# Package 'Glimma'

November 21, 2024

```
Title Interactive visualizations for gene expression analysis
Version 2.16.0
Description This package produces interactive visualizations for RNA-seq data analysis, utilizing out-
      put from limma, edgeR, or DESeq2. It produces interactive htmlwidgets versions of popu-
      lar RNA-seq analysis plots to enhance the exploration of analysis results by overlaying interac-
     tive features. The plots can be viewed in a web browser or embedded in notebook documents.
Encoding UTF-8
RoxygenNote 7.3.2
biocViews DifferentialExpression, GeneExpression, Microarray,
     ReportWriting, RNASeq, Sequencing, Visualization
Depends R (>= 4.0.0)
Imports htmlwidgets, edgeR, DESeq2, limma, SummarizedExperiment,
     stats, isonlite, methods, S4Vectors
Suggests testthat, knitr, rmarkdown, BiocStyle, IRanges,
     GenomicRanges, pryr, AnnotationHub, scRNAseq, scater, scran
License GPL-3
URL https://github.com/hasaru-k/GlimmaV2
BugReports https://github.com/hasaru-k/GlimmaV2/issues
VignetteBuilder knitr
git_url https://git.bioconductor.org/packages/Glimma
git_branch RELEASE_3_20
git_last_commit 3bb4025
git_last_commit_date 2024-10-29
Repository Bioconductor 3.20
Date/Publication 2024-11-20
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```

Type Package

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as.hexcol

Numeric to hex colour converter

# **Description**

Convert numbers and R colour strings into corresponding hex codes for colours

### Usage

```
as.hexcol(x)
```

# Arguments

Х

the colour value(s) to be converted to hex values.

### Value

hex codes for colours

buildXYData

XY Data Object Builder

# Description

Common processing steps for both MA, XY and volcano plots. Expects a dataframe, table, which contains two columns labelled xlab and ylab as well as a unique identifier column labelled gene.

# Usage

```
buildXYData(
  table,
  status,
  main,
  display.columns,
  anno,
  counts,
  xlab,
  ylab,
  status.cols,
  sample.cols,
  groups,
  transform.counts
)
```

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### **Arguments**

table dataframe containing xlab and ylab columns for plotting.

status vector of length nrow(x) indicating the status of each gene. By default genes in

the summary plot are coloured based on its differential expression status using an adjusted p-value cutoff of 5% by calling the limma::decideTests function, where the value of -1 marks down-regulated genes, 0 marks genes with no ex-

pression difference, and 1 marks up-regulated genes.

main character string for the main title of summary plot.

display.columns

character vector containing names of columns from anno from which to display

in mouseover tooltips and table.

anno dataframe with nrow(x) rows containing gene annotations.

counts numeric matrix with nrow(x) rows containing gene expression values. This

can be used to replace the gene counts from dge\$counts, i.e. you may have

log-rpkm values stored in a different object that you wish to use.

xlab character string for the x-axis label of summary plot.
ylab character string for the y-axis label of summary plot.

status.cols vector of length 3 containing valid CSS strings for colours associated with status

in the order of -1, 0 and 1.

sample.cols character vector of length ncol(counts) containing valid CSS strings for colours

associated with each sample to be displayed on the expression plot. If left un-

specified, samples will be coloured according to groups.

groups vector of length ncol(dge) representing categorisation of samples in expression

plot.

transform.counts

the type of transformation used on the counts - "logcpm" for using edgeR::cpm(counts, log=TRUE); "cpm" for edgeR::cpm(counts); "rpkm" for edgeR::rpkm(counts); "logrpkm" for edgeR::rpkm(counts, log=TRUE); and "none" for no transfor-

mation). Defaults to "logcpm".

#### Value

object for XY plot internal use

extractGroups extractGroups

### **Description**

Extracts the column named group from column data matrix of a SummarizedExperiment object if it is present. Otherwise return a vector of 1s.

### Usage

extractGroups(cdata)

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# Arguments

cdata

SummarizedExperiment column data matrix

# Value

groups column of data if present, otherwise 1

glBar

Glimma MD Plot

# Description

Create an interactive bar plot object.

# Usage

```
glBar(x, ...)
```

# Arguments

x the data.frame containing data to plot.

... additional arguments depending on input object type.

# Value

A chart object containing the information to create an interactive bar plot.

# Author(s)

Shian Su

# See Also

```
glBar.default
```

glBar.default

Glimma Bar Plot

# **Description**

Default method for interactive bar plot.

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# Usage

```
## Default S3 method:
glBar(
  Х,
  yval,
  names.arg = rownames(x),
  ndigits = NULL,
  signif = 6,
  xlab = NULL,
  ylab = yval,
  main = NULL,
  height = 400,
  width = 500,
  colval = NULL,
  annot = yval,
  flag = NULL,
  info = NULL,
)
```

# Arguments

X	the data.frame containing data to plot.
yval	the column name for the x-axis values.
names.arg	the column name for the label on each bar.
ndigits	the number of digits after the decimal to round to in the tooltip (overrides signif).
signif	the number of significant figures to display in the tooltip.
xlab	the label on the x-axis.
ylab	the label on the y-axis.
main	the title for the plot.
height	the height of the plot (in pixels).
width	the width of the plot (in pixels).
colval	the colours for each data point.
annot	the columns to display in the tooltip.
flag	the special flag to indicate special plot.
info	additional information for plotting.
	additional arguments.

# Value

A chart object containing the information to create an interactive bar plot.

### Author(s)

Shian Su

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glimma

Glimma: interactive graphics from limma

#### **Description**

The Glimma package provides iteractive versions of plots frequently used in the limma package. Currently the MDS and MD plots have been implemented. The functions can be used with both limma, edgeR and DESeq objecs.

#### **Main functions**

```
glMDSPlot, glMDPlot, glXYPlot
```

### Author(s)

Maintainer: Shian Su <su.s@wehi.edu.au>

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- · Hasaru Kariyawasam
- · Oliver Voogd
- · Matthew Ritchie
- · Charity Law

Other contributors:

- Stuart Lee [contributor]
- Isaac Virshup [contributor]

#### See Also

Useful links:

- https://github.com/hasaru-k/GlimmaV2
- Report bugs at https://github.com/hasaru-k/GlimmaV2/issues

glimmaMA

Glimma MA Plot

### **Description**

Generic function for drawing a two-panel interactive MA plot, a special case of the glimmaXY plot. The function invokes the following methods which depend on the class of the first argument:

- glimmaMA.MArrayLM for limma analysis
- glimmaMA.DGEExact for edgeR analysis, produced from exactTest
- glimmaMA.DGELRT for edgeR analysis, produced from glmLRT
- glimmaMA.DESeqDataSet for DESeq2 analysis

glimmaMD is an alias for glimmaMA.

#### **Usage**

```
glimmaMA(x, ...)
glimmaMD(x, ...)
```

#### **Arguments**

x the DE object to plot.

additional arguments affecting the plots produced. See specific methods for detailed arguments.

#### **Details**

The summary plot on the left represents gene-wise log-fold-change (logFC) on the y-axis versus average gene expression calculated as log-counts-per-million (logCPM) values. We call our summary plot an MA plot because this type of plot was originally referred to as an MA plot in the limma package, with the M-value representing logFC and A-value representing average expression - it has since been renamed to MD plot in the limma package. The expression plot on the right displays sample expression values for a single gene. Interactions with the htmlwidget include clicking on genes (points) in the summary plot to bring up associated sample expression values in the expression plot, as well as the summary statistics in the table below. Alternatively, users can interact with the table by clicking on genes (rows) to highlight genes in the summary plot, as well as bring up associated sample expression values in the expression plot. Briefly, other interactive features include a search box for the table, buttons to save plots and data (summary statistics and expression values), additional pop-up information when hovering on points in plots, and rescaling of the y-axis in the expression plot.

#### Value

htmlwidget object or NULL if html argument is specified.

### Author(s)

Hasaru Kariyawasam, Shian Su and Oliver Voogd

#### **Examples**

```
methods(glimmaMA) # show methods for glimmaMA
```

```
glimmaMA.DESeqDataSet Glimma MA Plot
```

### **Description**

Draws a two-panel interactive MA plot from an DESeqDataSet object. This is a special case of the glimmaXY plot.

#### **Usage**

```
## S3 method for class 'DESeqDataSet'
glimmaMA(
  Х,
  counts = DESeq2::counts(x),
  groups = extractGroups(colData(x)),
  status = NULL,
  anno = NULL,
  display.columns = NULL,
  status.cols = c("#1052bd", "silver", "#cc212f"),
  sample.cols = NULL,
  transform.counts = c("logcpm", "cpm", "rpkm", "logrpkm", "none"),
  main = "MA Plot",
  xlab = "logCPM",
  ylab = "logFC",
  html = NULL,
  width = 920,
  height = 920,
)
```

#### **Arguments**

x DESeqDataSet object from which summary statistics are extracted from to cre-

ate summary (left) plot.

counts numeric matrix with nrow(x) rows containing gene expression values.

groups vector/factor representing the experimental group for each sample; see extractGroups

for default value.

status vector of length nrow(x) indicating the status of each gene.

anno dataframe with nrow(x) rows containing gene annotations.

display.columns

character vector containing names of columns from anno from which to display

in mouseover tooltips and table.

status.cols vector of length 3 containing valid CSS strings for colours associated with status

in the order of -1, 0 and 1.

sample.cols character vector of length ncol(counts) containing valid CSS strings for colours

associated with each sample to be displayed on the expression plot. If left un-

specified, samples will be coloured according to groups.

transform.counts

the type of transformation used on the counts - "logcpm" for using edgeR::cpm(counts, log=TRUE); "cpm" for edgeR::cpm(counts); "rpkm" for edgeR::rpkm(counts); "logrpkm" for edgeR::rpkm(counts, log=TRUE); and "none" for no transfor-

mation). Defaults to "logcpm".

main character string for the main title of summary plot.

xlab character string for the x-axis label of summary plot.

ylab character string for the y-axis label of summary plot.

html character string for naming HTML file for exportation of widget. The extension

should be included in the file name e.g. "file.html".

width numeric value indicating width of widget in pixels.

height numeric value indicating width of height in pixels.

additional unused arguments.

#### **Details**

The summary plot on the left represents gene-wise log-fold-change (logFC) on the y-axis versus average gene expression calculated as log-counts-per-million (logCPM) values. We call our summary plot an MA plot because this type of plot was originally referred to as an MA plot in the limma package, with the M-value representing logFC and A-value representing average expression - it has since been renamed to MD plot in the limma package. The expression plot on the right displays sample expression values for a single gene. Interactions with the htmlwidget include clicking on genes (points) in the summary plot to bring up associated sample expression values in the expression plot, as well as the summary statistics in the table below. Alternatively, users can interact with the table by clicking on genes (rows) to highlight genes in the summary plot, as well as bring up associated sample expression values in the expression plot. Briefly, other interactive features include a search box for the table, buttons to save plots and data (summary statistics and expression values), additional pop-up information when hovering on points in plots, and rescaling of the y-axis in the expression plot.

#### Value

htmlwidget object or NULL if html argument is specified.

#### Author(s)

Hasaru Kariyawasam, Shian Su and Oliver Voogd

# See Also

