# Package 'datawizard’ 

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adjust Adjust data for the effect of other variable(s)

## Description

This function can be used to adjust the data for the effect of other variables present in the dataset. It is based on an underlying fitting of regressions models, allowing for quite some flexibility, such as including factors as random effects in mixed models (multilevel partialization), continuous variables as smooth terms in general additive models (non-linear partialization) and/or fitting these models under a Bayesian framework. The values returned by this function are the residuals of the regression models. Note that a regular correlation between two "adjusted" variables is equivalent to the partial correlation between them.

## Usage

adjust(
data,
effect $=$ NULL,
select = NULL,
exclude = NULL,
multilevel = FALSE,
additive = FALSE,
bayesian = FALSE,
keep_intercept = FALSE,
ignore_case = FALSE
)
data_adjust(
data,
effect $=$ NULL,
select $=$ NULL,
exclude = NULL,
multilevel = FALSE,
additive = FALSE,
bayesian = FALSE,
keep_intercept = FALSE,

```
        ignore_case = FALSE
)
```


## Arguments

## data

A data frame.
effect
select Variables that will be included when performing the required tasks. Can be either

- a variable specified as a literal variable name (e.g., column_name),
- a string with the variable name (e.g., "column_name"), or a character vector of variable names (e.g., c("col1", "col2", "col3")),
- a formula with variable names (e.g., $\sim$ column_1 + column_2),
- a vector of positive integers, giving the positions counting from the left (e.g. 1 or $c(1,3,5)$ ),
- a vector of negative integers, giving the positions counting from the right (e.g., -1 or $-1:-3$ ),
- one of the following select-helpers: starts_with(""), ends_with(""), contains(""), a range using : or regex (""),
- or a function testing for logical conditions, e.g. is.numeric() (or is.numeric), or any user-defined function that selects the variables for which the function returns TRUE (like: foo <- function(x) mean (x) > 3),
- ranges specified via literal variable names, select-helpers (except regex()) and (user-defined) functions can be negated, i.e. return non-matching elements, when prefixed with a -, e.g. -ends_with(""), -is.numeric or -Sepal.Width:Petal.Length. Note: Negation means that matches are excluded, and thus, the exclude argument can be used alternatively. For instance, select=-ends_with("Length") (with -) is equivalent to exclude=ends_with("Length") (no -). In case negation should not work as expected, use the exclude argument instead.

If NULL, selects all columns. Patterns that found no matches are silently ignored, e.g. find_columns(iris, select = c("Species", "Test")) will just return "Species".
exclude See select, however, column names matched by the pattern from exclude will be excluded instead of selected. If NULL (the default), excludes no columns.
multilevel If TRUE, the factors are included as random factors. Else, if FALSE (default), they are included as fixed effects in the simple regression model.
additive If TRUE, continuous variables as included as smooth terms in additive models. The goal is to regress-out potential non-linear effects.
bayesian If TRUE, the models are fitted under the Bayesian framework using rstanarm.
keep_intercept If FALSE (default), the intercept of the model is re-added. This avoids the centering around 0 that happens by default when regressing out another variable (see the examples below for a visual representation of this).
ignore_case Logical, if TRUE and when one of the select-helpers or a regular expression is used in select, ignores lower/upper case in the search pattern when matching against variable names.

## Value

A data frame comparable to data, with adjusted variables.

## Examples

```
adjusted_all <- adjust(attitude)
head(adjusted_all)
adjusted_one <- adjust(attitude, effect = "complaints", select = "rating")
head(adjusted_one)
adjust(attitude, effect = "complaints", select = "rating", bayesian = TRUE)
adjust(attitude, effect = "complaints", select = "rating", additive = TRUE)
attitude$complaints_LMH <- cut(attitude$complaints, 3)
adjust(attitude, effect = "complaints_LMH", select = "rating", multilevel = TRUE)
if (require("bayestestR")) {
    # Generate data
    data <- simulate_correlation(n = 100, r = 0.7)
    data$V2 <- (5 * data$V2) + 20 # Add intercept
    # Adjust
    adjusted <- adjust(data, effect = "V1", select = "V2")
    adjusted_icpt <- adjust(data, effect = "V1", select = "V2", keep_intercept = TRUE)
    # Visualize
    plot(data$V1, data$V2,
        pch = 19, col = "blue",
        ylim = c(min(adjusted$V2), max(data$V2)),
        main = "Original (blue), adjusted (green), and adjusted - intercept kept (red) data"
    )
    abline(lm(V2 ~ V1, data = data), col = "blue")
    points(adjusted$V1, adjusted$V2, pch = 19, col = "green")
    abline(lm(V2 ~ V1, data = adjusted), col = "green")
    points(adjusted_icpt$V1, adjusted_icpt$V2, pch = 19, col = "red")
    abline(lm(V2 ~ V1, data = adjusted_icpt), col = "red")
}
```

categorize
Recode (or "cut") data into groups of values.

## Description

This functions divides the range of variables into intervals and recodes the values inside these intervals according to their related interval. It is basically a wrapper around base R's cut(), providing a simplified and more accessible way to define the interval breaks (cut-off values).

```
Usage
    categorize(x, ...)
    data_cut(x, ...)
    \#\# S3 method for class 'numeric'
    categorize(
        x ,
        split = "median",
        n_groups = NULL,
        range \(=\) NULL,
        lowest = 1,
        labels = NULL,
        verbose = TRUE,
    )
    \#\# S3 method for class 'data.frame'
    categorize(
        x ,
        select \(=\) NULL,
        exclude \(=\) NULL,
        split = "median",
        n_groups = NULL,
        range = NULL,
        lowest = 1,
        labels = NULL,
        append = FALSE,
        ignore_case = FALSE,
        verbose = TRUE,
    )
```


## Arguments

x
. . .
split

A (grouped) data frame, numeric vector or factor.
not used.
Character vector, indicating at which breaks to split variables, or numeric values with values indicating breaks. If character, may be one of "median", "mean", "quantile", "equal_length", or "equal_range". "median" or "mean" will return dichotomous variables, split at their mean or median, respectively. "quantile" and "equal_length" will split the variable into n_groups groups, where each group refers to an interval of a specific range of values. Thus, the length of each interval will be based on the number of groups. "equal_range" also splits the variable into multiple groups, however, the length of the interval is given, and the number of resulting groups (and hence, the number of breaks) will be determined by how many intervals can be generated, based on the full range of the variable.

| n_groups | If split is "quantile" or "equal_length", this defines the number of re- <br> quested groups (i.e. resulting number of levels or values) for the recoded vari- <br> able(s). "quantile" will define intervals based on the distribution of the vari- <br> able, while "equal_length" tries to divide the range of the variable into pieces <br> of equal length. |
| :--- | :--- |
| range | If split = "equal_range", this defines the range of values that are recoded into <br> a new value. |
| lowest | Minimum value of the recoded variable(s). If NULL (the default), for numeric <br> variables, the minimum of the original input is preserved. For factors, the default <br> minimum is 1. For split = "equal_range", the default minimum is always 1, <br> unless specified otherwise in lowest. |
| labels $\quad$Character vector of value labels. If not NULL, categorize() will returns factors <br> instead of numeric variables, with labels used for labelling the factor levels. <br> Toggle warnings. <br> verbose <br> select$\quad$Variables that will be included when performing the required tasks. Can be <br> either |  | either

- a variable specified as a literal variable name (e.g., column_name),
- a string with the variable name (e.g., "column_name"), or a character vector of variable names (e.g., c("col1", "col2", "col3")),
- a formula with variable names (e.g., $\sim$ column_ $1+$ column_2),
- a vector of positive integers, giving the positions counting from the left (e.g. 1 or $c(1,3,5)$ ),
- a vector of negative integers, giving the positions counting from the right (e.g., -1 or -1 :-3),
- one of the following select-helpers: starts_with(""), ends_with(""), contains(""), a range using : or regex(""),
- or a function testing for logical conditions, e.g. is.numeric() (or is.numeric), or any user-defined function that selects the variables for which the function return TRUE (like: foo <- function $(x)$ mean $(x)>3$ ),
- ranges specified via literal variable names, select-helpers (except regex()) and (user-defined) functions can be negated, i.e. return non-matching elements, when prefixed with a -, e.g. -ends_with(""), -is.numeric or -Sepal.Width:Petal.Length. Note: Negation means that matches are excluded, and thus, the exclude argument can be used alternatively. For instance, select=-ends_with("Length") (with -) is equivalent to exclude=ends_with("Length") (no -). In case negation should not work as expected, use the exclude argument instead.
If NULL, selects all columns. Patterns that found no matches are silently ignored, e.g. find_columns(iris, select $=c($ "Species", "Test")) will just return "Species".
exclude See select, however, column names matched by the pattern from exclude will be excluded instead of selected. If NULL (the default), excludes no columns.
append Logical or string. If TRUE, recoded or converted variables get new column names and are appended (column bind) to $x$, thus returning both the original and the recoded variables. The new columns get a suffix, based on the calling function:
"_r" for recode functions, "_n" for to_numeric(), "_f" for to_factor(), or "_s" for slide(). If append=FALSE, original variables in $x$ will be overwritten by their recoded versions. If a character value, recoded variables are appended with new column names (using the defined suffix) to the original data frame.
ignore_case Logical, if TRUE and when one of the select-helpers or a regular expression is used in select, ignores lower/upper case in the search pattern when matching against variable names.


## Details

Splits and breaks (cut-off values): Breaks are in general exclusive, this means that these values indicate the lower bound of the next group or interval to begin. Take a simple example, a numeric variable with values from 1 to 9 . The median would be 5 , thus the first interval ranges from 1-4 and is recoded into 1 , while $5-9$ would turn into 2 (compare cbind( $1: 9$, categorize( $1: 9$ )) ). The same variable, using split = "quantile" and n_groups $=3$ would define breaks at 3.67 and 6.33 (see quantile ( $1: 9$, probs $=c(1 / 3,2 / 3)$ ) , which means that values from 1 to 3 belong to the first interval and are recoded into 1 (because the next interval starts at 3.67), 4 to 6 into 2 and 7 to 9 into 3 .

Recoding into groups with equal size or range: split = "equal_length" and split = "equal_range" try to divide the range of $x$ into intervals of similar (or same) length. The difference is that split = "equal_length" will divide the range of $x$ into $n \_g r o u p s$ pieces and thereby defining the intervals used as breaks (hence, it is equivalent to cut ( $x$, breaks = n_groups) ), while split = "equal_range" will cut $x$ into intervals that all have the length of range, where the first interval by defaults starts at 1 . The lowest (or starting) value of that interval can be defined using the lowest argument.

## Value

$x$, recoded into groups. By default $x$ is numeric, unless labels is specified. In this case, a factor is returned, where the factor levels (i.e. recoded groups are labelled accordingly.

## Selection of variables - the select argument

For most functions that have a select argument (including this function), the complete input data frame is returned, even when select only selects a range of variables. That is, the function is only applied to those variables that have a match in select, while all other variables remain unchanged. In other words: for this function, select will not omit any non-included variables, so that the returned data frame will include all variables from the input data frame.

## See Also

- Functions to rename stuff: data_rename(), data_rename_rows(), data_addprefix(), data_addsuffix()
- Functions to reorder or remove columns: data_reorder(), data_relocate(), data_remove()
- Functions to reshape, pivot or rotate data frames: data_to_long(), data_to_wide(), data_rotate()
- Functions to recode data: rescale(), reverse(), categorize(), change_code(), slide()
- Functions to standardize, normalize, rank-transform: center(), standardize(), normalize(), ranktransform(), winsorize()
- Split and merge data frames: data_partition(), data_merge()
- Functions to find or select columns: data_select(), data_find()
- Functions to filter rows: data_match(), data_filter()


## Examples

```
set.seed(123)
x <- sample(1:10, size = 50, replace = TRUE)
table(x)
# by default, at median
table(categorize(x))
# into 3 groups, based on distribution (quantiles)
table(categorize(x, split = "quantile", n_groups = 3))
# into 3 groups, user-defined break
table(categorize(x, split = c(3, 5)))
set.seed(123)
x <- sample(1:100, size = 500, replace = TRUE)
# into 5 groups, try to recode into intervals of similar length,
# i.e. the range within groups is the same for all groups
table(categorize(x, split = "equal_length", n_groups = 5))
# into 5 groups, try to return same range within groups
# i.e. 1-20, 21-40, 41-60, etc. Since the range of " }x\mathrm{ " is
# 1-100, and we have a range of 20, this results into 5
# groups, and thus is for this particular case identical
# to the previous result.
table(categorize(x, split = "equal_range", range = 20))
# return factor with value labels instead of numeric value
set.seed(123)
x <- sample(1:10, size = 30, replace = TRUE)
categorize(x, "equal_length", n_groups = 3)
categorize(x, "equal_length", n_groups = 3, labels = c("low", "mid", "high"))
```


## Description

Performs a grand-mean centering of data.

```
Usage
    center(x, ...)
    centre(x, ...)
    ## S3 method for class 'numeric'
    center(
        x,
        robust = FALSE,
        weights = NULL,
        reference = NULL,
        center = NULL,
        verbose = TRUE,
    )
    ## S3 method for class 'data.frame'
    center(
        x,
        select = NULL,
        exclude = NULL,
        robust = FALSE,
        weights = NULL,
        reference = NULL,
        center = NULL,
        force = FALSE,
        remove_na = c("none", "selected", "all"),
        append = FALSE,
        ignore_case = FALSE,
        verbose = TRUE,
    )
```


## Arguments

x
... Currently not used.
robust Logical, if TRUE, centering is done by subtracting the median from the variables. If FALSE, variables are centered by subtracting the mean.
weights
reference A data frame or variable from which the centrality and deviation will be computed instead of from the input variable. Useful for standardizing a subset or new data according to another data frame.

| center | Numeric value, which can be used as alternative to reference to define a refer- <br> ence centrality. If center is of length 1, it will be recycled to match the length <br> of selected variables for centering. Else, center must be of same length as the <br> number of selected variables. Values in center will be matched to selected vari- <br> ables in the provided order, unless a named vector is given. In this case, names <br> are matched against the names of the selected variables. |
| :--- | :--- |
| verbose | Toggle warnings and messages. <br> select |
|  | Variables that will be included when performing the required tasks. Can be <br> either |

- a variable specified as a literal variable name (e.g., column_name),
- a string with the variable name (e.g., "column_name"), or a character vector of variable names (e.g., c("col1", "col2", "col3")),
- a formula with variable names (e.g., $\sim$ column_1 + column_2),
- a vector of positive integers, giving the positions counting from the left (e.g. 1 or $c(1,3,5)$ ),
- a vector of negative integers, giving the positions counting from the right (e.g., -1 or $-1:-3$ ),
- one of the following select-helpers: starts_with(""), ends_with(""), contains(""), a range using : or regex(""),
- or a function testing for logical conditions, e.g. is.numeric() (or is .numeric), or any user-defined function that selects the variables for which the function returns TRUE (like: foo <- function(x) mean (x) > 3),
- ranges specified via literal variable names, select-helpers (except regex()) and (user-defined) functions can be negated, i.e. return non-matching elements, when prefixed with a -, e.g. -ends_with(""), -is.numeric or -Sepal.Width:Petal.Length. Note: Negation means that matches are excluded, and thus, the exclude argument can be used alternatively. For instance, select=-ends_with("Length") (with -) is equivalent to exclude=ends_with("Length") (no -). In case negation should not work as expected, use the exclude argument instead.
If NULL, selects all columns. Patterns that found no matches are silently ignored, e.g. find_columns(iris, select $=c($ "Species", "Test")) will just return "Species".
exclude See select, however, column names matched by the pattern from exclude will be excluded instead of selected. If NULL (the default), excludes no columns.
force Logical, if TRUE, forces centering of factors as well. Factors are converted to numerical values, with the lowest level being the value 1 (unless the factor has numeric levels, which are converted to the corresponding numeric value).
remove_na How should missing values (NA) be treated: if "none" (default): each column's standardization is done separately, ignoring NAs. Else, rows with NA in the columns selected with select / exclude ("selected") or in all columns ("all") are dropped before standardization, and the resulting data frame does not include these cases.
append Logical or string. If TRUE, centered variables get new column names (with the suffix "_c") and are appended (column bind) to $x$, thus returning both the orig-
inal and the centered variables. If FALSE, original variables in $x$ will be overwritten by their centered versions. If a character value, centered variables are appended with new column names (using the defined suffix) to the original data frame.
ignore_case Logical, if TRUE and when one of the select-helpers or a regular expression is used in select, ignores lower/upper case in the search pattern when matching against variable names.


## Value

The centered variables.

## Selection of variables - the select argument

For most functions that have a select argument (including this function), the complete input data frame is returned, even when select only selects a range of variables. That is, the function is only applied to those variables that have a match in select, while all other variables remain unchanged. In other words: for this function, select will not omit any non-included variables, so that the returned data frame will include all variables from the input data frame.

## Note

Difference between centering and standardizing: Standardized variables are computed by subtracting the mean of the variable and then dividing it by the standard deviation, while centering variables involves only the subtraction.

## See Also

If centering within-clusters (instead of grand-mean centering) is required, see demean(). For standardizing, see standardize().

## Examples

```
data(iris)
# entire data frame or a vector
head(iris$Sepal.Width)
head(center(iris$Sepal.Width))
head(center(iris))
head(center(iris, force = TRUE))
# only the selected columns from a data frame
center(anscombe, select = c("x1", "x3"))
center(anscombe, exclude = c("x1", "x3"))
# centering with reference center and scale
d <- data.frame(
    a = c(-2, -1, 0, 1, 2),
    b = c(3, 4, 5, 6, 7)
)
```

```
    # default centering at mean
    center(d)
    # centering, using 0 as mean
    center(d, center = 0)
    # centering, using -5 as mean
    center(d, center = -5)
```

    change_code
        Recode old values of variables into new values
    
## Description

This functions recodes old values into new values and can be used to to recode numeric or character vectors, or factors.

## Usage

```
change_code(x, ...)
data_recode(x, ...)
## S3 method for class 'numeric'
change_code(
        x,
        recode = NULL,
        default = NULL,
        preserve_na = TRUE,
        verbose = TRUE,
    )
    ## S3 method for class 'data.frame'
    change_code(
        x,
        select = NULL,
        exclude = NULL,
        recode = NULL,
        default = NULL,
        preserve_na = TRUE,
        append = FALSE,
        ignore_case = FALSE,
        verbose = TRUE,
    )
```


## Arguments

x
recode
default
preserve_na
verbose
select
exclude See select, however, column names matched by the pattern from exclude will be excluded instead of selected. If NULL (the default), excludes no columns.
append Logical or string. If TRUE, recoded or converted variables get new column names and are appended (column bind) to $x$, thus returning both the original and the recoded variables. The new columns get a suffix, based on the calling function: "_r" for recode functions, "_n" for to_numeric(), "_f" for to_factor(), or "_s" for slide(). If append=FALSE, original variables in $x$ will be overwritten by their recoded versions. If a character value, recoded variables are appended with new column names (using the defined suffix) to the original data frame.
ignore_case Logical, if TRUE and when one of the select-helpers or a regular expression is used in select, ignores lower/upper case in the search pattern when matching against variable names.

