

Package: kdry (via r-universe)

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Title K's ``Don't Repeat Yourself''-Collection

Version 0.0.2

Description A collection of personal helper functions to avoid redundancy in the spirit of the ``Don't repeat yourself'' principle of software development (https://en.wikipedia.org/wiki/Don%27t_repeat_yourself).

License GPL (>= 3)

URL <https://github.com/kapsner/kdry>

BugReports <https://github.com/kapsner/kdry/issues>

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dtr_matrix2df	<i>dtr_matrix2df</i>
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Description

Data transformation: Converts a `matrix` to `data.table` and encodes categorical variables as `factor`.

Usage

```
dtr_matrix2df(matrix, cat_vars = NULL)
```

Arguments

<code>matrix</code>	An R matrix object.
<code>cat_vars</code>	A character vector with colnames that should be converted to <code>factor</code> (default: <code>NULL</code>).

Value

A `data.table` is returned.

Examples

```
data("iris")
mat <- data.matrix(iris)
dataset <- dtr_matrix2df(mat)
str(dataset)

dataset <- dtr_matrix2df(mat, cat_vars = "Species")
str(dataset)
```

icolnames	<i>icolnames</i>
-----------	------------------

Description

Return colnames in a table with index numbers.

Usage

```
icolnames(df)
```

Arguments

df A data.frame object.

Value

A data.table with the two columns index and name is returned.

Examples

```
data("iris")
icolnames(iris)
```

list.append	<i>list.append</i>
-------------	--------------------

Description

Helper function to append an R list.

Usage

```
list.append(main_list, append_list, ...)
```

Arguments

`main_list` A list, to which another should be appended.
`append_list` A list to append to `main_list` (can be NULL or empty).
`...` Further arguments passed to `utils::modifyList()`.

Details

This function is a save wrapper around `utils::modifyLists` to combine lists as it checks for the input types and only appends the new list if its length is greater than 0.

Value

A list is returned.

See Also

[utils::modifyList\(\)](#)

Examples

```
l1 <- list("a" = 1, "b" = 2)
l2 <- list("c" = 3, "d" = 4)
list.append(l1, l2)
```

`list.update`

list.update

Description

Helper function to update items in an R list.

Usage

```
list.update(main_list, new_list, ...)
```

Arguments

`main_list` A list, which items should be updated.
`new_list` A list with new values of items from `main_list` that should be updated. All names of `new_list` must be present in `main_list`.
`...` Further arguments passed to `utils::modifyList()`.

Details

This function is a save wrapper around `utils::modifyLists` to update items in R lists as it checks for the input types and only accepts named lists.

Value

A list is returned.

See Also

`utils::modifyList()`

Examples

```
l1 <- list("a" = 1, "b" = 2)
l2 <- list("a" = 3, "b" = 4)
list.update(l1, l2)
```

`misc_argument_catcher` *misc_argument_catcher*

Description

Miscellaneous helper function to type-save catch arguments passed with R's ellipsis ("...").

Usage

```
misc_argument_catcher(...)
```

Arguments

... Named arguments passed to a function.

Details

This function aims at catching arguments that have been passed to an R function using R's ellipsis ("..."). Its purpose is to catch these arguments even in the case, if a list with arguments was provided to the ellipsis.

Value

A list is returned.

Examples

```
misc_argument_catcher(a = 1)
misc_argument_catcher(a = 1, b = 2, c = 3, d = "car")
misc_argument_catcher(list(a = 1, b = 2, c = 3, d = "car"))
misc_argument_catcher(list(a = 1, b = 2, c = 3, d = "car"), f = 9)
```

misc_duplicated_by_names
misc_duplicated_by_names

Description

Miscellaneous helper function to detect items in an object with duplicated names, e.g. in named vectors or named lists.

Usage

```
misc_duplicated_by_names(object, ...)
```

Arguments

object	An R object that has names.
...	Named arguments passed on to duplicated.

Value

Returns a logical vector of length(object) with TRUE indicating the identified items with duplicated names.

