# Package 'sistmr'

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Type Package	
<b>Title</b> A Collection of Utility Function from the Inserm/Inria SISTM Team	
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<b>Description</b> Functions common to members of the SISTM team.	
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BlandAltmanPlot

Bland-Altman plot function

#### **Description**

Bland-Altman plot function

## Usage

```
BlandAltmanPlot(
  var1,
  var2,
  with_gradient = FALSE,
  line_color = c("blue", "lightblue"),
  extremum_pctg = TRUE
)
```

## **Arguments**

var1 a vector of numerics for the 1rst group to be compared. var2 a vector of numerics for the 2nd group to be compared.

with\_gradient a logical indicating if you have a lot of measures, use with\_gradient=TRUE to

have gradient scale and not points. Default value is FALSE.

line\_color a vector of color for the three lines: average difference and upper and lower

limits of the confidence interval for the average difference.

extremum\_pctg a logical indicating if you want to add the percentage of points outside the con-

fidence interval for the upper and lower limits. Default is TRUE.

#### Value

```
a ggplot2 object
```

## **Examples**

```
library(ggplot2)

#Small sample
#Generate data
x <- rnorm(30)
y <- rnorm(30, mean = 5, sd = 3)

#Plotting
BlandAltmanPlot(var1 = x, var2 = y)
#Add color by group
gr <- c(rep("G1", 15), rep("G2", 15))
BlandAltmanPlot(var1 = x, var2 = y) + geom_point(aes(color = gr))</pre>
```

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```
#High sample
#Generate data
x <- rnorm(10000)
y <- rnorm(10000, mean = 5, sd = 3)
#Plotting with gradient
BlandAltmanPlot(var1 = x, var2 = y, with_gradient = TRUE)</pre>
```

multipleBoxplots

Multiple boxplots for many times

#### **Description**

Multiple boxplots for many times

#### Usage

```
multipleBoxplots(data, x_var, y_var, add_points = TRUE)
```

## **Arguments**

data a dataset from which the variable x\_var and y\_var should be taken.

x\_var corresponding to the x coordinates for the plot, it must be a factor to obtain

multiple boxplots.

y\_var corresponding to the y coordinates for the plot.

add\_points if you want to add points on boxplots. Default value is TRUE.

#### Value

```
a ggplot2 object
```

#### **Examples**

```
library(ggplot2)

#Generate data
x_ex <- factor(c(rep("J0", 10), rep("J7", 10), rep("J14", 10)), levels = c("J0", "J7", "J14"))
y_ex <- rnorm(30)

data_ex <- cbind.data.frame(x_ex, y_ex)

#Plotting
multipleBoxplots(data = data_ex, x_var = x_ex, y_var = y_ex)

multipleBoxplots(data = data_ex, x_var = x_ex, y_var = y_ex) +
labs(x = "Time", y = "Value") +
theme(legend.position = "none")</pre>
```

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normal\_distribution

**Functions** 

#### **Description**

**Functions** 

#### Usage

```
normal_distribution(vec)
```

## Arguments

vec

a vector

#### Value

a vector

sistmr

sistmr.

## Description

This package contains functions common to members of the SISTM team.

volcanoPlot

Volcano plot function

## Description

Volcano plot function

## Usage

```
volcanoPlot(
  log2fc,
  pValue,
  data,
  FDR_threshold = 0.05,
  LFC_threshold = log2(1.5),
  color = c("red", "black"),
  geneNames = NULL,
  nb_geneTags = 20,
  logTransformPVal = TRUE
)
```

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### Arguments

log2fc a magnitude of change (fold-change) in base log 2 corresponding to the x-axis.

pValue a statistical significance (p-value) corresponding to the y-axis.

data a data frame of differentially expressed results from which the variable log2fc,

pValue and geneNames (if it is used) should be taken.

FDR\_threshold a threshold of false discovery rate. LFC\_threshold a threshold of log fold change.

color a vector of two colors for significant or not significant points.

geneNames a vector of gene names if you want to put gene tags on the volcano plot. Default

is NULL.

nb\_geneTags number of tags for the significant genes if geneNames is not NULL. Default is

20 to obtain the 20 first significant genes.

logTransformPVal

If TRUE, the p-values will have a negative logarithm transformation (base 10).

Default is TRUE.

#### Value

```
a ggplot2 object
```

#### **Examples**

```
genes <- paste0("G", 1:500)
pval <- runif(500, max = 0.5)
log2FC <- runif(500, min = -4, max = 4)

data <- cbind.data.frame(genes, pval, log2FC)
rm(genes, pval, log2FC)
volcanoPlot(log2FC, pval, data, geneNames = genes)</pre>
```

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