## Details

This section describes the pattern of the recode arguments, which also provides some shortcuts, in particular when recoding numeric values.

- Single values

Single values either need to be wrapped in backticks (in case of numeric values) or "as is" (for character or factor levels). Example: recode=list ( $0^{`}=1, ` 1 `=2$ ) would recode 1 into 0 , and 2 into 1 . For factors or character vectors, an example is: recode=list ( $x=" a ", y=" b "$ ) (recode "a" into "x" and "b" into "y").

- Multiple values

Multiple values that should be recoded into a new value can be separated with comma. Example: recode $=1 \operatorname{ist}\left(` 1 `=c(1,4), 2^{`}=c(2,3)\right.$ ) would recode the values 1 and 4 into 1 , and 2 and 3 into 2 . It is also possible to define the old values as a character string, like: recode=list (`1`="1,4", `2`="2,3") For factors or character vectors, an example is: recode=list (x=c("a", "b"),

- Value range

Numeric value ranges can be defined using the : . Example: recode=list( ${ }^{\prime} 1 `=1: 3$, ${ }^{\prime} 2^{`}=4: 6$ ) would recode all values from 1 to 3 into 1 , and 4 to 6 into 2 .

- min and max
placeholder to use the minimum or maximum value of the (numeric) variable. Useful, e.g., when recoding ranges of values. Example: recode=list(`1`="min:10", `2`="11:max").
- default values

The default argument defines the default value for all values that have no match in the recodepairs. For example, recode=list( $\left.` 1 `=c(1,2), `{ }^{`}=c(3,4)\right)$, default=9 would recode values 1 and 2 into 1,3 and 4 into 2 , and all other values into 9 . If preserve_na is set to FALSE, NA (missing values) will also be recoded into the specified default value.

- Reversing and rescaling

See reverse() and rescale().

## Value

$x$, where old values are replaced by new values.

## Selection of variables - the select argument

For most functions that have a select argument (including this function), the complete input data frame is returned, even when select only selects a range of variables. That is, the function is only applied to those variables that have a match in select, while all other variables remain unchanged. In other words: for this function, select will not omit any non-included variables, so that the returned data frame will include all variables from the input data frame.

## Note

You can use options(data_recode_pattern = "old=new") to switch the behaviour of the recodeargument, i.e. recode-pairs are now following the pattern old values $=$ new values, e.g. if getOption("data_recode_pattern") is set to "old=new", then recode(' 1 '=0) would recode all 1 into 0 . The default for recode $(` 1 `=0)$ is to recode all 0 into 1 .

## See Also

- Functions to rename stuff: data_rename(), data_rename_rows(), data_addprefix(), data_addsuffix()
- Functions to reorder or remove columns: data_reorder(), data_relocate(), data_remove()
- Functions to reshape, pivot or rotate data frames: data_to_long(), data_to_wide(), data_rotate()
- Functions to recode data: rescale(), reverse(), categorize(), change_code(), slide()
- Functions to standardize, normalize, rank-transform: center(), standardize(), normalize(), ranktransform(), winsorize()
- Split and merge data frames: data_partition(), data_merge()
- Functions to find or select columns: data_select(), data_find()
- Functions to filter rows: data_match(), data_filter()


## Examples

```
# numeric ----------
set.seed(123)
x <- sample(c(1:4, NA), 15, TRUE)
table(x, useNA = "always")
out <- change_code(x, list(`0` = 1, `1` = 2:3, `2` = 4))
out
table(out, useNA = "always")
# to recode NA values, set preserve_na to FALSE
out <- change_code(
    x,
    list(`0` = 1, `1` = 2:3, `2`= 4, `9` = NA),
    preserve_na = FALSE
)
out
table(out, useNA = "always")
# preserve na ---------
out <- change_code(x, list(`0` = 1, `1` = 2:3), default = 77)
```

```
out
table(out, useNA = "always")
# recode na into default ----------
out <- change_code(
        x,
        list(`0` = 1, `1` = 2:3),
    default = 77,
    preserve_na = FALSE
)
out
table(out, useNA = "always")
# factors (character vectors are similar) ----------
set.seed(123)
x <- as.factor(sample(c("a", "b", "c"), 15, TRUE))
table(x)
out <- change_code(x, list(x = "a", y = c("b", "c")))
out
table(out)
out <- change_code(x, list(x = "a", y = "b", z = "c"))
out
table(out)
out <- change_code(x, list(y = "b,c"), default = 77)
# same as
# change_code(x, list(y = c("b", "c")), default = 77)
out
table(out)
# data frames ----------
set.seed(123)
d <- data.frame(
    x = sample(c(1:4,NA), 12, TRUE),
    y = as.factor(sample(c("a", "b", "c"), 12, TRUE)),
    stringsAsFactors = FALSE
)
change_code(
    d,
    recode = list(`0` = 1, `1` = 2:3, `2` = 4, x = "a", y = c("b", "c")),
    append = TRUE
)
# switch recode pattern to "old=new" -
options(data_recode_pattern = "old=new")
# numeric
```

```
set.seed(123)
x <- sample(c(1:4, NA), 15, TRUE)
table(x, useNA = "always")
out <- change_code(x, list(`1` = 0, `2:3` = 1, `4` = 2))
table(out, useNA = "always")
# factors (character vectors are similar)
set.seed(123)
x <- as.factor(sample(c("a", "b", "c"), 15, TRUE))
table(x)
out <- change_code(x, list(a = "x", `b, c` = "y"))
table(out)
# reset options
options(data_recode_pattern = NULL)
```

coerce_to_numeric Convert to Numeric (if possible)

## Description

Tries to convert vector to numeric if possible (if no warnings or errors). Otherwise, leaves it as is.

## Usage

```
coerce_to_numeric(x)
```


## Arguments

$x \quad$ A vector to be converted.

## Value

Numeric vector (if possible)

## Examples

```
coerce_to_numeric(c("1", "2"))
coerce_to_numeric(c("1", "2", "A"))
```

```
convert_na_to
Replace missing values in a variable or a data frame.
```


## Description

Replace missing values in a variable or a data frame.

## Usage

```
convert_na_to(x, ...)
## S3 method for class 'numeric'
convert_na_to(x, replacement = NULL, verbose = TRUE, ...)
## S3 method for class 'character'
convert_na_to(x, replacement = NULL, verbose = TRUE, ...)
    ## S3 method for class 'data.frame'
    convert_na_to(
        x,
    select = NULL,
    exclude = NULL,
    replacement = NULL,
    replace_num = replacement,
    replace_char = replacement,
    replace_fac = replacement,
    ignore_case = FALSE,
    verbose = TRUE,
    ..
    )
```


## Arguments

x
...
replacement Numeric or character value that will be used to replace NA.
verbose
select Variables that will be included when performing the required tasks. Can be either

- a variable specified as a literal variable name (e.g., column_name),
- a string with the variable name (e.g., "column_name"), or a character vector of variable names (e.g., c("col1", "col2", "col3")),
- a formula with variable names (e.g., $\sim$ column_1 + column_2),
- a vector of positive integers, giving the positions counting from the left (e.g. 1 or $c(1,3,5)$ ),
- a vector of negative integers, giving the positions counting from the right (e.g., -1 or $-1:-3$ ),
- one of the following select-helpers: starts_with(""), ends_with(""), contains(""), a range using : or regex(""),
- or a function testing for logical conditions, e.g. is.numeric() (or is.numeric), or any user-defined function that selects the variables for which the function returns TRUE (like: foo <- function $(x)$ mean $(x)>3$ ),
- ranges specified via literal variable names, select-helpers (except regex()) and (user-defined) functions can be negated, i.e. return non-matching elements, when prefixed with a -, e.g. -ends_with(""), -is.numeric or -Sepal.Width:Petal.Length. Note: Negation means that matches are excluded, and thus, the exclude argument can be used alternatively. For instance, select=-ends_with("Length") (with -) is equivalent to exclude=ends_with("Length") (no -). In case negation should not work as expected, use the exclude argument instead.
If NULL, selects all columns. Patterns that found no matches are silently ignored, e.g. find_columns(iris, select $=c($ "Species", "Test")) will just return "Species".
exclude See select, however, column names matched by the pattern from exclude will be excluded instead of selected. If NULL (the default), excludes no columns.
replace_num Value to replace NA when variable is of type numeric.
replace_char Value to replace NA when variable is of type character.
replace_fac Value to replace NA when variable is of type factor.
ignore_case Logical, if TRUE and when one of the select-helpers or a regular expression is used in select, ignores lower/upper case in the search pattern when matching against variable names.


## Value

$x$, where NA values are replaced by replacement.

## Selection of variables - the select argument

For most functions that have a select argument (including this function), the complete input data frame is returned, even when select only selects a range of variables. That is, the function is only applied to those variables that have a match in select, while all other variables remain unchanged. In other words: for this function, select will not omit any non-included variables, so that the returned data frame will include all variables from the input data frame.

## Examples

```
# Convert NA to 0 in a numeric vector
convert_na_to(
    c(9, 3, NA, 2, 3, 1, NA, 8),
    replacement = 0
)
```

```
# Convert NA to "missing" in a character vector
convert_na_to(
    c("a", NA, "d", "z", NA, "t"),
    replacement = "missing"
)
### For data frames
test_df <- data.frame(
    x = c(1, 2,NA),
    x2 = c(4, 5, NA),
    y = c("a", "b",NA)
)
# Convert all NA to 0 in numeric variables, and all NA to "missing" in
# character variables
convert_na_to(
    test_df,
    replace_num = 0,
    replace_char = "missing"
)
# Convert a specific variable in the data frame
convert_na_to(
    test_df,
    replace_num = 0,
    replace_char = "missing",
    select = "x"
)
# Convert all variables starting with "x"
convert_na_to(
    test_df,
    replace_num = 0,
    replace_char = "missing",
    select = starts_with("x")
)
# Convert NA to 1 in variable 'x2' and to 0 in all other numeric
# variables
convert_na_to(
    test_df,
    replace_num = 0,
    select = list(x2 = 1)
)
```


## Description

Convert non-missing values in a variable into missing values.

```
Usage
    convert_to_na(x, ...)
    ## S3 method for class 'numeric'
    convert_to_na(x, na = NULL, verbose = TRUE, ...)
    ## S3 method for class 'factor'
    convert_to_na(x, na = NULL, drop_levels = FALSE, verbose = TRUE, ...)
    ## S3 method for class 'data.frame'
    convert_to_na(
        x,
        select = NULL,
        exclude = NULL,
        na = NULL,
        drop_levels = FALSE,
        ignore_case = FALSE,
        verbose = TRUE,
    )
```


## Arguments

x
... Not used.
na Numeric, character vector or logical (or a list of numeric, character vectors or logicals) with values that should be converted to NA. Numeric values applied to numeric vectors, character values are used for factors, character vectors or date variables, and logical values for logical vectors.
verbose Toggle warnings.
drop_levels Logical, for factors, when specific levels are replaced by NA, should unused levels be dropped?
select Variables that will be included when performing the required tasks. Can be either

- a variable specified as a literal variable name (e.g., column_name),
- a string with the variable name (e.g., "column_name"), or a character vector of variable names (e.g., c("col1", "col2", "col3")),
- a formula with variable names (e.g., ~column_1 + column_2),
- a vector of positive integers, giving the positions counting from the left (e.g. 1 or c $(1,3,5)$ ),
- a vector of negative integers, giving the positions counting from the right (e.g., -1 or $-1:-3$ ),
- one of the following select-helpers: starts_with(""), ends_with(""), contains(""), a range using : or regex(""),
- or a function testing for logical conditions, e.g. is.numeric() (or is.numeric), or any user-defined function that selects the variables for which the function returns TRUE (like: foo <- function(x) mean (x) > 3),
- ranges specified via literal variable names, select-helpers (except regex()) and (user-defined) functions can be negated, i.e. return non-matching elements, when prefixed with a -, e.g. -ends_with(""), -is.numeric or -Sepal.Width:Petal.Length. Note: Negation means that matches are excluded, and thus, the exclude argument can be used alternatively. For instance, select=-ends_with("Length") (with -) is equivalent to exclude=ends_with("Length") (no -). In case negation should not work as expected, use the exclude argument instead.
If NULL, selects all columns. Patterns that found no matches are silently ignored, e.g. find_columns(iris, select = c("Species", "Test")) will just return "Species".
exclude See select, however, column names matched by the pattern from exclude will be excluded instead of selected. If NULL (the default), excludes no columns.
ignore_case Logical, if TRUE and when one of the select-helpers or a regular expression is used in select, ignores lower/upper case in the search pattern when matching against variable names.


## Value

$x$, where all values in na are converted to NA.

## Examples

```
x <- sample(1:6, size = 30, replace = TRUE)
x
# values 4 and 5 to NA
convert_to_na(x, na = 4:5)
# data frames
set.seed(123)
x <- data.frame(
    a = sample(1:6, size = 20, replace = TRUE),
    b = sample(letters[1:6], size = 20, replace = TRUE),
    c = sample(c(30:33, 99), size = 20, replace = TRUE)
)
# for all numerics, convert 5 to NA. Character/factor will be ignored.
convert_to_na(x, na = 5)
# for numerics, 5 to NA, for character/factor, "f" to NA
convert_to_na(x, na = list(6, "f"))
# select specific variables
convert_to_na(x, select = c("a", "b"), na = list(6, "f"))
```


## Description

Safe and intuitive functions to rename variables or rows in data frames. data_rename() will rename column names, i.e. it facilitates renaming variables data_addprefix() or data_addsuffix() add prefixes or suffixes to column names. data_rename_rows() is a convenient shortcut to add or rename row names of a data frame, but unlike row. names(), its input and output is a data frame, thus, integrating smoothly into a possible pipe-workflow.

```
Usage
    data_addprefix(
        data,
        pattern,
        select = NULL,
        exclude = NULL,
        ignore_case = FALSE,
        ...
    )
    data_addsuffix(
        data,
        pattern,
        select = NULL,
        exclude = NULL,
        ignore_case = FALSE,
    )
```

    data_rename(data, pattern \(=\) NULL, replacement \(=\) NULL, safe \(=\) TRUE, ...)
    data_rename_rows(data, rows = NULL)
    
## Arguments

$$
\begin{array}{ll}
\text { data } & \text { A data frame, or an object that can be coerced to a data frame. } \\
\text { pattern } & \begin{array}{l}
\text { Character vector. For data_rename(), indicates columns that should be se- } \\
\text { lected for renaming. Can be NULL (in which case all columns are selected). For } \\
\text { data_addprefix() or data_addsuffix(), a character string, which will be } \\
\text { added as prefix or suffix to the column names. }
\end{array} \\
\text { select } & \begin{array}{l}
\text { Variables that will be included when performing the required tasks. Can be } \\
\text { either }
\end{array}
\end{array}
$$

- a variable specified as a literal variable name (e.g., column_name),
- a string with the variable name (e.g., "column_name"), or a character vector of variable names (e.g., c("col1", "col2", "col3")),
- a formula with variable names (e.g., $\sim$ column_1 + column_2),
- a vector of positive integers, giving the positions counting from the left (e.g. 1 or $c(1,3,5)$ ),
- a vector of negative integers, giving the positions counting from the right (e.g., -1 or $-1:-3$ ),
- one of the following select-helpers: starts_with(""), ends_with(""), contains(""), a range using : or regex (""),
- or a function testing for logical conditions, e.g. is.numeric() (or is .numeric), or any user-defined function that selects the variables for which the function returns TRUE (like: foo <- function(x) mean (x) > 3),
- ranges specified via literal variable names, select-helpers (except regex ()) and (user-defined) functions can be negated, i.e. return non-matching elements, when prefixed with a -, e.g. -ends_with(""), -is.numeric or -Sepal.Width:Petal.Length. Note: Negation means that matches are excluded, and thus, the exclude argument can be used alternatively. For instance, select=-ends_with("Length") (with -) is equivalent to exclude=ends_with("Length") (no -). In case negation should not work as expected, use the exclude argument instead.
If NULL, selects all columns. Patterns that found no matches are silently ignored, e.g. find_columns(iris, select $=c(" S p e c i e s ", ~ " T e s t ")) ~ w i l l ~ j u s t ~ r e t u r n ~$ "Species".
exclude See select, however, column names matched by the pattern from exclude will be excluded instead of selected. If NULL (the default), excludes no columns.
ignore_case Logical, if TRUE and when one of the select-helpers or a regular expression is used in select, ignores lower/upper case in the search pattern when matching against variable names.
... Other arguments passed to or from other functions.
replacement Character vector. Indicates the new name of the columns selected in pattern. Can be NULL (in which case column are numbered in sequential order). If not NULL, pattern and replacement must be of the same length.
safe Do not throw error if for instance the variable to be renamed/removed doesn't exist.
rows Vector of row names.


## Value

A modified data frame.

## See Also

- Functions to rename stuff: data_rename(), data_rename_rows(), data_addprefix(), data_addsuffix()
- Functions to reorder or remove columns: data_reorder(), data_relocate(), data_remove()
- Functions to reshape, pivot or rotate data frames: data_to_long(), data_to_wide(), data_rotate()
- Functions to recode data: rescale(), reverse(), categorize(), change_code(), slide()
- Functions to standardize, normalize, rank-transform: center(), standardize(), normalize(), ranktransform(), winsorize()
- Split and merge data frames: data_partition(), data_merge()
- Functions to find or select columns: data_select(), data_find()
- Functions to filter rows: data_match(), data_filter()


## Examples

```
    # Add prefix / suffix to all columns
    head(data_addprefix(iris, "NEW_"))
    head(data_addsuffix(iris, "_OLD"))
    # Rename columns
    head(data_rename(iris, "Sepal.Length", "length"))
    # data_rename(iris, "FakeCol", "length", safe=FALSE) # This fails
    head(data_rename(iris, "FakeCol", "length")) # This doesn't
    head(data_rename(iris, c("Sepal.Length", "Sepal.Width"), c("length", "width")))
    # Reset names
    head(data_rename(iris, NULL))
    # Change all
    head(data_rename(iris, paste0("Var", 1:5)))
```

    data_extract Extract one or more columns or elements from an object
    
## Description

data_extract() (or its alias extract()) is similar to \$. It extracts either a single column or element from an object (e.g., a data frame, list), or multiple columns resp. elements.

