

# Package 'forestplot'

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**Title** Advanced Forest Plot Using 'grid' Graphics

**Description** A forest plot that allows for multiple confidence intervals per row, custom fonts for each text element, custom confidence intervals, text mixed with expressions, and more. The aim is to extend the use of forest plots beyond meta-analyses. This is a more general version of the original 'rmeta' package's forestplot() function and relies heavily on the 'grid' package.

**License** GPL-2

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

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

**Biarch** yes

**Depends** grid, magrittr, checkmate

**Suggests** testthat, abind, knitr, rmarkdown

**Encoding** UTF-8

**NeedsCompilation** no

**VignetteBuilder** knitr

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**Author** Max Gordon [aut, cre],  
Thomas Lumley [aut, ctb]

**Maintainer** Max Gordon <max@gforge.se>

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forestplot-package	<i>Package description</i>
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## Description

The forest plot function, [forestplot](#), is a more general version of the original **rmeta**-packages [forestplot](#) implementation. The aim is at using forest plots for more than just meta-analyses.

## Details

The forestplot:

1. Allows for multiple confidence intervals per row
2. Custom fonts for each text element
3. Custom confidence intervals
4. Text mixed with expressions
5. Legends both on top/left of the plot and within the graph
6. Custom line height including auto-adapt height
7. Graph width that auto-adapts
8. Flexible arguments
9. and more

## Additional functions

The [getTicks](#) tries to format ticks for plots in a nicer way. The major use is for exponentials where ticks are generated using the  $2^n$  since a doubling is a concept easy to grasp for less mathematical-savvy readers.

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forestplot	<i>Draws a forest plot</i>
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## Description

The *forestplot* is based on the **rmeta**-package's [forestplot](#) function. This function resolves some limitations of the original functions such as:

- Adding expressions: Allows use of expressions, e.g. `expression(beta)`
- Multiple bands: Using multiple confidence bands for the same label
- Autosize: Adapts to viewport (graph) size

## Usage

```
forestplot(...)
```

```
## Default S3 method:
```

```
forestplot(labeltext, mean, lower, upper, align,
  is.summary = FALSE, graph.pos = "right", hrzl_lines, clip = c(-Inf,
  Inf), xlab = "", zero = ifelse(xlog, 1, 0), graphwidth = "auto", colgap,
  lineheight = "auto", line.margin, col = fpColors(), txt_gp = fpTxtGp(),
  xlog = FALSE, xticks, xticks.digits = 2, grid = FALSE, lwd.xaxis,
  lwd.zero, lwd.ci, lty.ci = 1, ci.vertices, ci.vertices.height = 0.1,
  boxsize, mar = unit(rep(5, times = 4), "mm"), title, legend,
  legend_args = fpLegend(), new_page = getOption("forestplot_new_page",
  TRUE), fn.ci_norm = fpDrawNormalCI, fn.ci_sum = fpDrawSummaryCI,
  fn.legend, ...)
```

## Arguments

...	Passed on to the <code>fn.ci_norm</code> and <code>fn.ci_sum</code> arguments
labeltext	A list, matrix, vector or expression with the names of each row. The list should be wrapped in <code>m x n</code> number to resemble a matrix: <code>list(list("rowname 1 col 1", "rowname 2 col 1", ...))</code> . You can also provide a matrix although this cannot have expressions by design: <code>matrix(c("rowname 1 col 1", "rowname 2 col 1", "r1c2", "beta"), ncol=2)</code> . Use <code>NA:s</code> for blank spaces and if you provide a full column with <code>NA</code> then that column is a empty column that adds some space. <i>Note</i> : If you do not provide the <code>mean/lower/upper</code> arguments the function expects the label text to be a matrix containing the <code>labeltext</code> in the rownames and then columns for <code>mean</code> , <code>lower</code> , and <code>upper</code> .
mean	A vector or a matrix with the averages. You can also provide a 2D/3D matrix that is automatically converted to the <code>lower/upper</code> parameters. The values should be in exponentiated form if they follow this interpretation, e.g. use <code>exp(mean)</code> if you have the output from a logistic regression
lower	The lower bound of the confidence interval for the forestplot, needs to be the same format as the <code>mean</code> , i.e. matrix/vector of equal columns & length

