

# Package ‘Gmisc’

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**Title** Descriptive Statistics, Transition Plots, and More

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**Description** Tools for making the descriptive “Table 1” used in medical articles, a transition plot for showing changes between categories (also known as a Sankey diagram), flow charts by extending the grid package, a method for variable selection based on the SVD, Bézier lines with arrows complementing the ones in the 'grid' package, and more.

**License** GPL (>= 3)

**URL** <http://gforge.se>

**BugReports** <https://github.com/gforge/Gmisc/issues>

**Biarch** yes

**Depends** R (>= 3.2.0), Rcpp (>= 0.11.4), htmlTable (>= 1.13)

**Imports** grid, grDevices, graphics, lattice, Hmisc, forestplot, abind, methods, knitr, magrittr, rmarkdown, XML, utils, stats, checkmate, stringr

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Gmisc-package	<i>Collection of functions for plotting relations, generating tables, and more.</i>
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**Description**

This is a collection of functions that I've found useful in my research. The package is inspired by Frank Harrell's **Hmisc** package. The main focus is on tables, plots, and **knitr**-integration.

## Awesome tables

The `getDescriptionStatsBy` is a straight forward function that aims at helping you to generate descriptive table stratified by different variables. In other words, the function returns everything you need for generating a *Table 1* ready for publication. This function is accompanied by the `describeMean`, `describeMedian`, `describeProp`, and `describeFactors` functions.

The `mergeDesc` allows you to merge a set of outputs `getDescriptionStatsBy` into a `htmlTable` with the `rgroup` arguments automatically generated, see `vignette("descriptives", package="Gmisc")` for a detailed workflow description.

## Some fancy plots

The transition plot function, `transitionPlot`, is for descriptive purposes. It tries to illustrate the size of change between one state and the next, i.e. a transition. This is basically a graph of based upon `table(var1, var2)`.

The **Singular value decomposition** is a common method for reducing the number of variables. Unfortunately this compression can reduce the interpretability of the model. The `getSvdMostInfluential` function tries to remedy that by identifying the most influential elements from the V-matrix.

## Other stuff

The `insertRowAndKeepAttr` simply adds a row while remembering all the attributes previously set by using the `copyAllNewAttributes`. The `mergeLists` tries to merge lists that do not have identical elements.

---

bezierArrowGradient    *A bezier arrow with gradient*

---

## Description

This is an experimental addition to the original `bezierArrowSmp1` with the addition of a gradient in the center of the arrow that fades.

## Usage

```
bezierArrowGradient(x = c(0.2, 0.7, 0.3, 0.9), y = c(0.2, 0.2, 0.9,
  0.9), width = 0.05, clr = "#000000", default.units = "npc",
  align_2_axis = TRUE, grdt_type = c("triangle", "rectangle"),
  grdt_prop = 0.8, grdt_decrease_prop = 0.5, grdt_clr_prop = 0.7,
  grdt_line_width, grdt_clr = "#2F4F2F", vp = NULL, gp = gpar(),
  rm_intersect = 3L, ...)
```

**Arguments**

x	A numeric vector or unit object specifying x-locations of spline control points.
y	A numeric vector or unit object specifying y-locations of spline control points.
width	The width of the arrow, either a numeric single number or a unit. <b>Note:</b> The arrow does not rely on lwd but on actual width.
clr	The color of the arrow. This is the main color of the arrow and not the gradient color.
default.units	A string indicating the default units to use if x or y are only given as numeric vectors.
align_2_axis	Indicates if the arrow should be vertically/horizontally aligned. This is useful for instance if the arrow attaches to a box.
grdt_type	The type of growth and gradient that is to be used, currently it only supports triangle (I'm considering adding bezier curves but currently I'm a little tired of coding)
grdt_prop	The proportion of the full length that should be a the gradient. The gradient consists of three things: (1) the central band, (2) the slimming of the central band, (3) the color shift into the arrow color. <i>Note</i> that the the slimming and color proportions can be overlapping.
grdt_decrease_prop	The proportion of the gradient that should be decreasing, i.e. narrowing according to the grdt_type argument.
grdt_clr_prop	The proportion of the gradient that should be converging to the arrow color.
grdt_line_width	The width of the border line. If not specified it defaults to 5 % of the original width, note the gradient's width is thus 90 %.
grdt_clr	The color of the gradient.
vp	A Grid viewport object (or NULL).
gp	An object of class gpar, typically the output from a call to the function gpar. This is basically a list of graphical parameter settings.
rm_intersect	Set to 0 if you want to skip intersection removal, 1 only to remove left or 2 to only remove right. See details for why. @section Remove intersections: When the line is wide and the arrow has a narrow curve there may appear an empty triangle due to polygon cancellation (two polygons within the same are cancel out). This behaviour may be ugly and the function therefor tries to remove these. <i>Note:</i> it is expensive to check if there are the lines may intersect at one point, remove those unexpected, and then adjust the line to the new situation so that the top and bottom lines match. It can also cause some unexpected behaviour why you may want to remove this feature if the arrow behaves erratically.
...	Passed on to <a href="#">bezierArrowSmpl</a>

**Value**

grid::gList() A grob of [gList](#)-type

**Note**

The triangle section of the arrow is not currently included in the gradient.

**Examples**

```
library(grid)
grid.newpage()
arrowGrob <- bezierArrowGradient(x = c(.1,.3,.6,.9),
                                y = c(0.2, 0.2, 0.9, 0.9))
grid.draw(arrowGrob)
```

---

bezierArrowSmpl	<i>A simple bezier arrow</i>
-----------------	------------------------------

---

**Description**

This is an alternative to the grid packages [bezierGrob](#) with the advantage that it allows you to draw an arrow with a specific unit width. Note, it has only a end-arrow at this point.

**Usage**

```
bezierArrowSmpl(x = c(0.2, 0.7, 0.3, 0.9), y = c(0.2, 0.2, 0.9, 0.9),
               width = 0.05, clr = "#000000", default.units = "npc",
               arrow = list(), rez = 200, align_2_axis = TRUE, name = NULL,
               rm_intersect = 3L, gp = gpar(), vp = NULL)
```

**Arguments**

x	A numeric vector or unit object specifying x-locations of spline control points.
y	A numeric vector or unit object specifying y-locations of spline control points.
width	The width of the arrow, either a numeric single number or a unit. <b>Note:</b> The arrow does not rely on lwd but on actual width.
clr	The color of the arrow.
default.units	A string indicating the default units to use if x or y are only given as numeric vectors.
arrow	This is a list with all the <b>base</b> (width) and the desired <b>length</b> for the arrow. <b>Note:</b> This differs from the original <a href="#">bezierGrob</a> function.
rez	The resolution of the arrow. This specifies how many points to retrieve from the <a href="#">gnrlBezierPoints</a> function. Defaults to 200.
align_2_axis	Indicates if the arrow should be vertically/horizontally aligned. This is useful for instance if the arrow attaches to a box.

name	A character identifier.
rm_intersect	Set to 0 if you want to skip intersection removal, 1 only to remove left or 2 to only remove right. See details for why. @section Remove intersections: When the line is wide and the arrow has a narrow curve there may appear an empty triangle due to polygon cancellation (two polygons within the same are cancel out). This behaviour may be ugly and the function therefor tries to remove these. <i>Note:</i> it is expensive to check if there are the lines may intersect at one point, remove those unexpected, and then adjust the line to the new situation so that the top and bottom lines match. It can also cause some unexpected behaviour why you may want to remove this feature if the arrow behaves erratically.
gp	An object of class gpar, typically the output from a call to the function gpar. This is basically a list of graphical parameter settings.
vp	A Grid viewport object (or NULL).

**Value**

grid: :grob A grob of the class polygonGrob with attributes that correspond to the bezier points.

**Examples**

```
library(grid)
grid.newpage()
arrowGrob <- bezierArrowSmpl(x = c(.1,.3,.6,.9),
                             y = c(0.2, 0.2, 0.9, 0.9))
grid.draw(arrowGrob)
```

---

boxGrob	<i>Create a box with text</i>
---------	-------------------------------

---

**Description**

Creates a grob box with text inside it.

**Usage**

```
boxGrob(label, y = unit(0.5, "npc"), x = unit(0.5, "npc"), width,
        height, just = "center", bjust = "center",
        txt_gp = getOption("boxGrobTxt", default = gpar(color = "black")),
        box_gp = getOption("boxGrob", gpar(fill = "#D8F0D1")), name = NULL)

## S3 method for class 'box'
widthDetails(x)

## S3 method for class 'box'
```

```
heightDetails(x)

## S3 method for class 'box'
print(x, ...)

## S3 method for class 'box'
plot(x, ...)
```

### Arguments

label	The label to print - should be a number, text or expression.
y	The y position to put the box at. Can be either in npc (i.e. 0-1) or a <a href="#">unit</a> .
x	The x position to put the box at. Can be either in npc (i.e. 0-1) or a <a href="#">unit</a> .
width	The box autosizes but you can force by specifying the width
height	The box autosizes but you can force by specifying the height
just	The justification for the text: left, center or right.
bjust	The justification for the box: left, center, right, top or bottom. See the just option for the <a href="#">viewport</a>
txt_gp	The <a href="#">gpar</a> style to apply to the text. Set boxGrobTxt option if you want to customize all the boxes at once.
box_gp	The <a href="#">gpar</a> style to apply to the box. Set boxGrob option if you want to customize all the boxes at once.
name	a character identifier for the grob. Used to find the grob on the display list and/or as a child of another grob.
...	Passed to <a href="#">grid.draw</a>

### Value

A grob

### See Also

Other box-functions: [boxPropGrob](#), [connectGrob](#)

### Examples

```
library(grid)
grid.newpage()
boxGrob("My box")
```

---

 boxPropGrob

 Create a box with a color split
 

---

### Description

Creates a grob box with text inside it and a color split in the horizontal axes that allow indicating different proportions. The box can also have a title that spans the two color areas and that has its own background.

