatology*, 2, 5694.

van Buuren, S. (2012). *Flexible Imputation of Missing Data*. Boca Raton, FL: Chapman & Hall/CRC Press.

## Examples

```
data <- fdd
md.pattern(fdd)
```

---

fdgs

*Fifth Dutch growth study 2009*

---

## Description

Age, height, weight and region of 10030 children measured within the Fifth Dutch Growth Study 2009

## Format

fdgs is a data frame with 10030 rows and 8 columns:

**id** Person number  
**reg** Region (factor, 5 levels)  
**age** Age (years)  
**sex** Sex (boy, girl)  
**hgt** Height (cm)  
**wgt** Weight (kg)  
**hgt.z** Height Z-score  
**wgt.z** Weight Z-score

## Details

The data set contains data from children of Dutch descent (biological parents are born in the Netherlands). Children with growth-related diseases were excluded. The data were used to construct new growth charts of children of Dutch descent (Schonbeck 2013), and to calculate overweight and obesity prevalence (Schonbeck 2011).

Some groups were underrepresented. Multiple imputation was used to create synthetic cases that were used to correct for the nonresponse. See Van Buuren (2012), chapter 8 for details.

## Source

Schonbeck, Y., Talma, H., van Dommelen, P., Bakker, B., Buitendijk, S. E., Hirasing, R. A., van Buuren, S. (2011). Increase in prevalence of overweight in Dutch children and adolescents: A comparison of nationwide growth studies in 1980, 1997 and 2009. *PLoS ONE*, 6(11), e27608.

Schonbeck, Y., Talma, H., van Dommelen, P., Bakker, B., Buitendijk, S. E., Hirasing, R. A., \& van Buuren, S. (2013). The world's tallest nation has stopped growing taller: the height of Dutch children from 1955 to 2009. *Pediatric Research*, 73(3), 371-377.

van Buuren, S. (2012). *Flexible Imputation of Missing Data*. Boca Raton, FL: Chapman & Hall/CRC Press.

## Examples

```
data <- data(fdgs)
summary(data)
```

---

fico

*Fraction of incomplete cases among cases with observed*

---

## Description

FICO is an outbound statistic defined by the fraction of incomplete cases among cases with  $Y_j$  observed (White and Carlin, 2010).

## Usage

```
fico(data)
```

## Arguments

`data` A data frame or a matrix containing the incomplete data. Missing values are coded as NA's.

## Value

A vector of length `ncol(data)` of FICO statistics.

**Author(s)**

Stef van Buuren, 2012

**References**

van Buuren, S. (2012). *Flexible Imputation of Missing Data*. Boca Raton, FL: Chapman & Hall/CRC Press.

White, I.R., Carlin, J.B. (2010). Bias and efficiency of multiple imputation compared with complete-case analysis for missing covariate values. *Statistics in Medicine*, 29, 2920-2931.

**See Also**

[fluxplot](#), [flux](#), [md.pattern](#)

---

flux

*Influx and outflux of multivariate missing data patterns*

---

**Description**

Influx and outflux are statistics of the missing data pattern. These statistics are useful in selecting predictors that should go into the imputation model.

**Usage**

```
flux(data, local = names(data))
```

**Arguments**

data	A data frame or a matrix containing the incomplete data. Missing values are coded as NA's.
local	A vector of names of columns of data. The default is to include all columns in the calculations.

**Details**

Influx and outflux have been proposed by Van Buuren (2012), chapter 4.

Influx is equal to the number of variable pairs ( $Y_j$ ,  $Y_k$ ) with  $Y_j$  missing and  $Y_k$  observed, divided by the total number of observed data cells. Influx depends on the proportion of missing data of the variable. Influx of a completely observed variable is equal to 0, whereas for completely missing variables we have  $\text{influx} = 1$ . For two variables with the same proportion of missing data, the variable with higher influx is better connected to the observed data, and might thus be easier to impute.

Outflux is equal to the number of variable pairs with  $Y_j$  observed and  $Y_k$  missing, divided by the total number of incomplete data cells. Outflux is an indicator of the potential usefulness of  $Y_j$  for imputing other variables. Outflux depends on the proportion of missing data of the variable. Outflux of a completely observed variable is equal to 1, whereas outflux of a completely missing variable is equal to 0. For two variables having the same proportion of missing data, the variable with higher

outflux is better connected to the missing data, and thus potentially more useful for imputing other variables.

FICO is an outbound statistic defined by the fraction of incomplete cases among cases with  $Y_j$  observed (White and Carlin, 2010).

### Value

A data frame with `ncol(data)` rows and six columns: `pobs` = Proportion observed, `influx` = Influx, `outflux` = Outflux, `ainb` = Average inbound statistic, `aout` = Average outbound statistic, `fico` = Fraction of incomplete cases among cases with  $Y_j$  observed

### Author(s)

Stef van Buuren, 2012

### References

van Buuren, S. (2012). *Flexible Imputation of Missing Data*. Boca Raton, FL: Chapman & Hall/CRC Press.

White, I.R., Carlin, J.B. (2010). Bias and efficiency of multiple imputation compared with complete-case analysis for missing covariate values. *Statistics in Medicine*, 29, 2920-2931.

### See Also

[fluxplot](#), [md.pattern](#), [fico](#)

---

fluxplot

*Fluxplot of the missing data pattern*

---

### Description

Influx and outflux are statistics of the missing data pattern. These statistics are useful in selecting predictors that should go into the imputation model.

### Usage

```
fluxplot(data, local = names(data), plot = TRUE, labels = TRUE,
  xlim = c(0, 1), ylim = c(0, 1), las = 1, xlab = "Influx",
  ylab = "Outflux", main = paste("Influx-outflux pattern for",
  deparse(substitute(data))), eqsplot = TRUE, pty = "s", ...)
```

**Arguments**

<code>data</code>	A data frame or a matrix containing the incomplete data. Missing values are coded as NA's.
<code>local</code>	A vector of names of columns of data. The default is to include all columns in the calculations.
<code>plot</code>	Should a graph be produced?
<code>labels</code>	Should the points be labeled?
<code>xlim</code>	See par.
<code>ylim</code>	See par.
<code>las</code>	See par.
<code>xlab</code>	See par.
<code>ylab</code>	See par.
<code>main</code>	See par.
<code>eqsplot</code>	Should a square plot be produced?
<code>pty</code>	See par.
<code>...</code>	Further arguments passed to <code>plot()</code> or <code>eqsplot()</code> .

**Details**

Influx and outflux have been proposed by Van Buuren (2012), chapter 4.

Influx is equal to the number of variable pairs ( $Y_j$ ,  $Y_k$ ) with  $Y_j$  missing and  $Y_k$  observed, divided by the total number of observed data cells. Influx depends on the proportion of missing data of the variable. Influx of a completely observed variable is equal to 0, whereas for completely missing variables we have  $\text{influx} = 1$ . For two variables with the same proportion of missing data, the variable with higher influx is better connected to the observed data, and might thus be easier to impute.

Outflux is equal to the number of variable pairs with  $Y_j$  observed and  $Y_k$  missing, divided by the total number of incomplete data cells. Outflux is an indicator of the potential usefulness of  $Y_j$  for imputing other variables. Outflux depends on the proportion of missing data of the variable. Outflux of a completely observed variable is equal to 1, whereas outflux of a completely missing variable is equal to 0. For two variables having the same proportion of missing data, the variable with higher outflux is better connected to the missing data, and thus potentially more useful for imputing other variables.

**Value**

An invisible data frame with `ncol(data)` rows and six columns: `pobs` = Proportion observed, `influx` = Influx `outflux` = Outflux `ainb` = Average inbound statistic `aout` = Average outbound statistic `fico` = Fraction of incomplete cases among cases with  $Y_j$  observed

**Author(s)**

Stef van Buuren, 2012

## References

van Buuren, S. (2012). *Flexible Imputation of Missing Data*. Boca Raton, FL: Chapman & Hall/CRC Press.

White, I.R., Carlin, J.B. (2010). Bias and efficiency of multiple imputation compared with complete-case analysis for missing covariate values. *Statistics in Medicine*, 29, 2920-2931.

## See Also

[flux](#), [md.pattern](#), [fico](#)

---

getfit

*Extracts fit objects from mira object*

---

## Description

getfit returns the list of objects containing the repeated analysis results, or optionally, one of these fit objects.

## Usage

```
getfit(x, i = -1, simplify = FALSE)
```

## Arguments

x	An object of class <code>mira</code> , typically produced by a call to <code>with()</code> .
i	An integer between 1 and <code>x\$m</code> signalling the number of the repeated analysis. The default <code>i = -1</code> return a list with all analyses.
simplify	Should the return value be unlisted?

## Details

This function is shorthand notation for `x$analyses` and `x$analyses[[i]]`.

## Value

If `i = -1` an object containing all analyses, otherwise it returns the fitted object of the `i`'th repeated analysis.

## Author(s)

Stef van Buuren, March 2012.

## See Also

[mira](#), [with.mids](#)



## Examples

```
imp <- mice(nhanes)
fit <- with(imp, lm(bmi~chl+hyp))
getfit(fit)
getfit(fit, 2)
```

---

glm.mids

*Generalized linear model for mids object*

---

## Description

Applies `glm()` to a multiply imputed data set

## Usage

```
glm.mids(formula, family = gaussian, data, ...)
```

## Arguments

formula	a formula expression as for other regression models, of the form response ~ predictors. See the documentation of <a href="#">lm</a> and <a href="#">formula</a> for details.
family	The family of the glm model
data	An object of type <code>mids</code> , which stands for 'multiply imputed data set', typically created by function <code>mice()</code> .
...	Additional parameters passed to <a href="#">glm</a> .

## Details

This function is included for backward compatibility with V1.0. The function is superseded by [with.mids](#).

## Value

An objects of class `mira`, which stands for 'multiply imputed repeated analysis'. This object contains `data$m` distinct `glm` objects, plus some descriptive information.

## Author(s)

Stef van Buuren, Karin Groothuis-Oudshoorn, 2000

## References

Van Buuren, S., Groothuis-Oudshoorn, C.G.M. (2000) *Multivariate Imputation by Chained Equations: MICE V1.0 User's manual*. Leiden: TNO Quality of Life.

**See Also**

[with.mids](#), [glm](#), [mids](#), [mira](#)

**Examples**

```
imp <- mice(nhanes)

# logistic regression on the imputed data
fit <- glm.mids((hyp==2)~bmi+chl, data=imp, family = binomial)
fit
```

---

ibind

*Combine imputations fitted to the same data*

---

**Description**

This function combines two `mids` objects `x` and `y` into a single `mids` object. The two `mids` objects should have the same underlying multiple imputation model and should be fitted on exactly the same dataset. If the number of imputations in `x` is  $m(x)$  and in `y` is  $m(y)$  then the combination of both objects contains  $m(x)+m(y)$  imputations.

**Usage**

```
ibind(x, y)
```

**Arguments**

<code>x</code>	A <code>mids</code> object.
<code>y</code>	A <code>mids</code> object.

**Value**

An S3 object of class `mids`

**Author(s)**

Karin Groothuis-Oudshoorn, Stef van Buuren, 2009

**See Also**

[mids](#), [rbind.mids](#), [cbind.mids](#)

---

ic	<i>Select incomplete cases</i>
----	--------------------------------

---

**Description**

Extracts incomplete cases from a data set. The companion function for selecting the complete cases is [cc](#).

**Usage**

```
ic(x)
```

**Arguments**

x                    An R object. Methods are available for classes `mids`, `data.frame` and `matrix`. Also, x could be a vector.

**Value**

A vector, matrix or `data.frame` containing the data of the complete cases.

**Author(s)**

Stef van Buuren, 2017.

**See Also**

[cc](#), [ici](#)

**Examples**

```
ic(nhanes) # get the 12 rows with incomplete cases
ic(nhanes[1:10,]) # incomplete cases within the first ten rows
ic(nhanes[, c("bmi", "hyp")]) # restrict extraction to variables bmi and hyp
```

ici *Incomplete case indicator*

---

**Description**

This array is useful for extracting the subset of incomplete cases. The companion function `cci()` selects the complete cases.

**Usage**

```
ici(x)
```

**Arguments**

`x` An R object. Currently supported are methods for the following classes: `mids`.

**Value**

Logical vector indicating the incomplete cases,

**Author(s)**

Stef van Buuren, 2017.

**See Also**

[cci](#), [ic](#)

**Examples**

```
ici(nhanes) # indicator for 12 rows with incomplete cases
```

---

is.mads *Check for mads object*

---

**Description**

Check for mads object

**Usage**

```
is.mads(x)
```

**Arguments**

x                    An object

**Value**

A logical indicating whether x is an object of class mids

---

is.mids                    *Check for mids object*

---

**Description**

Check for mids object

**Usage**

is.mids(x)

**Arguments**

x                    An object

**Value**

A logical indicating whether x is an object of class mids

---

is.mipo                    *Check for mipo object*

---

**Description**

Check for mipo object

**Usage**

is.mipo(x)

**Arguments**

x                    An object

**Value**

A logical indicating whether x is an object of class mipo

---

<code>is.mira</code>	<i>Check for mira object</i>
----------------------	------------------------------

---

**Description**

Check for mira object

**Usage**

```
is.mira(x)
```

**Arguments**

<code>x</code>	An object
----------------	-----------

**Value**

A logical indicating whether `x` is an object of class `mira`

---

<code>leiden85</code>	<i>Leiden 85+ study</i>
-----------------------	-------------------------

---

**Description**

Subset of data from the Leiden 85+ study

**Format**

`leiden85` is a data frame with 956 rows and 336 columns.

**Details**

The data set concerns of subset of 956 members of a very old (85+) cohort in Leiden.

Multiple imputation of this data set has been described in Boshuizen et al (1998), Van Buuren et al (1999) and Van Buuren (2012), chapter 7.

The data set is not available as part of mice.

**Source**

Lagaay, A. M., van der Meij, J. C., Hijmans, W. (1992). Validation of medical history taking as part of a population based survey in subjects aged 85 and over. *Brit. Med. J.*, 304(6834), 1091-1092.

Izaks, G. J., van Houwelingen, H. C., Schreuder, G. M., Ligthart, G. J. (1997). The association between human leucocyte antigens (HLA) and mortality in community residents aged 85 and older. *Journal of the American Geriatrics Society*, 45(1), 56-60.

Boshuizen, H. C., Izaks, G. J., van Buuren, S., Ligthart, G. J. (1998). Blood pressure and mortality in elderly people aged 85 and older: Community based study. *Brit. Med. J.*, 316(7147), 1780-1784.

Van Buuren, S., Boshuizen, H.C., Knook, D.L. (1999) Multiple imputation of missing blood pressure covariates in survival analysis. *Statistics in Medicine*, 18, 681–694.

van Buuren, S. (2012). *Flexible Imputation of Missing Data*. Boca Raton, FL: Chapman & Hall/CRC Press.

---

lm.mids

*Linear regression for mids object*

---

**Description**

Applies `lm()` to multiply imputed data set

**Usage**

```
lm.mids(formula, data, ...)
```

**Arguments**

formula	a formula object, with the response on the left of a <code>~</code> operator, and the terms, separated by <code>+</code> operators, on the right. See the documentation of <code>lm</code> and <code>formula</code> for details.
data	An object of type 'mids', which stands for 'multiply imputed data set', typically created by a call to function <code>mice()</code> .
...	Additional parameters passed to <code>lm</code>

**Details**

This function is included for backward compatibility with V1.0. The function is superseded by `with.mids`.

**Value**

An objects of class `mira`, which stands for 'multiply imputed repeated analysis'. This object contains `data$m` distinct `lm` objects, plus some descriptive information.

**Author(s)**

Stef van Buuren, Karin Groothuis-Oudshoorn, 2000

## References

Van Buuren, S., Groothuis-Oudshoorn, K. (2011). mice: Multivariate Imputation by Chained Equations in R. *Journal of Statistical Software*, **45**(3), 1-67. <http://www.jstatsoft.org/v45/i03/>

## See Also

[lm](#), [mids](#), [mira](#)

## Examples

```
imp <- mice(nhanes)
fit <- lm.mids(bmi~hyp+chl, data = imp)
fit
```

---

mads-class

*Multivariate Amputed Data Set (mads)*

---

## Description

The mads object contains an amputed data set. The mads object is generated by the `ampute` function. The mads class of objects has methods for the following generic functions: `print`, `summary`, `bwplot` and `xyplot`.

## Contents

**call:** The function call.

**prop:** Proportion of cases with missing values. Note: even when the proportion is entered as the proportion of missing cells (when `bycases == TRUE`), this object contains the proportion of missing cases.

**patterns:** A data frame of size `#patterns` by `#variables` where 0 indicates a variable has missing values and 1 indicates a variable remains complete.

**freq:** A vector of length `#patterns` containing the relative frequency with which the patterns occur. For example, if the vector is `c(0.4, 0.4, 0.2)`, this means that of all cases with missing values, 40 percent is candidate for pattern 1, 40 percent for pattern 2 and 20 percent for pattern 3. The vector sums to 1.

**mech:** A string specifying the missingness mechanism, either "MCAR", "MAR" or "MNAR".

**weights:** A data frame of size `#patterns` by `#variables`. It contains the weights that were used to calculate the weighted sum scores. The weights may differ between patterns and between variables.

**cont:** Logical, whether probabilities are based on continuous logit functions or on discrete odds distributions.



**type:** A vector of strings containing the type of missingness for each pattern. Either "LEFT", "MID", "TAIL" or "RIGHT". The first type refers to the first pattern, the second type to the second pattern, etc.

**odds:** A matrix where #patterns defines the #rows. Each row contains the odds of being missing for the corresponding pattern. The amount of odds values defines in how many quantiles the sum scores were divided. The values are relative probabilities: a quantile with odds value 4 will have a probability of being missing that is four times higher than a quantile with odds 1. The #quantiles may differ between patterns, NA is used for cells remaining empty.

**amp:** A data frame containing the input data with NAs for the amputed values.

**cand:** A vector that contains the pattern number for each case. A value between 1 and #patterns is given. For example, a case with value 2 is candidate for missing data pattern 2.

**scores:** A list containing vectors with weighted sum scores of the candidates. The first vector refers to the candidates of the first pattern, the second vector refers to the candidates of the second pattern, etc. The length of the vectors differ because the number of candidates is different for each pattern.

**data:** The complete data set that was entered in ampute.

#### Note

Many of the functions of the mice package do not use the S4 class definitions, and instead rely on the S3 list equivalent `oldClass(obj) <- "mads"`.

#### Author(s)

Rianne Schouten, 2016

#### See Also

[ampute](#), Vignette titled "Multivariate Amputation using Ampute".

---

mammalsleep

*Mammal sleep data*

---

#### Description

Dataset from Allison and Cicchetti (1976) of 62 mammal species on the interrelationship between sleep, ecological, and constitutional variables. The dataset contains missing values on five variables.

#### Format

mammalsleep is a data frame with 62 rows and 11 columns:

**species** Species of animal

**bw** Body weight (kg)

**brw** Brain weight (g)

- sws** Slow wave ("nondreaming") sleep (hrs/day)
- ps** Paradoxical ("dreaming") sleep (hrs/day)
- ts** Total sleep (hrs/day) (sum of slow wave and paradoxical sleep)
- mls** Maximum life span (years)
- gt** Gestation time (days)
- pi** Predation index (1-5), 1 = least likely to be preyed upon
- sei** Sleep exposure index (1-5), 1 = least exposed (e.g. animal sleeps in a well-protected den), 5 = most exposed
- odi** Overall danger index (1-5) based on the above two indices and other information, 1 = least danger (from other animals), 5 = most danger (from other animals)

### Details

Allison and Cicchetti (1976) investigated the interrelationship between sleep, ecological, and constitutional variables. They assessed these variables for 39 mammalian species. The authors concluded that slow-wave sleep is negatively associated with a factor related to body size. This suggests that large amounts of this sleep phase are disadvantageous in large species. Also, paradoxical sleep (REM sleep) was associated with a factor related to predatory danger, suggesting that large amounts of this sleep phase are disadvantageous in prey species.

### Source

Allison, T., Cicchetti, D.V. (1976). Sleep in Mammals: Ecological and Constitutional Correlates. *Science*, 194(4266), 732-734.

### Examples

```
sleep <- data(mammalsleep)
```

---

md.pairs

*Missing data pattern by variable pairs*

---

### Description

Number of observations per variable pair.

### Usage

```
md.pairs(data)
```

**Arguments**

**data** A data frame or a matrix containing the incomplete data. Missing values are coded as NA.

**Details**

The four components in the output value is have the following interpretation:

**list('rr')** response-response, both variables are observed

**list('rm')** response-missing, row observed, column missing

**list('mr')** missing -response, row missing, column observed

**list('mm')** missing -missing, both variables are missing

**Value**

A list of four components named `rr`, `rm`, `mr` and `mm`. Each component is square numerical matrix containing the number observations within four missing data pattern.

**Author(s)**

Stef van Buuren, Karin Groothuis-Oudshoorn, 2009

**References**

Van Buuren, S., Groothuis-Oudshoorn, K. (2011). mice: Multivariate Imputation by Chained Equations in R. *Journal of Statistical Software*, **45**(3), 1-67. <http://www.jstatsoft.org/v45/i03/>

**Examples**

```
pat <- md.pairs(nhanes)
pat

# show that these four matrices decompose the total sample size
# for each pair
pat$rr + pat$rm + pat$mr + pat$mm

# percentage of usable cases to impute row variable from column variable
round(100*pat$mr/(pat$mr+pat$mm))
```

---

md.pattern	<i>Missing data pattern</i>
------------	-----------------------------

---

## Description

Display missing-data patterns.

## Usage

```
md.pattern(x)
```

## Arguments

x                    A data frame or a matrix containing the incomplete data. Missing values are coded as NA's.