See Also

[base::duplicated\(\)](#)

Examples

```
misc_duplicated_by_names(list(a = 1, a = 1))
```

misc_recursive_copy *misc_recursive_copy*

Description

Recursively copying directories and subdirectories.

Usage

```
misc_recursive_copy(source_dir, target_dir, force = FALSE)
```

Arguments

source_dir	A character string. The path to the directory to be copied.
target_dir	A character string. The target path.
force	A boolean. If FALSE (the default) a dialog is prompted to ask the user if the directories should be copied recursively. If TRUE, the dialog is not prompted and the source_dir is copied directly to target_dir.

Value

This function has no return value.

Examples

```
if (interactive()) {  
  d1 <- file.path(tempdir(), "folder1")  
  d2 <- file.path(d1, "folder2")  
  d3 <- file.path(tempdir(), "new_folder")  
  f1 <- file.path(d1, "file.one")  
  dir.create(d2, recursive = TRUE)  
  file.create(f1)  
  misc_recursive_copy(d1, d3)  
}
```

misc_subset_options *misc_subset_options*

Description

Miscellaneous helper function to subset R options by a keyword.

Usage

```
misc_subset_options(keyword)
```

Arguments

keyword	A character. The keyword to subset the R options.
---------	---

Details

This function subsets R's options() by a keyword. It returns a list of all available options that match with the keyword. The keyword is evaluated as a regular expression.

Value

A list is returned, containing the subset of R's options() that matches with the keyword.

Examples

```
misc_subset_options("default")
```

```
mlh_outsample_row_indices
      mlh_outsample_row_indices
```

Description

Machine learning helper function to convert a vector of (in- sample) row indices of a fold into out-of-sample row indices.

Usage

```
mlh_outsample_row_indices(fold_list, dataset_nrows, type = NULL)
```

Arguments

<code>fold_list</code>	A list of integer vectors that describe the row indices of cross-validation folds. The list must be named.
<code>dataset_nrows</code>	An integer. The number of rows in the dataset <code>dataset</code> . This parameter is required in order to compute the out-of-sample row indices.
<code>type</code>	A character. To be used if the out-of-sample row indices need to be formatted in a special manner (default: <code>NULL</code>). Currently, the only allowed value is <code>type = "glmnet"</code> in order to format the row indices as required by <code>glmnet::cv.glmnet</code> 's argument <code>foldid</code> .

Value

If `type = NULL`, returns a list of same length as `fold_list` with each item containing a vector of out-of-sample row indices. If `type = "glmnet"`, a `data.table` is returned with two columns and each row representing one observation of the dataset that is assigned to a specific test fold. The column `"fold_id"` should be passed further on to the argument `foldid` of `glmnet::cv.glmnet`.

Examples

```
fold_list <- list(
  "Fold1" = setdiff(seq_len(100), 1:33),
  "Fold2" = setdiff(seq_len(100), 66:100),
  "Fold3" = setdiff(seq_len(100), 34:65)
)
mlh_outsample_row_indices(fold_list, 100)
mlh_outsample_row_indices(fold_list, 100, "glmnet")
```

mlh_reshape	<i>mlh_reshape</i>
-------------	--------------------

Description

Machine learning helper function to reshape a matrix of predicted probabilities to classes.

Usage

```
mlh_reshape(object)
```

Arguments

object	A matrix with predicted probabilities for several classes. Each row must sum up to 1.
--------	---

Value

Returns a vector of type factor of the same length as rows in object, representing the class with the highest probability for each observation in object.

Examples

```
set.seed(123)
class_0 <- rbeta(100, 2, 4)
class_1 <- (1 - class_0) * 0.4
class_2 <- (1 - class_0) * 0.6
dataset <- cbind("0" = class_0, "1" = class_1, "2" = class_2)
mlh_reshape(dataset)
```

mlh_subset	<i>mlh_subset</i>
------------	-------------------

Description

Machine learning helper function to select a subset from a data matrix or a response vector.

Usage

```
mlh_subset(object, ids)
```

Arguments

object	A vector or a data matrix. Supports also subsetting of "Surv" objects.
ids	An integer vector specifying the indices that should be selected from the object.