### Usage

```
boxPropGrob(label, label_left, label_right, prop, y = unit(0.5, "npc"),
  x = unit(0.5, "npc"), width, height, just = "center",
  bjust = "center", txt_gp = getOption("boxPropGrobTxt", default =
  gpar(color = "black")), txt_left_gp = getOption("boxPropGrobLeftTxt",
  default = gpar(col = "black")),
  txt_right_gp = getOption("boxPropGrobRightTxt", default = gpar(col =
  "black")), box_left_gp = getOption("boxPropGrobLeft", default =
  gpar(fill = "#81BFD4")), box_right_gp = getOption("boxPropGrobRight",
  default = gpar(fill = "#D8F0D1")),
  box_highlight_gp = getOption("boxPropGrobHighlight", default =
  gpar(fill = "#ffffff55", col = NA)), name = NULL)
```

### Arguments

label	The label to print - should be a number, text or expression.
label_left	The label for the left area
label_right	The label for the right area
prop	The proportion to split along
y	The y position to put the box at. Can be either in npc (i.e. 0-1) or a <a href="#">unit</a> .
x	The x position to put the box at. Can be either in npc (i.e. 0-1) or a <a href="#">unit</a> .
width	The box autosizes but you can force by specifying the width
height	The box autosizes but you can force by specifying the height
just	The justification for the text: left, center or right.
bjust	The justification for the box: left, center, right, top or bottom. See the <a href="#">just</a> option for the <a href="#">viewport</a>
txt_gp	The <a href="#">gpar</a> style to apply to the text. Set <code>boxPropGrobTxt</code> option if you want to customize all the boxes at once.
txt_left_gp	The <a href="#">gpar</a> style to apply to the left text. Set <code>boxPropGrobLeftTxt</code> option if you want to customize all the boxes at once.
txt_right_gp	The <a href="#">gpar</a> style to apply to the right text. Set <code>boxPropGrobRightTxt</code> option if you want to customize all the boxes at once.



box_left_gp	The <code>gpar</code> style to apply to the left box. Set <code>boxPropGrobLeft</code> option if you want to customize all the boxes at once.
box_right_gp	The <code>gpar</code> style to apply to the right box. Set <code>boxPropGrobRight</code> option if you want to customize all the boxes at once.
box_highlight_gp	The <code>gpar</code> style to apply to the background of the main label. Set <code>boxPropGrobHighlight</code> option if you want to customize all the boxes at once.
name	a character identifier for the grob. Used to find the grob on the display list and/or as a child of another grob.

**Value**

A box grob

**See Also**

Other box-functions: [boxGrob](#), [connectGrob](#)

**Examples**

```
library(grid)
grid.newpage()
boxPropGrob("Main label", "Left text", "Right text", prop = .3)
```

calculateLinesAndArrow

*Gets offsetted lines*

**Description**

The function calculates new points according to the offset that lie to the left/right of the provided line.

**Usage**

```
calculateLinesAndArrow(x, y, offset, end_x = -1, end_y = -1,
  arrow_offset = -1, rm_intersect = 3L)
```

**Arguments**

x	A numeric vector containing all the x-elements
y	A numeric vector containing all the y-elements
offset	The offset to add to the line, can be a vector if you want to use different offsets.
end_x	The x end of the line where the arrow occurs (if < 0 arrow is skipped)
end_y	The y end of the line where the arrow occurs (if < 0 arrow is skipped)
arrow_offset	The offset to add to the arrow section if any (if <= 0 arrow is skipped)

`rm_intersect` Set to 0 if you want to skip intersection removal, 1 only to remove left or 2 to only remove right. See details for why.

@section Remove intersections:

When the line is wide and the arrow has a narrow curve there may appear an empty triangle due to polygon cancellation (two polygons within the same are cancel out). This behaviour may be ugly and the function therefor tries to remove these.

*Note:* it is expensive to check if there are the lines may intersect at one point, remove those unexpected, and then adjust the line to the new situation so that the top and bottom lines match. It can also cause some unexpected behaviour why you may want to remove this feature if the arrow behaves erratically.

### Value

`list(list(x = ..., y = ...))` Returns a list with the right/left lines that in turn lists with *x* and *y* elements

---

connectGrob

*Connect boxes with an arrow*

---

### Description

The function creates a grob that links two boxes together. It looks for which side it should attach the arrow, e.g. if the start is on top of the bottom it should attach to the bottom edge of the start box and then to the top at the end.

### Usage

```
connectGrob(start, end, type = c("vertical", "horizontal", "L", "-", "Z",
  "N"), subelmnt = c("right", "left"),
  lty_gp = getOption("connectGrob", default = gpar(fill = "black")),
  arrow_obj = getOption("connectGrobArrow", default = arrow(ends =
  "last", type = "closed")))
```

```
## S3 method for class 'connect_boxes'
print(x, ...)
```

```
## S3 method for class 'connect_boxes'
plot(x, ...)
```

### Arguments

`start` The start box  
`end` The end box

type	How the boxes are stacked. The L alternative generates a straight line up/down and then turns to right/left for connecting with the end. The - generates a straight horizontal arrow. The Z creates a horizontal line that looks like a Z with 90 degree turns. The option N allows for vertical lines.
subelmnt	If we have a split box we can specify the right/left x as the connector point.
lty_gp	The <a href="#">gpar</a> for the line. Set connectGrob option if you want to customize all the arrows at once.
arrow_objj	The arrow spec according to <a href="#">arrow</a> . Set connectGrobArrow option if you want to customize all the arrows at once.
x	The grob to print/plot
...	Passed to <a href="#">grid.draw</a>

### Details

The exact positions of the line is stored at the `attr(..., "line")`. If you want to draw your own custom line all you need to do is check which `attr(my_line, "line")$x` and `attr(my_line, "line")$y` you want to attach to and then create your own custom [linesGrob](#).

### Value

grob with an arrow

### See Also

Other box-functions: [boxGrob](#), [boxPropGrob](#)

### Examples

```
library(grid)
grid.newpage()

# Initiate the boxes that we want to connect
start <- boxGrob("Top", x=.5, y=.8)
end <- boxGrob("Bottom", x=.5, y=.2)
side <- boxPropGrob("Side", "Left", "Right", prop=.3, x=.2, y=.8)
sub_side_left <- boxGrob("Left", x = attr(side, "coords")$left_x, y = .5)
sub_side_right <- boxGrob("Right", x = attr(side, "coords")$right_x, y = .5)
exclude <- boxGrob("Exclude:\n - Too sick\n - Prev. surgery", x=.8, y=.5,just="left")

# Connect the boxes and print/plot them
connectGrob(start, end, "vertical")
connectGrob(start, side, "horizontal")
connectGrob(side, sub_side_left, "v", "l")
connectGrob(side, sub_side_right, "v", "r")
connectGrob(start, exclude, "L")

# Print the grobs
start
end
side
```

```
exclude
sub_side_left
sub_side_right
```

---

```
convertShowMissing    A function for converting a show_missing variable.
```

---

### Description

The variable is supposed to be directly compatible with `table(..., useNA=show_missing)`. It throws an error if not compatible. It is mostly useful for custom describe functions.

### Usage

```
convertShowMissing(show_missing)
```

### Arguments

```
show_missing    Boolean or "no", "ifany", "always"
```

### Details

*Deprecated:* This function will be deprecated as all functions now use the useNA style in order to comply with standard R naming.

### Value

```
string
```

---

```
coords    Get the box coordinates
```

---

### Description

Retrieves the boxes "coords" attribute.

### Usage

```
coords(box)
```

### Arguments

```
box    The boxGrobs
```

### Value

A list with the coordinates

**Examples**

```
box <- boxGrob("A test box")
coords(box)
```

---

copyAllNewAttributes *A simple thing to keep the attributes*

---

**Description**

Skips the attributes that the to object already has to avoid overwriting dim and other important attributes

**Usage**

```
copyAllNewAttributes(from, to, attr2skip = c(), attr2force = c())
```

**Arguments**

from	The from object
to	The to object
attr2skip	An optional lists of attributes that you may want to avoid having copied
attr2force	An optional lists of attributes that you may want to force copy even if they already exist in the new object

**Value**

object The to argument object

**Examples**

```
a <- "test"
attr(a, 'wow') <- 1000
b <- a
b <- copyAllNewAttributes(a, b)
print(attr(b, 'wow'))
```

---

descGetMissing      *Get missingness statistics*

---

### Description

This function calculates the missingness row for [describeMean](#), [describeMedian](#) and custom description functions. It will return invisibly when no missing values are present.