## Details

This function is useful for investigating any structure of missing observation in the data. In specific case, the missing data pattern could be (nearly) monotone. Monotonicity can be used to simplify the imputation model. See Schafer (1997) for details. Also, the missing pattern could suggest which variables could potentially be useful for imputation of missing entries.

## Value

A matrix with  $\text{ncol}(x)+1$  columns, in which each row corresponds to a missing data pattern (1=observed, 0=missing). Rows and columns are sorted in increasing amounts of missing information. The last column and row contain row and column counts, respectively.

## Author(s)

Stef van Buuren, Karin Groothuis-Oudshoorn, 2000

## References

Schafer, J.L. (1997), Analysis of multivariate incomplete data. London: Chapman&Hall.

Van Buuren, S., Groothuis-Oudshoorn, K. (2011). mice: Multivariate Imputation by Chained Equations in R. *Journal of Statistical Software*, **45**(3), 1-67. <http://www.jstatsoft.org/v45/i03/>

## Examples

```
md.pattern(nhanes)
#   age hyp bmi chl
# 13  1  1  1  1  0
#   1  1  1  0  1  1
#   3  1  1  1  0  1
```

```
# 1 1 0 0 1 2
# 7 1 0 0 0 3
# 0 8 9 10 27
```

---

mdc

*Graphical parameter for missing data plots.*


---

## Description

mdc returns colors used to distinguish observed, missing and combined data in plotting. `mice` theme return a partial list of named objects that can be used as a theme in `stripplot`, `bwplot`, `densityplot` and `xypplot`.

## Usage

```
mdc(r = "observed", s = "symbol", transparent = TRUE, cso = hcl(240,
  100, 40, 0.7), csi = hcl(0, 100, 40, 0.7), csc = "gray50",
  clo = hcl(240, 100, 40, 0.8), cli = hcl(0, 100, 40, 0.8),
  clc = "gray50")
```

## Arguments

<code>r</code>	A numerical or character vector. The numbers 1-6 request colors as follows: 1=cso, 2=csi, 3=csc, 4=clo, 5=cli and 6=clc. Alternatively, <code>r</code> may contain the strings 'observed', 'missing', or 'both', or abbreviations thereof.
<code>s</code>	A character vector containing the strings 'symbol' or 'line', or abbreviations thereof.
<code>transparent</code>	A logical indicating whether alpha-transparency is allowed. The default is TRUE.
<code>cso</code>	The symbol color for the observed data. The default is a transparent blue.
<code>csi</code>	The symbol color for the missing or imputed data. The default is a transparent red.
<code>csc</code>	The symbol color for the combined observed and imputed data. The default is a grey color.
<code>clo</code>	The line color for the observed data. The default is a slightly darker transparent blue.
<code>cli</code>	The line color for the missing or imputed data. The default is a slightly darker transparent red.
<code>clc</code>	The line color for the combined observed and imputed data. The default is a grey color.

## Details

This function eases consistent use of colors in plots. The default follows the Abayomi convention, which uses blue for observed data, red for missing or imputed data, and black for combined data.

**Value**

`mdc()` returns a vector containing color definitions. The length of the output vector is calculate from the length of `r` and `s`. Elements of the input vectors are repeated if needed.

**Author(s)**

Stef van Buuren, sept 2012.

**References**

Sarkar, Deepayan (2008) *Lattice: Multivariate Data Visualization with R*, Springer. <http://lmdvr.r-forge.r-project.org/>

**See Also**

[hcl](#), [rgb](#), [xyplot.mids](#), [xyplot](#), [trellis.par.set](#)

**Examples**

```
# all six colors
mdc(1:6)

# lines color for observed and missing data
mdc(c('obs','mis'), 'lin')
```

---

mice

---

*Multivariate Imputation by Chained Equations (MICE)*


---

**Description**

Generates Multivariate Imputations by Chained Equations (MICE)

**Usage**

```
mice(data, m = 5, method = vector("character", length = ncol(data)),
      predictorMatrix = (1 - diag(1, ncol(data))),
      visitSequence = (1:ncol(data))[apply(is.na(data), 2, any)],
      form = vector("character", length = ncol(data)),
      post = vector("character", length = ncol(data)), defaultMethod = c("pmm",
      "logreg", "polyreg", "polr"), maxit = 5, diagnostics = TRUE,
      printFlag = TRUE, seed = NA, imputationMethod = NULL,
      defaultImputationMethod = NULL, data.init = NULL, ...)
```

**Arguments**

<code>data</code>	A data frame or a matrix containing the incomplete data. Missing values are coded as NA.
<code>m</code>	Number of multiple imputations. The default is <code>m=5</code> .
<code>method</code>	Can be either a single string, or a vector of strings with length <code>ncol(data)</code> , specifying the elementary imputation method to be used for each column in data. If specified as a single string, the same method will be used for all columns. The default imputation method (when no argument is specified) depends on the measurement level of the target column and are specified by the <code>defaultMethod</code> argument. Columns that need not be imputed have the empty method <code>' '</code> . See details for more information.
<code>predictorMatrix</code>	A square matrix of size <code>ncol(data)</code> containing 0/1 data specifying the set of predictors to be used for each target column. Rows correspond to target variables (i.e. variables to be imputed), in the sequence as they appear in data. A value of <code>'1'</code> means that the column variable is used as a predictor for the target variable (in the rows). The diagonal of <code>predictorMatrix</code> must be zero. The default for <code>predictorMatrix</code> is that all other columns are used as predictors (sometimes called massive imputation). Note: For two-level imputation codes <code>'2'</code> and <code>'-2'</code> are also allowed.
<code>visitSequence</code>	A vector of integers of arbitrary length, specifying the column indices of the visiting sequence. The visiting sequence is the column order that is used to impute the data during one pass through the data. A column may be visited more than once. All incomplete columns that are used as predictors should be visited, or else the function will stop with an error. The default sequence <code>1:ncol(data)</code> implies that columns are imputed from left to right. It is possible to specify one of the keywords <code>'roman'</code> (left to right), <code>'arabic'</code> (right to left), <code>'monotone'</code> (sorted in increasing amount of missingness) and <code>'revmonotone'</code> (reverse of monotone). The keyword should be supplied as a string and may be abbreviated.
<code>form</code>	A vector of strings with length <code>ncol(data)</code> , specifying formulae. Each string is parsed and executed within the <code>sampler()</code> function to create terms for the predictor. The default is to do nothing, indicated by a vector of empty strings <code>' '</code> . The main value lies in the easy specification of interaction terms. The user must ensure that the set of variables in the formula match those in predictors.
<code>post</code>	A vector of strings with length <code>ncol(data)</code> , specifying expressions. Each string is parsed and executed within the <code>sampler()</code> function to postprocess imputed values. The default is to do nothing, indicated by a vector of empty strings <code>' '</code> .
<code>defaultMethod</code>	A vector of three strings containing the default imputation methods for numerical columns, factor columns with 2 levels, and columns with (unordered or ordered) factors with more than two levels, respectively. If nothing is specified, the following defaults will be used: <code>pmm</code> , predictive mean matching (numeric data) <code>logreg</code> , logistic regression imputation (binary data, factor with 2 levels) <code>polyreg</code> , polytomous regression imputation for unordered categorical data (factor $\geq 2$ levels) <code>polr</code> , proportional odds model for (ordered, $\geq 2$ levels)
<code>maxit</code>	A scalar giving the number of iterations. The default is 5.

<code>diagnostics</code>	A Boolean flag. If TRUE, diagnostic information will be appended to the value of the function. If FALSE, only the imputed data are saved. The default is TRUE.
<code>printFlag</code>	If TRUE, mice will print history on console. Use <code>print=FALSE</code> for silent computation.
<code>seed</code>	An integer that is used as argument by the <code>set.seed()</code> for offsetting the random number generator. Default is to leave the random number generator alone.
<code>imputationMethod</code>	Same as <code>method</code> argument. Included for backwards compatibility.
<code>defaultImputationMethod</code>	Same as <code>defaultMethod</code> argument. Included for backwards compatibility.
<code>data.init</code>	A data frame of the same size and type as <code>data</code> , without missing data, used to initialize imputations before the start of the iterative process. The default NULL implies that starting imputation are created by a simple random draw from the data. Note that specification of <code>data.init</code> will start the <code>m</code> Gibbs sampling streams from the same imputations.
<code>...</code>	Named arguments that are passed down to the elementary imputation functions.

## Details

Generates multiple imputations for incomplete multivariate data by Gibbs sampling. Missing data can occur anywhere in the data. The algorithm imputes an incomplete column (the target column) by generating 'plausible' synthetic values given other columns in the data. Each incomplete column must act as a target column, and has its own specific set of predictors. The default set of predictors for a given target consists of all other columns in the data. For predictors that are incomplete themselves, the most recently generated imputations are used to complete the predictors prior to imputation of the target column.

A separate univariate imputation model can be specified for each column. The default imputation method depends on the measurement level of the target column. In addition to these, several other methods are provided. You can also write their own imputation functions, and call these from within the algorithm.

The data may contain categorical variables that are used in a regressions on other variables. The algorithm creates dummy variables for the categories of these variables, and imputes these from the corresponding categorical variable. The extended model containing the dummy variables is called the padded model. Its structure is stored in the list component `pad`.

Built-in elementary imputation methods are:

- pmm** Predictive mean matching (any)
- norm** Bayesian linear regression (numeric)
- norm.nob** Linear regression ignoring model error (numeric)
- norm.boot** Linear regression using bootstrap (numeric)
- norm.predict** Linear regression, predicted values (numeric)
- mean** Unconditional mean imputation (numeric)
- 2l.norm** Two-level normal imputation (numeric)
- 2l.pan** Two-level normal imputation using pan (numeric)



**2lonly.mean** Imputation at level-2 of the class mean (numeric)  
**2lonly.norm** Imputation at level-2 by Bayesian linear regression (numeric)  
**2lonly.pmm** Imputation at level-2 by Predictive mean matching (any)  
**quadratic** Imputation of quadratic terms (numeric)  
**logreg** Logistic regression (factor, 2 levels)  
**logreg.boot** Logistic regression with bootstrap  
**polyreg** Polytomous logistic regression (factor,  $\geq 2$  levels)  
**polr** Proportional odds model (ordered,  $\geq 2$  levels)  
**lda** Linear discriminant analysis (factor,  $\geq 2$  categories)  
**cart** Classification and regression trees (any)  
**rf** Random forest imputations (any)  
**ri** Random indicator method for nonignorable data (numeric)  
**sample** Random sample from the observed values (any)  
**fastpmm** Experimental: Fast predictive mean matching using C++ (any)

These corresponding functions are coded in the `mice` library under names `mice.impute.method`, where `method` is a string with the name of the elementary imputation method name, for example `norm`. The `method` argument specifies the methods to be used. For the  $j$ 'th column, `mice()` calls the first occurrence of `paste('mice.impute.', method[j], sep='')` in the search path. The mechanism allows users to write customized imputation function, `mice.impute.myfunc`. To call it for all columns specify `method='myfunc'`. To call it only for, say, column 2 specify `method=c('norm', 'myfunc', 'logreg', ...)`.

*Passive imputation:* `mice()` supports a special built-in method, called passive imputation. This method can be used to ensure that a data transform always depends on the most recently generated imputations. In some cases, an imputation model may need transformed data in addition to the original data (e.g. `log`, `quadratic`, `recodes`, `interaction`, `sum scores`, and so on).

Passive imputation maintains consistency among different transformations of the same data. Passive imputation is invoked if `~` is specified as the first character of the string that specifies the elementary method. `mice()` interprets the entire string, including the `~` character, as the formula argument in a call to `model.frame(formula, data[!r[, j], ])`. This provides a simple mechanism for specifying deterministic dependencies among the columns. For example, suppose that the missing entries in variables `data$height` and `data$weight` are imputed. The body mass index (BMI) can be calculated within `mice` by specifying the string `'~I(weight/height^2)'` as the elementary imputation method for the target column `data$bmi`. Note that the `~` mechanism works only on those entries which have missing values in the target column. You should make sure that the combined observed and imputed parts of the target column make sense. An easy way to create consistency is by coding all entries in the target as `NA`, but for large data sets, this could be inefficient. Note that you may also need to adapt the default `predictorMatrix` to evade linear dependencies among the predictors that could cause errors like `Error in solve.default()` or `Error: system is exactly singular`. Though not strictly needed, it is often useful to specify `visitSequence` such that the column that is imputed by the `~` mechanism is visited each time after one of its predictors was visited. In that way, deterministic relation between columns will always be synchronized.

## Value

Returns an S3 object of class `mids` (multiply imputed data set)

**Author(s)**

Stef van Buuren <stef.vanbuuren@tno.nl>, Karin Groothuis-Oudshoorn <c.g.m.oudshoorn@utwente.nl>, 2000-2010, with contributions of Alexander Robitzsch, Gerko Vink, Shahab Jolani, Roel de Jong, Jason Turner, Lisa Doove, John Fox, Frank E. Harrell, and Peter Malewski.

**References**

- Van Buuren, S., Groothuis-Oudshoorn, K. (2011). mice: Multivariate Imputation by Chained Equations in R. *Journal of Statistical Software*, **45**(3), 1-67. <http://www.jstatsoft.org/v45/i03/>
- van Buuren, S. (2012). *Flexible Imputation of Missing Data*. Boca Raton, FL: Chapman & Hall/CRC Press.
- Van Buuren, S., Brand, J.P.L., Groothuis-Oudshoorn C.G.M., Rubin, D.B. (2006) Fully conditional specification in multivariate imputation. *Journal of Statistical Computation and Simulation*, **76**, 12, 1049–1064.
- Van Buuren, S. (2007) Multiple imputation of discrete and continuous data by fully conditional specification. *Statistical Methods in Medical Research*, **16**, 3, 219–242.
- Van Buuren, S., Boshuizen, H.C., Knook, D.L. (1999) Multiple imputation of missing blood pressure covariates in survival analysis. *Statistics in Medicine*, **18**, 681–694.
- Brand, J.P.L. (1999) *Development, implementation and evaluation of multiple imputation strategies for the statistical analysis of incomplete data sets*. Dissertation. Rotterdam: Erasmus University.

**See Also**

[mids](#), [with.mids](#), [set.seed](#), [complete](#)

**Examples**

```
# do default multiple imputation on a numeric matrix
imp <- mice(nhanes)
imp

# list the actual imputations for BMI
imp$imputations$bmi

# first completed data matrix
complete(imp)

# imputation on mixed data with a different method per column
mice(nhanes2, meth=c('sample','pmm','logreg','norm'))
```

---

mice.impute.2l.norm     *Imputation by a two-level normal model*

---

### Description

Imputes univariate missing data using a two-level normal model

### Usage

```
mice.impute.2l.norm(y, ry, x, type, intercept = TRUE, ...)
```

### Arguments

y	Incomplete data vector of length n
ry	Vector of missing data pattern (FALSE=missing, TRUE=observed)
x	Matrix (n x p) of complete covariates.
type	Vector of length ncol(x) identifying random and class variables. Random variables are identified by a '2'. The class variable (only one is allowed) is coded as '-2'. Random variables also include the fixed effect.
intercept	Logical determining whether the intercept is automatically added.
...	Other named arguments.

### Details

Implements the Gibbs sampler for the linear multilevel model with heterogeneous with-class variance (Kasim and Raudenbush, 1998). Imputations are drawn as an extra step to the algorithm. For simulation work see Van Buuren (2011).

The random intercept is automatically added in `mice.impute.2l.norm()`. A model within a random intercept can be specified by `mice(..., intercept = FALSE)`.

### Value

A vector of length `nmi`s with imputations.

### Note

Added June 25, 2012: The currently implemented algorithm does not handle predictors that are specified as fixed effects (`type=1`). When using `mice.impute.2l.norm()`, the current advice is to specify all predictors as random effects (`type=2`).

Warning: The assumption of heterogeneous variances requires that in every class at least one observation has a response in `y`.

### Author(s)

Roel de Jong, 2008

## References

- Kasim RM, Raudenbush SW. (1998). Application of Gibbs sampling to nested variance components models with heterogeneous within-group variance. *Journal of Educational and Behavioral Statistics*, 23(2), 93–116.
- Van Buuren, S., Groothuis-Oudshoorn, K. (2011). mice: Multivariate Imputation by Chained Equations in R. *Journal of Statistical Software*, 45(3), 1-67. <http://www.jstatsoft.org/v45/i03/>
- Van Buuren, S. (2011) Multiple imputation of multilevel data. In Hox, J.J. and Roberts, J.K. (Eds.), *The Handbook of Advanced Multilevel Analysis*, Chapter 10, pp. 173–196. Milton Park, UK: Routledge.

---

mice.impute.2l.pan      *Imputation by a two-level normal model using pan*

---

## Description

Imputes univariate missing data using a two-level normal model with homogeneous within group variances. Aggregated group effects (i.e. group means) can be automatically created and included as predictors in the two-level regression (see argument type). This function needs the pan package.

## Usage

```
mice.impute.2l.pan(y, ry, x, type, intercept = TRUE, paniter = 500,
  groupcenter.slope = FALSE, ...)
```

## Arguments

y	Incomplete data vector of length n
ry	Vector of missing data pattern (FALSE=missing, TRUE=observed)
x	Matrix (n x p) of complete covariates.
type	Vector of length ncol(x) identifying random and class variables. Random effects are identified by a '2'. The group variable (only one is allowed) is coded as '-2'. Random effects also include the fixed effect. If for a covariates X1 group means shall be calculated and included as further fixed effects choose '3'. In addition to the effects in '3', specification '4' also includes random effects of X1.
intercept	Logical determining whether the intercept is automatically added.
paniter	Number of iterations in pan. Default is 500.
groupcenter.slope	If TRUE, in case of group means (type is '3' or '4') group mean centering for these predictors are conducted before doing imputations. Default is FALSE.
...	Other named arguments.

**Details**

Implements the Gibbs sampler for the linear two-level model with homogeneous within group variances which is a special case of a multivariate linear mixed effects model (Schafer & Yucel, 2002). For a two-level imputation with heterogeneous within-group variances see [mice.impute.2l.norm](#). The random intercept is automatically added in `mice.impute.2l.norm()`.

**Value**

A vector of length `nmiss` with imputations.

**Author(s)**

Alexander Robitzsch (Federal Institute for Education Research, Innovation, and Development of the Austrian School System, Salzburg, Austria), <a.robitzsch@bifie.at>

**References**

Schafer J L, Yucel RM (2002). Computational strategies for multivariate linear mixed-effects models with missing values. *Journal of Computational and Graphical Statistics*. **11**, 437-457.

Van Buuren, S., Groothuis-Oudshoorn, K. (2011). mice: Multivariate Imputation by Chained Equations in R. *Journal of Statistical Software*, **45**(3), 1-67. <http://www.jstatsoft.org/v45/i03/>

**See Also**

[mice.impute.2l.norm](#)

**Examples**

```
#####
# simulate some data
# two-level regression model with fixed slope

# number of groups
G <- 250
# number of persons
n <- 20
# regression parameter
beta <- .3
# intraclass correlation
rho <- .30
# correlation with missing response
rho.miss <- .10
# missing proportion
missrate <- .50
y1 <- rep( rnorm( G , sd = sqrt( rho ) ) , each=n ) + rnorm(G*n , sd = sqrt( 1 - rho ))
x <- rnorm( G*n )
y <- y1 + beta * x
dfr0 <- dfr <- data.frame( "group" = rep(1:G , each=n ) , "x" = x , "y" = y )
dfr[ rho.miss * x + rnorm( G*n , sd = sqrt( 1 - rho.miss ) ) < qnorm( missrate ) , "y" ] <- NA
```

```

#....
# empty imputation in mice
imp0 <- mice( as.matrix(dfr) , maxit=0 )
predM <- imp0$predictorMatrix
impM <- imp0$method

#...
# specify predictor matrix and imputationMethod
predM1 <- predM
predM1["y","group"] <- -2
predM1["y","x"] <- 1      # fixed x effects imputation
impM1 <- impM
impM1["y"] <- "2l.pan"

# multilevel imputation
imp1 <- mice( as.matrix( dfr ) , m = 1 , predictorMatrix = predM1 ,
             imputationMethod = impM1 , maxit=1 )
# multilevel analysis
library(lme4)
mod <- lmer( y ~ ( 1 + x | group) + x , data = complete(imp1) )
summary(mod)

#####
# Examples of predictorMatrix specification

# random x effects
# predM1["y","x"] <- 2

# fixed x effects and group mean of x
# predM1["y","x"] <- 3

# random x effects and group mean of x
# predM1["y","x"] <- 4

```

---

mice.impute.2lonly.mean

*Imputation of the mean within the class*

---

## Description

Imputes the mean of within the class

## Usage

```
mice.impute.2lonly.mean(y, ry, x, type, ...)
```

**Arguments**

y	Incomplete data vector of length n
ry	Vector of missing data pattern (FALSE=missing, TRUE=observed)
x	Matrix (n x p) of complete covariates.
type	Vector of length ncol(x) identifying random and class variables. The class variable (only one is allowed) is coded as '-2'.
...	Other named arguments.

**Details**

Observed values in y are averaged within the class, and replicated to the missing y within that class. If there are no observed data in the class, all entries of the class are set to NaN. This function is primarily useful for repairing incomplete data that are constant within the class, but that vary over the classes.

**Value**

A vector of length nmi s with imputations.

**Author(s)**

Gerko Vink, Stef van Buuren, 2013

---

mice.impute.2lonly.norm

*Imputation at level 2 by Bayesian linear regression*

---

**Description**

Imputes univariate missing data at level 2 using Bayesian linear regression analysis. Variables are level 1 are aggregated at level 2. The group identifier at level 2 must be indicated by type=-2 in the predictorMatrix.

