

# Package ‘MoffittFunctions’

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**Title** Moffitt Biostat Functions

**Version** 0.1.0

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**Description** Statistical, data processing, and annotation functions for Moffitt Biostat.

**License** GPL-3

**LazyData** true

**Imports** knitr, devtools, data.table, kableExtra, coin, Exact

**Suggests** testthat

**RoxygenNote** 6.0.1

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cor_test	<i>Correlation Test for Two Variables</i>
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## Description

A wrapper for cor.test function, except if spearman selected and ties in at least one variable, in which case this is a wrapper for coin::spreaman\_test in with approximate method.

## Usage

```
cor_test(x, y, method = c("pearson", "kendall", "spearman"),
  seed = 68954857, B = 10000, exact = TRUE, verbose = FALSE)
```

**Arguments**

x	numeric vector (can include NA values)
y	numeric vector (can include NA values)
method	a character string indicating which correlation coefficient is to be used for the test. One of "pearson", "kendall", or "spearman", can be abbreviated
seed	seed (only used if method = "spearman")
B	number of reps (only used if method = "spearman")
exact	if no ties should you do exact (TRUE) or asymptotic (FALSE) method. Exact method is the default (only used if method = "spearman" (and no ties) or "kendall")
verbose	a logical variable indicating if warnings and messages should be displayed

**Details**

To always get reproducible results when using approximate method we need to set seed inside of the call, and order the data

**Value**

spearman\_test pvalue

**Examples**

```
set.seed(5432322)
x <- rnorm(20,0,3)
y <- x + rnorm(20,0,5)
cor_test(x,y, method = 'pearson')
cor_test(x,y, method = 'kendall')
cor_test(x,y, method = 'spearman')
```

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paste\_tbl\_grp

*Pasting Together Information for Two Groups*

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**Description**

Paste together information, often statistics, from two groups. There are two predefined combinations: mean(sd) and median[min,max], but user may also paste any single measure together.

**Usage**

```
paste_tbl_grp(data, vars_to_paste = "all", first_name = "Group1",
  second_name = "Group2", sep_val = " vs. ", na_str_out = "---",
  alternative = c("two.sided", "less", "greater"), digits = 0,
  trailing_zeros = TRUE, keep_all = TRUE, verbose = FALSE)
```

## Arguments

data	input dataset. User must use consistent naming throughout, <b>with an underscore</b> to separate the group names from the measures (i.e. Group1_mean and Group2_mean). There also must be two columns with column names that exactly match the input for first_name and second_name (i.e. 'Group1' and 'Group2'), which are used to form the Comparison variable.
vars_to_paste	vector of names of common measures to paste together. Can be the predefined 'median_min_max' or 'mean_sd', or any variable as long as they have matching columns for each group (i.e. Group1_MyMeasure and Group2_MyMeasure). Multiple measures can be requested. Default: "all" will run 'median_min_max' and 'mean_sd', as well as any pairs of columns in the proper format.
first_name	name of first group (string before '_'). Default is 'Group1'.
second_name	name of second group (string before '_'). Default is 'Group2'.
sep_val	value to be pasted between the two measures. Default is ' vs. '.
na_str_out	the character to replace missing values with.
alternative	a character string specifying the alternative hypothesis, must be one of "two.sided" (default), "greater" or "less". Will be used to determine the character to be pasted between the group names (Comparison variable). Specifying "two.sided" will use the sep_val input.
digits	integer indicating the number of decimal places to round to before pasting for numeric variables. Default is 0.
trailing_zeros	logical indicating if trailing zeros should be included (i.e. 0.100 instead of 0.1). Note if set to TRUE output is a character vector.
keep_all	logical indicating if all remaining, unpasted variables in data should be returned with the pasted variables. Default TRUE.
verbose	a logical variable indicating if warnings and messages should be displayed. Default FALSE.