## Usage

```
data_extract(data, select, ...)
```

\#\# S3 method for class 'data.frame'
data_extract(
data,
select,
name = NULL,
extract = "all",
as_data_frame = FALSE,
ignore_case = FALSE,
verbose = TRUE,
)

## Arguments

data
select Variables that will be included when performing the required tasks. Can be either

- a variable specified as a literal variable name (e.g., column_name),
- a string with the variable name (e.g., "column_name"), or a character vector of variable names (e.g., c("col1", "col2", "col3")),
- a formula with variable names (e.g., ~column_1 + column_2),
- a vector of positive integers, giving the positions counting from the left (e.g. 1 or $c(1,3,5)$ ),
- a vector of negative integers, giving the positions counting from the right (e.g., -1 or $-1:-3$ ),
- one of the following select-helpers: starts_with(""), ends_with(""), contains(""), a range using : or regex(""),
- or a function testing for logical conditions, e.g. is.numeric() (or is.numeric), or any user-defined function that selects the variables for which the function returns TRUE (like: foo <- function(x) mean (x) > 3),
- ranges specified via literal variable names, select-helpers (except regex ()) and (user-defined) functions can be negated, i.e. return non-matching elements, when prefixed with a -, e.g. -ends_with(""), -is.numeric or -Sepal.Width:Petal.Length. Note: Negation means that matches are excluded, and thus, the exclude argument can be used alternatively. For instance, select=-ends_with("Length") (with -) is equivalent to exclude=ends_with("Length") (no -). In case negation should not work as expected, use the exclude argument instead.
If NULL, selects all columns. Patterns that found no matches are silently ignored, e.g. find_columns(iris, select $=c($ "Species", "Test")) will just return "Species".
$\ldots \quad$ For use by future methods.
name An optional argument that specifies the column to be used as names for the vector elements after extraction. Must be specified either as literal variable name (e.g., column_name) or as string ("column_name"). name will be ignored when a data frame is returned.
extract String, indicating which element will be extracted when select matches multiple variables. Can be "all" (the default) to return all matched variables, "first" or "last" to return the first or last match, or "odd" and "even" to return all odd-numbered or even-numbered matches. Note that "first" or "last" return a vector (unless as_data_frame = TRUE), while "all" can return a vector (if only one match was found) or a data frame (for more than one match). Type safe return values are only possible when extract is "first" or "last" (will always return a vector) or when as_data_frame = TRUE (always returns a data frame).
as_data_frame Logical, if TRUE, will always return a data frame, even if only one variable was matched. If FALSE, either returns a vector or a data frame. See extract for details.
ignore_case Logical, if TRUE and when one of the select-helpers or a regular expression is used in select, ignores lower/upper case in the search pattern when matching against variable names.
verbose Toggle warnings.


## Details

data_extract () can be used to select multiple variables or pull a single variable from a data frame. Thus, the return value is by default not type safe - data_extract() either returns a vector or a data frame.

Extracting single variables (vectors): When select is the name of a single column, or when select only matches one column, a vector is returned. A single variable is also returned when extract is either "first or "last". Setting as_data_frame to TRUE overrides this behaviour and always returns a data frame.

Extracting a data frame of variables: When select is a character vector containing more than one column name (or a numeric vector with more than one valid column indices), or when select uses one of the supported select-helpers that match multiple columns, a data frame is returned. Setting as_data_frame to TRUE always returns a data frame.

## Value

A vector (or a data frame) containing the extracted element, or NULL if no matching variable was found.

## Examples

```
# single variable
data_extract(mtcars, cyl, name = gear)
data_extract(mtcars, "cyl", name = gear)
data_extract(mtcars, -1, name = gear)
data_extract(mtcars, cyl, name = 0)
data_extract(mtcars, cyl, name = "row.names")
# selecting multiple variables
head(data_extract(iris, starts_with("Sepal")))
head(data_extract(iris, ends_with("Width")))
head(data_extract(iris, 2:4))
# select first of multiple variables
data_extract(iris, starts_with("Sepal"), extract = "first")
# select first of multiple variables, return as data frame
head(data_extract(iris, starts_with("Sepal"), extract = "first", as_data_frame = TRUE))
```

```
data_group Create a grouped data frame
```


## Description

This function is comparable to dplyr: :group_by(), but just following the datawizard function design. data_ungroup() removes the grouping information from a grouped data frame.

## Usage

data_group( x , select $=$ NULL, exclude = NULL, ignore_case = FALSE, verbose = TRUE,
)
data_ungroup (x, verbose = TRUE, ...)

## Arguments

## x

select Variables that will be included when performing the required tasks. Can be either

- a variable specified as a literal variable name (e.g., column_name),
- a string with the variable name (e.g., "column_name"), or a character vector of variable names (e.g., c("col1", "col2", "col3")),
- a formula with variable names (e.g., ~column_1 + column_2),
- a vector of positive integers, giving the positions counting from the left (e.g. 1 or c(1, 3, 5)),
- a vector of negative integers, giving the positions counting from the right (e.g., -1 or $-1:-3$ ),
- one of the following select-helpers: starts_with(""), ends_with(""), contains(""), a range using : or regex (""),
- or a function testing for logical conditions, e.g. is.numeric() (or is.numeric), or any user-defined function that selects the variables for which the function returns TRUE (like: foo <- function (x) mean (x) > 3),
- ranges specified via literal variable names, select-helpers (except regex ()) and (user-defined) functions can be negated, i.e. return non-matching elements, when prefixed with a -, e.g. -ends_with(""), -is.numeric or -Sepal.Width:Petal.Length. Note: Negation means that matches are excluded, and thus, the exclude argument can be used alternatively. For instance, select=-ends_with("Length") (with -) is equivalent to exclude=ends_with("Length")
(no -). In case negation should not work as expected, use the exclude argument instead.
If NULL, selects all columns. Patterns that found no matches are silently ignored, e.g. find_columns(iris, select $=c($ "Species", "Test")) will just return "Species".
exclude See select, however, column names matched by the pattern from exclude will be excluded instead of selected. If NULL (the default), excludes no columns.
ignore_case Logical, if TRUE and when one of the select-helpers or a regular expression is used in select, ignores lower/upper case in the search pattern when matching against variable names.
verbose Toggle warnings.
... Arguments passed down to other functions. Mostly not used yet.


## Value

A grouped data frame, i.e. a data frame with additional information about the grouping structure saved as attributes.

## Examples

```
data(efc)
if (requireNamespace("poorman")) {
    suppressPackageStartupMessages(library(poorman, quietly = TRUE))
    # total mean
    efc %>%
        summarize(mean_hours = mean(c12hour, na.rm = TRUE))
    # mean by educational level
    efc %>%
        data_group(c172code) %>%
        summarize(mean_hours = mean(c12hour, na.rm = TRUE))
}
```

data_match Return filtered or sliced data frame, or row indices

## Description

Return a filtered (or sliced) data frame or row indices of a data frame that match a specific condition. data_filter() works like data_match(), but works with logical expressions or row indices of a data frame to specify matching conditions.

## Usage

data_match(x, to, match = "and", return_indices = FALSE, drop_na = TRUE, ...)
data_filter(x, filter, ...)

## Arguments

| X | A data frame. |
| :---: | :---: |
| to | A data frame matching the specified conditions. Note that if match is a value other than "and", the original row order might be changed. See 'Details'. |
| match | String, indicating with which logical operation matching conditions should be combined. Can be "and" (or "\&"), "or" (or "\|") or "not" (or "!"). |
| return_indices | Logical, if FALSE, return the vector of rows that can be used to filter the original data frame. If FALSE (default), returns directly the filtered data frame instead of the row indices. |
| drop_na | Logical, if TRUE, missing values (NAs) are removed before filtering the data. This is the default behaviour, however, sometimes when row indices are requested (i.e. return_indices=TRUE), it might be useful to preserve NA values, so returned row indices match the row indices of the original data frame. |
|  | Not used. |
| filter | A logical expression indicating which rows to keep, or a numeric vector indicating the row indices of rows to keep. |

## Details

For data_match(), if match is either "or" or "not", the original row order from $\times$ might be changed. If preserving row order is required, use data_filter() instead.

```
# mimics subset() behaviour, preserving original row order
head(data_filter(mtcars[c("mpg", "vs", "am")], vs == 0 | am == 1))
#> mpg vs am
#> Mazda RX4 21.0 0 1
#> Mazda RX4 Wag 21.0 0 1
#> Datsun 710 22.8 1 1
#> Hornet Sportabout 18.7 0 0
#> Duster 360 14.3 0 0
#> Merc 450SE 16.4 0 0
# re-sorting rows
head(data_match(mtcars[c("mpg", "vs", "am")],
    data.frame(vs = 0, am = 1),
    match = "or"))
#> mpg vs am
#> Mazda RX4 21.0 0 1
#> Mazda RX4 Wag 21.0 0 1
#> Hornet Sportabout 18.7 0 0
#> Duster 360 14.3 0 0
#> Merc 450SE 16.4 0 0
#> Merc 450SL 17.3 0 0
```

While data_match() works with data frames to match conditions against, data_filter () is basically a wrapper around subset(subset = <filter>). However, unlike subset(), it preserves label attributes and is useful when working with labelled data.

## Value

A filtered data frame, or the row indices that match the specified configuration.

## See Also

- Functions to rename stuff: data_rename(), data_rename_rows(), data_addprefix(), data_addsuffix()
- Functions to reorder or remove columns: data_reorder(), data_relocate(), data_remove()
- Functions to reshape, pivot or rotate data frames: data_to_long(), data_to_wide(), data_rotate()
- Functions to recode data: rescale(), reverse(), categorize(), change_code(), slide()
- Functions to standardize, normalize, rank-transform: center(), standardize(), normalize(), ranktransform(), winsorize()
- Split and merge data frames: data_partition(), data_merge()
- Functions to find or select columns: data_select(), data_find()
- Functions to filter rows: data_match(), data_filter()


## Examples

```
data_match(mtcars, data.frame(vs = 0, am = 1))
data_match(mtcars, data.frame(vs = 0, am = c(0, 1)))
# observations where "vs" is NOT 0 AND "am" is NOT 1
data_match(mtcars, data.frame(vs = 0, am = 1), match = "not")
# equivalent to
data_filter(mtcars, vs != 0 & am != 1)
# observations where EITHER "vs" is 0 OR "am" is 1
data_match(mtcars, data.frame(vs = 0, am = 1), match = "or")
# equivalent to
data_filter(mtcars, vs == 0 | am == 1)
# slice data frame by row indices
data_filter(mtcars, 5:10)
```

data_merge $\quad$ Merge (join) two data frames, or a list of data frames

## Description

Merge (join) two data frames, or a list of data frames. However, unlike base R's merge(), data_merge() offers a few more methods to join data frames, and it does not drop data frame nor column attributes.

## Usage

```
data_merge(x, ...)
data_join(x, ...)
\#\# S3 method for class 'data.frame'
data_merge(x, y, join = "left", by = NULL, id = NULL, verbose = TRUE, ...)
\#\# S3 method for class 'list'
data_merge(x, join = "left", by = NULL, id = NULL, verbose = TRUE, ...)
```


## Arguments

| $x, y$ | A data frame to merge. $x$ may also be a list of data frames that will be merged. |
| :--- | :--- |
| Note that the list-method has no y argument. |  |
| $\ldots$ | Not used. |
| join | Character vector, indicating the method of joining the data frames. Can be <br>  <br> "full", "left" (default), "right", "inner", "anti", "semi" or "bind". See <br> details below. |
| by | Specifications of the columns used for merging. |
| id | Optional name for ID column that will be created to indicate the source data <br> frames for appended rows. Only applies if join = "bind". <br> verbose |
|  | Toggle warnings. |

## Details

Merging data frames: Merging data frames is performed by adding rows (cases), columns (variables) or both from the source data frame (y) to the target data frame (x). This usually requires one or more variables which are included in both data frames and that are used for merging, typically indicated with the by argument. When by contains a variable present in both data frames, cases are matched and filtered by identical values of by in $x$ and $y$.

Left- and right-joins: Left- and right joins usually don't add new rows (cases), but only new columns (variables) for existing cases in $x$. For join = "left" or join = "right" to work, by must indicate one or more columns that are included in both data frames. For join = "left", if by is an identifier variable, which is included in both $x$ and $y$, all variables from $y$ are copied to $x$, but only those cases from $y$ that have matching values in their identifier variable in $x$ (i.e. all cases in $x$ that are also found in $y$ get the related values from the new columns in $y$ ). If there is no match between identifiers in $x$ and $y$, the copied variable from $y$ will get a NA value for this particular case. Other variables that occur both in $x$ and $y$, but are not used as identifiers (with by), will be renamed to avoid multiple identical variable names. Cases in $y$ where values from the identifier have no match in x's identifier are removed. join = "right" works in a similar way as join = "left", just that only cases from $x$ that have matching values in their identifier variable in $y$ are chosen.

In base $R$, these are equivalent to merge $(x, y$, all. $x=\operatorname{TRUE})$ and merge $(x, y, a l l . y=$ TRUE $)$.

Full joins: Full joins copy all cases from y to $x$. For matching cases in both data frames, values for new variables are copied from $y$ to $x$. For cases in $y$ not present in $x$, these will be added as new rows to $x$. Thus, full joins not only add new columns (variables), but also might add new rows (cases).

In base $R$, this is equivalent to merge $(x, y, a l l=T R U E)$.
Inner joins: Inner joins merge two data frames, however, only those rows (cases) are kept that are present in both data frames. Thus, inner joins usually add new columns (variables), but also remove rows (cases) that only occur in one data frame.

In base $R$, this is equivalent to merge $(x, y)$.
Binds: join = "bind" row-binds the complete second data frame $y$ to $x$. Unlike simple rbind(), which requires the same columns for both data frames, join = "bind" will bind shared columns from y to x , and add new columns from y to x .

## Value

A merged data frame.

## See Also

- Functions to rename stuff: data_rename(), data_rename_rows(), data_addprefix(), data_addsuffix()
- Functions to reorder or remove columns: data_reorder(), data_relocate(), data_remove()
- Functions to reshape, pivot or rotate data frames: data_to_long(), data_to_wide(), data_rotate()
- Functions to recode data: rescale(), reverse(), categorize(), change_code(), slide()
- Functions to standardize, normalize, rank-transform: center(), standardize(), normalize(), ranktransform(), winsorize()
- Split and merge data frames: data_partition(), data_merge()
- Functions to find or select columns: data_select(), data_find()
- Functions to filter rows: data_match(), data_filter()


## Examples

```
x <- data.frame(a = 1:3, b = c("a", "b", "c"), c = 5:7, id = 1:3)
y <- data.frame(c = 6:8, d = c("f", "g", "h"), e = 100:102, id = 2:4)
X
y
# "by" will default to all shared columns, i.e. "c" and "id". new columns
# "d" and "e" will be copied from "y" to "x", but there are only two cases
# in "x" that have the same values for "c" and "id" in "y". only those cases
# have values in the copied columns, the other case gets "NA".
data_merge(x, y, join = "left")
```

```
# we change the id-value here
x <- data.frame(a = 1:3, b = c("a", "b", "c"), c = 5:7, id = 1:3)
y <- data.frame(c = 6:8, d = c("f", "g", "h"), e = 100:102, id = 3:5)
x
y
# no cases in "y" have the same matching "c" and "id" as in "x", thus
# copied variables from "y" to "x" copy no values, all get NA.
data_merge(x, y, join = "left")
# one case in "y" has a match in "id" with "x", thus values for this
# case from the remaining variables in "y" are copied to "x", all other
# values (cases) in those remaining variables get NA
data_merge(x, y, join = "left", by = "id")
data(mtcars)
x <- mtcars[1:5, 1:3]
y <- mtcars[28:32, 4:6]
# add ID common column
x$id <- 1:5
y$id <- 3:7
# left-join, add new variables and copy values from y to x,
# where "id" values match
data_merge(x, y)
# right-join, add new variables and copy values from x to y,
# where "id" values match
data_merge(x, y, join = "right")
# full-join
data_merge(x, y, join = "full")
data(mtcars)
x <- mtcars[1:5, 1:3]
y <- mtcars[28:32, c(1, 4:5)]
# add ID common column
x$id <- 1:5
y$id <- 3:7
# left-join, no matching rows (because columns "id" and "disp" are used)
# new variables get all NA values
data_merge(x, y)
# one common value in "mpg", so one row from y is copied to x
data_merge(x, y, by = "mpg")
# only keep rows with matching values in by-column
data_merge(x, y, join = "semi", by = "mpg")
```

```
# only keep rows with non-matching values in by-column
data_merge(x, y, join = "anti", by = "mpg")
# merge list of data frames. can be of different rows
x <- mtcars[1:5, 1:3]
y <- mtcars[28:31, 3:5]
z <- mtcars[11:18, c(1, 3:4, 6:8)]
x$id <- 1:5
y$id <- 4:7
z$id <- 3:10
data_merge(list(x, y, z), join = "bind", by = "id", id = "source")
```

data_partition Partition data

## Description

Creates data partitions (for instance, a training and a test set) based on a data frame that can also be stratified (i.e., evenly spread a given factor) using the group argument.

## Usage

```
data_partition(
        data,
        proportion = 0.7,
        group = NULL,
        seed = NULL,
        row_id = ".row_id",
        verbose = TRUE,
        training_proportion = proportion,
    )
```


## Arguments

| data | A data frame, or an object that can be coerced to a data frame. |
| :--- | :--- |
| proportion | Scalar (between 0 and 1) or numeric vector, indicating the proportion(s) of the <br> training set(s). The sum of proportion must not be greater than 1. The remain- <br> ing part will be used for the test set. |
| group | A character vector indicating the name(s) of the column(s) used for stratified <br> partitioning. |
| seed | A random number generator seed. Enter an integer (e.g. 123) so that the random <br> sampling will be the same each time you run the function. |
| row_id | Character string, indicating the name of the column that contains the row-id's. |
| verbose | Toggle messages and warnings. |

training_proportion
Deprecated, please use proportion.
... Other arguments passed to or from other functions.

## Value

A list of data frames. The list includes one training set per given proportion and the remaining data as test set. List elements of training sets are named after the given proportions (e.g., \$p_0.7), the test set is named \$test.

## See Also

- Functions to rename stuff: data_rename(), data_rename_rows(), data_addprefix(), data_addsuffix()
- Functions to reorder or remove columns: data_reorder(), data_relocate(), data_remove()
- Functions to reshape, pivot or rotate data frames: data_to_long(), data_to_wide(), data_rotate()
- Functions to recode data: rescale(), reverse(), categorize(), change_code(), slide()
- Functions to standardize, normalize, rank-transform: center(), standardize(), normalize(), ranktransform(), winsorize()
- Split and merge data frames: data_partition(), data_merge()
- Functions to find or select columns: data_select(), data_find()
- Functions to filter rows: data_match(), data_filter()


## Examples

```
data(iris)
out <- data_partition(iris, proportion = 0.9)
out$test
nrow(out$p_0.9)
# Stratify by group (equal proportions of each species)
out <- data_partition(iris, proportion = 0.9, group = "Species")
out$test
# Create multiple partitions
out <- data_partition(iris, proportion = c(0.3, 0.3))
lapply(out, head)
# Create multiple partitions, stratified by group - 30% equally sampled
# from species in first training set, 50% in second training set and
# remaining 20% equally sampled from each species in test set.
out <- data_partition(iris, proportion = c(0.3, 0.5), group = "Species")
lapply(out, function(i) table(i$Species))
```


## Description

This functions imports data from various file types. It is a small wrapper around haven: : read_spss(), haven:: read_stata(), haven: :read_sas(), readxl: :read_excel() and data.table::fread() resp. readr: : read_delim() (the latter if package data.table is not installed). Thus, supported file types for importing data are data files from SPSS, SAS or Stata, Excel files or text files (like '.csv' files). All non-supported file types are passed to rio::import().

## Usage

```
    data_read(path, path_catalog = NULL, encoding = NULL, verbose = TRUE, ...)
```


## Arguments

path Character string, the file path to the data file.
path_catalog Character string, path to the catalog file. Only relevant for SAS data files.
encoding The character encoding used for the file. Usually not needed.
verbose Toggle warnings and messages.
... Arguments passed to the related read_*() function.