**Usage**

```
mice.impute.2lonly.norm(y, ry, x, type, ...)
```

**Arguments**

y	Incomplete data vector of length n
ry	Vector of missing data pattern (FALSE=missing, TRUE=observed)
x	Matrix (n x p) of complete covariates. Only numeric variables are permitted for usage of this function.
type	Group identifier must be specified by '-2'. Predictors must be specified by '1'.
...	Other named arguments.

**Details**

This function allows in combination with [mice.impute.2l.pan](#) switching regression imputation between level 1 and level 2 as described in Yucel (2008) or Gelman and Hill (2007, p. 541).

**Value**

A vector of length `nmis` with imputations.

**Author(s)**

Alexander Robitzsch (Federal Institute for Education Research, Innovation, and Development of the Austrian School System, Salzburg, Austria), <a.robitzsch@bifie.at>

**References**

Gelman, A. and Hill, J. (2007). *Data analysis using regression and multilevel/hierarchical models*. Cambridge, Cambridge University Press.

Yucel, RM (2008). Multiple imputation inference for multivariate multilevel continuous data with ignorable non-response. *Philosophical Transactions of the Royal Society A*, **366**, 2389-2404.

**See Also**

[mice.impute.norm](#), [mice.impute.2lonly.pmm](#), [mice.impute.2l.pan](#)

**Examples**

```
#####
# simulate some data
# x,y ... level 1 variables
# v,w ... level 2 variables

G <- 250          # number of groups
n <- 20           # number of persons
beta <- .3        # regression coefficient
rho <- .30        # residual intraclass correlation
rho.miss <- .10   # correlation with missing response
missrate <- .50   # missing proportion
y1 <- rep( rnorm( G , sd = sqrt( rho ) ) , each=n ) + rnorm(G*n , sd = sqrt( 1 - rho ))
w <- rep( round( rnorm(G) , 2 ) , each=n )
v <- rep( round( runif( G , 0 , 3 ) ) , each=n )
x <- rnorm( G*n )
y <- y1 + beta * x + .2 * w + .1 * v
dfr0 <- dfr <- data.frame( "group" = rep(1:G , each=n ) , "x" = x , "y" = y , "w" = w , "v" = v )
dfr[ rho.miss * x + rnorm( G*n , sd = sqrt( 1 - rho.miss ) ) < qnorm( missrate ) , "y" ] <- NA
dfr[ rep( rnorm(G) , each=n ) < qnorm( missrate ) , "w" ] <- NA
dfr[ rep( rnorm(G) , each=n ) < qnorm( missrate ) , "v" ] <- NA

#...
# empty mice imputation
imp0 <- mice( as.matrix(dfr) , maxit=0 )
```



```

predM <- imp0$predictorMatrix
impM <- imp0$method

#...
# multilevel imputation
predM1 <- predM
predM1[c("w","y","v"),"group"] <- -2
predM1["y","x"] <- 1      # fixed x effects imputation
impM1 <- impM
impM1[c("y","w","v")] <- c("2l.pan" , "2lonly.norm" , "2lonly.pmm" )

# y ... imputation using pan
# w ... imputation at level 2 using norm
# v ... imputation at level 2 using pmm

imp1 <- mice( as.matrix( dfr ) , m = 1 , predictorMatrix = predM1 ,
              imputationMethod = impM1 , maxit=1 , paniter=500)

```

---

mice.impute.2lonly.pmm

*Imputation at level 2 by predictive mean matching*

---

## Description

Imputes univariate missing data at level 2 using predictive mean matching. Variables are level 1 are aggregated at level 2. The group identifier at level 2 must be indicated by type=-2 in the predictorMatrix.

## Usage

```
mice.impute.2lonly.pmm(y, ry, x, type, ...)
```

## Arguments

y	Incomplete data vector of length n
ry	Vector of missing data pattern (FALSE=missing, TRUE=observed)
x	Matrix (n x p) of complete covariates. Only numeric variables are permitted for usage of this function.
type	Group identifier must be specified by '-2'. Predictors must be specified by '1'.
...	Other named arguments.

## Details

This function allows in combination with [mice.impute.2l.pan](#) switching regression imputation between level 1 and level 2 as described in Yucel (2008) or Gelman and Hill (2007, p. 541).

**Value**

A vector of length `nmis` with imputations.

**Author(s)**

Alexander Robitzsch (Federal Institute for Education Research, Innovation, and Development of the Austrian School System, Salzburg, Austria), <a.robitzsch@bifie.at>

**References**

Gelman, A. and Hill, J. (2007). *Data analysis using regression and multilevel/hierarchical models*. Cambridge, Cambridge University Press.

Yucel, RM (2008). Multiple imputation inference for multivariate multilevel continuous data with ignorable non-response. *Philosophical Transactions of the Royal Society A*, **366**, 2389-2404.

**See Also**

[mice.impute.pmm](#), [mice.impute.2lonly.norm](#), [mice.impute.2l.pan](#)

**Examples**

```
#####
# simulate some data
# x,y ... level 1 variables
# v,w ... level 2 variables

G <- 250          # number of groups
n <- 20           # number of persons
beta <- .3        # regression coefficient
rho <- .30        # residual intraclass correlation
rho.miss <- .10   # correlation with missing response
missrate <- .50   # missing proportion
y1 <- rep( rnorm( G , sd = sqrt( rho ) ) , each=n ) + rnorm(G*n , sd = sqrt( 1 - rho ))
w <- rep( round( rnorm(G) , 2 ) , each=n )
v <- rep( round( runif( G , 0 , 3 ) ) , each=n )
x <- rnorm( G*n )
y <- y1 + beta * x + .2 * w + .1 * v
dfr0 <- dfr <- data.frame( "group" = rep(1:G , each=n ) , "x" = x , "y" = y , "w" = w , "v" = v )
dfr[ rho.miss * x + rnorm( G*n , sd = sqrt( 1 - rho.miss ) ) < qnorm( missrate ) , "y" ] <- NA
dfr[ rep( rnorm(G) , each=n ) < qnorm( missrate ) , "w" ] <- NA
dfr[ rep( rnorm(G) , each=n ) < qnorm( missrate ) , "v" ] <- NA

#...
# empty mice imputation
imp0 <- mice( as.matrix(dfr) , maxit=0 )
predM <- imp0$predictorMatrix
impM <- imp0$method

#...
# multilevel imputation
```

```

predM1 <- predM
predM1[c("w","y","v"),"group"] <- -2
predM1["y","x"] <- 1          # fixed x effects imputation
impM1 <- impM
impM1[c("y","w","v")] <- c("2l.pan" , "2lonly.norm" , "2lonly.pmm" )

# y ... imputation using pan
# w ... imputation at level 2 using norm
# v ... imputation at level 2 using pmm

imp1 <- mice( as.matrix( dfr ) , m = 1 , predictorMatrix = predM1 ,
              imputationMethod = impM1 , maxit=1 , paniter=500)

```

---

mice.impute.cart      *Imputation by classification and regression trees*

---

## Description

Imputes univariate missing data using classification and regression trees.

## Usage

```
mice.impute.cart(y, ry, x, minbucket = 5, cp = 1e-04, ...)
```

## Arguments

y	Numeric vector with incomplete data
ry	Response pattern of y (TRUE = observed, FALSE = missing)
x	Design matrix with length(y) rows and p columns containing complete covariates.
minbucket	The minimum number of observations in any terminal node used. See <a href="#">rpart.control</a> for details.
cp	Complexity parameter. Any split that does not decrease the overall lack of fit by a factor of cp is not attempted. See <a href="#">rpart.control</a> for details.
...	Other named arguments passed down to <code>rpart()</code> .

## Details

Imputation of y by classification and regression trees. The procedure is as follows:

1. Fit a classification or regression tree by recursive partitioning;
2. For each y<sub>m</sub>i's, find the terminal node they end up according to the fitted tree;
3. Make a random draw among the member in the node, and take the observed value from that draw as the imputation.

**Value**

Numeric vector of length `sum(!ry)` with imputations

**Author(s)**

Lisa Doove, Stef van Buuren, Elise Dusseldorp, 2012

**References**

Doove, L.L., van Buuren, S., Dusseldorp, E. (2014), Recursive partitioning for missing data imputation in the presence of interaction Effects. *Computational Statistics & Data Analysis*, 72, 92-104.

Breiman, L., Friedman, J. H., Olshen, R. A., and Stone, C. J. (1984), *Classification and regression trees*, Monterey, CA: Wadsworth & Brooks/Cole Advanced Books & Software.

Van Buuren, S.(2012), *Flexible imputation of missing data*, Boca Raton, FL: Chapman & Hall/CRC.

**See Also**

[mice](#), [mice.impute.rf](#), [rpart](#), [rpart.control](#)

**Examples**

```
require(rpart)
require(lattice)

imp <- mice(nhanes2, meth = "cart", minbucket = 4)
plot(imp)
```

---

`mice.impute.fastpmm`    *Imputation by fast predictive mean matching*

---

**Description**

Imputes univariate missing data using fast predictive mean matching

**Usage**

```
mice.impute.fastpmm(y, ry, x, donors = 5, type = 1, ridge = 1e-05,
  version = "", ...)
```

**Arguments**

<code>y</code>	Numeric vector with incomplete data
<code>ry</code>	Response pattern of <code>y</code> (TRUE=observed, FALSE=missing)
<code>x</code>	Design matrix with <code>length(y)</code> rows and <code>p</code> columns containing complete covariates.

donors	The size of the donor pool among which a draw is made. The default is donors = 5. Setting donors = 1 always selects the closest match. Values between 3 and 10 provide the best results. Note: The default was changed from 3 to 5 in version 2.19, based on simulation work by Tim Morris.
type	Type of matching distance. The default choice type = 1 calculates the distance between the predicted value of jobs and the drawn values of ymis. Other choices are type = 0 (distance between predicted values) and type = 2 (distance between drawn values). The current version supports only type = 1.
ridge	The ridge penalty applied in .norm.draw() to prevent problems with multicollinearity. The default is ridge = 1e-05, which means that 0.01 percent of the diagonal is added to the cross-product. Larger ridges may result in more biased estimates. For highly noisy data (e.g. many junk variables), set ridge = 1e-06 or even lower to reduce bias. For highly collinear data, set ridge = 1e-04 or higher.
version	A character variable indicating the version. Currently unused.
...	Other named arguments.

## Details

Imputation of  $y$  by predictive mean matching, based on Rubin (1987, p. 168, formulas a and b). The procedure is as follows:

1. Estimate beta and sigma by linear regression
2. Draw beta\* and sigma\* from the proper posterior
3. Compute predicted values for jobs beta and ymis beta\*
4. For each ymis, find donors observations with closest predicted values, randomly sample one of these, and take its observed value in  $y$  as the imputation.
5. Ties are broken by making a random draw among ties. Note: The matching is done on predicted  $y$ , NOT on observed  $y$ .

## Value

Numeric vector of length `sum(!ry)` with imputations

## Note

The `mice.impute.fastpmm()` function is an experimental version of the standard `mice.impute.pmm()` function. In `mice 2.22` both are equivalent. In future versions of `mice` the `mice.impute.fastpmm()` function may be subject to additional optimizations. This is an experimental feature.

## Author(s)

Stef van Buuren, Karin Groothuis-Oudshoorn, 2000, 2012

## References

- Little, R.J.A. (1988), Missing data adjustments in large surveys (with discussion), *Journal of Business Economics and Statistics*, 6, 287–301.
- Rubin, D.B. (1987). *Multiple imputation for nonresponse in surveys*. New York: Wiley.
- Van Buuren, S., Brand, J.P.L., Groothuis-Oudshoorn C.G.M., Rubin, D.B. (2006) Fully conditional specification in multivariate imputation. *Journal of Statistical Computation and Simulation*, **76**, 12, 1049–1064.
- Van Buuren, S., Groothuis-Oudshoorn, K. (2011). mice: Multivariate Imputation by Chained Equations in R. *Journal of Statistical Software*, **45**(3), 1-67. <http://www.jstatsoft.org/v45/i03/>

## See Also

[mice.impute.pmm](#)

---

mice.impute.lda	<i>Imputation by linear discriminant analysis</i>
-----------------	---

---

## Description

Imputes univariate missing data using linear discriminant analysis

## Usage

```
mice.impute.lda(y, ry, x, ...)
```

## Arguments

y	Incomplete data vector of length n
ry	Vector of missing data pattern (FALSE=missing, TRUE=observed)
x	Matrix (n x p) of complete covariates.
...	Other named arguments.

## Details

Imputation of categorical response variables by linear discriminant analysis. This function uses the Venables/Ripley functions `lda()` and `predict.lda()` to compute posterior probabilities for each incomplete case, and draws the imputations from this posterior.

## Value

A vector of length `nmi.s` with imputations.

## Warning

The function does not incorporate the variability of the discriminant weight, so it is not 'proper' in the sense of Rubin. For small samples and rare categories in the y, variability of the imputed data could therefore be somewhat underestimated.

**Note**

This function can be called from within the Gibbs sampler by specifying 'lda' in the method argument of `mice()`. This method is usually faster and uses fewer resources than calling the function `mice.impute.polyreg`.

**Author(s)**

Stef van Buuren, Karin Groothuis-Oudshoorn, 2000

**References**

Van Buuren, S., Groothuis-Oudshoorn, K. (2011). `mice`: Multivariate Imputation by Chained Equations in R. *Journal of Statistical Software*, **45**(3), 1-67. <http://www.jstatsoft.org/v45/i03/>

Brand, J.P.L. (1999). Development, Implementation and Evaluation of Multiple Imputation Strategies for the Statistical Analysis of Incomplete Data Sets. Ph.D. Thesis, TNO Prevention and Health/Erasmus University Rotterdam. ISBN 90-74479-08-1.

Venables, W.N. & Ripley, B.D. (1997). Modern applied statistics with S-PLUS (2nd ed). Springer, Berlin.

**See Also**

`mice`, `link{mice.impute.polyreg}`, `lda`

---

`mice.impute.logreg`      *Imputation by logistic regression*

---

**Description**

Imputes univariate missing data using logistic regression.

**Usage**

```
mice.impute.logreg(y, ry, x, ...)
```

**Arguments**

<code>y</code>	Incomplete data vector of length <code>n</code>
<code>ry</code>	Vector of missing data pattern of length <code>n</code> (FALSE=missing, TRUE=observed)
<code>x</code>	Matrix ( <code>n x p</code> ) of complete covariates.
<code>...</code>	Other named arguments.

## Details

Imputation for binary response variables by the Bayesian logistic regression model (Rubin 1987, p. 169-170). The Bayesian method consists of the following steps:

1. Fit a logit, and find  $(\hat{\beta}, V(\hat{\beta}))$
2. Draw BETA from  $N(\hat{\beta}, V(\hat{\beta}))$
3. Compute predicted scores for m.d., i.e.  $\text{logit}^{-1}(X \text{ BETA})$
4. Compare the score to a random (0,1) deviate, and impute.

The method relies on the standard `glm.fit` function. Warnings from `glm.fit` are suppressed. Perfect prediction is handled by the data augmentation method.

## Value

A vector of length `nmi.s` with imputations (0 or 1).

## Author(s)

Stef van Buuren, Karin Groothuis-Oudshoorn, 2000, 2011

## References

- Van Buuren, S., Groothuis-Oudshoorn, K. (2011). `mice`: Multivariate Imputation by Chained Equations in R. *Journal of Statistical Software*, **45**(3), 1-67. <http://www.jstatsoft.org/v45/i03/>
- Brand, J.P.L. (1999). Development, Implementation and Evaluation of Multiple Imputation Strategies for the Statistical Analysis of Incomplete Data Sets. Ph.D. Thesis, TNO Prevention and Health/Erasmus University Rotterdam. ISBN 90-74479-08-1.
- Venables, W.N. & Ripley, B.D. (1997). Modern applied statistics with S-Plus (2nd ed). Springer, Berlin.
- White, I., Daniel, R. and Royston, P (2010). Avoiding bias due to perfect prediction in multiple imputation of incomplete categorical variables. *Computational Statistics and Data Analysis*, 54:22672275.

## See Also

[mice](#), [glm](#), [glm.fit](#)

---

`mice.impute.logreg.boot`

*Imputation by logistic regression using the bootstrap*

---

## Description

Imputes univariate missing data using logistic regression by a bootstrapped logistic regression model. The bootstrap method draws a simple bootstrap sample with replacement from the observed data `y[ry]` and `x[ry, ]`. Perfect prediction is handled by the data augmentation method.



**Usage**

```
mice.impute.logreg.boot(y, ry, x, ...)
```

**Arguments**

y	Incomplete data vector of length n
ry	Vector of missing data pattern of length n (FALSE=missing, TRUE=observed)
x	Matrix (n x p) of complete covariates.
...	Other named arguments.

**Value**

A vector of length `nmi` with imputations (0 or 1).

**Author(s)**

Stef van Buuren, Karin Groothuis-Oudshoorn, 2000, 2011

**References**

Van Buuren, S., Groothuis-Oudshoorn, K. (2011). `mice`: Multivariate Imputation by Chained Equations in R. *Journal of Statistical Software*, **45**(3), 1-67. <http://www.jstatsoft.org/v45/i03/>

Brand, J.P.L. (1999). Development, Implementation and Evaluation of Multiple Imputation Strategies for the Statistical Analysis of Incomplete Data Sets. Ph.D. Thesis, TNO Prevention and Health/Erasmus University Rotterdam. ISBN 90-74479-08-1.

Venables, W.N. & Ripley, B.D. (1997). *Modern applied statistics with S-Plus* (2nd ed). Springer, Berlin.

White, I., Daniel, R. and Royston, P (2010). Avoiding bias due to perfect prediction in multiple imputation of incomplete categorical variables. *Computational Statistics and Data Analysis*, 54:22672275.

**See Also**

[mice](#), [glm](#), [glm.fit](#)

---

mice.impute.mean	<i>Imputation by the mean</i>
------------------	-------------------------------

---

**Description**

Imputes the arithmetic mean of the observed data

**Usage**

```
mice.impute.mean(y, ry, x = NULL, ...)
```

**Arguments**

y	Incomplete data vector of length n
ry	Vector of missing data pattern (FALSE=missing, TRUE=observed)
x	Matrix (n x p) of complete covariates.
...	Other named arguments.

**Value**

A vector of length nmi s with imputations.

**Warning**

Imputing the mean of a variable is almost never appropriate. See Little and Rubin (1987).

**Author(s)**

Stef van Buuren, Karin Groothuis-Oudshoorn, 2000

**References**

Van Buuren, S., Groothuis-Oudshoorn, K. (2011). mice: Multivariate Imputation by Chained Equations in R. *Journal of Statistical Software*, **45**(3), 1-67. <http://www.jstatsoft.org/v45/i03/>

Little, R.J.A. and Rubin, D.B. (1987). *Statistical Analysis with Missing Data*. New York: John Wiley and Sons.

**See Also**

[mice](#), [mean](#)

---

mice.impute.midastouch

*Predictive Mean Matching with distance aided selection of donors*

---

**Description**

Imputes univariate missing data using predictive mean matching.

**Usage**

```
mice.impute.midastouch(y, ry, x, ridge = 1e-05,  
midas.kappa = NULL, outout = TRUE, neff = NULL, debug = NULL, ...)
```

**Arguments**

<code>y</code>	Numeric vector with incomplete data
<code>ry</code>	Response pattern of <code>y</code> (TRUE=observed, FALSE=missing)
<code>x</code>	Design matrix with <code>length(y)</code> rows and <code>p</code> columns containing complete covariates.
<code>ridge</code>	The ridge penalty applied to prevent problems with multicollinearity. The default is <code>ridge = 1e-05</code> , which means that 0.001 percent of the diagonal is added to the cross-product. Larger ridges may result in more biased estimates. For highly noisy data (e.g. many junk variables), set <code>ridge = 1e-06</code> or even lower to reduce bias. For highly collinear data, set <code>ridge = 1e-04</code> or higher.
<code>midas.kappa</code>	Scalar. If NULL (default) then the optimal kappa gets selected automatically. Alternatively, the user may specify a scalar. Siddique and Belin 2008 find <code>midas.kappa = 3</code> to be sensible.
<code>outout</code>	Logical. If TRUE (default) one model is estimated for each donor (leave-one-out principle). For speedup choose <code>outout = FALSE</code> , which estimates one model for all observations leading to in-sample predictions for the donors and out-of-sample predictions for the recipients. Mind the inappropriateness, though.
<code>neff</code>	FOR EXPERTS. Null or character string. The name of an existing environment in which the effective sample size of the donors for each loop (CE iterations times multiple imputations) is supposed to be written. The effective sample size is necessary to compute the correction for the total variance as originally suggested by Parzen, Lipsitz and Fitzmaurice 2005. The objectname is <code>midastouch.neff</code> .
<code>debug</code>	FOR EXPERTS. Null or character string. The name of an existing environment in which the input is supposed to be written. The objectname is <code>midastouch.inputlist</code> .
<code>...</code>	Other named arguments.

**Details**

Imputation of `y` by predictive mean matching, based on Rubin (1987, p. 168, formulas a and b) and Siddique and Belin 2008. The procedure is as follows:

1. Draw a bootstrap sample from the donor pool.
2. Estimate a beta matrix on the bootstrap sample by the leave one out principle.
3. Compute type II predicted values for `yobs` (`nobs x 1`) and `ymis` (`nmis x nobs`).
4. Calculate the distance between all `yobs` and the corresponding `ymis`.
5. Convert the distances in drawing probabilities.
6. For each recipient draw a donor from the entire pool while considering the probabilities from the model.
7. Take its observed value in `y` as the imputation.

