

Package ‘MAGeCKFlute’

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Type Package

Title Integrative Analysis Pipeline for Pooled CRISPR Functional Genetic Screens

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Description CRISPR (clustered regularly interspaced short palindrome repeats) coupled with nuclease Cas9 (CRISPR/Cas9) screens represent a promising technology to systematically evaluate gene functions. Data analysis for CRISPR/Cas9 screens is a critical process that includes identifying screen hits and exploring biological functions for these hits in downstream analysis. We have previously developed two algorithms, MAGeCK and MAGeCK-VISPR, to analyze CRISPR/Cas9 screen data in various scenarios. These two algorithms allow users to perform quality control, read count generation and normalization, and calculate beta score to evaluate gene selection performance. In downstream analysis, the biological functional analysis is required for understanding biological functions of these identified genes with different screening purposes. Here, We developed MAGeCKFlute for supporting downstream analysis. MAGeCKFlute provides several strategies to remove potential biases within sgRNA-level read counts and gene-level beta scores. The downstream analysis with the package includes identifying essential, non-essential, and target-associated genes, and performing biological functional category analysis, pathway enrichment analysis and protein complex enrichment analysis of these genes. The package also visualizes genes in multiple ways to benefit users exploring screening data. Collectively, MAGeCKFlute enables accurate identification of essential, non-essential, and targeted genes, as well as their related biological functions. This vignette explains the use of the package and demonstrates typical workflows.

License GPL (>=3)

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Suggests knitr, testthat

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arrangePathview *Kegg pathway view and arrange grobs on page*

Description

Kegg pathway view and arrange grobs on page.

Usage

```
arrangePathview(genelist, pathways = c(), top = 4, ncol = 2,
  title = NULL, sub = NULL, organism = "hsa", view_allpath = FALSE,
  output = ".", path.archive = ".", kegg.native = TRUE,
  verbose = TRUE)
```

Arguments

- genelist a data frame with columns of ENTREZID, Control and Treatment. The columns of Control and Treatment represent gene score in Control and Treatment sample.
- pathways character vector, the KEGG pathway ID(s), usually 5 digit, may also include the 3 letter KEGG species code.
- top integer, specifying how many top enriched pathways to be visualized.
- ncol integer, specifying how many column of figures to be arranged in each page.
- title optional string, or grob.
- sub optional string, or grob.
- organism character, either the kegg code, scientific name or the common name of the target species. This applies to both pathway and gene.data or cpd.data. When KEGG ortholog pathway is considered, species="ko". Default species="hsa", it is equivalent to use either "Homo sapiens" (scientific name) or "human" (common name).
- view_allpath boolean, specifying whether view all pathways. Default view_allpath='FALSE', and only plot top enriched pathways.

output	Path to save plot to.
path.archive	character, the directory of KEGG pathway data file (.xml) and image file (.png). Users may supply their own data files in the same format and naming convention of KEGG's (species code + pathway id, e.g. hsa04110.xml, hsa04110.png etc) in this directory. Default kegg.dir="." (current working directory).
kegg.native	logical, whether to render pathway graph as native KEGG graph (.png) or using graphviz layout engine (.pdf). Default kegg.native=TRUE.
verbose	Boolean

Value

plot on the current device

Author(s)

Wubing Zhang

See Also

[KeggPathwayView](#)

Examples

```
data(mle.gene_summary)
# Read beta score from gene summary table in MAGECK MLE results
dd = ReadBeta(mle.gene_summary)
colnames(dd)[2:3] = c("Control", "Treatment")
arrangePathview(dd, "hsa00534", title=NULL, sub=NULL, organism="hsa")
```

BarView

Bar plot

Description

Bar plot

Usage

```
BarView(df, x = "x", y = "y", fill = "#FC6665", bar.width = 0.8,
        position = "dodge", dodge.width = 0.8, main = NA, xlab = NULL,
        ylab = NA, ...)
```

Arguments

df	A data frame.
x	A character, specifying the x-axis.
y	A character, specifying the x-axis.
fill	A character, specifying the fill color.
bar.width	A numeric, specifying the width of bar.

position	"dodge" (default), "stack", "fill".
dodge.width	A numeric, set the width in position_dodge.
main	A character, specifying the figure title.
xlab	A character, specifying the title of x-axis.
ylab	A character, specifying the title of y-axis.
...	Other parameters in geom_bar

Value

An object created by ggplot, which can be assigned and further customized.

Author(s)

Wubing Zhang

Examples

```
mdata = data.frame(group=letters[1:5], count=sample(1:100,5))
BarView(mdata, x = "group", y = "count")
```

BatchRemove	<i>Batch effect removal</i>
-------------	-----------------------------

Description

Batch effect removal

Usage

```
BatchRemove(mat, batchMat, log2trans = FALSE, pca = TRUE,
  positive = FALSE, cluster = FALSE, outdir = NULL)
```

Arguments

mat	A data frame, each row is a gene, and each column is a sample.
batchMat	A data frame, the first column should be 'Samples' (matched colnames of mat) and the second column is 'Batch'. The remaining columns could be Covariates.
log2trans	Boolean, specifying whether do logarithmic transformation before batch removal.
pca	Boolean, specifying whether return pca plot.
positive	Boolean, specifying whether all values should be positive.
cluster	Boolean, specifying whether perform hierarchical clustering.
outdir	Output directory for hierarchical cluster tree.

Value

A list contains two objects, including data and p.

Author(s)

Wubing Zhang

See Also[ComBat](#)**Examples**

```
edata = matrix(c(rnorm(2000, 5), rnorm(2000, 8)), 1000)
colnames(edata) = paste0("s", 1:4)
batchMat = data.frame(sample = colnames(edata), batch = rep(1:2, each = 2))
edata1 = BatchRemove(edata, batchMat)
print(edata1$p)
```

ConsistencyView*Visualize the estimate cell cycle compared to control.*

Description

Estimate cell cycle time in different samples by linear fitting of beta scores.

Usage

```
ConsistencyView(beta, ctrlname, treatname, main = NULL,
  filename = NULL, width = 5, height = 4, ...)
```

Arguments

beta	Data frame, which has columns of ctrlname and other samples.
ctrlname	A character, specifying the names of control samples.
treatname	A character, specifying the name of treatment samples.
main	As in 'plot'.
filename	Figure file name to create on disk. Default filename="NULL", which means no output.
width	As in ggsave.
height	As in ggsave.
...	Other available parameters in ggsave.

Value

An object created by ggplot, which can be assigned and further customized.

Author(s)

Wubing Zhang

Examples

```
data(mle.gene_summary)
# Read beta score from gene summary table in MAGeCK MLE results
dd = ReadBeta(mle.gene_summary)
ConsistencyView(dd, ctrlname = "dms0", treatname = "plx")
```

countsummary

Count summary data generated by running MAGeCK count

Description

The summary of QC values at count level

Usage

```
data("countsummary")
```

Format

A data frame with 4 observations on 13 variables.

References

<https://www.ncbi.nlm.nih.gov/pubmed/25494202> <https://www.ncbi.nlm.nih.gov/pubmed/25476604>

Examples

```
data("countsummary")
head(countsummary)
```

CutoffCalling

Quantile of normal distribution.

Description

Compute cutoff from a normal-distributed vector.

Usage

```
CutoffCalling(d, scale = 1)
```

Arguments

d A numeric vector.

scale Boolean or numeric, specifying how many standard deviation will be used as cutoff.

Value

A numeric value.

Examples

```
CutoffCalling(rnorm(10000))
```

DensityDiffView	<i>Density plot</i>
-----------------	---------------------

Description

Plot the density of beta score deviations.

Usage

```
DensityDiffView(beta, ctrlname = "Control", treatname = "Treatment",
  main = NULL, filename = NULL, width = 5, height = 4, ...)
```

Arguments

beta	Data frame, including ctrlname and treatname as columns.
ctrlname	A character, specifying the name of control sample.
treatname	A character, specifying the name of treatment sample.
main	As in 'plot'.
filename	Figure file name to create on disk. Default filename="NULL", which means no output.
width	As in ggsave.
height	As in ggsave.
...	Other parameters in ggsave.

Value

An object created by ggplot, which can be assigned and further customized.

Author(s)

Wubing Zhang

Examples

```
data(mle.gene_summary)
# Read beta score from gene summary table in MAGeCK MLE results
dd = ReadBeta(mle.gene_summary)
# Density plot of beta score deviation between control and treatment
DensityDiffView(dd, ctrlname = "dms0", treatname = "plx")
```

DensityView

Density plot for gene beta scores in Control and Treatment

Description

Plot the density of gene beta scores in two samples.

Usage

```
DensityView(beta, samples = NULL, main = NULL, xlab = "Beta Score",  
            filename = NULL, width = 5, height = 4, ...)
```

Arguments

beta	Data frame, including samples as columns.
samples	Character, specifying sample names in beta.
main	As in 'plot'.
xlab	As in 'plot'.
filename	Figure file name to create on disk. Default filename="NULL", which means don't save the figure on disk.
width	As in ggsave.
height	As in ggsave.
...	Other available parameters in ggsave.