### Usage

```
descGetMissing(x, html = TRUE, number_first = TRUE,
  percentage_sign = TRUE, language = "en", useNA.digits = 1, ...)
```

### Arguments

x	The variable that you want the statistics for
html	If HTML compatible output should be used. If FALSE it outputs LaTeX formatting
number_first	If the number should be given or if the percentage should be presented first. The second is encapsulated in parentheses (). This is only used together with the useNA variable.
percentage_sign	If you want to suppress the percentage sign you can set this variable to FALSE. You can also choose something else that the default % if you so wish by setting this variable. Note, this is only used when combined with the missing information.
language	The ISO-639-1 two-letter code for the language of interest. Currently only english is distinguished from the ISO format using a ',' as the separator in the <a href="#">txtInt</a> function.
useNA.digits	The number of digits to use for the missing percentage, defaults to the overall digits.
...	Passed on to <a href="#">describeFactors</a>

### Value

vector A vector with the missing estimate

---

describeFactors      *Describes factor variables*

---

### Description

A function that returns a description of proportions in a factor that contains the number of times a level occurs and the percentage

### Usage

```
describeFactors(x, html = TRUE, digits = 1, digits.nonzero = NA,
  number_first = TRUE, useNA = c("ifany", "no", "always"),
  useNA.digits = digits, horizontal_proportions,
  percentage_sign = TRUE, language = "en", ...)
```

### Arguments

x	The variable that you want the statistics for
html	If HTML compatible output should be used. If FALSE it outputs LaTeX formatting
digits	The number of decimals used
digits.nonzero	The number of decimals used for values that are close to zero
number_first	If the number should be given or if the percentage should be presented first. The second is encapsulated in parentheses (). This is only used together with the useNA variable.
useNA	This indicates if missing should be added as a separate row below all other. See <a href="#">table</a> for useNA-options. <i>Note:</i> defaults to ifany and not "no" as <a href="#">table</a> does.
useNA.digits	The number of digits to use for the missing percentage, defaults to the overall digits.
horizontal_proportions	Is only active if useNA since this is the only case of a proportion among continuous variables. This is default NULL and indicates that the proportions are to be interpreted in a vertical manner. If we want the data to be horizontal, i.e. the total should be shown and then how these differ in the different groups then supply the function with the total number in each group, i.e. if done in a by manner as in <a href="#">getDescriptionStatsBy</a> it needs to provide the number before the by() command.
percentage_sign	If you want to suppress the percentage sign you can set this variable to FALSE. You can also choose something else that the default % if you so wish by setting this variable. Note, this is only used when combined with the missing information.
language	The ISO-639-1 two-letter code for the language of interest. Currently only english is distinguished from the ISO format using a ',' as the separator in the <a href="#">txtInt</a> function.
...	Passed on to <a href="#">txtInt</a>

**Value**

A string formatted for printing either latex by HTML

**See Also**

[getDescriptionStatsBy](#)

Other descriptive functions: [describeMean](#), [describeMedian](#), [describeProp](#), [getDescriptionStatsBy](#), [getPvalWilcox](#)

**Examples**

```
set.seed(1)
describeFactors(sample(50, x=c("A","B", "C"), replace=TRUE))

n <- 500
my_var <- factor(sample(size=n, x=c("A","B", "C", NA), replace=TRUE))
my_exp <- rbinom(n=n, size=1, prob=0.2)
total <- table(my_var, useNA="ifany")
by(my_var,
    INDICES=my_exp,
    FUN=describeFactors,
    useNA="ifany",
    horizontal_proportions = total)
```

---

describeMean

*Describe the mean*

---

**Description**

A function that returns a description of a continuous variable using the mean together with the standard deviation. The standard deviation is used as it is "industry standard" to use mean with standard deviation and not because it's the only option.

**Usage**

```
describeMean(x, html = TRUE, digits = 1, digits.nonzero = NA,
  number_first = TRUE, useNA = c("ifany", "no", "always"),
  useNA.digits = digits, percentage_sign = TRUE, plusmin_str,
  language = "en", ...)
```

**Arguments**

<code>x</code>	The variable that you want the statistics for
<code>html</code>	If HTML compatible output should be used. If FALSE it outputs LaTeX formatting
<code>digits</code>	The number of decimals used



digits.nonzero	The number of decimals used for values that are close to zero
number_first	If the number should be given or if the percentage should be presented first. The second is encapsulated in parentheses (). This is only used together with the useNA variable.
useNA	This indicates if missing should be added as a separate row below all other. See <a href="#">table</a> for useNA-options. <i>Note:</i> defaults to ifany and not "no" as <a href="#">table</a> does.
useNA.digits	The number of digits to use for the missing percentage, defaults to the overall digits.
percentage_sign	If you want to suppress the percentage sign you can set this variable to FALSE. You can also choose something else that the default % if you so wish by setting this variable. Note, this is only used when combined with the missing information.
plusmin_str	Provide if you want anything other than the plus minus sign suited for the given output format.
language	The ISO-639-1 two-letter code for the language of interest. Currently only english is distinguished from the ISO format using a ',' as the separator in the <a href="#">txtInt</a> function.
...	Passed on to <a href="#">describeFactors</a>

**Value**

string Returns a string formatted for either LaTeX or HTML

**See Also**

[getDescriptionStatsBy](#)

Other descriptive functions: [describeFactors](#), [describeMedian](#), [describeProp](#), [getDescriptionStatsBy](#), [getPvalWilcox](#)

**Examples**

```
describeMean(1:10)
describeMean(c(1:10, NA), useNA="always")
describeMean(c(1:10, NA), useNA="no")
```

---

describeMedian	<i>A function that returns a description median that contains the interquartile range or the full range</i>
----------------	---

---

**Description**

A function that returns a description median that contains the interquartile range or the full range

**Usage**

```
describeMedian(x, iqr = TRUE, html = TRUE, digits = 1,
  digits.nonzero = NA, number_first = TRUE, useNA = c("ifany", "no",
  "always"), useNA.digits = digits, percentage_sign = TRUE,
  language = "en", ...)
```

**Arguments**

x	The variable that you want the statistics for
iqr	If interquartile range should be used
html	If HTML compatible output should be used. If FALSE it outputs LaTeX formatting
digits	The number of decimals used
digits.nonzero	The number of decimals used for values that are close to zero
number_first	If the number should be given or if the percentage should be presented first. The second is encapsulated in parentheses (). This is only used together with the useNA variable.
useNA	This indicates if missing should be added as a separate row below all other. See <a href="#">table</a> for useNA-options. <i>Note:</i> defaults to ifany and not "no" as <a href="#">table</a> does.
useNA.digits	The number of digits to use for the missing percentage, defaults to the overall digits.
percentage_sign	If you want to suppress the percentage sign you can set this variable to FALSE. You can also choose something else that the default % if you so wish by setting this variable. Note, this is only used when combined with the missing information.
language	The ISO-639-1 two-letter code for the language of interest. Currently only english is distinguished from the ISO format using a ',' as the separator in the <a href="#">txtInt</a> function.
...	Passed on to <a href="#">describeFactors</a>

**Value**

string A string formatted for either LaTeX or HTML

**See Also**

[getDescriptionStatsBy](#)

Other descriptive functions: [describeFactors](#), [describeMean](#), [describeProp](#), [getDescriptionStatsBy](#), [getPvalWilcox](#)

**Examples**

```
describeMedian(1:10)
describeMedian(c(1:10, NA), useNA="ifany")
```

---

describeProp	<i>A function that returns a description proportion that contains the number and the percentage</i>
--------------	---

---

### Description

A function that returns a description proportion that contains the number and the percentage

### Usage

```
describeProp(x, html = TRUE, digits = 1, digits.nonzero = NA,
  number_first = TRUE, useNA = c("ifany", "no", "always"),
  useNA.digits = digits, default_ref, percentage_sign = TRUE,
  language = "en", ...)
```

### Arguments

x	The variable that you want the statistics for
html	If HTML compatible output should be used. If FALSE it outputs LaTeX formatting
digits	The number of decimals used
digits.nonzero	The number of decimals used for values that are close to zero
number_first	If the number should be given or if the percentage should be presented first. The second is encapsulated in parentheses (). This is only used together with the useNA variable.
useNA	This indicates if missing should be added as a separate row below all other. See <a href="#">table</a> for useNA-options. <i>Note</i> : defaults to ifany and not "no" as <a href="#">table</a> does.
useNA.digits	The number of digits to use for the missing percentage, defaults to the overall digits.
default_ref	The default reference, either first, the level name or a number within the levels. If left out it defaults to the first value.
percentage_sign	If you want to suppress the percentage sign you can set this variable to FALSE. You can also choose something else that the default % if you so wish by setting this variable. Note, this is only used when combined with the missing information.
language	The ISO-639-1 two-letter code for the language of interest. Currently only english is distinguished from the ISO format using a ',' as the separator in the <a href="#">txtInt</a> function.
...	Passed on to <a href="#">describeFactors</a>

### Value

string A string formatted for either LaTeX or HTML

**See Also**

Other descriptive functions: [describeFactors](#), [describeMean](#), [describeMedian](#), [getDescriptionStatsBy](#), [getPvalWilcox](#)

**Examples**

```
describeProp(factor(sample(50, x=c("A","B", NA), replace=TRUE)))
```

---

distance	<i>Get the distance between boxes</i>
----------	---------------------------------------

---

**Description**

Retrieves the distance between two boxes as absolute "mm" units.

**Usage**

```
distance(box1, box2, type = c("vertical", "horizontal"), half = FALSE)
```

**Arguments**

box1	The first boxGrob
box2	The second boxGrob
type	Wheter we should retrieve the vertical or horizontal difference
half	If set to true it returns half the distance. This is convenient when postioning boxes between eachother.