## Value

A data frame.

## Supported file types

data_read() is a wrapper around the haven, data.table, readr readxl and rio packages. Currently supported file types are .txt, .csv, .xls, .xlsx, .sav, .por, .dta and .sas (and related files). All other file types are passed to rio: :import().

## Compressed files (zip) and URLs

data_read() can also read the above mentioned files from URLs or from inside zip-compressed files. Thus, path can also be a URL to a file like "http://www. url.com/file.csv". When path points to a zip-compressed file, and there are multiple files inside the zip-archive, then the first supported file is extracted and loaded.

## General behaviour

data_read() detects the appropriate read_*() function based on the file-extension of the data file. Thus, in most cases it should be enough to only specify the path argument. However, if more control is needed, all arguments in . . . are passed down to the related read_*() function.

## Differences to other packages that read foreign data formats

data_read() is most comparable to rio: :import(). For data files from SPSS, SAS or Stata, which support labelled data, variables are converted into their most appropriate type. The major difference to rio: :import() is that data_read() automatically converts variables into factors, unless the variables are only partially labelled, in which case variables are converted to numerics. Character vectors are preserved. Hence, variables, where all values are labelled, will be converted into factors, where imported value labels will be set as factor levels. Else, if a variable has no value labels or less value labels than values, the variable is either converted into numeric or character vector. Value labels are then preserved as "labels" attribute.

```
data_relocate Relocate (reorder) columns of a data frame
```


## Description

data_relocate() will reorder columns to specific positions, indicated by before or after. data_reorder() will instead move selected columns to the beginning of a data frame. Finally, data_remove() removes columns from a data frame. All functions support select-helpers that allow flexible specification of a search pattern to find matching columns, which should be reordered or removed.

## Usage

```
    data_relocate(
        data,
        select,
        before = NULL,
        after = NULL,
        ignore_case = FALSE,
        verbose = TRUE,
    )
    data_reorder(data, select, ignore_case = FALSE, verbose = TRUE, ...)
    data_remove(data, select, ignore_case = FALSE, verbose = FALSE, ...)
```


## Arguments

data A data frame.
select Variables that will be included when performing the required tasks. Can be either

- a variable specified as a literal variable name (e.g., column_name),
- a string with the variable name (e.g., "column_name"), or a character vector of variable names (e.g., c("col1", "col2", "col3")),
- a formula with variable names (e.g., $\sim$ column_1 + column_2),
- a vector of positive integers, giving the positions counting from the left (e.g. 1 or $c(1,3,5)$ ),
- a vector of negative integers, giving the positions counting from the right (e.g., -1 or $-1:-3$ ),
- one of the following select-helpers: starts_with(""), ends_with(""), contains(""), a range using : or regex (""),
- or a function testing for logical conditions, e.g. is.numeric() (or is.numeric), or any user-defined function that selects the variables for which the function returns TRUE (like: foo <- function ( $x$ ) mean $(x)>3$ ),
- ranges specified via literal variable names, select-helpers (except regex()) and (user-defined) functions can be negated, i.e. return non-matching elements, when prefixed with a -, e.g. -ends_with(""), -is.numeric or -Sepal.Width:Petal.Length. Note: Negation means that matches are excluded, and thus, the exclude argument can be used alternatively. For instance, select=-ends_with("Length") (with -) is equivalent to exclude=ends_with("Length") (no -). In case negation should not work as expected, use the exclude argument instead.
If NULL, selects all columns. Patterns that found no matches are silently ignored, e.g. find_columns(iris, select $=c($ "Species", "Test")) will just return "Species".
before, after Destination of columns. Supplying neither will move columns to the left-hand side; specifying both is an error. Can be a character vector, indicating the name of the destination column, or a numeric value, indicating the index number of the destination column. If -1 , will be added before or after the last column.
ignore_case Logical, if TRUE and when one of the select-helpers or a regular expression is used in select, ignores lower/upper case in the search pattern when matching against variable names.
verbose Toggle warnings.
... Arguments passed down to other functions. Mostly not used yet.


## Value

A data frame with reordered columns.

## See Also

- Functions to rename stuff: data_rename(), data_rename_rows(), data_addprefix(), data_addsuffix()
- Functions to reorder or remove columns: data_reorder(), data_relocate(), data_remove()
- Functions to reshape, pivot or rotate data frames: data_to_long(), data_to_wide(), data_rotate()
- Functions to recode data: rescale(), reverse(), categorize(), change_code(), slide()
- Functions to standardize, normalize, rank-transform: center(), standardize(), normalize(), ranktransform(), winsorize()
- Split and merge data frames: data_partition(), data_merge()
- Functions to find or select columns: data_select(), data_find()
- Functions to filter rows: data_match(), data_filter()


## Examples

```
\# Reorder columns
head(data_relocate(iris, select = "Species", before = "Sepal.Length"))
head(data_relocate(iris, select = "Species", before = "Sepal.Width"))
head(data_relocate(iris, select = "Sepal.Width", after = "Species"))
\# same as
head(data_relocate(iris, select = "Sepal.Width", after = -1))
\# reorder multiple columns
head(data_relocate(iris, select = c("Species", "Petal.Length"), after = "Sepal.Width"))
\# same as
head(data_relocate(iris, select = c("Species", "Petal.Length"), after = 2))
\# Reorder columns
head(data_reorder(iris, c("Species", "Sepal.Length")))
head(data_reorder(iris, c("Species", "dupa"))) \# Safe for non-existing cols
\# Remove columns
head(data_remove(iris, "Sepal.Length"))
head(data_remove(iris, starts_with("Sepal")))
```

data_restoretype Restore the type of columns according to a reference data frame

## Description

Restore the type of columns according to a reference data frame

## Usage

data_restoretype(data, reference = NULL, ...)

## Arguments

$$
\begin{array}{ll}
\text { data } & \text { A data frame to pivot. } \\
\text { reference } & \text { A reference data frame from which to find the correct column types. } \\
\ldots & \text { Currently not used. }
\end{array}
$$

## Value

A data frame with columns whose types have been restored based on the reference data frame.

## Examples

```
data <- data.frame(
    Sepal.Length = c("1", "3", "2"),
    Species = c("setosa", "versicolor", "setosa"),
    New = c("1", "3", "4")
```

)
fixed <- data_restoretype(data, reference = iris)
summary (fixed)
data_rotate Rotate a data frame

## Description

This function rotates a data frame, i.e. columns become rows and vice versa. It's the equivalent of using t () but restores the data.frame class, preserves attributes and prints a warning if the data type is modified (see example).

## Usage

data_rotate(data, rownames $=$ NULL, colnames $=$ FALSE, verbose $=$ TRUE)
data_transpose(data, rownames $=$ NULL, colnames $=$ FALSE, verbose $=$ TRUE)

## Arguments

| data | A data frame. |
| :--- | :--- |
| rownames | Character vector (optional). If not NULL, the data frame's rownames will be <br> added as (first) column to the output, with rownames being the name of this <br> column. |
| colnames | Logical or character vector (optional). If TRUE, the values of the first column in <br> x will be used as column names in the rotated data frame. If a character vector, <br> values from that column are used as column names. |
| verbose | Toggle warnings. |

## Value

A (rotated) data frame.

## See Also

- Functions to rename stuff: data_rename(), data_rename_rows(), data_addprefix(), data_addsuffix()
- Functions to reorder or remove columns: data_reorder (), data_relocate(), data_remove()
- Functions to reshape, pivot or rotate data frames: data_to_long(), data_to_wide(), data_rotate()
- Functions to recode data: rescale(), reverse(), categorize(), change_code(), slide()
- Functions to standardize, normalize, rank-transform: center(), standardize(), normalize(), ranktransform(), winsorize()
- Split and merge data frames: data_partition(), data_merge()
- Functions to find or select columns: data_select(), data_find()
- Functions to filter rows: data_match(), data_filter()


## Examples

$x<-\operatorname{mtcars}[1: 3,1: 4]$
x
data_rotate(x)
data_rotate(x, rownames = "property")
\# use values in 1. column as column name
data_rotate ( $x$, colnames $=$ TRUE)
data_rotate(x, rownames = "property", colnames = TRUE)
\# warn that data types are changed
str(data_rotate(iris[1:4, ]))
\# use either first column or specific column for column names
$x<-\operatorname{data} . f r a m e(a=1: 5, b=11: 15, c=1 e t t e r s[1: 5], d=r n o r m(5))$
data_rotate ( x , colnames $=$ TRUE)
data_rotate(x, colnames = "c")
data_tabulate Create frequency tables of variables

## Description

This function creates frequency tables of variables, including the number of levels/values as well as the distribution of raw, valid and cumulative percentages.

## Usage

```
data_tabulate(x, ...)
## Default S3 method:
data_tabulate(x, drop_levels = FALSE, name = NULL, verbose = TRUE, ...)
## S3 method for class 'data.frame'
data_tabulate(
    x,
    select = NULL,
    exclude = NULL,
    ignore_case = FALSE,
    collapse = FALSE,
    drop_levels = FALSE,
    verbose = TRUE,
)
```


## Arguments

x
...
drop_levels Logical, if TRUE, factor levels that do not occur in the data are included in the table (with frequency of zero), else unused factor levels are dropped from the frequency table.
name Optional character string, which includes the name that is used for printing.
verbose
select
exclude See select, however, column names matched by the pattern from exclude will be excluded instead of selected. If NULL (the default), excludes no columns.
ignore_case Logical, if TRUE and when one of the select-helpers or a regular expression is used in select, ignores lower/upper case in the search pattern when matching against variable names.
collapse Logical, if TRUE collapses multiple tables into one larger table for printing. This affects only printing, not the returned object.

## Value

A data frame, or a list of data frames, with one frequency table as data frame per variable.

## Selection of variables - the select argument

For most functions that have a select argument (including this function), the complete input data frame is returned, even when select only selects a range of variables. That is, the function is only applied to those variables that have a match in select, while all other variables remain unchanged. In other words: for this function, select will not omit any non-included variables, so that the returned data frame will include all variables from the input data frame.

## Examples

```
data(efc)
# vector/factor
data_tabulate(efc$c172code)
# data frame
data_tabulate(efc, c("e42dep", "c172code"))
# grouped data frame
if (requireNamespace("poorman")) {
    suppressPackageStartupMessages(library(poorman, quietly = TRUE))
    efc %>%
        group_by(c172code) %>%
        data_tabulate("e16sex")
    # collapse tables
    efc %>%
        group_by(c172code) %>%
        data_tabulate("e16sex", collapse = TRUE)
}
# for larger N's (> 100000), a big mark is automatically added
set.seed(123)
x <- sample(1:3, 1e6, TRUE)
data_tabulate(x, name = "Large Number")
# to remove the big mark, use "print(..., big_mark = "")"
print(data_tabulate(x), big_mark = "")
```

data_to_long Reshape (pivot) data from wide to long

## Description

This function "lengthens" data, increasing the number of rows and decreasing the number of columns. This is a dependency-free base-R equivalent of tidyr: :pivot_longer().

## Usage

```
data_to_long(
    data,
    select = "all",
    names_to = "name",
    names_prefix = NULL,
    names_sep = NULL,
    names_pattern = NULL,
    values_to = "value",
    values_drop_na = FALSE,
    rows_to = NULL,
    ignore_case = FALSE,
    regex = FALSE,
    ...,
    cols,
    colnames_to
)
reshape_longer(
    data,
    select = "all",
    names_to = "name",
    names_prefix = NULL,
    names_sep = NULL,
    names_pattern = NULL,
    values_to = "value",
    values_drop_na = FALSE,
    rows_to = NULL,
    ignore_case = FALSE,
    regex = FALSE,
    ...,
    cols,
    colnames_to
)
```


## Arguments

## data

A data frame to pivot.
select Variables that will be included when performing the required tasks. Can be either

- a variable specified as a literal variable name (e.g., column_name),
- a string with the variable name (e.g., "column_name"), or a character vector of variable names (e.g., c("col1", "col2", "col3")),
- a formula with variable names (e.g., ~column_1 + column_2),
- a vector of positive integers, giving the positions counting from the left (e.g. 1 or $c(1,3,5)$ ),
- a vector of negative integers, giving the positions counting from the right (e.g., -1 or $-1:-3$ ),
- one of the following select-helpers: starts_with(""), ends_with(""), contains(""), a range using : or regex(""),
- or a function testing for logical conditions, e.g. is.numeric() (or is.numeric), or any user-defined function that selects the variables for which the function returns TRUE (like: foo <- function $(x)$ mean $(x)>3$ ),
- ranges specified via literal variable names, select-helpers (except regex()) and (user-defined) functions can be negated, i.e. return non-matching elements, when prefixed with a -, e.g. -ends_with(""), -is.numeric or -Sepal.Width:Petal.Length. Note: Negation means that matches are excluded, and thus, the exclude argument can be used alternatively. For instance, select=-ends_with("Length") (with -) is equivalent to exclude=ends_with("Length") (no -). In case negation should not work as expected, use the exclude argument instead.
If NULL, selects all columns. Patterns that found no matches are silently ignored, e.g. find_columns(iris, select $=c($ "Species", "Test")) will just return "Species".
names_to The name of the new column that will contain the column names.
names_prefix A regular expression used to remove matching text from the start of each variable name.
names_sep, names_pattern
If names_to contains multiple values, this argument controls how the column name is broken up. names_pattern takes a regular expression containing matching groups, i.e. "()".
values_to The name of the new column that will contain the values of the pivoted variables.
values_drop_na If TRUE, will drop rows that contain only NA in the values_to column. This effectively converts explicit missing values to implicit missing values, and should generally be used only when missing values in data were created by its structure.
rows_to The name of the column that will contain the row names or row numbers from the original data. If NULL, will be removed.
ignore_case Logical, if TRUE and when one of the select-helpers or a regular expression is used in select, ignores lower/upper case in the search pattern when matching against variable names.
regex Logical, if TRUE, the search pattern from select will be treated as regular expression. When regex $=$ TRUE, select must be a character string (or a variable containing a character string) and is not allowed to be one of the supported select-helpers or a character vector of length $>1$. regex $=$ TRUE is comparable to using one of the two select-helpers, select = contains("") or select $=$ regex (""), however, since the select-helpers may not work when called from inside other functions (see 'Details'), this argument may be used as workaround.
... Currently not used.
cols Identical to select. This argument is here to ensure compatibility with tidyr: : pivot_longer(). If both select and cols are provided, cols is used.
colnames_to Deprecated. Use names_to instead.


## Value

If a tibble was provided as input, reshape_longer() also returns a tibble. Otherwise, it returns a data frame.

## See Also

- Functions to rename stuff: data_rename(), data_rename_rows(), data_addprefix(), data_addsuffix()
- Functions to reorder or remove columns: data_reorder(), data_relocate(), data_remove()
- Functions to reshape, pivot or rotate data frames: data_to_long(), data_to_wide(), data_rotate()
- Functions to recode data: rescale(), reverse(), categorize(), change_code(), slide()
- Functions to standardize, normalize, rank-transform: center(), standardize(), normalize(), ranktransform(), winsorize()
- Split and merge data frames: data_partition(), data_merge()
- Functions to find or select columns: data_select(), data_find()
- Functions to filter rows: data_match(), data_filter()


## Examples

```
wide_data <- data.frame(replicate(5, rnorm(10)))
# Default behaviour (equivalent to tidyr::pivot_longer(wide_data, cols = 1:5))
data_to_long(wide_data)
# Customizing the names
data_to_long(wide_data,
    select = c(1, 2),
    names_to = "Column",
    values_to = "Numbers",
    rows_to = "Row"
)
# Full example
# -----------------
if (require("psych")) {
    data <- psych::bfi # Wide format with one row per participant's personality test
    # Pivot long format
    data_to_long(data,
        select = regex("\\d"), # Select all columns that contain a digit
        colnames_to = "Item",
        values_to = "Score",
        rows_to = "Participant"
    )
    if (require("tidyr")) {
        reshape_longer(
            tidyr::who,
            select = new_sp_m014:newrel_f65,
```

```
            names_to = c("diagnosis", "gender", "age"),
            names_pattern = "new_?(.*)_(.)(.*)",
            values_to = "count"
        )
        }
}
```

data_to_wide Reshape (pivot) data from long to wide

## Description

This function "widens" data, increasing the number of columns and decreasing the number of rows. This is a dependency-free base-R equivalent of tidyr: : pivot_wider().

## Usage

data_to_wide( data,
id_cols = NULL,
values_from = "Value",
names_from = "Name",
names_sep = "_",
names_prefix = "",
names_glue = NULL,
values_fill = NULL,
verbose = TRUE,
...,
colnames_from,
rows_from,
sep
)
reshape_wider(
data,
id_cols = NULL,
values_from = "Value",
names_from = "Name",
names_sep = "_",
names_prefix = "",
names_glue = NULL, values_fill = NULL, verbose = TRUE,
...,
colnames_from, rows_from,

```
    sep
```

)

## Arguments

| data | A data frame to pivot. |
| :---: | :---: |
| id_cols | The name of the column that identifies the rows. If NULL, it will use all the unique rows. |
| values_from | The name of the column that contains the values to be used as future variable values. |
| names_from | The name of the column that contains the levels to be used as future column names. |
| names_sep | If names_from or values_from contains multiple variables, this will be used to join their values together into a single string to use as a column name. |
| names_prefix | String added to the start of every variable name. This is particularly useful if names_from is a numeric vector and you want to create syntactic variable names. |
| names_glue | Instead of names_sep and names_prefix, you can supply a glue specification that uses the names_from columns to create custom column names. Note that the only delimiters supported by names_glue are curly brackets, $\{$ and \}. |
| values_fill | Optionally, a (scalar) value that will be used to replace missing values in the new columns created. |
| verbose | Toggle warnings. |
|  | Not used for now. |
| colnames_from | Deprecated. Use names_from instead. |
| rows_from | Deprecated. Use id_cols instead. |
| sep | Deprecated. Use names_sep instead. |

## Value

If a tibble was provided as input, reshape_wider () also returns a tibble. Otherwise, it returns a data frame.

