

# Package ‘usefun’

July 22, 2025

**Type** Package

**Title** A Collection of Useful Functions by John

**Version** 0.5.2

**Description** A set of general functions that I have used in various projects and other R packages. Miscellaneous operations on data frames, matrices and vectors, ROC and PR statistics.

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**URL** <https://github.com/bblodfon/usefun>

**BugReports** <https://github.com/bblodfon/usefun/issues>

**Imports** checkmate, dplyr (>= 1.1.2), graphics, precrec (>= 0.14.2), PRROC (>= 1.3.1), stats, tibble (>= 3.2.1), utils

**Suggests** combinat, covr, ggplot2, mlr3misc, readr, testthat

**Encoding** UTF-8

**RoxygenNote** 7.3.2

**NeedsCompilation** no

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**Repository** CRAN

**Date/Publication** 2024-09-14 23:50:02 UTC

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---

add\_row\_to\_ternary\_df *Add a row to a 3-valued (ternary) data.frame*

---

**Description**

Use this function on a `data.frame` object (with values only in the 3-element set  $\{-1,0,1\}$  ideally - specifying either a positive, negative or none/absent condition/state/result about something) and add an extra **first or last row vector** with zero values, where  $1$  and  $-1$  will be filled when the column names of the given `data.frame` match the values in the `values.pos` or `values.neg` vector parameters respectively.

**Usage**

```
add_row_to_ternary_df(
  df,
  values.pos,
  values.neg,
  pos = "first",
  row.name = NULL
)
```

**Arguments**

df	a data.frame object with values only in the the 3-element set {-1,0,1}. The column names should be node names (gene, protein names, etc.).
values.pos	a character vector whose elements are indicators of a positive state/condition and will be assigned a value of <i>1</i> . These elements <b>must be a subset of the column names</b> of the given df parameter. If empty, no values equal to <i>1</i> will be added to the new row.
values.neg	a character vector whose elements are indicators of a negative state/condition and will be assigned a value of <i>-1</i> . If empty, no values equal to <i>-1</i> will be added to the new row. These elements <b>must be a subset of the column names</b> of the given df parameter.
pos	string. The position where we should put the new row that will be generated. Two possible values: "first" (default) or "last".
row.name	string. The name of the new row that we will added. Default value: NULL.

**Value**

the df with one extra row, having elements from the {-1,0,1} set depending on values of values.pos and values.neg vectors.

**Examples**

```
df = data.frame(c(0,-1,0), c(0,1,-1), c(1,0,0))
colnames(df) = c("A","B","C")
df.new = add_row_to_ternary_df(df, values.pos = c("A"), values.neg = c("C"), row.name = "Hello!")
```

---

add_vector_to_df	<i>Add vector to a (n x 2) data frame</i>
------------------	---

---

**Description**

Given a vector, adds each value and its corresponding name to a data frame of 2 columns as new rows, where the name fills in the 1st column and the value the 2nd column.

**Usage**

```
add_vector_to_df(df, vec)
```

**Arguments**

```
df          data.frame, with n rows and 2 columns
vec         a vector
```

**Value**

a data.frame with additional rows and each element as a character.

**Examples**

```
df = data.frame(c(0,0,1), c(0,0,2))
vec = 1:3
names(vec) = c("a", "b", "c")

add_vector_to_df(df, vec)
```

---

```
binarize_to_thres      Binarize matrix to given threshold
```

---

**Description**

Simple function that checks every element of a given matrix (or data.frame) if it surpasses the given threshold either positively or negatively and it outputs 1 for that element, otherwise 0.

**Usage**

```
binarize_to_thres(mat, thres)
```

**Arguments**

```
mat          a matrix or data.frame object
thres        a positive numerical value
```

**Value**

a binarized matrix (values either 0 or 1): elements that have 1 correspond to values of mat that they were either larger than the threshold or smaller than it's negative.

**Examples**

```
mat = matrix(data = -4:4, nrow = 3, ncol = 3)
binarize_to_thres(mat, thres = 0.5)
binarize_to_thres(mat, thres = 2.5)
```

---

colors.100	<i>100 distinct colors</i>
------------	----------------------------

---

**Description**

100 as-much-as-possible distinct colors!

**Usage**

```
colors.100
```

**Format**

An object of class character of length 100.

