# Package 'bigPint'

October 16, 2020

```
Version 1.4.0
```

Title Big multivariate data plotted interactively

**Description** Methods for visualizing large multivariate datasets using static and interactive scatterplot matrices, parallel coordinate plots, volcano plots, and litre plots. Includes examples for visualizing RNA-sequencing datasets and differentially expressed genes.

```
License GPL-3
Depends R (>= 3.6.0)
Imports DelayedArray (>= 0.12.2), dplyr (>= 0.7.2), GGally (>= 1.3.2),
      ggplot2 (>= 2.2.1), graphics (>= 3.5.0), grDevices (>= 3.5.0),
      grid (>= 3.5.0), gridExtra (>= 2.3), hexbin (>= 1.27.1), Hmisc
      (>= 4.0.3), htmlwidgets (>= 0.9), methods (>= 3.5.2), plotly
      (>= 4.7.1), plyr (>= 1.8.4), RColorBrewer (>= 1.1.2), reshape
      (>= 0.8.7), shiny (>= 1.0.5), shinycssloaders (>= 0.2.0),
      shinydashboard (>= 0.6.1), stats (>= 3.5.0), stringr (>=
      1.3.1), SummarizedExperiment (>= 1.16.1), tidyr (>= 0.7.0),
      utils (>= 3.5.0)
VignetteBuilder knitr
Suggests BiocGenerics (>= 0.29.1), data.table (>= 1.11.8), EDASeq (>=
      2.14.0), edgeR (>= 3.22.2), gtools (>= 3.5.0), knitr (>= 1.13),
      matrixStats (>= 0.53.1), rmarkdown (>= 1.10), roxygen2 (>=
      3.0.0), RUnit (>= 0.4.32), tibble (>= 1.4.2),
biocViews Clustering, DataImport, DifferentialExpression,
      GeneExpression, MultipleComparison, Normalization,
      Preprocessing, QualityControl, RNASeq, Sequencing, Software,
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RoxygenNote 7.0.2
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```

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 $\verb|bigPint| bigPint| package|$ 

# Description

bigPint R API

# **Details**

See the README on GitHub

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convertSEPair	Convert SummarizedExperiment to contain defined treatment pair

## **Description**

Reduce a SummarizedExperiment object that initially contains more than two treatment groups to now only contain a user-specified subset pair of treatment groups. Note that this function is only necesary for users using the SummarizedExperiment input object (instead of the combination of data and dataMetrics input objects.)

# Usage

```
convertSEPair(dataSE, group1, group2)
```

#### **Arguments**

dataSE	SUMMARIZEDEXPERIMENT   Summarized experiment format that can be used in lieu of data and dataMetrics
group1	CHARACTER STRING   Name of one treatment group that will remain in the dataSE object
group2	CHARACTER STRING   Name of second treatment group that will remain in the dataSE object

#### Value

A new dataSE object that is a subset of the input dataSE in that it now only contains the user-specified pair of treatment groups.

# **Examples**

```
# Example: Read in example SummarizedExperiment object that contains three
# treatment groups (S1, S2, and S3). Reduce it to now only contain two
# treatment groups (S1 and S3).

data(se_soybean_cn_sub)
se_soybean_cn_sub_2 <- convertSEPair(se_soybean_cn_sub, "S1", "S3")</pre>
```

 ${\tt convertSESubsetGenes} \quad \textit{Convert Summarized Experiment to only contain defined genes}$ 

# Description

Reduce a SummarizedExperiment object so that it only now contains a subset of genes.

## Usage

```
convertSESubsetGenes(dataSE, geneList)
```

## **Arguments**

dataSE SUMMARIZEDEXPERIMENT | Summarized experiment format that can be

used in lieu of data and dataMetrics

geneList CHARACTER ARRAY | List of gene IDs to remain in dataSE

#### Value

A new dataSE object that is a subset of the input dataSE in that it now only contains the user-specified list of genes.

#### **Examples**

```
# Example: Read in example SummarizedExperiment object that originally
# contains 5604 genes. Reduce it to now only contain ten genes (specifically
# the ones with the lowest FDR).

suppressMessages(library(dplyr))
suppressMessages(library(SummarizedExperiment))
data(se_soybean_ir_sub)
geneList <- as.data.frame(rowData(se_soybean_ir_sub)) %>%
    arrange(N_P.FDR) %>% filter(row_number() <= 10)
geneList <- geneList[,1]
se_soybean_ir_sub_2 <- convertSESubsetGenes(se_soybean_ir_sub, geneList)</pre>
```

plotClusters

Plot static parallel coordinate clusters

## **Description**

Perform hierarchical clustering analysis and visualize results with parallel coordinate plots. Optionally, save gene IDs within each cluster to .rds files for later use.

### Usage

```
plotClusters(
 data,
 dataMetrics = NULL,
 dataSE = NULL,
  geneList = NULL,
  geneLists = NULL,
  threshVar = "FDR",
  threshVal = 0.05,
  clusterAllData = TRUE,
 nC = 4,
 colList = rainbow(nC),
 aggMethod = c("ward.D", "ward.D2", "single", "complete", "average", "mcquitty",
    "median", "centroid"),
 yAxisLabel = "Count",
 xAxisLabel = "Sample",
  lineSize = 0.1,
```

```
lineAlpha = 0.5,
vxAxis = FALSE,
outDir = tempdir(),
saveFile = TRUE,
verbose = FALSE
)
```

#### **Arguments**

data DATA FRAME | Read counts

dataMetrics LIST | Differential expression metrics; default NULL

dataSE SUMMARIZEDEXPERIMENT | Summarized experiment format that can be

used in lieu of data and dataMetrics; default NULL

geneList CHARACTER ARRAY | Array of ID values of genes to be drawn from data as

parallel coordinate lines. Use this parameter if you have predetermined genes to be drawn. These genes will be clustered. Otherwise, use dataMetrics, threshVar, and threshVal to create clusters to be overlaid as parallel coordinate lines; default

NULL. See package website for examples

geneLists LIST | List of ID values of genes already clustered to be #' drawn from data

as parallel coordinate lines. Each list item is an array of genes ID values that are already grouped as a cluster. Unlike the singular geneList object, the plural geneLists object is not be clustered. If you instead wish to cluster genes, use dataMetrics, threshVar, and threshVal or geneList to create clusters to be overlaid as parallel coordinate lines; default NULL. See package website for

examples

threshVar CHARACTER STRING | Name of column in dataMetrics object that is used to

threshold significance; default "FDR"

threshVal INTEGER | Maximum value to threshold significance from threshVar object;

default 0.05

 ${\tt clusterAllData} \ \ BOOLEAN\ [TRUE\ |\ FALSE]\ |\ Create\ clusters\ based\ on\ the\ whole\ dataset\ and$ 

then assign significant genes to those clusters; default is TRUE. If FALSE, create clusters based on just the significant genes. With either option, the side-by-side boxplot will represent the whole dataset (from data input) and the parallel coordinate lines will represent only the significant genes (those that pass threshVal

for threshVar)

nC INTEGER | Number of clusters; default 4

colList CHARACTER ARRAY | List of colors for each cluster; default is rainbow(nC)

aggMethod CHARACTER STRING ["ward.D" | "ward.D2" | "single" | "complete" | "aver-

age" | "mcquitty" | "median" | "centroid"] | The agglomeration method to be used

in the hierarchical clustering; default "ward.D"

yAxisLabel CHARACTER STRING | Vertical axis label; default "Count"

xAxisLabel CHARACTER STRING | Horizontal axis label; default "Sample"

1ineSize INTEGER | Size of plotted parallel coordinate lines; default 0.1

lineAlpha INTEGER | Alpha value of plotted parallel coordinate lines, default 0.5

vxAxis BOOLEAN [TRUE | FALSE] | Flip x-axis text labels to vertical orientation;

default FALSE

outDir CHARACTER STRING | Output directory to save all images; default tempdir()

saveFile BOOLEAN [TRUE | FALSE] | Save file to outDir; default TRUE

verbose

BOOLEAN [TRUE | FALSE] | Print each cluster from each cluster size into separate files and print the associated IDs of each cluster from each cluster size into separate .rds files; default is FALSE

#### Value

List of n elements each containing a grid of parallel coordinate plots, where n is the number of treatment pair combinations in the data object. If the saveFile parameter has a value of TRUE, then each grid of parallel coordinate plots is saved to the location specified in the outDir parameter as a JPG file. If the verbose parameter has a value of TRUE, then a JPG file for each parallel coordinate plot in each grid, RDS file containing the superimposed IDs for each parallel coordinate plot in each grid, and the JPG file of each grid of parallel coordinate plots is saved to the location specified in the outDir parameter.

#### See Also

hclust https://lindsayrutter.github.io/bigPint/articles/clusters.html