## See Also

- Functions to rename stuff: data_rename(), data_rename_rows(), data_addprefix(), data_addsuffix()
- Functions to reorder or remove columns: data_reorder(), data_relocate(), data_remove()
- Functions to reshape, pivot or rotate data frames: data_to_long(), data_to_wide(), data_rotate()
- Functions to recode data: rescale(), reverse(), categorize(), change_code(), slide()
- Functions to standardize, normalize, rank-transform: center(), standardize(), normalize(), ranktransform(), winsorize()
- Split and merge data frames: data_partition(), data_merge()
- Functions to find or select columns: data_select(), data_find()
- Functions to filter rows: data_match(), data_filter()


## Examples

```
data_long <- read.table(header = TRUE, text = "
    subject sex condition measurement
1 M control 7.9
1 \begin{tabular}{lll}
M & cond1 & 12.3
\end{tabular}
            1 M cond2 10.7
            2 F control 6.3
            2 F cond1 10.6
            2 F cond2 11.1
            F control 9.5
            3 F cond1 13.1
            3 F cond2 13.8
            4 control 11.5
            4 cond1 13.4
            4 cond2 12.9")
reshape_wider(
    data_long,
    id_cols = "subject",
    names_from = "condition",
    values_from = "measurement"
)
reshape_wider(
    data_long,
    id_cols = "subject",
    names_from = "condition",
    values_from = "measurement",
    names_prefix = "Var.",
    names_sep = "."
)
production <- expand.grid(
    product = c("A", "B"),
    country = c("AI", "EI"),
    year = 2000:2014
)
production <- data_filter(production, (product == "A" & country == "AI") | product == "B")
production$production <- rnorm(nrow(production))
reshape_wider(
    production,
    names_from = c("product", "country"),
    values_from = "production",
    names_glue = "prod_{product}_{country}"
)
```

demean $\quad$ Compute group-meaned and de-meaned variables

## Description

demean() computes group- and de-meaned versions of a variable that can be used in regression analysis to model the between- and within-subject effect. degroup() is more generic in terms of the centering-operation. While demean() always uses mean-centering, degroup() can also use the mode or median for centering.

## Usage

```
demean(
    x,
    select,
    group,
    suffix_demean = "_within",
    suffix_groupmean = "_between",
    add_attributes = TRUE,
    verbose = TRUE
)
degroup(
    x,
    select,
    group,
    center = "mean",
    suffix_demean = "_within",
    suffix_groupmean = "_between",
    add_attributes = TRUE,
    verbose = TRUE
)
detrend(
    x,
    select,
    group,
    center = "mean",
    suffix_demean = "_within",
    suffix_groupmean = "_between",
    add_attributes = TRUE,
    verbose = TRUE
    )
```


## Arguments

x
A data frame.

```
select Character vector (or formula) with names of variables to select that should be
    group- and de-meaned.
group Character vector (or formula) with the name of the variable that indicates the
    group- or cluster-ID.
suffix_demean, suffix_groupmean
    String value, will be appended to the names of the group-meaned and de-meaned
    variables of x. By default, de-meaned variables will be suffixed with "_wi thin"
    and grouped-meaned variables with "_between".
add_attributes Logical, if TRUE, the returned variables gain attributes to indicate the within- and
    between-effects. This is only relevant when printing model_parameters() - in
    such cases, the within- and between-effects are printed in separated blocks.
verbose Toggle warnings and messages.
center Method for centering. demean() always performs mean-centering, while degroup()
    can use center = "median" or center = "mode" for median- or mode-centering,
    and also "min" or "max".
```


## Details

Heterogeneity Bias: Mixed models include different levels of sources of variability, i.e. error terms at each level. When macro-indicators (or level-2 predictors, or higher-level units, or more general: group-level predictors that vary within and across groups) are included as fixed effects (i.e. treated as covariate at level-1), the variance that is left unaccounted for this covariate will be absorbed into the error terms of level-1 and level-2 (Bafumi and Gelman 2006; Gelman and Hill 2007, Chapter 12.6.): "Such covariates contain two parts: one that is specific to the higherlevel entity that does not vary between occasions, and one that represents the difference between occasions, within higher-level entities" (Bell et al. 2015). Hence, the error terms will be correlated with the covariate, which violates one of the assumptions of mixed models (iid, independent and identically distributed error terms). This bias is also called the heterogeneity bias (Bell et al. 2015). To resolve this problem, level-2 predictors used as (level-1) covariates should be separated into their "within" and "between" effects by "de-meaning" and "group-meaning": After demeaning time-varying predictors, "at the higher level, the mean term is no longer constrained by Level 1 effects, so it is free to account for all the higher-level variance associated with that variable" (Bell et al. 2015).

Panel data and correlating fixed and group effects: demean() is intended to create group- and de-meaned variables for panel regression models (fixed effects models), or for complex random-effect-within-between models (see Bell et al. 2015, 2018), where group-effects (random effects) and fixed effects correlate (see Bafumi and Gelman 2006). This can happen, for instance, when analyzing panel data, which can lead to Heterogeneity Bias. To control for correlating predictors and group effects, it is recommended to include the group-meaned and de-meaned version of timevarying covariates (and group-meaned version of time-invariant covariates that are on a higher level, e.g. level-2 predictors) in the model. By this, one can fit complex multilevel models for panel data, including time-varying predictors, time-invariant predictors and random effects.

Why mixed models are preferred over fixed effects models: A mixed models approach can model the causes of endogeneity explicitly by including the (separated) within- and betweeneffects of time-varying fixed effects and including time-constant fixed effects. Furthermore, mixed
models also include random effects, thus a mixed models approach is superior to classic fixedeffects models, which lack information of variation in the group-effects or between-subject effects. Furthermore, fixed effects regression cannot include random slopes, which means that fixed effects regressions are neglecting "cross-cluster differences in the effects of lower-level controls (which) reduces the precision of estimated context effects, resulting in unnecessarily wide confidence intervals and low statistical power" (Heisig et al. 2017).

Terminology: The group-meaned variable is simply the mean of an independent variable within each group (or id-level or cluster) represented by group. It represents the cluster-mean of an independent variable. The regression coefficient of a group-meaned variable is the between-subject-effect. The de-meaned variable is then the centered version of the group-meaned variable. De-meaning is sometimes also called person-mean centering or centering within clusters. The regression coefficient of a de-meaned variable represents the within-subject-effect.

De-meaning with continuous predictors: For continuous time-varying predictors, the recommendation is to include both their de-meaned and group-meaned versions as fixed effects, but not the raw (untransformed) time-varying predictors themselves. The de-meaned predictor should also be included as random effect (random slope). In regression models, the coefficient of the demeaned predictors indicates the within-subject effect, while the coefficient of the group-meaned predictor indicates the between-subject effect.

De-meaning with binary predictors: For binary time-varying predictors, there are two recommendations. First is to include the raw (untransformed) binary predictor as fixed effect only and the de-meaned variable as random effect (random slope). The alternative would be to add the de-meaned version(s) of binary time-varying covariates as additional fixed effect as well (instead of adding it as random slope). Centering time-varying binary variables to obtain within-effects (level 1) isn't necessary. They have a sensible interpretation when left in the typical 0/1 format (Hoffmann 2015, chapter 8-2.I). demean() will thus coerce categorical time-varying predictors to numeric to compute the de- and group-meaned versions for these variables, where the raw (untransformed) binary predictor and the de-meaned version should be added to the model.

De-meaning of factors with more than 2 levels: Factors with more than two levels are demeaned in two ways: first, these are also converted to numeric and de-meaned; second, dummy variables are created (binary, with $0 / 1$ coding for each level) and these binary dummy-variables are de-meaned in the same way (as described above). Packages like panelr internally convert factors to dummies before demeaning, so this behaviour can be mimicked here.

De-meaning interaction terms: There are multiple ways to deal with interaction terms of within- and between-effects. A classical approach is to simply use the product term of the demeaned variables (i.e. introducing the de-meaned variables as interaction term in the model formula, e.g. y ~ x_within * time_within). This approach, however, might be subject to bias (see Giesselmann \& Schmidt-Catran 2020).

Another option is to first calculate the product term and then apply the de-meaning to it. This approach produces an estimator "that reflects unit-level differences of interacted variables whose moderators vary within units", which is desirable if no within interaction of two time-dependent variables is required.

A third option, when the interaction should result in a genuine within estimator, is to "double de-mean" the interaction terms (Giesselmann \& Schmidt-Catran 2018), however, this is currently
not supported by demean(). If this is required, the wmb() function from the panelr package should be used.

To de-mean interaction terms for within-between models, simply specify the term as interaction for the select-argument, e.g. select = "a*b" (see 'Examples').

Analysing panel data with mixed models using lme4: A description of how to translate the formulas described in Bell et al. 2018 into R using lmer() from lme4 can be found in this vignette.

## Value

A data frame with the group-/de-meaned variables, which get the suffix "_between" (for the groupmeaned variable) and "_within" (for the de-meaned variable) by default.

## References

- Bafumi J, Gelman A. 2006. Fitting Multilevel Models When Predictors and Group Effects Correlate. In. Philadelphia, PA: Annual meeting of the American Political Science Association.
- Bell A, Fairbrother M, Jones K. 2019. Fixed and Random Effects Models: Making an Informed Choice. Quality \& Quantity (53); 1051-1074
- Bell A, Jones K. 2015. Explaining Fixed Effects: Random Effects Modeling of Time-Series Cross-Sectional and Panel Data. Political Science Research and Methods, 3(1), 133-153.
- Gelman A, Hill J. 2007. Data Analysis Using Regression and Multilevel/Hierarchical Models. Analytical Methods for Social Research. Cambridge, New York: Cambridge University Press
- Giesselmann M, Schmidt-Catran, AW. 2020. Interactions in fixed effects regression models. Sociological Methods \& Research, 1-28. https://doi.org/10.1177/0049124120914934
- Heisig JP, Schaeffer M, Giesecke J. 2017. The Costs of Simplicity: Why Multilevel Models May Benefit from Accounting for Cross-Cluster Differences in the Effects of Controls. American Sociological Review 82 (4): 796-827.
- Hoffman L. 2015. Longitudinal analysis: modeling within-person fluctuation and change. New York: Routledge


## See Also

If grand-mean centering (instead of centering within-clusters) is required, see center().

## Examples

```
data(iris)
iris$ID <- sample(1:4, nrow(iris), replace = TRUE) # fake-ID
iris$binary <- as.factor(rbinom(150, 1, .35)) # binary variable
x <- demean(iris, select = c("Sepal.Length", "Petal.Length"), group = "ID")
head(x)
```

```
x <- demean(iris, select = c("Sepal.Length", "binary", "Species"), group = "ID")
head(x)
# demean interaction term x*y
dat <- data.frame(
    a = c(1, 2, 3, 4, 1, 2, 3, 4),
    x = c(4, 3, 3, 4, 1, 2, 1, 2),
    y = c(1, 2, 1, 2, 4, 3, 2, 1),
    ID = c(1, 2, 3, 1, 2, 3, 1, 2)
)
demean(dat, select = c("a", "x*y"), group = "ID")
# or in formula-notation
demean(dat, select = ~ a + x * y, group = ~ID)
```

```
describe_distribution Describe a distribution
```


## Description

This function describes a distribution by a set of indices (e.g., measures of centrality, dispersion, range, skewness, kurtosis).

## Usage

describe_distribution(x, ...)
\#\# S3 method for class 'numeric'
describe_distribution(
x ,
centrality = "mean",
dispersion = TRUE,
iqr = TRUE,
range = TRUE,
quartiles = FALSE, ci = NULL,
iterations = 100,
threshold = 0.1,
verbose = TRUE,
)
\#\# S3 method for class 'factor'
describe_distribution(x, dispersion = TRUE, range = TRUE, verbose = TRUE, ...)
\#\# S3 method for class 'data.frame'

```
describe_distribution(
    X,
    select = NULL,
    exclude = NULL,
    centrality = "mean",
    dispersion = TRUE,
    iqr = TRUE,
    range = TRUE,
    quartiles = FALSE,
    include_factors = FALSE,
    ci = NULL,
    iterations = 100,
    threshold = 0.1,
    ignore_case = FALSE,
    verbose = TRUE,
)
```


## Arguments

x
...
centrality
dispersion
iqr Logical, if TRUE, the interquartile range is calculated (based on stats: :IQR(), using type $=6$ ).
range $\quad$ Return the range ( $\min$ and $\max$ ).
quartiles
ci
ci Confidence Interval (CI) level. Default is NULL, i.e. no confidence intervals are computed. If not NULL, confidence intervals are based on bootstrap replicates (see iterations). If centrality = "all", the bootstrapped confidence interval refers to the first centrality index (which is typically the median).
iterations The number of bootstrap replicates for computing confidence intervals. Only applies when ci is not NULL.
threshold For centrality = "trimmed" (i.e. trimmed mean), indicates the fraction ( 0 to 0.5 ) of observations to be trimmed from each end of the vector before the mean is computed.
verbose Toggle warnings and messages.
select Variables that will be included when performing the required tasks. Can be either

- a variable specified as a literal variable name (e.g., column_name),
- a string with the variable name (e.g., "column_name"), or a character vector of variable names (e.g., c("col1", "col2", "col3")),
- a formula with variable names (e.g., ~column_1 + column_2),
- a vector of positive integers, giving the positions counting from the left (e.g. 1 or c(1, 3,5)),
- a vector of negative integers, giving the positions counting from the right (e.g., -1 or $-1:-3$ ),
- one of the following select-helpers: starts_with(""), ends_with(""), contains(""), a range using : or regex(""),
- or a function testing for logical conditions, e.g. is.numeric() (or is .numeric), or any user-defined function that selects the variables for which the function returns TRUE (like: foo <- function (x) mean $(x)>3$ ),
- ranges specified via literal variable names, select-helpers (except regex()) and (user-defined) functions can be negated, i.e. return non-matching elements, when prefixed with a -, e.g. -ends_with(""), -is.numeric or -Sepal.Width:Petal.Length. Note: Negation means that matches are excluded, and thus, the exclude argument can be used alternatively. For instance, select=-ends_with("Length") (with -) is equivalent to exclude=ends_with("Length") (no -). In case negation should not work as expected, use the exclude argument instead.
If NULL, selects all columns. Patterns that found no matches are silently ignored, e.g. find_columns(iris, select = c("Species", "Test")) will just return "Species".
exclude See select, however, column names matched by the pattern from exclude will be excluded instead of selected. If NULL (the default), excludes no columns.
include_factors
Logical, if TRUE, factors are included in the output, however, only columns for range (first and last factor levels) as well as $n$ and missing will contain information.
ignore_case Logical, if TRUE and when one of the select-helpers or a regular expression is used in select, ignores lower/upper case in the search pattern when matching against variable names.


## Details

If $x$ is a data frame, only numeric variables are kept and will be displayed in the summary.
If $x$ is a list, the behavior is different whether $x$ is a stored list. If $x$ is stored (for example, describe_distribution(mylist) where mylist was created before), artificial variable names are used in the summary (Var_1, Var_2, etc.). If $x$ is an unstored list (for example, describe_distribution(list(mtcars $\$$ r then "mtcars\$mpg" is used as variable name.

## Value

A data frame with columns that describe the properties of the variables.

## Selection of variables - the select argument

For most functions that have a select argument (including this function), the complete input data frame is returned, even when select only selects a range of variables. That is, the function is only
applied to those variables that have a match in select, while all other variables remain unchanged. In other words: for this function, select will not omit any non-included variables, so that the returned data frame will include all variables from the input data frame.

## Note

There is also a plot()-method implemented in the see-package.

## Examples

```
describe_distribution(rnorm(100))
data(iris)
describe_distribution(iris)
describe_distribution(iris, include_factors = TRUE, quartiles = TRUE)
describe_distribution(list(mtcars$mpg, mtcars$cyl))
```

```
distribution_mode Compute mode for a statistical distribution
```


## Description

Compute mode for a statistical distribution

## Usage

distribution_mode(x)

## Arguments

## X

An atomic vector, a list, or a data frame.

## Value

The value that appears most frequently in the provided data. The returned data structure will be the same as the entered one.

## Examples

```
distribution_mode(c(1, 2, 3, 3, 4, 5))
distribution_mode(c(1.5, 2.3, 3.7, 3.7, 4.0, 5))
```


## Description

Selected variables from the EUROFAMCARE survey. Useful when testing on "real-life" data sets, including random missing values. This data set also has value and variable label attributes.

## Description

find_columns() returns column names from a data set that match a certain search pattern, while get_columns() returns the found data. data_select() is an alias for get_columns(), and data_find() is an alias for find_columns().

## Usage

find_columns(
data,
select $=$ NULL,
exclude $=$ NULL,
ignore_case = FALSE, regex = FALSE,
verbose = TRUE,
)
data_find(
data,
select $=$ NULL,
exclude = NULL,
ignore_case = FALSE, regex = FALSE, verbose $=$ TRUE,
. . .
)
get_columns(
data,
select $=$ NULL,
exclude = NULL,
ignore_case = FALSE,
regex = FALSE,

```
    verbose = TRUE,
    )
    data_select(
        data,
        select = NULL,
        exclude = NULL,
        ignore_case = FALSE,
        regex = FALSE,
        verbose = TRUE,
    ...
)
```


## Arguments

data A data frame
select Variables that will be included when performing the required tasks. Can be either

- a variable specified as a literal variable name (e.g., column_name),
- a string with the variable name (e.g., "column_name"), or a character vector of variable names (e.g., c("col1", "col2", "col3")),
- a formula with variable names (e.g., ~column_1 + column_2),
- a vector of positive integers, giving the positions counting from the left (e.g. 1 or $c(1,3,5)$ ),
- a vector of negative integers, giving the positions counting from the right (e.g., -1 or $-1:-3$ ),
- one of the following select-helpers: starts_with(""), ends_with(""), contains(""), a range using : or regex (""),
- or a function testing for logical conditions, e.g. is.numeric() (or is.numeric), or any user-defined function that selects the variables for which the function returns TRUE (like: foo <- function $(x)$ mean $(x)>3$ ),
- ranges specified via literal variable names, select-helpers (except regex()) and (user-defined) functions can be negated, i.e. return non-matching elements, when prefixed with a -, e.g. -ends_with(""), -is.numeric or -Sepal.Width:Petal.Length. Note: Negation means that matches are excluded, and thus, the exclude argument can be used alternatively. For instance, select=-ends_with("Length") (with -) is equivalent to exclude=ends_with("Length") (no -). In case negation should not work as expected, use the exclude argument instead.

If NULL, selects all columns. Patterns that found no matches are silently ignored, e.g. find_columns(iris, select = c("Species", "Test")) will just return "Species".
exclude See select, however, column names matched by the pattern from exclude will be excluded instead of selected. If NULL (the default), excludes no columns.

$$
\begin{array}{ll}
\text { ignore_case } & \begin{array}{l}
\text { Logical, if TRUE and when one of the select-helpers or a regular expression is } \\
\text { used in select, ignores lower/upper case in the search pattern when matching } \\
\text { against variable names. }
\end{array} \\
\text { regex } & \begin{array}{l}
\text { Logical, if TRUE, the search pattern from select will be treated as regular ex- } \\
\text { pression. When regex }=\text { TRUE, select must be a character string (or a variable } \\
\text { containing a character string) and is not allowed to be one of the supported }
\end{array} \\
\text { select-helpers or a character vector of length }>1 \text {. regex = TRUE is compara- } \\
\text { ble to using one of the two select-helpers, select = contains("") or select } \\
\text { = regex(""), however, since the select-helpers may not work when called from } \\
\text { inside other functions (see 'Details'), this argument may be used as workaround. } \\
\text { verbose } & \text { Toggle warnings. }
\end{array}
$$

... Arguments passed down to other functions. Mostly not used yet.