upper	The upper bound of the confidence interval for the forestplot, needs to be the same format as the mean, i.e. matrix/vector of equal columns & length
align	Vector giving alignment (l,r,c) for the table columns
is.summary	A vector indicating by TRUE/FALSE if the value is a summary value which means that it will have a different font-style
graph.pos	The position of the graph element within the table of text. The position can be $1 - (\text{ncol}(\text{labeltext}) + 1)$ . You can also choose set the positin to "left" or "right".
hrzl_lines	Add horizontal lines to graph. Can either be TRUE or a list of <a href="#">gpar</a> . See line section below for details.
clip	Lower and upper limits for clipping confidence intervals to arrows
xlab	x-axis label
zero	x-axis coordinate for zero line. If you provide a vector of length 2 it will print a rectangle instead of just a line.
graphwidth	Width of confidence interval graph, see <a href="#">unit</a> for details on how to utilize mm etc. The default is auto, that is it uses up whatever space that is left after adjusting for text size and legend
colgap	Sets the gap between columns, defaults to 6 mm but for relative widths. Note that the value should be in <a href="#">unit</a> (, "npc").
lineheight	Height of the graph. By default this is auto and adjusts to the space that is left after adjusting for x-axis size and legend. Sometimes it might be desireable to set the line height to a certain height, for instance if you have several forestplots you may want to standardize their line height, then you set this variable to a certain height, note this should be provided as a <a href="#">unit</a> object. A good option is to set the line height to <a href="#">unit</a> (2, "cm"). A third option is to set line height to "lines" and then you get 50 % more than what the text height is as your line height
line.margin	Set the margin between rows, provided in numeric or <a href="#">unit</a> format. When having multiple confidence lines per row setting the correct margin in order to visually separate rows
col	Set the colors for all the elements. See <a href="#">fpColors</a> for details
txt_gp	Set the fonts etc for all text elements. See <a href="#">fpTxtGp</a> for details
xlog	If TRUE, x-axis tick marks are to follow a logarithmic scale, e.g. for logistic regressoin (OR), survival estimates (HR), poisson regression etc. <i>Note:</i> This is an intentional break with the original <a href="#">forestplot</a> function as I've found that exponentiated ticks/clips/zero effect are more difficult to for non-statisticians and there are sometimes issues with rounding the tick marks properly.
xticks	Optional user-specified x-axis tick marks. Specify NULL to use the defaults, numeric(0) to omit the x-axis. By adding a labels-attribute, <code>attr(my_ticks, "labels") &lt;- ...</code> you can dictate the outputted text at each tick. If you specify a boolean vector then ticks indicated with FALSE wont be printed. Note that the labels have to be the same length as the main variable.
xticks.digits	The number of digits to allow in the x-axis if this is created by default

grid	If you want a discrete gray dashed grid at the level of the ticks you can set this parameter to TRUE. If you set the parameter to a vector of values lines will be drawn at the corresponding positions. If you want to specify the <code>gpar</code> of the lines then either directly pass a <code>gpar</code> object or set the <code>gp</code> attribute e.g. <code>attr(line_vector, "gp") &lt;- gpar(lty=2, col = "red")</code>
lwd.xaxis	lwd for the xaxis, see <code>gpar</code>
lwd.zero	lwd for the vertical line that gives the no-effect line, see <code>gpar</code>
lwd.ci	lwd for the confidence bands, see <code>gpar</code>
lty.ci	lty for the confidence bands, see <code>gpar</code>
ci.vertices	Set this to TRUE if you want the ends of the confidence intervals to be shaped as a T. This is set default to TRUE if you have any other line type than 1 since there is a risk of a dash occurring at the very end, i.e. showing incorrectly narrow confidence interval.
ci.vertices.height	The height of the vertices. Defaults to npc units corresponding to 10% of the row height. <i>Note that the arrows correspond to the vertices heights.</i>
boxsize	Override the default box size based on precision
mar	A numerical vector of the form <code>c(bottom, left, top, right)</code> of the type <code>unit</code>
title	The title of the plot if any
legend	Legend corresponding to the number of bars
legend_args	The legend arguments as returned by the <code>fpLegend</code> function.
new_page	If you want the plot to appear on a new blank page then set this to TRUE, by default it is TRUE. If you want to change this behavior for all plots then set the <code>options(forestplot_new_page = FALSE)</code>
fn.ci_norm	You can specify exactly how the line with the box is drawn for the normal (i.e. non-summary) confidence interval by changing this parameter to your own function or some of the alternatives provided in the package. It defaults to the box function <code>fpDrawNormalCI</code>
fn.ci_sum	Same as previous argument but for the summary outputs and it defaults to <code>fpDrawSummaryCI</code> .
fn.legend	What type of function should be used for drawing the legends, this can be a list if you want different functions. It defaults to a box if you have anything else than a single function or the number of columns in the <code>mean</code> argument