Value

An object created by ggplot, which can be assigned and further customized.

Author(s)

Wubing Zhang

See Also

[ViolinView](#)

Examples

```
data(mle.gene_summary)  
# Read beta score from gene summary table in MAGeCK MLE results  
dd = ReadBeta(mle.gene_summary)  
DensityView(dd, samples=c("dms", "plx"))  
#or  
DensityView(dd[, c("dms", "plx")])
```

enrich.GSE

*Gene set enrichment analysis***Description**

A universal gene set enrichment analysis tools

Usage

```
enrich.GSE(geneList, keytype = "Symbol", type = "GOBP",
           organism = "hsa", pvalueCutoff = 0.25, limit = c(2, 200),
           gmtpath = NULL, nPerm = 2000, by = "fgsea", verbose = TRUE)
```

Arguments

geneList	A order ranked numeric vector with geneid as names.
keytype	"Entrez" or "Symbol".
type	Molecular signatures for testing, available datasets include Pathway (KEGG, REACTOME, C2_CP), GO (GOBP, GOCC, GOMF), MSIGDB (C1, C2 (C2_CP (C2_CP_PID, C2_CP_BIOCARTA), C2_CGP), C3 (C3_MIR, C3_TFT), C4, C6, C7, HALLMARK) and Complex (CORUM). Any combination of them are also accessible (e.g. 'GOBP+GOMF+KEGG+REACTOME').
organism	'hsa' or 'mmu'.
pvalueCutoff	Pvalue cutoff.
limit	A two-length vector, specifying the minimal and maximal size of gene sets for enrichment analysis.
gmtpath	The path to customized gmt file.
nPerm	The number of permutations.
by	One of 'fgsea' or 'DOSE'
verbose	Boolean

Value

A enrichResult instance.

Author(s)

Wubing Zhang

See Also

[enrich.HGT](#)
[enrich.ORT](#)
[EnrichAnalyzer](#)
[enrichResult-class](#)

Examples

```

data(geneList, package = "DOSE")
## Not run:
  enrichRes = enrich.GSE(geneList, keytype = "entrez")
  head(slot(enrichRes, "result"))

## End(Not run)

```

enrich.HGT

*Do enrichment analysis using Hypergeometric test***Description**

Do enrichment analysis using Hypergeometric test

Usage

```

enrich.HGT(geneList, keytype = "Symbol", type = "GOBP",
  organism = "hsa", pvalueCutoff = 0.25, limit = c(2, 200),
  universe = NULL, gmtpath = NULL, verbose = TRUE)

```

Arguments

geneList	A numeric vector with gene as names.
keytype	"Entrez" or "Symbol".
type	Molecular signatures for testing, available datasets include Pathway (KEGG, REACTOME, C2_CP), GO (GOBP, GOCC, GOMF), MSIGDB (C1, C2 (C2_CP (C2_CP_PID, C2_CP_BIOCARTA), C2_CGP), C3 (C3_MIR, C3_TFT), C4, C6, C7, HALLMARK) and Complex (CORUM). Any combination of them are also accessible (e.g. 'GOBP+GOMF+KEGG+REACTOME').
organism	'hsa' or 'mmu'.
pvalueCutoff	Pvalue cutoff.
limit	A two-length vector, specifying the minimal and maximal size of gene sets for enrichment analysis.
universe	A character vector, specifying the background genelist, default is whole genome.
gmtpath	The path to customized gmt file.
verbose	Boolean

Value

A enrichResult instance.

Author(s)

Wubing Zhang

See Also

[enrich.GSE](#)
[enrich.ORT](#)
[EnrichAnalyzer](#)
[enrichResult-class](#)

Examples

```

data(geneList, package = "DOSE")
genes <- geneList[1:300]
enrichRes <- enrich.HGT(genes, type = "KEGG", keytype = "entrez")
head(slot(enrichRes, "result"))

```

enrich.ORT

Do enrichment analysis using over-representation test

Description

Do enrichment analysis using over-representation test

Usage

```

enrich.ORT(geneList, keytype = "Symbol", type = "GOBP",
  organism = "hsa", pvalueCutoff = 0.25, limit = c(2, 200),
  universe = NULL, gmtpath = NULL, verbose = TRUE)

```

Arguments

geneList	A numeric vector with gene as names.
keytype	"Entrez" or "Symbol".
type	Molecular signatures for testing, available datasets include Pathway (KEGG, REACTOME, C2_CP), GO (GOBP, GOCC, GOMF), MSIGDB (C1, C2 (C2_CP (C2_CP_PID, C2_CP_BIOCARTA), C2_CGP), C3 (C3_MIR, C3_TFT), C4, C6, C7, HALLMARK) and Complex (CORUM). Any combination of them are also accessible (e.g. 'GOBP+GOMF+KEGG+REACTOME').
organism	'hsa' or 'mmu'.
pvalueCutoff	Pvalue cutoff.
limit	A two-length vector, specifying the minimal and maximal size of gene sets for enrichment analysis.
universe	A character vector, specifying the background genelist, default is whole genome.
gmtpath	The path to customized gmt file.
verbose	Boolean

Value

A enrichedResult instance.

Author(s)

Wubing Zhang

See Also[enrich.HGT](#)[enrich.GSE](#)[EnrichAnalyzer](#)[enrichResult-class](#)**Examples**

```
data(geneList, package = "DOSE")
genes <- geneList[1:100]
enrichedRes <- enrich.ORT(genes, keytype = "entrez")
head(slot(enrichedRes, "result"))
```

EnrichAB*Enrichment analysis for Positive and Negative selection genes*

Description

Do enrichment analysis for selected genes, in which positive selection and negative selection are termed as GroupA and GroupB

Usage

```
EnrichAB(data, pvalue = 0.25, enrich_method = "ORT",
  organism = "hsa", limit = c(1, 120), filename = NULL,
  out.dir = ".", width = 6.5, height = 4, verbose = TRUE, ...)
```

Arguments

<code>data</code>	A data frame.
<code>pvalue</code>	Pvalue cutoff.
<code>enrich_method</code>	One of "ORT"(Over-Representing Test) and "HGT"(HyperGemetric test).
<code>organism</code>	"hsa" or "mmu".
<code>limit</code>	A two-length vector (default: c(1, 120)), specifying the min and max size of pathways for enrichent analysis.
<code>filename</code>	Suffix of output file name.
<code>out.dir</code>	Path to save plot to (combined with filename).
<code>width</code>	As in ggsave.
<code>height</code>	As in ggsave.
<code>verbose</code>	Boolean
<code>...</code>	Other available parameters in ggsave.

Value

A list containing enrichment results for each group genes. This list contains eight items, which contain subitems of gridPlot and enrichRes.

Author(s)

Wubing Zhang

EnrichAnalyzer *Enrichment analysis*

Description

Enrichment analysis

Usage

```
EnrichAnalyzer(geneList, keytype = "Symbol", type = "Pathway+GOBP",
  method = "HGT", organism = "hsa", pvalueCutoff = 0.25,
  limit = c(2, 200), universe = NULL, filter = FALSE,
  gmtpath = NULL, verbose = TRUE)
```

Arguments

geneList	A numeric vector with gene as names.
keytype	"Entrez" or "Symbol".
type	Molecular signatures for testing, available datasets include Pathway (KEGG, REACTOME, C2_CP), GO (GOBP, GOCC, GOMF), MSIGDB (C1, C2 (C2_CP (C2_CP_PID, C2_CP_BIOCARTA), C2_CGP), C3 (C3_MIR, C3_TFT), C4, C6, C7, HALLMARK) and Complex (CORUM). Any combination of them are also accessible (e.g. 'GOBP+GOMF+KEGG+REACTOME').
method	One of "ORT"(Over-Representing Test), "GSEA"(Gene Set Enrichment Analysis), and "HGT"(HyperGemetric test).
organism	'hsa' or 'mmu'.
pvalueCutoff	Pvalue cutoff.
limit	A two-length vector (default: c(2, 200)), specifying the minimal and maximal size of gene sets for enrichent analysis.
universe	A character vector, specifying the background genelist, default is whole genome.
filter	Boolean, specifying whether filter out redundancies from the enrichment results.
gmtpath	The path to customized gmt file.
verbose	Boolean

Value

enrichRes is an enrichResult instance.