## Details

Note that there are some limitations when calling this from inside other functions. The following will work as expected, returning all columns that start with "Sep":

```
foo <- function(data) {
    find_columns(data, select = starts_with("Sep"))
}
foo(iris)
```

However, this example won't work as expected!

```
foo <- function(data) {
    i <- "Sep"
    find_columns(data, select = starts_with(i))
}
foo(iris)
```

One workaround is to use the regex argument, which provides at least a bit more flexibility than exact matching. regex in its basic usage (as seen below) means that select behaves like the contains("") select-helper, but can also make the function more flexible by allowing to define complex regular expression pattern in select.

```
foo <- function(data) {
    i <- "Sep"
    find_columns(data, select = i, regex = TRUE)
}
foo(iris)
```


## Value

find_columns() returns a character vector with column names that matched the pattern in select and exclude, or NULL if no matching column name was found. get_columns() returns a data frame with matching columns.

## See Also

- Functions to rename stuff: data_rename(), data_rename_rows(), data_addprefix(), data_addsuffix()
- Functions to reorder or remove columns: data_reorder(), data_relocate(), data_remove()
- Functions to reshape, pivot or rotate data frames: data_to_long(), data_to_wide(), data_rotate()
- Functions to recode data: rescale(), reverse(), categorize(), change_code(), slide()
- Functions to standardize, normalize, rank-transform: center(), standardize(), normalize(), ranktransform(), winsorize()
- Split and merge data frames: data_partition(), data_merge()
- Functions to find or select columns: data_select(), data_find()
- Functions to filter rows: data_match(), data_filter()


## Examples

```
# Find columns names by pattern
find_columns(iris, starts_with("Sepal"))
find_columns(iris, ends_with("Width"))
find_columns(iris, regex("\\."))
find_columns(iris, c("Petal.Width", "Sepal.Length"))
# starts with "Sepal", but not allowed to end with "width"
find_columns(iris, starts_with("Sepal"), exclude = contains("Width"))
# find numeric with mean > 3.5
numeric_mean_35 <- function(x) is.numeric(x) && mean(x, na.rm = TRUE) > 3.5
find_columns(iris, numeric_mean_35)
```

format_text Convenient text formatting functionalities

## Description

Convenience functions to manipulate and format text.

## Usage

```
format_text(
    text,
    sep = ", ",
    last = " and ",
    width = NULL,
    enclose = NULL,
)
```

text_fullstop(text)

```
text_lastchar(text, \(\mathrm{n}=1\) )
text_concatenate(text, sep = ", ", last = " and ", enclose = NULL)
text_paste(text, text2 \(=\) NULL, sep \(=", "\), enclose \(=\) NULL, ...)
text_remove(text, pattern = "", ...)
text_wrap(text, width = NULL, ...)
```


## Arguments

| text, text2 | A character string. |
| :--- | :--- |
| sep | Separator. |
| last | Last separator. |
| width | Positive integer giving the target column width for wrapping lines in the output. <br> Can be "auto", in which case it will select $90 \backslash$ default width. |
| enclose | Character that will be used to wrap elements of text, so these can be, e.g., <br> enclosed with quotes or backticks. If NULL (default), text elements will not be <br> enclosed. |
| $\ldots$ | Other arguments to be passed to or from other functions. |
| n | The number of characters to find. |
| pattern | Character vector. For data_rename(), indicates columns that should be se- <br> lected for renaming. Can be NULL (in which case all columns are selected). For <br> data_addprefix() or data_addsuffix(), a character string, which will be <br> added as prefix or suffix to the column names. |

## Value

A character string.

## Examples

```
# Add full stop if missing
text_fullstop(c("something", "something else."))
# Find last characters
text_lastchar(c("ABC", "DEF"), n = 2)
# Smart concatenation
text_concatenate(c("First", "Second", "Last"))
text_concatenate(c("First", "Second", "Last"), last = " or ", enclose = "`")
# Remove parts of string
text_remove(c("one!", "two", "three!"), "!")
# Wrap text
```

```
long_text <- paste(rep("abc ", 100), collapse = "")
cat(text_wrap(long_text, width = 50))
# Paste with optional separator
text_paste(c("A", "", "B"), c("42", "42", "42"))
```

nhanes_sample Sample dataset from the National Health and Nutrition Examination
Survey

## Description

Selected variables from the National Health and Nutrition Examination Survey that are used in the example from Lumley (2010), Appendix E.

## References

Lumley T (2010). Complex Surveys: a guide to analysis using R. Wiley

```
normalize Normalize numeric variable to 0-1 range
```


## Description

Performs a normalization of data, i.e., it scales variables in the range $0-1$. This is a special case of rescale(). unnormalize() is the counterpart, but only works for variables that have been normalized with normalize().

## Usage

normalize(x, ...)
\#\# S3 method for class 'numeric'
normalize(x, include_bounds = TRUE, verbose = TRUE, ...)
\#\# S3 method for class 'data.frame'
normalize(
x ,
select $=$ NULL,
exclude = NULL,
include_bounds = TRUE,
ignore_case = FALSE, verbose = TRUE,
)

```
unnormalize(x, ...)
## S3 method for class 'numeric'
unnormalize(x, verbose = TRUE, ...)
## S3 method for class 'data.frame'
unnormalize(
    x,
    select = NULL,
    exclude = NULL,
    ignore_case = FALSE,
    verbose = TRUE,
)
```


## Arguments

x
... Arguments passed to or from other methods.
include_bounds Logical, if TRUE, return value may include 0 and 1. If FALSE, the return value is compressed, using Smithson and Verkuilen's (2006) formula ( $x *(n-1$ ) + $0.5) / n$, to avoid zeros and ones in the normalized variables. This can be useful in case of beta-regression, where the response variable is not allowed to include zeros and ones.
verbose Toggle warnings and messages on or off.
select Variables that will be included when performing the required tasks. Can be either

- a variable specified as a literal variable name (e.g., column_name),
- a string with the variable name (e.g., "column_name"), or a character vector of variable names (e.g., c("col1", "col2", "col3")),
- a formula with variable names (e.g., ~column_1 + column_2),
- a vector of positive integers, giving the positions counting from the left (e.g. 1 or $c(1,3,5)$ ),
- a vector of negative integers, giving the positions counting from the right (e.g., -1 or $-1:-3$ ),
- one of the following select-helpers: starts_with(""), ends_with(""), contains(""), a range using : or regex (""),
- or a function testing for logical conditions, e.g. is.numeric() (or is.numeric), or any user-defined function that selects the variables for which the function returns TRUE (like: foo <- function (x) mean (x) > 3),
- ranges specified via literal variable names, select-helpers (except regex()) and (user-defined) functions can be negated, i.e. return non-matching elements, when prefixed with a -, e.g. -ends_with(""), -is.numeric or -Sepal.Width:Petal.Length. Note: Negation means that matches are excluded, and thus, the exclude argument can be used alternatively. For instance, select=-ends_with("Length") (with -) is equivalent to exclude=ends_with("Length")
(no -). In case negation should not work as expected, use the exclude argument instead.
If NULL, selects all columns. Patterns that found no matches are silently ignored, e.g. find_columns(iris, select $=c($ "Species", "Test")) will just return "Species".
exclude See select, however, column names matched by the pattern from exclude will be excluded instead of selected. If NULL (the default), excludes no columns.
ignore_case Logical, if TRUE and when one of the select-helpers or a regular expression is used in select, ignores lower/upper case in the search pattern when matching against variable names.


## Details

- If $x$ is a matrix, normalization is performed across all values (not column- or row-wise). For column-wise normalization, convert the matrix to a data.frame.
- If $x$ is a grouped data frame (grouped_df), normalization is performed separately for each group.


## Value

A normalized object.

## Selection of variables - the select argument

For most functions that have a select argument (including this function), the complete input data frame is returned, even when select only selects a range of variables. That is, the function is only applied to those variables that have a match in select, while all other variables remain unchanged. In other words: for this function, select will not omit any non-included variables, so that the returned data frame will include all variables from the input data frame.

## References

Smithson M, Verkuilen J (2006). A Better Lemon Squeezer? Maximum-Likelihood Regression with Beta-Distributed Dependent Variables. Psychological Methods, 11(1), 54-71.

## See Also

Other transform utilities: ranktransform(), rescale(), reverse(), standardize()

## Examples

```
normalize(c(0, 1, 5, -5, -2))
normalize(c(0, 1, 5, -5, -2), include_bounds = FALSE)
head(normalize(trees))
```


## Description

Transform numeric values with the integers of their rank (i.e., 1st smallest, 2nd smallest, 3rd smallest, etc.). Setting the sign argument to TRUE will give you signed ranks, where the ranking is done according to absolute size but where the sign is preserved (i.e., $2,1,-3,4$ ).

## Usage

```
ranktransform(x, ...)
    ## S3 method for class 'numeric'
    ranktransform(x, sign = FALSE, method = "average", verbose = TRUE, ...)
    ## S3 method for class 'data.frame'
    ranktransform(
    x,
    select = NULL,
    exclude = NULL,
    sign = FALSE,
    method = "average",
    ignore_case = FALSE,
    ...
    )
```


## Arguments

x
...
sign
method
verbose
select Variables that will be included when performing the required tasks. Can be either

- a variable specified as a literal variable name (e.g., column_name),
- a string with the variable name (e.g., "column_name"), or a character vector of variable names (e.g., c("col1", "col2", "col3")),
- a formula with variable names (e.g., $\sim$ column_1 + column_2),
- a vector of positive integers, giving the positions counting from the left (e.g. 1 or $c(1,3,5)$ ),
- a vector of negative integers, giving the positions counting from the right (e.g., -1 or $-1:-3$ ),
- one of the following select-helpers: starts_with(""), ends_with(""), contains(""), a range using : or regex(""),
- or a function testing for logical conditions, e.g. is.numeric() (or is.numeric), or any user-defined function that selects the variables for which the function returns TRUE (like: foo <- function $(x)$ mean $(x)>3$ ),
- ranges specified via literal variable names, select-helpers (except regex()) and (user-defined) functions can be negated, i.e. return non-matching elements, when prefixed with a -, e.g. -ends_with(""), -is.numeric or -Sepal.Width:Petal.Length. Note: Negation means that matches are excluded, and thus, the exclude argument can be used alternatively. For instance, select=-ends_with("Length") (with -) is equivalent to exclude=ends_with("Length") (no -). In case negation should not work as expected, use the exclude argument instead.

If NULL, selects all columns. Patterns that found no matches are silently ignored, e.g. find_columns(iris, select $=c($ "Species", "Test")) will just return "Species".
exclude See select, however, column names matched by the pattern from exclude will be excluded instead of selected. If NULL (the default), excludes no columns.
ignore_case Logical, if TRUE and when one of the select-helpers or a regular expression is used in select, ignores lower/upper case in the search pattern when matching against variable names.

## Value

A rank-transformed object.

## Selection of variables - the select argument

For most functions that have a select argument (including this function), the complete input data frame is returned, even when select only selects a range of variables. That is, the function is only applied to those variables that have a match in select, while all other variables remain unchanged. In other words: for this function, select will not omit any non-included variables, so that the returned data frame will include all variables from the input data frame.

## See Also

Other transform utilities: normalize(), rescale(), reverse(), standardize()

## Examples

```
ranktransform(c(0, 1, 5, -5, -2))
ranktransform(c(0, 1, 5, -5, -2), sign = TRUE)
head(ranktransform(trees))
```

```
remove_empty Return or remove variables or observations that are completely miss-
ing
```


## Description

These functions check which rows or columns of a data frame completely contain missing values, i.e. which observations or variables completely have missing values, and either (1) returns their indices; or (2) removes them from the data frame.

## Usage

empty_columns(x)
empty_rows(x)
remove_empty_columns(x)
remove_empty_rows(x)
remove_empty (x)

## Arguments

$x \quad$ A data frame.

## Value

- For empty_columns() and empty_rows(), a numeric (named) vector with row or column indices of those variables that completely have missing values.
- For remove_empty_columns() and remove_empty_rows(), a data frame with "empty" columns or rows removed, respectively.
- For remove_empty, both empty rows and columns will be removed.


## Examples

```
tmp <- data.frame(
    a = c(1, 2, 3, NA, 5),
    b = c(1, NA, 3, NA, 5),
    c = c(NA,NA,NA,NA,NA),
    d = c(1,NA, 3,NA, 5)
)
tmp
# indices of empty columns or rows
empty_columns(tmp)
empty_rows(tmp)
```

```
# remove empty columns or rows
remove_empty_columns(tmp)
remove_empty_rows(tmp)
# remove empty columns and rows
remove_empty(tmp)
```

```
replace_nan_inf Convert infinite or NaN values into NA
```


## Description

Replaces all infinite (Inf and -Inf) or NaN values with NA.

## Usage

replace_nan_inf(data)

## Arguments

data
A vector or a data frame.

## Value

Data with Inf, -Inf, and NaN converted to NA.

## Examples

```
# a vector
x <- c(1, 2, NA, 3, NaN, 4, NA, 5, Inf, -Inf, 6, 7)
replace_nan_inf(x)
# a data frame
df <- data.frame(
    x = c(1,NA, 5, Inf, 2, NA),
    y = c(3, NaN, 4, -Inf, 6, 7),
    stringsAsFactors = FALSE
)
replace_nan_inf(df)
```


## Description

Rescale variables to a new range. Can also be used to reverse-score variables (change the keying/scoring direction).

## Usage

```
rescale(x, ...)
change_scale(x, ...)
## S3 method for class 'numeric'
rescale(x, to = c(0, 100), range = NULL, verbose = TRUE, ...)
    ## S3 method for class 'data.frame'
    rescale(
        x,
        select = NULL,
    exclude = NULL,
    to = c(0, 100),
    range = NULL,
    ignore_case = FALSE,
    ...
    )
```


## Arguments

x
...
to $\quad$ Numeric vector of length 2 giving the new range that the variable will have after rescaling. To reverse-score a variable, the range should be given with the maximum value first. See examples.
range Initial (old) range of values. If NULL, will take the range of the input vector (range (x)).
verbose Toggle warnings.
select Variables that will be included when performing the required tasks. Can be either

- a variable specified as a literal variable name (e.g., column_name),
- a string with the variable name (e.g., "column_name"), or a character vector of variable names (e.g., c("col1", "col2", "col3")),
- a formula with variable names (e.g., ~column_1 + column_2),
- a vector of positive integers, giving the positions counting from the left (e.g. 1 or $c(1,3,5)$ ),
- a vector of negative integers, giving the positions counting from the right (e.g., -1 or $-1:-3$ ),
- one of the following select-helpers: starts_with(""), ends_with(""), contains(""), a range using : or regex (""),
- or a function testing for logical conditions, e.g. is.numeric() (or is .numeric), or any user-defined function that selects the variables for which the function returns TRUE (like: foo <- function (x) mean $(x)>3$ ),
- ranges specified via literal variable names, select-helpers (except regex()) and (user-defined) functions can be negated, i.e. return non-matching elements, when prefixed with a -, e.g. -ends_with(""), -is.numeric or -Sepal.Width:Petal.Length. Note: Negation means that matches are excluded, and thus, the exclude argument can be used alternatively. For instance, select=-ends_with("Length") (with -) is equivalent to exclude=ends_with("Length") (no -). In case negation should not work as expected, use the exclude argument instead.
If NULL, selects all columns. Patterns that found no matches are silently ignored, e.g. find_columns(iris, select = c("Species", "Test")) will just return "Species".
exclude See select, however, column names matched by the pattern from exclude will be excluded instead of selected. If NULL (the default), excludes no columns.
ignore_case Logical, if TRUE and when one of the select-helpers or a regular expression is used in select, ignores lower/upper case in the search pattern when matching against variable names.


## Value

A rescaled object.

## Selection of variables - the select argument

For most functions that have a select argument (including this function), the complete input data frame is returned, even when select only selects a range of variables. That is, the function is only applied to those variables that have a match in select, while all other variables remain unchanged. In other words: for this function, select will not omit any non-included variables, so that the returned data frame will include all variables from the input data frame.

## See Also

Other transform utilities: normalize(), ranktransform(), reverse(), standardize()

## Examples

```
rescale(c(0, 1, 5, -5, -2))
rescale(c(0, 1, 5, -5, -2), to = c(-5, 5))
rescale(c(1, 2, 3, 4, 5), to = c(-2, 2))
```

```
# Specify the "theoretical" range of the input vector
rescale(c(1, 3, 4), to = c(0, 40), range = c(0, 4))
# Reverse-score a variable
rescale(c(1, 2, 3, 4, 5), to = c(5, 1))
rescale(c(1, 2, 3, 4, 5), to = c(2, -2))
# Data frames
head(rescale(iris, to = c(0, 1)))
head(rescale(iris, to = c(0, 1), select = "Sepal.Length"))
# One can specify a list of ranges
head(rescale(iris, to = list(
    "Sepal.Length" = c(0, 1),
    "Petal.Length" = c(-1, 0)
)))
```

```
rescale_weights Rescale design weights for multilevel analysis
```


## Description

Most functions to fit multilevel and mixed effects models only allow to specify frequency weights, but not design (i.e. sampling or probability) weights, which should be used when analyzing complex samples and survey data. rescale_weights() implements an algorithm proposed by Asparouhov (2006) and Carle (2009) to rescale design weights in survey data to account for the grouping structure of multilevel models, which then can be used for multilevel modelling.

## Usage

rescale_weights(data, group, probability_weights, nest = FALSE)

## Arguments

data A data frame.
group Variable names (as character vector, or as formula), indicating the grouping structure (strata) of the survey data (level-2-cluster variable). It is also possible to create weights for multiple group variables; in such cases, each created weighting variable will be suffixed by the name of the group variable.
probability_weights
Variable indicating the probability (design or sampling) weights of the survey data (level-1-weight).
nest Logical, if TRUE and group indicates at least two group variables, then groups are "nested", i.e. groups are now a combination from each group level of the variables in group.

## Details

Rescaling is based on two methods: For pweights_a, the sample weights probability_weights are adjusted by a factor that represents the proportion of group size divided by the sum of sampling weights within each group. The adjustment factor for pweights_b is the sum of sample weights within each group divided by the sum of squared sample weights within each group (see Carle (2009), Appendix B). In other words, pweights_a "scales the weights so that the new weights sum to the cluster sample size" while pweights_b "scales the weights so that the new weights sum to the effective cluster size".
Regarding the choice between scaling methods A and B, Carle suggests that "analysts who wish to discuss point estimates should report results based on weighting method A. For analysts more interested in residual between-group variance, method B may generally provide the least biased estimates". In general, it is recommended to fit a non-weighted model and weighted models with both scaling methods and when comparing the models, see whether the "inferential decisions converge", to gain confidence in the results.
Though the bias of scaled weights decreases with increasing group size, method A is preferred when insufficient or low group size is a concern.
The group ID and probably PSU may be used as random effects (e.g. nested design, or group and PSU as varying intercepts), depending on the survey design that should be mimicked.

## Value

data, including the new weighting variables: pweights_a and pweights_b, which represent the rescaled design weights to use in multilevel models (use these variables for the weights argument).