```
# The first set of five examples use data and dataMetrics
# objects as input. The last set of five examples create the same plots now
# using the SummarizedExperiment (i.e. dataSE) object input.
# Example 1: Perform hierarchical clustering of size four using the
# default agglomeration method "ward.D". Cluster only on the genes that have
\# FDR < 1e-7 (n = 113) and overlay these genes.
library(grid)
library(matrixStats)
library(ggplot2)
data(soybean_ir_sub)
soybean_ir_sub[,-1] <- log(soybean_ir_sub[-1]+1)</pre>
data(soybean_ir_sub_metrics)
colList = c("#00A600FF", rainbow(5)[c(1,4,5)])
ret <- plotClusters(data=soybean_ir_sub,</pre>
    dataMetrics = soybean_ir_sub_metrics, nC=4, colList = colList,
    clusterAllData = FALSE, threshVal = 1e-7, saveFile = FALSE)
grid.draw(ret[["N_P_4"]])
# Example 2: Perform the same analysis, only now create the four groups by
\# clustering on all genes in the data (n = 5,604). Then, overlay the genes
\# that have FDR < 1e-7 (n = 113) into their corresponding clusters.
ret <- plotClusters(data=soybean_ir_sub,</pre>
    dataMetrics = soybean_ir_sub_metrics, nC=4, colList = colList,
    clusterAllData = TRUE, threshVal = 1e-7, saveFile = FALSE)
grid.draw(ret[["N_P_4"]])
# Example 3: Perform the same analysis, only now overlay all genes in the
# data by keeping the dataMetrics object as its default value of NULL.
ret <- plotClusters(data=soybean_ir_sub, nC=4, colList = colList,</pre>
    clusterAllData = TRUE, saveFile = FALSE)
grid.draw(ret[["N_P_4"]])
```

```
# Example 4: Visualization of gene clusters is usually performed on
# standardized data. Here, hierarchical clustering of size four is performed
# using the agglomeration method "average" on standardized data. Only genes
# with FDR < 0.05 are used for the clustering. Only two of the three
# pairwise combinations of treatment groups (S1 and S2; S1 and S3) have any
\# genes with FDR < 0.05. The output plots for these two pairs are examined.
data(soybean_cn_sub)
data(soybean_cn_sub_metrics)
soybean_cn_sub_st <- as.data.frame(t(apply(as.matrix(soybean_cn_sub[,-1]),</pre>
soybean_cn_sub_st$ID <- as.character(soybean_cn_sub$ID)</pre>
soybean_cn_sub_st <- soybean_cn_sub_st[,c(length(soybean_cn_sub_st),</pre>
    1:length(soybean_cn_sub_st)-1)]
colnames(soybean_cn_sub_st) <- colnames(soybean_cn_sub)</pre>
nID <- which(is.nan(soybean_cn_sub_st[,2]))</pre>
soybean_cn_sub_st[nID,2:length(soybean_cn_sub_st)] <- 0</pre>
ret <- plotClusters(data=soybean_cn_sub_st,</pre>
    dataMetrics = soybean_cn_sub_metrics, nC=4,
    colList = c("#00A600FF", "#CC00FFFF", "red", "darkorange"),
    lineSize = 0.5, lineAlpha = 1, clusterAllData = FALSE,
    aggMethod = "average", yAxisLabel = "Standardized read count",
    saveFile = FALSE)
names(ret)
grid.draw(ret[["S1_S2_4"]])
grid.draw(ret[["S1_S3_4"]])
# Example 5: Run the same analysis, only now set the verbose parameter to
# value TRUE. This will save images of each individual cluster, .rds files
# that contain the IDs within each cluster, and images of the conglomerate
# clusters to outDir (default tempdir()).
## Not run:
plotClusters(data=soybean_cn_sub_st, dataMetrics = soybean_cn_sub_metrics,
  nC=4, colList = c("#00A600FF", "#CC00FFFF", "red", "darkorange"),
  lineSize = 0.5, lineAlpha = 1, clusterAllData = FALSE,
  aggMethod = "average", yAxisLabel = "Standardized read count",
  verbose = TRUE)
## End(Not run)
# Below are the same five examples, only now using the
# SummarizedExperiment (i.e. dataSE) object as input.
# Example 1: Perform hierarchical clustering of size four using the
# default agglomeration method "ward.D". Cluster only on the genes that have
\# FDR < 1e-7 (n = 113) and overlay these genes.
## Not run:
library(grid)
library(matrixStats)
library(ggplot2)
data(se_soybean_ir_sub)
assay(se_soybean_ir_sub) <- log(as.data.frame(assay(se_soybean_ir_sub))+1)</pre>
colList = c("#00A600FF", rainbow(5)[c(1,4,5)])
ret <- plotClusters(dataSE=se_soybean_ir_sub, nC=4, colList = colList,</pre>
    clusterAllData = FALSE, threshVal = 1e-7, saveFile = FALSE)
```

```
grid.draw(ret[["N_P_4"]])
## End(Not run)
## Not run:
# Example 2: Perform the same analysis, only now create the four groups by
\# clustering on all genes in the data (n = 5,604). Then, overlay the genes
# that have FDR < 1e-7 (n = 113) into their corresponding clusters.
ret <- plotClusters(dataSE=se_soybean_ir_sub, nC=4, colList = colList,</pre>
    clusterAllData = TRUE, threshVal = 1e-7, saveFile = FALSE)
grid.draw(ret[["N_P_4"]])
## End(Not run)
# Example 3: Perform the same analysis, only now overlay all genes in the
# data by setting the rowData() to NULL.
## Not run:
se_soybean_ir_sub_nm <- se_soybean_ir_sub</pre>
rowData(se_soybean_ir_sub_nm) <- NULL</pre>
ret <- plotClusters(dataSE=se_soybean_ir_sub_nm, nC=4, colList = colList,</pre>
    clusterAllData = TRUE, saveFile = FALSE)
grid.draw(ret[["N_P_4"]])
## End(Not run)
# Example 4: Visualization of gene clusters is usually performed on
# standardized data. Here, hierarchical clustering of size four is performed
# using the agglomeration method "average" on standardized data. Only genes
# with FDR < 0.05 are used for the clustering. Only two of the three
# pairwise combinations of treatment groups (S1 and S2; S1 and S3) have any
\# genes with FDR < 0.05. The output plots for these two pairs are examined.
## Not run:
data(se_soybean_cn_sub)
se_soybean_cn_sub_st = se_soybean_cn_sub
assay(se_soybean_cn_sub_st) <-as.data.frame(t(apply(as.matrix(as.data.frame(</pre>
    assay(se_soybean_cn_sub))), 1, scale)))
nID <- which(is.nan(as.data.frame(assay(se_soybean_cn_sub_st))[,1]))</pre>
assay(se_soybean_cn_sub_st)[nID,] <- 0</pre>
ret <- plotClusters(dataSE=se_soybean_cn_sub_st, nC=4,</pre>
    colList = c("#00A600FF", "#CC00FFFF", "red", "darkorange"),
    lineSize = 0.5, lineAlpha = 1, clusterAllData = FALSE,
    aggMethod = "average", yAxisLabel = "Standardized read count",
    saveFile = FALSE)
names(ret)
grid.draw(ret[["S1_S2_4"]])
grid.draw(ret[["S1_S3_4"]])
## End(Not run)
# Example 5: Run the same analysis, only now set the verbose parameter to
# value TRUE. This will save images of each individual cluster, .rds files
# that contain the IDs within each cluster, and images of the conglomerate
# clusters to outDir (default tempdir()).
```

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plotLitre

Plot static litre plots

# Description

Plot static litre plots.

# Usage

```
plotLitre(
   data = data,
   dataMetrics = NULL,
   dataSE = NULL,
   geneList = NULL,
   threshVar = "FDR",
   threshVal = 0.05,
   option = c("hexagon", "allPoints"),
   pointSize = 2,
   pointColor = "orange",
   xbins = 10,
   outDir = tempdir(),
   saveFile = TRUE
)
```

## **Arguments**

data	DATA FRAME   Read counts
dataMetrics	LIST   Differential expression metrics; default NULL
dataSE	SUMMARIZEDEXPERIMENT   Summarized experiment format that can be used in lieu of data and dataMetrics; default NULL
geneList	CHARACTER ARRAY   List of ID values of genes to be drawn from data as litre plots. Use this parameter if you have predetermined genes to be drawn. Otherwise, use dataMetrics, threshVar, and threshVal to create genes to be drawn; default NULL
threshVar	CHARACTER STRING   Name of column in dataMetrics object that is used to threshold significance; default "FDR"
threshVal	INTEGER   Maximum value to threshold significance from threshVar object; default $0.05$

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option	CHARACTER STRING ["hexagon"   "allPoints"]   The background of plot; default "hexagon"
pointSize	INTEGER   Size of plotted points; default 2
pointColor	CHARACTER STRING   Color of gene superimposed on litre plot; default "orange"
xbins	INTEGER   Number of bins partitioning the range of the plot; default 10
outDir	CHARACTER STRING   Output directory to save all plots; default tempdir()
saveFile	BOOLEAN [TRUE   FALSE]   Save file to outDir; default TRUE

#### Value

List of n elements of litre plots, where n is the number of genes determined to be superimposed through the dataMetrics or geneList parameter. If the saveFile parameter has a value of TRUE, then each of these litre plots is saved to the location specified in the outDir parameter as a JPG file.

