## Package 'HTGM'

April 2, 2025

```
Version 1.0
Date 2025-03-26
Title High Throughput 'GoMiner'
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Depends R (>= 4.2.0)
Imports minimalistGODB, GoMiner, grDevices, stats, gplots
LazyData true
LazyDataCompression xz
Description Two papers published in the early 2000's (Zeeberg, B.R., Feng, W., Wang, G. et al.
     (2003) <doi:10.1186/gb-2003-4-4-
     r28>) and (Zeeberg, B.R., Qin, H., Narashimhan, S., et al. (2005)
     <doi:10.1186/1471-2105-6-168>) implement 'GoMiner' and 'High Throughput GoMiner'
     ('HTGM') to map lists of genes to the Gene Ontology (GO) <https:
     //geneontology.org>. Until recently,
     these were hosted on a server at The National Cancer Institute (NCI). In order to continue
     providing these services to the bio-medical community, I have developed stand-alone versions.
     The current package 'HTGM' builds upon my recent package 'GoMiner'.
     The output of 'GoMiner' is a heatmap showing the relationship of a single list
     of genes and the significant categories into which they map. 'High Throughput GoMiner'
     ('HTGM') integrates the results of the individual 'GoMiner' analyses. The output of 'HTGM'
     is a heatmap showing the relationship of the significant categories derived from each gene list.
     The heatmap has only 2 axes, so the identity of the genes are unfortunately
     ``integrated out of the equation." Because the graphic for the heatmap is implemented
     in Scalable Vector Graphics (SVG) technology, it is relatively easy to hyperlink each
     picture element to the relevant list of genes. By clicking on the desired picture
     element, the user can recover the ``lost" genes.
License GPL (>= 2)
Encoding UTF-8
VignetteBuilder knitr
Suggests knitr, rmarkdown, testthat (>= 3.0.0)
RoxygenNote 7.3.2
```

## Config/testthat/edition 3

NeedsCompilation no

Repository CRAN

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Hous	eeping_Genes HTGM data set	

## Description

HTGM data set

## Usage

data(Housekeeping\_Genes)

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## Description

driver to invoke GoMiner for multiple studies, and integrate the results in a categories versus study hyperlinked heatmap

## Usage

```
HTGM(
  title = NULL,
  dir = tempdir(),
  sampleLists,
  GOGOA3,
  ONT,
  enrichThresh = 2,
  countThresh = 5,
  fdrThresh = 0.1,
  nrand = 100,
  verbose = TRUE
)
```

## Arguments

title	character string descriptive title
dir	character string full pathname to the directory acting as result repository
sampleLists	list of character vector of user-supplied genes of interest
GOGOA3	return value of subsetGOGOA()
ONT	$c("molecular\_function","cellular\_component","biological\_process")$
enrichThresh	numerical acceptance threshold for enrichment passed to GoMiner
countThresh	numerical acceptance threshold for gene count passed to GoMiner
fdrThresh	numerical acceptance threshold for fdr passed to GoMiner
nrand	integer number of randomizations passed to GoMiner
verbose	Boolean if TRUE, GoMiner will message some helpful diagnostics