Author(s)

Wubing Zhang

See Also

[enrich.GSE](#)
[enrich.ORT](#)
[enrich.HGT](#)
[enrichResult-class](#)

Examples

```
data(geneList, package = "DOSE")
## Not run:
  keggA = EnrichAnalyzer(geneList[1:500], keytype = "entrez")
  head(keggA@result)

## End(Not run)
```

EnrichedFilter*Simplify the enrichment results based on Jaccard index*

Description

Simplify the enrichment results based on Jaccard index

Usage

```
EnrichedFilter(enrichment = enrichment, cutoff = 0.8)
```

Arguments

enrichment A data frame of enrichment result.
cutoff A numeric, specifying the cutoff of Jaccard index between two pathways.

Value

A data frame.

Author(s)

Yihan Xiao

Examples

```
data(geneList, package = "DOSE")
## Not run:
  enrichRes <- enrich.HGT(geneList, keytype = "entrez")
  EnrichedFilter(enrichRes)

## End(Not run)
```

EnrichedGeneView *Visualize enriched pathways and genes in those pathways*

Description

Visualize enriched pathways and genes in those pathways

Usage

```
EnrichedGeneView(enrichment, geneList, rank_by = "p.adjust", top = 5,
  bottom = 0, keytype = "Symbol", gene_cutoff = c(-log2(1.5),
  log2(1.5)), custom_gene = NULL, charLength = 40, filename = NULL,
  width = 7, height = 5, ...)
```

Arguments

enrichment	A data frame of enrichment result or an enrichResult object.
geneList	A numeric geneList used in enrichment analysis.
rank_by	"p.adjust" or "NES", specifying the indices for ranking pathways.
top	An integer, specifying the number of positively enriched terms to show.
bottom	An integer, specifying the number of negatively enriched terms to show.
keytype	"Entrez" or "Symbol".
gene_cutoff	A two-length numeric vector, specifying cutoff for genes to show.
custom_gene	A character vector (gene names), customizing genes to show.
charLength	Integer, specifying max length of enriched term name to show as coordinate label.
filename	Figure file name to create on disk. Default filename="NULL", which means no output.
width	As in ggsave.
height	As in ggsave.
...	Other available parameters in ggsave.

Value

An object created by ggplot, which can be assigned and further customized.

Author(s)

Wubing Zhang

Examples

```
data(geneList, package = "DOSE")
## Not run:
  enrichRes <- enrich.GSE(geneList, keytype = "Entrez")
  EnrichedGeneView(enrichment=slot(enrichRes, "result"), geneList, keytype = "Entrez")

## End(Not run)
```

EnrichedView	<i>View enriched terms</i>
--------------	----------------------------

Description

Grid plot for enriched terms

Usage

```
EnrichedView(enrichment, rank_by = "pvalue", mode = 1, subset = NULL,
             top = 0, bottom = 0, x = "LogFDR", charLength = 40,
             filename = NULL, width = 7, height = 4, ...)
```

Arguments

enrichment	A data frame of enrichment result, with columns of ID, Description, p.adjust and NES.
rank_by	"pvalue" or "NES", specifying the indices for ranking pathways.
mode	1 or 2.
subset	A vector of pathway ids.
top	An integer, specifying the number of positively enriched terms to show.
bottom	An integer, specifying the number of negatively enriched terms to show.
x	Character, "NES", "LogP", or "LogFDR", indicating the variable on the x-axis.
charLength	Integer, specifying max length of enriched term name to show as coordinate lab.
filename	Figure file name to create on disk. Default filename="NULL".
width	As in ggsave.
height	As in ggsave.
...	Other available parameters in ggsave.

Value

An object created by ggplot, which can be assigned and further customized.

Author(s)

Wubing Zhang

See Also

[EnrichedView](#)

Examples

```
## Not run:
data(geneList, package = "DOSE")
enrichRes = enrich.GSE(geneList, organism="hsa")
EnrichedView(slot(enrichRes, "result"))

## End(Not run)
```

 EnrichSquare

Enrichment analysis for selected treatment related genes

Description

Do enrichment analysis for selected treatment related genes in 9-squares

Usage

```
EnrichSquare(beta, id = "Gene", keytype = "Symbol", x = "Control",
  y = "Treatment", pvalue = 0.05, enrich_method = "ORT",
  organism = "hsa", limit = c(1, 120), filename = NULL,
  out.dir = ".", width = 6.5, height = 4, verbose = TRUE, ...)
```

Arguments

beta	Data frame, with columns of "Gene", "group", and "Diff".
id	A character, indicating the gene column in the data.
keytype	A character, "Symbol" or "Entrez".
x	A character, indicating the x-axis in the 9-square scatter plot.
y	A character, indicating the y-axis in the 9-square scatter plot.
pvalue	Pvalue cutoff.
enrich_method	One of "ORT"(Over-Representing Test) and "HGT"(HyperGemetric test).
organism	"hsa" or "mmu".
limit	A two-length vector (default: c(1, 120)), specifying the min and max size of pathways for enrichent analysis.
filename	Suffix of output file name. NULL(default) means no output.
out.dir	Path to save plot to (combined with filename).
width	As in ggsave.
height	As in ggsave.
verbose	Boolean.
...	Other available parameters in ggsave.

Value

A list containing enrichment results for each group genes. Each item in the returned list has two sub items:

gridPlot	an object created by ggplot, which can be assigned and further customized.
enrichRes	a enrichResult instance.

Author(s)

Wubing Zhang

Description

Integrative analysis pipeline using the gene summary table in MAGeCK MLE results

Usage

```
FluteMLE(gene_summary, treatname, ctrlname = "Depmap",
         keytype = "Symbol", organism = "hsa", incorporateDepmap = FALSE,
         cell_lines = NA, lineages = "All", norm_method = "cell_cycle",
         posControl = NULL, omitEssential = TRUE, top = 10,
         toplabels = NA, scale_cutoff = 2, limit = c(0, 200),
         pvalueCutoff = 0.25, enrich_method = "ORT", proj = NA,
         width = 10, height = 7, outdir = ".", view_allpath = FALSE,
         verbose = TRUE)
```

Arguments

gene_summary	A data frame or a file path to gene summary file generated by MAGeCK-MLE.
treatname	A character vector, specifying the names of treatment samples.
ctrlname	A character vector, specifying the names of control samples. If there is no controls in your CRISPR screen, you can specify "Depmap" as ctrlname and set 'incorporateDepmap=TRUE'.
keytype	"Entrez" or "Symbol".
organism	"hsa" or "mmu".
incorporateDepmap	Boolean, indicating whether incorporate Depmap data into analysis.
cell_lines	A character vector, specifying the cell lines in Depmap to be considered.
lineages	A character vector, specifying the lineages in Depmap to be considered.
norm_method	One of "none", "cell_cycle" (default) or "loess".
posControl	A character vector, specifying a list of positive control gene symbols.
omitEssential	Boolean, indicating whether omit common essential genes from the downstream analysis.
top	An integer, specifying number of top selected genes to be labeled in rank figure.
toplabels	A character vector, specifying interested genes to be labeled in rank figure.
scale_cutoff	Boolean or numeric, specifying how many standard deviation will be used as cutoff.
limit	A two-length vector, specifying the minimal and maximal size of gene sets for enrichment analysis.
pvalueCutoff	A numeric, specifying pvalue cutoff of enrichment analysis, default 1.
enrich_method	One of "ORT"(Over-Representing Test) and "HGT"(HyperGometric test).
proj	A character, indicating the prefix of output file name, which can't contain special characters.

width	The width of summary pdf in inches.
height	The height of summary pdf in inches.
outdir	Output directory on disk.
view_allpath	Boolean, whether output all pathway view figures (time-consuming).
verbose	Boolean

Details

MAGeCK-MLE can be used to analyze screen data from multi-conditioned experiments. MAGeCK-MLE also normalizes the data across multiple samples, making them comparable to each other. The most important output of MAGeCK MLE is 'gene_summary' file, which includes the beta scores of multiple conditions and the associated statistics. The 'beta score' for each gene describes how the gene is selected: a positive beta score indicates a positive selection, and a negative beta score indicates a negative selection.

The downstream analysis includes identifying essential, non-essential, and target-associated genes, and performing biological functional category analysis and pathway enrichment analysis of these genes. The function also visualizes genes in the context of pathways to benefit users exploring screening data.

Value

All of the pipeline results is output into the `out.dir/MAGeCKFlute_proj`, which includes a pdf file and many folders. The pdf file 'FluteMLE_proj_norm_method.pdf' is the summary of pipeline results. For each section in this pipeline, figures and useful data are outputted to corresponding subfolders.

- QC: Quality control
- Selection: Positive selection and negative selection.
- Enrichment: Enrichment analysis for positive and negative selection genes.
- PathwayView: Pathway view for top enriched pathways.