```
# The first set of three examples use data and dataMetrics
# objects as input. The last set of three examples create the same plots now
# using the SummarizedExperiment (i.e. dataSE) object input.
# Example 1: Create litre plots for each of the 61 genes with FDR < 1e-10.
# Examine the first plot (gene "N_P_Glyma.19G168700.Wm82.a2.v1")
data(soybean_ir_sub)
soybean_ir_sub[,-1] <- log(soybean_ir_sub[,-1]+1)</pre>
data(soybean_ir_sub_metrics)
ret <- plotLitre(data = soybean_ir_sub,</pre>
    dataMetrics = soybean_ir_sub_metrics, threshVal = 1e-10,
    saveFile = FALSE)
length(ret)
names(ret)[1]
ret[[1]]
# Example 2: Create litre plots for each of the five most significant genes
\# (low FDR values). View plot for gene "N_P_Glyma.19G168700.Wm82.a2.v1".
geneList = soybean_ir_sub_metrics[["N_P"]][1:5,]$ID
ret <- plotLitre(data = soybean_ir_sub, geneList = geneList,</pre>
    pointColor = "deeppink")
names(ret)
ret[["N_P_Glyma.19G168700.Wm82.a2.v1"]]
# Example 3: Create one litre plot for each of the five most significant
# genes (low FDR values). View the plot for gene
\# "N_P_Glyma.19G168700.Wm82.a2.v1". Use points instead of the default
# hexagons as the background.
ret <- plotLitre(data = soybean_ir_sub, geneList = geneList,</pre>
    pointColor = "deeppink", option = "allPoints")
names(ret)
ret[["N_P_Glyma.19G168700.Wm82.a2.v1"]]
# Below are the same three examples, only now using the
# SummarizedExperiment (i.e. dataSE) object as input.
```

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```
# Example 1: Create litre plots for each of the 61 genes with FDR < 1e-10.
# Examine the first plot (gene "N_P_Glyma.19G168700.Wm82.a2.v1")
## Not run:
data(se_soybean_ir_sub)
assay(se\_soybean\_ir\_sub) <- log(as.data.frame(assay(se\_soybean\_ir\_sub))+1)
ret <- plotLitre(dataSE = se_soybean_ir_sub, threshVal = 1e-10,</pre>
    saveFile = FALSE)
length(ret)
names(ret)[1]
ret[[1]]
## End(Not run)
# Example 2: Create litre plots for each of the five most significant genes
# (low FDR values). View plot for gene "N_P_Glyma.19G168700.Wm82.a2.v1".
## Not run:
geneList <- as.data.frame(rowData(se_soybean_ir_sub)) %>%
    arrange(N_P.FDR) %>% filter(row_number() <= 5)</pre>
geneList <- geneList[,1]</pre>
ret <- plotLitre(dataSE = se_soybean_ir_sub, geneList = geneList,</pre>
    pointColor = "deeppink")
names(ret)
ret[["N_P_Glyma.19G168700.Wm82.a2.v1"]]
## End(Not run)
# Example 3: Create one litre plot for each of the five most significant
# genes (low FDR values). View the plot for gene
# "N_P_Glyma.19G168700.Wm82.a2.v1". Use points instead of the default
# hexagons as the background.
## Not run:
ret <- plotLitre(dataSE = se_soybean_ir_sub, geneList = geneList,</pre>
    pointColor = "deeppink", option = "allPoints")
names(ret)
ret[["N_P_Glyma.19G168700.Wm82.a2.v1"]]
## End(Not run)
```

plotLitreApp

Plot interactive litre plots

## **Description**

Plot interactive litre plots.

# Usage

```
plotLitreApp(
  data = data,
```

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```
dataMetrics = dataMetrics,
dataSE = NULL,
geneList = NULL,
pointColor = "orange",
option = c("hexagon", "allPoints")
```

#### **Arguments**

data DATA FRAME | Read counts

dataMetrics LIST | Differential expression metrics

dataSE SUMMARIZEDEXPERIMENT | Summarized experiment format that can be

used in lieu of data; default NULL

geneList CHARACTER ARRAY | List of gene IDs to be drawn onto the litre. Use this

parameter if you have predetermined subset of genes to be drawn. Otherwise, all genes in the data object can be superimposed on the litre plot; default NULL

pointColor CHARACTER STRING | Color of overlaid points on scatterplot matrix; default

"orange"

option CHARACTER STRING ["hexagon" | "allPoints"] | The background of plot; de-

fault "hexagon"; "allPoints" may be too slow depending on data

#### Value

A Shiny application that shows a litre plot background and allows users to superimpose the subset of genes determined to be superimposed through the dataMetrics or geneList parameter. The application allows users to order how to sequentially superimpose the genes by columns in the dataMetrics parameter.

```
# The first pair of examples use data and dataMetrics
# objects as input. The last pair of examples create the same plots now
# using the SummarizedExperiment (i.e. dataSE) object input.
# Example 1: Create an interactive litre plot for the logged data using
# default background of hexagons.
data(soybean_ir_sub)
data(soybean_ir_sub_metrics)
soybean_ir_sub_log <- soybean_ir_sub</pre>
soybean_ir_sub_log[,-1] <- log(soybean_ir_sub[,-1]+1)</pre>
app <- plotLitreApp(data = soybean_ir_sub_log,</pre>
    dataMetrics = soybean_ir_sub_metrics)
if (interactive()) {
    shiny::runApp(app, port = 1234, launch.browser = TRUE)
# Example 2: Repeat the same process, only now plot background data as
# individual points. Note this may be too slow now that all points are drawn
# in the background.
app <- plotLitreApp(data = soybean_ir_sub_log,</pre>
    dataMetrics = soybean_ir_sub_metrics, option = "allPoints",
    pointColor = "red")
```

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```
if (interactive()) {
    shiny::runApp(app)
# Below are the same pair of examples, only now using the
# SummarizedExperiment (i.e. dataSE) object as input.
# Example 1: Create an interactive litre plot for the logged data using
# default background of hexagons.
## Not run:
data(se_soybean_ir_sub)
se_soybean_ir_sub_log <- se_soybean_ir_sub</pre>
assay(se_soybean_ir_sub_log) <-</pre>
   log(as.data.frame(assay(se_soybean_ir_sub_log))+1)
app <- plotLitreApp(dataSE = se_soybean_ir_sub_log)</pre>
if (interactive()) {
    shiny::runApp(app, port = 1234, launch.browser = TRUE)
}
## End(Not run)
# Example 2: Repeat the same process, only now plot background data as
# individual points. Note this may be too slow now that all points are
# drawn in the background.
## Not run:
app <- plotLitreApp(dataSE = se_soybean_ir_sub_log, option = "allPoints",</pre>
    pointColor = "red")
if (interactive()) {
    shiny::runApp(app)
## End(Not run)
```

plotPCP

Plot static parallel coordinate plots

#### Description

Plot static parallel coordinate plots onto side-by-side boxplot of whole dataset.

