Package ‘gcplyr’

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Type Package

Title Manipulate and Analyze Growth Curve Data

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Description Easy import, manipulation, and model-free analysis of microbial growth curve data, as commonly output by plate readers. Tools for reshaping common plate reader outputs into 'tidy' formats and merging them with design information, making data easy to work with using 'gcplyr' and other packages. Also streamlines common growth curve processing steps, like smoothing and calculating derivatives, and facilitates model-free characterization and analysis of growth data. See methods at <https://mikeblazanin.github.io/gcplyr/>.

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https://github.com/mikeblazanin/gcplyr/

Depends R (>= 2.10)

Imports dplyr, stats, tidyr, tools, utils

Suggests cowplot, ggplot2, knitr, lubridate, mgcv, readxl, rmarkdown, testthat (>= 3.0.0), xlsx

VignetteBuilder knitr

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NeedsCompilation no

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 auc

Calculate area under the curve

Description

This function takes a vector of x and y values and returns a scalar for the area under the curve, calculated using the trapezoid rule.
Usage

```r
auc(x, y, xlim = NULL, blank = 0, na.rm = TRUE, neg.rm = FALSE)
```

Arguments

- `x`: Numeric vector of x values
- `y`: Numeric vector of y values
- `xlim`: Vector, of length 2, delimiting the x range over which the area under the curve should be calculated (where NA can be provided for the area to be calculated from the start or to the end of the data)
- `blank`: Value to be subtracted from y values before calculating area under the curve
- `na.rm`: a logical indicating whether missing values should be removed
- `neg.rm`: a logical indicating whether y values below zero should be treated as zeros. If FALSE, area under the curve for negative y values will be calculated normally, effectively subtracting from the returned value.

Details

This function is designed to be compatible for use within `dplyr::group_by` and `dplyr::summarize`

Value

A scalar for the total area under the curve

---

**block_tidydesign**  
*Turn tidydesign into block format*

Description

This function allows users to convert designs created with tidydesign into a block format for easy output to csv for inclusion in lab notebooks, etc in a human-readable format

Usage

```r
block_tidydesign(
    tidydesign,
    collapse = NULL,
    wellnames_sep = "_",
    wellnames_colname = "Well"
)
```
Arguments
- **tidydesign**: A tidydesign data.frame (e.g. as created by make_tidydesign)
- **collapse**: NULL or a string to use for concatenating design elements together. If NULL, each design column will be put into its own block. If a string, that string will be used to paste together all design elements and all design elements will be returned in a single block.
- **wellnames_sep**: A string used when concatenating rownames and column names to create well names.
- **wellnames_colname**: Header for newly-created column containing the well names.

Value
A list of blockdesign data.frames (if collapse is not NULL the list is of length 1)

---

## Description
Provided a vector of y values, this function returns either the plain or per-capita difference or derivative between sequential values.

## Usage
```
calc_deriv(
  y,
  x = NULL,
  return = "derivative",
  percapita = FALSE,
  x_scale = 1,
  blank = NULL,
  subset_by = NULL,
  window_width = NULL,
  window_width_n = NULL,
  trans_y = "linear",
  na.rm = TRUE
)
```

## Arguments
- **y**: Data to calculate difference or derivative of.
- **x**: Vector of x values provided as a simple numeric.
- **return**: One of c("difference", "derivative") for whether the differences in y should be returned, or the derivative of y with respect to x.
percapita: When percapita = TRUE, the per-capita difference or derivative is returned.

x_scale: Numeric to scale x by in derivative calculation. Set x_scale to the ratio of the the units of x to the desired units. E.g. if x is in seconds, but the desired derivative is in units of /minute, set x_scale = 60 (since there are 60 seconds in 1 minute).

blank: y-value associated with a "blank" where the density is 0. Is required when percapita = TRUE.

If a vector of blank values is specified, blank values are assumed to be in the same order as unique(subset_by).

subset_by: An optional vector as long as y will be split by the unique values of this vector and the derivative for each group will be calculated independently of the others. This provides an internally-implemented approach similar to dplyr::group_by and dplyr::mutate.

window_width_n, window_width: Set how many data points are used to determine the slope at each point. When both are NULL, calc_deriv calculates the difference or derivative of each point with the next point, appending NA at the end.

When one or both are specified, a linear regression is fit to all points in the window to determine the slope. window_width_n specifies the width of the window in number of data points. window_width specifies the width of the window in units of x.

When using window_width and window_width_n at the same time, windows are conservative. Points included in each window will meet both the window_width and the window_width_n.

trans_y: One of c("linear", "log") specifying the transformation of y-values.

'log' is only available when calculating per-capita derivatives using a fitting approach (when non-default values are specified for window_width or window_width_n).

For per-capita growth expected to be exponential or nearly-exponential, "log" is recommended, since exponential growth is linear when log-transformed. However, log-transformations must be used with care, since y-values at or below 0 will become undefined and results will be more sensitive to incorrect values of blank.

na.rm: logical whether NA's should be removed before analyzing.

Details:

For per-capita derivatives, trans_y = 'linear' and trans_y = 'log' approach the same value as time resolution increases.

For instance, let's assume exponential growth \( N = e^{rt} \) with per-capita growth rate \( r \).

With trans_y = 'linear', note that \( dN/dt = re^{rt} = rN \). So we can calculate per-capita growth rate as \( r = dN/dt * 1/N \).

With trans_y = 'log', note that \( log(N) = log(e^{rt}) = rt \). So we can calculate per-capita growth rate as the slope of a linear fit of \( log(N) \) against time, \( r = log(N)/t \).
Value
A vector of values for the plain (if percapita = FALSE) or per-capita (if percapita = TRUE) difference (if return = "difference") or derivative (if return = "derivative") between y values. Vector will be the same length as y, with NA values at the ends.

---

doubling_time  
*Calculate doubling time equivalent of per-capita growth rate*

---

Description
Provided a vector of per-capita growth rates, this function returns the vector of equivalent doubling times.

Usage
doubling_time(y, x_scale = 1)

Arguments
- y: Vector of per-capita derivative data to calculate the equivalent doubling time of
- x_scale: Numeric to scale per-capita derivative values by
  Set x_scale to the ratio of the the units of y to the desired units. E.g. if y is in per-second, but the desired doubling time is in minutes, x_scale = 60 (since there are 60 seconds in 1 minute).

Value
A vector of values for the doubling time equivalent to the per-capita growth rate supplied for y.

---

example_widedata  
*Example noisy growth curve data in wide format*

---

Description
A dataset containing example growth of 96 wells of simulated bacteria or bacteria and phages.
Wells A1...A8 through F1...F8 contain 48 different simulated bacterial strains growing alone. Wells G1...G8 through L1...L8 contain the same 48 bacterial strains in an identical layout, but this time growing in the presence of a phage.