## Details

User must use consistent naming throughout, with a underscore to separate the group names from the measures (i.e. Group1\_mean and Group2\_mean). There also must be columns defining the group names (i.e. Group1 and Group2), which are used to form the Comparison variable.

alternative included as a parameter so the direction can easily be seen in one-sided test. If "two.sided" is selected the value to be pasted between the two group names will be set to sep\_val, where "greater" will use ">" and "less" will use "<" as the pasting value.

## Value

data.frame with all the pasted values requested. Each name will have '\_comparison' at the end of the names (i.e. mean\_comparison, median\_comparison, ...)

## Examples

```
data(exampleData_BAMA)

descriptive_stats_by_group <- exampleData_BAMA[, .(
  Group1 = unique(group[group == 1]), Group2 = unique(group[group == 2]),
```

```

    Group1_n = length(magnitude[group == 1]), Group2_n = length(magnitude[group == 2]),
    Group1_mean = mean(magnitude[group == 1]), Group2_mean = mean(magnitude[group == 2]),
    Group1_sd = sd(magnitude[group == 1]), Group2_sd = sd(magnitude[group == 2]),
    Group1_median = median(magnitude[group == 1]), Group2_median = median(magnitude[group == 2]),
    Group1_min = min(magnitude[group == 1]), Group2_min = min(magnitude[group == 2]),
    Group1_max = max(magnitude[group == 1]), Group2_max = max(magnitude[group == 2])
  ), by = .(visitno, antigen)]

paste_tbl_grp(data = descriptive_stats_by_group, vars_to_paste = 'all', first_name = 'Group1', second_name = '
paste_tbl_grp(data = descriptive_stats_by_group, vars_to_paste = c("mean", "median_min_max"), alternative= "1
paste_tbl_grp(data = descriptive_stats_by_group, vars_to_paste = 'all', first_name = 'Group1', second_name = '

# Same example wit tidyverse (dplyr+tidyr) with some custom functions

library(dplyr)
library(tidyr)

q95_fun = function(x) quantile(x, 0.95)
N = function(x) length(x)

exampleData_BAMA %>%
  mutate(group = paste0("Group", group)) %>%
  group_by(group, visitno, antigen) %>%
  summarise_at("magnitude", funs(N, mean, sd, median, min, max, q95_fun)) %>%
  gather(variable, value, -(group:antigen)) %>% # these three chains create a wide dataset
  unite(temp, group, variable) %>%
  spread(temp, value) %>%
  mutate(Group1 = "Group 1", Group2 = "Group 2") %>%
  paste_tbl_grp()

```

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pretty\_pvalues

*Round and format a vector of p-values*


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## Description

pretty\_pvalues() takes a vector of p-values, rounds them to a specified digit amount, allows options for emphasizing p-values < the defined significance level, and returns a character for missing.

## Usage

```

pretty_pvalues(pvalues, digits = 3, bold = FALSE, italic = FALSE,
  background = NULL, sig_alpha = 0.05, missing_char = "---",
  include_p = FALSE, trailing_zeros = TRUE)

```

## Arguments

pvalues	numeric vector of raw p-values to be formatted
digits	number of digits to round to; values with zeros past this number of digits are truncated

<b>bold</b>	TRUE or FALSE: set to TRUE to bold p-values < the defined significance level
<b>italic</b>	TRUE or FALSE: set to TRUE to italicize p-values < the defined significance level
<b>background</b>	highlight color for p-values < the defined significance level. Default = NULL (no highlighting)
<b>sig_alpha</b>	the defined significance level. Default = 0.05
<b>missing_char</b>	character string that will replace missing values from the p-value vector. Default = "—"
<b>include_p</b>	TRUE or FALSE: set to TRUE to print "p = " before each p-value
<b>trailing_zeros</b>	TRUE or FALSE: default = TRUE, p-values are formatted with trailing zeros to the defined number of digits (i.e. 0.100 instead of 0.1 if digits = 3)

### Details

With this function, there are two things to be noted: Since the p-value vector formatting uses `cell_spec`, which generates raw HTML or LaTeX code, make sure you remember to put `escape = FALSE` into your kable code when generating your table. At the same time, you will need to escape special symbols manually. Additionally, `cell_spec` needs a way to know whether you want HTML or LaTeX output. You can specify it locally in the function or globally using `options(knitr.table.format = "latex")`. If you don't provide anything, this function will output as HTML by default.