Author(s)

Wubing Zhang

See Also

[FluteRRA](#)

Examples

```
data(mle.gene_summary)
## Not run:
# functional analysis for MAGeCK MLE results
FluteMLE(mle.gene_summary, treatname = "plx", ctrlname = "dms", proj = "PLX")

## End(Not run)
```

 FluteRRA

Downstream analysis based on MAGeCK-RRA result

Description

Integrative analysis pipeline using the gene summary table in MAGeCK RRA results

Usage

```
FluteRRA(gene_summary, sgrna_summary = NULL, keytype = "Symbol",
  organism = "hsa", incorporateDepmap = TRUE, cell_lines = NA,
  lineages = "All", omitEssential = TRUE, top = 5,
  toplabels = NULL, scale_cutoff = 2, limit = c(2, 200),
  pvalueCutoff = 0.25, proj = NA, width = 12, height = 6,
  outdir = ".", verbose = TRUE)
```

Arguments

gene_summary	A file path or a data frame of gene summary data.
sgrna_summary	A file path or a data frame of sgRNA summary data.
keytype	"Entrez" or "Symbol".
organism	"hsa" or "mmu".
incorporateDepmap	Boolean, indicating whether incorporate Depmap data into analysis.
cell_lines	A character vector, specifying the cell lines in Depmap to be considered.
lineages	A character vector, specifying the lineages in Depmap to be considered.
omitEssential	Boolean, indicating whether omit common essential genes from the downstream analysis.
top	An integer, specifying number of top selected genes to be labeled in rank figure.
toplabels	A character vector, specifying interested genes to be labeled in rank figure.
scale_cutoff	Boolean or numeric, specifying how many standard deviation will be used as cutoff.
limit	A two-length vector, specifying the minimal and maximal size of gene sets for enrichment analysis.
pvalueCutoff	A numeric, specifying pvalue cutoff of enrichment analysis, default 1.
proj	A character, indicating the prefix of output file name.
width	The width of summary pdf in inches.
height	The height of summary pdf in inches.
outdir	Output directory on disk.
verbose	Boolean

Details

MAGeCK RRA allows for the comparison between two experimental conditions. It can identify genes and sgRNAs are significantly selected between the two conditions. The most important output of MAGeCK RRA is the file 'gene_summary.txt'. MAGeCK RRA will output both the negative score and positive score for each gene. A smaller score indicates higher gene importance. MAGeCK RRA will also output the statistical value for the scores of each gene. Genes that are significantly positively and negatively selected can be identified based on the p-value or FDR.

The downstream analysis of this function includes identifying positive and negative selection genes, and performing biological functional category analysis and pathway enrichment analysis of these genes.

Value

All of the pipeline results is output into the `out.dir/proj_Results`, which includes a pdf file and a folder named 'RRA'.

Author(s)

Wubing Zhang

See Also

[FluteMLE](#)

Examples

```
data("rra.gene_summary")
data("rra.sgrna_summary")
## Not run:
  # Run the FluteRRA pipeline
  FluteRRA(rra.gene_summary, rra.sgrna_summary, proj="PLX", organism="hsa")

## End(Not run)
```

getCols

Map values to colors

Description

Map values to colors

Usage

```
getCols(x, palette = 1)
```

Arguments

x	A numeric vector.
palette	diverge, rainbow, sequential

Value

A vector of colors corresponding to input vector.

Author(s)

Wubing Zhang

getGeneAnn	<i>Retrieve gene annotations from the NCBI, HNSC, and Uniprot databases.</i>
------------	--

Description

Retrieve gene annotations from the NCBI, HNSC, and Uniprot databases.

Usage

```
getGeneAnn(org = "hsa", update = FALSE)
```

Arguments

org	Character, hsa (default), bta, cfa, mmu, ptr, rno, ssc are optional.
update	Boolean, indicating whether download current annotation.

Value

A data frame.

Author(s)

Wubing Zhang

Examples

```
## Not run:  
ann = getGeneAnn("hsa")  
head(ann)  
  
## End(Not run)
```

getOrg *Get the kegg code of specific mammalia organism.*

Description

Get the kegg code of specific mammalia organism.

Usage

```
getOrg(organism)
```

Arguments

organism Character, KEGG species code, or the common species name. For all potential values check: data(bods); bods. Default org="hsa", and can also be "human" (case insensitive).

Value

A list containing three elements:

org species
pkgannotation package name

Author(s)

Wubing Zhang

Examples

```
ann = getOrg("human")
print(ann$pkg)
```

getOrtAnn *Retrieve reference orthologs annotation.*

Description

Retrieve reference orthologs annotation.

Usage

```
getOrtAnn(fromOrg = "mmu", toOrg = "hsa", update = FALSE)
```

Arguments

fromOrg Character, hsa (default), bta, cfa, mmu, ptr, rno, ssc are optional.
toOrg Character, hsa (default), bta, cfa, mmu, ptr, rno, ssc are optional.
update Boolean, indicating whether download recent annotation from NCBI.

Value

A data frame.

Author(s)

Wubing Zhang

Examples

```
## Not run:
ann = getOrtAnn("mmu", "hsa")
head(ann)

## End(Not run)
```

 gsGetter

Extract pathway annotation from GMT file.

Description

Extract pathway annotation from GMT file.

Usage

```
gsGetter(gmtpath = NULL, type = "All", limit = c(0, Inf),
  organism = "hsa", update = FALSE)
```

Arguments

gmtpath	The path to customized gmt file.
type	Molecular signatures for testing, available datasets include Pathway (KEGG, REACTOME, C2_CP:PID, C2_CP:BIOCARTA), GO (GOBP, GOCC, GOMF), MSIGDB (C1, C2 (C2_CP (C2_CP:PID, C2_CP:BIOCARTA), C2_CGP), C3 (C3_MIR, C3_TFT), C4 (C4_CGN, C4_CM), C5 (C5_BP, C5_CC, C5_MF), C6, C7, H) and Complex (CORUM). Any combination of them are also accessible (e.g. 'GOBP+GOMF+KEGG+REACTOME').
limit	A two-length vector, specifying the minimal and maximal size of gene sets to load.
organism	'hsa' or 'mmu'.
update	Boolean, indicating whether update the gene sets from source database.

Value

A three-column data frame.

Author(s)

Wubing Zhang

Examples

```
gene2path = gsGetter(type = "REACTOME+CORUM")
head(gene2path)
```

hclustView

Cluster and view cluster tree

Description

Cluster and view cluster tree

Usage

```
hclustView(d, method = "average", label_cols = NULL, bar_cols = NULL,
  main = NA, xlab = NA, horiz = TRUE, ...)
```

Arguments

d	A dissimilarity structure as produced by dist.
method	The agglomeration method to be used. This should be (an unambiguous abbreviation of) one of "ward.D", "ward.D2", "single", "complete", "average" (= UPGMA), "mcquitty" (= WPGMA), "median" (= WPGMC) or "centroid" (= UPGMC).
label_cols	A vector to be used as label's colors for the dendrogram.
bar_cols	Either a vector or a matrix, which will be plotted as a colored bar.
main	As in 'plot'.
xlab	As in 'plot'.
horiz	Logical indicating if the dendrogram should be drawn horizontally or not.
...	Arguments to be passed to methods, such as graphical parameters (see par).

Value

Plot figure on open device.

Author(s)

Wubing Zhang

Examples

```
label_cols = rownames(USArrests)
hclustView(dist(USArrests), label_cols=label_cols, bar_cols=label_cols)
```

HeatmapView	<i>Draw heatmap</i>
-------------	---------------------

Description

Draw heatmap

Usage

```
HeatmapView(mat, limit = c(-2, 2),  
  colPal = rev(colorRampPalette(c("#c12603", "white", "#0073B6"), space =  
  "Lab")(199)), filename = NA, width = NA, height = NA, ...)
```

Arguments

mat	Matrix like object, each row is gene and each column is sample.
limit	Max value in heatmap
colPal	colorRampPalette.
filename	File path where to save the picture.
width	Manual option for determining the output file width in inches.
height	Manual option for determining the output file height in inches.
...	Other parameters in pheatmap.

Value

Invisibly a pheatmap object that is a list with components.

Author(s)

Wubing Zhang

Examples

```
data(mle.gene_summary)  
dd = ReadBeta(mle.gene_summary)  
gg = cor(dd[,2:ncol(dd)])  
HeatmapView(gg, display_numbers = TRUE)
```

IdentBarView	<i>Identical bar plot</i>
--------------	---------------------------

Description

Identical bar plot

Usage

```
IdentBarView(gg, x = "x", y = "y", fill = c("#CF3C2B", "#394E80"),  
  main = NULL, xlab = NULL, ylab = NULL, filename = NULL,  
  width = 5, height = 4, ...)
```

Arguments

gg	A data frame.
x	A character, indicating column (in countSummary) of x-axis.
y	A character, indicating column (in countSummary) of y-axis.
fill	A character, indicating fill color of all bars.
main	A character, specifying the figure title.
xlab	A character, specifying the title of x-axis.
ylab,	A character, specifying the title of y-axis.
filename	Figure file name to create on disk. Default filename="NULL", which means don't save the figure on disk.
width	As in ggsave.
height	As in ggsave.
...	Other available parameters in ggsave.