---

dec_to_bin	<i>Convert decimal number to its binary representation</i>
------------	--

---

**Description**

Get the binary representation of any decimal number from 0 to  $(2^{31}) - 1$ . Doesn't work for larger numbers.

**Usage**

```
dec_to_bin(decimal_num, bits = 32)
```

**Arguments**

decimal_num	decimal number between 0 and $(2^{31}) - 1$
bits	number of bits to keep in the result counting from the right. <b>Default value is 32.</b>

**Value**

a binary string representation of the given decimal number.

**Examples**

```
# representing 0
dec_to_bin(0,1)
dec_to_bin(0,10)
dec_to_bin(0,32)
dec_to_bin(0)
```

```
# representing 24
dec_to_bin(24,6)
dec_to_bin(24,21)
```

```
dec_to_bin(24)
dec_to_bin(24,3) # note that this will cut the returned result so be careful!
```

---

get\_common\_names      *Get the common names of two vectors*

---

## Description

This function prints and returns the common names of two vectors. The two vectors don't have to be the same length.

## Usage

```
get_common_names(vec1, vec2, vector.names.str = "nodes", with.gt = TRUE)
```

## Arguments

vec1	vector with names attribute
vec2	vector with names attribute
vector.names.str	string. Used for printing, it tell us what are the names of the two vectors (use plural form). Default value: "nodes".
with.gt	logical. Determines if the ">" sign will be appended for nice printing in an R notebook (use with the chunk option <i>results = 'asis'</i> ). Default value: TRUE.

## Value

the character vector of the common names. If there is only one name in common, the `vector.names.str` gets the last character stripped for readability. If there is no common names, it returns FALSE.

## See Also

[pretty\\_print\\_vector\\_values](#), [pretty\\_print\\_string](#)

## Examples

```
vec1 = c(1,1,1)
vec2 = c(1,2)
names(vec1) = c("a","b","c")
names(vec2) = c("c","b")

common.names = get_common_names(vec1, vec2)
```

---

get_common_values	<i>Get the common values of two vectors</i>
-------------------	---

---

### Description

This function prints and returns the common values of two vectors. The two vectors don't have to be the same length.

### Usage

```
get_common_values(vec1, vec2, vector.values.str = "nodes", with.gt = TRUE)
```

### Arguments

vec1	vector
vec2	vector
vector.values.str	string. Used for printing, it tell us what are the values of the two vectors (use plural form). Default value: "nodes".
with.gt	logical. Determines if the ">" sign will be appended for nice printing in an R notebook (use with the chunk option <i>results = 'asis'</i> ). Default value: TRUE.

### Value

the vector of the common values. If there is only one value in common, the `vector.values.str` gets the last character stripped for readability. If there are no common values, it returns NULL.

### See Also

[pretty\\_print\\_vector\\_values](#), [pretty\\_print\\_string](#)

### Examples

```
vec1 = c(1,2,3)
vec2 = c(3,4,1)

common.names = get_common_values(vec1, vec2)
```

---

get_parent_dir	<i>Retrieve the parent directory</i>
----------------	--------------------------------------

---

**Description**

Use this function to retrieve the parent directory from a string representing the full path of a file or a directory.

**Usage**

```
get_parent_dir(pathStr)
```

**Arguments**

pathStr	string. The name of the directory, can be a full path filename.
---------	---

**Value**

a string representing the parent directory. When a non-file path is used as input (or something along those lines :) then it returns the root ("/") directory.

**Examples**

```
get_parent_dir("/home/john")
get_parent_dir("/home/john/a.txt")
get_parent_dir("/home")
```

---

get_percentage_of_matches	<i>Get percentage of matches between two vectors</i>
---------------------------	--

---

**Description**

Use this function on two numeric vectors with the same names attribute (columns) and same length, in order to find the percentage of common elements (value matches between the two vectors). The same names for the two vectors ensures that their values are logically matched one-to-one.

**Usage**

```
get_percentage_of_matches(vec1, vec2)
```

**Arguments**

vec1	numeric vector with names attribute
vec2	numeric vector with names attribute



**Value**

the percentage of common values (exact matches) between the two vectors. Can only be a value between 0 (no common elements) and 1 (perfect element match). Note that *NaN* and *NA* values are allowed in the input vectors, but they will always count as a mismatch.