```
glimmaMA, glimmaMA.MArrayLM, glimmaMA.DGEExact, glimmaMA.DGELRT
```

# **Examples**

```
dge <- readRDS(
   system.file("RNAseq123/dge.rds", package = "Glimma"))

dds <- DESeq2::DESeqDataSetFromMatrix(
   countData = dge$counts,
   colData = dge$samples,
   rowData = dge$genes,
   design = ~group
)

dds <- DESeq2::DESeq(dds, quiet=TRUE)
glimmaMA(dds)</pre>
```

glimmaMA.DGEExact

Glimma MA Plot

#### **Description**

Draws a two-panel interactive MA plot from an DGEExact object. This is a special case of the glimmaXY plot.

#### Usage

```
## S3 method for class 'DGEExact'
glimmaMA(
  х,
  dge = NULL,
  counts = dge$counts,
  groups = dge$samples$group,
  status = edgeR::decideTests.DGEExact(x),
  anno = x$genes,
  display.columns = NULL,
  status.cols = c("#1052bd", "silver", "#cc212f"),
  sample.cols = NULL,
  p.adj.method = "BH",
  transform.counts = c("logcpm", "cpm", "rpkm", "logrpkm", "none"),
  main = paste(x$comparison[2], "vs", x$comparison[1]),
  xlab = "logCPM",
  ylab = "logFC",
  html = NULL,
  width = 920,
  height = 920,
)
```

#### **Arguments**

dge

counts

status

X	DGEExact object from which summary statistics are extracted from to create
	summary (left) plot.

DGEList object with nrow(x) rows from which expression values are extracted from to create expression (right) plot. Gene counts are taken from dge\$counts and sample groups from dge\$samples\$group. By default raw counts are trans-

formed to log-cpm values (see more in the transform.counts argument).

numeric matrix with nrow(x) rows containing gene expression values. This can be used to replace the gene counts from dge\$counts, i.e. you may have log-rpkm values stored in a different object that you wish to use.

groups vector of length ncol(dge) representing categorisation of samples in expression

plot.

vector of length nrow(x) indicating the status of each gene. By default genes in the summary plot are coloured based on its differential expression status using an adjusted p-value cutoff of 0.05 by calling the edgeR::decideTests() function, where the value of -1 marks down-regulated genes, 0 marks genes with no expression difference, and 1 marks up-regulated genes.

anno dataframe with nrow(x) rows containing gene annotations.

display.columns

character vector containing names of columns from anno from which to display

in mouseover tooltips and table.

status.cols vector of length 3 containing valid CSS strings for colours associated with status

in the order of -1, 0 and 1.

sample.cols character vector of length ncol(counts) containing valid CSS strings for colours

associated with each sample to be displayed on the expression plot. If left un-

specified, samples will be coloured according to groups.

p.adj.method character string specifying p-value adjustment method.

transform.counts

the type of transformation used on the counts - "logcpm" for using edgeR::cpm(counts, log=TRUE); "cpm" for edgeR::cpm(counts); "rpkm" for edgeR::rpkm(counts); "logrpkm" for edgeR::rpkm(counts, log=TRUE); and "none" for no transfor-

mation). Defaults to "logcpm".

main character string for the main title of summary plot.

xlab character string for the x-axis label of summary plot.

ylab character string for the y-axis label of summary plot.

html character string for naming HTML file for exportation of widget. The extension

should be included in the file name e.g. "file.html".

width numeric value indicating width of widget in pixels. height numeric value indicating width of height in pixels.

... additional unused arguments.

### **Details**

The summary plot on the left represents gene-wise log-fold-change (logFC) on the y-axis versus average gene expression calculated as log-counts-per-million (logCPM) values. We call our summary plot an MA plot because this type of plot was originally referred to as an MA plot in the limma package, with the M-value representing logFC and A-value representing average expression - it has since been renamed to MD plot in the limma package. The expression plot on the right displays sample expression values for a single gene. Interactions with the htmlwidget include clicking on genes (points) in the summary plot to bring up associated sample expression values in the expression plot, as well as the summary statistics in the table below. Alternatively, users can interact with the table by clicking on genes (rows) to highlight genes in the summary plot, as well as bring up associated sample expression values in the expression plot. Briefly, other interactive features include a search box for the table, buttons to save plots and data (summary statistics and expression values), additional pop-up information when hovering on points in plots, and rescaling of the y-axis in the expression plot.

# Value

htmlwidget object or NULL if html argument is specified.

#### Author(s)

Hasaru Kariyawasam, Shian Su and Oliver Voogd

# See Also

glimmaMA, glimmaMA.MArrayLM, glimmaMA.DGELRT, glimmaMA.DESeqDataSet

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#### **Examples**

```
dge <- readRDS(
   system.file("RNAseq123/dge.rds", package = "Glimma"))
design <- readRDS(
   system.file("RNAseq123/design.rds", package = "Glimma"))
contr.matrix <- readRDS(
   system.file("RNAseq123/contr.matrix.rds", package = "Glimma"))
dge <- edgeR::estimateDisp(dge, design)
gfit <- edgeR::glmFit(dge, design)
glrt <- edgeR::glmLRT(gfit, design, contrast = contr.matrix)
glimmaMA(glrt, dge = dge)</pre>
```

glimmaMA.DGELRT

Glimma MA Plot

#### **Description**

Draws a two-panel interactive MA plot from an DGELRT object. This is a special case of the glimmaXY plot.

### Usage

```
## S3 method for class 'DGELRT'
glimmaMA(
  Х,
  dge = NULL,
  counts = dge$counts,
  groups = dge$samples$group,
  status = edgeR::decideTests.DGEExact(x),
  anno = x$genes,
  display.columns = NULL,
  status.cols = c("#1052bd", "silver", "#cc212f"),
  sample.cols = NULL,
  p.adj.method = "BH",
  transform.counts = c("logcpm", "cpm", "rpkm", "logrpkm", "none"),
  main = paste(x$comparison[2], "vs", x$comparison[1]),
  xlab = "logCPM",
  ylab = "logFC",
  html = NULL,
  width = 920.
  height = 920,
)
```

#### **Arguments**

x DGELRT object from which summary statistics are extracted from to create summary (left) plot.

dge DGEList object with nrow(x) rows from which expression values are extracted from to create expression (right) plot. Gene counts are taken from dge\$counts and sample groups from dge\$samples\$group. By default raw counts are transformed to log-cpm values (see more in the transform.counts argument).

counts numeric matrix with nrow(x) rows containing gene expression values. This

can be used to replace the gene counts from dge\$counts, i.e. you may have

log-rpkm values stored in a different object that you wish to use.

groups vector of length ncol(dge) representing categorisation of samples in expression

plot.

status vector of length nrow(x) indicating the status of each gene. By default genes in

the summary plot are coloured based on its differential expression status using an adjusted p-value cutoff of 0.05 by calling the edgeR::decideTests() function, where the value of -1 marks down-regulated genes, 0 marks genes with no

expression difference, and 1 marks up-regulated genes.

anno dataframe with nrow(x) rows containing gene annotations.

display.columns

character vector containing names of columns from anno from which to display

in mouseover tooltips and table.

status.cols vector of length 3 containing valid CSS strings for colours associated with status

in the order of -1, 0 and 1.

sample.cols character vector of length ncol(counts) containing valid CSS strings for colours

associated with each sample to be displayed on the expression plot. If left un-

specified, samples will be coloured according to groups.

p.adj.method character string specifying p-value adjustment method.

 ${\it transform.} \, {\it counts} \,$ 

the type of transformation used on the counts - "logcpm" for using edgeR::cpm(counts, log=TRUE); "cpm" for edgeR::cpm(counts); "rpkm" for edgeR::rpkm(counts); "logrpkm" for edgeR::rpkm(counts, log=TRUE); and "none" for no transfor-

mation). Defaults to "logcpm".

main character string for the main title of summary plot.

xlab character string for the x-axis label of summary plot.

ylab character string for the y-axis label of summary plot.

html character string for naming HTML file for exportation of widget. The extension

should be included in the file name e.g. "file.html".

width numeric value indicating width of widget in pixels.
height numeric value indicating width of height in pixels.

... additional unused arguments.

#### **Details**

The summary plot on the left represents gene-wise log-fold-change (logFC) on the y-axis versus average gene expression calculated as log-counts-per-million (logCPM) values. We call our summary plot an MA plot because this type of plot was originally referred to as an MA plot in the limma package, with the M-value representing logFC and A-value representing average expression - it has since been renamed to MD plot in the limma package. The expression plot on the right displays sample expression values for a single gene. Interactions with the htmlwidget include clicking on genes (points) in the summary plot to bring up associated sample expression values in the expression plot, as well as the summary statistics in the table below. Alternatively, users can interact with

the table by clicking on genes (rows) to highlight genes in the summary plot, as well as bring up associated sample expression values in the expression plot. Briefly, other interactive features include a search box for the table, buttons to save plots and data (summary statistics and expression values), additional pop-up information when hovering on points in plots, and rescaling of the y-axis in the expression plot.

#### Value

htmlwidget object or NULL if html argument is specified.

#### Author(s)

Hasaru Kariyawasam, Shian Su and Oliver Voogd

#### See Also

```
glimmaMA, glimmaMA.MArrayLM, glimmaMA.DGEExact, glimmaMA.DESeqDataSet
```

glimmaMA.MArrayLM

Glimma MA Plot

### **Description**

Draws a two-panel interactive MA plot from an MArrayLM object. This is a special case of the glimmaXY plot.

# Usage