Value

An object created by ggplot, which can be assigned and further customized.

Author(s)

Wubing Zhang

Examples

```
data(countsummary)  
IdentBarView(countsummary, x="Label", y="Reads")
```

IncorporateDepmap	<i>Incorporate Depmap screen into analysis</i>
-------------------	--

Description

Incorporate Depmap screen into analysis

Usage

```
IncorporateDepmap(dd, symbol = "id", cell_lines = NA,
  lineages = "All", na.rm = FALSE)
```

Arguments

dd	A data frame.
symbol	A character, specifying the column name of gene symbols in the data frame.
cell_lines	A character vector, specifying the cell lines in Depmap to be considered.
lineages	A character vector, specifying the lineages in Depmap to be considered.
na.rm	Boolean, indicating whether removing NAs from the results.

Value

A data frame with Depmap column attached.

Author(s)

Wubing Zhang

Examples

```
dd.rra = ReadRRA(rra.gene_summary)
## Not run:
  dd.rra = IncorporateDepmap(dd.rra, cell_lines=rownames(depmap_similarity)[1:3])
  head(dd.rra)

## End(Not run)
```

KeggPathwayView	<i>Kegg pathway view</i>
-----------------	--------------------------

Description

Plot kegg pathway and color specific genes.

Usage

```
KeggPathwayView(gene.data = NULL, cpd.data = NULL, pathway.id,
  species = "hsa", kegg.dir = ".", cpd.idtype = "kegg",
  gene.idtype = "ENTREZ", gene.annotpkg = NULL, min.nnodes = 3,
  kegg.native = TRUE, map.null = TRUE, expand.node = FALSE,
  split.group = FALSE, map.symbol = TRUE, map.cpdname = TRUE,
  node.sum = "sum", discrete = list(gene = FALSE, cpd = FALSE),
  limit = list(gene = 1, cpd = 1), bins = list(gene = 10, cpd = 10),
  both.dirs = list(gene = TRUE, cpd = TRUE), trans.fun = list(gene =
  NULL, cpd = NULL), low = list(gene = "deepskyblue1", cpd = "blue"),
  mid = list(gene = "gray", cpd = "gray"), high = list(gene = "red",
  cpd = "yellow"), na.col = "transparent", verbose = TRUE, ...)
```

Arguments

gene.data	Either vector (single sample) or a matrix-like data (multiple sample). Vector should be numeric with gene IDs as names or it may also be character of gene IDs. Character vector is treated as discrete or count data. Matrix-like data structure has genes as rows and samples as columns. Row names should be gene IDs. Here gene ID is a generic concepts, including multiple types of gene, transcript and protein uniquely mappable to KEGG gene IDs. KEGG ortholog IDs are also treated as gene IDs as to handle metagenomic data. Check details for mappable ID types. Default gene.data=NULL.
cpd.data	The same as gene.data, except named with IDs mappable to KEGG compound IDs. Over 20 types of IDs included in ChEMBL database can be used here. Check details for mappable ID types. Default cpd.data=NULL. Note that gene.data and cpd.data can't be NULL simultaneously.
pathway.id	Character vector, the KEGG pathway ID(s), usually 5 digit, may also include the 3 letter KEGG species code.
species	Character, either the kegg code, scientific name or the common name of the target species. This applies to both pathway and gene.data or cpd.data. When KEGG ortholog pathway is considered, species="ko". Default species="hsa", it is equivalent to use either "Homo sapiens" (scientific name) or "human" (common name).
kegg.dir	Character, the directory of KEGG pathway data file (.xml) and image file (.png). Users may supply their own data files in the same format and naming convention of KEGG's (species code + pathway id, e.g. hsa04110.xml, hsa04110.png etc) in this directory. Default kegg.dir="." (current working directory).
cpd.idtype	Character, ID type used for the cpd.data. Default cpd.idtype="kegg" (include compound, glycan and drug accessions).
gene.idtype	Character, ID type used for the gene.data, case insensitive. Default gene.idtype="entrez", i.e. Entrez Gene, which are the primary KEGG gene ID for many common model organisms. For other species, gene.idtype should be set to "KEGG" as KEGG use other types of gene IDs. For the common model organisms, you may also specify other types of valid IDs. To check the ID list, do: data(gene.idtype.list); gene.idtype.list.
gene.annotpkg	Character, the name of the annotation package to use for mapping between other gene ID types including symbols and Entrez gene ID. Default gene.annotpkg=NULL.
min.nnodes	Integer, minimal number of nodes of type "gene","enzyme", "compound" or "ortholog" for a pathway to be considered. Default min.nnodes=3.

kegg.native	Logical, whether to render pathway graph as native KEGG graph (.png) or using graphviz layout engine (.pdf). Default kegg.native=TRUE.
map.null	Logical, whether to map the NULL gene.data or cpd.data to pathway. When NULL data are mapped, the gene or compound nodes in the pathway will be rendered as actually mapped nodes, except with NA-valued color. When NULL data are not mapped, the nodes are rendered as unmapped nodes. This argument mainly affects native KEGG graph view, i.e. when kegg.native=TRUE. Default map.null=TRUE.
expand.node	Logical, whether the multiple-gene nodes are expanded into single-gene nodes. Each expanded single-gene nodes inherits all edges from the original multiple-gene node. This option only affects graphviz graph view, i.e. when kegg.native=FALSE. This option is not effective for most metabolic pathways where it conflicts with converting reactions to edges. Default expand.node=FLASE.
split.group	Logical, whether split node groups are split to individual nodes. Each split member nodes inherits all edges from the node group. This option only affects graphviz graph view, i.e. when kegg.native=FALSE. This option also effects most metabolic pathways even without group nodes defined originally. For these pathways, genes involved in the same reaction are grouped automatically when converting reactions to edges unless split.group=TRUE. d split.group=FLASE.
map.symbol	Logical, whether map gene IDs to symbols for gene node labels or use the graphic name from the KGML file. This option is only effective for kegg.native=FALSE or same.layer=FALSE when kegg.native=TRUE. For same.layer=TRUE when kegg.native=TRUE, the native KEGG labels will be kept. Default map.symbol=TRUE.
map.cpdname	Logical, whether map compound IDs to formal names for compound node labels or use the graphic name from the KGML file (KEGG compound accessions). This option is only effective for kegg.native=FALSE. When kegg.native=TRUE, the native KEGG labels will be kept. Default map.cpdname=TRUE.
node.sum	Character, the method name to calculate node summary given that multiple genes or compounds are mapped to it. Poential options include "sum", "mean", "median", "max", "max.abs" and "random". Default node.sum="sum".
discrete	A list of two logical elements with "gene" and "cpd" as the names. This argument tells whether gene.data or cpd.data should be treated as discrete. Default dsicrete=list(gene=FALSE, cpd=FALSE), i.e. both data should be treated as continuous.
limit	A list of two numeric elements with "gene" and "cpd" as the names. This argument specifies the limit values for gene.data and cpd.data when converting them to pseudo colors. Each element of the list could be of length 1 or 2. Length 1 suggests discrete data or 1 directional (positive-valued) data, or the absolute limit for 2 directional data. Length 2 suggests 2 directional data. Default limit=list(gene=1, cpd=1).
bins	A list of two integer elements with "gene" and "cpd" as the names. This argument specifies the number of levels or bins for gene.data and cpd.data when converting them to pseudo colors. Default limit=list(gene=10, cpd=10).
both.dirs	A list of two logical elements with "gene" and "cpd" as the names. This argument specifies whether gene.data and cpd.data are 1 directional or 2 directional data when converting them to pseudo colors. Default limit=list(gene=TRUE, cpd=TRUE).
trans.fun	A list of two function (not character) elements with "gene" and "cpd" as the names. This argument specifies whether and how gene.data and cpd.data are

	transformed. Examples are log, abs or users' own functions. Default limit=list(gene=NULL, cpd=NULL).
low	A list of two colors with "gene" and "cpd" as the names.
mid	A list of two colors with "gene" and "cpd" as the names.
high	A list of two colors with "gene" and "cpd" as the names.
na.col	Color used for NA's or missing values in gene.data and cpd.data. d na.col="transparent".
verbose	Boolean
...	Extra arguments passed to keggview.native or keggview.graph function.

Details

The function `KeggPathwayView` is a revised version of `pathview` function in `pathview` package. `KeggPathwayView` maps and renders user data on relevant pathway graphs. `KeggPathwayView` is a stand alone program for pathway based data integration and visualization. It also seamlessly integrates with pathway and functional analysis tools for large-scale and fully automated analysis. `KeggPathwayView` provides strong support for data Integration. It works with: 1) essentially all types of biological data mappable to pathways, 2) over 10 types of gene or protein IDs, and 20 types of compound or metabolite IDs, 3) pathways for over 2000 species as well as KEGG orthology, 4) various data attributes and formats, i.e. continuous/discrete data, matrices/vectors, single/multiple samples etc. To see mappable external gene/protein IDs do: `data(gene.idtype.list)`, to see mappable external compound related IDs do: `data(rn.list); names(rn.list)`. `KeggPathwayView` generates both native KEGG view and Graphviz views for pathways. Currently only KEGG pathways are implemented. Hopefully, pathways from Reactome, NCI and other databases will be supported in the future.