**Examples**

```
vec1 = c(1, 2, 3, 2)
vec2 = c(20, 2, 2.5, 8)
vec3 = c(1, 2, 333, 222)
names.vec = c(seq(1,4))
names(vec1) = names.vec
names(vec2) = names.vec
names(vec3) = names.vec

match.1.2 = get_percentage_of_matches(vec1, vec2)
match.1.3 = get_percentage_of_matches(vec1, vec3)
```

---

get\_roc\_stats

*Generate ROC statistics*


---

**Description**

Use this function to generate the most useful statistics related to the generation of a basic ROC (Receiver Operating Characteristic) curve.

**Usage**

```
get_roc_stats(df, pred_col, label_col, direction = "<")
```

**Arguments**

df	a data.frame with (at least) two columns. See next two parameters for what values these two columns should have (which should match one to one).
pred_col	string. The name of the column of the df data.frame that has the prediction values. The values can be any numeric, negative, positive or zero. What matters is the <b>ranking</b> of these values which is clarified with the direction parameter.
label_col	string. The name of the column of the df data.frame that has the true positive labelings/observed classes for the prediction values. This column must have either 1 or 0 elements representing either a <i>positive</i> or <i>negative</i> classification label for the corresponding values.
direction	string. Can be either > or < (default value) and indicates the direction/ranking of the prediction values with respect to the positive class labeling (for a specific threshold). If <b>smaller</b> prediction values indicate the positive class/label use < whereas if <b>larger</b> prediction values indicate the positive class/label (e.g. probability of positive class), use >.

**Value**

A list with two elements:

- `roc_stats`: a tibble which includes the **thresholds** for the ROC curve and the **confusion matrix stats** for each threshold as follows: *TP* (#True Positives), *FN* (#False Negatives), *TN* (#True Negatives), *FP* (#False Positives), *FPR* (False Positive Rate - the x-axis values for the ROC curve) and *TPR* (True Positive Rate - the y-axis values for the ROC curve). Also included are the *dist-from-chance* (the vertical distance of the corresponding (FPR,TPR) point to the chance line or positive diagonal) and the *dist-from-0-1* (the euclidean distance of the corresponding (FPR,TPR) point from (0,1)).
- *AUC*: a number representing the Area Under the (ROC) Curve.

The returned results provide an easy way to compute two optimal *cutpoints* (thresholds) that dichotomize the predictions to positive and negative. The first is the *Youden index*, which is the maximum vertical distance from the ROC curve to the chance line or positive diagonal. The second is the point of the ROC curve closest to the (0,1) - the point of perfect differentiation. See examples below.

**Examples**

```
# load libraries
library(readr)
library(dplyr)

# load test tibble
test_file = system.file("extdata", "test_df.tsv", package = "usefun", mustWork = TRUE)
test_df = readr::read_tsv(test_file, col_types = "di")

# get ROC stats
res = get_roc_stats(df = test_df, pred_col = "score", label_col = "observed")

# Plot ROC with a legend showing the AUC value
plot(x = res$roc_stats$FPR, y = res$roc_stats$TPR,
     type = 'l', lwd = 2, col = '#377EB8', main = 'ROC curve',
     xlab = 'False Positive Rate (FPR)', ylab = 'True Positive Rate (TPR)')
legend('bottomright', legend = round(res$AUC, digits = 3),
      title = 'AUC', col = '#377EB8', pch = 19)
grid()
abline(a = 0, b = 1, col = '#FF726F', lty = 2)

# Get two possible cutoffs
youden_index_df = res$roc_stats %>%
  filter(dist_from_chance == max(dist_from_chance))
min_classification_df = res$roc_stats %>%
  filter(dist_from_0_1 == min(dist_from_0_1))
```

---

`get_stats_for_unique_values`*Get stats for unique values*

---

**Description**

Use this function on two vectors with same names attribute (column names), to find for each unique (numeric) value of the first vector, the average and standard deviation values of the second vector's values (matching is done by column name)

**Usage**

```
get_stats_for_unique_values(vec1, vec2)
```

**Arguments**

<code>vec1</code>	vector with names attribute
<code>vec2</code>	vector with names attribute

**Value**

A data.frame consisting of 3 column vectors. The data.frame size is  $n \times 3$ , where  $n$  is the number of unique values of `vec1` (rows). The columns vectors are:

1. the first input vector pruned to its unique values
2. a vector with the average values for each unique value of the first vector (the matching is done by column name)
3. a vector with the standard deviation values for each unique value of the first vector (the matching is done by column name)

**Examples**

```
vec1 = c(1, 2, 3, 2)
vec2 = c(20, 2, 2.5, 8)
names.vec = c(seq(1,4))
names(vec1) = names.vec
names(vec2) = names.vec

res = get_stats_for_unique_values(vec1, vec2)
```

---

get\_ternary\_class\_id    *Get ternary class id*

---

### Description

Helper function that checks if a *value* surpasses the given *threshold* either positively, negatively or not at all and returns a value indicating in which class (i.e. interval) it belongs.