```
## S3 method for class 'MArrayLM'
glimmaMA(
  Х,
  dge = NULL,
  counts = dge$counts,
  groups = dge$samples$group,
  coef = ncol(x$coefficients),
  status = limma::decideTests(x),
  anno = x$genes,
  display.columns = NULL,
  status.cols = c("#1052bd", "silver", "#cc212f"),
  sample.cols = NULL,
  p.adj.method = "BH",
  transform.counts = c("logcpm", "cpm", "rpkm", "logrpkm", "none"),
  main = colnames(x)[coef],
  xlab = "logCPM",
  ylab = "logFC",
  html = NULL,
  width = 920,
  height = 920,
)
```

#### **Arguments**

x MArrayLM object from which summary statistics are extracted from to create

summary (left) plot.

dge DGEList object with nrow(x) rows from which expression values are extracted

from to create expression (right) plot. Gene counts are taken from dge\$counts and sample groups from dge\$samples\$group. By default raw counts are transformed to log-cpm values (see more in the transform.counts argument).

counts numeric matrix with nrow(x) rows containing gene expression values. This

can be used to replace the gene counts from dge\$counts, i.e. you may have

log-rpkm values stored in a different object that you wish to use.

groups vector of length ncol(dge) representing categorisation of samples in expression

plot.

coef integer indicating the column in x from the summary plot is created.

status vector of length nrow(x) indicating the status of each gene. By default genes in

the summary plot are coloured based on its differential expression status using an adjusted p-value cutoff of 5% by calling the limma::decideTests function, where the value of -1 marks down-regulated genes, 0 marks genes with no ex-

pression difference, and 1 marks up-regulated genes.

anno dataframe with nrow(x) rows containing gene annotations.

display.columns

character vector containing names of columns from anno from which to display

in mouseover tooltips and table.

status.cols vector of length 3 containing valid CSS strings for colours associated with status

in the order of -1, 0 and 1.

sample.cols character vector of length ncol(counts) containing valid CSS strings for colours

associated with each sample to be displayed on the expression plot. If left un-

specified, samples will be coloured according to groups.

p.adj.method character string specifying p-value adjustment method.

transform.counts

the type of transformation used on the counts - "logcpm" for using edgeR::cpm(counts, log=TRUE); "cpm" for edgeR::cpm(counts); "rpkm" for edgeR::rpkm(counts); "logrpkm" for edgeR::rpkm(counts, log=TRUE); and "none" for no transfor-

mation). Defaults to "logcpm".

main character string for the main title of summary plot.

xlab character string for the x-axis label of summary plot.
ylab character string for the y-axis label of summary plot.

html character string for naming HTML file for exportation of widget. The extension

should be included in the file name e.g. "file.html".

width numeric value indicating width of widget in pixels.

height numeric value indicating width of height in pixels.

... additional unused arguments.

### Details

The summary plot on the left represents gene-wise log-fold-change (logFC) on the y-axis versus average gene expression calculated as log-counts-per-million (logCPM) values. We call our summary plot an MA plot because this type of plot was originally referred to as an MA plot in the limma

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package, with the M-value representing logFC and A-value representing average expression - it has since been renamed to MD plot in the 1imma package. The expression plot on the right displays sample expression values for a single gene. Interactions with the htmlwidget include clicking on genes (points) in the summary plot to bring up associated sample expression values in the expression plot, as well as the summary statistics in the table below. Alternatively, users can interact with the table by clicking on genes (rows) to highlight genes in the summary plot, as well as bring up associated sample expression values in the expression plot. Briefly, other interactive features include a search box for the table, buttons to save plots and data (summary statistics and expression values), additional pop-up information when hovering on points in plots, and rescaling of the y-axis in the expression plot.

#### Value

htmlwidget object or NULL if html argument is specified.

#### Author(s)

Hasaru Kariyawasam, Shian Su and Oliver Voogd

#### See Also

```
glimmaMA, glimmaMA.DGEExact, glimmaMA.DGELRT, glimmaMA.DESeqDataSet
```

# **Examples**

```
dge <- readRDS(
   system.file("RNAseq123/dge.rds", package = "Glimma"))
design <- readRDS(
   system.file("RNAseq123/design.rds", package = "Glimma"))
contr.matrix <- readRDS(
   system.file("RNAseq123/contr.matrix.rds", package = "Glimma"))
v <- limma::voom(dge, design)
vfit <- limma::lmFit(v, design)
vfit <- limma::contrasts.fit(vfit, contrasts = contr.matrix)
efit <- limma::eBayes(vfit)
glimmaMA(efit, dge = dge)</pre>
```

glimmaMDS

Glimma MDS Plot

#### **Description**

Generic function for drawing a two-panel interactive multidimensional scaling (MDS) plot. The function invokes the following methods which depend on the class of the first argument:

- glimmaMDS.DGEList for edgeR analysis
- glimmaMDS.DESeqDataSet for DESeq2 analysis
- glimmaMDS.default for all other object types

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### Usage

```
glimmaMDS(x, ...)
```

#### **Arguments**

x the matrix containing the gene expressions.

the additional arguments affecting the plot produced. See specific methods for detailed arguments.

#### **Details**

The left plot shows two MDS dimensions, with sample annotations displayed on hover. The right panel contains a bar plot of the eigenvalues of each dimension. The controls beneath the plots can be used to change the dimensions being displayed, and the scale, colour and shape of points. The interactive MDS plot allows users to adjust sample points by scale, colour and shape for multiple vectors associated with sample information. This is carried out most effectively when x\$samples includes an abundance of sample information, or when a data frame object is supplied to groups. If a simple character or factor vector is given to groups (with the default of continous.colour=FALSE), then sample points will have no scaling options, but can only be adjusted in colour and shape by groups and labels. Instead, if groups is a numeric vector (e.g. library size or expression level of a specific gene), then the plot can be scaled and coloured by the numeric values with continous.colour=TRUE. For more details, refer to limma::plotMDS.

#### Value

htmlwidget object or NULL if html argument is specified.

#### Author(s)

Hasaru Kariyawasam, Shian Su and Oliver Voogd

### **Examples**

```
dge <- readRDS(system.file("RNAseq123/dge.rds", package = "Glimma"))
glimmaMDS(dge)

# using DESeqDataSet
dds <- DESeq2::DESeqDataSetFromMatrix(
    countData = dge$counts,
    colData = dge$samples,
    rowData = dge$genes,
    design = ~group
)
glimmaMDS(dds)

# using matrix object
expr <- edgeR::cpm(dge, log = TRUE)
glimmaMDS(expr)</pre>
```

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glimmaMDS.default Glimma MDS Plot

#### **Description**

Draws a two-panel interactive MDS plot.

#### Usage

```
## Default S3 method:
glimmaMDS(
    x,
    groups = as.character(rep(1, ncol(x))),
    labels = as.character(seq_len(ncol(x))),
    continuous.colour = FALSE,
    top = 500,
    gene.selection = c("pairwise", "common"),
    html = NULL,
    width = 900,
    height = 500,
    ...
)
```

# **Arguments**

the matrix containing the gene expressions.

groups vector or data frame object with associated sample information such as experi-

mental groups. The information is displayed in mouseover tooltips, and appropriate vector(s) can be used to adjust the plot using scale\_by, colour\_by and

shape\_by drop-down boxes of the widget.

labels character vector of sample names or labels.

continuous.colour

TRUE if continuous colour schemes should be used. Defaults to FALSE where

distinct colour schemes are used.

top integer indiating number of top genes used to calculate pairwise distances.

gene.selection character string specifying how genes are selected from the plot - "pairwise" if

most variable genes are to be chosen for each pair of samples, or "common" to

select the same genes for all comparisons.

html character string for naming HTML file or exportation of widget. The extension

should be included in the file name e.g. "file.hml".

width numeric value indicating width of widget in pixels.

height numeric value indicating width of widget in pixels.

... additional unused arguments.

#### **Details**

The left plot shows two MDS dimensions, with sample annotations displayed on hover. The right panel contains a bar plot of the eigenvalues of each dimension. The controls beneath the plots can be used to change the dimensions being displayed, and the scale, colour and shape of points. The interactive MDS plot allows users to adjust sample points by scale, colour and shape for multiple vectors associated with sample information. This is carried out most effectively when x\$samples includes an abundance of sample information, or when a data frame object is supplied to groups. If a simple character or factor vector is given to groups (with the default of continous.colour=FALSE), then sample points will have no scaling options, but can only be adjusted in colour and shape by groups and labels. Instead, if groups is a numeric vector (e.g. library size or expression level of a specific gene), then the plot can be scaled and coloured by the numeric values with continous.colour=TRUE. For more details, refer to limma::plotMDS.

#### Value

htmlwidget object or NULL if html argument is specified.

#### Author(s)

Hasaru Kariyawasam, Shian Su and Oliver Voogd

#### See Also

```
glimmaMDS, glimmaMDS.DGEList, glimmaMDS.DESeqDataSet
```

#### **Examples**

```
dge <- readRDS(system.file("RNAseq123/dge.rds", package = "Glimma"))
expr <- edgeR::cpm(dge, log = TRUE)
glimmaMDS(expr)</pre>
```

```
{\tt glimmaMDS.DESeqDataSet}
```

Glimma MDS Plot

### **Description**

Draws a two-panel interactive MDS plot using a DESeqDataset x. Transforms counts using edgeR::cpm(DESeq2::count log = TRUE, prior.count = prior.count).

#### **Usage**

```
## S3 method for class 'DESeqDataSet'
glimmaMDS(
    x,
    groups = as.data.frame(SummarizedExperiment::colData(x)),
    labels = rownames(SummarizedExperiment::colData(x)),
    continuous.colour = FALSE,
    top = 500,
    gene.selection = c("pairwise", "common"),
    prior.count = 2,
```

```
html = NULL,
width = 900,
height = 500,
...
)
```

#### **Arguments**

x DESeqDataSet object containing gene counts.

groups vector or data frame object with associated sample information such as experi-

mental groups. The information is displayed in mouseover tooltips, and appropriate vector(s) can be used to adjust the plot using scale\_by, colour\_by and

shape\_by drop-down boxes of the widget.

labels character vector of sample names or labels.

continuous.colour

TRUE if continuous colour schemes should be used. Defaults to FALSE where

distinct colour schemes are used.

top integer indiating number of top genes used to calculate pairwise distances.

gene. selection character string specifying how genes are selected from the plot - "pairwise" if

most variable genes are to be chosen for each pair of samples, or "common" to

select the same genes for all comparisons.

prior.count integer indicating the average count to be added to each observation to avoid

taking log of zero when raw counts are transformed to log-counts-per-million

values (using edgeR::cpm function).

html character string for naming HTML file or exportation of widget. The extension

should be included in the file name e.g. "file.hml".

width numeric value indicating width of widget in pixels.
height numeric value indicating width of widget in pixels.

... additional unused arguments.

#### **Details**

The left plot shows two MDS dimensions, with sample annotations displayed on hover. The right panel contains a bar plot of the eigenvalues of each dimension. The controls beneath the plots can be used to change the dimensions being displayed, and the scale, colour and shape of points. The interactive MDS plot allows users to adjust sample points by scale, colour and shape for multiple vectors associated with sample information. This is carried out most effectively when x\$samples includes an abundance of sample information, or when a data frame object is supplied to groups. If a simple character or factor vector is given to groups (with the default of continous.colour=FALSE), then sample points will have no scaling options, but can only be adjusted in colour and shape by groups and labels. Instead, if groups is a numeric vector (e.g. library size or expression level of a specific gene), then the plot can be scaled and coloured by the numeric values with continous.colour=TRUE. For more details, refer to limma::plotMDS.

#### Value

htmlwidget object or NULL if html argument is specified.

#### Author(s)

Hasaru Kariyawasam, Shian Su and Oliver Voogd

#### See Also

```
glimmaMDS, glimmaMDS.default, glimmaMDS.DGEList
```

#### **Examples**

```
dge <- readRDS(system.file("RNAseq123/dge.rds", package = "Glimma"))
dds <- DESeq2::DESeqDataSetFromMatrix(
  countData = dge$counts,
  colData = dge$samples,
  rowData = dge$genes,
  design = ~group
)
glimmaMDS(dds)</pre>
```

glimmaMDS.DGEList

Glimma MDS Plot

### **Description**

Draws a two-panel interactive MDS plot using a DGEList x. Transforms counts using edgeR::cpm(x, log=TRUE, prior.count = prior.count).

### Usage

