

Package: moal (via r-universe)

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Title Multi Omic Analysis at Lab

Version 1.2.2

Description Multi Omic Analysis at Lab.

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Depends foreach, ggplot2, igraph, magrittr, R (>= 4.3)

Imports broom, colourvalues, dendextend, doParallel, dplyr, fgsea, forcats, ggforce, ggpubr, ggrepel, gplots, graphics, grDevices, gridExtra, limma, moalannotensg, moalannotensp, moalannotenst, moalannotgene, moalstringbdr, moalstringdbhs, moalstringdbmm, moalstringdboa, moalstringdbrn, moalstringdbss, parallel, plyr, Rgraphviz, rlang, scales, stats, stringr, tidyselect, utils

Suggests knitr, rmarkdown

VignetteBuilder knitr

Encoding UTF-8

Roxygen list(markdown = TRUE)

RoxygenNote 8.0.0

Config/pak/sysreqs

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annot	<i>Annotation function for Symbol, NCBI or Ensembl IDs</i>
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Description

Annotation function for Symbol, NCBI or Ensembl IDs

Usage

```
annot(
  symbollist = NULL,
  species = NULL,
  ortholog = F,
  dboutput = "ncbi",
  idtype = NULL
)
```

Arguments

symbollist	character list of IDs or Symbols
species	character species 'hs' 'mm' 'rn' 'dr' 'ss' (see details for complete list)
ortholog	logical if TRUE return homo sapiens ortholog for choosen species
dboutput	character database used for output 'ncbi'(default) or 'ebi'
idtype	character database ID accepted: 'SYMBOL'(default), 'GENE', 'ENST', 'ENSG', 'ENSP'

Details

Use `moal:::orthoinfo` to see complete species list

Value

data.frame

Author(s)

Florent Dumont florent.dumont@universite-paris-saclay.fr

Examples

```
# not run
# annot(Symbol)
```

ena

Gene set enrichment analysis and interaction network

Description

Gene set enrichment analysis and interaction network

Usage

```
ena(
  omicdata = NULL,
  gmtfiles = NULL,
  species = "hs",
  dat = NULL,
  factor = NULL,
  filtergeneset = NULL,
  threshold = 1,
  topdeg = 100,
  rangedeg = NULL,
  topena = 50,
  topgeneset = 50,
  intmaxdh = 5000,
  nodesize = 0.6,
  bg = 25000,
  doena = TRUE,
  gsearank = "logfc",
  gseatail = "twotail",
  layout = 1,
  mings = 5,
  maxgs = 700,
  overlapmin = 2,
  addratioena = TRUE,
  addenrankbarplot = TRUE,
  dotopnetwork = TRUE,
  dotopgenesetnetwork = FALSE,
  dogmtgenesetnetwork = FALSE,
  dotopheatmap = TRUE,
  dotopgenesetheatmap = TRUE,
  dogmtgenesetheatmap = TRUE,
  path = NULL,
  dirname = NULL,
  dopar = TRUE
)
```

Arguments

omicdata	character data.frame see details
gmtfiles	character gmt files list path
species	character hs mm rn dr ss
dat	data.frame file paths
factor	factor factor for heatmap color
filtergeneset	character list to filter MSigDB geneset collection
threshold	numeric pval 0.05 fc 1.5 by default see details
topdeg	numeric top feature to plot on network
rangedeg	numeric top DEGs from 1 to topdeg by rangedeg to plot on network
topena	numeric top geneset for ena plot
topgeneset	numeric top geneset number to plot on network
intmaxdh	numeric maximum number of interaction to use for Davidson and Harel algorithm layout
nodesize	numeric change Symbol size
bg	numeric background used for functional analysis over-representation test
doena	logical do MSigDB enrichment analysis
gsearank	character to choose gsea rank type among fc (by default) logration logfc sqrt
gseatail	character to choose gsea twotail (by default) or onetail
layout	numeric for layout neetwork 1 fr by default 2 dh 3 tree 4 circle 5 grid 6 sphere
mings	numeric minimal size of a gene set
maxgs	numeric maximal size of a gene set
overlapmin	numeric minimal overlap to keep for gene set analysis
addratioena	logical if TRUE add overlap and geneset size on enrichment barplot
addenarankbarplot	logical if TRUE add ena barplot ranked by NES score
dotopnetwork	logical do top networks
dotopgenesetnetwork	logical do geneset networks
dogmtgenesetnetwork	logical do keyword networks
dotopheatmap	logical do top heatmap
dotopgenesetheatmap	logical do geneset heatmap
dogmtgenesetheatmap	logical do keyword heatmap
path	character for relative path of output directory
dirname	character name for output
dopar	logical TRUE for parallelization