### Usage

```
get_ternary_class_id(value, threshold)
```

### Arguments

value	numeric
threshold	numeric

### Value

an integer. There are 3 cases:

- 1: when *value* > *threshold*
- -1: when *value* < *-threshold*
- 0: otherwise

---

is\_between                    *Is value between two others?*

---

### Description

This function checks if a given value is inside an interval specified by two boundary values.

### Usage

```
is_between(value, low.thres, high.thres, include.high.value = FALSE)
```

### Arguments

value	numeric
low.thres	numeric. Lower boundary of the interval.
high.thres	numeric. Upper boundary of the interval.
include.high.value	logical. Whether the upper bound is included in the interval or not. Default value: FALSE.

**Value**

a logical specifying if the value is inside the interval [low.thres,high.thres) (default behaviour) or inside the interval [low.thres,high.thres] if include.high.value is TRUE.

**Examples**

```
is_between(3,2,4)
is_between(4,2,4)
is_between(4,2,4,include.high.value=TRUE)
```

---

is\_empty

*Is object empty?*

---

**Description**

A function to test whether an object is **empty**. It checks the length of the object, so it has different behaviour than [is.null](#).

**Usage**

```
is_empty(obj)
```

**Arguments**

obj            a general object

**Value**

a logical specifying if the object is NULL or not.

**Examples**

```
# TRUE
is_empty(NULL)
is_empty(c())

# FALSE
is_empty("")
is_empty(NA)
is_empty(NaN)
```

ldf\_arrange\_by\_rownames

*Rearrange a list of data frames by rownames*

---

## Description

Rearrange a list of data frames by rownames

## Usage

```
ldf_arrange_by_rownames(list_df)
```

## Arguments

`list_df` a (non-empty) list of `data.frame` objects. The data frames must have the same `colnames` attribute.

## Value

a rearranged list of data frames, where the names of the elements of the `list_df` (the 'ids' of the data frames) and the rownames of the data frames have switched places: the unique row names of the original list's combined data frames serve as names for the returned list of data frames, while the data frame 'ids' (names of the original list's elements) now serve as rownames for the data frames in the new list.

E.g. if in the given list there was a data.frame with id 'A': `a = list_df[["A"]]` and `rownames(a) = c("row1", "row2")`, then in the rearranged list there would be two data frames with ids "row1" and "row2", each of them having a row with name "A" where also these data rows would be the same as before: `list_df[["A"]][["row1", ] == returned_list[["row1"]][["A", ]` and `list_df[["A"]][["row2", ] == returned_list[["row2"]][["A", ]` respectively.

## Examples

```
df.1 = data.frame(matrix(data = 0, nrow = 3, ncol = 3,
  dimnames = list(c("row1", "row2", "row3"), c("C.1", "C.2", "C.3"))))
df.2 = data.frame(matrix(data = 1, nrow = 3, ncol = 3,
  dimnames = list(c("row1", "row2", "row4"), c("C.1", "C.2", "C.3"))))
list_df = list(df.1, df.2)
names(list_df) = c("zeros", "ones")
res_list_df = ldf_arrange_by_rownames(list_df)
```

---

make\_color\_bar\_plot    *Make a color bar plot*

---

### Description

Use this function when you want to visualize some numbers and their respective color values. Note that more than 42 colors won't be nice to see (too thin bars)!

### Usage

```
make_color_bar_plot(color.vector, number.vector, title, x.axis.label = "")
```

### Arguments

`color.vector`    vector of color values  
`number.vector`    vector of numeric values (same length with `color.vector`)  
`title`            string. The title of the barplot  
`x.axis.label`    string. The x-axis label. Default value: empty string

### Examples

```
color.vector = rainbow(10)  
number.vector = 1:10  
title = "First 10 rainbow() colors"  
make_color_bar_plot(color.vector, number.vector, title)
```

---

make\_multiple\_density\_plot  
                          *Multiple densities plot*

---

### Description

Combine many density distributions to one common plot.