## Details

See `vignette("forestplot")` for details.

## Value

NULL

## Multiple bands

Using multiple bands, i.e. multiple lines, per variable can be interesting when you want to compare different outcomes. E.g. if you want to compare survival specific to heart disease to overall survival for smoking it may be useful to have two bands on top of each other. Another useful implementation is to show crude and adjusted estimates as separate bands.

## Horizontal lines

The argument `hrzl_lines` can be either TRUE or a list with `gpar` elements:

- TRUE A line will be added based upon the `is.summary` rows. If the first line is a summary it
- `gpar` The same as above but the lines will be formatted according to the `gpar` element
- list The list must either be numbered, i.e. `list("2" = gpar(lty=1))`, or have the same length as the `NROW(mean) + 1`. If the list is numbered the numbers should not exceed the `NROW(mean) + 1`. The no. 1 row designates the top, i.e. the line above the first row, all other correspond to the row below. Each element in the list needs to be TRUE, NULL, or `gpar` element. The TRUE defaults to a standard line, the NULL skips a line, while `gpar` corresponds to the fully customized line. Apart from allowing standard `gpar` line descriptions, `lty`, `lwd`, `col`, and more you can also specify `gpar(columns = c(1:3, 5))` if you for instance want the line to skip a column.

## Known issues

The x-axis does not entirely respect the margin. Autosizing boxes is not always the best option, try to set these manually as much as possible.

## API-changes from `rmeta`-package's `forestplot`

- `xlog`: The `xlog` outputs the axis in `log()` format but the input data should be in `antilog/exp` format
- `col`: The corresponding function is `fpColors` for this package

