## Package 'KOGMWU'

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Type Package

Title Functional Summary and Meta-Analysis of Gene Expression Data

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#### Description

Rank-based tests for enrichment of KOG (euKaryotic Orthologous Groups) classes with upor down-regulated genes based on a continuous measure. The meta-analysis is based on correlation of KOG delta-ranks across datasets (delta-rank is the difference be-

tween mean rank of genes belonging to a KOG class and mean rank of all other genes). With binary measure (1 or 0 to indicate significant and non-significant genes), one-tailed Fisher's exact test for over-representation of each KOG class among significant genes will be performed.

License GPL-3

**Depends** R ( $\geq$  2.10), pheatmap

NeedsCompilation no

**Repository** CRAN

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KOGMWU-package

Functional summary and meta-analysis of gene expression data

#### Description

Rank-based tests for enrichment of KOG (euKaryotic Orthologous Groups) classes with up- or down-regulated genes based on a continuous measure. The meta-analysis is based on correlation of KOG delta-ranks across datasets (delta-rank is the difference between mean rank of genes belonging to a KOG class and mean rank of all other genes). With binary measure (1 or 0 to indicate significant and non-significant genes), one-tailed Fisher's exact test for over-representation of each KOG class among significant genes will be performed.

#### Details

Package:	KOGMWU
Type:	Package
Version:	1.2
Date:	2019-02-19
License:	GPL-3

The most important function is kog.mwu, which performs a series of Mann-Whitney U tests when given two data tables: one, containing measures of interest for each gene (for example, log fold-change), and another, listing the association of each gene with a KOG class. The KOG class annotations for a collection of genes can be obtained using eggNOG-mapper: http://eggnogdb.embl.de/#/app/emapper. To extract KOG annotations understood by this package out of the eggNOG-mapper output, see here: https://github.com/z0on/emapper\_to\_GOMWU\_KOGMWU

#### Author(s)

Mikhail V. Matz

Maintainer: Mikhail V. Matz <matz@utexas.edu>

#### References

Dixon, G. B., Davies, S. W., Aglyamova, G. V., Meyer, E., Bay, L. K. and Matz, M. V. Genomic determinants of coral heat tolerance across latitudes. Science 2015, 348:1460-1462. eggNOG-mapper to obtain KOG annotations: http://eggnogdb.embl.de/#/app/emapper To extract KOG annotations from eggNOG-mapper output: https://github.com/z0on/emapper\_to\_GOMWU\_KOGMWU

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#### Examples

```
## Not run:
data(adults.3dHeat.logFoldChange)
data(larvae.longTerm)
data(larvae.shortTerm)
data(gene2kog)
# Analyzing adult coral response to 3-day heat stress:
alfc.lth=kog.mwu(adults.3dHeat.logFoldChange,gene2kog)
alfc.lth
# coral larvae response to 5-day heat stress:
1.lth=kog.mwu(larvae.longTerm,gene2kog)
1.lth
# coral larvae response to 4-hour heat stress
1.sth=kog.mwu(larvae.shortTerm,gene2kog)
1.sth
# compiling a table of delta-ranks to compare these results:
ktable=makeDeltaRanksTable(list("adults.long"=alfc.lth,"larvae.long"=l.lth,"larvae.short"=l.sth))
# Making a heatmap with hierarchical clustering trees:
pheatmap(as.matrix(ktable),clustering_distance_cols="correlation")
# exploring correlations between datasets
pairs(ktable, lower.panel = panel.smooth, upper.panel = panel.cor)
# p-values of these correlations in the upper panel:
pairs(ktable, lower.panel = panel.smooth, upper.panel = panel.cor.pval)
# plotting individual delta-rank correlations:
corrPlot(x="adults.long",y="larvae.long",ktable)
corrPlot(x="larvae.short",y="larvae.long",ktable)
## End(Not run)
```

adults.3dHeat.logFoldChange

Heat stress response of adult coral

## Description

Acropora millepora (adult) response to three days of heat stress (31.5oC) log-fold-changes inferred using DESeq package from tag-based RNA-seq data.