### Value

Vector of transformed p-values for table output

### Examples

```
pvalue_example = c(1, 0.06, 0.0005, NA, 1e-6)

pretty_pvalues(pvalue_example, background = "pink")

pretty_pvalues(pvalue_example, digits = 4, missing_char = "missing", bold = TRUE)

# How to use pretty_pvalues in reports
raw_pvals <- c(0.00000007, .05000001, NaN, NA, 0.783)
pretty_pvals <- pretty_pvalues(raw_pvals, digits = 3, background = "green", italic = TRUE, bold = TRUE)
kableExtra::kable(pretty_pvals, format = "latex", escape = FALSE, col.names = c("P-values"))
```

---

round\_away\_0

*Rounding Using Round Away From 0 Method*


---

### Description

`round_away_0` takes a numeric vector, rounds them to a specified digit amount using the round away from 0 method for ties (i.e. 1.5). This is the SAS method for rounding.

### Usage

```
round_away_0(x, digits = 0, trailing_zeros = FALSE)
```

**Arguments**

- `x` numeric vector (can include NA values).
- `digits` positive integer of length 1 between 0 (default) and 14, giving the amount of digits to round to.
- `trailing_zeros` logical indicating if trailing zeros should included (i.e. 0.100 instead of 0.1). Note is set to TRUE output is a character vector

**Details**

`round_away_0` is not designed for use at precision levels  $\leq 1e-15$

**Value**

if `trailing_zeros = TRUE` returns a character vector of rounded values with trailing zeros, otherwise returns a numeric vector of rounded values.

**Examples**

```
vals_to_round = c(NA,-3.5:3.5,NA)
# [1] NA -3.5 -2.5 -1.5 -0.5 0.5 1.5 2.5 3.5 NA

# round() will round to even numbers when tied at X.5
round(vals_to_round)
# [1] NA -4 -2 -2 0 0 2 2 4 NA

# round_away_0() will round away from 0 when tied at X.5
round_away_0(vals_to_round)
# [1] NA -4 -3 -2 -1 1 2 3 4 NA

# Can force trailing zeros (will output character vector)
round_away_0(vals_to_round, digits = 2, trailing_zeros = TRUE)
```

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stat_paste	<i>Rounds and combines up to three numbers into table friendly presentation</i>
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**Description**

Takes in up to 3 numeric values, rounds each to a specified digit amount (if numeric), and then combines them accordingly.

**Usage**

```
stat_paste(stat1, stat2 = NULL, stat3 = NULL, digits = 0,
  trailing_zeros = TRUE, bound_char = c("(", "[", "{", "|"), sep = ", ",
  na_str_out = "---")
```

**Arguments**

stat1	first statistic to be pasted.
stat2	second statistic to be pasted (optional).
stat3	third statistic to be pasted (optional).
digits	positive integer of length 1 between 0 (default) and 14, giving the amount of digits to round to.
trailing_zeros	logical indicating if trailing zeros should included (i.e. 0.100 instead of 0.1). Note is set to TRUE output is a character vector
bound_char	the character to be used between stat1 and stat2/stat3. Available options are '(' (default), '[', '{', and 'l'.
sep	the string to be used between stat2 and stat3. The default is ', '.
na_str_out	the character to replace missing values with.

**Details**

One value provided - returns a rounded value or the missing character. Two values - returns stat1 (stat2). e.g., mean (sd) Three values - returns stat1 (stat2, stat3). e.g., estimate (95% lower, 95% upper) or median [min, max]

Currently the method does work with string variables, but of course rounding would not be relevant for strings.