**Value**

Numeric vector of length `sum(!ry)` with imputations

**Author(s)**

Philipp Gaffert, Florian Meinfelder, Volker Bosch 2015

**References**

Gaffert, P., Meinfelder, F., Bosch V. (2015) Towards an MI-proper Predictive Mean Matching, Discussion Paper. [https://www.uni-bamberg.de/fileadmin/uni/fakultaeten/sowi\\_lehrstuehle/statistik/Personen/Dateien\\_Florian/properPMM.pdf](https://www.uni-bamberg.de/fileadmin/uni/fakultaeten/sowi_lehrstuehle/statistik/Personen/Dateien_Florian/properPMM.pdf)

Little, R.J.A. (1988), Missing data adjustments in large surveys (with discussion), *Journal of Business Economics and Statistics*, 6, 287–301.

Parzen, M., Lipsitz, S. R., Fitzmaurice, G. M. (2005), A note on reducing the bias of the approximate bayesian bootstrap imputation variance estimator. *Biometrika* **92**, 4, 971–974.

Rubin, D.B. (1987), *Multiple imputation for nonresponse in surveys*. New York: Wiley.

Siddique, J., Belin, T.R. (2008), Multiple imputation using an iterative hot-deck with distance-based donor selection. *Statistics in medicine*, **27**, 1, 83–102

Van Buuren, S., Brand, J.P.L., Groothuis-Oudshoorn C.G.M., Rubin, D.B. (2006), Fully conditional specification in multivariate imputation. *Journal of Statistical Computation and Simulation*, **76**, 12, 1049–1064.

Van Buuren, S., Groothuis-Oudshoorn, K. (2011), mice: Multivariate Imputation by Chained Equations in R. *Journal of Statistical Software*, **45**, 3, 1–67. <http://www.jstatsoft.org/v45/i03/>

**Examples**

```
# do default multiple imputation on a numeric matrix
imp <- mice(nhanes, method = 'midastouch')
imp

# list the actual imputations for BMI
imp$imp$bmi

# first completed data matrix
complete(imp)

# imputation on mixed data with a different method per column
mice(nhanes2, method = c('sample', 'midastouch', 'logreg', 'norm'))
```

---

mice.impute.norm

*Imputation by Bayesian linear regression*

---

**Description**

Imputes univariate missing data using Bayesian linear regression analysis

**Usage**

```
mice.impute.norm(y, ry, x, ...)
```

**Arguments**

y	Incomplete data vector of length n
ry	Vector of missing data pattern (FALSE=missing, TRUE=observed)
x	Matrix (n x p) of complete covariates.
...	Other named arguments.

**Details**

Draws values of beta and sigma for Bayesian linear regression imputation of y given x according to Rubin p. 167.

**Value**

A vector of length nmis with imputations.

**Note**

Using mice.impute.norm for all columns is similar to Schafer's NORM method (Schafer, 1997).

**Author(s)**

Stef van Buuren, Karin Groothuis-Oudshoorn, 2000

**References**

- Van Buuren, S., Groothuis-Oudshoorn, K. (2011). mice: Multivariate Imputation by Chained Equations in R. *Journal of Statistical Software*, **45**(3), 1-67. <http://www.jstatsoft.org/v45/i03/>
- Brand, J.P.L. (1999) *Development, implementation and evaluation of multiple imputation strategies for the statistical analysis of incomplete data sets*. Dissertation. Rotterdam: Erasmus University.
- Schafer, J.L. (1997). *Analysis of incomplete multivariate data*. London: Chapman & Hall.

---

mice.impute.norm.boot *Imputation by linear regression, bootstrap method*

---

**Description**

Imputes univariate missing data using linear regression with bootstrap

**Usage**

```
mice.impute.norm.boot(y, ry, x, ridge = 1e-05, ...)
```

**Arguments**

y	Incomplete data vector of length n
ry	Vector of missing data pattern (FALSE=missing, TRUE=observed)
x	Matrix (n x p) of complete covariates.
ridge	Ridge parameter
...	Other named arguments.

**Details**

Draws a bootstrap sample from  $x[ry, ]$  and  $y[ry]$ , calculates regression weights and imputes with normal residuals. The ridge parameter adds a penalty term  $\text{ridge} \cdot \text{diag}(x^T x)$  to the variance-covariance matrix  $x^T x$ .

**Value**

A vector of length `nmi`s with imputations.

**Author(s)**

Stef van Buuren, 2011

**References**

Van Buuren, S., Groothuis-Oudshoorn, K. (2011). mice: Multivariate Imputation by Chained Equations in R. *Journal of Statistical Software*, **45**(3), 1-67. <http://www.jstatsoft.org/v45/i03/>

---

mice.impute.norm.nob *Imputation by linear regression (non Bayesian)*

---

**Description**

Imputes univariate missing data using linear regression analysis (non Bayesian version)

**Usage**

```
mice.impute.norm.nob(y, ry, x, ...)
```

**Arguments**

y	Incomplete data vector of length n
ry	Vector of missing data pattern (FALSE=missing, TRUE=observed)
x	Matrix (n x p) of complete covariates.
...	Other named arguments.

**Details**

This creates imputation using the spread around the fitted linear regression line of  $y$  given  $x$ , as fitted on the observed data.

**Value**

A vector of length `nmi`s with imputations.

**Warning**

The function does not incorporate the variability of the regression weights, so it is not 'proper' in the sense of Rubin. For small samples, variability of the imputed data is therefore underestimated.

**Note**

This function is provided mainly to allow comparison between proper and improper norm methods. Also, it may be useful to impute large data containing many rows.

**Author(s)**

Stef van Buuren, Karin Groothuis-Oudshoorn, 2000

**References**

Van Buuren, S., Groothuis-Oudshoorn, K. (2011). `mice`: Multivariate Imputation by Chained Equations in R. *Journal of Statistical Software*, **45**(3), 1-67. <http://www.jstatsoft.org/v45/i03/>

Brand, J.P.L. (1999). Development, Implementation and Evaluation of Multiple Imputation Strategies for the Statistical Analysis of Incomplete Data Sets. Ph.D. Thesis, TNO Prevention and Health/Erasmus University Rotterdam.

**See Also**

[mice](#), [mice.impute.norm](#)

---

`mice.impute.norm.predict`

*Imputation by linear regression, prediction method*

---

**Description**

Imputes univariate missing data using the predicted value from a linear regression

**Usage**

```
mice.impute.norm.predict(y, ry, x, ridge = 1e-05, ...)
```

**Arguments**

y	Incomplete data vector of length n
ry	Vector of missing data pattern (FALSE=missing, TRUE=observed)
x	Matrix (n x p) of complete covariates.
ridge	Ridge parameter
...	Other named arguments.

**Details**

Calculates regression weights from the observed data and return predicted values to as imputations. The ridge parameter adds a penalty term  $\text{ridge} \cdot \text{diag}(x^T x)$  to the variance-covariance matrix  $x^T x$ .

**Value**

A vector of length `nmi.s` with imputations.

**Author(s)**

Stef van Buuren, 2011

**References**

Van Buuren, S., Groothuis-Oudshoorn, K. (2011). mice: Multivariate Imputation by Chained Equations in R. *Journal of Statistical Software*, **45**(3), 1-67. <http://www.jstatsoft.org/v45/i03/>

---

mice.impute.passive    *Passive imputation*

---

**Description**

Derive a new variable based on the imputed data

**Usage**

```
mice.impute.passive(data, func)
```

**Arguments**

data	A data frame
func	A formula specifying the transformations on data



**Details**

Passive imputation is a special internal imputation function. Using this facility, the user can specify, at any point in the mice Gibbs sampling algorithm, a function on the imputed data. This is useful, for example, to compute a cubic version of a variable, a transformation like  $Q = W/H^2$  based on two variables, or a mean variable like  $(x_1+x_2+x_3)/3$ . The so derived variables might be used in other places in the imputation model. The function allows to dynamically derive virtually any function of the imputed data at virtually any time.

**Value**

The result of applying formula

**Author(s)**

Stef van Buuren, Karin Groothuis-Oudshoorn, 2000

**References**

Van Buuren, S., Groothuis-Oudshoorn, K. (2011). mice: Multivariate Imputation by Chained Equations in R. *Journal of Statistical Software*, **45**(3), 1-67. <http://www.jstatsoft.org/v45/i03/>

**See Also**

[mice](#)

---

mice.impute.pmm	<i>Imputation by predictive mean matching</i>
-----------------	---

---

**Description**

Imputes univariate missing data using predictive mean matching

**Usage**

```
mice.impute.pmm(y, ry, x, donors = 5, type = 1, ridge = 1e-05,
  version = "", ...)
```

**Arguments**

y	Numeric vector with incomplete data
ry	Response pattern of y (TRUE=observed, FALSE=missing)
x	Design matrix with length(y) rows and p columns containing complete covariates.
donors	The size of the donor pool among which a draw is made. The default is donors = 5. Setting donors = 1 always selects the closest match. Values between 3 and 10 provide the best results. Note: The default was changed from 3 to 5 in version 2.19, based on simulation work by Tim Morris.

type	Type of matching distance. The default choice <code>type = 1</code> calculates the distance between the predicted value of jobs and the drawn values of ymis. Other choices are <code>type = 0</code> (distance between predicted values) and <code>type = 2</code> (distance between drawn values). The current version supports only <code>type = 1</code> .
ridge	The ridge penalty applied in <code>.norm.draw()</code> to prevent problems with multicollinearity. The default is <code>ridge = 1e-05</code> , which means that 0.01 percent of the diagonal is added to the cross-product. Larger ridges may result in more biased estimates. For highly noisy data (e.g. many junk variables), set <code>ridge = 1e-06</code> or even lower to reduce bias. For highly collinear data, set <code>ridge = 1e-04</code> or higher.
version	A character variable indicating the version to be used. Specifying <code>version = "2.21"</code> calls <code>.pmm.match()</code> instead of the default <code>matcher()</code> function.
...	Other named arguments.

### Details

Imputation of y by predictive mean matching, based on Rubin (1987, p. 168, formulas a and b). The procedure is as follows:

1. Estimate beta and sigma by linear regression
2. Draw beta\* and sigma\* from the proper posterior
3. Compute predicted values for jobs beta and ymis beta\*
4. For each ymis, find donors observations with closest predicted values, randomly sample one of these, and take its observed value in y as the imputation.
5. Ties are broken by making a random draw among ties. Note: The matching is done on predicted y, NOT on observed y.

### Value

Numeric vector of length `sum(!ry)` with imputations

### Note

Since `mice 2.22` the standard `mice.impute.pmm()` calls the much faster `matcher()` function instead of `.pmm.match()`. Since `matcher()` uses its own random generator, results cannot be exactly reproduced. In case where you want the old `.pmm.match()`, specify `mice(..., version = "2.21")`.

### Author(s)

Stef van Buuren, Karin Groothuis-Oudshoorn, 2000, 2012

### References

- Little, R.J.A. (1988), Missing data adjustments in large surveys (with discussion), *Journal of Business Economics and Statistics*, 6, 287–301.
- Rubin, D.B. (1987). *Multiple imputation for nonresponse in surveys*. New York: Wiley.

Van Buuren, S., Brand, J.P.L., Groothuis-Oudshoorn C.G.M., Rubin, D.B. (2006) Fully conditional specification in multivariate imputation. *Journal of Statistical Computation and Simulation*, **76**, 12, 1049–1064.

Van Buuren, S., Groothuis-Oudshoorn, K. (2011). mice: Multivariate Imputation by Chained Equations in R. *Journal of Statistical Software*, **45**(3), 1-67. <http://www.jstatsoft.org/v45/i03/>

mice.impute.polr      *Imputation by polytomous regression - ordered*

## Description

Imputes missing data in a categorical variable using polytomous regression

## Usage

```
mice.impute.polr(y, ry, x, nnet.maxit = 100, nnet.trace = FALSE,
  nnet.MaxNWts = 1500, ...)
```

## Arguments

y	Incomplete data vector of length n
ry	Vector of missing data pattern (FALSE=missing, TRUE=observed)
x	Matrix (n x p) of complete covariates.
nnet.maxit	Tuning parameter for nnet().
nnet.trace	Tuning parameter for nnet().
nnet.MaxNWts	Tuning parameter for nnet().
...	Other named arguments.

## Details

By default, ordered factors with more than two levels are imputed by `mice.impute.polr`.

The function `mice.impute.polr()` imputes for ordered categorical response variables by the proportional odds logistic regression (polr) model. The function repeatedly applies logistic regression on the successive splits. The model is also known as the cumulative link model.

The algorithm of `mice.impute.polr` uses the function `polr()` from the MASS package.

In order to avoid bias due to perfect prediction, the algorithm augment the data according to the method of White, Daniel and Royston (2010).

The call to `polr` might fail, usually because the data are very sparse. In that case, `multinom` is tried as a fallback, and a record is written to the `loggedEvents` component of the `mids` object.

## Value

A vector of length `nmi.s` with imputations.

**Author(s)**

Stef van Buuren, Karin Groothuis-Oudshoorn, 2000-2010

**References**

Van Buuren, S., Groothuis-Oudshoorn, K. (2011). mice: Multivariate Imputation by Chained Equations in R. *Journal of Statistical Software*, **45**(3), 1-67. <http://www.jstatsoft.org/v45/i03/>

Brand, J.P.L. (1999) *Development, implementation and evaluation of multiple imputation strategies for the statistical analysis of incomplete data sets*. Dissertation. Rotterdam: Erasmus University.

White, I.R., Daniel, R. Royston, P. (2010). Avoiding bias due to perfect prediction in multiple imputation of incomplete categorical variables. *Computational Statistics and Data Analysis*, **54**, 2267-2275.

Venables, W.N. & Ripley, B.D. (2002). *Modern applied statistics with S-Plus (4th ed)*. Springer, Berlin.

**See Also**

[mice](#), [multinom](#), [polr](#)

---

mice.impute.polyreg    *Imputation by polytomous regression - unordered*

---

**Description**

Imputes missing data in a categorical variable using polytomous regression

**Usage**

```
mice.impute.polyreg(y, ry, x, nnet.maxit = 100, nnet.trace = FALSE,
  nnet.MaxNWts = 1500, ...)
```

**Arguments**

y	Incomplete data vector of length n
ry	Vector of missing data pattern (FALSE=missing, TRUE=observed)
x	Matrix (n x p) of complete covariates.
nnet.maxit	Tuning parameter for nnet().
nnet.trace	Tuning parameter for nnet().
nnet.MaxNWts	Tuning parameter for nnet().
...	Other named arguments.

## Details

By default, unordered factors with more than two levels are imputed by `mice.impute.polyreg()`. The function `mice.impute.polyreg()` imputes categorical response variables by the Bayesian polytomous regression model. See J.P.L. Brand (1999), Chapter 4, Appendix B.

The method consists of the following steps:

1. Fit categorical response as a multinomial model
2. Compute predicted categories
3. Add appropriate noise to predictions.

The algorithm of `mice.impute.polyreg` uses the function `multinom()` from the `nnet` package.

In order to avoid bias due to perfect prediction, the algorithm augment the data according to the method of White, Daniel and Royston (2010).

## Value

A vector of length `nmi.s` with imputations.

## Author(s)

Stef van Buuren, Karin Groothuis-Oudshoorn, 2000-2010

## References

Van Buuren, S., Groothuis-Oudshoorn, K. (2011). `mice`: Multivariate Imputation by Chained Equations in R. *Journal of Statistical Software*, **45**(3), 1-67. <http://www.jstatsoft.org/v45/i03/>

Brand, J.P.L. (1999) *Development, implementation and evaluation of multiple imputation strategies for the statistical analysis of incomplete data sets*. Dissertation. Rotterdam: Erasmus University.

White, I.R., Daniel, R. Royston, P. (2010). Avoiding bias due to perfect prediction in multiple imputation of incomplete categorical variables. *Computational Statistics and Data Analysis*, **54**, 2267-2275.

Venables, W.N. & Ripley, B.D. (2002). *Modern applied statistics with S-Plus (4th ed)*. Springer, Berlin.

## See Also

[mice](#), [multinom](#), [polr](#)

---

mice.impute.quadratic *Imputation of quadratic terms*

---

### Description

Imputes univariate missing data of incomplete variable that appears as both main effect and quadratic effect in the complete-data model.

### Usage

```
mice.impute.quadratic(y, ry, x, ...)
```

### Arguments

y	Incomplete data vector of length n
ry	Vector of missing data pattern (FALSE=missing, TRUE=observed)
x	Matrix (n x p) of complete covariates.
...	Other named arguments.

### Details

This implements polynomial combination method. First, the polynomial combination  $Z = Y \beta_1 + Y^2 \beta_2$  is formed.  $Z$  is imputed by predictive mean matching, followed by a decomposition of the imputed data  $Z$  into components  $Y$  and  $Y^2$ . See Van Buuren (2012, pp. 139-141) and Vink et al (2012) for more details. The method ensures that 1) the imputed data for  $Y$  and  $Y^2$  are mutually consistent, and 2) that provides unbiased estimates of the regression weights in a complete-data linear regression that use both  $Y$  and  $Y^2$ .

### Value

A vector of length `nmi`s with imputations.

### Note

There are two situations to consider. If only the linear term  $Y$  is present in the data, calculate the quadratic term  $YY$  after imputation. If both the linear term  $Y$  and the the quadratic term  $YY$  are variables in the data, then first impute  $Y$  by calling `mice.impute.quadratic()` on  $Y$ , and then impute  $YY$  by passive imputation as `meth["YY"] <- "~I(Y^2)"`. See example section for details. Generally, we would like  $YY$  to be present in the data if we need to preserve quadratic relations between  $YY$  and any third variables in the multivariate incomplete data that we might wish to impute.

### Author(s)

Gerko Vink (University of Utrecht), <g.vink@uu.nl>

## References

van Buuren, S. (2012). *Flexible Imputation of Missing Data*. Boca Raton, FL: Chapman & Hall/CRC Press.

Vink, G., Frank, L.E., van Buuren, S. (2012). Multiple Imputation of Squares. *Sociological Methods & Research*, accepted for publication.

## See Also

[mice.impute.pmm](#)

## Examples

```
require(lattice)

# Create Data
B1=.5
B2=.5
X<-rnorm(1000)
XX<-X^2
e<-rnorm(1000, 0, 1)
Y <- B1*X+B2*XX+e
dat <- data.frame(x=X, xx=XX, y=Y)

# Impose 25 percent MCAR Missingness
dat[0 == rbinom(1000, 1, 1-.25), 1:2] <- NA

# Prepare data for imputation
ini <- mice(dat, maxit=0)
meth <- c("quadratic", "~I(x^2)", "")
pred <- ini$pred
pred[,"xx"] <- 0

# Impute data
imp <- mice(dat, meth=meth, pred=pred)

# Pool results
pool(with(imp, lm(y~x+xx)))

# Plot results
stripplot(imp)
plot(dat$x, dat$xx, col=mdc(1), xlab="x", ylab="xx")
points(complete(imp,1)$x[is.na(dat$x)], complete(imp,1)$xx[is.na(dat$x)], col=mdc(2))
```

**Description**

Imputes univariate missing data using random forests.

**Usage**

```
mice.impute.rf(y, ry, x, ntree = 10, ...)
```

**Arguments**

<code>y</code>	Numeric vector with incomplete data
<code>ry</code>	Response pattern of <code>y</code> (TRUE = observed, FALSE = missing)
<code>x</code>	Design matrix with <code>length(y)</code> rows and <code>p</code> columns containing complete covariates.
<code>ntree</code>	The number of trees to grow. The default is 10.
<code>...</code>	Other named arguments passed down to <code>randomForest()</code> and <code>randomForest:::randomForest.default</code>

**Details**

Imputation of `y` by random forests. The method calls `randomForrest()` which implements Breiman's random forest algorithm (based on Breiman and Cutler's original Fortran code) for classification and regression. See Appendix A.1 of Doove et al. (2014) for the definition of the algorithm used. An alternative implementation was independently developed by Shah et al (2014), and is available in the package `CALIBERrfimpute`. Simulations by Shah (Feb 13, 2014) suggested that the quality of the imputation for 10 and 100 trees was identical, so `mice 2.22` changed the default number of trees from `ntree = 100` to `ntree = 10`.

**Value**

Numeric vector of length `sum(!ry)` with imputations

**Author(s)**

Lisa Doove, Stef van Buuren, Elise Dusseldorp, 2012

**References**

Doove, L.L., van Buuren, S., Dusseldorp, E. (2014), Recursive partitioning for missing data imputation in the presence of interaction Effects. *Computational Statistics & Data Analysis*, 72, 92-104.

Shah, A.D., Bartlett, J.W., Carpenter, J., Nicholas, O., Hemingway, H. (2014), Comparison of random forest and parametric imputation models for imputing missing data using MICE: A CALIBER study. *American Journal of Epidemiology*, doi: 10.1093/aje/kwt312.

Van Buuren, S.(2012), *Flexible imputation of missing data*, Boca Raton, FL: Chapman & Hall/CRC.

**See Also**

[mice](#), [mice.impute.cart](#), [randomForest](#), [mice.impute.rfcat](#), [mice.impute.rfcont](#)



## Examples

```
library("lattice")

imp <- mice(nhanes2, meth = "rf", ntree = 3)
plot(imp)
```

---

mice.impute.ri

*Imputation by the random indicator method for nonignorable data*

---

## Description

Imputes univariate missing data using the random indicator method. This method estimates an offset between the distribution of the observed and missing data using an algorithm that iterates over the response model and the imputation model.

## Usage

```
mice.impute.ri(y, ry, x, ri.maxit = 10, ...)
```

## Arguments

y	Incomplete data vector of length n
ry	Vector of missing data pattern (FALSE=missing, TRUE=observed)
x	Matrix (n x p) of complete covariates.
ri.maxit	Number of inner iterations
...	Other named arguments passed down to .norm.draw()

## Value

A vector of length nmi s with imputations.