## Author(s)

Max Gordon, Thomas Lumley

## See Also

Other forestplot functions: `fpColors`, `fpDrawNormalCI`, `fpLegend`

## Examples

```
#####
# Simple examples of how to do a forestplot #
#####

ask <- par(ask=TRUE)

# A basic example, create some fake data
```

```
row_names <- list(list("test = 1", expression(test >= 2)))
test_data <- data.frame(coef=c(1.59, 1.24),
                        low=c(1.4, 0.78),
                        high=c(1.8, 1.55))
forestplot(row_names,
            test_data$coef,
            test_data$low,
            test_data$high,
            zero = 1,
            cex = 2,
            lineheight = "auto",
            xlab = "Lab axis txt")

# Print two plots side by side using the grid
# package's layout option for viewports
grid.newpage()
pushViewport(viewport(layout = grid.layout(1, 2)))
pushViewport(viewport(layout.pos.col = 1))
forestplot(row_names,
            test_data$coef,
            test_data$low,
            test_data$high,
            zero = 1,
            cex = 2,
            lineheight = "auto",
            xlab = "Lab axis txt",
            new_page = FALSE)
popViewport()
pushViewport(viewport(layout.pos.col = 2))
forestplot(row_names,
            test_data$coef,
            test_data$low,
            test_data$high,
            zero = 1,
            cex = 2,
            lineheight = "auto",
            xlab = "Lab axis txt",
            new_page = FALSE)
popViewport(2)

# An advanced test
test_data <- data.frame(coef1=c(1, 1.59, 1.3, 1.24),
                        coef2=c(1, 1.7, 1.4, 1.04),
                        low1=c(1, 1.3, 1.1, 0.99),
                        low2=c(1, 1.6, 1.2, 0.7),
                        high1=c(1, 1.94, 1.6, 1.55),
                        high2=c(1, 1.8, 1.55, 1.33))

col_no <- grep("coef", colnames(test_data))
row_names <- list(
  list("Category 1", "Category 2", "Category 3", expression(Category >= 4)),
  list("ref",
```

```

substitute(expression(bar(x) == val),
             list(val = round(rowMeans(test_data[2, col_no]), 2))),
substitute(expression(bar(x) == val),
             list(val = round(rowMeans(test_data[3, col_no]), 2))),
substitute(expression(bar(x) == val),
             list(val = round(rowMeans(test_data[4, col_no]), 2)))
)

coef <- with(test_data, cbind(coef1, coef2))
low <- with(test_data, cbind(low1, low2))
high <- with(test_data, cbind(high1, high2))
forestplot(row_names, coef, low, high,
            title="Cool study",
            zero = c(0.98, 1.02),
            grid = structure(c(2^-0.5, 2^0.5), gp = gpar(col = "steelblue", lty=2)),
            boxsize=0.25,
            col=fpColors(box=c("royalblue", "gold"),
                          line=c("darkblue", "orange"),
                          summary=c("darkblue", "red")),
            xlab="The estimates",
            new_page = TRUE,
            legend=c("Treatment", "Placebo"),
            legend_args = fpLegend(pos = list("topright"),
                                    title="Group",
                                    r = unit(.1, "snpc"),
                                    gp = gpar(col="#CCCCCC", lwd=1.5)))

# An example of how the exponential works
test_data <- data.frame(coef=c(2.45, 0.43),
                        low=c(1.5, 0.25),
                        high=c(4, 0.75),
                        boxsize=c(0.5, 0.5))
row_names <- cbind(c("Name", "Variable A", "Variable B"),
                  c("HR", test_data$coef))
test_data <- rbind(rep(NA, 3), test_data)

forestplot(labeltext = row_names,
            test_data[,c("coef", "low", "high")],
            is.summary=c(TRUE, FALSE, FALSE),
            boxsize = test_data$boxsize,
            zero = 1,
            xlog = TRUE,
            col = fpColors(lines="red", box="darkred"))

par(ask=ask)
# See vignette for a more detailed description
# vignette("forestplot", package="forestplot")

```



## Description

This function encapsulates all the colors that are used in the [forestplot](#) function. As there are plenty of color options this function gathers them all in one place.

## Usage

```
fpColors(all.elements, box = "black", lines = "gray", summary = "black",
         zero = "lightgray", text = "black", axes = "black",
         hrz_lines = "black")
```

## Arguments

<code>all.elements</code>	A color for all the elements. If set to NULL then it's set to the <code>par("fg")</code> color
<code>box</code>	The color of the box indicating the estimate
<code>lines</code>	The color of the confidence lines
<code>summary</code>	The color of the summary
<code>zero</code>	The color of the zero line
<code>text</code>	The color of the text
<code>axes</code>	The color of the x-axis at the bottom
<code>hrz_lines</code>	The color of the horizontal lines

## Details

If you have several values per row in a forestplot you can set a color to a vector where the first value represents the first line/box, second the second line/box etc. The vectors are only valid for the `box` & `lines` options.

This function is a copy of the [meta.colors](#) function in the **rmeta** package.

## Value

list A list with the elements:

<code>box</code>	the color of the box/marker
<code>lines</code>	the color of the lines
<code>summary</code>	the color of the summary
<code>zero</code>	the color of the zero vertical line
<code>text</code>	the color of the text
<code>axes</code>	the color of the axes

## Author(s)

Max Gordon, Thomas Lumley

## See Also

Other forestplot functions: [forestplot](#), [fpDrawNormalCI](#), [fpLegend](#)