**Value**

a unit with "mm"

**Examples**

```
box1 <- boxGrob("A test box", y=.8)
box2 <- boxGrob("Another test box", y=.2)
distance(box1, box2, "v")
```

---

docx_document	<i>Formatter wrapper for <a href="#">html_document</a>, facilitates easier porting to docx</i>
---------------	--

---

## Description

This function adds the option of having adaptations needed for seamless integration with MS Word for importing html-documents in the .docx-format. The advantage of html documents is the ability to create advanced formatting frequently needed in medical publications and that is available in the [htmlTable](#) function. You can view [the series](#) for more details regarding how to achieve fast-track-publishing (ftp) together with knitr.

## Usage

```
docx_document(..., self_contained = FALSE, mathjax = NULL,
  theme = NULL, highlight = NULL, css = "rmarkdown/docx.css",
  h1_style = "margin: 24pt 0pt 0pt 0pt;",
  other_h_style = "margin: 10pt 0pt 0pt 0pt;", remove_scripts = TRUE,
  force_captions = FALSE, css_max_width)
```

## Arguments

...	Passed onto <a href="#">html_document</a> .
self_contained	Overrides the default TRUE for <a href="#">html_document</a> to FALSE as LibreOffice hangs on long lines such as the base64 images included in the self-contained version.
mathjax	The advanced mathjax does not work with with Word/LibreOffice.
theme	No theme should be used for the output as the custom CSS should take care of everything.
highlight	By default turn off highlighting as scripts are difficult to import. This does though work somewhat OK when copy-pasting from the web-browser.
css	The CSS if other than the default within the package
h1_style	You can choose any css style formatting here that you want to be applied to all h1 elements. Note: this is only applied if LibreOffice_adapt is TRUE.
other_h_style	This is the formatting applied to any other h elements not included to the first. Note: this is only applied if LibreOffice_adapt is TRUE.
remove_scripts	TRUE if <code>&lt;script&gt;&lt;/script&gt;</code> tags are to be removed. These are usually not compatible with Word-processors and should therefore in most cases be stripped from the document.
force_captions	Since <code>out.width</code> and <code>out.height</code> remove the option of having captions this allows a workaround through some processing via the XML-package
css_max_width	The max width of the body element. Defaults to "40em" if not specified. Any CSS-compliant width format works.

## Details

If you want to get equations into Word the currently best way is to use the [word\\_document](#) format.

## Value

R Markdown output format to pass to [render](#)

## Author(s)

Max Gordon

## Examples

```
# Possible yaml configuration at the top of the Rmd doc
## Not run:
---
title: "Test"
author: "Max Gordon"
output:
  Gmisc: :docx_document
---

## End(Not run)
```

---

fastDoCall

*An alternative to the internal do.call*

---

## Description

The `do.call` can be somewhat slow, especially when working with large objects. This function is based upon the suggestions from Hadley Wickham on the R mailing list, see [here](#). Also thanks to *Tommy* at StackOverflow for [suggesting](#) how to handle double and triple colon operators, `:::`, further enhancing the function.

## Usage

```
fastDoCall(what, args, quote = FALSE, envir = parent.frame())
```

## Arguments

<code>what</code>	either a function or a non-empty character string naming the function to be called.
<code>args</code>	a <i>list</i> of arguments to the function call. The names attribute of <code>args</code> gives the argument names.
<code>quote</code>	a logical value indicating whether to quote the arguments.
<code>envir</code>	an environment within which to evaluate the call. This will be most useful if <code>what</code> is a character string and the arguments are symbols or quoted expressions.

**Note**

While the function attempts to do most of what `do.call` can it has limitations. It can currently not parse the example code from the original function: `do.call(paste, list(as.name("A"), as.name("B")), quote = TRUE)` and the functionality of `quote` has not been thoroughly tested.

**Examples**

```
fastDoCall("complex", list(imaginary = 1:3))

## if we already have a list (e.g. a data frame)
## we need c() to add further arguments
tmp <- expand.grid(letters[1:2], 1:3, c("+", "-"))
fastDoCall("paste", c(tmp, sep = ""))

## examples of where objects will be found.
A <- 2
f <- function(x) print(x^2)
env <- new.env()
assign("A", 10, envir = env)
assign("f", f, envir = env)
f <- function(x) print(x)
f(A) # 2
fastDoCall("f", list(A)) # 2
fastDoCall("f", list(A), envir = env) # 4
fastDoCall(f, list(A), envir = env) # 2
fastDoCall("f", list(quote(A)), envir = env) # 100
fastDoCall(f, list(quote(A)), envir = env) # 10
fastDoCall("f", list(as.name("A")), envir = env) # 100

eval(call("f", A)) # 2
eval(call("f", quote(A))) # 2
eval(call("f", A), envir = env) # 4
eval(call("f", quote(A)), envir = env) # 100
```

---

 figCapNo

*Adds a figure caption number*


---

**Description**

The function relies on `options("fig_caption_no")` in order to keep track of the last number. If you want to force the caption function to skip captions while still using it in the knitr `fig.cap` option then simply set `options(fig_caption_no = FALSE)`

**Usage**

```
figCapNo(str, roman = getOption("fig_caption_no_roman", FALSE),
  sprintf_str = getOption("fig_caption_no_sprintf", "Fig. %s: %s"))
```

**Arguments**

str	The string that is to be prepended with string
roman	Whether or not to use roman numbers instead of arabic. Can also be set through <code>options(fig_caption_no_roman = TRUE)</code>
sprintf_str	An <code>sprintf</code> formatted string where the first argument is reserved for the string generated by the counter and the second one is for the caption text. Can also be set through <code>options(fig_caption_no_sprintf = TRUE)</code>

**See Also**

Other figure caption functions: [figCapNoLast](#), [figCapNoNext](#)

**Examples**

```
## Not run:
```{r, fig.cap=figCapNo("My nice plot")}
plot(1:10 + rnorm(10), 1:10)
```

## End(Not run)
org_opts <- options(fig_caption_no = 2,
                    fig_caption_no_sprintf = "Figure %s: %s")
figCapNo("A plot with caption number = 3")

org_opts <- options(fig_caption_no = TRUE)
figCapNo("A plot with caption number = 1")

# Use default setting
options(fig_caption_no_sprintf = NULL)
figCapNo("A plot with caption number = 2")

# Return the original settings
options(org_opts)
```

---

figCapNoLast

*Gets the last figure caption number*

---

**Description**

The function relies on `options("fig_caption_no")` in order to keep track of the last number.

**Usage**

```
figCapNoLast(roman = getOption("fig_caption_no_roman", FALSE))
```

**Arguments**

|       |  |
|-------|--|
| roman | Whether or not to use roman numbers instead of arabic. Can also be set through <code>options(fig_caption_no_roman = TRUE)</code> |
|-------|--|



**See Also**

Other figure caption functions: [figCapNoNext](#), [figCapNo](#)

**Examples**

```
org_opts <- options(fig_caption_no=1)
figCapNoLast()
options(org_opts)
```

---

|              |  |
|--------------|--|
| figCapNoNext | <i>Gets the next figure caption number</i> |
|--------------|--|

---

**Description**

The function relies on `options("fig_caption_no")` in order to keep track of the last number.

**Usage**

```
figCapNoNext(roman = getOption("fig_caption_no_roman", FALSE))
```

**Arguments**

|       |  |
|-------|--|
| roman | Whether or not to use roman numbers instead of arabic. Can also be set through <code>options(fig_caption_no_roman = TRUE)</code> |
|-------|--|

**See Also**

Other figure caption functions: [figCapNoLast](#), [figCapNo](#)

**Examples**

```
org_opts <- options(fig_caption_no=1)
figCapNoNext()
options(org_opts)
```

---

getBezierAdj4Arrw      *Gets the bezier points adjusted for an arrow*

---

### Description

Gets the bezier points adjusted for an arrow

### Usage

```
getBezierAdj4Arrw(x, y, arrow_length, length_out = 100)
```

### Arguments

|              |   |
|--------------|---|
| x            | The x start and end points  |
| y            | The spline control points   |
| arrow_length | The desired length of the arrow   |
| length_out   | Increases the resolution for the final bezier points, i.e. generating more fine-grained intervals |

### Value

list

---

getDescriptionStatsBy      *Creating of description statistics*

---

### Description

A function that returns a description statistic that can be used for creating a publication "table 1" when you want it by groups. The function identifies if the variable is a continuous, binary or a factored variable. The format is inspired by NEJM, Lancet & BMJ.