The argument `low`, `mid`, and `high` specifies the color spectra to code `gene.data` and `cpd.data`. When data are 1 directional (TRUE value in `both.dirs`), only `mid` and `high` are used to specify the color spectra. Default spectra (low-mid-high) "green"- "gray"- "red" and "blue"- "gray"- "yellow" are used for `gene.data` and `cpd.data` respectively. The values for 'low, mid, high' can be given as color names ('red'), plot color index (2=red), and HTML-style RGB, ("#FF0000"=red).

Value

The result returned by `KeggPathwayView` function is a named list corresponding to the input pathway ids. Each element (for each pathway itself is a named list, with 2 elements ("plot.data.gene", "plot.data.cpd"). Both elements are `data.frame` or `NULL` depends on the corresponding input data `gene.data` and `cpd.data`. These `data.frames` record the plot data for mapped gene or compound nodes: rows are mapped genes/compounds, columns are:

<code>kegg.names</code>	standard KEGG IDs/Names for mapped nodes. It's Entrez Gene ID or KEGG Compound Accessions.
<code>labels</code>	Node labels to be used when needed.
<code>all.mapped</code>	All molecule (gene or compound) IDs mapped to this node.
<code>type</code>	node type, currently 4 types are supported: "gene", "enzyme", "compound" and "ortholog".
<code>x</code>	x coordinate in the original KEGG pathway graph.
<code>y</code>	y coordinate in the original KEGG pathway graph.
<code>width</code>	node width in the original KEGG pathway graph.
<code>height</code>	node height in the original KEGG pathway graph.
<code>other columns</code>	columns of the mapped gene/compound data and corresponding pseudo-color codes for individual samples

Author(s)

Wubing Zhang

Examples

```
#load data
data(mle.gene_summary)
dd = ReadBeta(mle.gene_summary)
gene.data = dd$plx
names(gene.data) = rownames(dd)

pv.out <- KeggPathwayView(gene.data, pathway.id = "04110",
  species = "hsa", out.suffix = "gse16873", kegg.native = TRUE)
```

MapRatesView	<i>View mapping ratio</i>
--------------	---------------------------

Description

View mapping ratio of each sample

Usage

```
MapRatesView(countSummary, Label = "Label", Reads = "Reads",
  Mapped = "Mapped", filename = NULL, width = 5, height = 4, ...)
```

Arguments

countSummary	A data frame, which contains columns of ‘Label’, ‘Reads’, and ‘Mapped’
Label	A character, indicating column (in countSummary) of sample names.
Reads	A character, indicating column (in countSummary) of total reads.
Mapped	A character, indicating column (in countSummary) of mapped reads.
filename	Figure file name to create on disk. Default filename="NULL", which means don't save the figure on disk.
width	As in ggsave.
height	As in ggsave.
...	Other available parameters in ggsave.

Value

An object created by ggplot, which can be assigned and further customized.

Author(s)

Wubing Zhang

Examples

```
data(countsummary)
MapRatesView(countsummary)
```

MAView

*MAplot of gene beta scores***Description**

MAplot of gene beta scores in Control vs Treatment

Usage

```
MAView(beta, ctrlname = "Control", treatname = "Treatment",
       main = NULL, show.statistics = TRUE, add.smooth = TRUE, lty = 1,
       smooth.col = "red", plot.method = c("loess", "lm", "glm", "gam"),
       filename = NULL, width = 5, height = 4, ...)
```

Arguments

beta	Data frame, including ctrlname and treatname as columns.
ctrlname	Character vector, specifying the name of control sample.
treatname	Character vector, specifying the name of treatment sample.
main	As in plot.
show.statistics	Show statistics .
add.smooth	Whether add a smooth line to the plot.
lty	Line type for smooth line.
smooth.col	Color of smooth line.
plot.method	A string specifying the method to fit smooth line, which should be one of "loess" (default), "lm", "glm" and "gam".
filename	Figure file name to create on disk. Default filename="NULL", which means don't save the figure on disk.
width	As in ggsave.
height	As in ggsave.
...	Other available parameters in function 'ggsave'.

Value

An object created by ggplot, which can be assigned and further customized.

Author(s)

Wubing Zhang

Examples

```
data(mle.gene_summary)
# Read beta score from gene summary table in MAGECK MLE results
dd = ReadBeta(mle.gene_summary)
MAView(dd, ctrlname = "dms0", treatname = "plx")
```

mle.gene_summary	<i>Gene summary table in MAGeCK MLE results</i>
------------------	---

Description

The gene summary results generated by running MAGeCK MLE on CRISPR screens.

Usage

```
data("mle.gene_summary")
```

Format

A data frame.

References

<https://www.ncbi.nlm.nih.gov/pubmed/25494202> <https://www.ncbi.nlm.nih.gov/pubmed/26673418>

Examples

```
data("mle.gene_summary")  
head(mle.gene_summary)
```

noEnrichPlot	<i>Blank figure</i>
--------------	---------------------

Description

Blank figure

Usage

```
noEnrichPlot(main = "No enriched terms")
```

Arguments

`main` The title of figure.

Value

An object created by ggplot, which can be assigned and further customized.

Author(s)

Wubing Zhang

normalize.loess *normalize.loess*

Description

Loess normalization method.

Usage

```
normalize.loess(mat, subset = sample(1:(dim(mat)[1]), min(c(5000,
  nrow(mat))))), epsilon = 10^-2, maxit = 1, log.it = FALSE,
  verbose = TRUE, span = 2/3, family.loess = "symmetric", ...)
```

Arguments

mat	A matrix with columns containing the values of the chips to normalize.
subset	A subset of the data to fit a loess to.
epsilon	A tolerance value (supposed to be a small value - used as a stopping criterion).
maxit	Maximum number of iterations.
log.it	Logical. If TRUE it takes the log2 of mat.
verbose	Logical. If TRUE displays current pair of chip being worked on.
span	Parameter to be passed the function loess
family.loess	Parameter to be passed the function loess . "gaussian" or "symmetric" are acceptable values for this parameter.
...	Any of the options of <code>normalize.loess</code> you would like to modify (described above).

Value

A matrix similar as mat.

Author(s)

Wubing Zhang

See Also

[loess](#)

[NormalizeBeta](#)

Examples

```
beta = ReadBeta(mle.gene_summary)
beta_loess = normalize.loess(beta[,c("dms0", "plx")])
```

NormalizeBeta	<i>Normalize gene beta scores</i>
---------------	-----------------------------------

Description

Two normalization methods are available. `cell_cycle` method normalizes gene beta scores based on positive control genes in CRISPR screening. `loess` method normalizes gene beta scores using loess.

Usage

```
NormalizeBeta(beta, id = 1, method = "cell_cycle", posControl = NULL,  
             samples = NULL)
```

Arguments

<code>beta</code>	Data frame.
<code>id</code>	An integer specifying the column of gene.
<code>method</code>	Character, one of 'cell_cycle'(default) and 'loess'. or character string giving the name of the table column containing the gene names.
<code>posControl</code>	A character vector, specifying a list of positive control genes.
<code>samples</code>	Character vector, specifying the sample names in <i>beta</i> columns. If NULL (default), take all <i>beta</i> columns as samples.

Details

In CRISPR screens, cells treated with different conditions (e.g., with or without drug) may have different proliferation rates. So it's necessary to normalize the proliferation rate based on defined positive control genes among samples. After normalization, the beta scores are comparable across samples. `loess` is another optional normalization method, which is used to normalize array data before.

Value

A data frame with same format as input data *beta*.

Author(s)

Wubing Zhang

Examples

```
data(mle.gene_summary)  
# Read beta score from gene summary table in MAGeCK MLE results  
dd = ReadBeta(mle.gene_summary)  
#Cell Cycle normalization  
dd_essential = NormalizeBeta(dd, samples=c("dms0", "plx"), method="cell_cycle")  
head(dd_essential)  
  
#Optional loess normalization  
dd_loess = NormalizeBeta(dd, samples=c("dms0", "plx"), method="loess")
```

```
head(dd_loess)
```

OmitCommonEssential *Omit common essential genes based on depmap data*

Description

Omit common essential genes based on depmap data

Usage

```
OmitCommonEssential(dd, symbol = "id", lineages = "All",  
  dependency = -0.5)
```

Arguments

dd	A data frame.
symbol	A character, specifying the column name of gene symbols in the data frame.
lineages	A character vector, specifying the lineages used for common essential gene selection.
dependency	A numeric, specifying the threshold for common essential gene selection.