### Usage

```
make_multiple_density_plot(  
  densities,  
  legend.title,  
  title,  
  x.axis.label,  
  legend.size = 1  
)
```

**Arguments**

densities	a list, each element holding the results from executing the <code>density</code> function to a (different) vector. Note that you need to provide a name for each list element for the legend (see example).
legend.title	string. The legend title.
title	string. The plot title.
x.axis.label	string. The x-axis label.
legend.size	numeric. Default value: 1.

**Examples**

```
mat = matrix(rnorm(60), ncol=20)
densities = apply(mat, 1, density)
names(densities) = c("1st", "2nd", "3rd")
make_multiple_density_plot(densities, legend.title = "Samples",
  x.axis.label = "", title = "3 Normal Distribution Samples")
```

---

mat\_equal

*Matrix equality*


---

**Description**

Check if two matrices are equal. Equality is defined by both of them being matrices in the first place, having the same dimensions as well as the same elements.

**Usage**

```
mat_equal(x, y)
```

**Arguments**

x, y	matrices
------	----------

**Value**

a logical specifying if the two matrices are equal or not.



---

normalize\_to\_range      *Range normalization*

---

**Description**

Normalize a vector, matrix or data.frame of numeric values in a specified range.

**Usage**

```
normalize_to_range(x, range = c(0, 1))
```

**Arguments**

x	vector, matrix or data.frame with at least two different elements
range	vector of two elements specifying the desired normalized range. Default value is c(0,1)

**Value**

the normalized data

**Examples**

```
vec = 1:10
normalize_to_range(vec)
normalize_to_range(vec, range = c(-1,1))

mat = matrix(c(0,2,1), ncol = 3, nrow = 4)
normalize_to_range(mat, range = c(-5,5))
```

---

outersect      *Outersect*

---

**Description**

Performs set *outersection* on two vectors. The opposite operation from intersect!

**Usage**

```
outersect(x, y)
```

**Arguments**

x, y	vectors
------	---------

**Value**

a vector of the non-common elements of x and y.

**See Also**

[intersect](#)

**Examples**

```
x = 1:10
y = 2:11

# c(1,11)
outersect(x,y)
```

---

partial\_permut

*Get partial permutation of a vector*

---

**Description**

Get partial permutation of a vector

**Usage**

```
partial_permut(x, exp_sim = 0)
```

**Arguments**

x	a vector with at least 2 elements
exp_sim	a value between 0 and 1 indicating the level of <i>expected similarity</i> between the input and output vector. Default value is <b>0</b> (random permutation).

**Value**

a partially (random) permuted vector. If `exp_sim = 0` then the result is equal to `sample(x)` (a random permutation). If `exp_sim = 1` then the result is always the same as the input vector. For `exp_sim` values between 0 and 1 we randomly sample a subset of the input vector inversely proportionate to the `exp_sim` value (e.g. `exp_sim = 0.8` => 20% of the elements) and randomly permute these elements only.

**Examples**

```
set.seed(42)
partial_permut(x = LETTERS, exp_sim = 0)
partial_permut(x = LETTERS, exp_sim = 0.5)
partial_permut(x = LETTERS, exp_sim = 0.9)
```

---

powerset_icounts	<i>Powerset Intersection Table</i>
------------------	------------------------------------

---

## Description

This function computes the intersection of elements for all possible combinations of the provided sets of IDs. A typical use case is in a cohort of patients with incomplete data across multiple data types. This function helps determine how many patients have complete data for specific combinations of data types, allowing you to find the optimal combinations for analysis.

## Usage

```
powerset_icounts(ids)
```

## Arguments

ids	list() A named list, each element being a numeric vector of ids.
-----	---

## Value

A tibble with columns:

- set\_combo: name for combo set/vector
- num\_subsets: number of subsets in the combo set
- common\_ids: vector of common ids in the combo set
- count: number of common ids

## Examples

```
library(dplyr)
ids = list(a = 1:3, b = 2:5, c = 1:4, d = 3:6, e = 2:6)
res = powerset_icounts(ids)

res |>
  filter(num_subsets >= 2, count > 2) |>
  arrange(desc(count), desc(num_subsets))
```

pr.boot

*Bootstrap Confidence Intervals for Precision-Recall Curves***Description**

This functions calculates bootstrap percentile CIs for PR curves using **precrec**. These can then be used in a plotting function, see example.