#### Usage

data("adults.3dHeat.logFoldChange")

#### Format

A data frame with 44363 observations on the following 2 variables.

gene gene id, a factor with 44363 levels

1fc log fold-change, a numeric vector

corr	Plot
------	------

Plots a pairwise correlation with linear regression line

## Description

Plots Pearson's correlation between two columns in a dataframe, identified by column names. Also plots linear regression line and lists the correlation coefficient (r) and cor.test p-value.

## Usage

corrPlot(x, y, data, ...)

## Arguments

х	Name of the column to form X axis
У	Name of the column to form Y axis
data	The dataframe containing the two columns
	Additional options for plot()

#### References

Dixon GB, Davies SW, Aglyamova GA, Meyer E, Bay LK and Matz MV (2015) Genomic determinants of coral heat tolerance across latitudes.

gene2kog

KOG class annotations

#### Description

KOG class annotations for Acropora millerpoa transcriptome.

#### Usage

data("gene2kog")

## kog.ft

## Format

A data frame with 16175 observations on the following 2 variables.

- V1 a factor with 16175 levels
- V2 a factor with 23 levels

#### Source

https://dl.dropboxusercontent.com/u/37523721/amillepora\_transcriptome\_july2014.zip

#### References

Transcriptome assembly: Moya et al (2012), Mol Ecol 21:2440-2454. Transcriptome annotation: Dixon GB, Davies SW, Aglyamova GA, Meyer E, Bay LK and Matz MV (2015) Genomic determinants of coral heat tolerance across latitudes.

kog.ft

One-tailed Fisher's exact test for KOG enrichment.

#### Description

Accessory function to kog.mwu()

#### Usage

kog.ft(gos)

#### Arguments

gos A dataframe with three columns, 'seq' (gene id),'term' (KOG class) and 'value' (either 0 or 1, indicating significance).

## Value

A dataframe with three columns: 'term', 'nseqs', 'pval' and 'padj'

## Author(s)

Mikhail V. Matz

## References

Dixon GB, Davies SW, Aglyamova GA, Meyer E, Bay LK and Matz MV (2015) Genomic determinants of coral heat tolerance across latitudes.

kog.mwu

#### Description

Determines whether some KOG classes are significantly enriched with up- or down-regulated genes (Mann-Whitney U test for continuous measure), or whether some KOG classes are significantly over-represented among "significant" genes (one-tailed Fisher's exact test for binary measure, 0 or 1).

#### Usage

```
kog.mwu(data, gene2kog, Alternative = "t")
```

#### Arguments

data	Two-column dataframe: gene id, measure of significance.
gene2kog	Two-column dataframe of gene annotations: gene id, KOG class. The gene list can be longer or shorter than the first column in the 'data' item.
Alternative	Tailedness of the Mann-Whitney U test: two-tailed ("t"), greater ("g"), or less ("l")

#### Details

The measure can be continuous (such as log fold change), in which case Mann-Whitney U test will be performed, or binary (1 or 0: significant or not), in which case Fisher's exact test will be performed. The KOG class annotations for a collection of genes can be obtained using Weizhng Li's lab KOG BLAST server.

#### Value

For continuous measure, a dataframe with three columns: term : KOG class nseqs : Number of genes in this class delta.rank : Difference between the mean rank of genes belonging to this KOG class and all other genes pval : p-value of the Mann-Whitney U test padj : p-value adjusted using Benjamini-Hochberg 1995 "fdr" procedure

For binary measure, the output is similar but does not contain the delta.rank column.

#### Author(s)

Mikhail V. Matz <matz@utexas.edu>

#### References

Dixon GB, Davies SW, Aglyamova GA, Meyer E, Bay LK and Matz MV (2015) Genomic determinants of coral heat tolerance across latitudes. Weizhong Li's KOG BLAST server: http://weizhonglab.ucsd.edu/metagenomic-analysis/server/kog/