```
## S3 method for class 'DGEList'
glimmaMDS(
    x,
    groups = x$samples,
    labels = rownames(x$samples),
    continuous.colour = FALSE,
    top = 500,
    gene.selection = c("pairwise", "common"),
    prior.count = 2,
    html = NULL,
    width = 900,
    height = 500,
    ...
)
```

### **Arguments**

x DGEList object containing gene counts in x\$counts.

groups

vector or data frame object with associated sample information such as experimental groups. The information is displayed in mouseover tooltips, and appropriate vector(s) can be used to adjust the plot using scale\_by, colour\_by and shape\_by drop-down boxes of the widget.

labels character vector of sample names or labels.

continuous.colour

TRUE if continuous colour schemes should be used. Defaults to FALSE where distinct colour schemes are used.

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top integer indiating number of top genes used to calculate pairwise distances.

gene.selection character string specifying how genes are selected from the plot - "pairwise" if

most variable genes are to be chosen for each pair of samples, or "common" to

select the same genes for all comparisons.

prior.count integer indicating the average count to be added to each observation to avoid

taking log of zero when raw counts are transformed to log-counts-per-million

values (using edgeR::cpm function).

html character string for naming HTML file or exportation of widget. The extension

should be included in the file name e.g. "file.hml".

width numeric value indicating width of widget in pixels.

height numeric value indicating width of widget in pixels.

... additional unused arguments.

#### **Details**

The left plot shows two MDS dimensions, with sample annotations displayed on hover. The right panel contains a bar plot of the eigenvalues of each dimension. The controls beneath the plots can be used to change the dimensions being displayed, and the scale, colour and shape of points. The interactive MDS plot allows users to adjust sample points by scale, colour and shape for multiple vectors associated with sample information. This is carried out most effectively when x\$samples includes an abundance of sample information, or when a data frame object is supplied to groups. If a simple character or factor vector is given to groups (with the default of continous.colour=FALSE), then sample points will have no scaling options, but can only be adjusted in colour and shape by groups and labels. Instead, if groups is a numeric vector (e.g. library size or expression level of a specific gene), then the plot can be scaled and coloured by the numeric values with continous.colour=TRUE. For more details, refer to limma::plotMDS.

### Value

htmlwidget object or NULL if html argument is specified.

## Author(s)

Hasaru Kariyawasam, Shian Su and Oliver Voogd

#### See Also

```
glimmaMDS, glimmaMDS.default, glimmaMDS.DESeqDataSet
```

#### **Examples**

```
dge <- readRDS(system.file("RNAseq123/dge.rds", package = "Glimma"))
glimmaMDS(dge)</pre>
```

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glimmaVolcano

Glimma Volcano Plot

#### **Description**

Generic function for drawing a two-panel interactive volcano plot, a special case of the glimmaXY plot. The function invokes the following methods which depend on the class of the first argument:

- glimmaVolcano.MArrayLM for limma analysis
- glimmaVolcano.DGEExact for edgeR analysis, produced from exactTest
- glimmaVolcano.DGELRT for edgeR analysis, produced from glmLRT
- glimmaVolcano.DESeqDataSet for DESeq2 analysis

# Usage

```
glimmaVolcano(x, ...)
```

# **Arguments**

x the DE object to plot.

... additional arguments affecting the plots produced. See specific methods for detailed arguments.

#### **Details**

The summary plot on the left represents gene-wise log-fold-change (logFC) on the x-axis versus -log10(pvalue). The expression plot on the right displays sample expression values for a single gene. Interactions with the htmlwidget include clicking on genes (points) in the summary plot to bring up associated sample expression values in the expression plot, as well as the summary statistics in the table below. Alternatively, users can interact with the table by clicking on genes (rows) to highlight genes in the summary plot, as well as bring up associated sample expression values in the expression plot. Briefly, other interactive features include a search box for the table, buttons to save plots and data (summary statistics and expression values), additional pop-up information when hovering on points in plots, and rescaling of the y-axis in the expression plot.

### Value

htmlwidget object or NULL if html argument is specified.

# Author(s)

Hasaru Kariyawasam, Shian Su and Oliver Voogd

# **Examples**

```
dge <- readRDS(
   system.file("RNAseq123/dge.rds", package = "Glimma"))
design <- readRDS(
   system.file("RNAseq123/design.rds", package = "Glimma"))
contr.matrix <- readRDS(
   system.file("RNAseq123/contr.matrix.rds", package = "Glimma"))</pre>
```

```
v <- limma::voom(dge, design)
vfit <- limma::lmFit(v, design)
vfit <- limma::contrasts.fit(vfit, contrasts = contr.matrix)
efit <- limma::eBayes(vfit)
glimmaVolcano(efit, dge = dge)</pre>
```

```
glimmaVolcano.DESeqDataSet
```

Glimma Volcano Plot

### **Description**

Draws a two-panel interactive volcano plot from an DESeqDataSet object. This is a special case of the glimmaXY plot.

### Usage

```
## S3 method for class 'DESeqDataSet'
glimmaVolcano(
  х,
  counts = DESeq2::counts(x),
  groups = extractGroups(colData(x)),
  status = NULL,
  anno = NULL,
  display.columns = NULL,
  status.cols = c("#1052bd", "silver", "#cc212f"),
  sample.cols = NULL,
  transform.counts = c("logcpm", "cpm", "rpkm", "none"),
  main = "Volcano Plot",
  xlab = "logFC",
  ylab = "negLog10PValue",
  html = NULL,
  width = 920,
  height = 920,
)
```

### **Arguments**

X	DESeqDataSet object from which summary statistics are extracted from to create summary (left) plot.
counts	numeric matrix with nrow(x) rows containing gene expression values.
groups	$vector/factor\ representing\ the\ experimental\ group\ for\ each\ sample;\ see\ extract{\sf Groups}$ for default value.
status	vector of length nrow(x) indicating the status of each gene.
anno	dataframe with nrow(x) rows containing gene annotations.

display.columns

character vector containing names of columns from anno from which to display

in mouseover tooltips and table.

status.cols vector of length 3 containing valid CSS strings for colours associated with status

in the order of -1, 0 and 1.

sample.cols character vector of length ncol(counts) containing valid CSS strings for colours

associated with each sample to be displayed on the expression plot. If left un-

specified, samples will be coloured according to groups.

transform.counts

the type of transformation used on the counts - "logcpm" for using edgeR::cpm(counts, log=TRUE); "cpm" for edgeR::cpm(counts); "rpkm" for edgeR::rpkm(counts);

"logrpkm" for edgeR::rpkm(counts, log=TRUE); and "none" for no transfor-

mation). Defaults to "logcpm".

main character string for the main title of summary plot.

xlab character string for the x-axis label of summary plot.

ylab character string for the y-axis label of summary plot.

html character string for naming HTML file for exportation of widget. The extension

should be included in the file name e.g. "file.html".

width numeric value indicating width of widget in pixels.

height numeric value indicating width of height in pixels.

... additional unused arguments.

### Details

The summary plot on the left represents gene-wise log-fold-change (logFC) on the x-axis versus -log10(pvalue). The expression plot on the right displays sample expression values for a single gene. Interactions with the htmlwidget include clicking on genes (points) in the summary plot to bring up associated sample expression values in the expression plot, as well as the summary statistics in the table below. Alternatively, users can interact with the table by clicking on genes (rows) to highlight genes in the summary plot, as well as bring up associated sample expression values in the expression plot. Briefly, other interactive features include a search box for the table, buttons to save plots and data (summary statistics and expression values), additional pop-up information when hovering on points in plots, and rescaling of the y-axis in the expression plot.

#### Value

htmlwidget object or NULL if html argument is specified.

### Author(s)

Hasaru Kariyawasam, Shian Su and Oliver Voogd

#### See Also

glimmaVolcano, glimmaVolcano.MArrayLM, glimmaVolcano.DGEExact, glimmaVolcano.DGELRT

#### **Examples**

```
dge <- readRDS(
   system.file("RNAseq123/dge.rds", package = "Glimma"))

dds <- DESeq2::DESeqDataSetFromMatrix(
   countData = dge$counts,
   colData = dge$samples,
   rowData = dge$genes,
   design = ~group
)

dds <- DESeq2::DESeq(dds, quiet=TRUE)
glimmaVolcano(dds)</pre>
```

glimmaVolcano.DGEExact

Glimma Volcano Plot

#### **Description**

Draws a two-panel interactive volcano plot from an DGEExact object. This is a special case of the glimmaXY plot.

### Usage

```
## S3 method for class 'DGEExact'
glimmaVolcano(
  х,
  dge = NULL,
  counts = dge$counts,
  groups = dge$samples$group,
  status = edgeR::decideTests.DGEExact(x),
  anno = x$genes,
  display.columns = NULL,
  status.cols = c("#1052bd", "silver", "#cc212f"),
  sample.cols = NULL,
  p.adj.method = "BH",
  transform.counts = c("logcpm", "cpm", "rpkm", "none"),
  main = paste(x$comparison[2], "vs", x$comparison[1]),
  xlab = "logFC",
  ylab = "negLog10PValue",
  html = NULL,
  width = 920,
  height = 920,
)
```

#### **Arguments**

x DGEExact object from which summary statistics are extracted from to create summary (left) plot.

dge DGEList object with nrow(x) rows from which expression values are extracted

from to create expression (right) plot. Gene counts are taken from dge\$counts and sample groups from dge\$samples\$group. By default raw counts are transformed to log-cpm values (see more in the transform.counts argument).

counts numeric matrix with nrow(x) rows containing gene expression values. This

can be used to replace the gene counts from dge\$counts, i.e. you may have

log-rpkm values stored in a different object that you wish to use.

groups vector of length ncol(dge) representing categorisation of samples in expression

plot.

status vector of length nrow(x) indicating the status of each gene. By default genes in

the summary plot are coloured based on its differential expression status using an adjusted p-value cutoff of 0.05 by calling the edgeR::decideTests() function, where the value of -1 marks down-regulated genes, 0 marks genes with no

expression difference, and 1 marks up-regulated genes.

anno dataframe with nrow(x) rows containing gene annotations.

display.columns

character vector containing names of columns from anno from which to display

in mouseover tooltips and table.

status.cols vector of length 3 containing valid CSS strings for colours associated with status

in the order of -1, 0 and 1.

sample.cols character vector of length ncol (counts) containing valid CSS strings for colours

associated with each sample to be displayed on the expression plot. If left un-

specified, samples will be coloured according to groups.

p.adj.method character string specifying p-value adjustment method.

transform.counts

the type of transformation used on the counts - "logcpm" for using edgeR::cpm(counts,

log=TRUE); "cpm" for edgeR::cpm(counts); "rpkm" for edgeR::rpkm(counts);
"logrpkm" for edgeR::rpkm(counts, log=TRUE); and "none" for no transfor-

mation). Defaults to "logcpm".

main character string for the main title of summary plot.

xlab character string for the x-axis label of summary plot.

ylab character string for the y-axis label of summary plot.

html character string for naming HTML file for exportation of widget. The extension

should be included in the file name e.g. "file.html".

width numeric value indicating width of widget in pixels. height numeric value indicating width of height in pixels.

... additional unused arguments.

#### **Details**

The summary plot on the left represents gene-wise log-fold-change (logFC) on the x-axis versus -log10(pvalue). The expression plot on the right displays sample expression values for a single gene. Interactions with the htmlwidget include clicking on genes (points) in the summary plot to bring up associated sample expression values in the expression plot, as well as the summary statistics in the table below. Alternatively, users can interact with the table by clicking on genes (rows) to highlight genes in the summary plot, as well as bring up associated sample expression values in the expression plot. Briefly, other interactive features include a search box for the table, buttons to save plots and data (summary statistics and expression values), additional pop-up information when hovering on points in plots, and rescaling of the y-axis in the expression plot.

#### Value

htmlwidget object or NULL if html argument is specified.

#### Author(s)

Hasaru Kariyawasam, Shian Su and Oliver Voogd

#### See Also

glimmaVolcano,glimmaVolcano.MArrayLM,glimmaVolcano.DGELRT,glimmaVolcano.DESeqDataSet

### **Examples**