**Examples**

```
ask <- par(ask=TRUE)

# An example of how the exponential works
test_data <- data.frame(coef=c(2.45, 0.43),
                       low=c(1.5, 0.25),
                       high=c(4, 0.75),
                       boxsize=c(0.5, 0.5))
row_names <- cbind(c("Name", "Variable A", "Variable B"),
                  c("HR", test_data$coef))
test_data <- rbind(rep(NA, 3), test_data)

forestplot(labeltext = row_names,
           test_data[,c("coef", "low", "high")],
           is.summary=c(TRUE, FALSE, FALSE),
           boxsize = test_data$boxsize,
           zero = 1,
           xlog = TRUE,
           col = fpColors(lines="#990000", box="#660000", zero = "darkblue"),
           new_page = TRUE)

par(ask=ask)
```

---

fpDrawNormalCI

*Draw standard confidence intervals*


---

**Description**

A function that is used to draw the different confidence intervals for the non-summary lines. Use the fpDrawNormalCI function as a template if you want to make your own funky line + marker.

**Usage**

```
fpDrawNormalCI(lower_limit, estimate, upper_limit, size, y.offset = 0.5,
               clr.line, clr.marker, lwd, lty = 1, vertices, vertices.height = 0.1, ...)
```

```
fpDrawDiamondCI(lower_limit, estimate, upper_limit, size, y.offset = 0.5,
                clr.line, clr.marker, lwd, lty = 1, vertices, vertices.height = 0.1, ...)
```

```
fpDrawCircleCI(lower_limit, estimate, upper_limit, size, y.offset = 0.5,
               clr.line, clr.marker, lwd, lty = 1, vertices, vertices.height = 0.1, ...)
```

```
fpDrawPointCI(lower_limit, estimate, upper_limit, size, y.offset = 0.5,
               clr.line, clr.marker, lwd, lty = 1, vertices, vertices.height = 0.1,
               pch = 1, ...)
```

```
fpDrawSummaryCI(lower_limit, estimate, upper_limit, size, col, y.offset = 0.5,
                 ...)
```

```
fpDrawBarCI(lower_limit, estimate, upper_limit, size, col, y.offset = 0.5,
...)
```

### Arguments

lower_limit	The lower limit of the confidence line. A native numeric variable that can actually be outside the boundaries. If you want to see if it is outside then convert it to 'npc' and see if the value ends up more than 1 or less than 0. Here's how you do the conversion: <code>convertX(unit(upper_limit, "native"), "npc", valueOnly = TRUE)</code> and the <code>convertX</code> together with <code>unit</code> is needed to get the right values while you need to provide the <code>valueOnly</code> as you cannot compare a unit object.
estimate	The estimate indicating the placement of the actual box. Note, this can also be outside bounds and is provided in a numeric format the same way as the <code>lower_limit</code> .
upper_limit	The upper limit of the confidence line. See <code>lower_limit</code> for details.
size	The actual size of the box/diamond/marker. This provided in the 'npc' format to generate a perfect marker. Although you can provide it alternative units as well, this is useful for the legends to work nicely.
y.offset	If you have multiple lines they need an offset in the y-direction.
clr.line	The color of the line.
clr.marker	The color of the estimate marker
lwd	Line width, see <a href="#">gpar</a>
lty	Line type, see <a href="#">gpar</a>
vertices	Set this to TRUE if you want the ends of the confidence intervals to be shaped as a T. This is set default to TRUE if you have any other line type than 1 since there is a risk of a dash occurring at the very end, i.e. showing incorrectly narrow confidence interval.
vertices.height	The height of the vertices. Defaults to npc units corresponding to 10% of the row height.
...	Allows additional parameters for sibling functions
pch	Type of point see <a href="#">grid.points</a> for details
col	The color of the summary object

### Value

void The function outputs the line using grid compatible functions and does not return anything.

### Author(s)

Max Gordon, Thomas Lumley

### See Also

Other forestplot functions: [forestplot](#), [fpColors](#), [fpLegend](#)

