Package 'easylabel'

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```
Type Package
```

Title Interactive Scatter Plot and Volcano Plot Labels

Version 0.2.8

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Description Interactive labelling of scatter plots, volcano plots and Manhattan plots using a 'shiny' and 'plotly' interface. Users can hover over points to see where specific points are located and click points on/off to easily label them. Labels can be dragged around the plot to place them optimally. Plots can be exported directly to PDF for publication.

```
Language en-gb
```

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Encoding UTF-8

LazyData true

biocViews

Suggests AnnotationDbi, knitr, org.Hs.eg.db, qvalue, rmarkdown

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NeedsCompilation no

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R topics documented:

Index																													15
	volc2	•	 •	•	 •	٠	•	•		•	•	•	•	•	 •	٠	•	•	•	•	•		 	•	•	•	•	•	14
	volc1																												
	qqplot2																												
	qqplot														 							 	 						12
	gg_qqplot														 							 	 						11
	easyVolcano														 							 	 						10
	easyMAplot														 							 	 						9
	easyManhattan														 							 	 						7
	easylabel																					 	 						2

easylabel

Interactive scatter plot labels

Description

Interactive labelling of scatter plots using shiny/plotly interface.

Usage

```
easylabel(
 data,
  Х,
 у,
  labs = NULL,
  startLabels = NULL,
  cex.text = 0.72,
  col = NULL,
  colScheme = NULL,
  alpha = 1,
  shape = NULL,
  shapeScheme = 21,
  size = 8,
  sizeRange = c(4, 80),
  xlab = x,
  ylab = y,
 xlim = NULL,
 ylim = NULL,
 xticks = NULL,
  yticks = NULL,
  showOutliers = TRUE,
  outlier_shape = 5,
  outline_col = "white",
  outline_lwd = 0.5,
  plotly_filter = NULL,
```