Usage
example_widedata
**Format**

A dataframe with 97 rows and 97 variables:

- **time** time, in seconds, since growth curve began
- **A1, A2...H11, H12** bacterial density in the given well

**Details**

Bacterial populations exhibit diauxic growth as they approach their carrying capacity, and they also evolve resistance in the face of selection from the phage population.

This data includes some simulated noise to approximate the noise generated during data collection by plate readers.

---

**example_widedata_noiseless**

*Example growth curve data in wide format*

---

**Description**

A dataset containing example growth of 96 wells of simulated bacteria or bacteria and phages.

Wells A1...A8 through F1...F8 contain 48 different simulated bacterial strains growing alone. Wells G1...G8 through L1...L8 contain the same 48 bacterial strains in an identical layout, but this time growing in the presence of a phage.

**Usage**

*example_widedata_noiseless*

**Format**

A dataframe with 97 rows and 97 variables:

- **time** time, in seconds, since growth curve began
- **A1, A2...H11, H12** bacterial density in the given well

**Details**

Bacterial populations exhibit diauxic growth as they approach their carrying capacity, and they also evolve resistance in the face of selection from the phage population.

This data does not include any simulated noise.
ExtremaFunctions

Find local extrema of a numeric vector

Description

These functions take a vector of y values and identify local extrema.

Usage

```r
find_local_extrema(
  y,
  x = NULL,
  window_width = NULL,
  window_width_n = NULL,
  window_height = NULL,
  return = "index",
  return_maxima = TRUE,
  return_minima = TRUE,
  return_endpoints = TRUE,
  subset = NULL,
  na.rm = TRUE,
  width_limit = NULL,
  width_limit_n = NULL,
  height_limit = NULL
)

first_maxima(
  y,
  x = NULL,
  window_width = NULL,
  window_width_n = NULL,
  window_height = NULL,
  return = "index",
  return_endpoints = TRUE,
  ...
)

first_minima(
  y,
  x = NULL,
  window_width = NULL,
  window_width_n = NULL,
  window_height = NULL,
  return = "index",
  return_endpoints = TRUE,
  ...
)
```
ExtremaFunctions

Arguments

- **y**
  - Numeric vector of y values in which to identify local extrema
- **x**
  - Optional numeric vector of corresponding x values
- **window_width**
  - Width of the window (in units of x) used to search for local extrema. A narrower width will be more sensitive to narrow local maxima/minima, while a wider width will be less sensitive to local maxima/minima.
- **window_width_n**
  - The maximum number of data points a single extrema-search step is allowed to take. For example, when maxima-finding, the function will not pass a valley consisting of more than window_width_n data points. A smaller window_width_n will be more sensitive to narrow local maxima/minima, while a larger window_width_n will be less sensitive to narrow local maxima/minima.
- **window_height**
  - The maximum change in y a single extrema-search step is allowed to take. For example, when maxima-finding, the function will not pass a valley deeper than window_height. A smaller window_height will be more sensitive to shallow local maxima/minima, while a larger window_height will be less sensitive to shallow maxima/minima.
- **return**
  - One of c("index", "x", "y"), determining whether the function will return the index, x value, or y value associated with the identified extremas
- **return_maxima, return_minima**
  - logical for which classes of local extrema to return
- **return_endpoints**
  - Should the first and last values in y be included if they are in the returned vector of extrema?
- **subset**
  - A vector of logical values indicating which x and y values should be included (TRUE) or excluded (FALSE). If return = "index", index will be for the whole vector and not the subset of the vector
- **na_rm**
  - logical whether NA's should be removed before analyzing
- **width_limit**
  - Deprecated, use window_width instead
- **width_limit_n**
  - Deprecated, use window_width_n instead
- **height_limit**
  - Deprecated, use window_height instead
- **...**
  - (for first_maxima and first_minima), other parameters to pass to find_local_extrema

Details

For find_local_extrema, one of window_width, window_width_n, or window_height must be provided.

For first_minima and first_maxima, if none of window_width, window_width_n, or window_height are provided, window_width_n is set to 20

If multiple of window_width, window_width_n, or window_height are provided, steps are limited conservatively (a single step must meet all criteria)

This function is designed to be compatible for use within dplyr::group_by and dplyr::summarize

In the case of exact ties in y values within a window, only the first local extrema is returned.
find_local_extrema returns a vector corresponding to all the found local extrema.
first_maxima returns only the first maxima, so is a shortcut for find_local_extrema(return_maxima = TRUE, return_minima = FALSE)[1]
first_minima returns only the first minima, so is a shortcut for find_local_extrema(return_maxima = FALSE, return_minima = TRUE)[1]
If return = "index", the returned value(s) are the indices corresponding to local extrema in the data
If return = "x", the returned value(s) are the x value(s) corresponding to local extrema in the data
If return = "y", the returned value(s) are the y value(s) corresponding to local extrema in the data

extr_val
Extract parts of an object

Description
A wrapper for [ with handling of NA's for use in dplyr::summarize()

Usage
extr_val(x, i, allNA_NA = TRUE, na.rm = TRUE)

Arguments

x object from which to extract element(s)
i index specifying element to extract.
allNA_NA logical indicating whether NA should be returned when all(is.na(i)) == TRUE.
na.rm a logical indicating whether missing index values should be removed.

Value
If all_NA = FALSE and na.rm = FALSE, identical to x[i].
If all_NA = FALSE and na.rm = TRUE, identical to x[i[!is.na(i)]]).
If all_NA = TRUE, identical to x[i] unless all(is.na(i)) == TRUE, in which case returns NA
first_peak

Find the first local maxima of a numeric vector

Description
This function has been deprecated in favor of the identical new function first_maxima

Usage
first_peak(
  y,
  x = NULL,
  window_width = NULL,
  window_width_n = NULL,
  window_height = NULL,
  return = "index",
  return_endpoints = TRUE,
  ...
)

Arguments
y Numeric vector of y values in which to identify local extrema
x Optional numeric vector of corresponding x values
window_width Width of the window (in units of x) used to search for local extrema. A narrower width will be more sensitive to narrow local maxima/minima, while a wider width will be less sensitive to local maxima/minima.
window_width_n The maximum number of data points a single extrema-search step is allowed to take. For example, when maxima-finding, the function will not pass a valley consisting of more than window_width_n data points.
  A smaller window_width_n will be more sensitive to narrow local maxima/minima, while a larger window_width_n will be less sensitive to narrow local maxima/minima.
  If not provided, defaults to ~0.2*length(y)
window_height The maximum change in y a single extrema-search step is allowed to take. For example, when maxima-finding, the function will not pass a valley deeper than window_height.
  A smaller window_height will be more sensitive to shallow local maxima/minima, while a larger window_height will be less sensitive to shallow maxima/minima.
return One of c("index", "x", "y"), determining whether the function will return the index, x value, or y value associated with the first maxima in y values
return_endpoints Should the first or last value in y be allowed to be returned?
... Other parameters to pass to find_local_extrema
Details

This function takes a vector of y values and returns the index (by default) of the first local maxima. It serves as a shortcut for `find_local_extrema(return_maxima = TRUE, return_minima = FALSE)[1]`

If none of `window_width`, `window_width_n`, or `window_height` are provided, default value of `window_width_n` will be used.