### Usage

```
getDescriptionStatsBy(x, by, digits = 1, digits.nonzero = NA,
  html = TRUE, numbers_first = TRUE, statistics = FALSE,
  statistics.sig_lim = 10^-4, statistics.two_dec_lim = 10^-2,
  statistics.suppress_warnings = TRUE, useNA = c("ifany", "no",
  "always"), useNA.digits = digits, continuous_fn = describeMean,
  prop_fn = describeProp, factor_fn = describeFactors,
  show_all_values = FALSE, hrzl_prop = FALSE, add_total_col,
  total_col_show_perc = TRUE, use_units = FALSE, default_ref,
  NEJMstyle = FALSE, percentage_sign = TRUE, header_count,
  missing_value = "-", names_of_missing = NULL, ...)
```

**Arguments**

|   |  |
|---|--|
| <code>x</code>                            | The variable that you want the statistics for  |
| <code>by</code>                           | The variable that you want to split into different columns   |
| <code>digits</code>                       | The number of decimals used  |
| <code>digits.nonzero</code>               | The number of decimals used for values that are close to zero  |
| <code>html</code>                         | If HTML compatible output should be used. If FALSE it outputs LaTeX formatting   |
| <code>numbers_first</code>                | If the number should be given or if the percentage should be presented first. The second is encapsulated in parentheses ().  |
| <code>statistics</code>                   | Add statistics, fisher test for proportions and Wilcoxon for continuous variables. See details below for more customization.   |
| <code>statistics.sig_lim</code>           | The significance limit for < sign, i.e. p-value 0.0000312 should be < 0.0001 with the default setting.   |
| <code>statistics.two_dec_lim</code>       | The limit for showing two decimals. E.g. the p-value may be 0.056 and we may want to keep the two decimals in order to emphasize the proximity to the all-mighty 0.05 p-value and set this to $10^{-2}$ . This allows that a value of 0.0056 is rounded to 0.006 and this makes intuitive sense as the 0.0056 level as this is well below the 0.05 value and thus not as interesting to know the exact proximity to 0.05. <i>Disclaimer:</i> The 0.05-limit is really silly and debated, unfortunately it remains a standard and this package tries to adapt to the current standards in order to limit publication associated issues. |
| <code>statistics.suppress_warnings</code> | Hide warnings from the statistics function.  |
| <code>useNA</code>                        | This indicates if missing should be added as a separate row below all other. See <a href="#">table</a> for useNA-options. <i>Note:</i> defaults to ifany and not "no" as <a href="#">table</a> does.   |
| <code>useNA.digits</code>                 | The number of digits to use for the missing percentage, defaults to the overall digits.  |
| <code>continuous_fn</code>                | The method to describe continuous variables. The default is <a href="#">describeMean</a> .   |
| <code>prop_fn</code>                      | The method used to describe proportions, see <a href="#">describeProp</a> .  |
| <code>factor_fn</code>                    | The method used to describe factors, see <a href="#">describeFactors</a> .   |
| <code>show_all_values</code>              | This is by default false as for instance if there is no missing and there is only one variable then it is most sane to only show one option as the other one will just be a complement to the first. For instance sex - if you know gender then automatically you know the distribution of the other sex as it's 100 % - other %. To choose which one you want to show then set the <code>default_ref</code> parameter.  |
| <code>hrzl_prop</code>                    | This is default FALSE and indicates that the proportions are to be interpreted in a vertical manner. If we want the data to be horizontal, i.e. the total should be shown and then how these differ in the different groups then set this to TRUE.   |
| <code>add_total_col</code>                | This adds a total column to the resulting table. You can also specify if you want the total column "first" or "last" in the column order.  |

|                     |  |
|---------------------|--|
| total_col_show_perc | This is by default true but if requested the percentages are suppressed as this sometimes may be confusing.  |
| use_units           | If the Hmisc package's units() function has been employed it may be interesting to have a column at the far right that indicates the unit measurement. If this column is specified then the total column will appear before the units (if specified as last). You can also set the value to "name" and the units will be added to the name as a parenthesis, e.g. Age (years). |
| default_ref         | The default reference, either first, the level name or a number within the levels. If left out it defaults to the first value.   |
| NEJMstyle           | Adds - no (%) at the end to proportions  |
| percentage_sign     | If you want to suppress the percentage sign you can set this variable to FALSE. You can also choose something else that the default % if you so wish by setting this variable.   |
| header_count        | Set to TRUE if you want to add a header count, e.g. Smoking; No. 25 observations, where there is a new line after the factor name. If you want a different text for the second line you can specifically use the <code>sprintf</code> formatting, e.g. "No. %s patients".  |
| missing_value       | Value that is substituted for empty cells. Defaults to "-"   |
| names_of_missing    | Optional character vector containing the names of returned statistics, in case all returned values for a given by level are missing. Defaults to NULL  |
| ...                 | Currently only used for generating warnings of deprecated call parameters.   |

### Value

Returns a vector if vars wasn't specified and it's a continuous or binary statistic. If vars was a matrix then it appends the result to the end of that matrix. If the x variable is a factor then it does not append and you get a warning.

### Customizing statistics

You can specify what function that you want for statistic by providing a function that takes two arguments x and by and returns a p-value. There are a few functions already prepared for this see [getPvalAnova](#), [getPvalChiSq](#), [getPvalFisher](#), [getPvalKruskal](#), [getPvalWilcox](#). The default functions used are [getPvalFisher](#) and [getPvalWilcox](#) (unless the by argument has more than three unique levels where it defaults to [getPvalAnova](#)).

If you want the function to select functions depending on the type of input you can provide a list with the names 'continuous', 'proportion', 'factor' and the function will choose accordingly. If you fail to define a certain category it will default to the above.

You can also use a custom function that returns a string with the attribute 'colname' set that will be appended to the results instead of the p-value column. to the results instead of the p-value column.

### See Also

Other descriptive functions: [describeFactors](#), [describeMean](#), [describeMedian](#), [describeProp](#), [getPvalWilcox](#)

**Examples**

```

data(mtcars)
# For labelling we use the label()
# function from the Hmisc package
library(Hmisc)

label(mtcars$mpg) <- "Gas"
units(mtcars$mpg) <- "Miles/(US) gallon"

label(mtcars$wt) <- "Weight"
units(mtcars$wt) <- "10<sup>3</sup> kg" # not sure the unit is correct

mtcars$am <- factor(mtcars$am, levels=0:1, labels=c("Automatic", "Manual"))
label(mtcars$am) <- "Transmission"

mtcars$gear <- factor(mtcars$gear)
label(mtcars$gear) <- "Gears"

# Make up some data for making it slightly more interesting
mtcars$col <- factor(sample(c("red", "black", "silver"),
                          size=NROW(mtcars), replace=TRUE))
label(mtcars$col) <- "Car color"

mergeDesc(getDescriptionStatsBy(mtcars$mpg, mtcars$am,
                               header_count = TRUE,
                               use_units = TRUE),
          getDescriptionStatsBy(mtcars$wt, mtcars$am,
                               header_count = TRUE,
                               use_units = TRUE),
          htmlTable_args = list(caption = "Basic continuous stats from the mtcars dataset"))

t11 <- list()
t11[["Gear (3 to 5)"]] <- getDescriptionStatsBy(mtcars$gear, mtcars$am)
t11 <- c(t11,
        list(getDescriptionStatsBy(mtcars$col, mtcars$am)))

mergeDesc(t11,
          htmlTable_args = list(caption = "Factored variables"))

t1_no_units <- list()
t1_no_units[["Gas (mile/gallons)"]] <-
  getDescriptionStatsBy(mtcars$mpg, mtcars$am,
                        header_count = TRUE)
t1_no_units[["Weight (10<sup>3</sup> kg)"]] <-
  getDescriptionStatsBy(mtcars$wt, mtcars$am,
                        header_count = TRUE)
mergeDesc(t1_no_units, t11,
          # Remove the formatting for the groups
          htmlTable_args = list(css.rgroup = ""))

# A little more advanced

```

```

mtcars$mpg[sample(1:NROW(mtcars), size=5)] <- NA
getDescriptionStatsBy(mtcars$mpg, mtcars$am, statistics=TRUE)

# Do the horizontal version
getDescriptionStatsBy(mtcars$col, mtcars$am,
                      statistics=TRUE, hrzl_prop = TRUE)

mtcars$wt_with_missing <- mtcars$wt
mtcars$wt_with_missing[sample(1:NROW(mtcars), size=8)] <- NA
getDescriptionStatsBy(mtcars$wt_with_missing, mtcars$am, statistics=TRUE,
                      hrzl_prop = TRUE, total_col_show_perc = FALSE)

mtcars$col_with_missing <- mtcars$col
mtcars$col_with_missing[sample(1:NROW(mtcars), size=5)] <- NA
getDescriptionStatsBy(mtcars$col_with_missing, mtcars$am, statistics=TRUE,
                      hrzl_prop = TRUE, total_col_show_perc = FALSE)

## Not run:
## There is also a LaTeX wrapper
t11 <- list(
  getDescriptionStatsBy(mtcars$gear, mtcars$am),
  getDescriptionStatsBy(mtcars$col, mtcars$am))

latex(mergeDesc(t11),
      caption = "Factored variables",
      file="")

## End(Not run)

```

---

getPvalWilcox

*P-value extractors for [getDescriptionStatsBy](#)*


---

## Description

These functions are the base functions for getting the description p-values. You can provide your own functions but all functions should take two arguments and return a p-value (numeric, non-formatted)

## Usage

```
getPvalWilcox(x, by)
```

```
getPvalAnova(x, by)
```

```
getPvalFisher(x, by)
```

```
getPvalChiSq(x, by)
```

```
getPvalKruskal(x, by)
```

### Arguments

|    |                                     |
|----|-------------------------------------|
| x  | The main variable of interest       |
| by | The variable for the stratification |

### Value

numeric Returns the p-value from that particular test

### getPvalWilcox

Performs a two-sample two-sided Wilcoxon test (also known as the Mann-Whitney test), see [wilcox.test](#).

### getPvalAnova

Performs a standard Analysis of Variance model through [anova\(lm\(x ~ by\)\)](#)

### getPvalFisher

Performs Fisher's exact test through the [fisher.test](#).

### getPvalChiSq

Performs a standard Chi-Squares analysis through [chisq.test](#)

### getPvalKruskal

Performs a Kruskal-Wallis rank sum test through [kruskal.test](#)

### See Also

Other descriptive functions: [describeFactors](#), [describeMean](#), [describeMedian](#), [describeProp](#), [getDescriptionStatsBy](#)

### Examples

```
set.seed(123)
getPvalFisher(sample(letters[1:3], size = 100, replace = TRUE),
              sample(LETTERS[1:3], size = 100, replace = TRUE))
getPvalWilcox(rnorm(100),
              sample(LETTERS[1:2], size = 100, replace = TRUE))
```

---

getSvdMostInfluential *Gets the maximum contributor variables from svd()*

---

### Description

This function is inspired by Jeff Leeks Data Analysis course where he suggests that one way to use the `svd` is to look at the most influential rows for first columns in the V matrix.