## References

- Carle A.C. (2009). Fitting multilevel models in complex survey data with design weights: Recommendations. BMC Medical Research Methodology 9(49): 1-13
- Asparouhov T. (2006). General Multi-Level Modeling with Sampling Weights. Communications in Statistics - Theory and Methods 35: 439-460


## Examples

```
if (require("lme4")) {
    data(nhanes_sample)
    head(rescale_weights(nhanes_sample, "SDMVSTRA", "WTINT2YR"))
    # also works with multiple group-variables
    head(rescale_weights(nhanes_sample, c("SDMVSTRA", "SDMVPSU"), "WTINT2YR"))
    # or nested structures.
    x <- rescale_weights(
        data = nhanes_sample,
        group = c("SDMVSTRA", "SDMVPSU"),
        probability_weights = "WTINT2YR",
        nest = TRUE
    )
    head(x)
```

```
    nhanes_sample <- rescale_weights(nhanes_sample, "SDMVSTRA", "WTINT2YR")
    glmer(
        total ~ factor(RIAGENDR) * (log(age) + factor(RIDRETH1)) + (1 | SDMVPSU),
        family = poisson(),
        data = nhanes_sample,
        weights = pweights_a
    )
}
```

reshape_ci Reshape CI between wide/long formats

## Description

Reshape CI between wide/long formats.

## Usage

reshape_ci(x, ci_type = "CI")

## Arguments

x
A data frame containing columns named CI_low and CI_high (or similar, see ci_type).
ci_type String indicating the "type" (i.e. prefix) of the interval columns. Per easystats convention, confidence or credible intervals are named CI_low and CI_high, and the related ci_type would be "CI". If column names for other intervals differ, ci_type can be used to indicate the name, e.g. ci_type = "SI" can be used for support intervals, where the column names in the data frame would be SI_low and SI_high.

## Value

A data frame with columns corresponding to confidence intervals reshaped either to wide or long format.

## Examples

```
x <- data.frame(
    Parameter = c("Term 1", "Term 2", "Term 1", "Term 2"),
    CI = c(.8, .8, .9, .9),
    CI_low = c(.2, .3, .1, .15),
    CI_high = c(.5, .6, .8, .85),
    stringsAsFactors = FALSE
)
reshape_ci(x)
reshape_ci(reshape_ci(x))
```

reverse Reverse-Score Variables

## Description

Reverse-score variables (change the keying/scoring direction).

## Usage

```
reverse(x, ...)
reverse_scale(x, ...)
## S3 method for class 'numeric'
reverse(x, range = NULL, verbose = TRUE, ...)
    ## S3 method for class 'data.frame'
    reverse(
    x,
    select = NULL,
    exclude = NULL,
    range = NULL,
    ignore_case = FALSE,
    ...
    )
```


## Arguments

x
...
range Initial (old) range of values. If NULL, will take the range of the input vector (range (x)).
verbose Toggle warnings.
select Variables that will be included when performing the required tasks. Can be either

- a variable specified as a literal variable name (e.g., column_name),
- a string with the variable name (e.g., "column_name"), or a character vector of variable names (e.g., c("col1", "col2", "col3")),
- a formula with variable names (e.g., ~column_1 + column_2),
- a vector of positive integers, giving the positions counting from the left (e.g. 1 or $c(1,3,5)$ ),
- a vector of negative integers, giving the positions counting from the right (e.g., -1 or $-1:-3$ ),
- one of the following select-helpers: starts_with(""), ends_with(""), contains(""), a range using : or regex(""),
- or a function testing for logical conditions, e.g. is. numeric() (or is.numeric), or any user-defined function that selects the variables for which the function returns TRUE (like: foo <- function(x) mean (x) > 3),
- ranges specified via literal variable names, select-helpers (except regex ()) and (user-defined) functions can be negated, i.e. return non-matching elements, when prefixed with a -, e.g. -ends_with(""), -is.numeric or -Sepal.Width:Petal.Length. Note: Negation means that matches are excluded, and thus, the exclude argument can be used alternatively. For instance, select=-ends_with("Length") (with -) is equivalent to exclude=ends_with("Length") (no -). In case negation should not work as expected, use the exclude argument instead.
If NULL, selects all columns. Patterns that found no matches are silently ignored, e.g. find_columns(iris, select = c("Species", "Test")) will just return "Species".
exclude See select, however, column names matched by the pattern from exclude will be excluded instead of selected. If NULL (the default), excludes no columns.
ignore_case Logical, if TRUE and when one of the select-helpers or a regular expression is used in select, ignores lower/upper case in the search pattern when matching against variable names.


## Value

A reverse-scored object.

## Selection of variables - the select argument

For most functions that have a select argument (including this function), the complete input data frame is returned, even when select only selects a range of variables. That is, the function is only applied to those variables that have a match in select, while all other variables remain unchanged. In other words: for this function, select will not omit any non-included variables, so that the returned data frame will include all variables from the input data frame.

## See Also

Other transform utilities: normalize(), ranktransform(), rescale(), standardize()

## Examples

```
reverse(c(1, 2, 3, 4, 5))
reverse(c(-2, -1, 0, 2, 1))
# Specify the "theoretical" range of the input vector
reverse(c(1, 3, 4), range = c(0, 4))
# Factor variables
reverse(factor(c(1, 2, 3, 4, 5)))
reverse(factor(c(1, 2, 3, 4, 5)), range = 0:10)
# Data frames
```

```
head(reverse(iris))
head(reverse(iris, select = "Sepal.Length"))
```

```
rownames_as_column Tools for working with row names
```


## Description

Tools for working with row names

## Usage

```
rownames_as_column(x, var = "rowname")
    column_as_rownames(x, var = "rowname")
```


## Arguments

$x \quad$ A data frame.
var Name of column to use for rownames. For column_as_rownames(), this argument can be the variable name or the column number.

## Value

rownames_as_column() and column_as_rownames() both return a data frame.

## Examples

```
# Convert between row names and column ----------------------------------
test <- rownames_as_column(mtcars, var = "car")
test
head(column_as_rownames(test, var = "car"))
```

```
row_to_colnames Tools for working with column names
```


## Description

Tools for working with column names

## Usage

```
row_to_colnames(x, row = 1, na_prefix = "x", verbose = TRUE)
colnames_to_row(x, prefix = "x")
```


## Arguments

X
row Row to use as column names.
na_prefix Prefix to give to the column name if the row has an NA. Default is ' $x$ ', and it will be incremented at each NA (x1, x2, etc.).
verbose Toggle warnings.
prefix Prefix to give to the column name. Default is ' $x$ ', and it will be incremented at each column ( $\mathrm{x} 1, \mathrm{x} 2$, etc.).

## Value

row_to_colnames() and colnames_to_row() both return a data frame.

## Examples

```
# Convert a row to column names -----------------------------------
test <- data.frame(
    a = c("iso", 2, 5),
    b = c("year", 3, 6),
    c = c(NA, 5, 7)
)
test
row_to_colnames(test)
# Convert column names to row -----------------------------------
test <- data.frame(
        ARG = c("BRA", "FRA"),
        `1960` = c(1960, 1960),
        `2000` = c(2000, 2000)
    )
    test
    colnames_to_row(test)
```

    skewness Compute Skewness and (Excess) Kurtosis
    
## Description

Compute Skewness and (Excess) Kurtosis

## Usage

```
skewness(x, na.rm = TRUE, type = "2", iterations = NULL, verbose = TRUE, ...)
kurtosis(x, na.rm = TRUE, type = "2", iterations = NULL, verbose = TRUE, ...)
```

```
## S3 method for class 'parameters_kurtosis'
print(x, digits = 3, test = FALSE, ...)
## S3 method for class 'parameters_skewness'
print(x, digits = 3, test = FALSE, ...)
## S3 method for class 'parameters_skewness'
summary(object, test = FALSE, ...)
## S3 method for class 'parameters_kurtosis'
summary(object, test = FALSE, ...)
```


## Arguments

| x | A numeric vector or data.frame. |
| :--- | :--- |
| na.rm | Remove missing values. |
| type | Type of algorithm for computing skewness. May be one of 1 (or " " ", "I" <br> or "classic"), 2 (or "2", "II" or "SPSS" or "SAS") or 3 (or " " ", "III" or <br> "Minitab"). See 'Details'. |
| iterations | The number of bootstrap replicates for computing standard errors. If NULL (de- <br> fault), parametric standard errors are computed. |
| verbose | Toggle warnings and messages. |
| $\ldots$ | Arguments passed to or from other methods. |
| digits | Number of decimal places. |
| test | Logical, if TRUE, tests if skewness or kurtosis is significantly different from zero. |
| object | An object returned by skewness() or kurtosis(). |

## Details

Skewness: Symmetric distributions have a skewness around zero, while a negative skewness values indicates a "left-skewed" distribution, and a positive skewness values indicates a "rightskewed" distribution. Examples for the relationship of skewness and distributions are:

- Normal distribution (and other symmetric distribution) has a skewness of 0
- Half-normal distribution has a skewness just below 1
- Exponential distribution has a skewness of 2
- Lognormal distribution can have a skewness of any positive value, depending on its parameters
(https://en.wikipedia.org/wiki/Skewness)
Types of Skewness: skewness() supports three different methods for estimating skewness, as discussed in Joanes and Gill (1988):
- Type " 1 " is the "classical" method, which is $\mathrm{g} 1=\left(\operatorname{sum}\left((x-\operatorname{mean}(x))^{\wedge} 3\right) / n\right) /(\operatorname{sum}((x-$ mean $\left.\left.(x))^{\wedge} 2\right) / n\right)^{\wedge} 1.5$
- Type " 2 " first calculates the type-1 skewness, then adjusts the result: G1 $=\mathrm{g} 1 * \operatorname{sqrt}(\mathrm{n}$ * ( n $-1)) /(n-2)$. This is what SAS and SPSS usually return
- Type " 3 " first calculates the type-1 skewness, then adjusts the result: b1 = g1 * ( ( $1-1$ / $n))^{\wedge} 1.5$. This is what Minitab usually returns.

Kurtosis: The kurtosis is a measure of "tailedness" of a distribution. A distribution with a kurtosis values of about zero is called "mesokurtic". A kurtosis value larger than zero indicates a "leptokurtic" distribution with fatter tails. A kurtosis value below zero indicates a "platykurtic" distribution with thinner tails (https://en.wikipedia.org/wiki/Kurtosis).

Types of Kurtosis: kurtosis() supports three different methods for estimating kurtosis, as discussed in Joanes and Gill (1988):

- Type " 1 " is the "classical" method, which is $g 2=n * \operatorname{sum}\left((x-\operatorname{mean}(x))^{\wedge} 4\right) /(\operatorname{sum}((x-$ mean $\left.(x))^{\wedge} 2\right)^{\wedge} 2$ ) - 3 .
- Type "2" first calculates the type-1 kurtosis, than adjusts the result: $\mathrm{G} 2=((\mathrm{n}+1) * \mathrm{~g} 2+6)$ * $(n-1) /((n-2) *(n-3))$. This is what SAS and SPSS usually return
- Type " 3 " first calculates the type-1 kurtosis, than adjusts the result: $\mathrm{b} 2=(\mathrm{g} 2+3)$ * $(1-1 /$ $n)^{\wedge} 2-3$. This is what Minitab usually returns.

Standard Errors: It is recommended to compute empirical (bootstrapped) standard errors (via the iterations argument) than relying on analytic standard errors (Wright \& Herrington, 2011).

## Value

Values of skewness or kurtosis.

## References

- D. N. Joanes and C. A. Gill (1998). Comparing measures of sample skewness and kurtosis. The Statistician, 47, 183-189.
- Wright, D. B., \& Herrington, J. A. (2011). Problematic standard errors and confidence intervals for skewness and kurtosis. Behavior research methods, 43(1), 8-17.


## Examples

skewness(rnorm(1000))
kurtosis(rnorm(1000))

> slide

Shift numeric value range

## Description

This functions shifts the value range of a numeric variable, so that the new range starts at a given value.

## Usage

```
slide(x, ...)
    data_shift(x, ...)
    \#\# S3 method for class 'numeric'
    slide(x, lowest = 0, ...)
    \#\# S3 method for class 'data.frame'
    slide(
        x ,
        select \(=\) NULL,
        exclude \(=\) NULL,
        lowest = 0,
        append = FALSE,
        ignore_case = FALSE,
        verbose = TRUE,
    )
```


## Arguments

X
...
lowest Numeric, indicating the lowest (minimum) value when converting factors or character vectors to numeric values.
select Variables that will be included when performing the required tasks. Can be either

- a variable specified as a literal variable name (e.g., column_name),
- a string with the variable name (e.g., "column_name"), or a character vector of variable names (e.g., c("col1", "col2", "col3")),
- a formula with variable names (e.g., $\sim$ column_1 + column_2),
- a vector of positive integers, giving the positions counting from the left (e.g. 1 or $c(1,3,5)$ ),
- a vector of negative integers, giving the positions counting from the right (e.g., -1 or -1 : -3 ),
- one of the following select-helpers: starts_with(""), ends_with(""), contains(""), a range using : or regex(""),
- or a function testing for logical conditions, e.g. is. numeric() (or is.numeric), or any user-defined function that selects the variables for which the function returns TRUE (like: foo <- function ( $x$ ) mean $(x)>3$ ),
- ranges specified via literal variable names, select-helpers (except regex ()) and (user-defined) functions can be negated, i.e. return non-matching elements, when prefixed with a -, e.g. -ends_with(""), -is.numeric or -Sepal.Width:Petal.Length. Note: Negation means that matches are
excluded, and thus, the exclude argument can be used alternatively. For instance, select=-ends_with("Length") (with -) is equivalent to exclude=ends_with("Length") (no -). In case negation should not work as expected, use the exclude argument instead.
If NULL, selects all columns. Patterns that found no matches are silently ignored, e.g. find_columns(iris, select $=c($ "Species", "Test")) will just return "Species".
exclude See select, however, column names matched by the pattern from exclude will be excluded instead of selected. If NULL (the default), excludes no columns.
append Logical or string. If TRUE, recoded or converted variables get new column names and are appended (column bind) to $x$, thus returning both the original and the recoded variables. The new columns get a suffix, based on the calling function: "_r" for recode functions, "_n" for to_numeric(), "_f" for to_factor(), or "_s" for slide(). If append=FALSE, original variables in $x$ will be overwritten by their recoded versions. If a character value, recoded variables are appended with new column names (using the defined suffix) to the original data frame.
ignore_case Logical, if TRUE and when one of the select-helpers or a regular expression is used in select, ignores lower/upper case in the search pattern when matching against variable names.
verbose Toggle warnings.


## Value

$x$, where the range of numeric variables starts at a new value.

## Selection of variables - the select argument

For most functions that have a select argument (including this function), the complete input data frame is returned, even when select only selects a range of variables. That is, the function is only applied to those variables that have a match in select, while all other variables remain unchanged. In other words: for this function, select will not omit any non-included variables, so that the returned data frame will include all variables from the input data frame.

## See Also

- Functions to rename stuff: data_rename(), data_rename_rows(), data_addprefix(), data_addsuffix()
- Functions to reorder or remove columns: data_reorder(), data_relocate(), data_remove()
- Functions to reshape, pivot or rotate data frames: data_to_long(), data_to_wide(), data_rotate()
- Functions to recode data: rescale(), reverse(), categorize(), change_code(), slide()
- Functions to standardize, normalize, rank-transform: center(), standardize(), normalize(), ranktransform(), winsorize()
- Split and merge data frames: data_partition(), data_merge()
- Functions to find or select columns: data_select(), data_find()
- Functions to filter rows: data_match(), data_filter()


## Examples

```
# numeric
head(mtcars$gear)
head(slide(mtcars$gear))
head(slide(mtcars$gear, lowest = 10))
# data frame
sapply(slide(mtcars, lowest = 1), min)
sapply(mtcars, min)
```

```
smoothness
Quantify the smoothness of a vector
```


## Description

Quantify the smoothness of a vector

## Usage

smoothness(x, method = "cor", lag = 1, iterations = NULL, ...)

## Arguments

| x | Numeric vector (similar to a time series). |
| :--- | :--- |
| method | Can be "diff" (the standard deviation of the standardized differences) or "cor" <br> (default, lag-one autocorrelation). |
| lag | An integer indicating which lag to use. If less than 1, will be interpreted as <br> expressed in percentage of the length of the vector. |
| iterations | The number of bootstrap replicates for computing standard errors. If NULL (de- <br> fault), parametric standard errors are computed. |
| $\ldots$ | Arguments passed to or from other methods. |

## Value

Value of smoothness.

## References

https://stats.stackexchange.com/questions/24607/how-to-measure-smoothness-of-a-time-series-in-r

## Examples

```
x <- (-10:10)^3 + rnorm(21, 0, 100)
plot(x)
smoothness(x, method = "cor")
smoothness(x, method = "diff")
```


## Description

Performs a standardization of data (z-scoring), i.e., centering and scaling, so that the data is expressed in terms of standard deviation (i.e., mean $=0, \mathrm{SD}=1$ ) or Median Absolute Deviance (median $=0, \mathrm{MAD}=1$ ). When applied to a statistical model, this function extracts the dataset, standardizes it, and refits the model with this standardized version of the dataset. The normalize() function can also be used to scale all numeric variables within the 0-1 range.

For model standardization, see standardize. default().

## Usage

standardize(x, ...)
standardise(x, ...)
\#\# S3 method for class 'numeric'
standardize(
x ,
robust = FALSE,
two_sd = FALSE,
weights = NULL,
reference = NULL,
center = NULL,
scale = NULL,
verbose = TRUE,
)
\#\# S3 method for class 'factor'
standardize(
x ,
robust $=$ FALSE,
two_sd = FALSE,
weights = NULL,
force = FALSE,
verbose = TRUE,
)
\#\# S3 method for class 'data.frame'
standardize(
x ,
select $=$ NULL,

```
    exclude = NULL,
    robust = FALSE,
    two_sd = FALSE,
    weights = NULL,
    reference = NULL,
    center = NULL,
    scale = NULL,
    remove_na = c("none", "selected", "all"),
    force = FALSE,
    append = FALSE,
    ignore_case = FALSE,
    verbose = TRUE,
)
unstandardize(x, ...)
unstandardise(x, ...)
## S3 method for class 'numeric'
unstandardize(
    x,
    center = NULL,
    scale = NULL,
    reference = NULL,
    robust = FALSE,
    two_sd = FALSE,
)
## S3 method for class 'data.frame'
unstandardize(
    x,
    center = NULL,
    scale = NULL,
    reference = NULL,
    robust = FALSE,
    two_sd = FALSE,
    select = NULL,
    exclude = NULL,
    ...
)
```


## Arguments

x
A (grouped) data frame, a vector or a statistical model (for unstandardize() cannot be a model).
... Arguments passed to or from other methods.

| robust | Logical, if TRUE, centering is done by subtracting the median from the variables and dividing it by the median absolute deviation (MAD). If FALSE, variables are standardized by subtracting the mean and dividing it by the standard deviation (SD). |
| :---: | :---: |
| two_sd | If TRUE, the variables are scaled by two times the deviation (SD or MAD depending on robust). This method can be useful to obtain model coefficients of continuous parameters comparable to coefficients related to binary predictors, when applied to the predictors (not the outcome) (Gelman, 2008). |
| weights | Can be NULL (for no weighting), or: <br> - For model: if TRUE (default), a weighted-standardization is carried out. <br> - For data. frames: a numeric vector of weights, or a character of the name of a column in the data. frame that contains the weights. <br> - For numeric vectors: a numeric vector of weights. |
| reference | A data frame or variable from which the centrality and deviation will be computed instead of from the input variable. Useful for standardizing a subset or new data according to another data frame. |
| center, scale | - For standardize(): <br> Numeric values, which can be used as alternative to reference to define a reference centrality and deviation. If scale and center are of length 1 , they will be recycled to match the length of selected variables for standardization. Else, center and scale must be of same length as the number of selected variables. Values in center and scale will be matched to selected variables in the provided order, unless a named vector is given. In this case, names are matched against the names of the selected variables. <br> - For unstandardize(): center and scale correspond to the center (the mean / median) and the scale (SD / MAD) of the original non-standardized data (for data frames, should be named, or have column order correspond to the numeric column). However, one can also directly provide the original data through reference, from which the center and the scale will be computed (according to robust and two_sd). Alternatively, if the input contains the attributes center and scale (as does the output of standardize()), it will take it from there if the rest of the arguments are absent. |
| verbose | Toggle warnings and messages on or off. |
| force | Logical, if TRUE, forces recoding of factors and character vectors as well. |
| select | Variables that will be included when performing the required tasks. Can be either |
|  | - a variable specified as a literal variable name (e.g., column_name), <br> - a string with the variable name (e.g., "column_name"), or a character vector of variable names (e.g., c("col1", "col2", "col3")), <br> - a formula with variable names (e.g., ~column_1 + column_2), <br> - a vector of positive integers, giving the positions counting from the left (e.g. 1 or $c(1,3,5)$ ), |
|  | - a vector of negative integers, giving the positions counting from the right (e.g., -1 or -1 : -3 ), |

- one of the following select-helpers: starts_with(""), ends_with(""), contains(""), a range using : or regex(""),
- or a function testing for logical conditions, e.g. is.numeric() (or is .numeric), or any user-defined function that selects the variables for which the function returns TRUE (like: foo <- function (x) mean $(x)>3$ ),
- ranges specified via literal variable names, select-helpers (except regex()) and (user-defined) functions can be negated, i.e. return non-matching elements, when prefixed with a -, e.g. -ends_with(""), -is.numeric or -Sepal.Width:Petal.Length. Note: Negation means that matches are excluded, and thus, the exclude argument can be used alternatively. For instance, select=-ends_with("Length") (with -) is equivalent to exclude=ends_with("Length") (no -). In case negation should not work as expected, use the exclude argument instead.