**Value**

string of combined values

**Examples**

```
stat_paste(5.109293)
stat_paste(NA)
stat_paste(5.109293, 2.145)
stat_paste(5.109293, 1.7645, 8.0345)
stat_paste(NA, NA, NA)
stat_paste(5.109, "p < 0.001", digits = 3)
stat_paste(c(rep(5,5),NA),c(1:5,NA),c(1,NA,2,NA,3,NA),bound_char = '[')

library(data.table)
data(testData_BAMA)
testData_BAMA[,.(
  median_min_max = stat_paste(
    median(magnitude, na.rm = TRUE),
    min(magnitude, na.rm = TRUE),
    max(magnitude, na.rm = TRUE)
  )), by = .(antigen, visit, group)]
```

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two_samp_bin_test	<i>Binary (Response) Variable Compared to Binary (Group) Variable Test</i>
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---

### Description

Either Barnard, Fisher's, or Chi-sq test performed for unpaired data and McNemar's test for paired data

### Usage

```
two_samp_bin_test(x, y, method = NA, alternative = c("two.sided", "less",
"greater"), verbose = FALSE)
```

### Arguments

x	numeric vector (can include NA values).
y	vector with only 2 levels (can include NA values unless method = 'mcnemar').
method	what test to run ("barnard", "fisher", "chi.sq", "mcnemar"). No default so user must enter one of the four selections
alternative	a character string specifying the alternative hypothesis, must be one of "two.sided" (default), "greater" or "less". You can specify just the initial letter. Only "two.sided" available for method = 'chi.sq' or 'mcnemar'
verbose	a logical variable indicating if warnings and messages should be displayed.
...	parameters to pass to wilcox_test or t.test functions. For example the testing direction (alternative) in either call or the var.equal in the t.test function.

### Details

For one sided tests if y is a factor variable the level order is respected, otherwise the levels will set to alphabetical order (i.e. if alternative = less then testing  $a < b$ ).

If method = 'mcnemar' assumes the first observations of the first group matches the first observation of the second group, and so on. Also if method = 'mcnemar' then y must have the same number of samples for each level.

### Value

p-value for comparing x at the different levels of y.

### Examples

```
set.seed(5432322)
x <- c(sample(0:1,10,replace = TRUE, prob = c(.75,.25)), sample(0:1,10,replace = TRUE, prob = c(.25,.75)))
y <- c(rep('a', 10), rep('b', 10))
two_samp_bin_test(x,y, method = 'barnard')
two_samp_bin_test(x,y, 'fisher')
two_samp_bin_test(x,y, 'chi.sq')
two_samp_bin_test(x,y, 'mcnemar')
```



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two_samp_cont_test	<i>Continuous Variable Compared to Binary Variable Test</i>
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## Description

Either Wilcox or T-Test Performed, for unpaired or paired data

## Usage

```
two_samp_cont_test(x, y, method = c("wilcox", "t.test"), paired = FALSE,
  verbose = FALSE, ...)
```

## Arguments

x	numeric vector (can include NA values).
y	vector with only 2 levels (can include NA values unless paired = TRUE).
method	what test to run (wilcox or t-test).
paired	a logical indicating whether you want a paired test.
verbose	a logical variable indicating if warnings and messages should be displayed.
...	parameters to pass to wilcox_test or t.test functions. For example the testing direction (alternative) in either call or the var.equal in the t.test function.

## Details

Runs wilcox\_test() in the coin package, with "exact" distribution and mid-ranks ties method.

For one sided tests if y is a factor variable the level order is respected, otherwise the levels will set to alphabetical order (i.e. if alternative = less then testing  $a < b$ ).

If paired = TRUE assumes the first observations of the first group matches the first observation of the second group, and so on. Also if paired = TRUE then y must have the same number of samples for each level.

## Value

p-value for comparing x at the different levels of y.

## Examples

```
set.seed(5432322)
x <- c(rnorm(10,0,3), rnorm(10,3,3))
y <- c(rep('a', 10), rep('b', 10))
two_samp_cont_test(x = x, y = y, method = 'wilcox', paired = FALSE)
two_samp_cont_test(x = x, y = y, method = 'wilcox', paired = TRUE)
two_samp_cont_test(x = x, y = y, method = 't', paired = FALSE)
two_samp_cont_test(x = x, y = y, method = 't', paired = TRUE)
```

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