### Usage

```
getSvdMostInfluential(mtrx, quantile, similarity_threshold,  
  plot_selection = TRUE, plot_threshold = 0.05, varnames = NULL)
```

### Arguments

|                                   |   |
|-----------------------------------|---|
| <code>mtrx</code>                 | A matrix or data frame with the variables. Note: if it contains missing variables make sure to impute prior to this function as the <code>svd</code> can't handle missing values. |
| <code>quantile</code>             | The SVD D-matrix gives an estimate for the amount that is explained. This parameter is used for selecting the columns that have that quantile of explanation.                     |
| <code>similarity_threshold</code> | A quantile for how close other variables have to be in value to maximum contributor of that particular column. If you only want the maximum value then set this value to 1.       |
| <code>plot_selection</code>       | As this is all about variable exploring it is often interesting to see how the variables were distributed among the vectors   |
| <code>plot_threshold</code>       | The threshold of the plotted bars, measured as percent explained by the D-matrix. By default it is set to 0.05.   |
| <code>varnames</code>             | A vector with alternative names to the colnames   |

### Details

This function expands on that idea and adds the option of choosing more than just the most contributing variable for each row. For instance two variables may have a major impact on a certain component where the second variable has 95 important in that particular component it makes sense to include it in the selection.

It is of course useful when you have many continuous variables and you want to determine a subgroup to look at, i.e. finding the needle in the haystack.

### Value

Returns a list with vector with the column numbers that were picked in the "most\_influential" variable and the svd calculation in the "svd"



**Examples**

```

org_par <- par(ask=TRUE)
set.seed(12345);

# Simulate data with a pattern
dataMatrix <- matrix(rnorm(15*160),ncol=15)
colnames(dataMatrix) <-
  c(paste("Pos.3:", 1:3, sep=" #"),
    paste("Neg.Decr:", 4:6, sep=" #"),
    paste("No pattern:", 7:8, sep=" #"),
    paste("Pos.Incr:", 9:11, sep=" #"),
    paste("No pattern:", 12:15, sep=" #"))
for(i in 1:nrow(dataMatrix)){
  # flip a coin
  coinFlip1 <- rbinom(1,size=1,prob=0.5)
  coinFlip2 <- rbinom(1,size=1,prob=0.5)
  coinFlip3 <- rbinom(1,size=1,prob=0.5)

  # if coin is heads add a common pattern to that row
  if(coinFlip1){
    cols <- grep("Pos.3", colnames(dataMatrix))
    dataMatrix[i, cols] <- dataMatrix[i, cols] + 3
  }

  if(coinFlip2){
    cols <- grep("Neg.Decr", colnames(dataMatrix))
    dataMatrix[i, cols] <- dataMatrix[i, cols] - seq(from=5, to=15, length.out=length(cols))
  }

  if(coinFlip3){
    cols <- grep("Pos.Incr", colnames(dataMatrix))
    dataMatrix[i,cols] <- dataMatrix[i,cols] + seq(from=3, to=15, length.out=length(cols))
  }
}

# Illustrate data
heatmap(dataMatrix, Colv=NA, Rowv=NA, margins=c(7,2), labRow="")

svd_out <- svd(scale(dataMatrix))

library(lattice)
b_clr <- c("steelblue", "darkred")
key <- simpleKey(rectangles = TRUE, space = "top", points=FALSE,
  text=c("Positive", "Negative"))
key$rectangles$col <- b_clr

b1 <- barchart(as.table(svd_out$v[,1]),
  main="First column",
  horizontal=FALSE, col=ifelse(svd_out$v[,1] > 0,
    b_clr[1], b_clr[2]),
  ylab="Impact value",
  scales=list(x=list(rot=55, labels=colnames(dataMatrix), cex=1.1)),

```

```

key = key)

b2 <- barchart(as.table(svd_out$v[,2]),
  main="Second column",
  horizontal=FALSE, col=ifelse(svd_out$v[,2] > 0,
    b_clr[1], b_clr[2]),
  ylab="Impact value",
  scales=list(x=list(rot=55, labels=colnames(dataMatrix), cex=1.1)),
  key = key)

b3 <- barchart(as.table(svd_out$v[,3]),
  main="Third column",
  horizontal=FALSE, col=ifelse(svd_out$v[,3] > 0,
    b_clr[1], b_clr[2]),
  ylab="Impact value",
  scales=list(x=list(rot=55, labels=colnames(dataMatrix), cex=1.1)),
  key = key)

b4 <- barchart(as.table(svd_out$v[,4]),
  main="Fourth column",
  horizontal=FALSE, col=ifelse(svd_out$v[,4] > 0,
    b_clr[1], b_clr[2]),
  ylab="Impact value",
  scales=list(x=list(rot=55, labels=colnames(dataMatrix), cex=1.1)),
  key = key)

# Note that the fourth has the no pattern columns as the
# chosen pattern, probably partly because of the previous
# patterns already had been identified
print(b1, position=c(0,0.5,.5,1), more=TRUE)
print(b2, position=c(0.5,0.5,1,1), more=TRUE)
print(b3, position=c(0,0,.5,.5), more=TRUE)
print(b4, position=c(0.5,0,1,.5))

# Let's look at how well the SVD identifies
# the most influential columns
getSvdMostInfluential(dataMatrix,
  quantile=.8,
  similarity_threshold = .9,
  plot_threshold = .05,
  plot_selection = TRUE)

par(org_par)

```

**Description**

This is a general form of bezier line that can be used for cubic, quadratic, and more advanced Bézier lines.

**Usage**

```
gnrlBezierPoints(ctrl_points, length_out = 100L)
```

**Arguments**

`ctrl_points` The `ctrl_points` for the bezier control points. This should either be a matrix or a data.frame.

`length_out` The length of the return points, i.e. how fine detailed the points should be.

**Examples**

```
library(grid)
grid.newpage()
l <- gnrlBezierPoints(data.frame(x = c(.1, -.1, .7, 1, 1, 0.1),
                                y = c(.9, 0, 1, .8, .4, .1)),
                      length_out = 100)
grid.lines(l[,1], l[,2], gp=gpar(col="#550000", lwd = 4))

out_sizes <- 4:20
clrs <- colorRampPalette(c("orange", "darkblue"))(length(out_sizes))
for (i in out_sizes){
  l <- gnrlBezierPoints(data.frame(x = c(.1, -.1, .7, 1, 1, 0.1),
                                y = c(.9, 0, 1, .8, .4, .1)),
                        length_out = i)
  grid.lines(l[,1], l[,2],
            gp=gpar(col=clrs[which(i == out_sizes)]))
}
```

---

 has

---

*An R alternative to the lodash has in JavaScript*


---

**Description**

This is a handy function for checking if item exist in a nested structure

**Usage**

```
has(sourceList, path)
```

**Arguments**

`sourceList` The `list()/c()` that is to be searched for the element

`path` A string that can be separated by [,] or ., the string "elementname1.1.elementname" the validity of the path - it only separates and tries to address that element with '[[ ]'.

**Value**

Returns a boolean.

**See Also**

Other lodash similar functions: [retrieve](#)

**Examples**

```
has(list(a = list(b = 1)), "a.b")
```

---

`insertRowAndKeepAttr` *Insert a row into a matrix*

---

**Description**

Inserts a row and keeps the attributes [copyAllNewAttributes](#)

**Usage**

```
insertRowAndKeepAttr(m, r, v = NA, rName = "")
```

**Arguments**

|                    |   |
|--------------------|---|
| <code>m</code>     | matrix  |
| <code>r</code>     | row number where the new row should be inserted     |
| <code>v</code>     | optional values for the new row                     |
| <code>rName</code> | optional character string: the name of the new row. |

**Value**

`matrix` Returns a matrix with one more row than the provided matrix `m`

**Author(s)**

Max Gordon, Arne Henningsen

**Examples**

```
test <- matrix(1:4, ncol=2)
attr(test, 'wow') <- 1000
test <- insertRowAndKeepAttr(test, 2)
print(attr(test, 'wow'))
```

---

|           |  |
|-----------|--|
| mergeDesc | <i>Prepares a matrix for htmlTable from a list</i> |
|-----------|--|

---

### Description

By putting all the output from the [getDescriptionStatsBy](#) into a list, naming each element that we want in an rgroup we can automatically merge everything and create an object ready for the [htmlTable](#).

### Usage

```
mergeDesc(..., htmlTable_args = list(css.rgroup = ""))
```

### Arguments

... One or more elements coming from [getDescriptionStatsBy](#). You can also provide pure output from the [getDescriptionStatsBy](#) function and have the function merge this together with the ... argument. *Note* that all elements must have the same by argument or you will not be able to merge it into a list.

htmlTable\_args Any arguments that should be passed to [htmlTable](#) function. The default is to remove any css formatting for the rgroup.

### Value

matrix Returns a matrix object of class descList

### The rgroup value

The value for the rgroup is by default the name of the list element. If you have passed a list without a name for that particular element or if you have passed a matrix it will look for a label set by the `Hmisc::label` function. For those elements that have only one row no rgroup is set, and the naming sequence is the same as above but with an additional [rownames](#) if the previous two turn out empty. All this behaviour is exemplified in the example.