This function is designed to be compatible for use within `dplyr::group_by` and `dplyr::summarize`

Value

If `return = "index"`, a vector of indices corresponding to local extrema in the data
If `return = "x"`, a vector of x values corresponding to local extrema in the data
If `return = "y"`, a vector of y values corresponding to local extrema in the data

See Also

[first_maxima()]

Description

A function that converts base-26 Excel-style letters to numbers

Usage

`from_excel(x)`

Arguments

- `x` A vector of column names in Excel-style base-26 letter format (any values that are already in base-10 will be returned as-is)

Value

A vector of numbers in base-10

---

`from_excel` | *A function that converts base-26 Excel-style letters to numbers*
import_blockdesigns

**Description**

Function to import block-shaped designs from files and return tidy designs. This function acts as a wrapper to call `read_blocks`, `paste_blocks`, `trans_block_to_wide`, `trans_wide_to_tidy`, and `separate_tidy` in one go.

**Usage**

```r
import_blockdesigns(files, block_names = NULL, sep = NULL, ...)
```

**Arguments**

- `files`  
  Vector of filenames (as strings), each of which is a block-shaped designs file. Inputs can be .csv, .xls, or .xlsx

- `block_names`  
  Vector of names for design elements. These will be the resulting column names in the output data frame. Should be in the same order as files and/or same order as corresponding files themselves. If NULL, file names will be used as column names.

- `sep`  
  If block design files are already pasted, sep specifies the string separating design elements. If NULL, `import_blockdesigns` will assume no elements are already pasted together and attempt to find a character not used in the imported files to paste and later separate design elements.

- `...`  
  Other arguments to pass to `read_blocks`, `paste_blocks`, `trans_block_to_wide`, `trans_wide_to_tidy`, and `separate_tidy`. See Details for more information.

**Details**

Common arguments that you may want to provide via `...` include:

- `startrow`, `endrow`, `startcol`, `endcol`, `sheet` - specifying the location of design information inside files to `read_blocks`

- `wellnames_sep` - specifying what character (or "" for none) should be used when pasting together the rownames and column names. Note that this should be chosen to match the well names in your measures.

Note that `import_blockdesigns` cannot currently handle metadata specified via the `metadata` argument of `read_blocks`.

If you find yourself needing more control, you can run the steps manually, first reading with `read_blocks`, pasting as needed with `paste_blocks`, transforming to tidy with `trans_block_to_wide` and `trans_wide_to_tidy`, and separating as needed with `separate_tidy`s.
import_blockmeasures

Description

Function to import blockmeasures from files and return widemeasures. This function acts as a wrapper to call read_blocks, uninterleave, then trans_block_to_wide in one go.

Usage

import_blockmeasures(
  files,
  num_plates = 1,
  plate_names = NULL,
  wellnames_sep = "",
  ...
)

Arguments

files Vector of filenames (as strings), each of which is a block-shaped file containing measures data. File formats can be .csv, .xls, or .xlsx.
num_plates Number of plates. If multiple plates uninterleave will be used to separate blockmeasures into those plates accordingly.
plate_names (optional) Names to put onto the plates when output.
wellnames_sep String to use as separator for well names between rowname and column name.
...
Other arguments to pass to read_blocks, uninterleave, or widen_blocks.

Details

Common arguments that you may want to provide via ... include:
startrow, endrow, startcol, endcol, sheet - specifying the location of design information inside files to read_blocks.
metadata - specifying metadata to read_blocks.
See help for read_blocks for more details.
If you find yourself needing more control, you can run the steps manually, first reading with read_blocks, separating plates as needed with uninterleave, then transforming to wide with trans_block_to_wide.

Value

If num_plates = 1, a wide-shaped data.frame containing the measures data.
if num_plates is greater than one, a list of data.frame's, where each data.frame is wide-shaped.
Description

Lag time is calculated by projecting a tangent line at the point of maximum (per-capita) derivative backwards to find the time when it intersects with the starting y-value.

Usage

```r
lag_time(
  x = NULL,
  y = NULL,
  deriv = NULL,
  trans_y = "log",
  na.rm = TRUE,
  slope = NULL,
  x1 = NULL,
  y1 = NULL,
  y0 = NULL
)
```

Arguments

- **x**: Vector of x values (typically time)
- **y**: Vector of y values (typically density)
- **deriv**: Vector of derivative values (typically per-capita derivative)
- **trans_y**: One of `c("linear", "log")` specifying the transformation of y-values. 'log' is the default, producing calculations of lag time assuming a transition to exponential growth. 'linear' is available for alternate uses
- **na.rm**: a logical indicating whether missing values should be removed
- **slope**: Slope to project from x1,y1 to y0 (typically per-capita growth rate). If not provided, will be calculated as `max(deriv)`
- **x1**: x value (typically time) to project slope from. If not provided, will be calculated as `x[which.max(deriv)]`
- **y1**: y value (typically density) to project slope from. If not provided, will be calculated as `y[which.max(deriv)]`
- **y0**: y value (typically density) to find intersection of slope from x1, y1 with. If not provided, will be calculated as `min(y)`
Details

For most typical uses, simply supply \( x, y \), and \( \text{deriv} \) (using the per-capita derivative and \( \text{trans}_y = \log \)')

Advanced users may wish to use alternate values for the slope, origination point, or initial \( y \)-value. In that case, values can be supplied to \( \text{slope} \), \( x1 \), \( y1 \), and/or \( y0 \), which will override the default calculations. If and only if all of \( \text{slope} \), \( x1 \), \( y1 \), and \( y0 \) are provided, \( \text{lag}_\text{time} \) is vectorized on their inputs and will return a vector of lag time values.

This function is designed to be compatible for use within \texttt{dplyr::group_by} and \texttt{dplyr::summarize}.

Value

Typically a scalar of the lag time in units of \( x \). See Details for cases when value will be a vector.

\begin{verbatim}
make_design    Make design data.frame(s)
\end{verbatim}

Description

This is a function to easily input experimental design elements for later merging with read data.

