# Package 'EnhancedVolcano'

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Type Package

Title Publication-ready volcano plots with enhanced colouring and labeling

Version 1.24.0

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**Description** Volcano plots represent a useful way to visualise the results of differential expression analyses. Here, we present a highly-configurable function that produces publication-ready volcano plots. EnhancedVolcano will attempt to fit as many point labels in the plot window as possible, thus avoiding 'clogging' up the plot with labels that could not otherwise have been read. Other functionality allows the user to identify up to 4 different types of attributes in the same plot space via colour, shape, size, and shade parameter configurations.

License GPL-3

**Depends** ggplot2, ggrepel

**Imports** methods

Suggests ggalt, ggrastr, RUnit, BiocGenerics, knitr, DESeq2, pasilla, airway, org.Hs.eg.db, gridExtra, magrittr, rmarkdown

URL https://github.com/kevinblighe/EnhancedVolcano

**biocViews** RNASeq, GeneExpression, Transcription, DifferentialExpression, ImmunoOncology

VignetteBuilder knitr

RoxygenNote 7.1.2

git\_url https://git.bioconductor.org/packages/EnhancedVolcano

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EnhancedVolcano *Publication-ready volcano plots with enhanced colouring and label-ing.* 

#### Description

Volcano plots represent a useful way to visualise the results of differential expression analyses. Here, we present a highly-configurable function that produces publication-ready volcano plots [@EnhancedVolcano]. EnhancedVolcano will attempt to fit as many variable names in the plot window as possible, thus avoiding 'clogging' up the plot with labels that could not otherwise have been read.

#### Usage

```
EnhancedVolcano(
  toptable,
  lab.
  х,
  у,
  selectLab = NULL,
 xlim = c(min(toptable[[x]], na.rm = TRUE) - 1.5, max(toptable[[x]], na.rm = TRUE) +
    1.5),
  ylim = c(0, max(-log10(toptable[[y]]), na.rm = TRUE) + 5),
  xlab = bquote(~Log[2] ~ "fold change"),
  ylab = bquote(~-Log[10] ~ italic(P)),
  axisLabSize = 18,
  title = "Volcano plot",
  subtitle = bquote(italic(EnhancedVolcano)),
  caption = paste0("total = ", nrow(toptable), " variables"),
  titleLabSize = 18,
  subtitleLabSize = 14,
  captionLabSize = 14,
  pCutoff = 1e-05,
  pCutoffCol = y,
  FCcutoff = 1,
  cutoffLineType = "longdash",
  cutoffLineCol = "black",
  cutoffLineWidth = 0.4,
  pointSize = 2,
  labSize = 5,
  labCol = "black",
  labFace = "plain"
  boxedLabels = FALSE,
  parseLabels = FALSE,
  shape = 19,
  shapeCustom = NULL,
```

)

```
col = c("grey30", "forestgreen", "royalblue", "red2"),
colCustom = NULL,
colAlpha = 1/2,
colGradient = NULL,
colGradientBreaks = c(pCutoff, 1),
colGradientLabels = c("0", "1.0"),
colGradientLimits = c(0, 1),
legendLabels = c("NS", expression(Log[2] ~ FC), "p-value", expression(p - value ~ and
  ~ log[2] ~ FC)),
legendPosition = "top",
legendLabSize = 14,
legendIconSize = 5,
legendDropLevels = TRUE,
encircle = NULL,
encircleCol = "black",
encircleFill = "pink",
encircleAlpha = 3/4,
encircleSize = 2.5,
shade = NULL,
shadeFill = "grey",
shadeAlpha = 1/2,
shadeSize = 0.01,
shadeBins = 2,
drawConnectors = FALSE,
widthConnectors = 0.5,
typeConnectors = "closed",
endsConnectors = "first",
lengthConnectors = unit(0.01, "npc"),
colConnectors = "grey10",
max.overlaps = 15,
maxoverlapsConnectors = NULL,
min.segment.length = 0,
directionConnectors = "both",
arrowheads = TRUE,
hline = NULL,
hlineType = "longdash",
hlineCol = "black",
hlineWidth = 0.4,
vline = NULL,
vlineType = "longdash",
vlineCol = "black",
vlineWidth = 0.4,
gridlines.major = TRUE,
gridlines.minor = TRUE,
border = "partial",
borderWidth = 0.8,
borderColour = "black",
raster = FALSE
```

