Package: Grouphmap (via r-universe)

August 26, 2024

Type Package

Title 'Grouphmap' is an Automated One-Step Common Analysis of Batch Expression Profile

Version 1.0.0

Author Yuchen Sun [aut, cre]

Maintainer Yuchen Sun < yuchensun 2436@163.com>

Depends R (>= 4.2.0)

Description The 'Grouphmap' was implemented in R, an open-source programming environment, and was released under the provided website. The difference analysis is based on the 'limma' package, which can cover gene and protein expression profiles (Reference: Matthew E Ritchie, Belinda Phipson, Di Wu, Yifang Hu, Charity W Law, Wei Shi, Gordon K Smyth (2015) <doi:10.1093/nar/gkv007>). The GO enrichment analysis is based on the 'clusterProfiler' package and supports three common species: human, mouse, and yeast (Reference: Guangchuang Yu, Li-Gen Wang, Yanyan Han, Qing-Yu He (2012) <doi:10.1089/omi.2011.0118>). The results of batch difference analysis and enrichment analysis are output in separate folders for easy viewing and further visualization of the results during the process. The results returned a heatmap in R and exported to 3 folders named DEG, go, and merge.

License Artistic-2.0

Encoding UTF-8

Imports limma, clusterProfiler, dplyr, org.Mm.eg.db, pheatmap, stringr, stats, utils, grDevices

RoxygenNote 7.2.3

References Matthew E Ritchie, Belinda Phipson, Di Wu, Yifang Hu, Charity W Law, Wei Shi, Gordon K Smyth (2015) <doi:10.1093/nar/gkv007>; Guangchuang Yu, Li-Gen Wang, Yanyan Han, Qing-Yu He (2012) <doi:10.1089/omi.2011.0118>

NeedsCompilation no

2 ghmap

Date/Publication 2023-09-04 09:50:02 UTC **Repository** https://syc2436.r-universe.dev **RemoteUrl** https://github.com/cran/Grouphmap

RemoteRef HEAD

RemoteSha d7842c6f64f44d00df9cf89041fa24a734d2fed9

Contents

	ghmap	2
Index		3
ghmap	Grouphmap: Automated one-step common analysis of Batch expression profile	_

Description

Grouphmap: Automated one-step common analysis of Batch expression profile

Usage

```
ghmap(Path, conRep, treRep, OrgDb, TYPE, UP = TRUE, cutree, TOP)
```

Arguments

Path	"File storage path." Two groups are one file, and the control group should before
	the treatment group.
conRep	Number of repetitions in control group.

treRep Number of repetitions in control group.

Number of repetitions in treatment group.

OrgDb org.Mm.eg.db, org.Hs.eg.db, and org.Sc.sgd.db. Please library().

TYPE "SYMBOL","ENSEMBOL"...
UP up is TRUE and down is FALSE

cutree heatmap can be devided multiple modules that make the functional difference

and similarity of those group to be obvious

TOP the numeric. Such as 10 is the top 10 of GO analysis in each file.

Value

p1

Examples

```
library(org.Mm.eg.db)
Path<-system.file("extdata", package = "Grouphmap")
ghmap(Path,2,2,org.Mm.eg.db,"ENSEMBL",FALSE,2,10)</pre>
```

Index

ghmap, 2