#### kog.mwut

#### Examples

```
## Not run:
data(adults.3dHeat.logFoldChange)
data(larvae.longTerm)
data(larvae.shortTerm)
data(gene2kog)
# Analyzing adult coral response to 3-day heat stress:
alfc.lth=kog.mwu(adults.3dHeat.logFoldChange,gene2kog)
alfc.lth
# coral larvae response to 5-day heat stress:
1.lth=kog.mwu(larvae.longTerm,gene2kog)
1.lth
# coral larvae response to 4-hour heat stress
l.sth=kog.mwu(larvae.shortTerm,gene2kog)
1.sth
# compiling a table of delta-ranks to compare these results:
ktable=makeDeltaRanksTable(list("adults.long"=alfc.lth,"larvae.long"=l.lth,"larvae.short"=l.sth))
# Making a heatmap with hierarchical clustering trees:
pheatmap(as.matrix(ktable),clustering_distance_cols="correlation")
# exploring correlations between datasets
pairs(ktable, lower.panel = panel.smooth, upper.panel = panel.cor)
# p-values of these correlations in the upper panel:
pairs(ktable, lower.panel = panel.smooth, upper.panel = panel.cor.pval)
# plotting individual delta-rank correlations:
corrPlot(x="adults.long",y="larvae.long",ktable)
corrPlot(x="larvae.short",y="larvae.long",ktable)
## End(Not run)
```

kog.mwut

Mann-Whitney U test for KOG enrichment.

## Description

Accessory function to kog.mwu()

#### Usage

kog.mwut(gos, Alternative = "t")

## Arguments

gos	A dataframe with three columns, 'seq' (gene id),'term' (KOG class) and 'value' (continuous measure, such as log fold-change).
Alternative	Tailedness of the MWU test: two-tailed ("t"), greater-than ("g"), or less-than ("l")

#### Value

A dataframe with three columns: 'term', 'nseqs', 'delta.rank', 'pval' and 'padj'

#### Author(s)

Mikhail V. Matz

#### References

Dixon GB, Davies SW, Aglyamova GA, Meyer E, Bay LK and Matz MV (2015) Genomic determinants of coral heat tolerance across latitudes.

larvae.longTerm Long-term heat stress response of coral larvae

## Description

Acropora millepora (larvae) response to five days of heat stress (31.5oC) log-fold-changes inferred using DESeq package from tag-based RNA-seq data from Meyer et al Mol Ecol 2011,17:3599-3616

## Usage

```
data("larvae.longTerm")
```

#### Format

A data frame with 31844 observations on the following 2 variables.

gene gene id, a factor with 31844 levels

1fc log fold-change, a numeric vector

larvae.shortTerm Short-term heat stress response of coral larvae

#### Description

Acropora millepora (larvae) response to four hours of heat stress (31.5oC) log-fold-changes inferred using DESeq package from tag-based RNA-seq data from Meyer et al Mol Ecol 2011,17:3599-3616.

#### Usage

```
data("larvae.shortTerm")
```

## Format

A data frame with 32307 observations on the following 2 variables.

gene gene id, a factor with 32307 levels

1fc log fold-change, a numeric vector

makeDeltaRanksTable Make a combined delta-ranks table from several kog.mwu() results.

#### Description

Extracts delta ranks from several kog.mwu() result tables and combines them into a single dataframe for heat map plotting and correlation analysis.

#### Usage

```
makeDeltaRanksTable(11)
```

#### Arguments

11

A list of dataframes output by kog.mwu() function.

## Value

A dataframe of delta-ranks (rows - KOG classes, columns - delta-ranks in different datasets).

## Author(s)

Mikhail V. Matz

## References

Dixon GB, Davies SW, Aglyamova GA, Meyer E, Bay LK and Matz MV (2015) Genomic determinants of coral heat tolerance across latitudes.

panel.cor

#### Description

works as upper.panel or lower.panel argument of pairs() (package graphics).

## Usage

```
panel.cor(x, y, digits=2, cex.cor)
```

## Arguments

х	x element of the pairs() matrix
У	y element of the pairs() matrix
digits	number of non-zero digits to leave at the end
cex.cor	scaling factor for displayed text

#### References

cannibalized from an example in ?pairs (package graphics)

panel.cor.pval	accessory function for pairs() to display pvalue of the Pearson corre-
	lation

## Description

works as upper.panel or lower.panel argument of pairs() (package graphics). Displays pvalues better than 0.1.

## Usage

```
panel.cor.pval(x, y, digits = 2, cex.cor, p.cut=0.1)
```

## Arguments

х	x element of the pairs() matrix
У	y element of the pairs() matrix
digits	number of non-zero digits to leave at the end
cex.cor	scaling factor for displayed text
p.cut	p-value cutoff

## References

cannibalized from an example in ?pairs (package graphics)

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