# Package 'meaca'

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Type Package
Title Mixed-effects Enrichment Analysis with Correlation Adjusted (MEACA)
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<b>Description</b> This package documents the functions used when performing MEACA gene set enrichment analysis.
<b>Depends</b> R (>= 3.2.1)
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imports MASS, mvtnorm, tibble
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R topics documented:
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btw\_gene\_corr

Calculate sample (Pearson) correlations among gene clusters

#### **Description**

Calculate sample (Pearson) correlations among gene clusters

#### Usage

```
btw_gene_corr(expression_data, trt, go_term, standardize = T)
```

#### **Arguments**

expression\_data

the expressoin matrix.

trt treatment indicators, 1 for treatment, 0 for control group go\_term an indicator vector. 1 for genes in the test set, 0 otherwise

standardize whether the data should be standaridzed

#### Value

a \$1 \times 3\$ data frame containing values for \$\rho\_1\$, \$\rho\_3\$ and \$\rho\_2\$ respectively.

testSetCor Average correlation for genes in the test set

interCor Average correlation between genes in the test set and those not in the test set

backSetCor Average correlations for genes not in the test set.

estimate\_sigma

Estimate sample covariance.

#### **Description**

Estimate sample covariance and calculate the gene-level statistics

#### Usage

```
estimate_sigma(expression_data, trt)
```

#### Arguments

expression\_data

the expression matrix.

trt sample labels. 0 for control and 1 for treatment

#### Value

a list

sigma a covariance matrix

t\_val a vector of gene level test statistics

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#### **Description**

meaca for testing multiple gene sets.

#### Usage

```
meaca_multiple(expression_data, trt, geneset, standardize = T,
    min_set_size = 5, fdr_method = "BH")
```

#### **Arguments**

expression\_data

the expressoin matrix.

trt treatment labels.

geneset gene sets to be tested, an object from read\_gene\_set.

standardize whether the data should be standaridzed.

min\_set\_size the minimum number of genes contained for a gene set to be considered.

fdr\_method which method is ued to adjust the p values. see arguments in function p.adjust.

#### Value

a data frame

meaca_single	meaca-single.	

#### Description

meaca for single gene set test.

#### Usage

```
meaca_single(expression_data, trt, go_term, standardize = F)
```

#### **Arguments**

expression\_data

the expressoin matrix.

trt treatment indicators, 1 for treatment, 0 for control group.
go\_term an indicator vector. 1 for genes in the test set, 0 otherwise.

standardize whether the data should be standaridzed.

#### Value

a list

stat the test statistic

p1 chi-square test p value

status "up" or "down", the direction of differential expression

p2 two-sided test p-value using normal distribution

#### **Examples**

simulate\_expression\_data

Simulate expression data.

#### **Description**

simulate normally distributed expression data with desired DE probabilities for genes in the test set and for those not in the test set..

#### Usage

```
simulate_expression_data(size, n_gene, n_test, prop, de_mu, de_sd, rho1, rho2,
  rho3, data_gen_method = "chol", seed = 123)
```

#### **Arguments**

size number of samples to be simulatedn\_gene total number of genes to be simulatedn\_test number of genes in the test set.

prop a vector of length 2, proportion of DE genes within go term and outside go\_term,

corresponding to \$p\_t\$ and \$p\_b\$.

de\_mu, de\_sd if the gene is DE, delta  $\sim N(de_mu, de_sd)$ 

rho1 a scalar, correlation between two test genes (i.e.,  $\rho_1$  in the paper)

rho2 a scalar, correlation between two background genes (i.e.,  $\rho_2$  in the paper) rho3 correlation between a test gene and a background gene (i.e.,  $\rho_3$  in the paper)

 ${\tt data\_gen\_method}$ 

data generation method; if 'data\_gen\_method = MASS', then mvrnorm is used,

otherwise see function rmvnorm

seed the seed used for simulation (for reproducibility purpose)

#### Value

a list

data a expression matrix of  $m \times n$  where m is the number of genes and n is the

number of samples.

trt sample labels of length n, 1 for treatment and 0 for control.
go\_term gene labels of length m, 1 for go\_term genes and 0 otherwise.

sigma true covariance matrix upon which data is simulated.

#### **Examples**

standardize\_expression\_data

standardize expression data, with method described in the paper.

#### Description

Standardize the expression data.

#### Usage

```
standardize_expression_data(expression_data, trt)
```

#### **Arguments**

expression\_data

the expression matrix.

trt sample labels. 0 for control and 1 for treatment

#### Value

a matrix of the same dimension as input data.

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