If NULL, selects all columns. Patterns that found no matches are silently ignored, e.g. find_columns(iris, select = c("Species", "Test")) will just return "Species".
exclude See select, however, column names matched by the pattern from exclude will be excluded instead of selected. If NULL (the default), excludes no columns.
remove_na How should missing values (NA) be treated: if "none" (default): each column's standardization is done separately, ignoring NAs. Else, rows with NA in the columns selected with select / exclude ("selected") or in all columns ("all") are dropped before standardization, and the resulting data frame does not include these cases.
append Logical or string. If TRUE, standardized variables get new column names (with the suffix " $\quad z$ ") and are appended (column bind) to $x$, thus returning both the original and the standardized variables. If FALSE, original variables in $x$ will be overwritten by their standardized versions. If a character value, standardized variables are appended with new column names (using the defined suffix) to the original data frame.
ignore_case Logical, if TRUE and when one of the select-helpers or a regular expression is used in select, ignores lower/upper case in the search pattern when matching against variable names.

## Value

The standardized object (either a standardize data frame or a statistical model fitted on standardized data).

## Selection of variables - the select argument

For most functions that have a select argument (including this function), the complete input data frame is returned, even when select only selects a range of variables. That is, the function is only applied to those variables that have a match in select, while all other variables remain unchanged. In other words: for this function, select will not omit any non-included variables, so that the returned data frame will include all variables from the input data frame.

## Note

When $x$ is a vector or a data frame with remove_na = "none"), missing values are preserved, so the return value has the same length / number of rows as the original input.

## See Also

See center () for grand-mean centering of variables.
Other transform utilities: normalize(), ranktransform(), rescale(), reverse()
Other standardize: standardize.default()

## Examples

```
d <- iris[1:4, ]
# vectors
standardise(d$Petal.Length)
# Data frames
# overwrite
standardise(d, select = c("Sepal.Length", "Sepal.Width"))
# append
standardise(d, select = c("Sepal.Length", "Sepal.Width"), append = TRUE)
# append, suffix
standardise(d, select = c("Sepal.Length", "Sepal.Width"), append = "_std")
# standardizing with reference center and scale
d <- data.frame(
    a = c(-2, -1, 0, 1, 2),
    b = c(3, 4, 5, 6, 7)
)
# default standardization, based on mean and sd of each variable
standardize(d) # means are 0 and 5, sd ~ 1.581139
# standardization, based on mean and sd set to the same values
standardize(d, center = c(0, 5), scale = c(1.581, 1.581))
# standardization, mean and sd for each variable newly defined
standardize(d, center = c(3, 4), scale = c(2, 4))
# standardization, taking same mean and sd for each variable
standardize(d, center = 1, scale = 3)
```


## Description

Performs a standardization of data (z-scoring) using standardize() and then re-fits the model to the standardized data.

Standardization is done by completely refitting the model on the standardized data. Hence, this approach is equal to standardizing the variables before fitting the model and will return a new model object. This method is particularly recommended for complex models that include interactions or transformations (e.g., polynomial or spline terms). The robust (default to FALSE) argument enables a robust standardization of data, based on the median and the MAD instead of the mean and the SD.

```
Usage
    ## Default S3 method:
    standardize(
        x,
        robust = FALSE,
        two_sd = FALSE,
        weights = TRUE,
        verbose = TRUE,
        include_response = TRUE,
    )
```


## Arguments

x
robust
two_sd
weights If TRUE (default), a weighted-standardization is carried out.
verbose Toggle warnings and messages on or off.
include_response
If TRUE (default), the response value will also be standardized. If FALSE, only the predictors will be standardized.

- Note that for GLMs and models with non-linear link functions, the response value will not be standardized, to make re-fitting the model work.
- If the model contains an stats: :offset(), the offset variable(s) will be standardized only if the response is standardized. If two_sd = TRUE, offsets are standardized by one-sd (similar to the response).
- (For mediate models, the include_response refers to the outcome in the y model; m model's response will always be standardized when possible).
... Arguments passed to or from other methods.

Value
A statistical model fitted on standardized data

## Generalized Linear Models

Standardization for generalized linear models (GLM, GLMM, etc) is done only with respect to the predictors (while the outcome remains as-is, unstandardized) - maintaining the interpretability of the coefficients (e.g., in a binomial model: the exponent of the standardized parameter is the OR of a change of 1 SD in the predictor, etc.)

## Dealing with Factors

standardize(model) or standardize_parameters(model, method = "refit") do not standardize categorical predictors (i.e. factors) / their dummy-variables, which may be a different behaviour compared to other R packages (such as Im.beta) or other software packages (like SPSS). To mimic such behaviours, either use standardize_parameters(model, method = "basic") to obtain posthoc standardized parameters, or standardize the data with standardize (data, force = TRUE) before fitting the model.

## Transformed Variables

When the model's formula contains transformations (e.g. y $\sim \exp (X)$ ) the transformation effectively takes place after standardization (e.g., $\exp (\operatorname{scale}(X))$ ). Since some transformations are undefined for none positive values, such as $\log ()$ and $\operatorname{sqrt}()$, the relevel variables are shifted (post standardization) by $Z-\min (Z)+1$ or $Z-\min (Z)$ (respectively).

## See Also

Other standardize: standardize()

## Examples

```
model <- lm(Infant.Mortality ~ Education * Fertility, data = swiss)
coef(standardize(model))
```

    to_factor Convert data to factors
    
## Description

Convert data to factors

## Usage

```
to_factor (x, ...)
data_to_factor(x, ...)
\#\# S3 method for class 'data.frame'
to_factor (
        x,
        select \(=\) NULL,
        exclude \(=\) NULL,
        ignore_case = FALSE,
        append = FALSE,
        verbose = TRUE,
)
```


## Arguments

x
...
select Variables that will be included when performing the required tasks. Can be either

- a variable specified as a literal variable name (e.g., column_name),
- a string with the variable name (e.g., "column_name"), or a character vector of variable names (e.g., c("col1", "col2", "col3")),
- a formula with variable names (e.g., ~column_1 + column_2),
- a vector of positive integers, giving the positions counting from the left (e.g. 1 or $c(1,3,5)$ ),
- a vector of negative integers, giving the positions counting from the right (e.g., -1 or $-1:-3$ ),
- one of the following select-helpers: starts_with(""), ends_with(""), contains(""), a range using : or regex (""),
- or a function testing for logical conditions, e.g. is. numeric() (or is.numeric), or any user-defined function that selects the variables for which the function returns TRUE (like: foo <- function(x) mean (x) > 3),
- ranges specified via literal variable names, select-helpers (except regex ()) and (user-defined) functions can be negated, i.e. return non-matching elements, when prefixed with a -, e.g. -ends_with(""), -is.numeric or -Sepal.Width:Petal.Length. Note: Negation means that matches are excluded, and thus, the exclude argument can be used alternatively. For instance, select=-ends_with("Length") (with -) is equivalent to exclude=ends_with("Length") (no -). In case negation should not work as expected, use the exclude argument instead.

If NULL, selects all columns. Patterns that found no matches are silently ignored, e.g. find_columns(iris, select = c("Species", "Test")) will just return "Species".

| exclude | See select, however, column names matched by the pattern from exclude will be excluded instead of selected. If NULL (the default), excludes no columns. |
| :---: | :---: |
| ignore_case | Logical, if TRUE and when one of the select-helpers or a regular expression is used in select, ignores lower/upper case in the search pattern when matching against variable names. |
| append | Logical or string. If TRUE, recoded or converted variables get new column names and are appended (column bind) to $x$, thus returning both the original and the recoded variables. The new columns get a suffix, based on the calling function: "_r" for recode functions, "_n" for to_numeric(), "_f" for to_factor(), or "_s" for slide(). If append=FALSE, original variables in $x$ will be overwritten by their recoded versions. If a character value, recoded variables are appended with new column names (using the defined suffix) to the original data frame. |
| verbose | Toggle warnings. |

## Details

Convert data to numeric by converting characters to factors and factors to either numeric levels or dummy variables. The "counterpart" to convert variables into numeric is to_numeric().

## Value

A factor, or a data frame of factors.

## Selection of variables - the select argument

For most functions that have a select argument (including this function), the complete input data frame is returned, even when select only selects a range of variables. That is, the function is only applied to those variables that have a match in select, while all other variables remain unchanged. In other words: for this function, select will not omit any non-included variables, so that the returned data frame will include all variables from the input data frame.

## Examples

```
str(to_factor(iris))
```

\# use labels as levels
data(efc)
str (efc\$c172code)
head(to_factor(efc\$c172code))

```
to_numeric Convert data to numeric
```


## Description

Convert data to numeric by converting characters to factors and factors to either numeric levels or dummy variables. The "counterpart" to convert variables into factors is to_factor().

## Usage

```
to_numeric (x, ...)
data_to_numeric(x, ...)
\#\# S3 method for class 'data.frame'
to_numeric(
        x ,
        select \(=\) NULL,
        exclude \(=\) NULL,
        dummy_factors = TRUE,
        preserve_levels = FALSE,
        lowest = NULL,
        append = FALSE,
        ignore_case = FALSE,
        verbose = TRUE,
)
```


## Arguments

x
...
select Variables that will be included when performing the required tasks. Can be either

- a variable specified as a literal variable name (e.g., column_name),
- a string with the variable name (e.g., "column_name"), or a character vector of variable names (e.g., c("col1", "col2", "col3")),
- a formula with variable names (e.g., ~column_1 + column_2),
- a vector of positive integers, giving the positions counting from the left (e.g. 1 or c(1, 3, 5)),
- a vector of negative integers, giving the positions counting from the right (e.g., -1 or $-1:-3$ ),
- one of the following select-helpers: starts_with(""), ends_with(""), contains(""), a range using : or regex(""),
- or a function testing for logical conditions, e.g. is.numeric() (or is.numeric), or any user-defined function that selects the variables for which the function returns TRUE (like: foo <- function(x) mean (x) > 3),
- ranges specified via literal variable names, select-helpers (except regex()) and (user-defined) functions can be negated, i.e. return non-matching elements, when prefixed with a -, e.g. -ends_with(""), -is.numeric or -Sepal.Width:Petal.Length. Note: Negation means that matches are excluded, and thus, the exclude argument can be used alternatively. For instance, select=-ends_with("Length") (with -) is equivalent to exclude=ends_with("Length") (no -). In case negation should not work as expected, use the exclude argument instead.

If NULL, selects all columns. Patterns that found no matches are silently ignored, e.g. find_columns(iris, select = c("Species", "Test")) will just return "Species".
exclude See select, however, column names matched by the pattern from exclude will be excluded instead of selected. If NULL (the default), excludes no columns.
dummy_factors Transform factors to dummy factors (all factor levels as different columns filled with a binary $0-1$ value).
preserve_levels
Logical, only applies if $x$ is a factor. If TRUE, and $x$ has numeric factor levels, these will be converted into the related numeric values. If this is not possible, the converted numeric values will start from 1 to number of levels.
lowest Numeric, indicating the lowest (minimum) value when converting factors or character vectors to numeric values.
append Logical or string. If TRUE, recoded or converted variables get new column names and are appended (column bind) to $x$, thus returning both the original and the recoded variables. The new columns get a suffix, based on the calling function: "_r" for recode functions, "_n" for to_numeric(), "_f" for to_factor(), or "_s" for slide(). If append=FALSE, original variables in $x$ will be overwritten by their recoded versions. If a character value, recoded variables are appended with new column names (using the defined suffix) to the original data frame.
ignore_case Logical, if TRUE and when one of the select-helpers or a regular expression is used in select, ignores lower/upper case in the search pattern when matching against variable names.
verbose Toggle warnings.

## Value

A data frame of numeric variables.

## Selection of variables - select argument

For most functions that have a select argument the complete input data frame is returned, even when select only selects a range of variables. However, for to_numeric(), factors might be converted into dummies, thus, the number of variables of the returned data frame no longer match the input data frame. Hence, when select is used, only those variables (or their dummies) specified in select will be returned. Use append=TRUE to also include the original variables in the returned data frame.

## Examples

```
to_numeric(head(ToothGrowth))
to_numeric(head(ToothGrowth), dummy_factors = FALSE)
# factors
x <- as.factor(mtcars$gear)
to_numeric(x, dummy_factors = FALSE)
to_numeric(x, dummy_factors = FALSE, preserve_levels = TRUE)
```


## Description

This function prepares objects for visualisation by returning a list of layers with data and geoms that can be easily plotted using for instance ggplot2.
If the see package is installed, the call to visualization_recipe() can be replaced by plot(), which will internally call the former and then plot it using ggplot. The resulting plot can be customized ad-hoc (by adding ggplot's geoms, theme or specifications), or via some of the arguments of visualisation_recipe() that control the aesthetic parameters.
See the specific documentation page for your object's class:

- modelbased: https://easystats.github.io/modelbased/reference/visualisation_recipe. estimate_predicted.html
- correlation: https://easystats.github.io/correlation/reference/visualisation_recipe. easycormatrix.html


## Usage

visualisation_recipe(x, ...)

## Arguments

$x \quad$ An easystats object.
$\ldots \quad$ Other arguments passed to other functions.
weighted_mean Weighted Mean, Median, SD, and MAD

## Description

Weighted Mean, Median, SD, and MAD

## Usage

```
    weighted_mean(x, weights \(=\) NULL, verbose \(=\) TRUE, ...)
    weighted_median( \(x\), weights \(=\) NULL, verbose \(=\) TRUE, ...)
    weighted_sd(x, weights = NULL, verbose = TRUE, ...)
    weighted_mad( x , weights \(=\) NULL, constant \(=1.4826\), verbose \(=\) TRUE,.. .)
```


## Arguments

X
weights A numerical vector of weights the same length as $x$ giving the weights to use for elements of $x$.
verbose Show warning when weights are negative? If weights $=$ NULL, $x$ is passed to the non-weighted function.
... arguments to be passed to or from methods.
constant scale factor.

## Examples

```
## GPA from Siegel 1994
x <- c(3.7, 3.3, 3.5, 2.8)
wt <- c(5, 5, 4, 1) / 15
weighted_mean(x, wt)
weighted_median(x, wt)
weighted_sd(x, wt)
weighted_mad(x, wt)
```

winsorize Winsorize data

## Description

Winsorize data

## Usage

```
winsorize(data, ...)
## S3 method for class 'numeric'
winsorize(
        data,
        threshold = 0.2,
        method = "percentile",
        robust = FALSE,
        verbose = TRUE,
    )
```


## Arguments

data data frame or vector.
... Currently not used.
threshold The amount of winsorization, depends on the value of method:

- For method = "percentile": the amount to winsorize from each tail.
- For method = "zscore": the number of SD/MAD-deviations from the mean/median (see robust)
- For method = "raw": a vector of length 2 with the lower and upper bound for winsorization.
method One of "percentile" (default), "zscore", or "raw".
robust Logical, if TRUE, winsorizing through the "zscore" method is done via the median and the median absolute deviation (MAD); if FALSE, via the mean and the standard deviation.
verbose Toggle warnings.


## Details

Winsorizing or winsorization is the transformation of statistics by limiting extreme values in the statistical data to reduce the effect of possibly spurious outliers. The distribution of many statistics can be heavily influenced by outliers. A typical strategy is to set all outliers (values beyond a certain threshold) to a specified percentile of the data; for example, a $90 \backslash$ to the 5 th percentile, and data above the 95 th percentile set to the 95 th percentile. Winsorized estimators are usually more robust to outliers than their more standard forms.

## Value

A data frame with winsorized columns or a winsorized vector.

## See Also

- Functions to rename stuff: data_rename(), data_rename_rows(), data_addprefix(), data_addsuffix()
- Functions to reorder or remove columns: data_reorder(), data_relocate(), data_remove()
- Functions to reshape, pivot or rotate data frames: data_to_long(), data_to_wide(), data_rotate()
- Functions to recode data: rescale(), reverse(), categorize(), change_code(), slide()
- Functions to standardize, normalize, rank-transform: center(), standardize(), normalize(), ranktransform(), winsorize()
- Split and merge data frames: data_partition(), data_merge()
- Functions to find or select columns: data_select(), data_find()
- Functions to filter rows: data_match(), data_filter()


## Examples

```
hist(iris$Sepal.Length, main = "Original data")
hist(winsorize(iris$Sepal.Length, threshold = 0.2),
    xlim = c(4, 8), main = "Percentile Winsorization"
)
hist(winsorize(iris$Sepal.Length, threshold = 1.5, method = "zscore"),
    xlim = c(4, 8), main = "Mean (+/- SD) Winsorization"
)
hist(winsorize(iris$Sepal.Length, threshold = 1.5, method = "zscore", robust = TRUE),
    xlim = c(4, 8), main = "Median (+/- MAD) Winsorization"
)
hist(winsorize(iris$Sepal.Length, threshold = c(5, 7.5), method = "raw"),
    xlim = c(4, 8), main = "Raw Thresholds"
)
\# Also works on a data frame:
winsorize(iris, threshold \(=0.2\) )
```


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