## Author(s)

Shahab Jolani (University of Utrecht) <s.jolani@uu.nl>

## References

Jolani, S. (2012). *Dual Imputation Strategies for Analyzing Incomplete Data*. Dissertation. University of Utrecht, Dec 7 2012. <http://dspace.library.uu.nl/handle/1874/257547>

---

mice.impute.sample     *Imputation by simple random sampling*

---

### Description

Imputes a random sample from the observed y data

### Usage

```
mice.impute.sample(y, ry, x = NULL, ...)
```

### Arguments

y	Incomplete data vector of length n
ry	Vector of missing data pattern (FALSE=missing, TRUE=observed)
x	Matrix (n x p) of complete covariates.
...	Other named arguments.

### Details

This function takes a simple random sample from the observed values in y, and returns these as imputations.

### Value

A vector of length nmi s with imputations.

### Author(s)

Stef van Buuren, Karin Groothuis-Oudshoorn, 2000

### References

van Buuren S and Groothuis-Oudshoorn K (2011). mice: Multivariate Imputation by Chained Equations in R. *Journal of Statistical Software*, **45**(3), 1-67. <http://www.jstatsoft.org/v45/i03/>

---

mice.mids

*Multivariate Imputation by Chained Equations (Iteration Step)*


---

**Description**

Takes a mids object, and produces a new object of class mids.

**Usage**

```
mice.mids(obj, maxit = 1, diagnostics = TRUE, printFlag = TRUE, ...)
```

**Arguments**

obj	An object of class mids, typically produced by a previous call to <code>mice()</code> or <code>mice.mids()</code>
maxit	The number of additional Gibbs sampling iterations.
diagnostics	A Boolean flag. If TRUE, diagnostic information will be appended to the value of the function. If FALSE, only the imputed data are saved. The default is TRUE.
printFlag	A Boolean flag. If TRUE, diagnostic information during the Gibbs sampling iterations will be written to the command window. The default is TRUE.
...	Named arguments that are passed down to the elementary imputation functions.

**Details**

This function enables the user to split up the computations of the Gibbs sampler into smaller parts. This is useful for the following reasons:

- RAM memory may become easily exhausted if the number of iterations is large. Returning to prompt/session level may alleviate these problems.
- The user can compute customized convergence statistics at specific points, e.g. after each iteration, for monitoring convergence. - For computing a 'few extra iterations'.

Note: The imputation model itself is specified in the `mice()` function and cannot be changed with `mice.mids`. The state of the random generator is saved with the mids object.

**Author(s)**

Stef van Buuren, Karin Groothuis-Oudshoorn, 2000

**References**

Van Buuren, S., Groothuis-Oudshoorn, K. (2011). `mice`: Multivariate Imputation by Chained Equations in R. *Journal of Statistical Software*, **45**(3), 1-67. <http://www.jstatsoft.org/v45/i03/>

**See Also**

[complete](#), [mice](#), [set.seed](#), [mids](#)

## Examples

```
imp1 <- mice(nhanes,maxit=1)
imp2 <- mice.mids(imp1)

# yields the same result as
imp <- mice(nhanes,maxit=2)

# for example:
#
# > imp$imp$bmi[1,]
#   1   2   3   4   5
# 1 30.1 35.3 33.2 35.3 27.5
# > imp2$imp$bmi[1,]
#   1   2   3   4   5
# 1 30.1 35.3 33.2 35.3 27.5
#
```

---

mice.theme

*Set the theme for the plotting Trellis functions*

---

## Description

The `mice.theme()` function sets default choices for Trellis plots that are built into **mice**.

## Usage

```
mice.theme(transparent = TRUE, alpha.fill = 0.3)
```

## Arguments

`transparent` A logical indicating whether alpha-transparency is allowed. The default is TRUE.  
`alpha.fill` A numerical values between 0 and 1 that indicates the default alpha value for fills.

## Value

`mice.theme()` returns a named list that can be used as a theme in the functions in **lattice**. By default, the `mice.theme()` function sets `transparent <- TRUE` if the current device `.Device` supports semi-transparent colors.

## Author(s)

Stef van Buuren 2011

---

mids-class	<i>Multiply imputed data set (mids)</i>
------------	---

---

## Description

The `mids` object contains a multiply imputed data set. The `mids` object is generated by the `mice()` and `mice.mids()` functions. The `mids` class of objects has methods for the following generic functions: `print`, `summary`, `plot`.

## Slots

`.Data`: Object of class "list" containing the following slots:

`call`: The call that created the object.

`data`: A copy of the incomplete data set.

`m`: The number of imputations.

`nmis`: An array containing the number of missing observations per column.

`imp`: A list of `ncol(data)` components with the generated multiple imputations. Each part of the list is a `nmis[j]` by `m` matrix of imputed values for variable `j`.

`method`: A vector of strings of length `ncol(data)` specifying the elementary imputation method per column.

`predictorMatrix`: A square matrix of size `ncol(data)` containing integers specifying the predictor set.

`visitSequence`: The sequence in which columns are visited.

`post`: A vector of strings of length `ncol(data)` with commands for post-processing

`seed`: The seed value of the solution.

`iteration`: Last Gibbs sampling iteration number.

`lastSeedValue`: The most recent seed value.

`chainMean`: A list of `m` components. Each component is a `length(visitSequence)` by `maxit` matrix containing the mean of the generated multiple imputations. The array can be used for monitoring convergence. Note that observed data are not present in this mean.

`chainVar`: A list with similar structure of `chainMean`, containing the covariances of the imputed values.

`loggedEvents`: A `data.frame` with six columns containing warnings, corrective actions, and other inside info.

`pad`: A list containing various settings of the padded imputation model, i.e. the imputation model after creating dummy variables. Normally, this list is only useful for error checking. List members are `pad$data` (data padded with columns for factors), `pad$predictorMatrix` (predictor matrix for the padded data), `pad$method` (imputation methods applied to the padded data), the vector `pad$visitSequence` (the visit sequence applied to the padded data), `pad$post` (post-processing commands for padded data) and `categories` (a matrix containing descriptive information about the padding operation).

**loggedEvents:** A matrix with six columns containing a record of automatic removal actions. It is NULL is no action was made. At initialization the program does the following three actions: 1. A variable that contains missing values, that is not imputed and that is used as a predictor is removed, 2. a constant variable is removed, and 3. a collinear variable is removed. During iteration, the program does the following actions: 1. one or more variables that are linearly dependent are removed (for categorical data, a 'variable' corresponds to a dummy variable), and 2. proportional odds regression imputation that does not converge and is replaced by polyreg. Column `it` is the iteration number at which the record was added, `im` is the imputation number, `co` is the column number in the data, `dep` is the name of the name of the dependent variable, `meth` is the imputation method used, and `out` is a (possibly long) character vector with the names of the altered or removed predictors.

### Note

Many of the functions of the `mice` package do not use the S4 class definitions, and instead rely on the S3 list equivalent `oldClass(obj) <- "mids"`.

### Author(s)

Stef van Buuren, Karin Groothuis-Oudshoorn, 2000

### References

van Buuren S and Groothuis-Oudshoorn K (2011). `mice`: Multivariate Imputation by Chained Equations in R. *Journal of Statistical Software*, **45**(3), 1-67. <http://www.jstatsoft.org/v45/i03/>

### See Also

[mice](#), [mira](#), [mipo](#)

---

mids2mplus

*Export mids object to Mplus*

---

### Description

Converts a `mids` object into a format recognized by Mplus, and writes the data and the Mplus input files

### Usage

```
mids2mplus(imp, file.prefix = "imp", path = getwd(), sep = "\t",
  dec = ".", silent = FALSE)
```

**Arguments**

<code>imp</code>	The <code>imp</code> argument is an object of class <code>mids</code> , typically produced by the <code>mice()</code> function.
<code>file.prefix</code>	A character string describing the prefix of the output data files.
<code>path</code>	A character string containing the path of the output file. By default, files are written to the current R working directory.
<code>sep</code>	The separator between the data fields.
<code>dec</code>	The decimal separator for numerical data.
<code>silent</code>	A logical flag stating whether the names of the files should be printed.

**Details**

This function automates most of the work needed to export a `mids` object to `Mplus`. The function writes the multiple imputation datasets, the file that contains the names of the multiple imputation data sets and an `Mplus` input file. The `Mplus` input file has the proper file names, so in principle it should run and read the data without alteration. `Mplus` will recognize the data set as a multiply imputed data set, and do automatic pooling in procedures where that is supported.

**Value**

The return value is `NULL`.

**Author(s)**

Gerko Vink, 2011.

**See Also**

[mids](#), [mids2spss](#)

---

mids2spss

*Export mids object to SPSS*

---

**Description**

Converts a `mids` object into a format recognized by SPSS, and writes the data and the SPSS syntax files.

**Usage**

```
mids2spss(imp, filedat = "midsdata.txt", filesps = "readmids.sps",  
path = getwd(), sep = "\t", dec = ".", silent = FALSE)
```

**Arguments**

<code>imp</code>	The <code>imp</code> argument is an object of class <code>mids</code> , typically produced by the <code>mice()</code> function.
<code>filedat</code>	A character string describing the name of the output data file.
<code>filesps</code>	A character string describing the name of the output syntax file.
<code>path</code>	A character string containing the path of the output file. The value in <code>path</code> is appended to <code>filedat</code> and <code>filesps</code> . By default, files are written to the current R working directory. If <code>path=NULL</code> then no file path appending is done.
<code>sep</code>	The separator between the data fields.
<code>dec</code>	The decimal separator for numerical data.
<code>silent</code>	A logical flag stating whether the names of the files should be printed.

**Details**

This function automates most of the work needed to export a `mids` object to SPSS. It uses a modified version of `writeForeignSPSS()` from the `foreign` package. The modified version allows for a choice of the field and decimal separators, and makes some improvements to the formatting, so that the generated syntax file is amenable to the `INCLUDE` statement in SPSS.

Below are some things to pay attention to.

The SPSS syntax file has the proper file names and separators set, so in principle it should run and read the data without alteration. SPSS is more strict than R with respect to the paths. Always use the full path, otherwise SPSS may not be able to find the data file.

Factors in R translate into categorical variables in SPSS. The internal coding of factor levels used in R is exported. This is generally acceptable for SPSS. However, when the data are to be combined with existing SPSS data, watch out for any changes in the factor levels codes. The `read.spss()` in package `foreign` for reading `.sav` uses its own internal numbering scheme 1, 2, 3, . . . for the levels of a factor. Consequently, changes in factor code can cause discrepancies in factor level when re-imported to SPSS. The solution is to manually recode the factor level in SPSS.

SPSS will recognize the data set as a multiply imputed data set, and do automatic pooling in procedures where that is supported. Note however that pooling is an extra option only available to those who licence the `MISSING VALUES` module. Without this licence, SPSS will still recognize the structure of the data, but not do any pooling.

**Value**

The return value is `NULL`.

**Author(s)**

Stef van Buuren, dec 2010.

**See Also**

[mids](#)



---

mipo-class	<i>Multiply imputed pooled analysis (mipo)</i>
------------	--

---

### Description

The mipo object is generated by the `pool` function from a `link[=mira-class]{mira}` object. The mipo class of objects has methods for the following generic functions: `print`, `summary`.

### Slots

`.Data`: Object of class "list" containing the following slots:

`call`: The call that created the mipo object.

`call1`: The call that created the mira object that was used in `call`.

`call2`: The call that created the mids object that was used in `call1`.

`data`: A copy of the incomplete data set.

`nmis`: An array containing the number of missing observations per column.

`m`: Number of multiple imputations.

`qhat`: An  $m$  by  $npar$  matrix containing the complete data estimates for the  $npar$  parameters of the  $m$  complete data analyses.

`u`: An  $m$  by  $npar$  by  $npar$  array containing the variance-covariance matrices of the estimates of the  $m$  complete data analyses.

`qbar`: The average of complete data estimates. The multiple imputation estimate.

`ubar`: The average of the variance-covariance matrix of the complete data estimates.

`b`: The between imputation variance-covariance matrix for the estimates.

`t`: The total variance-covariance matrix for the estimates.

`r`: Relative increases in variance due to missing data.

`dfcom`: Degrees of freedom in the hypothetically complete data: the sample size minus the number of free parameters.

`df`: Degrees of freedom associated with the t-statistics.

`fmi`: Fraction of missing information.

`lambda`: Proportion of the variation attributable to the missing data:  $(b+b/m)/t$ .

### Note

The functions of the `mice` package do not use the S4 class definitions, and instead rely on the S3 list equivalent `oldClass(obj) <- "mipo"`.

### Author(s)

Stef van Buuren, Karin Groothuis-Oudshoorn, 2000

## References

van Buuren S and Groothuis-Oudshoorn K (2011). mice: Multivariate Imputation by Chained Equations in R. *Journal of Statistical Software*, **45**(3), 1-67. <http://www.jstatsoft.org/v45/i03/>

## See Also

[pool](#), [mids](#), [mira](#)

---

mira-class

*Multiply imputed repeated analyses (mira)*

---

## Description

The mira object is generated by the `with.mids()` function. The `as.mira()` function takes the results of repeated complete-data analysis stored as a list, and turns it into a mira object that can be pooled. Pooling requires that `coef()` and `vcov()` methods are available for fitted object. The mira class of objects has methods for the following generic functions: `print`, `summary`.

## Slots

#'

Object of class "list" containing the following slots:

`.Data1`: The call that created the object.

`call1`: The call that created the mids object that was used in call.

`nmis`: An array containing the number of missing observations per column.

`analyses`: A list of `m` components containing the individual fit objects from each of the `m` complete data analyses.

## Note

Many of the functions of the mice package do not use the S4 class definitions, and instead rely on the S3 list equivalent `oldClass(obj) <- "mira"`.

## Author(s)

Stef van Buuren, Karin Groothuis-Oudshoorn, 2000

## References

van Buuren S and Groothuis-Oudshoorn K (2011). mice: Multivariate Imputation by Chained Equations in R. *Journal of Statistical Software*, **45**(3), 1-67. <http://www.jstatsoft.org/v45/i03/>

## See Also

[with.mids](#), [mids](#), [mipo](#)

---

ncc	<i>Number of complete cases</i>
-----	---------------------------------

---

**Description**

Calculates the number of complete cases.

**Usage**

```
ncc(x)
```

**Arguments**

x An R object. Currently supported are methods for the following classes: `mids`, `data.frame` and `matrix`. Also, x can be a vector.

**Value**

Number of elements in x with complete data.

**Author(s)**

Stef van Buuren, 2017

**See Also**

[nic](#), [cci](#)

**Examples**

```
ncc(nhanes) # 13 complete cases
```

---

nelsonaalen	<i>Cumulative hazard rate or Nelson-Aalen estimator</i>
-------------	---

---

**Description**

Calculates the cumulative hazard rate (Nelson-Aalen estimator)

**Usage**

```
nelsonaalen(data, timevar, statusvar)
```

**Arguments**

data	A data frame containing the data.
timevar	The name of the time variable in data.
statusvar	The name of the event variable, e.g. death in data.

**Details**

This function is useful for imputing variables that depend on survival time. White and Royston (2009) suggested using the cumulative hazard to the survival time  $H_0(T)$  rather than  $T$  or  $\log(T)$  as a predictor in imputation models. See section 7.1 of Van Buuren (2012) for an example.

**Value**

A vector with `nrow(data)` elements containing the Nelson-Aalen estimates of the cumulative hazard function.

**Author(s)**

Stef van Buuren, 2012

**References**

- White, I. R., Royston, P. (2009). Imputing missing covariate values for the Cox model. *Statistics in Medicine*, 28(15), 1982-1998.
- van Buuren, S. (2012). *Flexible Imputation of Missing Data*. Boca Raton, FL: Chapman & Hall/CRC Press.

**Examples**

```
require(MASS)

leuk$status <- 1 ## no censoring occurs in leuk data (MASS)
ch <- nelsonaalen(leuk, time, status)
plot(x = leuk$time, y = ch, ylab='Cumulative hazard', xlab='Time')

### See example on http://www.engineeredsoftware.com/lmar/pe\_cum\_hazard\_function.htm
time <- c(43, 67, 92, 94, 149, rep(149,7))
status <- c(rep(1,5),rep(0,7))
eng <- data.frame(time, status)
ch <- nelsonaalen(eng, time, status)
plot(x = time, y = ch, ylab='Cumulative hazard', xlab='Time')
```

---

nhanes

*NHANES example - all variables numerical*

---

### Description

A small data set with non-monotone missing values.

### Format

A data frame with 25 observations on the following 4 variables.

**age** Age group (1=20-39, 2=40-59, 3=60+)

**bmi** Body mass index (kg/m\*\*2)

**hyp** Hypertensive (1=no,2=yes)

**chl** Total serum cholesterol (mg/dL)

### Details

A small data set with all numerical variables. The data set `nhanes2` is the same data set, but with `age` and `hyp` treated as factors.

### Source

Schafer, J.L. (1997). *Analysis of Incomplete Multivariate Data*. London: Chapman & Hall. Table 6.14.

### See Also

[nhanes2](#)

### Examples

```
imp <- mice(nhanes) # create 5 imputed data sets
complete(imp)      # print the first imputed data set
```

---

nhanes2

*NHANES example - mixed numerical and discrete variables*

---

### Description

A small data set with non-monotone missing values.

### Format

A data frame with 25 observations on the following 4 variables.

**age** Age group (1=20-39, 2=40-59, 3=60+)

**bmi** Body mass index (kg/m\*\*2)

**hyp** Hypertensive (1=no,2=yes)

**chl** Total serum cholesterol (mg/dL)

### Details

A small data set with missing data and mixed numerical and discrete variables. The data set `nhanes` is the same data set, but with all data treated as numerical.

### Source

Schafer, J.L. (1997). *Analysis of Incomplete Multivariate Data*. London: Chapman & Hall. Table 6.14.

### See Also

[nhanes](#)

### Examples

```
imp <- mice(nhanes2)      # create 5 imputed data sets
complete(imp)            # print the first imputed data set
```

---

nic	<i>Number of incomplete cases</i>
-----	-----------------------------------

---

**Description**

Calculates the number of incomplete cases.

**Usage**

```
nic(x)
```

**Arguments**

x                    An R object. Currently supported are methods for the following classes: `mids`, `data.frame` and `matrix`. Also, x can be a vector.

**Value**

Number of elements in x with incomplete data.

**Author(s)**

Stef van Buuren, 2017

**See Also**

[ncc](#), [cci](#)

**Examples**

```
nic(nhanes) # the remaining 12 rows
nic(nhanes[,c("bmi","hyp")]) # number of cases with incomplete bmi and hyp
```

---

norm.draw	<i>Draws values of beta and sigma by Bayesian linear regression</i>
-----------	---

---

**Description**

This function draws random values of beta and sigma under the Bayesian linear regression model as described in Rubin (1987, p. 167). This function can be called by user-specified imputation functions.

**Usage**

```
norm.draw(y, ry, x, ridge = 1e-05, ...)
```

```
.norm.draw(y, ry, x, ridge = 1e-05, ...)
```

**Arguments**

y	Incomplete data vector of length n
ry	Vector of missing data pattern (FALSE=missing, TRUE=observed)
x	Matrix (n x p) of complete covariates.
ridge	A small numerical value specifying the size of the ridge used. The default value <code>ridge = 1e-05</code> represents a compromise between stability and unbiasedness. Decrease ridge if the data contain many junk variables. Increase ridge for highly collinear data.
...	Other named arguments.

**Value**

A list containing components `coef` (least squares estimate), `beta` (drawn regression weights) and `sigma` (drawn value of the residual standard deviation).

**Author(s)**

Stef van Buuren, Karin Groothuis-Oudshoorn, 2000

**References**

Rubin, D.B. (1987). *Multiple imputation for nonresponse in surveys*. New York: Wiley.

---

pattern

*Datasets with various missing data patterns*

---

**Description**

Four simple datasets with various missing data patterns

**Format**

**list("pattern1")** Data with a univariate missing data pattern  
**list("pattern2")** Data with a monotone missing data pattern  
**list("pattern3")** Data with a file matching missing data pattern  
**list("pattern4")** Data with a general missing data pattern

**Details**

Van Buuren (2012) uses these four artificial datasets to illustrate various missing data patterns.



**Source**

van Buuren, S. (2012). *Flexible Imputation of Missing Data*. Boca Raton, FL: Chapman & Hall/CRC Press.

**Examples**

```
require(lattice)
require(MASS)

pattern4

data <- rbind(pattern1, pattern2, pattern3, pattern4)
mdpat <- cbind(expand.grid(rec = 8:1, pat = 1:4, var = 1:3), r=as.numeric(as.vector(is.na(data))))

types <- c("Univariate", "Monotone", "File matching", "General")
tp41 <- levelplot(r~var+rec|as.factor(pat), data=mdpat,
  as.table=TRUE, aspect="iso",
  shrink=c(0.9),
  col.regions = mdc(1:2),
  colorkey=FALSE,
  scales=list(draw=FALSE),
  xlab="", ylab="",
  between = list(x=1,y=0),
  strip = strip.custom(bg = "grey95", style = 1,
    factor.levels = types))
print(tp41)

md.pattern(pattern4)
p <- md.pairs(pattern4)
p

### proportion of usable cases
p$mr/(p$mr+p$mm)

### outbound statistics
p$rm/(p$rm+p$rr)

fluxplot(pattern2)
```

**Description**

Trace line plots, also called stream plots or history plots, portray the value of an estimate against the iteration number. The estimate can be anything that you can calculate, but typically are chosen

as parameter of scientific interest. The `plot` method for a `mids` object plots the mean and standard deviation of the imputed (not observed) values against the iteration number for each of the  $m$  replications. By default, the function plots the development of the mean and standard deviation for each incomplete variable. On convergence, the streams should intermingle and be free of any trend.