```
width = 800,
 height = 600,
  showgrid = FALSE,
  zeroline = TRUE,
 hline = NULL,
 vline = NULL,
 mgp = c(1.8, 0.5, 0),
 Ltitle = "",
 Rtitle = "",
 LRtitle_side = 1,
 labelDir = "radial",
  labCentre = NULL,
  lineLength = 75,
  text_col = "black",
  line_col = "black",
  rectangles = FALSE,
  rect_col = "white",
  border_col = "black",
  padding = 3,
  border_radius = 5,
  showLegend = TRUE,
  legendxy = c(1.02, 1),
  filename = NULL,
  panel.last = NULL,
  fullGeneNames = FALSE,
 AnnotationDb = NULL,
  custom_annotation = NULL,
 output_shiny = TRUE,
)
```

Arguments

data	Dataset (data.frame or data.table) to use for plot.
x	Specifies column of x coordinates in data.
у	Specifies column of y coordinates in data.
labs	Specifies the column in data with label names for points. Label names do not have to be unique. If NULL defaults to rownames(data).
startLabels	Vector of initial labels. With a character vector, labels are identified in the column specified by labs. With a numeric vector, points to be labelled are referred to by row number.
cex.text	Font size for labels. Default 0.72 to match plotly font size. See text().
col	Specifies which column in data affects point colour. Must be categorical. If it is not a factor, it will be coerced to a factor.
colScheme	A single colour or a vector of colours for points.
alpha	Alpha value for transparency of points.

shape Specifies which column in data controls point shapes. If not a factor, will be

coerced to a factor.

shapeScheme A single symbol for points or a vector of symbols. See pch in points().

size Either a single value for size of points (default 8), or specifies which column in

data affects point size for bubble charts.

sizeRange Range of size of points for bubble charts.

xlab x axis title. Accepts expressions when exporting base graphics. Set cex.lab to

alter the font size of the axis titles (default 1). Set cex.axis to alter the font size

of the axis numbering (default 1).

ylab y axis title. Accepts expressions when exporting base graphics.

xlim The x limits (x1, x2) of the plot.

ylim The y limits of the plot.

xticks List of custom x axis ticks and labels specified as a list of two named vectors at

= ... and labels = Another method is to use xaxp as a vector of the form c(x1, x2, n) giving the coordinates of the extreme tick marks and the number

of intervals between tick-marks.

yticks List of custom y axis ticks and labels specified as a list of two named vectors at

= ... and labels = Another method is to use yaxp as a vector of the form c(y1, y2, n) giving the coordinates of the extreme tick marks and the number

of intervals between tick-marks.

showOutliers Logical whether to show outliers on the margins of the plot.

outlier_shape Symbol for outliers.

outline_col Colour of symbol outlines. Set to NA for no outlines.

outline_lwd Line width of symbol outlines.

plotly_filter Refers to a column of logical values in data used to filter rows to reduce the

number of points shown by plotly. We recommend using this for datasets with >100,000 rows. When saving to pdf, the full original dataset is still plotted. This is useful for plots with millions of points such as Manhattan plots where a subset

of points to be labelled is already known.

width Width of the plot in pixels. Saving to pdf scales 100 pixels to 1 inch.

height Height of the plot in pixels.

showgrid Either logical whether to show gridlines, or a character value where "x" means

showing x axis gridlines and "y" means showing y axis gridlines.

zeroline Logical whether to show lines at x = 0 and y = 0.

hline Adds horizontal lines at values of y.
vline Adds vertical lines at values of x.

mgp The margin line for the axis title, axis labels and axis line. See par().

Ltitle A character or expression (see plotmath) value specifying text for left side title.

Size of font can be changed using cex.lab.

Rtitle A character or expression value specifying text for right side title. Size of font

can be changed using cex.lab.

LRtitle_side On which side of the plot for Ltitle and Rtitle (1 = bottom, 3 = top). See

mtext().

labelDir Initial label line directions. Options include 'radial' (default) for radial lines

around the centre of the plot, 'origin' for radial lines around the origin, 'horiz' for horizontal and 'vert' for vertical, 'xellipse' and 'yellipse' for near-horizontal and near-vertical lines arranged in an elliptical way around the centre, 'rect' for rectilinear lines (a mix of horizontal and vertical), 'x' for diagonal lines, 'oct'

for lines in 8 directions around the centre.

labCentre Coordinates in x/y units of the central point towards which radial labels con-

verge. Defaults to the centre of the plot.

lineLength Initial length of label lines in pixels.

text_col Colour of label text. If set to "match" label text will match the colour of each

point.

line_col Colour of label lines. If set to "match" label line will match the colour of each

point.

rectangles Logical whether to show rectangles around labels (not supported by plotly).

rect_col Colour for filling rectangles (not supported by plotly). If set to "match" rectan-

gle fill colour will match the colour of each point.

border_col Colour of rectangle borders (not supported by plotly). Use border_col = NA to

omit borders. If set to "match" rectangle border colour will match the colour of

each point.

padding Amount of padding in pixels around label text.

border_radius Amount of roundedness in pixels to apply to label rectangles (not supported by

plotly).

showLegend Logical whether to show or hide the legend.

legendxy Vector of coordinates for the position of the legend. Coordinates are in plotly

paper reference with c(0,0) being the bottom left corner and c(1,1) being the top right corner of the plot window. Plotly has unusual behaviour in that the x coordinate always aligns the left side of the legend. However, the y coordinate aligns the top, middle or bottom of the legend dependent on whether it is in the top, middle or bottom 1/3 of the plot window. So c(1,0) positions the legend in the bottom right corner outside the right margin of the plot, while c(1,0.5)

centre aligns the legend around the centre of y axis.

filename Filename for saving plots to pdf in a browser. Rstudio opens its own pdf file.

panel.last An expression to be evaluated after plotting has taken place but before the axes,

title and box are added. This can be useful for adding extra titles, legends or trend lines. Currently only works when saving plots using base graphics and

does not work with plotly. See plot.default

fullGeneNames Logical whether to expand gene symbols using Bioconductor AnnotationDbi

package. With multiple matches, returns first value only. See AnnotationDbi::mapIds().

AnnotationDb Annotation database to use when expanding gene symbols. Defaults to human

gene database AnnotationDb = org. Hs.eg.db.

custom_annotation

List of annotations to be added via plotly::layout().

output_shiny

Logical whether to output a shiny app. If FALSE a plotly figure will be returned.

. .

Further graphical parameters passed to plot() when saving via base graphics. The most useful for most users are likely to be cex.lab which alters axis title font size (default 1, see par()), cex.axis which alters axis numbering font size (default 1), and panel.last which allows additional plotting functions to be called after the main plot has been plotted but before the labels and label lines are drawn, which will allow the addition of trend lines, extra titles or legends for example (see plot.default()).

Details

Instructions:

- Hover over and click on/off genes which you want to label.
- When you have selected all your chosen genes, then drag gene names to move label positions.
- Click the save button to export a PDF in base graphics.
- The Table tab shows a table view of the dataset to help with annotation.

To export an SVG from plotly:

- Switch to SVG when finalised (only do this at last moment as otherwise editing is very slow).
- Press camera button in modebar to save image as SVG.

Value

By default no return value. If output_shiny = FALSE or the shiny button 'Export plotly & exit' is pressed, a plotly figure is returned.

See Also