## Usage

```
plotPCP(
  data,
  dataMetrics = NULL,
  dataSE = NULL,
  geneList = NULL,
  threshVar = "FDR",
  threshVal = 0.05,
  lineSize = 0.1,
  lineColor = "orange",
```

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```
vxAxis = FALSE,
outDir = tempdir(),
saveFile = TRUE,
hover = FALSE
)
```

## **Arguments**

data	DATA FRAME   Read counts
dataMetrics	LIST   Differential expression metrics; If both geneList and dataMetrics are NULL, then no genes will be overlaid onto the side-by-side boxplot; default NULL
dataSE	SUMMARIZEDEXPERIMENT   Summarized experiment format that can be used in lieu of data and dataMetrics; default NULL
geneList	CHARACTER ARRAY   List of gene IDs to be drawn onto the scatterplot matrix of all data. If this parameter is defined, these will be the overlaid genes to be drawn. After that, dataMetrics, threshVar, and threshVal will be considered for overlaid genes. If both geneList and dataMetrics are NULL, then no genes will be overlaid onto the side-by-side boxplot; default NULL
threshVar	CHARACTER STRING   Name of column in dataMetrics object that is used to threshold significance; default "FDR"
threshVal	INTEGER   Maximum value to threshold significance from threshVar object; default $0.05$
lineSize	INTEGER   Line width of parallel coordinate lines; default 0.1
lineColor	CHARACTER STRING   Color of parallel coordinate lines; default "orange"
vxAxis	BOOLEAN [TRUE   FALSE]   Flip x-axis text labels to vertical orientation; default FALSE
outDir	CHARACTER STRING   Output directory to save all plots; default tempdir()
saveFile	BOOLEAN [TRUE   FALSE]   Save file to outDir; default TRUE
hover	BOOLEAN [TRUE $\mid$ FALSE] $\mid$ Allow to hover over points to identify IDs; default FALSE

#### Value

List of n elements of parallel coordinate plots, where n is the number of treatment pair combinations in the data object. The background of each plot is a side-by-side boxplot of the full data object, and the parallel coordinate lines on each plot are the subset of genes determined to be superimposed through the dataMetrics or geneList parameter. If the saveFile parameter has a value of TRUE, then each parallel coordinate plot is saved to the location specified in the outDir parameter as a JPG file.

```
# The first set of four examples use data and dataMetrics
# objects as input. The last set of four examples create the same plots now
# using the SummarizedExperiment (i.e. dataSE) object input.

# Example 1: Plot the side-by-side boxplots of the whole dataset without
# overlaying any metrics data by keeping the dataMetrics parameter its
# default value of NULL.
```

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```
data(soybean_ir_sub)
soybean_ir_sub[,-1] = log(soybean_ir_sub[,-1] + 1)
ret <- plotPCP(data = soybean_ir_sub, saveFile = FALSE)</pre>
ret[[1]]
# Example 2: Overlay genes with FDR < 1e-4 as orange parallel coordinate
# lines.
data(soybean_ir_sub_metrics)
ret <- plotPCP(data = soybean_ir_sub, dataMetrics = soybean_ir_sub_metrics,</pre>
    threshVal = 1e-4, saveFile = FALSE)
ret[[1]]
# Example 3: Overlay the ten most significant genes (lowest FDR values) as
# blue parallel coordinate lines.
geneList = soybean_ir_sub_metrics[["N_P"]][1:10,]$ID
ret <- plotPCP(data = soybean_ir_sub, geneList = geneList, lineSize = 0.3,</pre>
    lineColor = "blue", saveFile = FALSE)
ret[[1]]
# Example 4: Repeat this same procedure, only now set the hover parameter to
# TRUE to allow us to hover over blue parallel coordinate lines and
# determine their individual IDs.
ret <- plotPCP(data = soybean_ir_sub, geneList = geneList, lineSize = 0.3,</pre>
    lineColor = "blue", saveFile = FALSE, hover = TRUE)
ret[[1]]
# Below are the same four examples, only now using the
# SummarizedExperiment (i.e. dataSE) object as input.
# Example 1: Plot the side-by-side boxplots of the whole dataset without
# overlaying any metrics. We prevent overlaying metrics by setting the
# rowData() to NULL.
## Not run:
data(se_soybean_ir_sub)
se_soybean_ir_sub[,-1] <- log(se_soybean_ir_sub[,-1]+1)</pre>
se_soybean_ir_sub_nm <- se_soybean_ir_sub</pre>
rowData(se_soybean_ir_sub_nm) <- NULL</pre>
ret <- plotPCP(dataSE = se_soybean_ir_sub_nm, saveFile = FALSE)</pre>
ret[[1]]
## End(Not run)
## Not run:
# Example 2: Overlay genes with FDR < 1e-4 as orange parallel coordinate
# lines.
ret <- plotPCP(dataSE = se_soybean_ir_sub, threshVal = 1e-4,
    saveFile = FALSE)
ret[[1]]
## End(Not run)
# Example 3: Overlay the ten most significant genes (lowest FDR values) as
```

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```
# blue parallel coordinate lines.
## Not run:
geneList <- as.data.frame(rowData(se_soybean_ir_sub)) %>%
    arrange(N_P.FDR) %>% filter(row_number() <= 10)</pre>
geneList <- geneList[,1]</pre>
ret <- plotPCP(dataSE = se_soybean_ir_sub, geneList = geneList,</pre>
    lineSize = 0.3, lineColor = "blue", saveFile = FALSE)
ret[[1]]
## End(Not run)
# Example 4: Repeat this same procedure, only now set the hover parameter to
# TRUE to allow us to hover over blue parallel coordinate lines and
# determine their individual IDs.
## Not run:
ret <- plotPCP(data = soybean_ir_sub, geneList = geneList, lineSize = 0.3,</pre>
    lineColor = "blue", saveFile = FALSE, hover = TRUE)
ret[[1]]
## End(Not run)
```

plotPCPApp

Plot interactive parallel coordinate plots

# Description

Plot interactive parallel coordinate plots.

## Usage

```
plotPCPApp(data = data, dataSE = NULL, pointColor = "orange")
```

## **Arguments**

data DATA FRAME | Read counts for parallel coordinate lines

dataSE SUMMARIZEDEXPERIMENT | Summarized experiment format that can be

used in lieu of data; default NULL

pointColor CHARACTER STRING | Color of overlaid points on scatterplot matrix; default

"orange"

#### Value

A Shiny application that shows a parallel coordinate plot and allows users to draw rectangular areas across samples and remove genes that are not inside these areas. The user can download a file that contains the gene IDs that remain.

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```
# The first example uses data and dataMetrics objects as
# input. The last example creates the same plot now using the
# SummarizedExperiment (i.e. dataSE) object input.
# Example: Create interactive parallel coordinate plot for genes that have
\# FDR < 0.01 and logFC < -4. Standardize genes to have an average of zero
# and a standard deviation of one.
data(soybean_ir_sub)
data(soybean_ir_sub_metrics)
# Create standardized version of data
library(matrixStats)
soybean_ir_sub_st = as.data.frame(t(apply(as.matrix(soybean_ir_sub[,-1]), 1,
soybean_ir_sub_st$ID = as.character(soybean_ir_sub$ID)
soybean_ir_sub_st[,c(length(soybean_ir_sub_st),
        1:length(soybean_ir_sub_st)-1)]
colnames(soybean_ir_sub_st) = colnames(soybean_ir_sub)
nID = which(is.nan(soybean_ir_sub_st[,2]))
soybean_ir_sub_st[nID,2:length(soybean_ir_sub_st)] = 0
library(dplyr, warn.conflicts = FALSE)
plotGenes = filter(soybean_ir_sub_metrics[["N_P"]], FDR < 0.01,</pre>
        logFC < -4) %>% select(ID)
pcpDat = filter(soybean_ir_sub_st, ID %in% plotGenes[,1])
app <- plotPCPApp(data = pcpDat, pointColor = "purple")</pre>
if (interactive()) {
        shiny::runApp(app, display.mode = "normal")
# Below is the same example, only now using the
# SummarizedExperiment (i.e. dataSE) object as input.
# Example: Create interactive parallel coordinate plot for genes that have
\# FDR < 0.01 and logFC < -4. Standardize genes to have an average of zero
# and a standard deviation of one.
## Not run:
data(se_soybean_ir_sub)
# Create standardized version of data
library(matrixStats)
se_soybean_ir_sub_st = se_soybean_ir_sub
assay(se\_soybean\_ir\_sub\_st) < -as.data.frame(t(apply(as.matrix(as.data.frame(t(apply(as.matrix(as.data.frame(t(apply(as.matrix(as.data.frame(t(apply(as.matrix(as.data.frame(t(apply(as.matrix(as.data.frame(t(apply(as.matrix(as.data.frame(t(apply(as.matrix(as.data.frame(t(apply(as.matrix(as.data.frame(t(apply(as.matrix(as.data.frame(t(apply(as.matrix(as.data.frame(t(apply(as.matrix(as.data.frame(t(apply(as.matrix(as.data.frame(t(apply(as.matrix(as.data.frame(t(apply(as.matrix(as.data.frame(t(apply(as.data.frame(t(apply(as.data.frame(t(apply(as.data.frame(t(apply(as.data.frame(t(apply(as.data.frame(t(apply(as.data.frame(t(apply(as.data.frame(t(apply(as.data.frame(t(apply(as.data.frame(t(apply(as.data.frame(t(apply(as.data.frame(t(apply(as.data.frame(t(apply(as.data.frame(t(apply(as.data.frame(t(apply(as.data.frame(t(apply(as.data.frame(t(apply(as.data.frame(t(apply(as.data.frame(t(apply(as.data.frame(t(apply(as.data.frame(t(apply(as.data.frame(t(apply(as.data.frame(t(apply(as.data.frame(t(apply(as.data.frame(t(apply(as.data.frame(t(apply(as.data.frame(t(apply(as.data.frame(t(apply(as.data.frame(t(apply(as.data.frame(t(apply(as.data.frame(t(apply(as.data.frame(t(apply(as.data.frame(t(apply(as.data.frame(t(apply(as.data.frame(t(apply(as.data.frame(t(apply(as.data.frame(t(apply(as.data.frame(t(apply(as.data.frame(t(apply(as.data.frame(t(apply(as.data.frame(t(apply(as.data.frame(t(apply(as.data.frame(t(apply(as.data.frame(t(apply(as.data.frame(t(apply(as.data.frame(t(apply(as.data.frame(t(apply(as.data.frame(t(apply(as.data.frame(t(apply(a).frame(t(apply(as.data.frame(t(apply(a).frame(t(apply(as.data.frame(t(apply(as.data.frame(t(apply(as.data.frame(t(apply(as.data.frame(t(apply(a).frame(t(apply(a).frame(t(apply(a).frame(t(apply(a).frame(t(apply(a).frame(t(a).frame(t(a).frame(t(a).frame(t(a).frame(t(a).frame(t(a).frame(t(a).frame(t(a).frame(t(a).frame(t(a).frame(t(a).frame(t(a).frame(t(a).frame(t(a).frame(t(a).frame(t(a).frame(t(a).frame(t(a).frame(t(a).frame(t(a).frame(t(a).frame(t(a).frame(t(a).frame(t(a).frame(t(
        assay(se_soybean_ir_sub))), 1, scale)))
nID <- which(is.nan(as.data.frame(assay(se_soybean_ir_sub_st))[,1]))</pre>
assay(se_soybean_ir_sub_st)[nID,] <- 0</pre>
# To subset our SummarizedExperiment data by a list of genes, we can
# invoke the convertSESubsetGenes() method.
library(dplyr, warn.conflicts = FALSE)
geneList <- as.data.frame(rowData(se_soybean_ir_sub_st)) %>%
```

```
filter(N_P.FDR <= 0.01) %>% filter(N_P.logFC < -4)
geneList <- geneList[,1]
pcpDat <- convertSESubsetGenes(se_soybean_ir_sub_st, geneList)
app <- plotPCPApp(dataSE = pcpDat, pointColor = "purple")
if (interactive()) {
    shiny::runApp(app, display.mode = "normal")
}
## End(Not run)</pre>
```