```
dge <- readRDS(
   system.file("RNAseq123/dge.rds", package = "Glimma"))
design <- readRDS(
   system.file("RNAseq123/design.rds", package = "Glimma"))
contr.matrix <- readRDS(
   system.file("RNAseq123/contr.matrix.rds", package = "Glimma"))
dge <- edgeR::estimateDisp(dge, design)
gfit <- edgeR::glmFit(dge, design)
glrt <- edgeR::glmLRT(gfit, design, contrast = contr.matrix)
glimmaVolcano(glrt, dge = dge)</pre>
```

glimmaVolcano.DGELRT Glimma Volcano Plot

### **Description**

Draws a two-panel interactive volcano plot from an DGELRT object. This is a special case of the glimmaXY plot.

### Usage

```
## S3 method for class 'DGELRT'
glimmaVolcano(
    x,
    dge = NULL,
    counts = dge$counts,
    groups = dge$samples$group,
    status = edgeR::decideTests.DGEExact(x),
    anno = x$genes,
    display.columns = NULL,
    status.cols = c("#1052bd", "silver", "#cc212f"),
    sample.cols = NULL,
    p.adj.method = "BH",
    transform.counts = c("logcpm", "cpm", "rpkm", "none"),
    main = paste(x$comparison[2], "vs", x$comparison[1]),
    xlab = "logFC",
```

```
ylab = "negLog10PValue",
html = NULL,
width = 920,
height = 920,
...
)
```

#### **Arguments**

x DGELRT object from which summary statistics are extracted from to create

summary (left) plot.

dge DGEList object with nrow(x) rows from which expression values are extracted

from to create expression (right) plot. Gene counts are taken from dge\$counts and sample groups from dge\$samples\$group. By default raw counts are transformed to log-cpm values (see more in the transform.counts argument).

counts numeric matrix with nrow(x) rows containing gene expression values. This

can be used to replace the gene counts from dge\$counts, i.e. you may have

log-rpkm values stored in a different object that you wish to use.

groups vector of length ncol(dge) representing categorisation of samples in expression

plot.

status vector of length nrow(x) indicating the status of each gene. By default genes in

the summary plot are coloured based on its differential expression status using an adjusted p-value cutoff of 0.05 by calling the edgeR::decideTests() function, where the value of -1 marks down-regulated genes, 0 marks genes with no

expression difference, and 1 marks up-regulated genes.

anno dataframe with nrow(x) rows containing gene annotations.

display.columns

character vector containing names of columns from anno from which to display

in mouseover tooltips and table.

status.cols vector of length 3 containing valid CSS strings for colours associated with status

in the order of -1, 0 and 1.

sample.cols character vector of length ncol (counts) containing valid CSS strings for colours

associated with each sample to be displayed on the expression plot. If left un-

specified, samples will be coloured according to groups.

p.adj.method character string specifying p-value adjustment method.

transform.counts

the type of transformation used on the counts - "logcpm" for using edgeR::cpm(counts,

log=TRUE); "cpm" for edgeR::cpm(counts); "rpkm" for edgeR::rpkm(counts);
"logrpkm" for edgeR::rpkm(counts, log=TRUE); and "none" for no transfor-

mation). Defaults to "logcpm".

main character string for the main title of summary plot.

xlab character string for the x-axis label of summary plot.

ylab character string for the y-axis label of summary plot.

html character string for naming HTML file for exportation of widget. The extension

should be included in the file name e.g. "file.html".

width numeric value indicating width of widget in pixels.

height numeric value indicating width of height in pixels.

... additional unused arguments.

#### **Details**

The summary plot on the left represents gene-wise log-fold-change (logFC) on the x-axis versus -log10(pvalue). The expression plot on the right displays sample expression values for a single gene. Interactions with the htmlwidget include clicking on genes (points) in the summary plot to bring up associated sample expression values in the expression plot, as well as the summary statistics in the table below. Alternatively, users can interact with the table by clicking on genes (rows) to highlight genes in the summary plot, as well as bring up associated sample expression values in the expression plot. Briefly, other interactive features include a search box for the table, buttons to save plots and data (summary statistics and expression values), additional pop-up information when hovering on points in plots, and rescaling of the y-axis in the expression plot.

#### Value

htmlwidget object or NULL if html argument is specified.

#### Author(s)

Hasaru Kariyawasam, Shian Su and Oliver Voogd

#### See Also

 ${\tt glimmaVolcano,glimmaVolcano.MArrayLM,glimmaVolcano.DGEExact,glimmaVolcano.DESeqDataSet}$ 

```
glimmaVolcano.MArrayLM

Glimma Volcano Plot
```

#### **Description**

Draws a two-panel interactive volcano plot from an MArrayLM object. This is a special case of the glimmaXY plot.

#### Usage

```
## S3 method for class 'MArrayLM'
glimmaVolcano(
 Х,
 dge = NULL,
 counts = dge$counts,
  groups = dge$samples$group,
  coef = ncol(x$coefficients),
  status = limma::decideTests(x),
  anno = x$genes,
  display.columns = NULL,
  status.cols = c("#1052bd", "silver", "#cc212f"),
  sample.cols = NULL,
  p.adj.method = "BH"
  transform.counts = c("logcpm", "cpm", "rpkm", "none"),
 main = colnames(x)[coef],
 xlab = "logFC",
 ylab = "negLog10PValue",
```

```
html = NULL,
width = 920,
height = 920,
...
)
```

### **Arguments**

x MArrayLM object from which summary statistics are extracted from to create

summary (left) plot.

dge DGEList object with nrow(x) rows from which expression values are extracted

from to create expression (right) plot. Gene counts are taken from dge\$counts and sample groups from dge\$samples\$group. By default raw counts are transformed to log-cpm values (see more in the transform.counts argument).

counts numeric matrix with nrow(x) rows containing gene expression values. This

can be used to replace the gene counts from dge\$counts, i.e. you may have

log-rpkm values stored in a different object that you wish to use.

groups vector of length ncol(dge) representing categorisation of samples in expression

plot.

coef integer indicating the column in x from the summary plot is created.

status vector of length nrow(x) indicating the status of each gene. By default genes in

the summary plot are coloured based on its differential expression status using an adjusted p-value cutoff of 5% by calling the limma::decideTests function, where the value of -1 marks down-regulated genes, 0 marks genes with no ex-

pression difference, and 1 marks up-regulated genes.

anno dataframe with nrow(x) rows containing gene annotations.

display.columns

character vector containing names of columns from anno from which to display

in mouseover tooltips and table.

status.cols vector of length 3 containing valid CSS strings for colours associated with status

in the order of -1, 0 and 1.

sample.cols character vector of length ncol (counts) containing valid CSS strings for colours

associated with each sample to be displayed on the expression plot. If left un-

specified, samples will be coloured according to groups.

p.adj.method character string specifying p-value adjustment method.

transform.counts

the type of transformation used on the counts - "logcpm" for using edgeR::cpm(counts,

log=TRUE); "cpm" for edgeR::cpm(counts); "rpkm" for edgeR::rpkm(counts);
"logrpkm" for edgeR::rpkm(counts, log=TRUE); and "none" for no transfor-

mation). Defaults to "logcpm".

main character string for the main title of summary plot.

xlab character string for the x-axis label of summary plot. ylab character string for the y-axis label of summary plot.

html character string for naming HTML file for exportation of widget. The extension

should be included in the file name e.g. "file.html".

width numeric value indicating width of widget in pixels. height numeric value indicating width of height in pixels.

... additional unused arguments.

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#### **Details**

The summary plot on the left represents gene-wise log-fold-change (logFC) on the x-axis versus -log10(pvalue). The expression plot on the right displays sample expression values for a single gene. Interactions with the htmlwidget include clicking on genes (points) in the summary plot to bring up associated sample expression values in the expression plot, as well as the summary statistics in the table below. Alternatively, users can interact with the table by clicking on genes (rows) to highlight genes in the summary plot, as well as bring up associated sample expression values in the expression plot. Briefly, other interactive features include a search box for the table, buttons to save plots and data (summary statistics and expression values), additional pop-up information when hovering on points in plots, and rescaling of the y-axis in the expression plot.

#### Value

htmlwidget object or NULL if html argument is specified.

#### Author(s)

Hasaru Kariyawasam, Shian Su and Oliver Voogd

#### See Also

glimmaVolcano,glimmaVolcano.DGEExact,glimmaVolcano.DGELRT,glimmaVolcano.DESeqDataSet

glimmaXY

Glimma XY Plot

### **Description**

Draws a two-panel interactive XY scatter plot.

# Usage

```
glimmaXY(
 Х,
 у,
 xlab = "x",
 ylab = "y",
 dge = NULL,
 counts = dge$counts,
  groups = dge$samples$group,
  status = rep(0, length(x)),
 anno = NULL,
 display.columns = NULL,
  status.cols = c("#1052bd", "silver", "#cc212f"),
  sample.cols = NULL,
  transform.counts = c("logcpm", "cpm", "rpkm", "none"),
 main = "XY Plot",
 html = NULL,
 width = 920,
 height = 920
)
```

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### **Arguments**

x numeric vector of values to plot on the x-axis of the summary plot.
y numeric vector of values to plot on the y-axis of the summary plot.

xlab character string for the x-axis label of summary plot. ylab character string for the y-axis label of summary plot.

dge DGEList object with length(x) rows from which expression values are ex-

tracted from to create expression (right) plot. Gene counts are taken from dge\$counts

and sample groups from dge\$samples\$group.

counts numeric matrix with length(x) rows containing gene expression values. This

can be used to replace raw gene counts from dge\$counts with transformed counts

e.g. logCPM or logRPKM values.

groups vector of length ncol(counts) representing categorisation of samples in ex-

pression plot.

status vector of length length(x) indicating the status of each gene. A value of -1

marks a down-regulated gene, 0 marks a gene with no expression difference,

and 1 marks an up-regulated gene.

anno dataframe with length(x) rows containing gene annotations.

display.columns

character vector containing names of columns from anno from which to display

in mouseover tooltips and table.

status.cols vector of length 3 containing valid CSS strings for colours associated with status

in the order of -1, 0 and 1.

sample.cols character vector of length ncol (counts) containing valid CSS strings for colours

associated with each sample to be displayed on the expression plot. If left un-

specified, samples will be coloured according to groups.

transform.counts

the type of transformation used on the counts - "logcpm" for using edgeR::cpm(counts, log=TRUE); "cpm" for edgeR::cpm(counts); "rpkm" for edgeR::rpkm(counts);

"logrpkm" for edgeR::rpkm(counts, log=TRUE); and "none" for no transfor-

mation). Defaults to "logcpm".

main character string for the main title of summary plot.

html character string for naming HTML file for exportation of widget. The extension

should be included in the file name e.g. "file.html".

width numeric value indicating width of widget in pixels.

height numeric value indicating width of height in pixels.

#### **Details**

The summary plot on the left displays the x and y values specified. The expression plot on the right displays sample expression values for a single gene. Interactions with the htmlwidget include clicking on genes (points) in the summary plot to bring up associated sample expression values in the expression plot, as well as the summary statistics in the table below. Alternatively, users can interact with the table by clicking on genes (rows) to highlight genes in the summary plot, as well as bring up associated sample expression values in the expression plot. Briefly, other interactive features include a search box for the table, buttons to save plots and data (summary statistics and expression values), additional pop-up information when hovering on points in plots, and rescaling of the y-axis in the expression plot.

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#### Value

htmlwidget object or NULL if html argument is specified.

#### Author(s)

Hasaru Kariyawasam, Shian Su and Oliver Voogd

### **Examples**