### Usage

```
## S3 method for class 'mids'
plot(x, y = NULL, theme = mice.theme(), layout = c(2, 3),
     type = "l", col = 1:10, lty = 1, ...)
```

### Arguments

<code>x</code>	An object of class <code>mids</code>
<code>y</code>	A formula that specifies which variables, stream and iterations are plotted. If omitted, all streams, variables and iterations are plotted.
<code>theme</code>	The trellis theme to applied to the graphs. The default is <code>mice.theme()</code> .
<code>layout</code>	A vector of length 2 given the number of columns and rows in the plot. The default is <code>c(2, 3)</code> .
<code>type</code>	Parameter <code>type</code> of <a href="#">panel.xyplot</a> .
<code>col</code>	Parameter <code>col</code> of <a href="#">panel.xyplot</a> .
<code>lty</code>	Parameter <code>lty</code> of <a href="#">panel.xyplot</a> .
<code>...</code>	Extra arguments for <a href="#">xyplot</a> .

### Value

An object of class `"trellis"`.

### Author(s)

Stef van Buuren 2011

### See Also

[mice](#), [mids](#), [xyplot](#)

---

<code>pool</code>	<i>Multiple imputation pooling</i>
-------------------	------------------------------------

---

### Description

Pools the results of  $m$  repeated complete data analysis

### Usage

```
pool(object, method = "smallsample")
```

**Arguments**

object	An object of class <code>mira</code> , produced by <code>with.mids()</code> or <code>as.mira()</code>
method	A string describing the method to compute the degrees of freedom. The default value is "smallsample", which specifies the is Barnard-Rubin adjusted degrees of freedom (Barnard and Rubin, 1999) for small samples. Specifying a different string produces the conventional degrees of freedom as in Rubin (1987).

**Details**

The function averages the estimates of the complete data model, computes the total variance over the repeated analyses, and computes the relative increase in variance due to nonresponse and the fraction of missing information. The function relies on the availability of

1. the estimates of the model, typically present as 'coefficients' in the fit object
2. an appropriate estimate of the variance-covariance matrix of the estimates per analyses (estimated by `vcov`).

The function pools also estimates obtained with `lme()` and `lmer()`, BUT only the fixed part of the model.

**Value**

An object of class `mipo`, which stands for 'multiple imputation pooled outcome'.

**Author(s)**

Stef van Buuren, Karin Groothuis-Oudshoorn, 2009

**References**

- Barnard, J. and Rubin, D.B. (1999). Small sample degrees of freedom with multiple imputation. *Biometrika*, 86, 948-955.
- Rubin, D.B. (1987). *Multiple Imputation for Nonresponse in Surveys*. New York: John Wiley and Sons.
- van Buuren S and Groothuis-Oudshoorn K (2011). mice: Multivariate Imputation by Chained Equations in R. *Journal of Statistical Software*, 45(3), 1-67. <http://www.jstatsoft.org/v45/i03/>
- Pinheiro, J.C. and Bates, D.M. (2000). *Mixed-Effects Models in S and S-PLUS*. Berlin: Springer.

**See Also**

[with.mids](#), [as.mira](#), [vcov](#)

**Examples**

```

# which vcov methods can R find
methods(vcov)

#
imp <- mice(nhanes)
fit <- with(data=imp,exp=lm(bmi~hyp+chl))
pool(fit)

#Call: pool(object = fit)
#
#Pooled coefficients:
#(Intercept)      hyp      chl
# 22.01313    -1.45578    0.03459
#
#Fraction of information about the coefficients missing due to nonresponse:
#(Intercept)      hyp      chl
#  0.29571      0.05639    0.38759
#> summary(pool(fit))
#           est      se      t      df Pr(>|t|)   lo 95   hi 95 missing
#(Intercept) 22.01313 4.94086  4.4553 12.016 0.000783 11.24954 32.77673      NA
#hyp         -1.45578 2.26789 -0.6419 20.613 0.528006 -6.17752  3.26596      8
#chl          0.03459 0.02829  1.2228  9.347 0.251332 -0.02904  0.09822     10
#           fmi
#(Intercept) 0.29571
#hyp         0.05639
#chl         0.38759
#

```

---

pool.compare

*Compare two nested models fitted to imputed data*

---

**Description**

Compares two nested models after  $m$  repeated complete data analysis

**Usage**

```
pool.compare(fit1, fit0, data = NULL, method = "Wald")
```

**Arguments**

`fit1` An object of class 'mira', produced by `with.mids()`.

`fit0` An object of class 'mira', produced by `with.mids()`. The model in `fit0` should be a submodel of `fit1`. Moreover, the variables of the submodel should be the first variables of the larger model and in the same order as in the submodel.

data	In case of method 'likelihood' it is necessary to pass also the original mids object to the data argument. Default value is NULL, in case of method='Wald'.
method	A string describing the method to compare the two models. Two kind of comparisons are included so far: 'Wald' and 'likelihood'.

### Details

The function is based on the article of Meng and Rubin (1992). The Wald-method can be found in paragraph 2.2 and the likelihood method can be found in paragraph 3. One could use the Wald method for comparison of linear models obtained with e.g. `lm` (in `with.mids()`). The likelihood method should be used in case of logistic regression models obtained with `glm` in `with.mids()`. It is assumed that `fit1` contains the larger model and the model in `fit0` is fully contained in `fit1`. In case of `method='Wald'`, the null hypothesis is tested that the extra parameters are all zero.

### Value

A list containing several components. Component `call` is that call to the `pool.compare` function. Component `call11` is the call that created `fit1`. Component `call12` is the call that created the imputations. Component `call01` is the call that created `fit0`. Component `call02` is the call that created the imputations. Component `method` is the method used to compare two models: 'Wald' or 'likelihood'. Component `nmis` is the number of missing entries for each variable. Component `m` is the number of imputations. Component `qhat1` is a matrix, containing the estimated coefficients of the  $m$  repeated complete data analyses from `fit1`. Component `qhat0` is a matrix, containing the estimated coefficients of the  $m$  repeated complete data analyses from `fit0`. Component `ubar1` is the mean of the variances of `fit1`, formula (3.1.3), Rubin (1987). Component `ubar0` is the mean of the variances of `fit0`, formula (3.1.3), Rubin (1987). Component `qbar1` is the pooled estimate of `fit1`, formula (3.1.2) Rubin (1987). Component `qbar0` is the pooled estimate of `fit0`, formula (3.1.2) Rubin (1987). Component `Dm` is the test statistic. Component `rm` is the relative increase in variance due to nonresponse, formula (3.1.7), Rubin (1987). Component `df1`:  $df1 =$  under the null hypothesis it is assumed that  $D_m$  has an F distribution with  $(df1, df2)$  degrees of freedom. Component `df2`:  $df2$ . Component `pvalue` is the P-value of testing whether the larger model is statistically different from the smaller submodel.

### Author(s)

Karin Groothuis-Oudshoorn and Stef van Buuren, 2009

### References

- Li, K.H., Meng, X.L., Raghunathan, T.E. and Rubin, D. B. (1991). Significance levels from repeated p-values with multiply-imputed data. *Statistica Sinica*, 1, 65-92.
- Meng, X.L. and Rubin, D.B. (1992). Performing likelihood ratio tests with multiple-imputed data sets. *Biometrika*, 79, 103-111.
- van Buuren S and Groothuis-Oudshoorn K (2011). `mice`: Multivariate Imputation by Chained Equations in R. *Journal of Statistical Software*, 45(3), 1-67. <http://www.jstatsoft.org/v45/i03/>

**See Also**

[lm.mids](#), [glm.mids](#), [vcov](#),

**Examples**

```
### To compare two linear models:
imp <- mice(nhanes2)
mi1 <- with(data=imp, expr=lm(bmi~age+hyp+chl))
mi0 <- with(data=imp, expr=lm(bmi~age+hyp))
pc <- pool.compare(mi1, mi0, method='Wald')
pc$spvalue
#           [,1]
#[1,] 0.000293631
#

### Comparison of two general linear models (logistic regression).
## Not run:
imp <- mice(boys, maxit=2)
fit0 <- with(imp, glm(gen>levels(gen)[1] ~ hgt+hc,family=binomial))
fit1 <- with(imp, glm(gen>levels(gen)[1] ~ hgt+hc+reg,family=binomial))
pool.compare(fit1, fit0, method='likelihood', data=imp)
## End(Not run)
```

---

pool.r.squared

*Pooling: R squared*

---

**Description**

Pools  $R^2$  of  $m$  repeated complete data models.

**Usage**

```
pool.r.squared(object, adjusted = FALSE)
```

**Arguments**

object	An object of class 'mira', produced by <code>lm.mids</code> or <code>with.mids</code> with <code>lm</code> as modelling function.
adjusted	A logical value. If <code>adjusted=TRUE</code> then the adjusted $R^2$ is calculated. The default value is <code>FALSE</code> .

**Details**

The function pools the coefficients of determination  $R^2$  or the adjusted coefficients of determination ( $R^2_a$ ) obtained with the `lm` modelling function. For pooling it uses the Fisher  $z$ -transformation.

**Value**

Returns a 1x4 table with components. Component `est` is the pooled  $R^2$  estimate. Component `lo95` is the 95 % lower bound of the pooled  $R^2$ . Component `hi95` is the 95 % upper bound of the pooled  $R^2$ . Component `fmi` is the fraction of missing information due to nonresponse.

**Author(s)**

Karin Groothuis-Oudshoorn and Stef van Buuren, 2009

**References**

Harel, O (2009). The estimation of  $R^2$  and adjusted  $R^2$  in incomplete data sets using multiple imputation, *Journal of Applied Statistics*, 36:1109-1118.

Rubin, D.B. (1987). *Multiple Imputation for Nonresponse in Surveys*. New York: John Wiley and Sons.

van Buuren S and Groothuis-Oudshoorn K (2011). `mice`: Multivariate Imputation by Chained Equations in R. *Journal of Statistical Software*, **45**(3), 1-67. <http://www.jstatsoft.org/v45/i03/>

**See Also**

[pool, pool.scalar](#)

**Examples**

```
imp<-mice(nhanes)

fit<-lm.mids(chl~age+hyp+bmi,imp)
pool.r.squared(fit)
pool.r.squared(fit,adjusted=TRUE)

#fit<-lm.mids(chl~age+hyp+bmi,imp)
#
#> pool.r.squared(fit)
#      est      lo 95      hi 95      fmi
#R^2 0.5108041 0.1479687 0.7791927 0.3024413
#
#> pool.r.squared(fit,adjusted=TRUE)
#      est      lo 95      hi 95      fmi
#adj R^2 0.4398066 0.08251427 0.743172 0.3404165
#
```

---

 pool.scalar

*Multiple imputation pooling: univariate version*


---

**Description**

Pools univariate estimates of  $m$  repeated complete data analysis

**Usage**

```
pool.scalar(Q, U, n = 99999, k = 1, method = "smallsample")
```

**Arguments**

Q	A vector of univariate estimates of $m$ repeated complete data analyses.
U	A vector containing the corresponding $m$ variances of the univariate estimates.
n	A number providing the sample size. If nothing is specified, a large sample $n = 99999$ is assumed.
k	A number indicating the number of parameters to be estimated. By default, $k = 1$ is assumed.
method	A string indicating the method to calculate the degrees of freedom. If <code>method = "smallsample"</code> (the default) then the Barnard-Rubin adjustment for small degrees of freedom is used. Otherwise, the method from Rubin (1987) is used.

**Details**

The function averages the univariate estimates of the complete data model, computes the total variance over the repeated analyses, and computes the relative increase in variance due to nonresponse and the fraction of missing information.

**Value**

Returns a list with components. Component `m` is the number of imputations. Component `qhat` contains the  $m$  univariate estimates of repeated complete data analyses. Component `u` contains the corresponding  $m$  variances of the univariate estimates. Component `qbar` is the pooled univariate estimate, formula (3.1.2) Rubin (1987). Component `ubar` is the mean of the variances (i.e. the pooled within-imputation variance), formula (3.1.3) Rubin (1987). Component `b` is the between-imputation variance, formula (3.1.4) Rubin (1987). Component `t` is the total variance of the pooled estimated, formula (3.1.5) Rubin (1987). Component `r` is the relative increase in variance due to nonresponse, formula (3.1.7) Rubin (1987). Component `df` is the degrees of freedom for `t` reference distribution, formula (3.1.6) Rubin (1987) or method of Barnard-Rubin (1999) (if `method = "smallsample"`). Component `fmi` is the fraction missing information due to nonresponse, formula (3.1.10) Rubin (1987). Component `lambda` is the proportion of variation due to nonresponse, formula (2.24) Van Buuren (2012).

**Author(s)**

Karin Groothuis-Oudshoorn and Stef van Buuren, 2009



**References**

Rubin, D.B. (1987). *Multiple Imputation for Nonresponse in Surveys*. New York: John Wiley and Sons.

**See Also**

[pool](#)

**Examples**

```
imp <- mice(nhanes)
m <- imp$m
Q <- rep(NA, m)
U <- rep(NA, m)
for (i in 1:m) {
  Q[i] <- mean(complete(imp, i)$bmi)
  U[i] <- var(complete(imp, i)$bmi) / nrow(nhanes) # (standard error of estimate)^2
}
pool.scalar(Q, U, method = "rubin") # Rubin 1987
pool.scalar(Q, U, n = nrow(nhanes), k = 1) # Barnard-Rubin 1999
```

---

popmis

*Hox pupil popularity data with missing popularity scores*

---

**Description**

Hox pupil popularity data with some missing popularity scores

**Format**

A data frame with 2000 rows and 7 columns:

**pupil** Pupil number within school  
**school** School number  
**popular** Pupil popularity with 848 missing entries  
**sex** Pupil gender  
**texp** Teacher experience (years)  
**const** Constant intercept term  
**teachpop** Teacher popularity

**Details**

The original, complete dataset was generated by Joop Hox as an example of well-behaved multilevel data set. The distributed data contains missing data in pupil popularity.

**Source**

Hox, J. J. (2002) *Multilevel analysis. Techniques and applications*. Mahwah, NJ: Lawrence Erlbaum.

**Examples**

```
popmis[1:3,]
```

---

pops

*Project on preterm and small for gestational age infants (POPS)*

---

**Description**

Subset of data from the POPS study, a national, prospective study on preterm children, including all liveborn infants <32 weeks gestational age and/or <1500 g from 1983 (n = 1338).

**Format**

pops is a data frame with 959 rows and 86 columns. pops.pred is the 86 by 86 binary predictor matrix used for specifying the multiple imputation model.

**Details**

The data set concerns of subset of 959 children that survived up to the age of 19 years.

Hille et al (2005) divided the 959 survivors into three groups: Full responders (examined at an outpatient clinic and completed the questionnaires, n = 596), postal responders (only completed the mailed questionnaires, n = 109), non-responders (did not respond to any of the mailed requests or telephone calls, or could not be traced, n = 254).

Compared to the postal and non-responders, the full response group consists of more girls, contains more Dutch children, has higher educational and social economic levels and has fewer handicaps. The responders form a highly selective subgroup in the total cohort.

Multiple imputation of this data set has been described in Hille et al (2007) and Van Buuren (2012), chapter 8.

**Source**

Hille, E. T. M., Elbertse, L., Bennebroek Gravenhorst, J., Brand, R., Verloove-Vanhorick, S. P. (2005). Nonresponse bias in a follow-up study of 19-year-old adolescents born as preterm infants. *Pediatrics*, 116(5):662666.

Hille, E. T. M., Weisglas-Kuperus, N., Van Goudoever, J. B., Jacobusse, G. W., Ens-Dokkum, M. H., De Groot, L., Wit, J. M., Geven, W. B., Kok, J. H., De Kleine, M. J. K., Kollee, L. A. A., Mulder, A. L. M., Van Straaten, H. L. M., De Vries, L. S., Van Weissenbruch, M. M., Verloove-Vanhorick, S. P. (2007). Functional outcomes and participation in young adulthood for very preterm and very

low birth weight infants: The Dutch project on preterm and small for gestational age infants at 19 years of age. *Pediatrics*, 120(3):587595.

van Buuren, S. (2012). *Flexible Imputation of Missing Data*. Boca Raton, FL: Chapman & Hall/CRC Press.

## Examples

```
pops <- data(pops)
```

---

potthoffroy

*Potthoff-Roy data*

---

## Description

Data from Potthoff-Roy (1964) with repeated measures on dental fissures.

## Format

tbs is a data frame with 27 rows and 6 columns:

**id** Person number

**sex** Sex M/F

**d8** Distance at age 8 years

**d10** Distance at age 10 years

**d12** Distance at age 12 years

**d14** Distance at age 14 years

## Details

This data set is the famous Potthoff-Roy data, used to demonstrate MANOVA on repeated measure data. Potthoff and Roy (1964) published classic data on a study in 16 boys and 11 girls, who at ages 8, 10, 12, and 14 had the distance (mm) from the center of the pituitary gland to the pteryomaxillary fissure measured. Changes in pituitary-ptyeryomaxillary distances during growth is important in orthodontic therapy. The goals of the study were to describe the distance in boys and girls as simple functions of age, and then to compare the functions for boys and girls. The data have been reanalyzed by many authors including Jennrich and Schluchter (1986), Little and Rubin (1987), Pinheiro and Bates (2000), Verbeke and Molenberghs (2000) and Molenberghs and Kenward (2007). See Chapter 9 of Van Buuren (2012) for a challenging exercise using these data.

**Source**

Potthoff, R. F., Roy, S. N. (1964). A generalized multivariate analysis of variance model usefully especially for growth curve problems. *Biometrika*, 51(3), 313-326.

Little, R. J. A., Rubin, D. B. (1987). *Statistical Analysis with Missing Data*. New York: John Wiley & Sons.

Van Buuren, S. (2012). *Flexible Imputation of Missing Data*. Boca Raton, FL: Chapman & Hall/CRC Press.

**Examples**

```
### create missing values at age 10 as in Little and Rubin (1987)

phr <- potthoffroy
idmis <- c(3,6,9,10,13,16,23,24,27)
phr[idmis, 4] <- NA
phr

md.pattern(phr)
```

---

print.mads

*Print a mads object*


---

**Description**

Print a mads object

**Usage**

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

**Arguments**

x	Object of class mads
...	Other parameters passed down to print.default()

**Value**

NULL

**See Also**

[mads](#)

---

print.mids	<i>Print a mids object</i>
------------	----------------------------

---

**Description**

Print a mids object

Print a mira object

Print a mipo object

**Usage**

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

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

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

**Arguments**

x	Object of class mids, mira or mipo
...	Other parameters passed down to print.default()

**Value**

NULL

NULL

NULL

**See Also**

[mids](#)

[mira](#)

[mipo](#)

---

 quickpred

*Quick selection of predictors from the data*


---

### Description

Selects predictors according to simple statistics

### Usage

```
quickpred(data, mincor = 0.1, minpuc = 0, include = "", exclude = "",
  method = "pearson")
```

### Arguments

data	Matrix or data frame with incomplete data.
mincor	A scalar, numeric vector (of size <code>ncol(data)</code> ) or numeric matrix (square, of size <code>ncol(data)</code> ) specifying the minimum threshold(s) against which the absolute correlation in the data is compared.
minpuc	A scalar, vector (of size <code>ncol(data)</code> ) or matrix (square, of size <code>ncol(data)</code> ) specifying the minimum threshold(s) for the proportion of usable cases.
include	A string or a vector of strings containing one or more variable names from <code>names(data)</code> . Variables specified are always included as a predictor.
exclude	A string or a vector of strings containing one or more variable names from <code>names(data)</code> . Variables specified are always excluded as a predictor.
method	A string specifying the type of correlation. Use 'pearson' (default), 'kendall' or 'spearman'. Can be abbreviated.

### Details

This function creates a predictor matrix using the variable selection procedure described in Van Buuren et al. (1999, p. 687–688). The function is designed to aid in setting up a good imputation model for data with many variables.

Basic workings: The procedure calculates for each variable pair (i.e. target-predictor pair) two correlations using all available cases per pair. The first correlation uses the values of the target and the predictor directly. The second correlation uses the (binary) response indicator of the target and the values of the predictor. If the largest (in absolute value) of these correlations exceeds `mincor`, the predictor will be added to the imputation set. The default value for `mincor` is 0.1.

In addition, the procedure eliminates predictors whose proportion of usable cases fails to meet the minimum specified by `minpuc`. The default value is 0, so predictors are retained even if they have no usable case.

Finally, the procedure includes any predictors named in the `include` argument (which is useful for background variables like age and sex) and eliminates any predictor named in the `exclude` argument. If a variable is listed in both `include` and `exclude` arguments, the `include` argument takes precedence.

Advanced topic: mincor and minpuc are typically specified as scalars, but vectors and squares matrices of appropriate size will also work. Each element of the vector corresponds to a row of the predictor matrix, so the procedure can effectively differentiate between different target variables. Setting a high values for can be useful for auxiliary, less important, variables. The set of predictor for those variables can remain relatively small. Using a square matrix extends the idea to the columns, so that one can also apply cellwise thresholds.

### Value

A square binary matrix of size `ncol(data)`.