The rgroup value can be overridden by simply specifying a custom rgroup when calling the [htmlTable](#) function.

### The colnames of the matrix

The function chooses the [colnames](#) from the first element in the tlist.

### Examples

```
data(mtcars)
# For labelling we use the label()
# function from the Hmisc package
library(Hmisc)

label(mtcars$mpg) <- "Gas"
```

```

units(mtcars$mpg) <- "Miles/(US) gallon"

label(mtcars$wt) <- "Weight"
units(mtcars$wt) <- "10<sup>3</sup> kg" # not sure the unit is correct

mtcars$am <- factor(mtcars$am, levels=0:1, labels=c("Automatic", "Manual"))
label(mtcars$am) <- "Transmission"

mtcars$gear <- factor(mtcars$gear)
label(mtcars$gear) <- "Gears"

# Make up some data for making it slightly more interesting
mtcars$col <- factor(sample(c("red", "black", "silver"),
                          size=NROW(mtcars), replace=TRUE))
label(mtcars$col) <- "Car color"

mergeDesc(getDescriptionStatsBy(mtcars$mpg, mtcars$am,
                               header_count = TRUE,
                               use_units = TRUE),
          getDescriptionStatsBy(mtcars$wt, mtcars$am,
                               header_count = TRUE,
                               use_units = TRUE),
          htmlTable_args = list(caption = "Basic continuous stats from the mtcars dataset"))

tll <- list()
tll[["Gear (3 to 5)"]] <- getDescriptionStatsBy(mtcars$gear, mtcars$am)
tll <- c(tll,
        list(getDescriptionStatsBy(mtcars$col, mtcars$am)))

mergeDesc(tll,
          htmlTable_args = list(caption = "Factored variables"))

tl_no_units <- list()
tl_no_units[["Gas (mile/gallons)"]] <-
  getDescriptionStatsBy(mtcars$mpg, mtcars$am,
                        header_count = TRUE)
tl_no_units[["Weight (10<sup>3</sup> kg)"]] <-
  getDescriptionStatsBy(mtcars$wt, mtcars$am,
                        header_count = TRUE)
mergeDesc(tl_no_units, tll,
          # Remove the formatting for the groups
          htmlTable_args = list(css.rgroup = ""))

# A little more advanced
mtcars$mpg[sample(1:NROW(mtcars), size=5)] <- NA
getDescriptionStatsBy(mtcars$mpg, mtcars$am, statistics=TRUE)

# Do the horizontal version
getDescriptionStatsBy(mtcars$col, mtcars$am,
                      statistics=TRUE, hrzl_prop = TRUE)

mtcars$wt_with_missing <- mtcars$wt

```

```

mtcars$wt_with_missing[sample(1:NROW(mtcars), size=8)] <- NA
getDescriptionStatsBy(mtcars$wt_with_missing, mtcars$am, statistics=TRUE,
                      hrz1_prop = TRUE, total_col_show_perc = FALSE)

mtcars$col_with_missing <- mtcars$col
mtcars$col_with_missing[sample(1:NROW(mtcars), size=5)] <- NA
getDescriptionStatsBy(mtcars$col_with_missing, mtcars$am, statistics=TRUE,
                      hrz1_prop = TRUE, total_col_show_perc = FALSE)

## Not run:
## There is also a LaTeX wrapper
tll <- list(
  getDescriptionStatsBy(mtcars$gear, mtcars$am),
  getDescriptionStatsBy(mtcars$col, mtcars$am))

latex(mergeDesc(tll),
      caption = "Factored variables",
      file="")

## End(Not run)

```

---

mergeLists

*Merging of multiple lists*


---

## Description

The merge allows for a recursive component where the lists are compared on the subelement. If one does not contain that element it will get NA in for those parameters.

## Usage

```
mergeLists(..., lapplyOutput = NULL)
```

## Arguments

|              |   |
|--------------|---|
| ...          | Any number of lists that you want to merge  |
| lapplyOutput | The <code>lapply</code> function outputs a number of lists and this is for specifically merging all of those. |

## Value

Returns a list with all the given lists.

## Examples

```

v1 <- list("a"=c(1,2), b="test 1", sublist=list(one=20:21, two=21:22))
v2 <- list("a"=c(3,4), b="test 2", sublist=list(one=10:11, two=11:12, three=1:2))
mergeLists(v1, v2)

```

---

|            |   |
|------------|---|
| prPasteVec | <i>Collapses a vector for throwing errors</i> |
|------------|---|

---

**Description**

The function collapses a vector into an output useful when throwing errors, e.g. 1:3 becomes '1', '2', '3'

**Usage**

```
prPasteVec(x)
```

**Arguments**

|   |            |
|---|------------|
| x | The vector |
|---|------------|

---

|          |   |
|----------|---|
| retrieve | <i>An R alternative to the lodash get in JavaScript</i> |
|----------|---|

---

**Description**

This is a handy function for retrieving items deep in a nested structure without causing error if not found

**Usage**

```
retrieve(sourceList, path, default = NA)
```

**Arguments**

|            |   |
|------------|---|
| sourceList | The list()/c() that is to be searched for the element   |
| path       | A string that can be separated by [,] or ., the string "elementname1.1.elementname" is equivalent to "elementname1[[1]]elementname". Note that the function doesn't check the validity of the path - it only separates and tries to address that element with '[[ ]'. |
| default    | The value to return if the element isn't found  |

**Value**

Returns a sub-element from sourceList or the default value.

**See Also**

Other lodash similar functions: [has](#)



**Examples**

```
source <- list(a = list(b = 1, `odd.name` = 'I hate . in names', c(1,2,3)))
retrieve(source, "a.b")
retrieve(source, "a.b.1")
retrieve(source, "a.odd\\.name")
retrieve(source, "a.not_in_list")
```

---

Transition-class      *A reference class for generating transition plots*

---

**Description**

This class simplifies the creating of transition plots. It also allows for advanced multi-column transitions.

**Details**

Transition plots are a type of *Sankey diagrams*. These are a specific type of flow diagram, in which the width of the arrows is shown proportionally to the flow quantity. See [Wikipedia](#) for details.

**Fields**

**id** Optional id. The render uses named viewports that require a unique id if multiple transition plots are combined. In order to avoid having overlapping graphs we need to generate a unique id for each viewport and thus this variable exists. If left empty it will create a counter that is stored in the `options` ("Gmisc.transitionClassCounter") and each viewport will have the name preceded with 'tc\_[0-9]+'. Set this if you intend to use `seekViewport`.

**transitions** This is a  $\geq 3$  dimensional array with the transitions. Should not be directly accessed.

**box\_width** The box width

**box\_txt** The texts of each box

**box\_label** Box labels

**box\_label\_pos** The label's positions, either "top"/"bottom"

**box\_label\_cex** The size of the box labels

**box\_cex** The fontsize multiplier for the text within the boxes

**arrow\_type** The type of arrow to use, defaults to "gradient", but can also be "simple". The corresponding functions are `bezierArrowGradient`, and `bezierArrowSmpl`. *Note* The `bezierGrob` ("grid") has been deprecated as it is no longer faster than the `bezierArrows` and there is a difference in design.

**arrow\_clr** The arrow color

**arrow\_rez** The resolution of the arrow

**vertical\_space** The space between the boxes

**fill\_clr** The box fill color

`clr_bar` Shows a color bar if there are proportions. Can be "none", "top", "bottom"  
`clr_bar_clrs` Extracts the colors for the colorbar from the `fill_clr` if none is provided  
`clr_bar_cex` The size of the ticks in the color bar  
`clr_bar_subspace` If little or no difference exists at the low/high proportions of the spectrum then it can be of interest to focus the color change to the center leaving the tails constant  
`clr_bar_labels` The labels of the color bars. Defaults to the dimnames for the proportions.  
`txt_clr` The text color within the boxes  
`title` The plot title if any  
`title_cex` The fontsize multiplier for the title  
`skip_shadows` Skip the shadow effect on the boxes  
`mar` The margins for the plot.  
`min_lwd` The minimum line width that is still shown. The pixels will most likely not have the same fine resolution as the data and you therefore may want to hide lines that are smaller than a certain amount.  
`max_lwd` The maximum line width to show  
`lwd_prop_type` The line can either be proportional to the "set" of transitions (group of two box columns), to "all" transitions, or to each "box". It defaults to "all".  
`data` Internal storage variable. Should not be accessed directly.

## Methods

`addClr(fill, txt)` Adds colors or extends existing one so that they match the transition matrix. The fill corresponds to the `fill_clr` and `txt` corresponds to the `txt_clr`. If the colors are missing and the transitions consist of only two columns the default colors will be used. If the matrix is being extended and these values are missing the values from the previous last column will be used for the default columns.

`addTransitions(mtrx, label, txt, fill_clr, txt_clr)` Add a transition matrix. The input has to be a numerical matrix between 2 and 3 dimensions. If you don't provide the `txt` field the box' text field will be deduced from the transition matrix' dimnames. The `fill_clr` and `txt_clr` are passed on to the `addClr` function.