Value

A data frame.

Author(s)

Wubing Zhang

Examples

```
dd.rra = ReadRRA(rra.gene_summary)  
dim(dd.rra)  
## Not run:  
  rra.omit = OmitCommonEssential(dd.rra)  
  dim(rra.omit)  
  
## End(Not run)
```

RankView

View the rank of gene points

Description

Rank all genes according to beta score deviation, and label top and bottom meaningful genes. Some other interested genes can be labeled too.

Usage

```
RankView(rankdata, genelist = NULL, top = 10, bottom = 10,  
         cutoff = NULL, main = NULL, filename = NULL, width = 5,  
         height = 4, ...)
```

Arguments

rankdata	Numeric vector, with gene as names.
genelist	Character vector, specifying genes to be labeled in figure.
top	Integer, specifying number of top genes to be labeled.
bottom	Integer, specifying number of bottom genes to be labeled.
cutoff	Numeric.
main	As in 'plot'.
filename	Figure file name to create on disk. Default filename="NULL", which means no output.
width	As in ggsave.
height	As in ggsave.
...	Other available parameters in function 'ggsave'.

Value

An object created by ggplot, which can be assigned and further customized.

Author(s)

Wubing Zhang

Examples

```
data(rra.gene_summary)  
rra = ReadRRA(rra.gene_summary)  
rankdata = rra$Score  
names(rankdata) = rra$id  
RankView(rankdata)
```

ReadBeta	<i>Read gene beta scores</i>
----------	------------------------------

Description

Read gene beta scores from file or data frame

Usage

```
ReadBeta(gene_summary)
```

Arguments

`gene_summary` A data frame or a file path to gene summary file generated by MAGeCK-MLE.

Value

A data frame, whose first column is Gene and other columns are comparisons.

Author(s)

Wubing Zhang

Examples

```
data(mle.gene_summary)
dd = ReadBeta(mle.gene_summary)
head(dd)
```

ReadGMT	<i>ReadGMT</i>
---------	----------------

Description

Parse gmt file to a data.frame
write data frame to a gmt file

Usage

```
ReadGMT(gmtpath, limit = c(0, Inf))

writeGMT(gene2path, gmtfile)
```

Arguments

`gmtpath` The path to gmt file.
`limit` A integer vector of length two, specifying the limit of geneset size.
`gene2path` A data frame. The columns should be Gene, Pathway ID, and Pathway Name.
`gmtfile` Path to gmt file.

Value

An data.frame, in which the first column is gene, and the second column is pathway name.
Output gmt file to local folder.

Author(s)

Wubing Zhang
Wubing Zhang

Examples

```
gene2path = gsGetter(type = "Complex")  
writeGMT(gene2path, "Protein_complex.gmt")
```

ReadRRA

Read gene summary file in MAGeCK-RRA results

Description

Read gene summary file in MAGeCK-RRA results

Usage

```
ReadRRA(gene_summary, score = c("lfc", "rra")[1])
```

Arguments

gene_summary A data frame or a file path to gene summary file generated by MAGeCK-RRA.
score "lfc" (default) or "rra", specifying the score type.

Details

If the score type is equal to lfc, then LFC will be returned. If the score type is rra, the log10 transformed RRA score will be returned. For FACS-based CRISPR screens, rra score is not recommended.

Value

A data frame including three columns, including "id", "LFC" and "FDR".

Author(s)

Wubing Zhang

Examples

```
data(rra.gene_summary)  
dd.rra = ReadRRA(rra.gene_summary)  
head(dd.rra)
```

ReadsgRRA

Read sgRNA summary in MAGeCK-RRA results

Description

Read sgRNA summary in MAGeCK-RRA results

Usage

```
ReadsgRRA(sgRNA_summary)
```

Arguments

`sgRNA_summary` A file path or a data frame of sgRNA summary data.

Value

A data frame.

Author(s)

Wubing Zhang

Examples

```
data(rra.sgrna_summary)
sgrra = ReadsgRRA(rra.sgrna_summary)
head(sgrra)
```

ResembleDepmap*Compute the similarity between customized CRISPR screen with Depmap screens*

Description

Compute the similarity between customized CRISPR screen with Depmap screens

Usage

```
ResembleDepmap(dd, symbol = "id", score = "Score", lineages = "All",
  method = c("pearson", "spearman", "kendall")[1])
```

Arguments

dd	A data frame.
symbol	A character, specifying the column name of gene symbols in the data frame.
score	A character, specifying the column name of gene essentiality score in the data frame.
lineages	A character vector, specifying the lineages used for common essential gene selection.
method	A character, indicating which correlation coefficient is to be used for the test. One of "pearson", "kendall", or "spearman".

Value

A data frame with correlation and test p.value.

Author(s)

Wubing Zhang

Examples

```
dd.rra = ReadRRA(rra.gene_summary)
## Not run:
  rra.omit = OmitCommonEssential(dd.rra)
  depmap_similarity = ResembleDepmap(rra.omit)
  head(depmap_similarity)

## End(Not run)
```

retrieve_gs

Update genesets from source database

Description

Update genesets from source database

Usage

```
retrieve_gs(type = c("KEGG", "REACTOME", "CORUM"), organism = "hsa")
```

Arguments

type	A vector of databases, such as KEGG, REACTOME, CORUM.
organism	'hsa' or 'mmu'.

Value

save data to local library.

Author(s)

Wubing Zhang

rra.gene_summary	<i>Gene summary data generated by running MAGeCK RRA</i>
------------------	--

Description

The gene summary results generated by running MAGeCK on CRISPR screens.

Usage

```
data("rra.gene_summary")
```

Format

A data frame.

References

<https://www.ncbi.nlm.nih.gov/pubmed/25494202> <https://www.ncbi.nlm.nih.gov/pubmed/25476604>

Examples

```
data("rra.gene_summary")  
head(rra.gene_summary)
```

rra.sgrna_summary	<i>sgRNA summary data generated by running MAGeCK RRA</i>
-------------------	---

Description

The sgRNA summary results generated by running 'mageck test' on CRISPR screens.

Usage

```
data("rra.sgrna_summary")
```

Format

A data frame.

References

<https://www.ncbi.nlm.nih.gov/pubmed/25494202> <https://www.ncbi.nlm.nih.gov/pubmed/25476604>

Examples

```
data(rra.sgrna_summary)  
head(rra.sgrna_summary)
```

ScatterView	<i>Scatter plot</i>
-------------	---------------------

Description

Scatter plot supporting groups.

Usage

```
ScatterView(data, x = "x", y = "y", label = 0, model = c("none",
  "ninesquare", "volcano", "rank")[1], x_cut = NULL, y_cut = NULL,
  slope = 1, intercept = NULL, auto_cut = FALSE,
  auto_cut_x = auto_cut, auto_cut_y = auto_cut,
  auto_cut_diag = auto_cut, groups = NULL, group_col = NULL,
  groupnames = NULL, label.top = TRUE, top = 0, toplabels = NULL,
  display_cut = FALSE, color = NULL, shape = 16, size = 1,
  main = NULL, xlab = x, ylab = y, legend.position = "none", ...)
```

Arguments

data	Data frame.
x	A character, specifying the x-axis.
y	A character, specifying the y-axis.
label	An integer or a character specifying the column used as the label, default value is 0 (row names).
model	One of "none" (default), "ninesquare", "volcano", and "rank".
x_cut	An one or two-length numeric vector, specifying the cutoff used for x-axis.
y_cut	An one or two-length numeric vector, specifying the cutoff used for y-axis.
slope	A numeric value indicating slope of the diagonal cutoff.
intercept	A numeric value indicating intercept of the diagonal cutoff.
auto_cut	Boolean, take 1.5 fold standard deviation as cutoff.
auto_cut_x	Boolean, take 1.5 fold standard deviation as cutoff on x-axis.
auto_cut_y	Boolean, take 1.5 fold standard deviation as cutoff on y-axis.
auto_cut_diag	Boolean, take 1.5 fold standard deviation as cutoff on diagonal.
groups	A character vector specifying groups. Optional groups include "top", "mid", "bottom", "left", "center", "right", "topleft", "topcenter", "topright", "midleft", "midcenter", "midright", "bottomleft", "bottomcenter", "bottomright".
group_col	A vector of colors for specified groups.
groupnames	A vector of group names to show on the legend.
label.top	Boolean, specifying whether label top hits.
top	Integer, specifying the number of top terms in the groups to be labeled.
toplabels	Character vector, specifying terms to be labeled.
display_cut	Boolean, indicating whether display the dashed line of cutoffs.
color	A character, specifying the column name of color in the data frame.

shape	A character, specifying the column name of shape in the data frame.
size	A character, specifying the column name of size in the data frame.
main	Title of the figure.
xlab	Title of x-axis
ylab	Title of y-axis.
legend.position	Position of legend, "none", "right", "top", "bottom", or a two-length vector indicating the position.
...	Other available parameters in function 'geom_text_repel'.