### Author(s)

Stef van Buuren, Aug 2009

### References

van Buuren, S., Boshuizen, H.C., Knook, D.L. (1999) Multiple imputation of missing blood pressure covariates in survival analysis. *Statistics in Medicine*, **18**, 681–694.

van Buuren, S. and Groothuis-Oudshoorn, K. (2011). mice: Multivariate Imputation by Chained Equations in R. *Journal of Statistical Software*, **45**(3), 1-67. <http://www.jstatsoft.org/v45/i03/>

### See Also

[mice](#), [mids](#)

### Examples

```
# default: include all predictors with absolute correlation over 0.1
quickpred(nhanes)

# all predictors with absolute correlation over 0.4
quickpred(nhanes, mincor=0.4)

# include age and bmi, exclude chl
quickpred(nhanes, mincor=0.4, inc=c('age','bmi'), exc='chl')

# only include predictors with at least 30% usable cases
quickpred(nhanes, minpuc=0.3)

# use low threshold for bmi, and high thresholds for hyp and chl
pred <- quickpred(nhanes, mincor=c(0,0.1,0.5,0.5))
pred

# use it directly from mice
imp <- mice(nhanes, pred=quickpred(nhanes, minpuc=0.25, include='age'))
```

---

<code>rbind.mids</code>	<i>Rowwise combination of a mids object.</i>
-------------------------	--

---

**Description**

Append mids objects by rows

**Usage**

```
rbind.mids(x, y, ...)
```

**Arguments**

<code>x</code>	A mids object.
<code>y</code>	A mids object or a <code>data.frame</code> , <code>matrix</code> , <code>factor</code> or <code>vector</code> .
<code>...</code>	Additional <code>data.frame</code> , <code>matrix</code> , <code>vector</code> or <code>factor</code> . These can be given as named arguments.

**Details**

This function combines two mids objects rowwise into a single mids object or combines a mids object and a vector, matrix, factor or dataframe rowwise into a mids object. The number of columns in the (incomplete) data `x$data` and `y` (or `y$data` if `y` is a mids object) should be equal. If `y` is a mids object then the number of imputations in `x` and `y` should be equal.

**Value**

An S3 object of class `mids`

**Note**

Component `call` is a vector, with first argument the `mice()` statement that created `x` and second argument the call to `rbind.mids()`. Component `data` is the rowwise combination of the (incomplete) data in `x` and `y`. Component `m` is equal to `x$m`. Component `nmis` is an array containing the number of missing observations per column, defined as `x$nmis + y$nmis`. Component `imp` is a list of `nvar` components with the generated multiple imputations. Each part of the list is a `nmis[j]` by `m` matrix of imputed values for variable `j`. If `y` is a mids object then `imp[[j]]` equals `rbind(x$imp[[j]], y$imp[[j]])`; otherwise the original data of `y` will be copied into this list, including the missing values of `y` then `y` is not imputed. Component `method` is a vector of strings of length(`nvar`) specifying the elementary imputation method per column defined as `x$method`. Component `predictorMatrix` is a square matrix of size `ncol(data)` containing the predictor set defined as `x$predictorMatrix`. Component `visitSequence` is the sequence in which columns are visited, defined as `x$visitSequence`. Component `seed` is the seed value of the solution, `x$seed`. Component `iteration` is the last Gibbs sampling iteration number, `x$iteration`. Component `lastSeedValue` is the most recent seed value, `x$lastSeedValue`. Component `chainMean` is set to `NA`. Component `chainVar` is set to `NA`. Component `pad` is set to `x$pad`, a list containing various settings of the padded imputation model, i.e. the imputation model after creating dummy variables. Component `loggedEvents` is set to `x$loggedEvents`.



**Author(s)**

Karin Groothuis-Oudshoorn, Stef van Buuren, 2009

**References**

van Buuren S and Groothuis-Oudshoorn K (2011). mice: Multivariate Imputation by Chained Equations in R. *Journal of Statistical Software*, **45**(3), 1-67. <http://www.jstatsoft.org/v45/i03/>

**See Also**

[cbind.mids](#), [ibind](#), [mids](#)

---

selfreport

*Self-reported and measured BMI*

---

**Description**

Dataset containing height and weight data (measured, self-reported) from two studies.

**Format**

A data frame with 2060 rows and 15 variables:

**src** Study, either kru1 or mgg (factor)  
**id** Person identification number  
**pop** Population, all NL (factor)  
**age** Age of respondent in years  
**sex** Sex of respondent (factor)  
**hm** Height measured (cm)  
**wm** Weight measured (kg)  
**hr** Height reported (cm)  
**wr** Weight reported (kg)  
**prg** Pregnancy (factor), all Not pregnant  
**edu** Educational level (factor)  
**etn** Ethnicity (factor)  
**web** Obtained through web survey (factor)  
**bm** BMI measured (kg/m<sup>2</sup>)  
**br** BMI reported (kg/m<sup>2</sup>)

## Details

This dataset combines two datasets: *krul* data (Krul, 2010) (1257 persons) and the *mgg* data (Van Keulen 2011; Van der Klauw 2011) (803 persons). The *krul* dataset contains height and weight (both measures and self-reported) from 1257 Dutch adults, whereas the *mgg* dataset contains self-reported height and weight for 803 Dutch adults. Section 7.3 in Van Buuren (2012) shows how the missing measured data can be imputed in the *mgg* data, so corrected prevalence estimates can be calculated.

## Source

Krul, A., Daanen, H. A. M., Choi, H. (2010). Self-reported and measured weight, height and body mass index (BMI) in Italy, The Netherlands and North America. *European Journal of Public Health*, 21(4), 414-419.

Van Keulen, H.M., Chorus, A.M.J., Verheijden, M.W. (2011). *Monitor Convenant Gezond Gewicht Nulmeting (determinanten van) beweeg- en eetgedrag van kinderen (4-11 jaar), jongeren (12-17 jaar) en volwassenen (18+ jaar)*. TNO/LS 2011.016. Leiden: TNO.

Van der Klauw, M., Van Keulen, H.M., Verheijden, M.W. (2011). *Monitor Convenant Gezond Gewicht Beweeg- en eetgedrag van kinderen (4-11 jaar), jongeren (12-17 jaar) en volwassenen (18+ jaar) in 2010 en 2011*. TNO/LS 2011.055. Leiden: TNO. (in Dutch)

van Buuren, S. (2012). *Flexible Imputation of Missing Data*. Boca Raton, FL: Chapman & Hall/CRC Press.

## Examples

```
md.pattern(selfreport[,c("age", "sex", "hm", "hr", "wm", "wr")])

### FIMD Section 7.3.5 Application

bmi <- function(h,w){return(w/(h/100)^2)}
init <- mice(selfreport,maxit=0)
meth <- init$meth
meth["bm"] <- "~bmi(hm,wm)"
pred <- init$pred
pred[,c("src","id","web","bm","br")] <- 0
imp <- mice(selfreport, pred=pred, meth=meth, seed=66573, maxit=2, m=1)
## imp <- mice(selfreport, pred=pred, meth=meth, seed=66573, maxit=20, m=10)

### Like FIMD Figure 7.6

cd <- complete(imp, 1)
xy <- xy.coords(cd$bm, cd$br-cd$bm)
plot(xy,col=mdc(2),xlab="Measured BMI",ylab="Reported - Measured BMI",
      xlim=c(17,45),ylim=c(-5,5), type="n",lwd=0.7)
polygon(x=c(30,20,30),y=c(0,10,10),col="grey95",border=NA)
polygon(x=c(30,40,30),y=c(0,-10,-10),col="grey95",border=NA)
abline(0,0,lty=2,lwd=0.7)

idx <- cd$src=="krul"
```

```
xyx <- xy; xyc$x <- xy$x[idx]; xyc$y <- xy$y[idx]
xys <- xy; xys$x <- xy$x[!idx]; xys$y <- xy$y[!idx]
points(xyc,col=mdc(1), cex=0.7)
points(xys,col=mdc(2), cex=0.7)
lines(lowess(xyc),col=mdc(4),lwd=2)
lines(lowess(xys),col=mdc(5),lwd=2)
text(1:4,x=c(40,28,20,32),y=c(4,4,-4,-4),cex=3)
box(lwd=1)
```

---

squeeze

*Squeeze the imputed values to be within specified boundaries.*

---

### Description

This function replaces any values in  $x$  that are lower than  $\text{bounds}[1]$  by  $\text{bounds}[1]$ , and replaces any values higher than  $\text{bounds}[2]$  by  $\text{bounds}[2]$ .

### Usage

```
squeeze(x, bounds = c(min(x[r]), max(x[r])), r = rep(TRUE, length(x)))
```

### Arguments

$x$	A numerical vector with values
$\text{bounds}$	A numerical vector of length 2 containing the lower and upper bounds. By default, the bounds are to the minimum and maximum values in $x$ .
$r$	A logical vector of length $\text{length}(x)$ that is used to select a subset in $x$ before calculating automatic bounds.

### Value

A vector of length  $\text{length}(x)$ .

### Author(s)

Stef van Buuren, 2011.

stripplot.mids

*Stripplot of observed and imputed data*

## Description

Plotting methods for imputed data using **lattice**. `stripplot` produces one-dimensional scatterplots. The function automatically separates the observed and imputed data. The functions extend the usual features of **lattice**.

## Usage

```
## S3 method for class 'mids'
stripplot(x, data, na.groups = NULL, groups = NULL,
  as.table = TRUE, theme = mice.theme(), allow.multiple = TRUE,
  outer = TRUE,
  drop.unused.levels = lattice::lattice.getOption("drop.unused.levels"),
  panel = lattice::lattice.getOption("panel.stripplot"),
  default.prepanel = lattice::lattice.getOption("prepanel.default.stripplot"),
  jitter.data = TRUE, horizontal = FALSE, ..., subscripts = TRUE,
  subset = TRUE)
```

## Arguments

- |                        |   |
|------------------------|---|
| <code>x</code>         | A <code>mids</code> object, typically created by <code>mice()</code> or <code>mice.mids()</code> .  |
| <code>data</code>      | <p>Formula that selects the data to be plotted. This argument follows the <b>lattice</b> rules for <i>formulas</i>, describing the primary variables (used for the per-panel display) and the optional conditioning variables (which define the subsets plotted in different panels) to be used in the plot.</p> <p>The formula is evaluated on the complete data set in the long form. Legal variable names for the formula include <code>names(x\$data)</code> plus the two administrative factors <code>.imp</code> and <code>.id</code>.</p> <p><b>Extended formula interface:</b> The primary variable terms (both the LHS <code>y</code> and RHS <code>x</code>) may consist of multiple terms separated by a '+' sign, e.g., <code>y1 + y2 ~ x   a * b</code>. This formula would be taken to mean that the user wants to plot both <code>y1 ~ x   a * b</code> and <code>y2 ~ x   a * b</code>, but with the <code>y1 ~ x</code> and <code>y2 ~ x</code> in <i>separate panels</i>. This behavior differs from standard <b>lattice</b>. <i>Only combine terms of the same type</i>, i.e. only factors or only numerical variables. Mixing numerical and categorical data occasionally produces odds labeling of vertical axis.</p> <p>For convenience, in <code>stripplot()</code> and <code>bwplot</code> the formula <code>y~.imp</code> may be abbreviated as <code>y</code>. This applies only to a single <code>y</code>, and does not (yet) work for <code>y1+y2~.imp</code>.</p> |
| <code>na.groups</code> | An expression evaluating to a logical vector indicating which two groups are distinguished (e.g. using different colors) in the display. The environment in which this expression is evaluated in the response indicator is <code>.na(x\$data)</code> .   |

The default `na.group = NULL` contrasts the observed and missing data in the LHS `y` variable of the display, i.e. groups created by `is.na(y)`. The expression `y` creates the groups according to `is.na(y)`. The expression `y1 & y2` creates groups by `is.na(y1) & is.na(y2)`, and `y1 | y2` creates groups as `is.na(y1) | is.na(y2)`, and so on.

<code>groups</code>	This is the usual <code>groups</code> arguments in <b>lattice</b> . It differs from <code>na.groups</code> because it evaluates in the completed data <code>data.frame(complete(x, "long", inc=TRUE))</code> (as usual), whereas <code>na.groups</code> evaluates in the response indicator. See <a href="#">xyplot</a> for more details. When both <code>na.groups</code> and <code>groups</code> are specified, <code>na.groups</code> takes precedence, and <code>groups</code> is ignored.
<code>as.table</code>	See <a href="#">xyplot</a> .
<code>theme</code>	A named list containing the graphical parameters. The default function <code>mice.theme</code> produces a short list of default colors, line width, and so on. The extensive list may be obtained from <code>trellis.par.get()</code> . Global graphical parameters like <code>col</code> or <code>cex</code> in high-level calls are still honored, so first experiment with the global parameters. Many setting consists of a pair. For example, <code>mice.theme</code> defines two symbol colors. The first is for the observed data, the second for the imputed data. The theme settings only exist during the call, and do not affect the trellis graphical parameters.
<code>allow.multiple</code>	See <a href="#">xyplot</a> .
<code>outer</code>	See <a href="#">xyplot</a> .
<code>drop.unused.levels</code>	See <a href="#">xyplot</a> .
<code>panel</code>	See <a href="#">xyplot</a> .
<code>default.prepanel</code>	See <a href="#">xyplot</a> .
<code>jitter.data</code>	See <a href="#">panel.xyplot</a> .
<code>horizontal</code>	See <a href="#">xyplot</a> .
<code>subscripts</code>	See <a href="#">xyplot</a> .
<code>subset</code>	See <a href="#">xyplot</a> .
<code>...</code>	Further arguments, usually not directly processed by the high-level functions documented here, but instead passed on to other functions.

## Details

The argument `na.groups` may be used to specify (combinations of) missingness in any of the variables. The argument `groups` can be used to specify groups based on the variable values themselves. Only one of both may be active at the same time. When both are specified, `na.groups` takes precedence over `groups`.

Use the `subset` and `na.groups` together to plots parts of the data. For example, select the first imputed data set by `subset=.imp==1`.

Graphical parameters like `col`, `pch` and `cex` can be specified in the arguments list to alter the plotting symbols. If `length(col)==2`, the color specification to define the observed and missing groups. `col[1]` is the color of the 'observed' data, `col[2]` is the color of the missing or imputed data.

A convenient color choice is `col=mdc(1:2)`, a transparent blue color for the observed data, and a transparent red color for the imputed data. A good choice is `col=mdc(1:2)`, `pch=20`, `cex=1.5`. These choices can be set for the duration of the session by running `mice.theme()`.

### Value

The high-level functions documented here, as well as other high-level Lattice functions, return an object of class "trellis". The `update` method can be used to subsequently update components of the object, and the `print` method (usually called by default) will plot it on an appropriate plotting device.

### Note

The first two arguments (`x` and `data`) are reversed compared to the standard Trellis syntax implemented in **lattice**. This reversal was necessary in order to benefit from automatic method dispatch.

In **mice** the argument `x` is always a `mids` object, whereas in **lattice** the argument `x` is always a formula.

In **mice** the argument `data` is always a formula object, whereas in **lattice** the argument `data` is usually a data frame.

All other arguments have identical interpretation.

### Author(s)

Stef van Buuren

### References

Sarkar, Deepayan (2008) *Lattice: Multivariate Data Visualization with R*, Springer. <http://lmdvr.r-forge.r-project.org/>

van Buuren S and Groothuis-Oudshoorn K (2011). `mice`: Multivariate Imputation by Chained Equations in R. *Journal of Statistical Software*, **45**(3), 1-67. <http://www.jstatsoft.org/v45/i03/>

### See Also

[mice](#), [xyplot](#), [densityplot](#), [bwplot](#), [Lattice](#) for an overview of the package, as well as [stripplot](#), [panel.stripplot](#), [print.trellis](#), [trellis.par.set](#)

### Examples

```
require(lattice)

imp <- mice(boys, maxit=1)

### stripplot, all numerical variables
## Not run: stripplot(imp)

### same, but with improved display
## Not run: stripplot(imp, col=c("grey",mdc(2)),pch=c(1,20))
```

```

### distribution per imputation of height, weight and bmi
### labeled by their own missingness
## Not run: stripplot(imp, hgt+wgt+bmi~.imp, cex=c(2,4), pch=c(1,20),jitter=FALSE,
layout=c(3,1))
## End(Not run)

### same, but labeled with the missingness of wgt (just four cases)
## Not run: stripplot(imp, hgt+wgt+bmi~.imp, na=wgt, cex=c(2,4), pch=c(1,20),jitter=FALSE,
layout=c(3,1))
## End(Not run)

### distribution of age and height, labeled by missingness in height
### most height values are missing for those around
### the age of two years
### some additional missings occur in region WEST
## Not run: stripplot(imp, age+hgt~.imp|reg, hgt, col=c(hcl(0,0,40,0.2), mdc(2)),pch=c(1,20))

### heavily jitted relation between two categorical variables
### labeled by missingness of gen
### aggregated over all imputed data sets
## Not run: stripplot(imp, gen~phb, factor=2, cex=c(8,1), hor=TRUE)

### circle fun
stripplot(imp, gen~.imp, na = wgt, factor = 2, cex = c(8.6),
          hor = FALSE, outer = TRUE, scales = "free", pch = c(1,19))

```

---

summary.mira

*Summary of a mira object*


---

## Description

Summary of a mira object

Summary of a mipo object

Summary of a mids object

Summary of a mads object

## Usage

```
## S3 method for class 'mira'
summary(object, ...)
```

```
## S3 method for class 'mipo'
summary(object, ...)
```

```
## S3 method for class 'mids'
summary(object, ...)
```

```
## S3 method for class 'mads'  
summary(object, ...)
```

**Arguments**

object	A mira object
...	Other parameters passed down to print() and summary()

**Value**

NULL  
A table containing summary statistics of the pooled analysis  
NULL  
NULL

**See Also**

[mira](#)  
[mipo](#)  
[mids](#)  
[mads](#)

---

supports.transparent *Supports semi-transparent foreground colors?*

---

**Description**

This function is used by mdc() to find out whether the current device supports semi-transparent foreground colors.

**Usage**

```
supports.transparent()
```

**Details**

The function calls the function dev.capabilities() from the package grDevices. The function return FALSE if the status of the current device is unknown.

**Value**

TRUE or FALSE

**See Also**

[mdc dev.capabilities](#)



## Examples

```
supports.transparent()
```

---

tbc	<i>Terneuzen birth cohort</i>
-----	-------------------------------

---

## Description

Data of subset of the Terneuzen Birth Cohort data on child growth.

## Format

tbc is a data frame with 3951 rows and 11 columns:

**id** Person number  
**occ** Occasion number  
**nocc** Number of occasions  
**first** Is this the first record for this person? (TRUE/FALSE)  
**typ** Type of data (all observed)  
**age** Age (years)  
**sex** Sex 1=M, 2=F  
**hgt.z** Height Z-score  
**wgt.z** Weight Z-score  
**bmi.z** BMI Z-score  
**ao** Adult overweight (0=no, 1=yes)

tbc.target is a data frame with 2612 rows and 3 columns:

**id** Person number  
**ao** Adult overweight (0=no, 1=yes)  
**bmi.z.jv** BMI Z-score as young adult (18-29 years)

## Details

This tbc data set is a random subset of persons from a much larger collection of data from the Terneuzen Birth Cohort. The total cohort comprises of 2604 unique persons, whereas the subset in tbc covers 306 persons. The tbc.target is an auxiliary data set containing two outcomes at adult age. For more details, see De Kroon et al (2008, 2010, 2011). The imputation methodology is explained in Chapter 9 of Van Buuren (2012).

**Source**

De Kroon, M. L. A., Renders, C. M., Kuipers, E. C., van Wouwe, J. P., van Buuren, S., de Jonge, G. A., Hirasing, R. A. (2008). Identifying metabolic syndrome without blood tests in young adults - The Terneuzen birth cohort. *European Journal of Public Health*, 18(6), 656-660.

De Kroon, M. L. A., Renders, C. M., Van Wouwe, J. P., Van Buuren, S., Hirasing, R. A. (2010). The Terneuzen birth cohort: BMI changes between 2 and 6 years correlate strongest with adult overweight. *PLoS ONE*, 5(2), e9155.

De Kroon, M. L. A. (2011). *The Terneuzen Birth Cohort. Detection and Prevention of Overweight and Cardiometabolic Risk from Infancy Onward*. Dissertation, Vrije Universiteit, Amsterdam. <http://dare.uvu.nl/handle/1871/23806>

Van Buuren, S. (2012). *Flexible Imputation of Missing Data*. Boca Raton, FL: Chapman & Hall/CRC Press.

**Examples**

```
data <- tbc
md.pattern(data)
```

---

version	<i>Echoes the package version number</i>
---------	--

---

**Description**

Echoes the package version number

**Usage**

```
version(pkg = "mice")
```

**Arguments**

pkg                    A character vector with the package name.

**Value**

A character vector containing the package name, version number and installed directory.

**Author(s)**

Stef van Buuren, Oct 2010

**Examples**

```
version()
version("base")
```

---

walking	<i>Walking disability data</i>
---------	--------------------------------

---

### Description

Two items YA and YB measuring walking disability in samples A, B and E.

### Format

A data frame with 890 rows on the following 5 variables:

**sex** Sex of respondent (factor)

**age** Age of respondent

**YA** Item administered in samples A and E (factor)

**YB** Item administered in samples B and E (factor)

**src** Source: Sample A, B or E (factor)

### Details

Example dataset to demonstrate imputation of two items (YA and YB). Item YA is administered to sample A and sample E, item YB is administered to sample B and sample E, so sample E acts as a bridge study. Imputation using a bridge study is better than simple equating or than imputation under independence.

Item YA corresponds to the HAQ8 item, and item YB corresponds to the GAR9 items from Van Buuren et al (2005). Sample E (as well as sample B) is the Euridiss study (n=292), sample A is the ERGOPLUS study (n=306).

See Van Buuren (2012) chapter 7 for more details on the imputation methodology.

### References

van Buuren, S., Eyres, S., Tennant, A., Hopman-Rock, M. (2005). Improving comparability of existing data by Response Conversion. *Journal of Official Statistics*, **21**(1), 53-72.

van Buuren, S. (2012). *Flexible Imputation of Missing Data*. Boca Raton, FL: Chapman & Hall/CRC.