**Usage**

```
pr.boot(
  labels,
  preds,
  boot.n = 10000,
  boot.stratified = TRUE,
  alpha = 0.1,
  ...
)
```

**Arguments**

labels	(numeric()) Vector of responses/labels (only two classes/values allowed: cases/positive class = 1 and controls/negative class = 0)
preds	(numeric()) Vector of prediction values. Higher values denote positive class.
boot.n	(numeric(1)) Number of bootstrap resamples. Default: 10000
boot.stratified	(logical(1)) Whether the bootstrap resampling is stratified (same number of cases/controls in each replicate as in the original sample) or not. It is advised to use stratified resampling when classes from labels are imbalanced. Default: TRUE.
alpha	(numeric(1)) Confidence level for bootstrap percentile interval (between 0 and 1). Default is 0.1, corresponding to 90% confidence intervals.
...	Other parameters to pass on to <code>precrec::evalmod</code> , except mode (set to rocpr) and raw_curves (set to TRUE). For example x_bins indicates the minimum number of recall points on the x-axis.

**Value**

A tibble with columns:

- recall: recall of original data
- precision: precision of original data

- `low_precision`: low value of the bootstrap confidence interval
- `high_precision`: high value of the bootstrap confidence interval

## References

Saito, Takaya, Rehmsmeier, Marc (2016). “Precrec: fast and accurate precision-recall and ROC curve calculations in R.” *Bioinformatics*, **33**(1), 145–147. doi:10.1093/bioinformatics/btw570.

## Examples

```
set.seed(42)
# imbalanced labels
labels = sample(c(0,1), 100, replace = TRUE, prob = c(0.8,0.2))
# predictions
preds = rnorm(100)

# get CIs for PR curve
pr_tbl = pr.boot(labels, preds, boot.n = 100, x_bins = 30) # default x_bin is 1000
pr_tbl

# draw PR curve + add the bootstrap percentile confidence bands
library(ggplot2)

pr_tbl |>
  ggplot(aes(x = recall, y = precision)) +
  geom_step() +
  ylim(c(0,1)) +
  geom_ribbon(aes(ymin = precision_low, ymax = precision_high), alpha = 0.2)
```

---

pr.test

*Compare two Precision-Recall curves*

---

## Description

Test the hypothesis that the true difference in PR AUCs is equal to 0. We implement the same bootstrap method based on the idea from `pROC::roc.test()`. The PR AUC is calculated using `PRROC::pr.curve()` with the interpolation method of Davis (2006).

## Usage

```
pr.test(
  labels,
  pred1,
  pred2,
  boot.n = 10000,
  boot.stratified = TRUE,
  alternative = "two.sided"
)
```

**Arguments**

labels	(numeric()) Vector of responses/labels (only two classes/values allowed: cases/positive class = 1 and controls/negative class = 0)
pred1	(numeric()) Vector of prediction values. Higher values denote positive class.
pred2	(numeric()) Vector of prediction values. Higher values denote positive class. Must have the same length as pred1.
boot.n	(numeric(1)) Number of bootstrap resamples. Default: 10000
boot.stratified	(logical(1)) Whether the bootstrap resampling is stratified (same number of cases/controls in each replicate as in the original sample) or not. It is advised to use stratified resampling when classes from labels are imbalanced. Default: TRUE.
alternative	(character(1)) Specifies the alternative hypothesis. Either "two.sided", "less" or "greater". Default: "two.sided".

**Value**

a list with the AUCs of the two original prediction vectors and the p-value of the bootstrap-based test.

**References**

Davis J, Goadrich M (2006). “The relationship between precision-recall and ROC curves.” *Proceedings of the 23rd International Conference on Machine Learning*, **148**(4), 233–240. doi:[10.1145/1143844.1143874](https://doi.org/10.1145/1143844.1143874).