```
easyVolcano(), easyMAplot()
```

Examples

```
# Simple example using mtcars dataset
data(mtcars)
# Launch easylabel Shiny app: only run this example in interactive R sessions
if (interactive()) {
  easylabel(mtcars, x = 'mpg', y = 'wt', col = 'cyl')
}
```

easyManhattan 7

easyManhattan

Interactive Manhattan plot labels

Description

Interactive labelling of Manhattan plots using 'shiny' and 'plotly' interface.

Usage

```
easyManhattan(
  data,
  chrom = "chrom",
 pos = "pos",
 p = p''
  labs = "rsid",
  startLabels = NULL,
 pcutoff = 5e-08,
  chromGap = NULL,
  chromCols = c("royalblue", "skyblue"),
  sigCol = "red",
  alpha = 0.7,
  labelDir = "horiz",
  xlab = "Chromosome position",
 ylab = expression("-log"[10] ~ "P"),
 xlim = NULL,
 ylim = NULL,
  outline_col = NA,
  shapeScheme = 16,
  size = 6,
 width = ifelse(transpose, 600, 1000),
 height = ifelse(transpose, 800, 600),
 lineLength = 60,
  npoints = max(c(nrow(data)/5, 1e+06)),
  nplotly = 1e+05,
  npeaks = NULL,
  span = 2e+07,
  transpose = FALSE,
  filename = NULL,
)
```

Arguments

data The dataset (data.frame or data.table) for the plot.

chrom The column of chomosome values in data.

pos The column of SNP positions in data.

8 easyManhattan

p The column of p values in data.

labs The column of labels in data.

startLabels Vector of initial labels. With a character vector, labels are identified in the col-

umn specified by labs. With a numeric vector, points to be labelled are referred

to by row number.

pcutoff Cut-off for p value significance. Defaults to 5E-08.

chromGap Size of gap between chromosomes along the x axis in base pairs. If NULL this is

automatically calculated dependent on the size of the genome. Default is around

3E07 for a human genome, and smaller for smaller genomes.

chromCols A vector of colours for points by chromosome. Colours are recycled dependent

on the length of the vector.

sigCol Colour for statistically significant points. Ignored if set to NA.

alpha Transparency for points.

labelDir Option for label lines. See easylabel().

x lab x axis title. Accepts expressions. ylab y axis title. Accepts expressions. xlim The x limits (x1, x2) of the plot.

ylim The y limits of the plot.

outline_col Colour of symbol outlines. Passed to easylabel().

shapeScheme A single symbol for points or a vector of symbols. Passed to easylabel().

size Specifies point size. Passed to easylabel().

width Width of the plot in pixels. Saving to pdf scales 100 pixels to 1 inch.

height Height of the plot in pixels.

lineLength Initial length of label lines in pixels.

npoints Maximum number of points to plot when saving the final plot to pdf. By default

plots with >1 million points are thinned to speed up plotting. Setting a value of

NA will plot all points.

nplotly Maximum number of points to display via plotly. We recommend the default

setting of 100,000 points (or fewer).

npeaks Number of peaks to label initially.

span a peak is defined as the most significant SNP within a window of width span

centred at that SNP.

transpose Logical whether to transpose the plot.

filename Filename for saving to pdf.

... Other arguments passed to easylabel().

Value

By default no return value. If output_shiny = FALSE or the shiny button 'Export plotly & exit' is pressed, a plotly figure is returned. See easylabel().

See Also

easylabel() easyVolcano()

easyMAplot 9

easyMAplot	Interactive MA plot labels

Description

Interactive labelling of MA plots using shiny/plotly interface.

Usage