### Examples

```
md.pattern(walking)

micemill <- function(n) {
  for (i in 1:n) {
    imp <- mice.mids(imp) # global assignment
    cors <- with(imp, cor(as.numeric(YA),
                        as.numeric(YB),
                        method="kendall"))
```

```

    tau <- rbind(tau, getfit(cors, s=TRUE)) # global assignment
  }
}

plotit <- function()
  matplot(x=1:nrow(tau),y=tau,
          ylab=expression(paste("Kendall's ",tau)),
          xlab="Iteration", type="l", lwd=1,
          lty=1:10,col="black")

tau <- NULL
imp <- mice(walking, max=0, m=10, seed=92786)
pred <- imp$pred
pred[,c("src","age","sex")] <- 0
imp <- mice(walking, max=0, m=3, seed=92786, pred=pred)
micemill(5)
plotit()

### to get figure 7.8 van Buuren (2012) use m=10 and micemill(20)

```

---

windspeed

*Subset of Irish wind speed data*


---

### Description

Subset of Irish wind speed data

### Format

A data frame with 433 rows and 6 columns containing the daily average wind speeds within the period 1961-1978 at meteorological stations in the Republic of Ireland. The data are a random sample from a larger data set.

**RochePt** Roche Point

**Rosslare** Rosslare

**Shannon** Shannon

**Dublin** Dublin

**Clones** Clones

**MalinHead** Malin Head

### Details

The original data set is much larger and was analyzed in detail by Haslett and Raftery (1989). Van Buuren et al (2006) used this subset to investigate the influence of extreme MAR mechanisms on the quality of imputation.

## References

Haslett, J. and Raftery, A. E. (1989). *Space-time Modelling with Long-memory Dependence: Assessing Ireland's Wind Power Resource (with Discussion)*. Applied Statistics 38, 1-50. <http://lib.stat.cmu.edu/datasets/wind.desc> and <http://lib.stat.cmu.edu/datasets/wind.data>

Van Buuren, S., Brand, J.P.L., Groothuis-Oudshoorn C.G.M., Rubin, D.B. (2006) Fully conditional specification in multivariate imputation. *Journal of Statistical Computation and Simulation*, **76**, 12, 1049–1064.

## Examples

```
windspeed[1:3,]
```

---

```
with.mids
```

*Evaluate an expression in multiple imputed datasets*

---

## Description

Performs a computation of each of imputed datasets in data.

## Usage

```
## S3 method for class 'mids'
with(data, expr, ...)
```

## Arguments

data	An object of type mids, which stands for 'multiply imputed data set', typically created by a call to function mice().
expr	An expression with a formula object, with the response on the left of a ~ operator, and the terms, separated by + operators, on the right. See the documentation of <a href="#">lm</a> and <a href="#">formula</a> for details.
...	Additional parameters passed to expr

## Value

A list object of S3 class mira

## Author(s)

Karin Oudshoorn, Stef van Buuren 2009-2012

## References

van Buuren S and Groothuis-Oudshoorn K (2011). mice: Multivariate Imputation by Chained Equations in R. *Journal of Statistical Software*, **45**(3), 1-67. <http://www.jstatsoft.org/v45/i03/>

## See Also

[mids](#), [mira](#), [pool](#), [pool.compare](#), [pool.r.squared](#)

## Examples

```
imp <- mice(nhanes2)
fit1 <- with(data=imp,exp=lm(bmi~age+hyp+chl))
fit2 <- with(data=imp,exp=glm(hyp~age+bmi+chl,family=binomial))
anova.imp <- with(data=imp,exp=anova(lm(bmi~age+hyp+chl)))
```

---

xyplot.mads

*Scatterplot of amputed and non-amputed data against weighted sum scores*

---

## Description

Plotting method to investigate relation between amputed data and the weighted sum scores. Based on [lattice](#). xyplot produces scatterplots. The function plots the variables against the weighted sum scores. The function automatically separates the amputed and non-amputed data to see the relation between the amputatoin and the weighted sum scores.

## Usage

```
## S3 method for class 'mads'
xyplot(x, yvar = NULL, which.pat = NULL,
       standardized = TRUE, layout = NULL, colors = c("#fc8d62", "#8da0cb"))
```

## Arguments

x	A mads object, typically created by <a href="#">ampute</a> .
yvar	A string or vector of variable names that needs to be plotted. As a default, all variables will be plotted.
which.pat	A scalar or vector indicating which patterns need to be plotted. As a default, all patterns are plotted.
standardized	Logical. Whether the scatterplots need to be created from standardized data or not. Default is TRUE.

layout	A vector of two values indicating how the scatterplots of one pattern should be divided over the plot. For example, <code>c(2, 3)</code> indicates that the scatterplots of six variables need to be placed on 3 rows and 2 columns. There are several defaults for different <code>#variables</code> . Note that for more than 9 variables, multiple plots will be created automatically.
colors	A vector of two RGB values defining the colors of the amputated and non-amputated data respectively. RGB values can be obtained with <a href="#">hcl</a> .

**Value**

A list containing the scatterplots. Note that a new pattern will always be shown in a new plot.

**Note**

The `mids` object contains all the information you need to make any desired plots. Check [mids-class](#) or the vignette *Multivariate Amputation using Ampute* to understand the contents of class object `mids`.

**Author(s)**

Rianne Schouten, 2016

**See Also**

[ampute](#), [bwplot](#), [Lattice](#) for an overview of the package, [mids-class](#)

---

xyplot.mids

*Scatterplot of observed and imputed data*


---

**Description**

Plotting methods for imputed data using **lattice**. `xyplot()` produces a conditional scatterplots. The function automatically separates the observed (blue) and imputed (red) data. The function extends the usual features of **lattice**.

**Usage**

```
## S3 method for class 'mids'
xyplot(x, data, na.groups = NULL, groups = NULL,
       as.table = TRUE, theme = mice.theme(), allow.multiple = TRUE,
       outer = TRUE,
       drop.unused.levels = lattice::lattice.getOption("drop.unused.levels"), ...,
       subscripts = TRUE, subset = TRUE)
```

**Arguments**

x	A mids object, typically created by <code>mice()</code> or <code>mice.mids()</code> .
data	<p>Formula that selects the data to be plotted. This argument follows the <b>lattice</b> rules for <i>formulas</i>, describing the primary variables (used for the per-panel display) and the optional conditioning variables (which define the subsets plotted in different panels) to be used in the plot.</p> <p>The formula is evaluated on the complete data set in the long form. Legal variable names for the formula include <code>names(x\$data)</code> plus the two administrative factors <code>.imp</code> and <code>.id</code>.</p> <p><b>Extended formula interface:</b> The primary variable terms (both the LHS <i>y</i> and RHS <i>x</i>) may consist of multiple terms separated by a '+' sign, e.g., <math>y1 + y2 \sim x \mid a * b</math>. This formula would be taken to mean that the user wants to plot both <math>y1 \sim x \mid a * b</math> and <math>y2 \sim x \mid a * b</math>, but with the <math>y1 \sim x</math> and <math>y2 \sim x</math> in <i>separate panels</i>. This behavior differs from standard <b>lattice</b>. <i>Only combine terms of the same type</i>, i.e. only factors or only numerical variables. Mixing numerical and categorical data occasionally produces odds labeling of vertical axis.</p>
na.groups	<p>An expression evaluating to a logical vector indicating which two groups are distinguished (e.g. using different colors) in the display. The environment in which this expression is evaluated in the response indicator <code>is.na(x\$data)</code>.</p> <p>The default <code>na.group = NULL</code> contrasts the observed and missing data in the LHS <i>y</i> variable of the display, i.e. groups created by <code>is.na(y)</code>. The expression <i>y</i> creates the groups according to <code>is.na(y)</code>. The expression <math>y1 \&amp; y2</math> creates groups by <code>is.na(y1) &amp; is.na(y2)</code>, and <math>y1 \mid y2</math> creates groups as <code>is.na(y1) \mid is.na(y2)</code>, and so on.</p>
groups	This is the usual groups arguments in <b>lattice</b> . It differs from <code>na.groups</code> because it evaluates in the completed data <code>data.frame(complete(x, "long", inc=TRUE))</code> (as usual), whereas <code>na.groups</code> evaluates in the response indicator. See <a href="#">xyplot</a> for more details. When both <code>na.groups</code> and <code>groups</code> are specified, <code>na.groups</code> takes precedence, and <code>groups</code> is ignored.
as.table	See <a href="#">xyplot</a> .
theme	A named list containing the graphical parameters. The default function <code>mice.theme</code> produces a short list of default colors, line width, and so on. The extensive list may be obtained from <code>trellis.par.get()</code> . Global graphical parameters like <code>col</code> or <code>cex</code> in high-level calls are still honored, so first experiment with the global parameters. Many setting consists of a pair. For example, <code>mice.theme</code> defines two symbol colors. The first is for the observed data, the second for the imputed data. The theme settings only exist during the call, and do not affect the trellis graphical parameters.
allow.multiple	See <a href="#">xyplot</a> .
outer	See <a href="#">xyplot</a> .
drop.unused.levels	See <a href="#">xyplot</a> .
subscripts	See <a href="#">xyplot</a> .
subset	See <a href="#">xyplot</a> .
...	Further arguments, usually not directly processed by the high-level functions documented here, but instead passed on to other functions.



## Details

The argument `na.groups` may be used to specify (combinations of) missingness in any of the variables. The argument `groups` can be used to specify groups based on the variable values themselves. Only one of both may be active at the same time. When both are specified, `na.groups` takes precedence over `groups`.

Use the `subset` and `na.groups` together to plots parts of the data. For example, select the first imputed data set by `subset=.imp==1`.

Graphical parameters like `col`, `pch` and `cex` can be specified in the arguments list to alter the plotting symbols. If `length(col)==2`, the color specification to define the observed and missing groups. `col[1]` is the color of the 'observed' data, `col[2]` is the color of the missing or imputed data. A convenient color choice is `col=mdc(1:2)`, a transparent blue color for the observed data, and a transparent red color for the imputed data. A good choice is `col=mdc(1:2)`, `pch=20`, `cex=1.5`. These choices can be set for the duration of the session by running `mice.theme()`.

## Value

The high-level functions documented here, as well as other high-level Lattice functions, return an object of class "trellis". The `update` method can be used to subsequently update components of the object, and the `print` method (usually called by default) will plot it on an appropriate plotting device.

## Note

The first two arguments (`x` and `data`) are reversed compared to the standard Trellis syntax implemented in **lattice**. This reversal was necessary in order to benefit from automatic method dispatch.

In **mice** the argument `x` is always a `mids` object, whereas in **lattice** the argument `x` is always a formula.

In **mice** the argument `data` is always a formula object, whereas in **lattice** the argument `data` is usually a data frame.

All other arguments have identical interpretation.

## Author(s)

Stef van Buuren

## References

Sarkar, Deepayan (2008) *Lattice: Multivariate Data Visualization with R*, Springer. <http://lmdvr.r-forge.r-project.org/>

van Buuren S and Groothuis-Oudshoorn K (2011). `mice`: Multivariate Imputation by Chained Equations in R. *Journal of Statistical Software*, **45**(3), 1-67. <http://www.jstatsoft.org/v45/i03/>

## See Also

`mice`, `stripplot`, `densityplot`, `bwplot`, `Lattice` for an overview of the package, as well as `xyplot`, `panel.xyplot`, `print.trellis`, `trellis.par.set`

**Examples**

```
require(lattice)

imp <- mice(boys, maxit=1)

### xyplot: scatterplot by imputation number
### observe the erroneous outlying imputed values
### (caused by imputing hgt from bmi)
xyplot(imp, hgt~age|.imp, pch=c(1,20),cex=c(1,1.5))

### same, but label with missingness of wgt (four cases)
xyplot(imp, hgt~age|.imp, na.group=wgt, pch=c(1,20),cex=c(1,1.5))
```

# Index

## \*Topic **classes**

- mids-class, 93
- mipo-class, 97
- mira-class, 98

## \*Topic **datagen**

- mice.impute.2l.norm, 59
- mice.impute.2lonly.mean, 62
- mice.impute.cart, 67
- mice.impute.fastpmm, 68
- mice.impute.lda, 70
- mice.impute.logreg, 71
- mice.impute.logreg.boot, 72
- mice.impute.mean, 73
- mice.impute.norm, 76
- mice.impute.norm.boot, 77
- mice.impute.norm.nob, 78
- mice.impute.norm.predict, 79
- mice.impute.passive, 80
- mice.impute.pmm, 81
- mice.impute.polr, 83
- mice.impute.polyreg, 84
- mice.impute.quadratic, 86
- mice.impute.rf, 87
- mice.impute.ri, 89
- mice.impute.sample, 90

## \*Topic **datasets**

- boys, 18
- fdd, 33
- fdgs, 35
- leiden85, 46
- mammalsleep, 49
- nhanes, 101
- nhanes2, 102
- pattern, 104
- popmis, 113
- pops, 114
- potthoffroy, 115
- selfreport, 121
- tbc, 129

- walking, 131

- windspeed, 132

## \*Topic **hplot**

- bwplot.mids, 21
- densityplot.mids, 29
- mdc, 53
- stripplot.mids, 124
- supports.transparent, 128
- xyplot.mids, 135

## \*Topic **htest**

- pool, 106
- pool.compare, 108
- pool.r.squared, 110
- pool.scalar, 112

## \*Topic **iteration**

- mice, 54
- mice.mids, 91

## \*Topic **manip**

- cbind.mids, 24
- complete, 27
- getfit, 40
- ibind, 42
- mids2mplus, 94
- mids2spss, 95
- rbind.mids, 120

## \*Topic **mice**

- mice.impute.midastouch, 74

## \*Topic **mids**

- as.mids, 16

## \*Topic **misc**

- fico, 36
- flux, 37
- fluxplot, 38
- nelsonaalen, 99
- quickpred, 118
- version, 130

## \*Topic **multivariate**

- glm.mids, 41
- lm.mids, 47

- with.mids, 133
- \*Topic **univar**
  - cc, 26
  - cci, 26
  - ic, 43
  - ici, 44
  - md.pairs, 50
  - md.pattern, 52
- .norm.draw (norm.draw), 103
- .pmm.match, 4
- 2l.norm (mice.impute.2l.norm), 59
- 2l.pan (mice.impute.2l.pan), 60
- 2lonly.mean (mice.impute.2lonly.mean), 62
- 2lonly.norm (mice.impute.2lonly.norm), 63
- 2lonly.pmm (mice.impute.2lonly.pmm), 65
- ampute, 5, 9–15, 20, 49, 134, 135
- ampute.continuous, 6, 8, 9
- ampute.default.freq, 6, 10
- ampute.default.odds, 6, 11, 14
- ampute.default.patterns, 6, 10, 11, 11, 12, 13, 15
- ampute.default.type, 6, 9, 10, 12
- ampute.default.weights, 6, 13
- ampute.discrete, 6, 8, 13
- ampute.mcar, 8, 14
- appendbreak, 15
- as.mids, 16
- as.mira, 17, 107
- boys, 18
- bwplot, 8, 20, 23, 32, 126, 135, 137
- bwplot (bwplot.mids), 21
- bwplot.mads, 19
- bwplot.mids, 21
- cart (mice.impute.cart), 67
- cbind.mids, 24, 42, 121
- cc, 26, 27, 43
- cci, 26, 26, 44, 99, 103
- complete, 27, 58, 91
- complete.cases, 27
- densityplot, 23, 32, 126, 137
- densityplot (densityplot.mids), 29
- densityplot.mids, 29
- dev.capabilities, 128
- extractBS, 32
- fastpmm (mice.impute.fastpmm), 68
- fdd, 33
- fdgs, 35
- fico, 36, 38, 40
- flux, 37, 37, 40
- fluxplot, 37, 38, 38
- formula, 41, 47, 133
- getfit, 40
- glm, 41, 42, 72, 73
- glm.fit, 72, 73
- glm.mids, 41, 110
- hazard (nelsonaalen), 99
- hcl, 54, 135
- ibind, 25, 42, 121
- ic, 43, 44
- ici, 26, 27, 43, 44
- ici, data.frame-method (ici), 44
- ici, matrix-method (ici), 44
- ici, mids-method (ici), 44
- is.mads, 44
- is.mids, 45
- is.mipo, 45
- is.mira, 46
- Lattice, 20, 23, 32, 126, 135, 137
- lattice, 20, 134
- lda, 71
- leiden85, 46
- lm, 41, 47, 48, 133
- lm.mids, 47, 110
- mads, 116, 128
- mads-class, 48
- mammalsleep, 49
- md.pairs, 50
- md.pattern, 6, 12, 37, 38, 40, 52
- mdc, 53, 128
- mean, 74
- mgg (selfreport), 121
- mice, 5, 7, 8, 23, 28, 32, 54, 68, 71–74, 79, 81, 84, 85, 88, 91, 94, 106, 119, 126, 137
- mice.impute.2l.norm, 59, 61
- mice.impute.2l.pan, 60, 64–66
- mice.impute.2lonly.mean, 62
- mice.impute.2lonly.norm, 63, 66

- mice.impute.2lonly.pmm, [64](#), [65](#)
- mice.impute.cart, [67](#), [88](#)
- mice.impute.fastpmm, [68](#)
- mice.impute.lda, [70](#)
- mice.impute.logreg, [71](#)
- mice.impute.logreg.boot, [72](#)
- mice.impute.mean, [73](#)
- mice.impute.midastouch, [74](#)
- mice.impute.norm, [64](#), [76](#), [79](#)
- mice.impute.norm.boot, [77](#)
- mice.impute.norm.nob, [78](#)
- mice.impute.norm.predict, [79](#)
- mice.impute.passive, [80](#)
- mice.impute.pmm, [66](#), [70](#), [81](#), [87](#)
- mice.impute.polr, [83](#)
- mice.impute.polyreg, [71](#), [84](#)
- mice.impute.quadratic, [86](#)
- mice.impute.rf, [68](#), [87](#)
- mice.impute.rfcat, [88](#)
- mice.impute.rfcont, [88](#)
- mice.impute.ri, [89](#)
- mice.impute.sample, [90](#)
- mice.mids, [91](#)
- mice.theme, [92](#)
- mids, [25](#), [28](#), [42](#), [48](#), [57](#), [58](#), [91](#), [95](#), [96](#), [98](#), [106](#), [117](#), [119](#), [121](#), [128](#), [134](#)
- mids (mids-class), [93](#)
- mids-class, [93](#)
- mids2mplus, [94](#)
- mids2spss, [95](#), [95](#)
- mipo, [94](#), [98](#), [117](#), [128](#)
- mipo (mipo-class), [97](#)
- mipo-class, [97](#)
- mira, [18](#), [40](#), [42](#), [48](#), [94](#), [98](#), [117](#), [128](#), [134](#)
- mira (mira-class), [98](#)
- mira-class, [98](#)
- multinom, [84](#), [85](#)
  
- na.omit, [26](#)
- ncc, [99](#), [103](#)
- nelsonaalen, [99](#)
- nhanes, [101](#), [102](#)
- nhanes2, [101](#), [102](#)
- nic, [99](#), [103](#)
- norm (mice.impute.norm), [76](#)
- norm.boot (mice.impute.norm.boot), [77](#)
- norm.draw, [103](#)
- norm.nob (mice.impute.norm.nob), [78](#)
  
- norm.predict  
(mice.impute.norm.predict), [79](#)
  
- panel.bwplot, [23](#)
- panel.densityplot, [32](#)
- panel.stripplot, [126](#)
- panel.xyplot, [106](#), [125](#), [137](#)
- pattern, [104](#)
- pattern1 (pattern), [104](#)
- pattern2 (pattern), [104](#)
- pattern3 (pattern), [104](#)
- pattern4 (pattern), [104](#)
- plot.mids, [105](#)
- pmm (mice.impute.pmm), [81](#)
- polr, [84](#), [85](#)
- pool, [97](#), [98](#), [106](#), [111](#), [113](#), [134](#)
- pool.compare, [108](#), [134](#)
- pool.r.squared, [110](#), [134](#)
- pool.scalar, [111](#), [112](#)
- popmis, [113](#)
- pops, [114](#)
- potthoffroy, [115](#)
- print, [23](#), [31](#), [126](#), [137](#)
- print.mads, [116](#)
- print.mids, [117](#)
- print.mipo (print.mids), [117](#)
- print.mira (print.mids), [117](#)
- print.trellis, [23](#), [32](#), [126](#), [137](#)
  
- quadratic (mice.impute.quadratic), [86](#)
- quickpred, [118](#)
  
- randomForest, [88](#)
- rbind.mids, [25](#), [42](#), [120](#)
- rgb, [54](#)
- ri (mice.impute.ri), [89](#)
- rpart, [68](#)
- rpart.control, [67](#), [68](#)
  
- selfreport, [121](#)
- set.seed, [58](#), [91](#)
- sleep (mammalsleep), [49](#)
- squeeze, [123](#)
- stripplot, [23](#), [32](#), [126](#), [137](#)
- stripplot (stripplot.mids), [124](#)
- stripplot.mids, [124](#)
- summary.mads (summary.mira), [127](#)
- summary.mids (summary.mira), [127](#)
- summary.mipo (summary.mira), [127](#)

summary.mira, [127](#)  
supports.transparent, [128](#)

tbc, [129](#)  
terneuzen (tbc), [129](#)  
transparent (supports.transparent), [128](#)  
trellis.par.set, [23](#), [32](#), [54](#), [126](#), [137](#)

update, [23](#), [31](#), [126](#), [137](#)

vcov, [107](#), [110](#)  
version, [130](#)

walking, [131](#)  
windspeed, [132](#)  
with.mids, [40–42](#), [47](#), [58](#), [98](#), [107](#), [133](#)

xyplot, [8](#), [22](#), [23](#), [30](#), [32](#), [54](#), [106](#), [125](#), [126](#),  
[136](#), [137](#)  
xyplot (xyplot.mids), [135](#)  
xyplot.mads, [134](#)  
xyplot.mids, [54](#), [135](#)