Usage

\begin{verbatim}
make_design(nrows = NULL, ncols = NULL, block_row_names = NULL, block_col_names = NULL, output_format = "tidy", wellnames_numeric = FALSE, wellnames_sep = ",", wellnames_colname = "Well", colnames_first = FALSE, lookup_tbl_start = 1, pattern_split = "", ...
)
\end{verbatim}

Arguments

- \texttt{nrows, ncols} Number of rows and columns in the plate data
- \texttt{block_row_names, block_col_names} Names of the rows, columns of the plate blockmeasures data
- \texttt{output_format} One of \texttt{c("blocks", "blocks_pasted", "wide", "tidy")} denoting the format of the resulting data.frame
  - For easy merging with \texttt{tidymeasures}, leave as default of 'tidy'.
  - For human-readability to confirm design is correct, choose 'blocks' or 'blocks_pasted'.
  - For writing to block-shaped file(s), choose 'blocks' or 'blocks_pasted'.

wellnames_numeric
If block_row_names or block_col_names are not specified, then names will be
generated automatically according to wellnames_numeric.
If wellnames_numeric is TRUE, rows and columns will be numbered with "R"
and "C" prefixes, respectively.
If wellnames_numeric is FALSE, rows will be lettered A through Z, while
columns will be numbered

wellnames_sep
A string used when concatenating rownames and column names to create well
names, when output_format = "wide" or output_format = "tidy"

wellnames_colname
Header for newly-created column containing the well names, when output_format
= "tidy"

colnames_first
When wellnames are created for output_format = "wide" or output_format
= "tidy" by paste-ing the rownames and column names, should the column
names come first.

lookup_tbl_start
Value in the lookup table for the split pattern values that corresponds to the first
value in the vector.
Lookup table by default is c(1,2,...,8,9,A,B,...,Z,a,b,...,y,z). If, for example,
lookup_tbl_start = "A", then the lookup table will now be c(A,B,...,Y,Z,a,b,...,y,z)

pattern_split
character to split pattern elements provided in ... by, if they're not already a
vector

... Each ... argument must be named, and must be a list with five elements:
1. a vector of the values
2. a vector of the rows the pattern should be applied to
3. a vector of the columns the pattern should be applied to
4. a string or vector denoting the pattern in which the values should be filled
   into the rows and columns specified.
   If it's a string, will be split by pattern_split. Pattern will be used as the
   indices of the values vector.
   0's refer to NA. The pattern will be recycled as necessary to fill all the wells of
   the rows and columns specified.
5. a logical for whether this pattern should be filled byrow

Details
Note that either nrows or block_row_names must be provided and that either ncols or block_col_names
must be provided

Value
Depends on output_format:
If output_format = "blocks", a list of data.frame's where each data.frame is block-shaped
containing the information for a single design element
If output_format = "blocks_pasted", a single data.frame containing the paste-ed information
for all design elements
If `output_format = "wide"`, a wide-shaped `data.frame` containing all the design elements.
If `output_format = "tidy"`, a tidy-shaped `data.frame` containing all the design elements.

**Examples**

```r
make_design(nrows = 8, ncols = 12,
  design_element_name = list(c("A", "B", "C"),
    2:7,
    2:11,
    "112301",
    TRUE))
```

```r
# To be reminded what arguments are needed, use make_designpattern:
make_design(nrows = 8, ncols = 12,
  design_element_name = make_designpattern(
    values = c("A", "B", "C"),
    rows = 2:7,
    cols = 2:11,
    pattern = "112301",
    byrow = TRUE))
```

---

**make_designpattern**

*Make design pattern*

**Description**

A helper function for use with `make_design`.

**Usage**

```r
make_designpattern(
  values,
  rows,
  cols,
  pattern = 1:length(values),
  byrow = TRUE
)
```

```r
mdp(values, rows, cols, pattern = 1:length(values), byrow = TRUE)
```

**Arguments**

- `values`: Vector of values to use.
- `rows`: Vector of rows where pattern applies.
- `cols`: Vector of cols where pattern applies.
- `pattern`: Numeric pattern itself, where numbers refer to entries in `values`.
- `byrow`: Logical for whether pattern should be created by row.
**make_tidydesign**

**Value**

\[
\text{list(values, rows, cols, pattern, byrow)}
\]

**See Also**

[gcplyr::make_design()]

**Examples**

```r
make_design(nrows = 8, ncols = 12,
    design_element_name = make_designpattern(
        values = c("A", "B", "C"),
        rows = 2:7,
        cols = 2:11,
        pattern = "112301",
        byrow = TRUE))
```

---

**Description**

This is a function to easily input experimental design elements for later merging with read data.

**Usage**

```r
make_tidydesign(
    nrows = NULL,
    ncols = NULL,
    block_row_names = NULL,
    block_col_names = NULL,
    wellnames_sep = "",
    wellnames_colname = "Well",
    wellnames_Excel = TRUE,
    lookup_tbl_start = 1,
    pattern_split = "",
    colnames_first = FALSE,
    ...
)
```

**Arguments**

- `nrows`, `ncols`  Number of rows and columns in the plate data
- `block_row_names`, `block_col_names`  Names of the rows, columns of the plate blockmeasures data
wellnames_sep  A string used when concatenating rownames and column names to create well names
wellnames_colname
    Header for newly-created column containing the well names
wellnames_Excel
    If block_row_names or block_col_names are not specified, should rows and columns be named using Excel-style base-26 lettering for rows and numbering for columns? If FALSE, rows and columns will be numbered with "R" and "C" prefix.
lookup_tbl_start
    Value in the lookup table for the split pattern values that corresponds to the first value in the vector.
    Lookup table by default is c(1,2,...,8,9,A,B,...Y,Z,a,b,...,y,z). If, for example, lookup_tbl_start = "A", then the lookup table will now be c(A,B,...Y,Z,a,b,...,y,z)
pattern_split  character to split pattern elements provided in ... by
colnames_first
    In the wellnames created by paste-ing the rownames and column names, should the column names come first
...  Each ... argument must be a list with five elements:
    1. a vector of the values
    2. a vector of the rows the pattern should be applied to
    3. a vector of the columns the pattern should be applied to
    4. a string of the pattern itself, where numbers refer to the indices in the values vector
    0's refer to NA
    This pattern will be split using pattern_split, which defaults to every character
    5. a logical for whether this pattern should be filled byrow

Details

Note that either nrows or block_row_names must be provided and that either ncols or block_col_names must be provided

Examples: my_example <- make_tidydesign(nrows = 8, ncols = 12, design_element_name = list(c("Value1", "Value2", "Value3"), rowstart:rowend, colstart:colend, "11122333000", TRUE) To make it easier to pass arguments, use make_designpattern: my_example <- make_tidydesign(nrows = 8, ncols = 12, design_element_name = make_designpattern(values = c("L", "G", "C"), rows = 2:7, cols = 2:11, pattern = "11223300", byrow = TRUE))

Value

a tidy-shaped data.frame containing all the design elements
merge_dfs

Collapse a list of dataframes, or merge two dataframes together

Description

This function is essentially a wrapper for dplyr::full_join. The most typical use of this function is to merge designs with measures data, or to use the collapse functionality of this function to merge a list of dataframes into a single dataframe. Merging is done by column-names that match between x and y.