**Examples**

```
set.seed(42)
# imbalanced labels
labels = sample(c(0,1), 20, replace = TRUE, prob = c(0.8,0.2))
# predictions
pred1 = rnorm(20)
pred2 = rnorm(20)
pr.test(labels, pred1, pred2, boot.n = 1000, boot.stratified = FALSE)
pr.test(labels, pred1, pred2, boot.n = 1000, boot.stratified = TRUE)
```

---

pretty\_print\_bold\_string  
*Pretty print a bold string*

---

**Description**

Prints a bold string only when `html.output` is enabled. Otherwise, it prints a normal string. The the ">" sign can be appended if nice output in an R notebook is desired.

**Usage**

```
pretty_print_bold_string(string, with.gt = TRUE, html.output = TRUE)
```

**Arguments**

<code>string</code>	a string
<code>with.gt</code>	logical. Determines if the ">" sign will be appended for nice printing in an R notebook. (use with the chunk option <i>results = 'asis'</i> ). Default value: TRUE.
<code>html.output</code>	logical. If TRUE, it encapsulates the string with the bold tags for an HTML document. Default value: TRUE.

**See Also**

[pretty\\_print\\_string](#)

---

pretty\_print\_name\_and\_value  
*Pretty print a name and value*

---

**Description**

Pretty print a name and value

**Usage**

```
pretty_print_name_and_value(name, value, with.gt = FALSE, with.comma = TRUE)
```

**Arguments**

<code>name</code>	string
<code>value</code>	string
<code>with.gt</code>	logical. Determines if the ">" sign will be appended for nice printing in an R notebook (use with the chunk option <i>results = 'asis'</i> ). Default value: FALSE.
<code>with.comma</code>	logical. Determines if the comma (,) character will be appended to the end of the output. Default value: TRUE.

## Examples

```
pretty_print_name_and_value("aName", "aValue", with.gt = TRUE)
pretty_print_name_and_value("aName", "aValue", with.comma = FALSE)
```

---

pretty\_print\_string    *Pretty print a string*

---

## Description

Nice printing of a string in an R notebook (default behaviour). Otherwise, it prints the string to the standard R output.

## Usage

```
pretty_print_string(string, with.gt = TRUE)
```

## Arguments

string	a string
with.gt	logical. Determines if the ">" sign will be appended for nice printing in an R notebook (use with the chunk option <i>results = 'asis'</i> ). Default value: TRUE.

## See Also

[cat](#)

---

pretty\_print\_vector\_names  
*Pretty printing of a vector's names attribute*

---

## Description

Pretty printing of a vector's names attribute

## Usage

```
pretty_print_vector_names(  
  vec,  
  vector.names.str = "nodes",  
  sep = ", ",  
  with.gt = TRUE  
)
```



**Arguments**

vec	vector
vector.names.str	string. It tell us what are the names of the vector (use plural form) in order to fill the print message. Default value: "nodes".
sep	string. The separator character to use to distinguish between the names values. Default value: ", ".
with.gt	logical. Determines if the ">" sign will be appended for nice printing in an R notebook (use with the chunk option <i>results = 'asis'</i> ). Default value: TRUE.

**See Also**

[pretty\\_print\\_string](#)

---

pretty\_print\_vector\_names\_and\_values

*Pretty printing of a vector's names and values*

---

**Description**

It outputs a vector's names and values in this format: *name1: value1, name2: value2,...*. You can choose how many elements to show in this format. Use with the chunk option *results = 'asis'* to get a nice printing in an R notebook.

**Usage**

```
pretty_print_vector_names_and_values(vec, n = -1)
```

**Arguments**

vec	vector with names attribute
n	the number of elements that you want to print in a nice way. Default value: -1 (pretty print all elements). For any $n < 1$ , all elements are printed.

**See Also**

[pretty\\_print\\_name\\_and\\_value](#)

pretty\_print\_vector\_values

*Pretty printing of a vector's values*

---

### Description

Pretty printing of a vector's values

### Usage

```
pretty_print_vector_values(  
  vec,  
  vector.values.str = "nodes",  
  sep = ", ",  
  with.gt = TRUE  
)
```

### Arguments

vec	vector
vector.values.str	string. It tell us what are the values of the vector (use plural form) in order to fill the print message. Default value: "nodes".
sep	string. The separator character to use to distinguish between the vector values. Default value: ", ".
with.gt	logical. Determines if the ">" sign will be appended for nice printing in an R notebook (use with the chunk option <i>results = 'asis'</i> ). Default value: TRUE.