```
easyMAplot(
  data,
  x = NULL,
  y = NULL,
  padj = NULL,
  fdrcutoff = 0.05,
  colScheme = c("darkgrey", "blue", "red"),
  hline = 0,
  labelDir = "yellipse",
  xlab = expression("log"[2] ~ " mean expression"),
  ylab = expression("log"[2] ~ " fold change"),
  filename = NULL,
  showCounts = TRUE,
  useQ = FALSE,
  ...
)
```

Arguments

data	The dataset for the plot. Automatically attempts to recognises DESeq2 and limma objects.
X	Name of the column containing mean expression. For DESeq2 and limma objects this is automatically set.
У	Name of the column containing log fold change. For DESeq2 and limma objects this is automatically set.
padj	Name of the column containing adjusted p values (optional). For DESeq2 and limma objects this is automatically set. If y is specified and padj is left blank or equal to y, nominal unadjusted p values are used for cut-off for significance.
fdrcutoff	Cut-off for FDR significance. Defaults to FDR < 0.05. Can be vector with multiple cut-offs. To use nominal P values instead of adjusted p values, set y but leave padj blank.
colScheme	Colour colScheme. Length must match either length(fdrcutoff) $+ 1$ to allow for non-significant genes, or match length(fdrcutoff) $* 2 + 1$ to accommodates asymmetric colour colSchemes for positive & negative fold change. (see examples).
hline	Vector of horizontal lines (default is $y = 0$).

10 easyVolcano

labelDir	Option for label lines. See easylabel().
xlab	x axis title. Accepts expressions.
ylab	y axis title. Accepts expressions.
filename	Filename for saving to pdf.
showCounts	Logical whether to show legend with number of differentially expressed genes.
useQ	Logical whether to convert nominal P values to q values. Requires the qvalue Bioconductor package.
	Other arguments passed to easylabel().

Value

By default no return value. If output_shiny = FALSE or the shiny button 'Export plotly & exit' is pressed, a plotly figure is returned. See easylabel().

See Also

```
easylabel() easyVolcano()
```

|--|

Description

Interactive labelling of volcano plots using shiny/plotly interface.

Usage

```
easyVolcano(
  data,
  x = NULL,
  y = NULL,
  padj = y,
  fdrcutoff = 0.05,
  fccut = NULL,
  colScheme = c("darkgrey", "blue", "red"),
  xlab = expression("log"[2] ~ " fold change"),
  ylab = expression("-log"[10] ~ " P"),
  filename = NULL,
  showCounts = TRUE,
  useQ = FALSE,
  ...
)
```

gg_qqplot 11

data	The dataset for the plot. Automatically attempts to recognises DESeq2 and limma objects.
X	Name of the column containing log fold change. For DESeq2 and limma objects this is automatically set.
у	Name of the column containing p values. For DESeq2 and limma objects this is automatically set.
padj	Name of the column containing adjusted p values (optional). If y is specified and padj is left blank or equal to y, nominal unadjusted p values are used for cut-off for significance instead of adjusted p values.
fdrcutoff	Cut-off for FDR significance. Defaults to FDR < 0.05 . If y is specified manually and padj is left blank then this refers to the cut-off for significant points using nominal unadjusted p values.
fccut	Optional vector of log fold change cut-offs.
colScheme	Colour scheme. If no fold change cut-off is set, 2 colours need to be specified. With a single fold change cut-off, 3 or 5 colours are required, depending on whether the colours are symmetrical about $x=0$. Accommodates asymmetric colour schemes with multiple fold change cut-offs (see examples).
xlab	x axis title. Accepts expressions.
ylab	y axis title. Accepts expressions.
filename	Filename for saving to pdf.
showCounts	Logical whether to show legend with number of differentially expressed genes.
useQ	Logical whether to convert nominal P values to q values. Requires the qvalue Bioconductor package.
	Other arguments passed to easylabel().

Value

By default no return value. If output_shiny = FALSE or the shiny button 'Export plotly & exit' is pressed, a plotly figure is returned. See easylabel().

See Also

easylabel() easyMAplot()

	gg_qqplot	Log QQ p-value plot (ggplot2)	
--	-----------	-------------------------------	--

Description

Produces a QQ plot via ggplot2. Requires a dataframe generated by qqplot().

12 qqplot

Usage