#### Value

returns the matrix of significant categories versus study

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#### **Examples**

```
## Not run:
# this example takes too long to run, and
# GOGOA3.RData is too large to include in the R package
# so I need to load it from a file that is not in the package.
# Since this is in a file in my own file system, I could not
# include this as a regular example in the package.
# This example is given in full detail in the package vignette.
# You can generate GOGOA3.RData using the package 'minimalistGODB'
# or you can retrieve it from https://github.com/barryzee/GO
# load("data/Housekeeping_Genes.RData")
# load("~/GODB_RDATA/GOGOA3.RData")
sampleList<-unique(as.matrix(Housekeeping_Genes[, "Gene.name"]))</pre>
n<-nrow(sampleList)</pre>
sampleLists<-list()</pre>
# test the effect of random sampling of the entire gene set
# this can give an idea of the quality of the GoMiner results
# when the complete gene set is yet to be determined
sampleLists[["1"]]<-sampleList[sample(n,n/2)]</pre>
sampleLists[["2"]]<-sampleList[sample(n,n/2)]</pre>
sampleLists[["3"]]<-sampleList[sample(n,n/2)]</pre>
sampleLists[["4"]]<-sampleList[sample(n,n/2)]</pre>
sampleLists[["5"]]<-sampleList[sample(n,n/2)]</pre>
sampleLists[["ALL"]]<-sampleList</pre>
m<-HTGM(title=NULL,dir=tempdir(),sampleLists,GOGOA3,ONT="biological_process",
enrichThresh=2,countThresh=5,fdrThresh=0.10,nrand=100,verbose=TRUE)
## End(Not run)
```

htgmM

htgmM

#### Description

generate FDR matrix of id versus cat

#### Usage

```
htgmM(1, fdrThresh)
```

## **Arguments**

l list of return values of GoMiner()
fdrThresh numerical acceptance threshold for fdr

#### Value

returns numeric matrix m containing FDR values

hyperGenes 5

#### **Examples**

```
# load("data/x_1.RData")
m<-htgmM(x_1,.1)</pre>
```

hyperGenes

hyperGenes

#### **Description**

populate subdirectory of hyperlinked gene lists

#### Usage

```
hyperGenes(1, dir)
```

#### **Arguments**

1 return value of GoMiner()

dir character string containing path name of results directory

#### Value

returns no value but has side effect of populating subdirectory of hyperlinked gene lists

## **Examples**

```
# x_l<-load("data/x_l.RData")
dir<-tempdir()
print(dir)
hyperGenes(x_l,dir)</pre>
```

hyperlinks

hyperlinks

## Description

driver to add gene list hyperlinks to the HTGM heatmap

#### Usage

```
hyperlinks(s, rownames, colnames)
```

6 pasteHyperlinks

#### **Arguments**

s character path name of the file containing the HTGM svg

rownames character vector of row names colnames character vector of column names

#### Value

returns the path name of the file containing the hyperlinked HTGM svg

#### **Examples**

```
#load("data/x_rn.RData")
#load("data/x_cn.RData")
#load("data/x_svg.RData")
s<-system.file("extdata","x_htgm.svg",package="HTGM")
hyperlinkedFileName<-hyperlinks(s,x_rn,x_cn)
print("hyperlinkedFileName")
print(hyperlinkedFileName)</pre>
```

pasteHyperlinks

pasteHyperlinks

#### **Description**

add gene list hyperlinks to the HTGM heatmap

#### Usage

```
pasteHyperlinks(str, c1, c2)
```

#### **Arguments**

str character a line from the svg that is to have a hyperlink inserted

c1 character list of row names c2 character list of column names

#### Value

returns a line of code to insert into svg

## **Examples**

```
#load("data/x_svgr.RData")
#load("data/x_rnr.RData")
#load("data/x_cnc.RData")
hl<-pasteHyperlinks(x_svgr,x_rnr,x_cnc)</pre>
```

x\_cn 7

x\_cn HTGM data set

## Description

HTGM data set

## Usage

data(x\_cn)

x\_cnc

HTGM data set

## Description

HTGM data set

## Usage

data(x\_cnc)

 $x_1$ 

HTGM data set

## Description

HTGM data set

## Usage

 $data(x_1)$ 

 $x\_{m}$ 

HTGM data set

## Description

HTGM data set

## Usage

data(x\_m)

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x\_rn HTGM data set

## Description

HTGM data set

## Usage

data(x\_rn)

x\_rnr

HTGM data set

## Description

HTGM data set

## Usage

data(x\_rnr)

x\_svg

HTGM data set

## Description

HTGM data set

## Usage

data(x\_svg)

x\_svgr

HTGM data set

## Description

HTGM data set

## Usage

data(x\_svgr)

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