Details

omicdata needs a data.frame with at list 4 column: rowID, (p-values,fold-change) x N and Symbol annotation.

Symbol list are accepted to make ORA enrichment analysis.

To generate heatmap dat and factor parameter are needed. dat accepted complete matrix with rowID for first column.

dat row IDs must match with omicdata row IDs.

Make MSigDB enrichment analysis using GSEA method for non filtering list as input (> 2000)

Make MSigDB Over-Representation enrichment analysis (ORA) using Fisher exact test for list < 2000

Generate STRINGDB interaction network and heatmap for top geneset according to topena par (80 by default)

Only features with p-values < 0.05 et fold-change > 1.1 are displayed on geneset heatmaps (threshold = 1 by default).

See omic function details to display all threshold

Value

file with enrichment analysis results

Author(s)

Florent Dumont florent.dumont@universite-paris-saclay.fr

Examples

```
# not run
# ena( omicdata , species = "mm")
```

heatmap

Heatmap

Description

To make a heatmap

Usage

```
heatmap(  
  dat = NULL,  
  factor = NULL,  
  method = "complete",  
  dendrogram = "both",  
  k = NULL,  
  labCol = "",
```

```

    cexCol = 0.5,
    labRow = "",
    cexRow = NULL,
    cexlegend = 0.65,
    keysize = 0.9,
    keycolor = c("darkgreen", "orange", "darkred"),
    parmar = c(5, 4, 5, 6),
    scale = "row"
  )

```

Arguments

dat	matrix numeric
factor	factor
method	character
dendrogram	character to display 'none', 'row', 'column' or 'both' (by default) dendrograms
k	numeric number of clusters to colorize for rows
labCol	character
cexCol	numeric
labRow	Character
cexRow	numeric
cexlegend	numeric
keysize	numeric
keycolor	character of 3 for low mid high value of the key
parmar	numeric 4 values for margin sizes
scale	numeric standardize row by default and column or none accepted

Details

To make a heatmap from a matrix or a data.frame

Value

no returned value

Author(s)

Florent Dumont florent.dumont@universite-paris-saclay.fr

Examples

```

# not run
# library(magrittr)
# data(sif1)
# data(mat1)
# mat1 %>% heatmap(sif1$F3)

```

input	<i>import tab file in data.frame</i>
-------	--------------------------------------

Description

import tab file in data.frame

Usage

```
input(filename, sep = "\t", quote = "")
```

Arguments

filename	character path to the file to read
sep	character for field separator
quote	character for field quote

Details

wrapper of read.table function for tabular separated files

Value

data.frame

Author(s)

Florent Dumont florent.dumont@universite-paris-saclay.fr

Examples

```
# not run  
# input( "filename" ) -> dt
```

norm	<i>Normalization function</i>
------	-------------------------------

Description

normalization and log2

Usage

```
norm(dat, method = NULL, log = TRUE)
```

Arguments

dat	data.frame
method	character apply quantile normalization by default see details
log	logical apply log base 2

Details

for see limma normalizeBetweenArrays method

Value

data.frame

Author(s)

Florent Dumont florent.dumont@universite-paris-saclay.fr

Examples

```
# not run  
# norm(dt)
```

omic

Omic bioanalysis workflow

Description

Omic function workflow description:

- Quality controls and unsupervised analysis: histogram, box plot, PCA and sample clustering.
- Supervised analysis: analysis of variance (ANOVA) and filter application.
- Unsupervised analysis for selected features: row clustering, PCA and pattern search across factor levels.
- Graph generation for selected feature: volcanoplots, heatmaps, lineplots, boxplots, PCA
- Functional analysis: MSigDB enrichment analysis and STRINGDB interaction network

See help("omic") section to test workflow with internal GEO data set GSE65055 and reproduce enrichment results for chromosome cytogenetic bands (doi: 10.1111/cge.12731)

Usage

```
omic(  
  dat = NULL,  
  sif = NULL,  
  annot = NULL,  
  species = "hs",  
  model = NULL,  
  paired = NULL,  
  nested = NULL,  
  batch = NULL,  
  addfactor = NULL,  
  doqc = TRUE,  
  threshold = c(1, 2, 3, 4, 9, 10, 11, 12),  
  padj = "none",  
  logratio = FALSE,  
  dopattern = TRUE,  
  dovenn = FALSE,  
  docluster = TRUE,  
  nc = c(2, 3, 6, 12),  
  maxclusterheatmap = 5000,  
  doheatmap = TRUE,  
  heatmapcluster = "row",  
  maxheatmap = 2000,  
  minheatmap = 3,  
  dovolcanoplot = TRUE,  
  nbgenevolc = 5,  
  dolineplot = TRUE,  
  doboxplotrow = TRUE,  
  doena = TRUE,  
  gsearank = "logfc",  
  gseatail = "twotail",  
  topdeg = 100,  
  topena = 50,  
  doenaora = FALSE,  
  gmtfiles = NULL,  
  filtergeneset = NULL,  
  bg = 25000,  
  dotopnetwork = TRUE,  
  dotopheatmap = TRUE,  
  layout = 2,  
  mings = 5,  
  maxgs = 700,  
  overlapmin = 2,  
  addenarankbarplot = TRUE,  
  dotopgenesetnetwork = FALSE,  
  dotopgenesetheatmap = TRUE,  
  dogmtgenesetnetwork = FALSE,  
  dogmtgenesetheatmap = TRUE,
```

```

crosscompint = FALSE,
sample = NULL,
seed = 123679,
dopar = NULL,
path = ".",
dirname = NULL,
zip = FALSE,
remove = FALSE
)

```

Arguments

<code>dat</code>	data.frame normalize data table with rowID for first column
<code>sif</code>	data.frame sample information file including model factors
<code>annot</code>	data.frame annotation with Symbol column for functional analysis
<code>species</code>	character available species: hs mm rn ss pt bt oa dr gg xt dm ce
<code>model</code>	character anova model factors (see details)
<code>paired</code>	character factor for paired design
<code>nested</code>	character factor for nested design
<code>batch</code>	character factor for batch effect design
<code>addfactor</code>	character additionnal factors
<code>doqc</code>	logical quality controls
<code>threshold</code>	numeric vector from 1 to 24 (see details)
<code>padj</code>	character fdr by default for Benjamini-Hochberg false discovery correction
<code>logratio</code>	logical change fc (by default) in log2ratio
<code>dopattern</code>	logical search relevant pattern across levels factor
<code>dovenn</code>	logical venn diagram
<code>docluster</code>	logical row hierarchical clustering using pearson correlation
<code>nc</code>	numeric number of clusters to cut in dendrogramm
<code>maxclusterheatmap</code>	numeric max row for cluster analysis
<code>doheatmap</code>	logical do heatmaps for all lists
<code>heatmapcluster</code>	character row clustering only by default both accepted
<code>maxheatmap</code>	numeric max rows for heatmap
<code>minheatmap</code>	numeric min rows for heatmap
<code>dovolcanoplot</code>	logical make volcanoplot for each threshold
<code>nbgenevolc</code>	numeric number of Symbol to display in volcanoplot
<code>dolineplot</code>	logical do lineplot for significant features
<code>doboxplotrow</code>	logical do boxplot for significant features with Kruskal
<code>doena</code>	logical msigdb enrichment analysis using gsea method without filtering