```
gg_qqplot(df, scheme = c("darkgrey", "royalblue"))
```

Arguments

df A dataframe generated by qqplot()

scheme Vector of 2 colours for plotting non-significant and significant SNPs

Value

A ggplot2 graphics plot object

qqplot

Log QQ p-value plot

Description

Fast function for generating a log quantile-quantile (QQ) p-value plot

Usage

```
qqplot(
  pval,
  fdr = NULL,
  fdr_cutoff = 0.05,
  scheme = c("darkgrey", "royalblue"),
  npoints = 5e+05,
  show_plot = TRUE,
  verbose = TRUE,
  ...
)
```

Arguments

pval A vector of p-values

fdr An optional vector of FDR values to save time if previously computed. If

not supplied, these will be calculated using p.adjust() using the Benjamini-

Hochberg method.

fdr_cutoff Cutoff for FDR significance

scheme Vector of 2 colours for plotting non-significant and significant SNPs

npoints Limits the number of non-significant points being plotted to speed up plotting.

See details. Set to NULL to plot all points.

show_plot Logical whether to produce a plot via base graphics or just return dataframe

ready for plotting.

verbose Whether to show messages

... Optional plotting arguments passed to qqplot2()

qqplot2

Details

Produces a fast QQ plot. Particularly useful for analyses with very large numbers of p-values (such as eQTL analysis) which can be slow to plot. The function looks first for all comparisons which reached FDR at the designated cut-off and ensures all of these points are plotted. Additional points which typically overlap substantially near the origin are thinned by random sampling. In this way the plot can be reduced from millions of points to 500,000 points with a plot which is indistinguishable from one with all points plotted. For comparison, set npoints to NULL to plot all points as usual.

Calling qqplot() will result in a base graphics plot. The plotting dataframe is returned invisibly, so users can save time when refining plots by saving the dataframe produced by qqplot() and then invoking qqplot2() to simply plot the points. Users who prefer ggplot2 can also pass the dataframe generated by qqplot() to gg_qqplot().

Value

Generates a plot using base graphics. Also returns a dataframe invisibly which can be used for downstream plotting via either qqplot2() or gg_qqplot().

See Also

```
qqplot2() gg_qqplot()
```

qqplot2

Log QQ p-value plot (2nd stage)

Description

Second stage plotting function which accepts dataframe generated by qqplot(). This can be used to avoid repeating computation of the QQ plot values.

Usage

```
qqplot2(df, scheme = c("darkgrey", "royalblue"), ...)
```

Arguments

df A dataframe generated by qqplot()

scheme Vector of 2 colours for plotting non-significant and significant SNPs

... Optional plotting arguments passed to plot()

Value

No return value. Produces a base graphics plot.

volc2

volc1

Example volcano data for vignette

Description

Example DESeq2 volcano data for vignette

Usage

volc1

Format

Data frame with 6 rows and 6 variables

volc2

Example volcano data for vignette

Description

Example limma volcano data for vignette

Usage

volc2

Format

Data frame with 6 rows and 6 variables

Index

```
* datasets
    volc1, 14
    volc2, 14
AnnotationDbi::mapIds(), 5
easylabel, \color{red} 2
easylabel(), 8, 10, 11
easyManhattan, 7
easyMAplot, 9
easyMAplot(), 6, 11
easyVolcano, 10
easyVolcano(), 6, 8, 10
gg_qqplot, 11
gg_qqplot(), 13
mtext(), 5
p.adjust(), 12
par(), 4, 6
plot(), 13
{\tt plot.default}, {\tt 5}
plot.default(),6
plotly::layout(), 5
plotmath, 4
points(), 4
qqplot, 12
qqplot(), 11–13
qqplot2, 13
qqplot2(), 12, 13
text(), 3
volc1, 14
volc2, 14
```