Usage

merge_dfs(
  x,
  y = NULL,
  by = NULL,
  drop = FALSE,
  collapse = FALSE,
  names_to = NA,
  ...
)

Arguments

x First data.frame, or list of data frames, to be joined
y Second data.frame, or list of data frames, to be joined
by A character vector of variables to join by, passed directly to dplyr::full_join
drop Should only complete_cases of the resulting data.frame be returned?
collapse A logical indicating whether x or y is a list containing data frames that should be merged together before being merged with the other
names_to Column name for where names(x) or names(y) will be entered in if collapse = TRUE.
  If a value of NA then names(x) or names(y) will not be put into a column in the returned data.frame
...
Other arguments to pass to dplyr::full_join

Value

Data.frame containing merged output of x and y
MinMaxGC  Maxima and Minima

Description

Returns the maxima and minima of the input values.

Usage

\[
\text{max\_gc}(\ldots, \text{na.rm} = \text{TRUE}, \text{allmissing\_NA} = \text{TRUE}) \\
\text{min\_gc}(\ldots, \text{na.rm} = \text{TRUE}, \text{allmissing\_NA} = \text{TRUE})
\]

Arguments

\[
\begin{align*}
... & \quad \text{numeric or character arguments} \\
\text{na.rm} & \quad \text{a logical indicating whether missing values should be removed.} \\
\text{allmissing\_NA} & \quad \text{a logical indicating whether NA should be returned when there are no non-missing arguments passed to min or max (often because na.rm = TRUE but all values are NA)}
\end{align*}
\]

Details

These functions are wrappers for min and max, with the additional argument allmissing\_NA.

Value

If allmissing\_NA = FALSE, identical to min or max.

If allmissing\_NA = TRUE, identical to min or max except that, in cases where min or max would return an infinite value and raise a warning because there are no non-missing arguments, min\_gc and max\_gc return NA

moving\_average  Moving average smoothing

Description

This function uses a moving average to smooth data

Usage

\[
\text{moving\_average}( \\
\quad \text{formula, } \\
\quad \text{data, } \\
\quad \text{window\_width\_n} = \text{NULL, } \\
\quad \text{window\_width} = \text{NULL, } \\
\quad \text{na.rm} = \text{TRUE})
\]
moving_median

Arguments

- **formula**: Formula specifying the numeric response (density) and numeric predictor (time).
- **data**: Dataframe containing variables in `formula`
- **window_width_n**: Number of data points wide the moving average window is (therefore, must be an odd number of points)
- **window_width**: Width of the moving average window (in units of \( x \))
- **na.rm**: logical whether NA's should be removed before analyzing

Value

Vector of smoothed data, with NA's appended at both ends

---

**moving_median**

*Moving median smoothing*

Description

This function uses a moving median to smooth data

Usage

```r
moving_median(
  formula,
  data,
  window_width_n = NULL,
  window_width = NULL,
  na.rm = TRUE
)
```

Arguments

- **formula**: Formula specifying the numeric response (density) and numeric predictor (time).
- **data**: Dataframe containing variables in `formula`
- **window_width_n**: Number of data points wide the moving median window is (therefore, must be an odd number of points)
- **window_width**: Width of the moving median window (in units of \( x \))
- **na.rm**: logical whether NA's should be removed before analyzing

Value

Vector of smoothed data, with NA's appended at both ends
**paste_blocks**

*Paste a list of blocks into a single block*

**Description**

This function uses `paste` to concatenate the same-location entries of a list of data.frames together (i.e. all the first row-first column values are pasted together, all the second row-first column values are pasted together, etc.)

**Usage**

```r
paste_blocks(blocks, sep = " ", nested_metadata = NULL)
```

**Arguments**

- `blocks`: Blocks, either a single data.frame or a list of data.frames
- `sep`: String to use as separator for output pasted values
- `nested_metadata`: A logical indicating the existence of nested metadata in the blockmeasures list, e.g. as is typically output by `read_blocks`. If NULL, will attempt to infer existence of nested metadata

**Value**

If `nested_metadata = TRUE` (or is inferred to be TRUE), a list containing a list containing: 1. a data.frame with the pasted data values from `blocks`, and 2. a vector with the pasted metadata values from `blocks`

If `nested_metadata = FALSE` (or is inferred to be FALSE), a list containing data.frame’s with the pasted values from `blocks`

---

**read_blocks**

*Read blockmeasures*

**Description**

A function that reads block measures into the R environment

**Usage**

```r
read_blocks(
  files, 
  extension = NULL,
  startrow = NULL,
  endrow = NULL,
  startcol = NULL,
```

---


endcol = NULL,
  sheet = NULL,
  metadata = NULL,
  block_names = NULL,
  header = NA,
  sider = NA,
  wellnames_numeric = FALSE,
  na.strings = c("NA", ""),
  ...

Arguments

files A vector of filepaths relative to the current working directory where each filepath is a single plate read

extension (optional) the extension of the files: "csv", "xls", or "xlsx", or "tbl" for use of read.table

If none provided, read_blocks will infer file extension from provided filenames. When extension is not "csv", "xls", or "xlsx" will use utils::read.table

startrow, endrow, startcol, endcol (optional) the rows and columns where the measures data are located in files. Can be a vector or list the same length as files, or a single value that applies to all files. Values can be numeric or a string that will be automatically converted to numeric by from_excel.

If not provided, data is presumed to begin on the first row and column of the file(s) and end on the last row and column of the file(s).

sheet (optional) If data is in .xls or .xlsx files, which sheet it is located on. Defaults to the first sheet if not specified

metadata (optional) non-spectrophotometric data that should be associated with each read blockmeasures. A named list where each item in the list is either: a vector of length 2, or a list containing two vectors.

In the former case, each vector should provide the row and column where the metadata is located in all of the blockmeasures input files.

In the latter case, the first vector should provide the rows where the metadata is located in each of the corresponding input files, and the second vector should provide the columns where the metadata is located in each of the corresponding input files. (This case is typically used when reading multiple blocks from a single file.)

block_names (optional) vector of names corresponding to each plate in files. If not provided, block_names are inferred from the filenames

header TRUE, FALSE, or NA, or a vector of such values, indicating whether the file(s) contains the column names as its first line. If header = NA will attempt to infer the presence of column names. If header = FALSE or no column names are inferred when header = NA, column names will be generated automatically according to wellnames_numeric
sider

TRUE, FALSE, or NA, or a vector of such values, indicating whether the file(s) contains the row names as its first line. If sider = NA will attempt to infer the presence of row names. If sider = FALSE or no row names are inferred when sider = NA, row names will be generated automatically according to wellnames_numeric.

wellnames_numeric

If row names and column names are not provided in the input dataframe as specified by header and sider, then names will be generated automatically according to wellnames_numeric.