`arrowWidths(set_no, add_width)` Retrieves the details regarding arrow sizes for each arrow within the transition group

`boxPositions(col)` The box positions as a list with scalars for the positions:

1. *x* The center x-position
2. *y* The center y-position
3. *right* The right edge
4. *left* The left edge
5. *top* The top edge
6. *bottom* The bottom edge
7. *height* The box height
8. *width* The box width
9. *unit* The unit used for the values (npc)

`boxSizes(col)` Gets the size of the boxes. The `col` argument should be either an integer or 'last'  
`getDim()` Gets the current dimensions of the transitions  
`getTransitionSet(no, reduce_dim = FALSE)` Gets a specific set of transitions. If the `reduce_dim` is set to `TRUE` it will only return a 2-dimensional matrix even if the original has a 3rd proportions dimension  
`getYProps(col)` Gets the proportions after removing the `vertical_space` between the boxes  
`initialize(transitions, label, txt, fill_clr, txt_clr, id, ...)` Set up a Transition object. The transitions should be a 2D or 3D matrix as defined in the `$addTransitions` section and not as later internally stored.  
`noCols()` Gets the number of columns, i.e. the number of transitions  
`noRows(no)` Gets the number of boxes in each row. If multiple rows the number of rows may differ between each transition matrix we therefore need to specify what transitions that we refer to. If no value is specified it returns all of them.  
`render(new_page = TRUE)` Call this to render the full graph. The `new_page` argument is for creating a new plot, set this to `FALSE` if you want to combine this plot with another or if you have additional viewports that you intend to use.  
`trnstnSizes(set_no)` Gets the transitions per box as a 2D matrix. For the proportions it also adds an attribute `attr('props', prop_mtrx)` that is a 2D matrix with the corresponding proportions.

## Examples

```

# Transitions
set.seed(1)
n <- 10
my_data <-
  data.frame(
    Var_a = sample(c("Test 1",
                    "Test 2",
                    "Test 3"),
                  size = n,
                  replace = TRUE,
                  prob = 3:1),
    Var_b = sample(c("Test 1",
                    "Test 2",
                    "Test 3"),
                  size = n,
                  replace = TRUE,
                  prob = 1:3))
mtrx <- with(my_data,
             table(Var_a, Var_b))

# Initialize the transition plot
transitions <-
  getRefClass("Transition")$new(mtrx, label=c("Before", "After"))

# Render the plot
transitions$render()
  
```

---

|                |                          |
|----------------|--------------------------|
| transitionPlot | <i>A transition plot</i> |
|----------------|--------------------------|

---

## Description

This plot's purpose is to illustrate how states change before and after. In my research I use it before surgery and after surgery but it can be used in any situation where you have a change from one state to another

## Usage

```
transitionPlot(transition_flow, type_of_arrow = c("grid", "simple",
  "gradient"), box_txt = rownames(transition_flow), tot_spacing = 0.2,
  box_width = 1/4, fill_start_box = "darkgreen",
  txt_start_clr = "white", fill_end_box = fill_start_box,
  txt_end_clr = txt_start_clr, cex = 2, min_lwd = if (type_of_arrow
  == "grid") 1 else unit(0.1, "mm"), max_lwd = if (type_of_arrow ==
  "grid") 6 else unit(5, "mm"), lwd_prop_total = TRUE,
  arrow_clr = "#000000", abs_arrow_width = FALSE,
  overlap_bg_clr = "#FFFFFF", overlap_order = 1:nrow(transition_flow),
  overlap_add_width = if (type_of_arrow == "grid") 1.5 else unit(1,
  "mm"), box_prop, mar = unit(rep(3, times = 4), "mm"), main = NULL,
  box_label = NULL, box_label_pos = "top", box_label_cex = cex,
  color_bar = TRUE, color_bar_cex = cex * 0.33, color_bar_labels,
  color_bar_subspace = NULL, new_page = FALSE)
```

## Arguments

|                 |   |
|-----------------|---|
| transition_flow | This should be a matrix with the size of the transitions. The unit for each cell should be number of observations, row/column-proportions will show incorrect sizes. The matrix needs to be square. The best way to generate this matrix is probably just do a <code>table(starting_state, end_state)</code> . The rows represent the starting positions, while the columns the end positions. I.e. the first row's third column is the number of observations that go from the first class to the third class. |
| type_of_arrow   | The types of arrow may be grid, simple, or gradient. Simple grid arrows are the <code>bezierGrob</code> arrows (not that pretty), simple is the <code>bezierArrowSmpl</code> that I've created to get a more exact control of the arrow position and width, while gradient corresponds to <code>bezierArrowGradient</code> allowing the arrow to have a fill color that slowly turns into the color of the arrow.   |
| box_txt         | The text to appear inside of the boxes. If you need line breaks then you need to manually add a <code>\n</code> inside the string.  |
| tot_spacing     | The proportion of the vertical space that is to be left empty. It is then split evenly between the boxes.   |
| box_width       | The width of the box. By default the box is one fourth of the plot width.   |

|                   |   |
|-------------------|---|
| fill_start_box    | The fill color of the start boxes. This can either be a single value or a vector if you desire different colors for each box. If you specify box_prop then this has to be a 2 column matrix.  |
| txt_start_clr     | The text color of the start boxes. This can either be a single value or a vector if you desire different colors for each box. If you specify box_prop then this has to be a 2 column matrix.  |
| fill_end_box      | The fill color of the end boxes. This can either be a single value or a vector if you desire different colors for each box. If you specify box_prop then this has to be a 2 column matrix.  |
| txt_end_clr       | The text color of the end boxes. This can either be a single value or a vector if you desire different colors for each box. If you specify box_prop then this has to be a 2 column matrix.  |
| cex               | The cex <a href="#">gpar</a> of the text  |
| min_lwd           | The minimum width of the line that we want to illustrate the transition with.   |
| max_lwd           | The maximum width of the line that we want to illustrate the transition with.   |
| lwd_prop_total    | The width of the lines may be proportional to either the other flows from that box, or they may be related to all flows. This is a boolean parameter that is set to true by default, i.e. relating to all flows.  |
| arrow_clr         | The color of the arrows. Usually black, can be a vector indicating each arrow from first to last arrow (counting from the top). If the vector is of the same length as the boxes then all box arrows will have the same color (that is all the arrows stemming from the left boxes) |
| abs_arrow_width   | The width can either be absolute, i.e. each arrow headed for a box has the exact same width. The alternative is that the width is related to the line width.  |
| overlap_bg_clr    | In order to enhance the 3D perspective and to make it easier to follow arrows the arrows have a background color to separate them from those underneath.  |
| overlap_order     | The order from first->last for the lines. This means that the last line will be on top while the first one will appear at the bottom. This should be provided as a vector.  |
| overlap_add_width | The width of the white cross-over line. You can specify this as a scalar multiplication of the current line width. In case of non-grid arrows then you can also have this as a unit which is recommended as it looks better. If the scalar is < 1 then the overlap is ignored.      |
| box_prop          | If you want the boxes to have proportions indicating some other factors then input a matrix with quantiles for the proportions. Note the size must be <code>nrow(transition_flow) x 2</code> .  |
| mar               | A numerical vector of the form <code>c(bottom, left, top, right)</code> of the type <code>unit()</code>   |
| main              | The title of the plot if any, default NULL  |
| box_label         | A vector of length 2 if you want to label each box column   |
| box_label_pos     | The position of the label, either 'top' or 'bottom'   |
| box_label_cex     | The cex of the label, defaults to the default cex   |
| color_bar         | If you have proportions inside the <code>transition_flow</code> variable then the <code>color_bar</code> will automatically appear at the bottom unless you set this to FALSE   |

color\_bar\_cex The size of the tick labels for the color bar

color\_bar\_labels The labels of the two proportions that make up the color bar. Defaults to the labels of the third dimension for the transition\_flow argument.

color\_bar\_subspace If there is little or no difference at the low/high proportions of the spectrum then it can be of interest to focus the color change to the center leaving the tails constant

new\_page If you want the plot to appear on a new blank page then set this to TRUE, by default it is FALSE.

### Value

void

### Examples

```
# This example does not run since it
# takes a little while to assemble the
# arrows and RMD Check complains that this
# is more than allowed for
library(grid)
par_org <- par(ask=TRUE)
# Settings
no_boxes <- 3
# Generate test setting
transition_matrix <- matrix(NA, nrow=no_boxes, ncol=no_boxes)
transition_matrix[1,] <- 200*c(.5, .25, .25)
transition_matrix[2,] <- 540*c(.75, .10, .15)
transition_matrix[3,] <- 340*c(0, .2, .80)

grid.newpage()
transitionPlot(transition_matrix,
  box_txt = c("First", "Second", "Third"),
  type_of_arrow = "simple",
  min_lwd = unit(1, "mm"),
  max_lwd = unit(6, "mm"),
  overlap_add_width = unit(1, "mm"))

# Setup proportions
box_prop <- cbind(c(1,0,0.5), c(.52,.2,.8))
# From the Set2 Colorbrewer
start_box_clr <- c("#8DA0CB", "#FC8D62")
# Darken the colors slightly
end_box_clr <- c(colorRampPalette(c(start_box_clr[1], "#000000"))(10)[2],
  colorRampPalette(c(start_box_clr[2], "#000000"))(10)[2])
# Create a new grid
grid.newpage()
transitionPlot(transition_matrix, box_prop=box_prop,
  fill_start_box=start_box_clr, fill_end_box=end_box_clr,
```

```
txt_start_clr = c("#FFFFFF", "#000000"), txt_end_clr = c("#FFFFFF", "#000000"),  
  box_txt = c("First", "Second", "Third"),  
  type_of_arrow = "gradient",  
  min_lwd = unit(1, "mm"),  
  max_lwd = unit(10, "mm"),  
  overlap_add_width = unit(1, "mm"))  
par(par_org)
```

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