### See Also

[pretty\\_print\\_string](#)

---

print\_empty\_line

*Print an empty line*

---

### Description

Print an empty line

### Usage

```
print_empty_line(html.output = FALSE)
```

**Arguments**

html.output      logical. If TRUE, it outputs an empty line for an HTML document, else an empty line for the standard R output. Default value: FALSE.

**See Also**

[cat](#)

---

prune\_and\_reorder\_vector

*Prune and reorder vector elements*

---

**Description**

Given two vectors, the first one's elements are pruned and reordered according to the common values of the second vector and the elements' names (*attribute*) of the first. If there no common such values, an empty vector is returned.

**Usage**

```
prune_and_reorder_vector(vec, filter.vec)
```

**Arguments**

vec                      a vector with names attribute  
filter.vec                a character vector whose values will be used to filter the vec elements

**Value**

the pruned and re-arranged vector.

**Examples**

```
vec = c(1,2,3)
names(vec) = c("a", "b", "c")

filter.vec1 = c("a")
prune_and_reorder_vector(vec, filter.vec1)

filter.vec2 = c("c", "ert", "b")
prune_and_reorder_vector(vec, filter.vec2)
```

---

prune\_columns\_from\_df *Prune single-value columns from a data frame*

---

**Description**

Given a `data.frame` and an integer value, it checks whether there is a column vector whose values match the given one. If so, it prunes that single-valued column from the `data.frame`

**Usage**

```
prune_columns_from_df(df, value)
```

**Arguments**

<code>df</code>	<code>data.frame</code>
<code>value</code>	an integer value

**Value**

the column-pruned `data.frame`

**Examples**

```
df = data.frame(c(0,0,0), c(0,1,0), c(1,0,0))
prune_columns_from_df(df, value = 0)
```

---

prune\_rows\_from\_df *Prune single-value rows from a data frame*

---

**Description**

Given a `data.frame` and an integer value, it checks whether there is a row vector whose values match the given one. If so, it prunes that single-valued row from the `data.frame`

**Usage**

```
prune_rows_from_df(df, value)
```

**Arguments**

<code>df</code>	<code>data.frame</code>
<code>value</code>	an integer value

**Value**

the row-pruned data.frame

**Examples**

```
df = data.frame(c(0,0,0), c(0,1,0), c(1,0,0))
prune_rows_from_df(df, value = 0)
```

---

remove\_commented\_and\_empty\_lines

*Remove commented and empty lines*

---

**Description**

Removes empty or commented lines from a character vector (each element being a line)

**Usage**

```
remove_commented_and_empty_lines(lines)
```

**Arguments**

lines            a character vector, usually the result from using the [readLines](#) function

**Value**

a character vector of the pruned lines

---

save\_df\_to\_file

*Save data frame to a specified file*

---

**Description**

Function for saving a data.frame to a specified file. Column and row names are written by default and the *tab* is used as a delimiter.

**Usage**

```
save_df_to_file(df, file)
```

**Arguments**

df                data.frame

file              string. The name of the file, can be a full path.

---

save_mat_to_file	<i>Save matrix to a specified file</i>
------------------	--

---

**Description**

Function for saving a matrix to a specified file. Uses the [save\\_df\\_to\\_file](#) function.

**Usage**

```
save_mat_to_file(mat, file)
```

**Arguments**

mat	matrix
file	string. The name of the file, can be a full path.

---

save_vector_to_file	<i>Save vector to a specified file</i>
---------------------	--

---

**Description**

Function for saving a vector with or without its row names to a specified file. By default the *tab* is used as a delimiter.

**Usage**

```
save_vector_to_file(vector, file, with.row.names = FALSE)
```

**Arguments**

vector	vector
file	string. The name of the file, can be a full path.
with.row.names	logical. If TRUE, then the names(vector) will be included in the output file. Default value: FALSE.

---

specify_decimal	<i>Specify decimal</i>
-----------------	------------------------

---

**Description**

Use this function to transform a given decimal number to the desired precision by choosing the number of digits after the decimal point.

**Usage**

```
specify_decimal(number, digits.to.keep)
```

**Arguments**

number            numeric

digits.to.keep    numeric. Refers to the digits to keep after decimal point '.'. This value should be 15 or less.

**Value**

the pruned number in string format

**Examples**

```
# 0.123  
specify_decimal(0.1233213, 3)
```

---

usefun	<i>usefun</i>
--------	---------------

---

**Description**

A collection of useful functions by John

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