## Arguments

toptable	A data-frame of test statistics (if not, a data frame, an attempt will be made to convert it to one). Requires at least the following: column for variable names (can be rownames); a column for log2 fold changes; a column for nominal or adjusted p-value.
lab	A column name in toptable containing variable names. Can be rownames(toptable).
x	A column name in toptable containing log2 fold changes.
У	A column name in toptable containing nominal or adjusted p-values.
selectLab	A vector containing a subset of lab.
xlim	Limits of the x-axis.
ylim	Limits of the y-axis.
xlab	Label for x-axis.
ylab	Label for y-axis.
axisLabSize	Size of x- and y-axis labels.
title	Plot title.
subtitle	Plot subtitle.
caption	Plot caption.
titleLabSize	Size of plot title.
subtitleLabSiz	
	Size of plot subtitle.
	Size of plot caption.
pCutoff	Cut-off for statistical significance. A horizontal line will be drawn at -log10(pCutoff).
pCutoffCol	Column name of statistical significance values to be used as the cut-off. A typ- ical usage situation would be to pass nominal [un-adjusted] p-values as 'y', but adjusted p-values as pCutoffCol. In this way, a plot is generated via - log10(unadjusted p-value), but cut-offs based on adjusted p-values.
FCcutoff	Cut-off for absolute log2 fold-change. Vertical lines will be drawn at the nega- tive and positive values of log2FCcutoff.
cutoffLineType	Line type for FCcutoff and pCutoff ('blank', 'solid', 'dashed', 'dotted', 'dot- dash', 'longdash', 'twodash').
cutoffLineCol	Line colour for FCcutoff and pCutoff.
cutoffLineWidt	
	Line width for FCcutoff and pCutoff.
pointSize	Size of plotted points for each variable. Can be a single value or a vector of sizes.
labSize	Size of labels for each variable.
labCol	Colour of labels for each variable.
labFace	Font face of labels for each variable.
boxedLabels	Logical, indicating whether or not to draw labels in boxes.
parseLabels	Logical, indicating whether or not to parse expressions in labels
shape	Shape of the plotted points. Either a single value for all points, or 4 values corresponding to the default 4 legend labels specified by legendLabels.

shapeCustom	Named vector / key-value pairs that will over-ride the default shape scheme. The order must match that of toptable. Names / keys relate to groups / categories;	
	values relate to shape encodings.	
col	Colour shading for plotted points, corresponding to the default 4 legend labels specified by legendLabels.	
colCustom	Named vector / key-value pairs that will over-ride the default colour scheme. The order must match that of toptable. Names / keys relate to groups / cate- gories; values relate to colour.	
colAlpha	Alpha for purposes of controlling colour transparency of variable points.	
colGradient	If activated, over-rides the default discrete colour scheme and replaces it with a continous scheme that shades based on nominal or adjusted p-value specified by y. For example, c('red2', 'blue2').	
colGradientBrea		
colGradientLab	Break-points for the two colours specified by colGradient.	
COIGLAUIGHTEAD	Labels for the break-points specified by colGradientBreaks.	
colGradientLim		
	Limits of the colour scheme specified by colGradient, i.e., max and min possible p-values.	
legendLabels	Plot legend text labels.	
legendPosition	Position of legend ('top', 'bottom', 'left', 'right').	
legendLabSize	Size of plot legend text.	
-	Size of plot legend icons / symbols.	
legendDropLevels		
anainala	Logical, drop unused factor levels from legend.	
encircle encircleCol	A vector of variable names to encircle. Requires installation of package ggalt. Colour of the encircled line.	
encircleCol		
	Colour fill of the encircled region.	
encircleAlpha encircleSize	Alpha for purposes of controlling colour transparency of encircled region. Line width of the encircled line.	
shade	A vector of variable names to shade.	
shadeFill		
shadeAlpha	Colour of shaded regions. Alpha for purposes of controlling colour transparency of shaded region.	
shadeSize	Size of the shade contour lines.	
shadeBins	Number of bins for the density of the shade.	
	Logical, indicating whether or not to connect plot labels to their corresponding	
widthConnectors	points by line connectors.	
withconnectors	Line width of connectors.	
typeConnectors	Have the arrow head open ('open') or filled ('closed')?	
	Which end of connectors to draw arrow head? ('last', 'first', 'both').	
lengthConnectors		
	Length (size) of the connector arrowheads.	
colConnectors	Line colour of connectors and line segments.	