```
dge <- readRDS(
   system.file("RNAseq123/dge.rds", package = "Glimma"))
design <- readRDS(
   system.file("RNAseq123/design.rds", package = "Glimma"))
contr.matrix <- readRDS(
   system.file("RNAseq123/contr.matrix.rds", package = "Glimma"))
v <- limma::voom(dge, design)
vfit <- limma::lmFit(v, design)
vfit <- limma::contrasts.fit(vfit, contrasts = contr.matrix)
efit <- limma::eBayes(vfit)
glimmaXY(efit$Amean, efit$coefficients)</pre>
```

glimmaXYWidget

GlimmaXY HTMLWidget Wrapper

### **Description**

Passes packaged data to JS interface for rendering.

### Usage

```
glimmaXYWidget(xData, width, height, html)
```

#### **Arguments**

xData packaged data object returned from buildXYData()

width htmlwidget element width in pixels height htmlwidget element height in pixels

html name of HTML file (including extension) to export widget into rather than dis-

playing the widget; NULL by default.

#### Value

htmlwidget object for XY plot internal use

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glimma\_plot

Glimma plot manager

# Description

Core glimma plot manager. Generates environment for glimma plots.

### Usage

```
glimma_plot(
    ...,
    layout = c(1, 1),
    path = getwd(),
    folder = "glimma-plots",
    html = "index",
    overwrite = TRUE,
    launch = TRUE
)
```

# Arguments

... the jschart or jslink objects for processing.

layout the numeric vector representing the number of rows and columns in plot window.

path the path in which the folder will be created.

folder the name of the fold to save html file to.

html the name of the html file to save plots to.

overwrite the option to overwrite existing folder if it already exists.

launch TRUE to launch plot after call.

#### Value

Generates interactive plots based on filling layout row by row from left to right.

gllink

Plot linkages

# Description

Helper function for writing the link properties in interactive Glimma plots

### Usage

```
gllink(
  from,
  to,
  src = "none",
  dest = "none",
  flag = "none",
  both = FALSE,
  info = "none"
)
```

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# **Arguments**

from	the index of the plot from which the event is dispatched.
to	the index of the plot which receives the event and performs an action.
src	the action that is performed in the "from" plot.
dest	the action that is performed in the "to" plot.
flag	indicates special links for particular chart types.
both	creates symmetric links whereby the "dest" action in "to" also triggers the "src" action in "from".
info	additional info for creating the link.

# Value

a link object containing the plot linking information.

glMDPlot	Glimma MD Plot	
----------	----------------	--

# Description

Draw an interactive MD plot

# Usage

```
glMDPlot(x, ...)
```

# **Arguments**

x the DE object to plot.

additional arguments affecting the plots produced. See specific methods for detailed arguments.

# Value

Draws a two-panel interactive MD plot in an html page. The left plot shows the log-fold-change vs average expression. The right plot shows the expression levels of a particular gene of each sample. Hovering over points on left plot will plot expression level for corresponding gene, clicking on points will fix the expression plot to gene. Clicking on rows on the table has the same effect as clicking on the corresponding gene in the plot.

# Author(s)

Shian Su

#### See Also

glMDPlot.default,glMDPlot.DGELRT,glMDPlot.DGEExact,glMDPlot.MArrayLM,glMDPlot.DESeqDataSet

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glMDPlot.default Glimma MD Plot

# **Description**

Draw an interactive MD plot from a data.frame

# Usage

```
## Default S3 method:
glMDPlot(
  Х,
  xval,
  yval,
  counts = NULL,
  anno = NULL,
  groups = NULL,
  samples = NULL,
  status = rep(0, nrow(x)),
  transform = FALSE,
  main = "",
  xlab = xval,
  ylab = yval,
  side.main = "GeneID",
  side.xlab = "Group",
  side.ylab = "Expression",
  side.log = FALSE,
  side.gridstep = ifelse(!transform || side.log, FALSE, 0.5),
  jitter = 30,
  display.columns = side.main,
  cols = c("#00bfff", "#858585", "#ff3030"),
  sample.cols = rep("#1f77b4", ncol(counts)),
  path = getwd(),
  folder = "glimma-plots",
  html = "MD-Plot",
  launch = TRUE,
  . . .
)
```

Х	the data.frame object containing expression and fold change values.
xval	the column to plot on x axis of left plot.
yval	the column to plot on y axis of left plot.
counts	the matrix of expression values, with samples in columns.
anno	the data.frame containing gene annotations.
groups	the factor containing experimental groups of the samples.
samples	the names of the samples.

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status vector giving the control status of data point, of same length as the number of

rows of object. If NULL, then all points are plotted in the default colour.

transform TRUE if counts should be log-cpm transformed.

main the title for the left plot.

xlab the label on the x axis for the left plot.
ylab the label on the y axis for the left plot.

side.main the column containing mains for right plot.

side.xlab label for x axis on right plot. side.ylab label for y axis on right plot.

side.log TRUE to plot expression on the right plot on log scale.

side.gridstep intervals along which to place grid lines on y axis. Currently only available for

linear scale.

jitter the amount of jitter to apply to the samples in the expressions plot.

display.columns

character vector containing names of columns to display in mouseover tooltips

and table.

cols vector of strings denoting colours corresponding to control status -1, 0 and 1.

(may be R named colours or Hex values)

sample.cols vector of strings denoting colours for each sample point on the expression plot.

path the path in which the folder will be created.

folder the name of the fold to save html file to.

html the name of the html file to save plots to.

launch TRUE to launch plot after call.

additional arguments to be passed onto the MD plot. (main, xlab, ylab can be

set for the left plot)

# Value

Draws a two-panel interactive MD plot in an html page. The left plot shows the log-fold-change vs average expression. The right plot shows the expression levels of a particular gene of each sample. Hovering over points on left plot will plot expression level for corresponding gene, clicking on points will fix the expression plot to gene. Clicking on rows on the table has the same effect as clicking on the corresponding gene in the plot.

### Author(s)

Shian Su

```
{\tt glMDPlot.DESeqDataSet} \quad \textit{Glimma MD Plot}
```

# **Description**

Draw an interactive MD plot from a DESeqDataSet object

### Usage

```
## S3 method for class 'DESeqDataSet'
glMDPlot(
  х,
  counts = NULL,
  anno,
  groups,
  samples = NULL,
  status = rep(0, nrow(x)),
  transform = FALSE,
  main = "",
  xlab = "Mean Expression",
  ylab = "log-fold-change",
  side.xlab = "Group",
  side.ylab = "logMean",
  side.log = FALSE,
  side.gridstep = ifelse(!transform || side.log, FALSE, 0.5),
  jitter = 30,
  side.main = "GeneID",
  display.columns = NULL,
  cols = c("#00bfff", "#858585", "#ff3030"),
  sample.cols = rep("#1f77b4", ncol(x)),
  path = getwd(),
  folder = "glimma-plots",
  html = "MD-Plot",
  launch = TRUE,
)
```

Χ	the DESeqDataSet object.
counts	the matrix of expression values, with samples in columns.
anno	the data.frame containing gene annotations.
groups	the factor containing experimental groups of the samples.
samples	the names of the samples.
status	vector giving the control status of data point, of same length as the number of rows of object. If NULL, then all points are plotted in the default colour.
transform	TRUE if counts should be log-cpm transformed.
main	the title for the left plot.
xlab	label for x axis on left plot.

ylab	label for y axis on left plot.
side.xlab	label for x axis on right plot.
side.ylab	label for y axis on right plot.

side.log TRUE to plot expression on the right plot on log scale.

side.gridstep intervals along which to place grid lines on y axis. Currently only available for

linear scale.

jitter the amount of jitter to apply to the samples in the expressions plot.

side.main the column containing mains for right plot.

display.columns

character vector containing names of columns to display in mouseover tooltips

and table.

cols vector of strings denoting colours corresponding to control status -1, 0 and 1.

(may be R named colours or Hex values)

sample.cols vector of strings denoting colours for each sample point on the expression plot.

the path in which the folder will be created.

folder the name of the fold to save html file to.

html the name of the html file to save plots to.

launch TRUE to launch plot after call.

... additional arguments to be passed onto the MD plot. (main, xlab, ylab can be

set for the left plot)

#### Value

Draws a two-panel interactive MD plot in an html page. The left plot shows the log-fold-change vs average expression. The right plot shows the expression levels of a particular gene of each sample. Hovering over points on left plot will plot expression level for corresponding gene, clicking on points will fix the expression plot to gene. Clicking on rows on the table has the same effect as clicking on the corresponding gene in the plot.

# Author(s)

Shian Su

glMDPlot.DESeqResults Glimma MD Plot

# **Description**

Draw an interactive MD plot from a DESeqResults object

### Usage