<code>gsearank</code>	character to choose gsea rank type among fc (by default) logrotation logfc sqrt
<code>gseatail</code>	character to choose gsea twotail (by default) or onetail
<code>topdeg</code>	numeric top DEGs number to plot on network
<code>topena</code>	numeric top geneset for ena plot
<code>doenaora</code>	logical msigdb enrichment analysis using ora method for diff list
<code>gmtfiles</code>	character gmt files list path
<code>filtergeneset</code>	character regular expression to filter collection geneset (e.g. "reactomelfft")
<code>bg</code>	numeric background used for functional analysis over-representation test
<code>dotopnetwork</code>	logical do top networks
<code>dotopheatmap</code>	logical do top heatmap
<code>layout</code>	numeric for layout neetwork 1 fr by default 2 dh 3 tree 4 circle 5 grid 6 sphere
<code>mings</code>	numeric minimal size of a gene set
<code>maxgs</code>	numeric maximal size of a gene set
<code>overlapmin</code>	numeric minimal overlap to keep for gene set analysis
<code>addenarankbarplot</code>	logical if TRUE add ena barplot ranked by NES score
<code>dotopgenesetnetwork</code>	logical do geneset networks
<code>dotopgenesetheatmap</code>	logical do geneset heatmap
<code>dogmtgenesetnetwork</code>	logical do keyword networks
<code>dogmtgenesetheatmap</code>	logical do keyword heatmap
<code>crosscompint</code>	logical add cross comparison to results for interaction model
<code>sample</code>	numeric analysis using random subset
<code>seed</code>	numeric seed for random function
<code>dopar</code>	numeric core number
<code>path</code>	character results directory path
<code>dirname</code>	character results directory name
<code>zip</code>	logical compress results directory if TRUE
<code>remove</code>	logical remove uncompress results directory if TRUE

Details

Use `moal::env()` to load required libraries before `moal::omic()` (see example)

Use `input()` function to import and analyse your own data starting from tsv file (or csv with `sep = ","`)

`dat` must have one IDs columns in the same order than annotations.

Use `annot()` function for annotation with Symbol, NCBI, Ensembl IDs.

sif must contains column with description sample corresponding to anova factor analysis.

sif rows must have the same number of samples in the same order that in the dat table.

Experimental design examples for model parameters:

- 1-way anova: model = "TREATMENT"
- 2-ways anova: model = "PHENOTYPE+TREATMENT"
- 2-ways anova with interaction: model = "TREATMENT+TIME+TREATMENT*TIME"
- 2-ways anova with paired factor: model = "TREATMENT", paired = "CASE"
- 2-ways anova with batch factor: model = "TREATMENT", batch = "BATCH"
- 2-ways anova with nested factor: model = "TREATMENT", nested = "CASEinTREATMENT"
- 3-ways or 4-ways anova (without interaction): model = "PHENOTYPE+TREATMENT+AGE"

For paired, batch and nested design, remove batch effect from limma package are used to calculate fold-change

Use dopar = 2 to decrease computing resources.

Use sample for random subset analysis.

To see complete threshold list: `moal:::thresholdlist %>% lapply("[",c(1,2)) %>% unlist %>% matrix(ncol=2,byrow = T) %>% data.frame %>% setNames(c("pval","fc"))`

Annotation updates: 22-04-2025 for gene and ensembl, MSigDB 2024.1.Hs, StringDB 12.0

Value

omic results directory

Author(s)