If wellnames_numeric is TRUE, rows and columns will be numbered with "R" and "C" prefixes, respectively.

If wellnames_numeric is FALSE, rows will be lettered A through Z, while columns will be numbered.

na.strings

A character vector of strings which are to be interpreted as NA values by utils::read.csv, readxl::read_xls, readxl::read_xlsx, or utils::read.table.

... Other arguments passed to utils::read.csv, readxl::read_xls, readxl::read_xlsx, or utils::read.table.

Details

For metadata, read_blocks can handle an arbitrary number of additional pieces of information to extract from each blockcurve file as metadata. These pieces of information are specified as a named list of vectors where each vector is the c(row, column) where the information is to be pulled from in the input files.

This metadata is returned as the second list element of each blockcurve, e.g.:

```
[2][2] date-time #1
[2][3] temp #1

[2][2] date-time #2
[2][3] temp #2
```

Calling uninterleave on the output of read_blocks works on block data and the associated metadata because uninterleave operates on the highest level entries of the list (the [[1]] [[2]] level items), leaving the meta-data associated with the block data.

trans_block_to_wide integrates this metadata into the wide-shaped dataframe it produces.

Value

A list where each entry is a list containing the block data frame followed by the block_names (or filenames, if block_names is not provided) and any specified metadata.
read_tidys

Description

A function that imports tidy-shaped files into R. Largely acts as a wrapper for \texttt{utils::read.csv, readxl::read_xls}, \texttt{readxl::read_xls}, or \texttt{readxl::read_xlsx}, but can handle multiple files at once and has additional options for taking subsets of rows/columns rather than the entire file and for adding filename or run names as an added column in the output.

Usage

```r
read_tidys(
  files,
  extension = NULL,
  startrow = NULL,
  endrow = NULL,
  startcol = NULL,
  endcol = NULL,
  sheet = NULL,
  run_names = NULL,
  names_to_col = NULL,
  na.strings = c("NA", ""),
  ...)
```

Arguments

- `files`: A vector of filepaths (relative to current working directory) where each one is a tidy-shaped data file.
- `extension`: (optional) the extension of the files: "csv", "xls", or "xlsx" for use of \texttt{read.table}. If none provided, \texttt{read_tidys} will infer file extension from provided filenames. When extension is not "csv", "xls", or "xlsx" will use \texttt{utils::read.table}.
- `startrow`, `endrow`, `startcol`, `endcol`: (optional) the rows and columns where the data are located in `files`. Can be a vector or list the same length as `files`, or a single value that applies to all files. Values can be numeric or a string that will be automatically converted to numeric by \texttt{from_excel}. If not provided, data is presumed to begin on the first row and column of the file(s) and end on the last row and column of the file(s).
- `sheet`: The sheet of the input files where data is located (if input files are .xls or .xlsx). If not specified defaults to the first.
- `run_names`: Names to give the tidy files read in. By default uses the file names if not specified. These names may be added to the resulting data frame depending on the value of the `names_to_col` argument.
names_to_col: Should the run names (provided in run_names or inferred from files) be added as a column to the output? If names_to_col is TRUE, they will be added with the column name "run_name". If names_to_col is FALSE, they will not be added. If names_to_col is a string, they will be added and the column name will be the string specified for names_to_col. If names_to_col is NULL, they only will be added if there are multiple tidy data.frames being read. In which case, the column name will be "run_name".

na.strings: A character vector of strings which are to be interpreted as NA values by utils::read.csv, readxl::read_xls, readxl::read_xlsx, or utils::read.table...

... Other arguments passed to utils::read.csv, readxl::read_xls, readxl::read_xlsx, or utils::read.table sheet

Details

startrow, endrow, startcol, endcol, sheet and extension can either be a single value that applies for all files or vectors or lists the same length as files. Note that the startrow is always assumed to be a header.

Value

A dataframe containing a single tidy data.frame, or A list of tidy-shaped data.frames named by filename

---

read_wides: Read wides

Description

A function that imports widemeasures in files into the R environment

Usage

```r
read_wides(
  files,
  extension = NULL,
  startrow = NULL,
  endrow = NULL,
  startcol = NULL,
  endcol = NULL,
  header = TRUE,
  sheet = NULL,
  run_names = NULL,
  names_to_col = "file",
  metadata = NULL,
  na.strings = c("NA", ""),
  ...)
```
read_wides

Arguments

files
A vector of filepaths (relative to current working directory) where each one is a widemeasures set of data.

extension
(optional) the extension of the files: "csv", "xls", or "xlsx", or "tbl" for use of read.table

If none provided, read_wides will infer file extension from provided filenames. When extension is not "csv", "xls", or "xlsx" will use utils::read.table

startrow, endrow, startcol, endcol
(optional) the rows and columns where the data are located in files.

Can be a vector or list the same length as files, or a single value that applies to all files. Values can be numeric or a string that will be automatically converted to numeric by from_excel.

If not provided, data is presumed to begin on the first row and column of the file(s) and end on the last row and column of the file(s).

header
logical for whether there is a header to the data. If FALSE columns are simply numbered. If TRUE is the row above startrow (if startrow is specified) or the first row of the input files (if startrow is not specified)

sheet
The sheet of the input files where data is located (if input files are .xls or .xlsx).

If not specified defaults to the first sheet

run_names
Names to give the widemeasures read in. By default uses the file names if not specified

names_to_col
Should the run names (provided in run_names or inferred from files) be added as a column to the widemeasures? If names_to_col is NULL, they will not be.

If names_to_col is a string, that string will be the column header for the column where the names will be stored

metadata
(optional) non-spectrophotometric data that should be associated with each read widemeasures. A named list where each item in the list is either: a vector of length 2, or a list containing two vectors.

In the former case, each vector should provide the row and column where the metadata is located in all of the blockmeasures input files.

In the latter case, the first vector should provide the rows where the metadata is located in each of the corresponding input files, and the second vector should provide the columns where the metadata is located in each of the corresponding input files. (This case is typically used when reading multiple blocks from a single file.)

na.strings
A character vector of strings which are to be interpreted as NA values by utils::read.csv, readxl::read_xls, readxl::read_xlsx, or utils::read.table

... Other arguments passed to utils::read.csv, readxl::read_xls, readxl::read_xlsx, or utils::read.table

Details

startrow, endrow, startcol, endcol, timecol, sheet and extension can either be a single value that applies for all files or vectors or lists the same length as files,
separate_tidy

Separate a column into multiple columns

Description

This function is primarily a wrapper for tidyr::separate, which turns a single character column into multiple columns.