plotSM

Plot static scatterplot matrices

#### **Description**

Plot static scatterplot matrix. Optionally, superimpose differentially expressed genes (DEGs) onto scatterplot matrix.

## Usage

```
plotSM(
 data = data,
 dataMetrics = NULL,
 dataSE = NULL,
  geneList = NULL,
  threshVar = "FDR",
  threshVal = 0.05,
 option = c("allPoints", "foldChange", "orthogonal", "hexagon"),
  xbins = 10,
  threshFC = 3,
  threshOrth = 3,
  pointSize = 0.5,
 pointColor = "orange",
 outDir = tempdir(),
  saveFile = TRUE
)
```

# **Arguments**

data

DATA FRAME | Read counts

LIST | Differential expression metrics; default NULL

dataSE

SUMMARIZEDEXPERIMENT | Summarized experiment format that can be used in lieu of data and dataMetrics; default NULL

geneList

CHARACTER ARRAY | List of gene IDs to be drawn onto the scatterplot matrix of all data. Use this parameter if you have predetermined genes to be drawn. Otherwise, use dataMetrics, threshVar, and threshVal to create genes to be drawn onto the scatterplot matrix; default NULL; used in "hexagon" and "allPoints"

threshVar

CHARACTER STRING | Name of column in dataMetrics object that is used to threshold significance; default "FDR"; used in all options

threshVal	INTEGER   Maximum value to threshold significance from threshVar object;
	default 0.05; used in all options
option	CHARACTER STRING ["foldChange"   "orthogonal"   "hexagon"   "allPoints"]   The type of plot; default "allPoints"
xbins	INTEGER   Number of bins partitioning the range of the plot; default 10; used in option "hexagon"
threshFC	INTEGER   Threshold of fold change; default 3; used in option "foldChange"
threshOrth	INTEGER   Threshold of orthogonal distance; default 3; used in option "orthogonal"
pointSize	INTEGER   Size of plotted points; default 0.5; used for DEGs in "hexagon" and "allPoints" and used for all points in "foldChange" and "orthogonal"
pointColor	CHARACTER STRING   Color of overlaid points on scatterplot matrix; default "orange"; used for DEGs in "hexagon" and "allPoints" and used for all points in "foldChange" and "orthogonal"
outDir	CHARACTER STRING   Output directory to save all plots; default tempdir(); used in all options
saveFile	BOOLEAN [TRUE   FALSE]   Save file to outDir; default TRUE; used in all options

#### **Details**

There are seven options:

- "foldChange": Plots DEGs onto scatterplot matrix of fold changes
- "orthogonal": Plots DEGs onto scatterplot matrix of orthogonal distance
- "hexagon": Plot DEGs onto scatterplot matrix of hexagon binning
- "allPoints": Plot DEGs onto scatterplot matrix of all data points

#### Value

List of n elements of scatterplot matrices, where n is the number of treatment pair combinations in the data object. The subset of genes that are superimposed are determined through the dataMetrics or geneList parameter. If the saveFile parameter has a value of TRUE, then each of these scatterplot matrices is saved to the location specified in the outDir parameter as a JPG file.

```
# The first set of six examples use data and dataMetrics
# objects as input. The last set of six examples create the same plots now
# using the SummarizedExperiment (i.e. dataSE) object input.

# Read in data and metrics (need for first set of six examples)
data(soybean_cn_sub)
data(soybean_cn_sub_metrics)
data(soybean_ir_sub)
data(soybean_ir_sub_metrics)

# Create standardized version of data (need for first set of six examples)
library(matrixStats)
library(ggplot2)
```

```
soybean_cn_sub_st <- as.data.frame(t(apply(as.matrix(soybean_cn_sub[,-1]),</pre>
    1, scale)))
soybean_cn_sub_st$ID <- as.character(soybean_cn_sub$ID)</pre>
soybean_cn_sub_st <- soybean_cn_sub_st[,c(length(soybean_cn_sub_st),</pre>
    1:length(soybean_cn_sub_st)-1)]
colnames(soybean_cn_sub_st) <- colnames(soybean_cn_sub)</pre>
nID <- which(is.nan(soybean_cn_sub_st[,2]))</pre>
soybean_cn_sub_st[nID,2:length(soybean_cn_sub_st)] <- 0</pre>
# Example 1: Plot scatterplot matrix of points. Saves three plots to outDir
# because saveFile equals TRUE by default.
## Not run:
plotSM(soybean_cn_sub, soybean_cn_sub_metrics)
## End(Not run)
# Example 2: Plot scatterplot matrix of points. Return list of plots so user
# can tailor them (such as add title) and does not save to outDir because
# saveFile equals FALSE.
ret <- plotSM(soybean_cn_sub, soybean_cn_sub_metrics, pointColor = "pink",</pre>
    saveFile = FALSE)
# Determine names of plots in returned list
names(ret)
ret[["S1_S2"]] + ggtitle("S1 versus S2")
ret[["S1_S3"]] + ggtitle("S1 versus S3")
ret[["S2_S3"]] + ggtitle("S2 versus S3")
# Example 3: Plot standardized data as scatterplot matrix of points.
ret <- plotSM(soybean_cn_sub_st, soybean_cn_sub_metrics,</pre>
    pointColor = "#00C379", saveFile = FALSE)
ret[[1]] + xlab("Standardized read counts") +
ylab("Standardized read counts")
# Example 4: Plot scatterplot matrix of hexagons.
ret <- plotSM(soybean_cn_sub, soybean_cn_sub_metrics, option = "hexagon",</pre>
    xbins = 5, pointSize = 0.1, saveFile = FALSE)
ret[[2]]
# Example 5: Plot scatterplot matrix of orthogonal distance on the logged
# data, first without considering the metrics dataset and then considering
# it.
soybean_ir_sub[,-1] <- log(soybean_ir_sub[,-1] + 1)</pre>
ret <- plotSM(soybean_ir_sub, option = "orthogonal", threshOrth = 2.5,</pre>
    pointSize = 0.2, saveFile = FALSE)
ret <- plotSM(soybean_ir_sub, soybean_ir_sub_metrics, option = "orthogonal",</pre>
    threshOrth = 2.5, pointSize = 0.2, saveFile = FALSE)
ret[[1]]
# Example 6: Plot scatterplot matrix of fold change.
ret <- plotSM(soybean_cn_sub, soybean_cn_sub_metrics, option = "foldChange",</pre>
```

```
threshFC = 0.5, pointSize = 0.2, saveFile = FALSE)
ret[[1]]
# Below are the same six examples, only now using the
# SummarizedExperiment (i.e. dataSE) object as input.
# Read in data and metrics (need for first set of six examples)
data(se_soybean_cn_sub)
data(se_soybean_ir_sub)
# Create standardized version of data (need for first set of six examples)
library(matrixStats)
library(ggplot2)
library(SummarizedExperiment)
se_soybean_cn_sub_st = se_soybean_cn_sub
assay(se_soybean_cn_sub_st, withDimnames=FALSE) <-as.data.frame(t(</pre>
    apply(as.matrix(as.data.frame(assay(se_soybean_cn_sub))), 1, scale)))
nID <- which(is.nan(as.data.frame(assay(se_soybean_cn_sub_st))[,1]))</pre>
assay(se_soybean_cn_sub_st, withDimnames=FALSE)[nID,] <- 0</pre>
# Example 1: Plot scatterplot matrix of points. Saves three plots to outDir
# because saveFile equals TRUE by default.
## Not run:
plotSM(dataSE = se_soybean_cn_sub)
## End(Not run)
# Example 2: Plot scatterplot matrix of points. Return list of plots so user
# can tailor them (such as add title) and does not save to outDir because
# saveFile equals FALSE.
## Not run:
ret <- plotSM(dataSE = se_soybean_cn_sub, pointColor = "pink",</pre>
    saveFile = FALSE)
# Determine names of plots in returned list
names(ret)
ret[["S1_S2"]] + ggtitle("S1 versus S2")
ret[["S1_S3"]] + ggtitle("S1 versus S3")
ret[["S2_S3"]] + ggtitle("S2 versus S3")
## End(Not run)
# Example 3: Plot standardized data as scatterplot matrix of points.
## Not run:
ret <- plotSM(dataSE = se_soybean_cn_sub_st, pointColor = "#00C379",</pre>
    saveFile = FALSE)
ret[[1]] + xlab("Standardized read counts") +
ylab("Standardized read counts")
## End(Not run)
# Example 4: Plot scatterplot matrix of hexagons.
## Not run:
ret <- plotSM(dataSE = se_soybean_cn_sub, option = "hexagon", xbins = 5,</pre>
```