Florent Dumont florent.dumont@universite-paris-saclay.fr

Examples

```
# # Test workflow with internal GEO data set GSE65055
# # and reproduce enrichment results for chromosome cytogenetic bands (doi: 10.1111/cge.12731)
# # loading libraries:
# library(moal);moal::env()
# # loading data:
# moal:::GSE65055normdata -> dat
# moal:::GSE65055sampledata -> sif
# # Ordering factors for pairwise comparisons which compute contrast p-values and fold-changes.
# sif$ANEUPLOIDY %>% ordered(c("Control","T13","T18","T21")) -> sif$ANEUPLOIDY
# sif$TISSUE %>% as.factor -> sif$TISSUE
# # annotation
# dat$rowID %>% moal:::annot(species="hs",idtype="GENE",dboutput="ncbi") -> annot
# # omic analysis
# moal:::omic(dat,sif,annot,species="hs",model="ANEUPLOIDY",batch="TISSUE",dirname="GSE65055")
```

output	<i>export data.frame in tab file</i>
--------	--------------------------------------

Description

export data.frame in tab file

Usage

```
output(dt, filename)
```

Arguments

dt	data.frame
filename	character

Author(s)

Florent Dumont florent.dumont@universite-paris-saclay.fr

Examples

```
# not run  
# output( dt )
```

qc	<i>Quality Controls</i>
----	-------------------------

Description

Descriptive analysis applied on column:

- histogram, boxplot, hierarchical clustering and PCA for column

Usage

```
qc(  
  dat,  
  sif = NULL,  
  dooutputinput = FALSE,  
  dohisto = TRUE,  
  doboxplot = TRUE,  
  dohc = TRUE,  
  dopca = TRUE,  
  breaks = 70,  
  dirname = NULL,  
  path = "."  
)
```

Arguments

dat	data.frame first column for rowID column
sif	data.frame sample information file
dooutputinput	logical if TRUE (by default) export input data
dohisto	logical if TRUE (by default) do histogram
doboxplot	logical if TRUE (by default) do boxplot
dohc	logical if TRUE (by default)do hierarchical clustering
dopca	logical if TRUE (by default) do PCA
breaks	numeric break number for histogramm function
dirname	character
path	character

Details

dat row must be equal to sif row

Value

directory including analysis pdf plots

Author(s)

Florent Dumont florent.dumont@universite-paris-saclay.fr

Examples

```
# not run
# qc(dat,sif)
```

venn

Venn diagramm

Description

To make a Venn diagramm of 2, 3 or 4 lists

Usage

```
venn(
  list = NULL,
  listnames = NULL,
  returnlist = F,
  title = "Venn Diagram",
  plot = T,
  export = F,
```

```
    path = ".",
    dirname = "venn"
  )
```

Arguments

list	list of 2 , 3 or 4 character vector
listnames	character list names to display on graph
returnlist	logical
title	character title to display on graph
plot	logical to display the plot or not
export	logical export lists in a directory
path	character
dirname	character name of the directory created when export = T

Value

venn plot and new lists generated by venn.

Author(s)

Florent Dumont florent.dumont@universite-paris-saclay.fr

Examples

```
# library(magrittr)
# list(
#   c(letters[6:20] , letters[25] ) ,
#   letters[1:15] ,
#   c( letters[2:5] , letters[8:23] ) ) %>% moal::venn(.)
```

volcanoplot

Volcanoplot

Description

Do volcanoplot

Usage

```
volcanoplot(
  dat = NULL,
  pval = 0.05,
  fc = 1.5,
  topgenename = TRUE,
  topgenenamen = 5,
```

```
    genenamelist = NULL,  
    genenamesize = 2,  
    title = "Volcanoplot"  
  )
```

Arguments

dat	data.frame table with 4 columns (see details)
pval	numeric p-value threshold
fc	numeric fold-change threshold
topgenename	logical display gene label TRUE by default
topgenenamen	numeric increase number of gene label
genenamelist	character vector of gene list to label
genenamesize	numeric label size for gene name
title	character

Details

dat parameter must have 4 columns: rowID , p_AvsB , fc_AvsF and Symbol.

Value

no returned value

Author(s)

Florent Dumont florent.dumont@universite-paris-saclay.fr

Examples

```
# not run  
# data.frame(rowID,p_AvsB,fc_AvsB,Symbol) -> dat  
# volcanoplot(dat)
```

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