Usage

separate_tidy(data, col, into = NULL, sep = " ", coerce_NA = TRUE, ...)

Arguments

data A data frame

col Column name or position

into A character vector of the new column names. Use NA to omit the variable in the output.
If NULL, separate_gc will attempt to infer the new column names from the column name of col

sep Separator between columns passed to tidyr::separate:
If character, sep is interpreted as a regular expression.
If numeric, sep is interpreted as character positions to split at. Positive values start at 1 at the far-left of the string; negative values start at -1 at the far-right of the string. The length of sep should be one less than into

coerce_NA logical dictating if "NA" strings will be coerced into NA values after separating.

... Other arguments passed to tidyr::separate

Value

A data frame containing new columns in the place of col
smooth_data

Description

This function calls other functions to smooth growth curve data

Usage

smooth_data(
  ...,  
  x = NULL, 
  y = NULL, 
  sm_method, 
  subset_by = NULL, 
  return_fitobject = FALSE 
)

Arguments

... Arguments passed to stats::loess, mgcv::gam, moving_average, or moving_median. Typically includes tuning parameter(s), which in some cases are required. See Details for more information.

x An (optional) vector of predictor values to smooth along (e.g. time)

y A vector of response values to be smoothed (e.g. density). If NULL, formula and data *must* be provided via ...

sm_method Argument specifying which smoothing method should be used to smooth data. Options include "moving-average", "moving-median", "loess", and "gam"

subset_by An optional vector as long as y. y will be split by the unique values of this vector and the derivative for each group will be calculated independently of the others. This provides an internally-implemented approach similar to dplyr::group_by and dplyr::mutate

return_fitobject logical indicating whether entire object returned by fitting function should be returned. If FALSE, just fitted values are returned.

Details

For moving_average and moving_median, passing window_width or window_width_n via ... is required. window_width sets the width of the moving window in units of x, while window_width_n sets the width in units of number of data points. Larger values for either will produce more "smoothed" data.

For loess, the span argument sets the fraction of data points that should be included in each calculation. It's typically best to specify, since the default of 0.75 is often too large for growth curves data. Larger values of span will produce more more "smoothed" data
For `gam`, both arguments to `gam` and `s` can be provided via `. . .`. Most frequently, the `k` argument to `s` sets the number of "knots" the spline-fitting can use. Smaller values will be more "smoothed".

When using `sm_method = "gam"`, advanced users may also modify other parameters of `s()`, including the smoothing basis `bs`. These bases can be thin plate (`bs = "tp"`, the default), cubic regressions (`bs = "cr"`), or many other options (see `?mcgv::s`). I recommend leaving the default thin plate regressions, whose main drawback is that they are computationally intensive to calculate. For growth curves data, this is unlikely to be relevant.

As an alternative to passing `y`, for more advanced needs with `loess` or `gam`, `formula` and `data` can be passed to `smooth_data` via the `. . .` argument (in lieu of `y`).

In this case, the `formula` should specify the response (e.g. density) and predictors. For `gam` smoothing, the formula should typically be of the format: `y ~ s(x)`, which uses `mcgv::s` to smooth the data. The `data` argument should be a `data.frame` containing the variables in the formula. In such cases, `subset_by` can still be specified as a vector as long as `nrow(data)`

**Value**

If `return_fitobject == FALSE`:

A vector, the same length as `y`, with the now-smoothed `y` values

If `return_fitobject == TRUE`:

A list the same length as `unique(subset_by)` where each element is an object of the same class as returned by the smoothing method (typically a named list-like object)

Varies by method, but always with a first element named 'fitted' containing the smoothed values of the response variable, and a second element named 'residuals' containing the residuals of the fitted values and the input values

**Note: the first version released after Sep 1, 2023 will change ** **this behavior** to instead maintain the class of the object returned by the smoothing method. After the change, not all methods will return an object with 'fitted' as the first element and 'residuals' as the second element.

---

**solve_linear**

*Return missing information about a line*

**Description**

Takes a set of inputs that is sufficient information to infer a line and then returns information not provided (either the slope, an x point on the line, or a y point on the line)

**Usage**

```r
solve_linear(
  x1, y1,
  x2 = NULL, y2 = NULL,
  x3 = NULL, y3 = NULL,
)```

```r```
ThresholdFunctions

thresholds

m = NULL,
   named = TRUE
)

Arguments

x1, y1 A point on the line
x2, y2 An additional point on the line
x3, y3 An additional point on the line
m The slope of the line
named logical indicating whether the returned value(s) should be named according to what they are (m, x2, y2, x3, or y3)

Details

Note that there is no requirement that \( x_1 < x_2 < x_3 \): the points can be in any order along the line.
solve.linear works with vectors of all inputs to solve multiple lines at once, where the \( i \)th element of each argument corresponds to the \( i \)th output. Note that all lines must be missing the same information. Input vectors will be recycled as necessary.

Value

A named vector with the missing information from the line:
If \( m \) and \( x_2 \) are provided, \( y_2 \) will be returned
If \( m \) and \( y_2 \) are provided, \( x_2 \) will be returned
If \( x_2 \) and \( y_2 \) are provided, but neither \( x_3 \) nor \( y_3 \) are provided, \( m \) will be returned
If \( x_2 \) and \( y_2 \) are provided and one of \( x_3 \) or \( y_3 \) are provided, the other (\( y_3 \) or \( x_3 \)) will be returned

ThresholdFunctions Find point(s) when a numeric vector crosses some threshold

Description

These functions take a vector of \( y \) values and identify points where the \( y \) values cross some threshold \( y \) value.

Usage

find_threshold_crosses(
y,
x = NULL,
threshold,
return = "index",
return_rising = TRUE,
return_falling = TRUE,
ThresholdFunctions

return_endpoints = TRUE,
subset = NULL,
na.rm = TRUE
)

first_below(
  y,
  x = NULL,
  threshold,
  return = "index",
  return_endpoints = TRUE,
  ...
)

first_above(
  y,
  x = NULL,
  threshold,
  return = "index",
  return_endpoints = TRUE,
  ...
)

Arguments

y  Numeric vector of y values in which to identify threshold crossing event(s)
x  Optional numeric vector of corresponding x values
threshold  Threshold y value of interest
return  One of c("index", "x"), determining whether the function will return the index or x value associated with the threshold-crossing event.

If index, it will refer to the data point immediately after the crossing event.

If x, it will use linear interpolation and the data points immediately before and after the threshold-crossing to return the exact x value when the threshold crossing occurred

return_rising  logical for whether crossing events where y rises above threshold should be returned

return_falling  logical for whether crossing events where y falls below threshold should be returned

return_endpoints  logical for whether startpoint should be returned when the startpoint is above threshold and return_rising = TRUE, or when the startpoint is below threshold and return_falling = TRUE

subset  A vector of logical values indicating which x and y values should be included (TRUE) or excluded (FALSE).