Value

Returns the specified subset of the object.

Examples

```
data("iris")
mlh_subset(iris, c(1:30))
mlh_subset(iris[, 5], c(1:30))
```

```
pch_check_available_cores
      pch_check_available_cores
```

Description

Parallel computing helper function to check for the available cores.

Usage

```
pch_check_available_cores(ncores = -1L)
```

Arguments

`ncores` An integer. A number of cores requested for parallel computing (default: -1L).

Value

The function returns an integer that indicates the number of cores available. If `ncores <= parallel::detectCores()` the function returns `ncores`. If `ncores > parallel::detectCores()`, the function returns `parallel::detectCores() - 1L`.

Examples

```
pch_check_available_cores(2)
```

pch_clean_up *pch_clean_up*

Description

Parallel computing helper function to clean up the parallel backend.

Usage

```
pch_clean_up(cl)
```

Arguments

`cl` A cluster object of class `c("SOCKcluster", "cluster")`.

Value

The function returns nothing. Internally, it calls `parallel::stopCluster()` and `foreach::registerDoSEQ()`.

See Also

[parallel::stopCluster\(\)](#), [foreach::registerDoSEQ\(\)](#)

Examples

```
if (require("doParallel") && require("foreach")) {
  cl <- pch_register_parallel(pch_check_available_cores(2))
  pch_clean_up(cl)
}
```

pch_register_parallel *pch_register_parallel*

Description

Parallel computing helper function to register a parallel backend.

Usage

```
pch_register_parallel(ncores)
```

Arguments

`ncores` An integer. A number of cores requested for parallel computing (default: `-1L`).

Value

The function returns a object of class c("SOCKcluster", "cluster"), created with `parallel::makePSOCKcluster()`.

See Also

`parallel::makePSOCKcluster()`, `doParallel::registerDoParallel()`

Examples

```
if (require("doParallel") && require("foreach")) {
  cl <- pch_register_parallel(pch_check_available_cores(2))
  pch_clean_up(cl)
}
```

`plt_parallel_coordinates`

plt_parallel_coordinates

Description

Parallel coordinates plot

Usage

```
plt_parallel_coordinates(
  data,
  cols = NULL,
  color_variable = NULL,
  color_args = list(alpha = 0.6, begin = 0.1, end = 0.9, option = "inferno", direction =
    1),
  line_jitter = list(w = 0.04, h = 0.04),
  text_label_size = 3.5
)
```

Arguments

<code>data</code>	A <code>data.table</code> object with the columns containing the parameters to be plotted with the parallel coordinates plot.
<code>cols</code>	A character vector with column names to subset data (default: <code>NULL</code>).
<code>color_variable</code>	A character. The name of the column to be used to colorize the lines of the plot (default: <code>NULL</code>).
<code>color_args</code>	A list with parameters for the color gradient (see details).
<code>line_jitter</code>	A list with the elements <code>w</code> and <code>h</code> to define a line jitter (default: <code>list(w = 0.04, h = 0.04)</code>), which are passed further on to define the position of <code>ggplot2::geom_line()</code> .
<code>text_label_size</code>	A numeric value to define the size of the text annotations (default: 3.5).

Details

The color gradient of the plotted lines can be defined with a list provided to the argument `color_args`. Its default values are `alpha = 0.6`, `begin = .1`, `end = .9`, `option = "inferno"`, and `direction = 1` and are passed further on to `ggplot2::scale_color_viridis_c()`. The implementation to display categorical variables is still experimental.

Value

Returns a parallel coordinates plot as `ggplot2` object.

See Also

[ggplot2::scale_color_viridis_c\(\)](#)

Examples

```
if (require("ggplot2")) {  
  data("iris")  
  plt_parallel_coordinates(  
    data = data.table::as.data.table(iris[, -5]),  
    cols = colnames(iris)[c(-1, -5)],  
    color_variable = "Sepal.Length"  
  )  
}
```

rep_frac_pct

rep_frac_pct

Description

Reporting helper function: computes and formats the relative percentage of a fraction.