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```
pointSize = 0.1, saveFile = FALSE)
ret[[2]]
## End(Not run)
# Example 5: Plot scatterplot matrix of orthogonal distance on the logged
# data, first without considering the metrics dataset and then considering
# it.
## Not run:
assay(se_soybean_ir_sub) <- log(as.data.frame(assay(se_soybean_ir_sub))+1)</pre>
ret <- plotSM(dataSE = se_soybean_ir_sub, option = "orthogonal",</pre>
    threshOrth = 2.5, pointSize = 0.2, saveFile = FALSE)
ret[[1]]
## End(Not run)
# Example 6: Plot scatterplot matrix of fold change.
## Not run:
ret <- plotSM(dataSE = se_soybean_cn_sub, option = "foldChange",</pre>
    threshFC = 0.5, pointSize = 0.2, saveFile = FALSE)
## End(Not run)
```

plotSMApp

Plot interactive scatterplot matrices

# Description

Plot interactive scatterplot matrices.

## Usage

```
plotSMApp(data = data, dataSE = NULL, xbins = 10)
```

#### **Arguments**

data DATA FRAME | Read counts

dataSE SUMMARIZEDEXPERIMENT | Summarized experiment format that can be

used in lieu of data; default NULL

xbins INTEGER | Number of bins partitioning the range of the plot; default 10

#### Value

A Shiny application that shows a scatterplot matrix with hexagon bins and allows users to click on hexagon bins to determine how many genes they each contain. The user can download a file that contains the gene IDs that are located in the clicked hexagon bin.

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#### **Examples**

```
# The first example uses data and dataMetrics objects as
# input. The last example creates the same plots now using the
# SummarizedExperiment (i.e. dataSE) object input.
# Example: Create interactive scatterplot matrix for first two treatment
# groups of data.
data(soybean_cn_sub)
soybean_cn_sub <- soybean_cn_sub[,1:7]</pre>
app <- plotSMApp(data=soybean_cn_sub)</pre>
if (interactive()) {
    shiny::runApp(app)
}
# Below are the same example, only now using the
# SummarizedExperiment (i.e. dataSE) object as input.
# Example: Create interactive scatterplot matrix for first two treatment
\mbox{\tt\#} groups of data. When working with the SummarizedExperiment data,
\mbox{\#} we can summon the method convertSEPair() to subset our input data
# to only a pair of treatment groups.
## Not run:
data(se_soybean_cn_sub)
se_soybean_cn_sub <- convertSEPair(se_soybean_cn_sub, "S1", "S2")</pre>
app <- plotSMApp(dataSE=se_soybean_cn_sub)</pre>
if (interactive()) {
    shiny::runApp(app)
}
## End(Not run)
```

plotVolcano

Plot static volcano plot

## **Description**

Plot static volcano plot.

# Usage

```
plotVolcano(
  data = data,
  dataMetrics = dataMetrics,
  dataSE = NULL,
  geneList = NULL,
  threshVar = "FDR",
  threshVal = 0.05,
  option = c("hexagon", "allPoints"),
  logFC = "logFC",
  PValue = "PValue",
```

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```
xbins = 10,
pointSize = 0.5,
pointColor = "orange",
outDir = tempdir(),
saveFile = TRUE,
hover = FALSE
)
```

## **Arguments**

data	DATA FRAME   Read counts
dataMetrics	LIST   Differential expression metrics. This object must contain one column with magnitude changes (for the logFC parameter) and one column with statistical values (for the PValue parameter), unless geneList is not NULL
dataSE	SUMMARIZEDEXPERIMENT   Summarized experiment format that can be used in lieu of data and dataMetrics; default NULL
geneList	CHARACTER ARRAY   List of gene IDs to be drawn onto the scatterplot matrix of all data. Use this parameter if you have predetermined subset of genes to be superimposed. Otherwise, dataMetrics, threshVar, and threshVal will be used to create genes to be superimposed onto the volcano plot; default NULL
threshVar	CHARACTER STRING   Name of column in dataMetrics object that is used to determine genes to be overlaid; default "FDR"
threshVal	INTEGER   Maximum value to threshold significance from threshVar object; default 0.05
option	CHARACTER STRING ["hexagon"   "allPoints"]   The background of plot; default "hexagon"
logFC	CHARACTER STRING   Name of column in dataMetrics object that contains log fold change values; default "logFC"
PValue	CHARACTER STRING   Name of column in dataMetrics object that contains p-values; default "PValue"
xbins	INTEGER   Number of bins partitioning the range of the plot; default 10
pointSize	INTEGER   Size of plotted points; default 0.5
pointColor	CHARACTER STRING   Color of overlaid points on scatterplot matrix; default "orange"
outDir	CHARACTER STRING   Output directory to save all plots; default tempdir()
saveFile	BOOLEAN [TRUE   FALSE]   Save file to outDir; default TRUE
hover	BOOLEAN [TRUE   FALSE]   Allow to hover over points to identify IDs; default FALSE

# Value

List of n elements of volcano plots, where n is the number of treatment pair combinations in the data object. The subset of genes that are superimposed are determined through the dataMetrics or geneList parameter. If the saveFile parameter has a value of TRUE, then each of these volcano plots is saved to the location specified in the outDir parameter as a JPG file.