If return = "index", index will be for the whole vector and not the subset of the vector
to_excel

na.rm logical whether NA's should be removed before analyzing. If return = 'index', indices will refer to the original y vector *including* NA values

... (for first_above and first_below) other arguments to pass to find_threshold_crosses

Details

This function is designed to be compatible for use within dplyr::group_by and dplyr::summarize

Value

find_threshold_crosses returns a vector corresponding to all the threshold crossings.

first_above returns only the first time the y values rise above the threshold, so is a shortcut for find_threshold_crosses(return_rising = TRUE, return_falling = FALSE)[1]

first_below returns only the first time the y values fall below the threshold, so is a shortcut for find_threshold_crosses(return_rising = FALSE, return_falling = TRUE)[1]

If return = "index", the returned value(s) are the indices immediately following threshold crossing(s)

If return = "x", the returned value(s) are the x value(s) corresponding to threshold crossing(s)

If no threshold-crossings are detected that meet the criteria, will return NA

---

to_excel A function that converts numbers into base-26 Excel-style letters

Description

A function that converts numbers into base-26 Excel-style letters

Usage

to_excel(x)

Arguments

x A vector of numbers in base-10

Value

A vector of letters in Excel-style base-26 format
trans_block_to_wide  Transform blocks to wides

Description

Takes blocks and returns them in a wide format

Usage

```r
trans_block_to_wide(
  blocks,
  wellnames_sep = "",
  nested_metadata = NULL,
  colnames_first = FALSE
)
```

Arguments

- **blocks**: Blocks, either a single data.frame or a list of data.frames
- **wellnames_sep**: String to use as separator for well names between rowname and column name (ordered according to `colnames_first`)
- **nested_metadata**: A logical indicating the existence of nested metadata in the blockmeasures list, e.g. as is typically output by `read_blocks`. If NULL, will attempt to infer existence of nested metadata
- **colnames_first**: In the wellnames created by `paste`-ing the rownames and column names, should the column names come first

Value

A single widemeasures data.frame

trans_wide_to_tidy  Pivot widemeasures longer

Description

Essentially a wrapper for `tidyrr::pivot_longer` that works on both a single widemeasures as well as a list of widemeasures
Usage

trans_wide_to_tidy(
    wides,
    data_cols = NA,
    id_cols = NA,
    names_to = "Well",
    values_to = "Measurements",
    values_to_numeric = TRUE,
    ...
)

Arguments

wides A single widemeasures data.frame, or a list of widemeasures data.frame’s

data_cols, id_cols Specifies which columns have data vs are ID’s (in tidyr::pivot_longer parlance). Each can be a single vector (which will be applied for all widemeasures) or a list of vectors, with each vector corresponding to the same-index widemeasure in widemeasures

Entries that are NA in the list will not be used

If neither data_cols nor id_cols are specified, user must provide arguments to tidyr::pivot_longer via ... for at least the cols argument and these arguments provided via ... will be used for all widemeasures data.frame’s

names_to, values_to Specifies the output column names created by tidyr::pivot_longer. Each can be provided as vectors the same length as widemeasures. Note that if neither data_cols nor id_cols

values_to_numeric logical indicating whether values will be coerced to numeric. See below for when this may be overridden by arguments passed in ...

...

Other functions to be passed to tidyr::pivot_longer. Note that including values_transform here will override the behavior of values_to_numeric

Value

Pivoted longer data.frame (if widemeasures is a single data.frame) or list of pivoted longer data.frame’s (if widemeasures is a list of data.frame’s)

uninterleave Uninterleave list

Description

Uninterleave a list that is actually interleaved elements from multiple sources and uninterleaves them into the separate sources. For instance, a list of blockmeasures that actually corresponds to two different plates can be split into two lists, each of the blockmeasures corresponding to a single plate.
Usage

uninterleave(interleaved_list, n)

Arguments

interleaved_list

A list of R objects

n

How many output sub lists there should be (i.e. how many groups the interleaved list should be divided into)

Value

A list of lists of R objects

WhichMinMaxGC

Where is the Min() or Max() or first TRUE or FALSE?

Description

Determines the location, i.e. index, of the (first) minimum or maximum of a numeric (or logical) vector.

Usage

which_min_gc(x, empty_NA = TRUE)

which_max_gc(x, empty_NA = TRUE)

Arguments

x

numeric (logical, integer, or double) vector or an R object for which the internal coercion to double works whose min or max is searched for.

empty_NA

logical, indicating if an empty value should be returned as NA (the default) or as integer(0) (the same as which.min and which.max).

Details

These functions are wrappers for which.min and which.max, with the additional argument empty_NA.

Value

If empty_NA = FALSE, identical to which.min or which.max

If empty_NA = TRUE, identical to which.min or which.max except that, in cases where which.min or which.max would return integer(0), which_min_gc and which_max_gc return NA
write_blocks

Write block designs to csv

Description

This function writes block-shaped lists (as created by read_blocks or make_design) to csv files, including both data and metadata in a variety of output formats.

Usage

write_blocks(
  blocks,
  file,
  output_format = "multiple",
  block_name_location = NULL,
  paste_sep = "_",
  filename_sep = "_",
  na = "",
  ...
)

Arguments

blocks  list of block-shaped data to be written to file
file    NULL, a character string naming a file to write to, or a vector of character strings naming files to write to.
         A file name is required when output_format = "single"
         A file name can be specified when output_format = "pasted", or file can be set to NULL as long as block_name_location = "filename" (where pasted block_name metadata will be used for the file name)
         File names can be specified when output_format = "multiple", or file can be set to NULL as long as block_name_location = "filename" (where the block_name metadata will be used for the file names)
output_format One of "single", "pasted", "multiple".
         "single" will write all blocks into a single csv file, with an empty row between successive blocks.
         "pasted" will paste all blocks together using a paste_sep, and then write that now-pasted block to a single csv file.
         "multiple" will write each block to its own csv file.
block_name_location
         Either NULL, 'filename' or 'file'.
         If NULL, block_name_location will be automatically selected based on output_format.
         For output_format = 'single' and output_format = 'pasted', block_name_location defaults to 'file'. For output_format = 'multiple', block_name_location defaults to 'filename'
write_blocks

If 'filename', the block_name metadata will be used as the output file name(s) when no file name(s) are provided, or appended to file name(s) when they have been provided.

If 'file', the block_name metadata will be included as a row in the output file.

- **paste_sep**: When `output_format = 'pasted'`, what character will be used to paste together blocks.

- **filename_sep**: What character will be used to paste together filenames when `block_name_location = 'filename'`.

- **na**: The string to use for missing values in the data.

- **...**: Other arguments passed to `write.table`

**Value**

Nothing, but R objects are written to files
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