**Examples**

```

ask <- par(ask=TRUE)

test_data <- data.frame(coef1=c(1, 1.59, 1.3, 1.24),
                       coef2=c(1, 1.7, 1.4, 1.04))

test_data$low1 <- test_data$coef1 - 1.96*c(0, .2, .1, .15)
test_data$high1 <- test_data$coef1 + 1.96*c(0, .2, .1, .15)

test_data$low2 <- test_data$coef2 - 1.96*c(0, .1, .15, .2)
test_data$high2 <- test_data$coef2 + 1.96*c(0, .1, .15, .2)

col_no <- grep("coef", colnames(test_data))
row_names <- list(
  list("Category 1", "Category 2", "Category 3", expression(Category >= 4)),
  list("ref",
       substitute(expression(bar(x) == val),
                   list(val = round(rowMeans(test_data[2, col_no]), 2))),
       substitute(expression(bar(x) == val),
                   list(val = round(rowMeans(test_data[3, col_no]), 2))),
       substitute(expression(bar(x) == val),
                   list(val = round(rowMeans(test_data[4, col_no]), 2))))
)

coef <- with(test_data, cbind(coef1, coef2))
low <- with(test_data, cbind(low1, low2))
high <- with(test_data, cbind(high1, high2))

# Change all to diamonds
forestplot(row_names, coef, low, high,
           fn.ci_norm=fpDrawDiamondCI,
           title="Cool study",
           zero = 1, boxsize=0.25,
           col=fpColors(box=c("royalblue", "gold"),
                        line=c("darkblue", "orange"),
                        summary=c("darkblue", "red")),
           xlab="The estimates",
           new_page = TRUE,
           legend=c("Treatment", "Placebo"),
           legend_args = fpLegend(title="Group",
                                   pos = list("topright", inset=.1),
                                   r=unit(.1, "snpc"),
                                   gp = gpar(col="#CCCCCC", lwd=1.5)))

# Change first to diamonds
forestplot(row_names, coef, low, high,
           fn.ci_norm=c("fpDrawDiamondCI",
                       rep("fpDrawNormalCI",
                           times=nrow(coef)-1)),
           title="Cool study",
           zero = 1, boxsize=0.25,
           col=fpColors(box=c("royalblue", "gold"),

```

```

        line=c("darkblue", "orange"),
        summary=c("darkblue", "red")),
xlab="The estimates",
new_page = TRUE,
legend=c("Treatment", "Placebo"),
legend_args = fpLegend(title="Group",
                        pos = list("topright", inset=.1),
                        r=unit(.1, "snpc"),
                        gp = gpar(col="#CCCCCC", lwd=1.5)))

# You can also use a list with the actual functions
# as long as it is formatted [[row]][[column]]
# Note: if you have a non-square input then
# the software will reformat [[col]][[row]]
# to [[row]][[col]]
forestplot(row_names, coef, low, high,
           fn.ci_norm=list(list(fpDrawDiamondCI, fpDrawCircleCI),
                           list(fpDrawNormalCI, fpDrawNormalCI),
                           list(fpDrawNormalCI, fpDrawCircleCI),
                           list(fpDrawNormalCI, fpDrawNormalCI)),
           title="Cool study",
           zero = 1, boxsize=0.25,
           col=fpColors(box=c("royalblue", "gold"),
                        line=c("darkblue", "orange"),
                        summary=c("darkblue", "red")),
           xlab="The estimates",
           new_page = TRUE,
           legend=c("Treatment", "Placebo"),
           legend_args = fpLegend(title="Group",
                                   pos = list("topright", inset=.1),
                                   r=unit(.1, "snpc"),
                                   gp = gpar(col="#CCCCCC", lwd=1.5)))

par(ask=ask)

```

---

fpLegend

*A function for the legend used in forestplot()*


---

## Description

This function encapsulates all the legend options that are used in the `forestplot` function. This is in order to limit the crowding among the arguments for the `forestplot` call.

## Usage

```

fpLegend(pos = "top", gp = NULL, r = unit(0, "snpc"),
         padding = unit(ifelse(!is.null(gp), 3, 0), "mm"), title = NULL)