Usage

```
rep_frac_pct(  
  count,  
  count_reference,  
  digits = 2,  
  na.rm = TRUE,  
  brackets = c("round", "square"),  
  suffix = TRUE  
)
```

Arguments

count	A numeric. The numerator.
count_reference	A numeric. The denominator.
digits	An integer indicating the number of decimal places.
na.rm	A logical indicating if missings should be removed from x before computing the distributional parameters (default: TRUE).
brackets	A character. Either "round" (default) or "square" to indicate the type of brackets to surround the relative count.
suffix	A character which is placed between the lower and the upper confidence bound in the formatted output.

Value

A character with the formatted output.

See Also

[stats::median](#), [stats::quantile](#), [Hmisc::wtd.quantile\(\)](#)

Examples

```
rep_frac_pct(count = 40, count_reference = 200)
rep_frac_pct(count = 40, count_reference = 200, brackets = "square")
rep_frac_pct(40, 200, brackets = "square", suffix = FALSE)
```

rep_mean_sd

rep_mean_sd

Description

Reporting helper function: computes and formats mean and standard deviation from a numeric vector.

Usage

```
rep_mean_sd(
  x,
  digits = 2,
  na.rm = TRUE,
  sd_brackets = c("round", "square"),
  sd_prefix = TRUE,
  weighted = FALSE,
  weights = NA
)
```

Arguments

<code>x</code>	A numeric vector.
<code>digits</code>	An integer indicating the number of decimal places.
<code>na.rm</code>	A logical indicating if missings should be removed from <code>x</code> before computing the distributional parameters (default: TRUE).
<code>sd_brackets</code>	A character. Either "round" (default) or "square" to indicate the type of brackets to surround the standard deviation in the formatted output.
<code>sd_prefix</code>	A logical. If TRUE (default), the standard deviation is prefixed with a plus-minus sign ("±").
<code>weighted</code>	A logical. If TRUE, a weighted mean and standard deviation are calculated (default: FALSE).
<code>weights</code>	A vector with the weights (if <code>weighted = TRUE</code> ; default: NA) passed further on to <code>stats::weighted.mean()</code> and <code>Hmisc::wtd.var()</code> .

Value

A character with the formatted output.

See Also

`mean()`, `stats::sd()`, `stats::weighted.mean()`, `Hmisc::wtd.var()`

Examples

```
set.seed(123)
x <- rnorm(1000)
rep_mean_sd(x)
rep_mean_sd(rep(1, 10))
rep_mean_sd(x, sd_brackets = "square")
rep_mean_sd(x, sd_brackets = "square", sd_prefix = FALSE)
```

rep_median_ci

rep_median_ci

Description

Reporting helper function: computes and formats median and confidence interval from a numeric vector.

Usage

```
rep_median_ci(
  x,
  conf_int,
  digits = 2,
  na.rm = TRUE,
  collapse = "to",
  iqr_brackets = c("round", "square"),
  iqr_prefix = TRUE,
  weighted = FALSE,
  weights = NA
)
```

Arguments

<code>x</code>	A numeric vector.
<code>conf_int</code>	A numeric between 0 and 100 to indicate the confidence interval that should be computed.
<code>digits</code>	An integer indicating the number of decimal places.
<code>na.rm</code>	A logical indicating if missings should be removed from <code>x</code> before computing the distributional parameters (default: TRUE).
<code>collapse</code>	A character which is placed between the lower and the upper confidence bound in the formatted output.
<code>iqr_brackets</code>	A character. Either "round" (default) or "square" to indicate the type of brackets to surround the confidence interval in the formatted output.
<code>iqr_prefix</code>	A logical. If TRUE (default), the confidence interval is prefixed with "IQR: ".
<code>weighted</code>	A logical. If TRUE, a weighted median and confidence interval are calculated (default: FALSE).
<code>weights</code>	A numeric vector of weights passed further on to <code>Hmisc::wtd.quantile()</code> if <code>weighted = TRUE</code> (default: NA).

Value

A character with the formatted output.