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```
# The first set of four examples use data and dataMetrics objects as
# input. The last set of four examples create the same plots now
# using the SummarizedExperiment (i.e. dataSE) object input.
# Example 1: Plot volcano plot with default settings for overlaid points
\# (FDR < 0.05).
data(soybean_ir_sub)
data(soybean_ir_sub_metrics)
ret <- plotVolcano(soybean_ir_sub, soybean_ir_sub_metrics, pointSize = 1,</pre>
    saveFile = FALSE)
ret[[1]]
# Example 2: Plot volcano plot and overlay points with PValue < 1e-15.
ret <- plotVolcano(soybean_ir_sub, soybean_ir_sub_metrics,</pre>
    pointColor = "red", pointSize = 1, threshVar = "PValue",
    threshVal = 1e-15, saveFile = FALSE)
ret[[1]]
# Example 3: Plot volcano plot and overlay points with PValue < 1e-15. This
# time, plot all points (instead of hexagons) for the background.
ret <- plotVolcano(soybean_ir_sub, soybean_ir_sub_metrics,</pre>
    pointColor = "red", pointSize = 1, threshVar = "PValue"
    threshVal = 1e-15, option = "allPoints", saveFile = FALSE)
ret[[1]]
# Example 4: Plot volcano plot with points in background and overlay points
# with PValue < 1e-15. This time, use a value of TRUE for the hover
# parameter so that you can hover over overlaid points and determine their
# IDs.
ret <- plotVolcano(soybean_ir_sub, soybean_ir_sub_metrics,</pre>
    pointColor = "red", pointSize = 1, threshVar = "PValue",
    threshVal = 1e-15, option = "allPoints", saveFile = FALSE,
    hover = TRUE)
ret[[1]]
# Below are the same four examples, only now using the
# SummarizedExperiment (i.e. dataSE) object as input.
# Example 1: Plot volcano plot with default settings for overlaid points
\# (FDR < 0.05).
## Not run:
data(se_soybean_ir_sub)
ret <- plotVolcano(dataSE = se_soybean_ir_sub, pointSize = 1,</pre>
    saveFile = FALSE)
ret[[1]]
## End(Not run)
# Example 2: Plot volcano plot and overlay points with PValue < 1e-15.
```

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```
## Not run:
ret <- plotVolcano(dataSE = se_soybean_ir_sub, pointColor = "red",</pre>
    pointSize = 1, threshVar = "PValue", threshVal = 1e-15,
    saveFile = FALSE)
ret[[1]]
## End(Not run)
# Example 3: Plot volcano plot and overlay points with PValue < 1e-15. This
# time, plot all points (instead of hexagons) for the background.
## Not run:
ret <- plotVolcano(dataSE = se_soybean_ir_sub, pointColor = "red",</pre>
    pointSize = 1, threshVar = "PValue", threshVal = 1e-15,
    option = "allPoints", saveFile = FALSE)
ret[[1]]
## End(Not run)
# Example 4: Plot volcano plot with points in background and overlay points
# with PValue < 1e-15. This time, use a value of TRUE for the hover
# parameter so that you can hover overlaid points and determine their
# IDs.
## Not run:
ret <- plotVolcano(dataSE = se_soybean_ir_sub, pointColor = "red",</pre>
    pointSize = 1, threshVar = "PValue", threshVal = 1e-15,
    option = "allPoints", saveFile = FALSE, hover = TRUE)
ret[[1]]
## End(Not run)
```

plotVolcanoApp

Plot interactive volcano plots

# Description

Plot interactive volcano plots.

# Usage

```
plotVolcanoApp(
  data = data,
  dataMetrics = dataMetrics,
  dataSE = NULL,
  option = c("hexagon", "allPoints"),
  pointColor = "orange"
)
```

### **Arguments**

data

DATA FRAME | Read counts

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dataMetrics LIST | Differential expression metrics. This object must contain one column named "logFC" and one column named "PValue".

dataSE SUMMARIZEDEXPERIMENT | Summarized experiment format that can be used in lieu of data; default NULL

option CHARACTER STRING ["hexagon" | "allPoints"] | The background of plot; default "hexagon"

pointColor CHARACTER STRING | Color of overlaid points on scatterplot matrix; default "orange"

#### Value

A Shiny application that shows a volcano plot and allows users to overlay genes depending on two values, usually a statistical value (such as P-value) and a magnitude change value (such as log fold change). The user can download a file that contains the gene IDs that pass these thresholds.

```
# The first pair of examples use data and dataMetrics objects as input.
# The last pair of examples create the same plots now using the
# SummarizedExperiment (i.e. dataSE) object input.
# Example 1: Create interactive volcano plot of logged data using hexagon
# bins for the background.
data(soybean_cn_sub)
data(soybean_cn_sub_metrics)
app <- plotVolcanoApp(data = soybean_cn_sub,</pre>
    dataMetrics = soybean_cn_sub_metrics)
if (interactive()) {
    shiny::runApp(app)
}
# Example 2: Create interactive volcano plot of logged data using points for
# the background.
app <- plotVolcanoApp(data = soybean_cn_sub,</pre>
    dataMetrics = soybean_cn_sub_metrics, option = "allPoints",
    pointColor = "magenta")
if (interactive()) {
    shiny::runApp(app)
}
# Below is the same pair of examples, only now using the
# SummarizedExperiment (i.e. dataSE) object as input.
\mbox{\#} Example 1: Create interactive volcano plot of logged data using hexagon
# bins for the background.
## Not run:
data(se_soybean_cn_sub)
app <- plotVolcanoApp(dataSE = se_soybean_cn_sub)</pre>
if (interactive()) {
    shiny::runApp(app)
```

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se\_soybean\_cn\_sub

Normalized and subsetted soybean cotyledon SummarizedExperiment

### **Description**

This dataset contains normalized RNA-sequencing read counts from soybean cotyledon across three time stages of development. Early stage cotyledons were collected four days after planting and were green but closed. Middle stage cotyledons were collected while green and open, soon after the plant generated its first set of unifoliate leaves. Late stage cotyledons were collected immediately after the initiation of yellowing and shrinking. To save on size, this example dataset was generated by obtaining a random subset of 1 out of 10 genes from the original resource.

## Usage

```
data(se_soybean_cn_sub)
```

#### **Format**

- (1) A DelayedMatrix with 7,332 rows and 10 variables and 1 row per gene. Accessible by assay(se\_soybean\_cn\_sub).
- (2) A List with 1 list element per treatment group combination and 1 row per gene. Accessible by rowData(se\_soybean\_cn\_sub).

## **Details**

Normalized and subsetted soybean cotyledon SummarizedExperiment object

- assay(se\_soybean\_cn\_sub) Structure:
- S1.1 early stage replicate 1 normalized read counts
- S1.2 early stage replicate 2 normalized read counts
- S1.3 early stage replicate 3 normalized read counts
- S2.1 middle stage replicate 1 normalized read counts
- S2.2 middle stage replicate 2 normalized read counts
- S2.3 middle stage replicate 3 normalized read counts
- S3.1 late stage replicate 1 normalized read counts
- S3.2 late stage replicate 2 normalized read counts

se\_soybean\_ir\_sub

- S3.3 late stage replicate 3 normalized read counts
- rowData(se\_soybean\_cn\_sub) list element Structure:
- · ID gene name
- logFC log fold change
- logCPM log counts per million
- LR likelihood ratio
- PValue p-value
- FDR FDR value

#### References

Brown AV, Hudson KA (2015) Developmental profiling of gene expression in soybean trifoliate leaves and cotyledons. BMC Plant Biol 15:169

#### See Also

soybean\_cn\_sub and soybean\_cn\_sub\_metrics

se\_soybean\_ir\_sub

Raw and subsetted soybean leaves iron-metabolism SummarizedExperiment

# Description

This dataset contains raw RNA-sequencing read counts from a soybean dataset that compared leaves that were exposed to iron-rich (iron -postive) soil conditions versus leaves that were exposed to iron-poor (iron-negative) soil conditions. The data was collected 120 minutes after iron conditions were initiated. To save on size, this example dataset was generated by obtaining a random subset of 1 out of 10 genes from the original resource.

# Usage

```
data(se_soybean_ir_sub)
```

#### Format

- (1) A DelayedMatrix with 5,604 rows and 7 variables and 1 row per gene. Accessible by assay(se\_soybean\_ir\_sub).
- (2) A List with 1 list element per treatment group combination and 1 row per gene. Accessible by rowData(se\_soybean\_ir\_sub).

## **Details**

Raw and subsetted soybean leaves data SummarizedExperiment object

- assay(se\_soybean\_ir\_sub) Structure:
- N.1 iron-negative condition replicate 1 raw read counts
- N.2 iron-negative condition replicate 2 raw read counts
- N.3 iron-negative condition replicate 3 raw read counts

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- P.1 iron-positive condition replicate 1 raw read counts
- P.2 iron-positive condition replicate 2 raw read counts
- P.3 iron-positive condition replicate 3 raw read counts
- rowData(se\_soybean\_ir\_sub) list element Structure:
- · ID gene name
- logFC log fold change
- logCPM log counts per million
- · LR likelihood ratio
- PValue p-value
- FDR FDR value

#### References

Moran Lauter AN, Graham MA. NCBI SRA bioproject accession: PRJNA318409.