max.overlaps	Equivalent of max.overlaps in ggrepel. Set to 'Inf' to always display all labels when drawConnectors = TRUE.
maxoverlapsConn	
	See max.overlaps.
min.segment.len	gth
	When drawConnectors = TRUE, specifies the minimum length of the connector line segments.
directionConnec	tors
	direction in which to draw connectors. 'both', 'x', or 'y'.
arrowheads	Logical, indicating whether or not to draw arrow heads or or just have straight lines.
hline	Draw one or more horizontal lines passing through this/these values on y-axis. For single values, only a single numerical value is necessary. For multiple lines, pass these as a vector, e.g., $c(60,90)$ .
hlineType	Line type for hline ('blank', 'solid', 'dashed', 'dotted', 'dotdash', 'longdash', 'twodash').
hlineCol	Colour of hline.
hlineWidth	Width of hline.
vline	Draw one or more vertical lines passing through this/these values on x-axis. For single values, only a single numerical value is necessary. For multiple lines, pass these as a vector, e.g., $c(60,90)$ .
vlineType	Line type for vline ('blank', 'solid', 'dashed', 'dotted', 'dotdash', 'longdash', 'twodash').
vlineCol	Colour of vline.
vlineWidth gridlines.major	Width of vline.
	Logical, indicating whether or not to draw major gridlines.
gridlines.minor	
	Logical, indicating whether or not to draw minor gridlines.
border	Add a border for just the x and y axes ('partial') or the entire plot grid ('full')?
borderWidth	Width of the border on the x and y axes.
borderColour	Colour of the border on the x and y axes.
raster	Logical, indicating whether to rasterize the geom_point layer. Requires instal- lation of ggrastr.

#### Details

Volcano plots represent a useful way to visualise the results of differential expression analyses. Here, we present a highly-configurable function that produces publication-ready volcano plots [@EnhancedVolcano]. EnhancedVolcano will attempt to fit as many variable names in the plot window as possible, thus avoiding 'clogging' up the plot with labels that could not otherwise have been read.

#### Value

A ggplot2 object.

#### Author(s)

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#### EnhancedVolcano

#### Examples

```
library('pasilla')
pasCts <- system.file('extdata', 'pasilla_gene_counts.tsv',</pre>
  package='pasilla', mustWork=TRUE)
pasAnno <- system.file('extdata', 'pasilla_sample_annotation.csv',</pre>
 package='pasilla', mustWork=TRUE)
cts <- as.matrix(read.csv(pasCts,sep='\t',row.names='gene_id'))</pre>
coldata <- read.csv(pasAnno, row.names=1)</pre>
coldata <- coldata[,c('condition','type')]</pre>
rownames(coldata) <- sub('fb', '', rownames(coldata))</pre>
cts <- cts[, rownames(coldata)]</pre>
library('DESeq2')
dds <- DESeqDataSetFromMatrix(countData = cts,</pre>
 colData = coldata,
  design = ~ condition)
featureData <- data.frame(gene=rownames(cts))</pre>
mcols(dds) <- DataFrame(mcols(dds), featureData)</pre>
dds <- DESeq(dds)
res <- results(dds)</pre>
EnhancedVolcano(res,
  lab = rownames(res),
  x = 'log2FoldChange',
  y = 'pvalue',
  pCutoff = 10e-4,
  FCcutoff = 1.333,
  xlim = c(-5.5, 5.5),
  ylim = c(0, -log10(10e-12)),
  pointSize = 1.5,
  labSize = 2.5,
  title = 'DESeq2 results',
  subtitle = 'Differential expression',
  caption = 'FC cutoff, 1.333; p-value cutoff, 10e-4',
  legendPosition = "right",
  legendLabSize = 14,
  col = c('grey30', 'forestgreen', 'royalblue', 'red2'),
  colAlpha = 0.9,
  drawConnectors = TRUE,
  hline = c(10e-8),
  widthConnectors = 0.5)
```

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