Value

An object created by ggplot, which can be assigned and further customized.

Author(s)

Wubing Zhang

Examples

```
dd = ReadBeta(mle.gene_summary)
ScatterView(dd, x = "dms0", y = "plx", label = "Gene",
x_cut = 1, y_cut = 1, groups = "topright", top = 5, display_cut = TRUE)
```

Selector	<i>Select signatures from candidate list (according to the consistence in most samples).</i>
----------	--

Description

Select signatures from candidate list (according to the consistence in most samples).

Usage

```
Selector(mat, cutoff = 0, type = "<", select = 0.8)
```

Arguments

mat	Data matrix, each row is candidates (genes), each column is samples.
cutoff	Cutoff to define the signatures.
type	Direction to select signatures.
select	Proportion of samples in which signature is selected.

Value

An list containing two elements, first is selected signature and second is a ggplot object.

Examples

```
mat = matrix(rnorm(1000*30), 1000, 30)
rownames(mat) = paste0("Gene", 1:1000)
colnames(mat) = paste0("Sample", 1:30)
hits = Selector(mat, select = 0.68)
print(hits$p)
```

sgRankView

*View sgRNA rank.***Description**

View sgRNA rank.

Usage

```
sgRankView(df, gene = NULL, top = 3, bottom = 3, neg_ctrl = NULL,
  binwidth = 0.3, interval = 0.1, bg.col = "gray90",
  filename = NULL, width = 5, height = 3.5, ...)
```

Arguments

df	A data frame, which contains columns of 'sgrna', 'Gene', and 'LFC'.
gene	Character vector, specifying genes to be plotted.
top	Integer, specifying number of top genes to be plotted.
bottom	Integer, specifying number of bottom genes to be plotted.
neg_ctrl	A vector specifying negative ctrl genes.
binwidth	A numeric value specifying the bar width.
interval	A numeric value specifying the interval length between each bar.
bg.col	A character value specifying the background color.
filename	Figure file name to create on disk. Default filename="NULL", which means no output.
width	As in ggsave.
height	As in ggsave.
...	Other available parameters in function 'ggsave'.

Value

An object created by ggplot.

Author(s)

Yihan Xiao

Examples

```
data(rra.sgrna_summary)
sgrra = ReadsgRRA(rra.sgrna_summary)
sgRankView(sgrra)
```

SquareView

*Scatter plot of 9-Square***Description**

Plot a scatter plot with Control beta score as x-axis and Treatment beta score as y-axis, and colored treatment related genes.

Usage

```
SquareView(beta, ctrlname = "Control", treatname = "Treatment",
  label = 0, label.top = TRUE, top = 5, genelist = c(),
  x_cutoff = NULL, y_cutoff = NULL, intercept = NULL,
  groups = c("midleft", "topcenter", "midright", "bottomcenter"),
  groupnames = paste0("Group", 1:length(groups)), main = NULL,
  filename = NULL, width = 6, height = 4, ...)
```

Arguments

beta	Data frame, including columns of <i>ctrlname</i> and <i>treatname</i> , with Gene Symbol as rowname.
ctrlname	A character, specifying the names of control samples.
treatname	A character, specifying the name of treatment samples.
label	An integer or a character specifying the column used as the label, default value is 0 (row names).
label.top	Boolean, whether label the top selected genes, default label the top 10 genes in each group.
top	Integer, specifying the number of top selected genes to be labeled. Default is 5.
genelist	Character vector, specifying labeled genes.
x_cutoff	An one or two-length numeric vector, specifying the cutoff used for x-axis.
y_cutoff	An one or two-length numeric vector, specifying the cutoff used for y-axis.
intercept	An one or two-length numeric vector, specifying the intercept of diagonal.
groups	A character vector, specifying which group to be colored. Optional groups include "topleft", "topcenter", "topright", "midleft", "midright", "bottomleft", "bottomcenter", "bottomright".
groupnames	A character vector, specifying group names.
main	As in 'plot'.
filename	Figure file name to create on disk. Default filename="NULL", which means don't save the figure on disk.
width	As in ggsave.
height	As in ggsave.
...	Other available parameters in function 'ggsave'.

Value

An object created by ggplot, which can be assigned and further customized.

Author(s)

Wubing Zhang

See Also[ScatterView](#)**Examples**

```
data(mle.gene_summary)
# Read beta score from gene summary table in MAGECK MLE results
dd = ReadBeta(mle.gene_summary)
SquareView(dd, ctrlname = "dms0", treatname = "plx", label = "Gene")
```

TransGeneID

*Gene ID conversion between ENTREZID and SYMBOL***Description**

Gene ID conversion between ENTREZID and SYMBOL

Usage

```
TransGeneID(genes, fromType = "Symbol", toType = "Entrez",
  organism = "hsa", fromOrg = organism, toOrg = organism,
  ensemblHost = "www.ensembl.org", update = FALSE)
```

Arguments

genes	A character vector, input genes to be converted.
fromType	The input ID type, one of "entrez", "symbol"(default), "hgnc", "ensembl", "full-name" and "uniprotswissprot"; you can also input other valid attribute names for biomaRt. Look at the code in examples to check valid attributes.
toType	The output ID type, similar to 'fromType'.
organism	"hsa"(default), "mmu", "bta", "cfa", "ptr", "rno", and "ssc" are optional.
fromOrg	"hsa", "mmu", "bta", "cfa", "ptr", "rno", and "ssc" are optional (Only used when transform gene ids between organisms).
toOrg	"hsa"(default), "mmu", "bta", "cfa", "ptr", "rno", and "ssc" are optional (Only used when transform gene ids between organisms).
ensemblHost	String, specifying ensembl host, you can use 'listEnsemblArchives()' to show all available Ensembl archives hosts.
update	Boolean, specifying whether update built-in gene annotation (needs network and takes time).

Value

A character vector, named by unique input gene ids.

Author(s)

Wubing Zhang

Examples

```
TransGeneID("HLA-A", organism="hsa")
TransGeneID("H2-K1", toType="Symbol", fromOrg = "mmu", toOrg = "hsa")
```

ViolinView

*Violin plot***Description**

Plots the violin of beta scores in Control and Treatment samples.

Usage

```
ViolinView(beta, samples = NULL, main = NULL, ylab = "Beta Score",
  filename = NULL, width = 5, height = 4, ...)
```

Arguments

beta	Data frame, , including samples as columns.
samples	Character, specifying the name of samples to be compared.
main	As in 'plot'.
ylab	As in 'plot'.
filename	Figure file name to create on disk. Default filename="NULL", which means don't save the figure on disk.
width	As in ggsave.
height	As in ggsave.
...	Other available parameters in function 'ggsave'.

Value

An object created by ggplot, which can be assigned and further customized.

Author(s)

Wubing Zhang

See Also

[DensityView](#)

Examples

```

data(mle.gene_summary)
# Read beta score from gene summary table in MAGECK MLE results
dd = ReadBeta(mle.gene_summary)
ViolinView(dd, samples=c("dms0", "plx"))
#or
ViolinView(dd[, c("dms0", "plx")])

```

VolcanoView

*Volcano View***Description**

Volcano plot

Usage

```

VolcanoView(df, x = "logFC", y = "adj.P.Val", Label = NA, top = 5,
  topnames = NULL, x_cutoff = log2(1.5), y_cutoff = 0.05,
  mycolour = c("gray80", "#e41a1c", "#377eb8"), alpha = 0.6,
  force = 0.1, main = NULL, xlab = "Log2 Fold Change",
  ylab = "-Log10(Adjust.P)", filename = NULL, width = 4,
  height = 2.5, ...)

```

Arguments

df	Data frame
x	Colname of df specifying x-axis in Volcano figure, 'logFC' (default).
y	Colname of df specifying y-axis in Volcano figure, 'adj.P.Val' (default), which will be plot after log10 transformation.
Label	Colname of df specifying labeled terms in Volcano figure.
top	Integer, the number of top significant terms to be labeled.
topnames	Character vector, indicating interested terms to be labeled.
x_cutoff	Cutoff of x-axis.
y_cutoff	Cutoff of y-axis.
mycolour	A color vector, specifying colors of non-significant, significant up and down-regulated genes.
alpha	Parameter in ggplot.
force	Parameter for geom_text_repel.
main	Title of volcano figure.
xlab	Label of x-axis in figure.
ylab	Label of y-axis in figure.
filename	Figure file name to create on disk. Default filename="NULL", which means don't save the figure on disk.
width	Width of figure.
height	Height of figure.
...	Other available parameters in ggsave.

Value

An object created by ggplot, which can be assigned and further customized.

Author(s)

Wubing Zhang

Examples

```
data(rra.gene_summary)
rra = ReadRRA(rra.gene_summary)
VolcanoView(rra, x = "Score", y = "FDR", Label = "id")
```

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