```

**Arguments**

pos	The position of the legend, either at the "top" or the "right" unless positioned inside the plot. If you want the legend to be positioned inside the plot then you have to provide a list with the same x & y qualities as <a href="#">legend</a> . For instance if you want the legend to be positioned at the top right corner then use <code>pos = list("topright")</code> - this is equivalent to <code>pos = list(x=1, y=1)</code> . If you want to have a distance from the edge of the graph then add a inset to the list, e.g. <code>pos = list("topright", "inset"=.1)</code> - the inset should be either a <a href="#">unit</a> element or a value between 0 and 1. The default is to have the boxes aligned vertical, if you want them to be in a line then you can specify the "align" option, e.g. <code>pos = list("topright", "inset"=.1, "align"="horizontal")</code>
gp	The <a href="#">gpar</a> options for the legend. If you want the background color to be light grey then use <code>gp = gpar(fill = "lightgrey")</code> . If you want a border then set the col argument: <code>gp = gpar(fill = "lightgrey", col="black")</code> . You can also use the lwd and lty argument as usual, <code>gp = gpar(lwd=2, lty=1)</code> , will result in a black border box of line type 1 and line width 2.
r	The box can have rounded edges, check out <a href="#">grid.roundrect</a> . The r option should be a <a href="#">unit</a> object. This is by default <code>unit(0, "snpc")</code> but you can choose any value that you want. The "snpc" unit is the preferred option.
padding	The padding for the legend box, only used if box is drawn. This is the distance from the border to the text/boxes of the legend.
title	The title of the legend if any

**Value**

`list` Returns a list with all the elements

**See Also**

Other forestplot functions: [forestplot](#), [fpColors](#), [fpDrawNormalCI](#)

---

fpTxtGp

*Get font settings for forestplot*


---

**Description**

This function generates all the [gpar\(\)](#) elements for the different text elements within the graph. Elements not specified inherit their default settings from the label argument.

**Usage**

```
fpTxtGp(label, summary, xlab, title, ticks, legend, legend.title, cex = 1)
```

**Arguments**

label	The text labels (see details below)
summary	The summary labels (see details below)
xlab	The xlab text
title	The plot title
ticks	The ticks associated with the xlab
legend	The legend text
legend.title	The legend title
cex	The font size

**Value**

A list of the fpTxtGp class

**List arguments for label/summary**

You can provide a list of elements for the label and summary in order to specify separate elements. If you provide a list in one dimension the gpar elements are assumed to follow the columns. If you provide a list of 2 dimensions the structure assumes is `list[[row]][[column]]` and the number of elements should correspond to the number of labels for the label argument, i.e. without the rows marked as summary elements. The same goes for summary arguments.

**Examples**

```
fpTxtGp(label=gpar(fontfamily="HersheySerif"))
```

---

getTicks	<i>Ticks for plot axis</i>
----------	----------------------------

---

**Description**

Gets the ticks in a formatted version. This is since I'm not always that fond of just pretty(1:10/5). In exponential form the ticks are determined from the 2-base, meaning that you get an intuitive feeling for when the value is doubled.

**Usage**

```
getTicks(low, high = low, clip = c(-Inf, Inf), exp = FALSE, digits = 0)
```

**Arguments**

low	lower bound, can be a single number or a vector
high	upper bound - optional, you can just have all data in the low variable
clip	if the ci are clipped
exp	If the value should be in exponential form (default)
digits	Number of digits - used in exp mode

**Details**

This function is far from perfect and I recommend specifying yourself the ticks that you want.

**Value**

vector Returns a vector with the ticks

**Examples**

```
test_data <- data.frame(coef=c(2, 0.5),
  low=c(1.5, 0.05),
  high=c(3, 0.75),
  boxsize=c(0.5, 0.5))

# Exponential form where the exponent base is 2 for easier understanding
getTicks(low = test_data$low,
  high = test_data$high,
  clip=c(-Inf, Inf),
  exp=TRUE)

# Non exponential form with using pretty
getTicks(low = test_data$low,
  high = test_data$high,
  clip=c(-Inf, Inf),
  exp=FALSE)

# A very simple example
getTicks(1:5*2.33,
  exp=FALSE)

# A slightly more advanced exponential version
getTicks(1:10*.33,
  digits=2,
  exp=TRUE)
```

---

HRQoL

*Regression coefficients and confidence intervals from HRQoL study*

---

**Description**

The data is a list containing the Swedish and the Danish coefficients for health related quality of life (HRQoL) 1 year after total hip arthroplasty surgery. The age is modelled as a spline and is therefore presented as a contrast.

**Author(s)**

Max Gordon <max@gforge.se>



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