See Also

[stats::median](#), [stats::quantile](#), [Hmisc::wtd.quantile\(\)](#)

Examples

```
set.seed(123)
x <- rnorm(1000)
rep_median_ci(x, conf_int = 95)
rep_median_ci(rep(1, 10), conf_int = 95)
rep_median_ci(x, conf_int = 95, collapse = "-")
rep_median_ci(x, iqr_brackets = "square", conf_int = 50)
```

rep_median_iqr	<i>rep_median_iqr</i>
----------------	-----------------------

Description

Reporting helper function: computes and formats median and interquartile range from a numeric vector.

Usage

```
rep_median_iqr(  
  x,  
  digits = 2,  
  na.rm = TRUE,  
  collapse = "to",  
  iqr_brackets = c("round", "square"),  
  iqr_prefix = TRUE  
)
```

Arguments

x	A numeric vector.
digits	An integer indicating the number of decimal places.
na.rm	A logical indicating if missings should be removed from x before computing the distributional parameters (default: TRUE).
collapse	A character which is placed between the lower and the upper confidence bound in the formatted output.
iqr_brackets	A character. Either "round" (default) or "square" to indicate the type of brackets to surround the confidence interval in the formatted output.
iqr_prefix	A logical. If TRUE (default), the confidence interval is prefixed with "IQR: ".

Details

This is just a special case of [rep_median_ci\(\)](#) with the parameter `conf_int` set to 50.

Value

A character with the formatted output.

See Also

[rep_median_ci\(\)](#)

Examples

```
set.seed(123)
x <- rnorm(1000)
rep_median_iqr(x)
rep_median_iqr(rep(1, 10))
rep_median_iqr(x, collapse = "-")
rep_median_iqr(x, iqr_brackets = "square")
rep_median_iqr(x, iqr_brackets = "square", iqr_prefix = FALSE)
rep_median_iqr(x, collapse = ";", iqr_prefix = FALSE)
```

rep_pval

rep_pval

Description

Reporting helper function: formats p-value.

Usage

```
rep_pval(p, threshold = 0.001, digits = 3L)
```

Arguments

<code>p</code>	The p-value that should be formatted.
<code>threshold</code>	A threshold to indicate that only "< threshold" is printed as output (default: 0.001).
<code>digits</code>	The number of digits of the formatted p-value (digits).

Details

If the p-value is lower than the threshold, the output of the function is "< threshold". Otherwise, the p-value is formatted to the number of digits.

Value

A character with the formatted p-value.

Examples

```
rep_pval(0.032)
rep_pval(0.00032)
```

rep_sum_pct	<i>rep_sum_pct</i>
-------------	--------------------

Description

Reporting helper function: computes and formats the relative percentage of a count.

Usage

```
rep_sum_pct(  
  count,  
  count_reference,  
  digits = 2,  
  na.rm = TRUE,  
  brackets = c("round", "square"),  
  suffix = TRUE  
)
```

Arguments

count	A numeric. The numerator.
count_reference	A numeric. The denominator.
digits	An integer indicating the number of decimal places.
na.rm	A logical indicating if missings should be removed from x before computing the distributional parameters (default: TRUE).
brackets	A character. Either "round" (default) or "square" to indicate the type of brackets to surround the relative count.
suffix	A character which is placed between the lower and the upper confidence bound in the formatted output.

Value

A character with the formatted output.

See Also

[stats::median](#), [stats::quantile](#), [Hmisc:wtd.quantile\(\)](#)

Examples

```
rep_sum_pct(count = 40, count_reference = 200)  
rep_sum_pct(count = 40, count_reference = 200, brackets = "square")  
rep_sum_pct(40, 200, brackets = "square", suffix = FALSE)
```

sts_normalize	<i>sts_normalize</i>
---------------	----------------------

Description

Statistic helper function to normalize a continuous variable between zero and one.

Usage

```
sts_normalize(x, na.rm = FALSE)
```

Arguments

x	A vector of type numeric.
na.rm	A logical to indicate, if missings should be removed.

Value

Returns a vector of same length as x with values normalized between zero and one. If x contains missings and na.rm = TRUE, the missings are removed before normalization; otherwise, a vector of NA is returned.

Examples

```
sts_normalize(1:100)
```

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