## See Also

soybean\_ir\_sub and soybean\_ir\_sub\_metrics

soybean\_cn

Normalized soybean cotyledon data

## **Description**

This dataset contains normalized RNA-sequencing read counts from soybean cotyledon across three time stages of development. Early stage cotyledons were collected four days after planting and were green but closed. Middle stage cotyledons were collected while green and open, soon after the plant generated its first set of unifoliate leaves. Late stage cotyledons were collected immediately after the initiation of yellowing and shrinking.

## **Format**

a RData instance, 1 row per gene

# Details

Normalized soybean cotyledon data

- · ID gene name
- S1.1 early stage replicate 1 normalized read counts
- S1.2 early stage replicate 2 normalized read counts
- S1.3 early stage replicate 3 normalized read counts
- S2.1 middle stage replicate 1 normalized read counts
- S2.2 middle stage replicate 2 normalized read counts
- S2.3 middle stage replicate 3 normalized read counts
- S3.1 late stage replicate 1 normalized read counts
  S3.2 late stage replicate 2 normalized read counts
- S3.3 late stage replicate 3 normalized read counts

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#### References

Brown AV, Hudson KA (2015) Developmental profiling of gene expression in soybean trifoliate leaves and cotyledons. BMC Plant Biol 15:169

soybean\_cn\_metrics

Normalized soybean cotyledon metrics

#### **Description**

This data contains metrics for normalized RNA-sequencing read counts from soybean cotyledon across three time stages of development. Early stage cotyledons were collected four days after planting and were green but closed. Middle stage cotyledons were collected while green and open, soon after the plant generated its first set of unifoliate leaves. Late stage cotyledons were collected immediately after the initiation of yellowing and shrinking. The metrics include the log fold change, log counts per million, likelihood ratio, p-values, and FDR values for all genes and all pairwise combinations of treatment groups.

#### **Format**

a RData instance, 1 list per treatment group combination and 1 row per gene

#### **Details**

Normalized soybean cotyledon metrics

- · ID gene name
- logFC log fold change
- logCPM log counts per million
- LR likelihood ratio
- PValue p-value
- FDR FDR value

# See Also

soybean\_cn for information about the treatment groups

soybean\_cn\_sub

Normalized and subsetted soybean cotyledon data

### **Description**

This dataset contains normalized RNA-sequencing read counts from soybean cotyledon across three time stages of development. Early stage cotyledons were collected four days after planting and were green but closed. Middle stage cotyledons were collected while green and open, soon after the plant generated its first set of unifoliate leaves. Late stage cotyledons were collected immediately after the initiation of yellowing and shrinking. To save on size, this example dataset was generated by obtaining a random subset of 1 out of 10 genes from the original resource.

## Usage

```
data(soybean_cn_sub)
```

#### **Format**

A data frame with 7,332 rows and 10 variables. 1 row per gene.

#### **Details**

Normalized and subsetted soybean cotyledon data

- ID gene name
- S1.1 early stage replicate 1 normalized read counts
- S1.2 early stage replicate 2 normalized read counts
- S1.3 early stage replicate 3 normalized read counts
- S2.1 middle stage replicate 1 normalized read counts
- S2.2 middle stage replicate 2 normalized read counts
- S2.3 middle stage replicate 3 normalized read counts
- S3.1 late stage replicate 1 normalized read counts
- S3.2 late stage replicate 2 normalized read counts
- S3.3 late stage replicate 3 normalized read counts

#### References

Brown AV, Hudson KA (2015) Developmental profiling of gene expression in soybean trifoliate leaves and cotyledons. BMC Plant Biol 15:169

#### See Also

soybean\_cn from which this dataset is subsetted

```
soybean_cn_sub_metrics
```

Normalized and subsetted soybean cotyledon metrics

## **Description**

This data contains metrics for normalized RNA-sequencing read counts from soybean cotyledon across three time stages of development. Early stage cotyledons were collected four days after planting and were green but closed. Middle stage cotyledons were collected while green and open, soon after the plant generated its first set of unifoliate leaves. Late stage cotyledons were collected immediately after the initiation of yellowing and shrinking. The metrics include the log fold change, log counts per million, likelihood ratio, p-values, and FDR values for all genes and all pairwise combinations of treatment groups. To save on size, this example dataset was generated by obtaining a random subset of 1 out of 10 genes from the original resource.

### Usage

```
data(soybean_cn_sub_metrics)
```

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#### **Format**

a RData instance, List with one list element per treatment group combination and 1 row per gene

#### **Details**

Normalized and subsetted soybean cotyledon metrics

- ID gene name
- logFC log fold change
- logCPM log counts per million
- LR likelihood ratio
- PValue p-value
- FDR FDR value

#### See Also

soybean\_cn\_sub for information about the treatment groups

soybean\_ir

Raw soybean leaves iron-metabolism data

## **Description**

This dataset contains raw RNA-sequencing read counts from a soybean dataset that compared leaves that were exposed to iron-rich (iron -postive) soil conditions versus leaves that were exposed to iron-poor (iron-negative) soil conditions. The data was collected 120 minutes after iron conditions were initiated.

# **Format**

a RData instance, 1 row per gene

## **Details**

Raw soybean leaves data

- ID gene name
- N.1 iron-negative condition replicate 1 raw read counts
- N.2 iron-negative condition replicate 2 raw read counts
- N.3 iron-negative condition replicate 3 raw read counts
- P.1 iron-positive condition replicate 1 raw read counts
- P.2 iron-positive condition replicate 2 raw read counts
- P.3 iron-positive condition replicate 3 raw read counts

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soybean\_ir\_metrics

Raw soybean leaves iron-metabolism metrics

## **Description**

This data contains metrics for raw RNA-sequencing read counts from a soybean dataset that compared leaves that were exposed to iron-rich (iron-postive) soil conditions versus leaves that were exposed to iron -poor (iron-negative) soil conditions. The data was collected 120 minutes after iron conditions were initiated. The metrics include the log fold change and the p-values for all genes and all pairwise combinations of treatment groups.

#### **Format**

a RData instance, 1 list per treatment group combination and 1 row per gene

#### **Details**

Raw soybean leaves metrics

- ID gene name
- · logFC log fold change
- PValue p-value

#### See Also

soybean\_ir for information about the treatment groups

soybean\_ir\_sub

Raw and subsetted soybean leaves iron-metabolism data

#### **Description**

This dataset contains raw RNA-sequencing read counts from a soybean dataset that compared leaves that were exposed to iron-rich (iron -postive) soil conditions versus leaves that were exposed to iron-poor (iron-negative) soil conditions. The data was collected 120 minutes after iron conditions were initiated. To save on size, this example dataset was generated by obtaining a random subset of 1 out of 10 genes from the original resource.

## Usage

```
data(soybean_ir_sub)
```

## **Format**

a RData instance, 1 row per gene

#### **Details**

Raw and subsetted soybean leaves data

- · ID gene name
- N.1 iron-negative condition replicate 1 raw read counts
- N.2 iron-negative condition replicate 2 raw read counts
- N.3 iron-negative condition replicate 3 raw read counts
- P.1 iron-positive condition replicate 1 raw read counts
- P.2 iron-positive condition replicate 2 raw read counts
- P.3 iron-positive condition replicate 3 raw read counts

#### References

Moran Lauter AN, Graham MA. NCBI SRA bioproject accession: PRJNA318409.

#### See Also

soybean\_ir from which this dataset is subsetted

```
soybean_ir_sub_metrics
```

Raw and subsetted soybean leaves iron-metabolism metrics

# Description

This data contains metrics for raw RNA-sequencing read counts from a soybean dataset that compared leaves that were exposed to iron-rich (iron-postive) soil conditions versus leaves that were exposed to iron-poor (iron-negative) soil conditions. The data was collected 120 minutes after iron conditions were initiated. The metrics include the log fold change and the p-values for all genes and all pairwise combinations of treatment groups. To save on size, this example dataset was generated by obtaining a random subset of 1 out of 10 genes from the original resource.

#### Usage

```
data(soybean_ir_sub_metrics)
```

### Format

a RData instance, 1 list per treatment group combination and 1 row per gene

# **Details**

Raw and subsetted soybean leaves iron-metabolism metrics

- ID gene name
- logFC log fold change
- PValue p-value

# References

Moran Lauter AN, Graham MA. NCBI SRA bioproject accession: PRJNA318409.

# See Also

soybean\_ir\_sub for information about the treatment groups

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