```
## S3 method for class 'DESeqResults'
glMDPlot(
  Х,
  counts = NULL,
  anno,
  groups,
  samples = NULL,
  status = rep(0, nrow(x)),
  transform = FALSE,
 main = "",
  xlab = "Mean Expression",
  ylab = "log-fold-change",
  side.xlab = "Group",
  side.ylab = "Expression",
  side.log = FALSE,
  side.gridstep = ifelse(!transform || side.log, FALSE, 0.5),
  jitter = 30,
  side.main = "GeneID",
  display.columns = NULL,
  cols = c("#00bfff", "#858585", "#ff3030"),
  sample.cols = rep("#1f77b4", ncol(counts)),
  path = getwd(),
  folder = "glimma-plots",
  html = "MD-Plot",
  launch = TRUE,
)
```

X	the DESeqResults object.
counts	the matrix of expression values, with samples in columns.
anno	the data.frame containing gene annotations.
groups	the factor containing experimental groups of the samples.
samples	the names of the samples.
status	vector giving the control status of data point, of same length as the number of rows of object. If NULL, then all points are plotted in the default colour.
transform	TRUE if counts should be log-cpm transformed.
main	the title for the left plot.
xlab	label for x axis on left plot.
ylab	label for y axis on left plot.
side.xlab	label for x axis on right plot.
side.ylab	label for y axis on right plot.
side.log	TRUE to plot expression on the right plot on log scale.
side.gridstep	intervals along which to place grid lines on y axis. Currently only available for linear scale.
jitter	the amount of jitter to apply to the samples in the expressions plot.

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side.main the column containing mains for right plot.

display.columns

character vector containing names of columns to display in mouseover tooltips

and table.

cols vector of strings denoting colours corresponding to control status -1, 0 and 1.

(may be R named colours or Hex values)

sample.cols vector of strings denoting colours for each sample point on the expression plot.

path the path in which the folder will be created.

folder the name of the fold to save html file to.

html the name of the html file to save plots to.

launch TRUE to launch plot after call.

additional arguments to be passed onto the MD plot. (main, xlab, ylab can be

set for the left plot)

#### Value

Draws a two-panel interactive MD plot in an html page. The left plot shows the log-fold-change vs average expression. The right plot shows the expression levels of a particular gene of each sample. Hovering over points on left plot will plot expression level for corresponding gene, clicking on points will fix the expression plot to gene. Clicking on rows on the table has the same effect as clicking on the corresponding gene in the plot.

# Author(s)

Shian Su

glMDPlot.DGEExact Glimma MD Plot

### **Description**

Draw an interactive MD plot from a DGELRT objet

# Usage

```
## S3 method for class 'DGEExact'
glMDPlot(
    x,
    counts = NULL,
    anno = NULL,
    groups = NULL,
    samples = NULL,
    status = rep(0, nrow(x)),
    transform = FALSE,
    main = "",
    xlab = "Average log CPM",
    ylab = "log-fold-change",
    side.xlab = "Group",
    side.ylab = "Expression",
```

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```
side.log = FALSE,
side.gridstep = ifelse(!transform || side.log, FALSE, 0.5),
p.adj.method = "BH",
jitter = 30,
side.main = "GeneID",
display.columns = NULL,
cols = c("#00bfff", "#858585", "#ff3030"),
sample.cols = rep("#1f77b4", ncol(counts)),
path = getwd(),
folder = "glimma-plots",
html = "MD-Plot",
launch = TRUE,
...
)
```

### **Arguments**

x the DGEExact object.

counts the matrix of expression values, with samples in columns.

anno the data.frame containing gene annotations.

groups the factor containing experimental groups of the samples.

samples the names of the samples.

status vector giving the control status of data point, of same length as the number of

rows of object. If NULL, then all points are plotted in the default colour.

transform TRUE if counts should be log-cpm transformed.

main the title for the left plot.

xlab label for x axis on left plot.

ylab label for y axis on left plot.

side.xlab label for x axis on right plot.

side.ylab label for y axis on right plot.

side.log TRUE to plot expression on the right plot on log scale.

side.gridstep intervals along which to place grid lines on y axis. Currently only available for

linear scale.

p.adj.method character vector indicating multiple testing correction method. See p.adjust

for available methods. (defaults to "BH")

jitter the amount of jitter to apply to the samples in the expressions plot.

side.main the column containing mains for right plot.

display.columns

character vector containing names of columns to display in mouseover tooltips

and table.

cols vector of strings denoting colours corresponding to control status -1, 0 and 1.

(may be R named colours or Hex values)

sample.cols vector of strings denoting colours for each sample point on the expression plot.

path the path in which the folder will be created.

folder the name of the fold to save html file to.

html the name of the html file to save plots to.

launch TRUE to launch plot after call.

... additional arguments to be passed onto the MD plot. (main, xlab, ylab can be

set for the left plot)

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### Value

Draws a two-panel interactive MD plot in an html page. The left plot shows the log-fold-change vs average expression. The right plot shows the expression levels of a particular gene of each sample. Hovering over points on left plot will plot expression level for corresponding gene, clicking on points will fix the expression plot to gene. Clicking on rows on the table has the same effect as clicking on the corresponding gene in the plot.

#### Author(s)

Shian Su

glMDPlot.DGELRT

Glimma MD Plot

# **Description**

Draw an interactive MD plot from a DGELRT object

# Usage

```
## S3 method for class 'DGELRT'
glMDPlot(
  х,
  counts = NULL,
  anno = NULL,
  groups = NULL,
  samples = NULL,
  status = rep(0, nrow(x)),
  transform = FALSE,
  main = "",
  xlab = "Average log CPM",
  ylab = "log-fold-change",
  side.xlab = "Group",
  side.ylab = "Expression",
  side.log = FALSE,
  side.gridstep = ifelse(!transform || side.log, FALSE, 0.5),
  p.adj.method = "BH",
  jitter = 30,
  side.main = "GeneID",
  display.columns = NULL,
  cols = c("#00bfff", "#858585", "#ff3030"),
  sample.cols = rep("#1f77b4", ncol(counts)),
  path = getwd(),
  folder = "glimma-plots",
  html = "MD-Plot",
  launch = TRUE,
)
```

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#### **Arguments**

x the DGELRT object.

counts the matrix of expression values, with samples in columns.

anno the data.frame containing gene annotations.

groups the factor containing experimental groups of the samples.

samples the names of the samples.

status vector giving the control status of data point, of same length as the number of

rows of object. If NULL, then all points are plotted in the default colour.

transform TRUE if counts should be log-cpm transformed.

main the title for the left plot.

xlab label for x axis on left plot.

ylab label for y axis on left plot.

side.xlab label for x axis on right plot.

side.ylab label for y axis on right plot.

side.log TRUE to plot expression on the right plot on log scale.

side.gridstep intervals along which to place grid lines on y axis. Currently only available for

inear scale.

p.adj.method character vector indicating multiple testing correction method. See p.adjust

for available methods. (defaults to "BH")

jitter the amount of jitter to apply to the samples in the expressions plot.

side.main the column containing mains for right plot.

display.columns

character vector containing names of columns to display in mouseover tooltips

and table.

vector of strings denoting colours corresponding to control status -1, 0 and 1.

(may be R named colours or Hex values)

sample.cols vector of strings denoting colours for each sample point on the expression plot.

path the path in which the folder will be created.

folder the name of the fold to save html file to.

html the name of the html file to save plots to.

launch TRUE to launch plot after call.

... additional arguments to be passed onto the MD plot. (main, xlab, ylab can be

set for the left plot)

#### Value

Draws a two-panel interactive MD plot in an html page. The left plot shows the log-fold-change vs average expression. The right plot shows the expression levels of a particular gene of each sample. Hovering over points on left plot will plot expression level for corresponding gene, clicking on points will fix the expression plot to gene. Clicking on rows on the table has the same effect as clicking on the corresponding gene in the plot.

#### Author(s)

Shian Su

gIMDPlot.MArrayLM 47

glMDPlot.MArrayLM Glimma MD Plot

### **Description**

Draw an interactive MD plot from a MArrayLM object

#### Usage

```
## S3 method for class 'MArrayLM'
glMDPlot(
  х,
  counts = NULL,
  anno = NULL,
  groups = NULL,
  samples = NULL,
  status = rep(0, nrow(x)),
  transform = FALSE,
  main = "",
  xlab = "Average log CPM",
  ylab = "log-fold-change",
  side.main = "GeneID",
  side.xlab = "Group",
  side.ylab = "Expression",
  side.log = FALSE,
  side.gridstep = ifelse(!transform || side.log, FALSE, 0.5),
  coef = ncol(x$coefficients),
  p.adj.method = "BH",
  jitter = 30,
  display.columns = NULL,
  cols = c("#00bfff", "#858585", "#ff3030"),
  sample.cols = rep("#1f77b4", ncol(counts)),
  path = getwd(),
  folder = "glimma-plots",
  html = "MD-Plot",
  launch = TRUE,
)
```

X	the MArrayLM object.
counts	the matrix of expression values, with samples in columns.
anno	the data frame containing gene annotations.
groups	the factor containing experimental groups of the samples.
samples	the names of the samples.
status	vector giving the control status of data point, of same length as the number of rows of object. If NULL, then all points are plotted in the default colour.
transform	TRUE if counts should be log-cpm transformed.

main	the title for the left plot.
xlab	label for x axis on left plot.
ylab	label for y axis on left plot.
side.main	the column containing mains for right plot.
side.xlab	label for x axis on right plot.
side.ylab	label for y axis on right plot.
side.log	TRUE to plot expression on the right plot on log scale.
side.gridstep	intervals along which to place grid lines on y axis. Currently only available for linear scale.
coef	integer or character index vector indicating which column of object to plot.
p.adj.method	character vector indicating multiple testing correction method. See $p.adjust$ for available methods. (defaults to "BH")
jitter	the amount of jitter to apply to the samples in the expressions plot.
display.column	s
	character vector containing names of columns to display in mouseover tooltips and table.
cols	vector of strings denoting colours corresponding to control status -1, $0$ and $1$ . (may be R named colours or Hex values)
sample.cols	vector of strings denoting colours for each sample point on the expression plot.
path	the path in which the folder will be created.
folder	the name of the fold to save html file to.
html	the name of the html file to save plots to.
launch	TRUE to launch plot after call.

# Value

Draws a two-panel interactive MD plot in an html page. The left plot shows the log-fold-change vs average expression. The right plot shows the expression levels of a particular gene of each sample. Hovering over points on left plot will plot expression level for corresponding gene, clicking on points will fix the expression plot to gene. Clicking on rows on the table has the same effect as clicking on the corresponding gene in the plot.

set for the left plot)

additional arguments to be passed onto the MD plot. (main, xlab, ylab can be

# Author(s)

Shian Su

glMDRmd 49

glMDRmd

glMDPlot Rmarkdown link and instructions

# **Description**

When run inside of a text-block of Rmarkdown document using 'r ...' this produces a link and instructions about the usage of the interactive plots.

# Usage

```
glMDRmd(html = "MD-Plot")
```

### **Arguments**

html

name of the HTML page containing plots from glMDPlot.

#### Value

None

#### See Also

glMDPlot

### **Examples**

glMDRmd()

glMDSPlot

Glimma MDS Plot

### **Description**

Draw an interactive MD plot from a DGEList object with distances calculated from most variable genes.

# Usage

```
glMDSPlot(x, ...)
```

# **Arguments**

x the matrix containing the gene expressions.

... additional arguments.

### Value

Draws a two-panel interactive MDS plot in an html page. The left panel contains the plot between two MDS dimensions, with annotations displayed on hover. The right panel contains a bar plot of the eigenvalues of each dimension, clicking on any of the bars will plot the corresponding dimension against the next dimension.

50 glMDSPlot.default

### Author(s)

Shian Su, Gordon Smyth

### See Also

```
glMDSPlot.default, glMDSPlot.DGEList
```

glMDSPlot.default

Glimma MDS Plot

### **Description**

Draw an interactive MD plot from a DGEList object with distances calculated from most variable

### Usage

```
## Default S3 method:
glMDSPlot(
  х,
  top = 500,
  labels = seq_cols(x),
  groups = rep(1, ncol(x)),
  gene.selection = c("pairwise", "common"),
  main = "MDS Plot",
  path = getwd(),
  folder = "glimma-plots",
  html = "MDS-Plot",
  launch = TRUE,
)
```

# **Arguments**

top

the matrix containing the gene expressions. the number of top most variable genes to use.

labels the labels for each sample.

the experimental group to which samples belong. groups

"pairwise" if most variable genes are to be chosen for each pair of samples or gene.selection

"common" to select the same genes for all comparisons.

main the title of the plot.

the path in which the folder will be created. path folder the name of the fold to save html file to. html the name of the html file to save plots to.

TRUE to launch plot after call. launch

additional arguments. . . .

### Value

Draws a two-panel interactive MDS plot in an html page. The left panel contains the plot between two MDS dimensions, with annotations displayed on hover. The right panel contains a bar plot of the eigenvalues of each dimension, clicking on any of the bars will plot the corresponding dimension against the next dimension.

# Author(s)

Shian Su, Gordon Smyth

```
{\tt glMDSPlot.DESeqDataSet}
```

Glimma MDS Plot

# Description

Draw an interactive MD plot from a DGEList object with distances calculated from most variable genes.

# Usage

```
## S3 method for class 'DESeqDataSet'
glMDSPlot(
    x,
    top = 500,
    labels = NULL,
    groups = NULL,
    gene.selection = c("pairwise", "common"),
    prior.count = 0.25,
    main = "MDS Plot",
    path = getwd(),
    folder = "glimma-plots",
    html = "MDS-Plot",
    launch = TRUE,
    ...
)
```

X	the DESeqDataSet containing the gene expressions.
top	the number of top most variable genes to use.
labels	the labels for each sample.
groups	the experimental group to which samples belong.
gene.selection	"pairwise" if most variable genes are to be chosen for each pair of samples or "common" to select the same genes for all comparisons.
prior.count	average count to be added to each observation to avoid taking log of zero. Used only if log=TRUE.
main	the title of the plot.
path	the path in which the folder will be created.

52 gIMDSPlot.DGEList

```
folder the name of the fold to save html file to.

html the name of the html file to save plots to.

launch TRUE to launch plot after call.
```

TRUE to faulten plot after car

... additional arguments.

#### Value

Draws a two-panel interactive MDS plot in an html page. The left panel contains the plot between two MDS dimensions, with annotations displayed on hover. The right panel contains a bar plot of the eigenvalues of each dimension, clicking on any of the bars will plot the corresponding dimension against the next dimension.

# Author(s)

Shian Su, Gordon Smyth

glMDSPlot.DGEList Glimma MDS Plot

# **Description**

Draw an interactive MD plot from a DGEList object with distances calculated from most variable genes.

### Usage

```
## S3 method for class 'DGEList'
glMDSPlot(
    x,
    top = 500,
    labels = NULL,
    groups = rep(1, ncol(x)),
    gene.selection = c("pairwise", "common"),
    prior.count = 2,
    main = "MDS Plot",
    path = getwd(),
    folder = "glimma-plots",
    html = "MDS-Plot",
    launch = TRUE,
    ...
)
```

### **Arguments**

x the DGEList containing the gene expressions.

top the number of top most variable genes to use.

labels the labels for each sample.

groups the experimental group to which samples belong.

gene.selection "pairwise" if most variable genes are to be chosen for each pair of samples or "common" to select the same genes for all comparisons.

glScatter 53

prior.count average count to be added to each observation to avoid taking log of zero. Used

only if log=TRUE.

main the title of the plot.

path the path in which the folder will be created.

folder the name of the fold to save html file to.

html the name of the html file to save plots to.

launch TRUE to launch plot after call.

... additional arguments.

### Value

Draws a two-panel interactive MDS plot in an html page. The left panel contains the plot between two MDS dimensions, with annotations displayed on hover. The right panel contains a bar plot of the eigenvalues of each dimension, clicking on any of the bars will plot the corresponding dimension against the next dimension.

# Author(s)

Shian Su, Gordon Smyth

glScatter

Glimma Scatter Plot

# Description

Create an interactive scatter plot object

# Usage

```
glScatter(x, ...)
```

# **Arguments**

x the data.frame containing data to plot.

... additional arguments depending on input object type.

### Value

A chart object containing the information to create an interactive scatter plot.

# Author(s)

Shian Su

54 glScatter.default

glScatter.default Glimma Scatter Plot

# **Description**

Default method for creating an interactive scatter plot

### Usage

```
## Default S3 method:
glScatter(
  х,
  xval = "x",
  yval = "y",
  idval = NULL,
  point.size = 2,
  x.jitter = 0,
  y.jitter = 0,
  ndigits = NULL,
  signif = 6,
  log = "",
  xgrid = FALSE,
  ygrid = FALSE,
  xstep = FALSE,
  ystep = FALSE,
  xlab = xval,
  ylab = yval,
  main = NULL,
  height = 400,
  width = 500,
  colval = NULL,
  annot = c(xval, yval),
  annot.lab = NULL,
  flag = NULL,
  info = NULL,
  hide = FALSE,
  disable = NULL,
)
```

```
x the data.frame containing data to plot.

xval the column name for the x-axis values.

yval the column name for the y-axis values.

idval the column name for unique identifiers.

point.size the size of the data points.

x.jitter the amount of jittering to add to values along the x axis.

y.jitter the amount of jittering to add to values along the y axis.
```

glTable 55

ndigits	the number of digits after the decimal to round to in the tooltip (overrides signif).
signif	the number of significant figures to display in the tooltip.
log	a character string which contains "x" if the x axis is to be logarithmic, "y" if the y axis is to be logarithmic and "xy" or "yx" if both axes are to be logarithmic.
xgrid	TRUE if grid lines should be placed along x axis.
ygrid	TRUE if grid lines should be placed y axis.
xstep	the interval at which to set grid lines along the x axis.
ystep	the interval at which to set grid lines along the y axis.
xlab	the label on the x-axis.
ylab	the label on the y-axis.
main	the title for the plot.
height	the height of the plot (in pixels).
width	the width of the plot (in pixels).
colval	the colours for each data point.
annot	the columns to display in the tooltip.
annot.lab	alternative labels for the values displayed in the tooltip.
flag	the special flag to indicate special plot.
info	additional information for plotting.
hide	TRUE to hide the plot when page starts.
disable	the events to disable, options are "click", "hover", "zoom".
• • •	additional arguments.

### Value

A chart object containing the information to create an interactive scatter plot.

# Author(s)

Shian Su

glTable	Glimma Table

# Description

Create a table using the data from a chart.

# Usage

```
glTable(target, columns)
```

# Arguments

the index of the plot from which data is drawn.

columns of data to plot.

# Value

a input object containing the input field information.

56 glXYPlot

gltablink

Plot linkages

# **Description**

Helper function for writing the link properties in interactive Glimma plots

# Usage

```
gltablink(from, to, action = "none", info = "none")
```

# **Arguments**

from the index of the source table.

to the index of the plot which receives the event and performs an action.

action the action that is performed in the plot. info additional info for creating the link.

### Value

a link object containing the plot linking information.

glXYPlot

Glimma XY Plot

# Description

Draw an interactive XY plot with multiple panels

# Usage

```
glXYPlot(
  Х,
  у,
  counts = NULL,
  groups = NULL,
  samples = NULL,
  status = rep(0, nrow(data)),
  anno = NULL,
  display.columns = NULL,
  xlab = "x",
  ylab = "y",
  side.main = "GeneID",
  side.xlab = "Group",
  side.ylab = "Expression",
  sample.cols = rep("#1f77b4", length(groups)),
  cols = c("#00bfff", "#858585", "#ff3030"),
  jitter = 30,
  path = getwd(),
```

glXYPlot 57

```
folder = "glimma-plots",
html = "XY-Plot",
launch = TRUE,
...
)
```

# **Arguments**

x a numeric vector of values to plot on the x-axis of the summary plot. y a numeric vector of values to plot on the y-axis of the summary plot.

counts the matrix containing all counts, the column order should correspond to the order

of the x and y vectors.

groups the factor containing experimental groups of the samples.

samples the names of the samples.

status vector giving the control status of data point, of same length as the number of

rows of object. If NULL, then all points are plotted in the default colour

anno the data.frame containing gene annotations.

display.columns

character vector containing names of columns to display in mouseover tooltips

and table.

xlab the label on the x axis for the left plot.
ylab the label on the y axis for the left plot.
side.main the column containing mains for right plot.
side.xlab the label on the x axis for the right plot.
side.ylab the label on the y axis for the right plot.

sample.cols vector of strings denoting colours for each sample point on the expression plot. vector of strings denoting colours corresponding to control status -1, 0 and 1.

(may be R named colours or Hex values)

jitter the amount of jitter to apply to the samples in the expressions plot.

path the path in which the folder will be created.
folder the name of the fold to save html file to.
html the name of the html file to save plots to.

launch TRUE to launch plot after call.

... additional arguments to be passed onto the MD plot. (main, etc. can be set for

the left plot)

#### Value

Draws a two-panel interactive XY scatter plot in an html page. The left plot shows the x and y values specified. The right plot shows the expression levels of a particular gene in each sample. Hovering over points on left plot will plot expression level for the corresponding gene, clicking on points will fix the expression plot to that gene. Clicking on rows on the table has the same effect as clicking on the corresponding gene in the plot. This function generates a display that is similar in style to glMDPlot, except that it provides more flexibility in what the user can provide.

### Author(s)

Charity Law and Shian Su

58 makeJson

# **Examples**

is.hex

Hexcode colours

# Description

Check if string(s) are valid hex colour representation

# Usage

```
is.hex(x)
```

# **Arguments**

Х

the colour value(s) to check.

# Value

Logical vector indicating if strings(s) are valid hex representations

 ${\sf makeJson}$ 

JSON converter for R objects

# Description

Function to generate json strings from

# Usage

```
makeJson(x, ...)
```

# **Arguments**

x the object to be converted into JSON

... additional arguments

# Value

```
a stringified JSON object.
```

makeJson.data.frame 59

makeJson.data.frame

JSON converter for data frames

# **Description**

Function to create a JSON from a data.frame

### Usage

```
## S3 method for class 'data.frame'
makeJson(df, convert.logical = TRUE, dataframe = c("rows", "columns"))
```

# **Arguments**

df the data.frame to be converted into JSON convert.logical

whether to convert logicals into strings "TRUE" and "FALSE"

dataframe how to encode data.frame objects: must be one of 'rows', 'columns'

### Value

a stringified JSON, the data.frame is encoded as a vector of objects, with each column being one object with keys corresponding to column names.

makeJson.jschart

JSON converter for chart objects

# Description

Function to make json object from a chart, ignoring the json property

# Usage

```
## S3 method for class 'jschart'
makeJson(chart)
```

### **Arguments**

chart

the chart object to be converted into JSON

# Value

a stringified JSON object containing the chart data.

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