## **PEPPER**

February 2, 2018

convert.probe.to.gene.expression

Convert probe level expression to gene level

## Description

Convert probe level expression to gene level

## Usage

```
convert.probe.to.gene.expression(expr, gene.mapping, selection.method = "iqr")
```

## **Arguments**

expr Expression matrix.

gene.mapping Probe-gene mapping.

selection.method

How to handle multiple probes corresponding to the same gene. Defaults to iqr (probe with highest IQR). Other options var (highest variance), med (median of the probes)

## Value

a data frame containin gene level expression data

## **Examples**

```
#expr.gene = convert.probe.to.gene.expression(expr, gene.mapping)
```

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#### **Description**

Convert sample mapping to case control from conditions

#### Usage

```
convert.sample.mapping.to.case.control(sample.mapping, states.case,
   states.control, out.file = NULL)
```

#### Arguments

```
sample.mapping Sample-condition mapping for the data set.

states.case Conditions to be used as case

states.control Conditions to be used as background

out.file File name to write the converted mapping
```

#### Value

a data frame containin sample mapping

## **Examples**

```
#sample.mapping = convert.sample.mapping.to.case.control(sample.mapping,
# states.control = c("healthy donor"),
# states.case = c("tuberculosis", "latent tuberculosis infection"))
```

fetch.expression.data Fetch expression data from GEO / or given folder

#### **Description**

Fetch expression data from GEO / or given folder

## Usage

```
fetch.expression.data(geo.id, sample.mapping.column = "characteristics_ch1",
  do.log2 = NULL, probe.conversion = NULL, conversion.mapping = NULL,
  conversion.mapping.function = NULL, output.dir = paste(geo.id, "/", sep =
  ""), geo.id.sub = NULL, reprocess = NULL)
```

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#### **Arguments**

geo.id GEO id.

do.log2 Apply log2 transformation to the expression values. Defaults to TRUE.

probe.conversion

Convert probe level expression to gene level using provided annotation label (uses platform specific annotations downloaded from GEO). Defaults to NULL (no conversion). In case of multiple probes, probe with absolute max value is

chosen.

conversion.mapping

Mapping of platform specific ids to user provided ids

conversion.mapping.function

Function to process the mapped name such that it matches with the ids provided

in conversion.map

output.dir Directory where all files will be written.

geo.id.sub GEO id for the sub-set (e.g., specific to the platform).

#### Value

A list containing 3 data frames: expression matrix, sample mapping, gene mapping.

#### **Examples**

```
gds.data = fetch.expression.data("GDS4966", do.log2=F, probe.conversion="Gene ID")
expr = gds.data$expr
sample.mapping = gds.data$sample.mapping
```

find.de.genes.limma

Find differentially expressed genes using LIMMA

## Description

Find differentially expressed genes using LIMMA

#### Usage

```
find.de.genes.limma(expr, sample.mapping, states, out.file = NULL,
   state.background = NULL, adjust.method = "BH", cutoff = 0.05)
```

## **Arguments**

expr expression matrix.

sample.mapping Sample - condition mapping.

states Conditions to be considered as case.

out.file File to write output. If NULL (default) not used.

state.background

Condition to be considered as control.

adjust.method Multiple hypothesis testing correction method. Defaults to BH.

cutoff Adjust p-value cutoff. Defaults to 0.05

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#### Value

Data frame with results.

find.de.genes

Find differentially expressed genes

#### **Description**

Find differentially expressed genes

#### Usage

```
find.de.genes(expr, sample.mapping, states, method = "limma",
  out.file = NULL, state.background = NULL, adjust.method = "BH",
  cutoff = 0.05, functional.enrichment = NULL)
```

#### **Arguments**

```
expression matrix.
expr
sample.mapping Sample - condition mapping.
states
                  Conditions to be considered as case.
method
                  Differential expression analysis method: limma (Default) | sam | welch.
out.file
                  File to write output. If NULL (default) not used.
state.background
                  Condition to be considered as control.
adjust.method
                 Multiple hypothesis testing correction method. Defaults to BH.
cutoff
                  Adjust p-value cutoff. Defaults to 0.05
functional.enrichment
```

#### Value

data frame with results

## Examples

GO or KEGG based functional enrichment analysis

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find.de.genes.sam

Find differentially expressed genes using SAM

#### **Description**

Find differentially expressed genes using SAM

#### Usage

```
find.de.genes.sam(expr, sample.mapping, states, out.file = NULL,
    state.background = NULL, adjust.method = "BH", cutoff = 0.05)
```

#### Arguments

expr expression matrix.

sample.mapping Sample - condition mapping.

states Conditions to be considered as case.

out.file File to write output. If NULL (default) not used.

state.background

Condition to be considered as control.

adjust.method Multiple hypothesis testing correction method. Defaults to BH.

cutoff Adjust p-value cutoff. Defaults to 0.05

#### Value

Data frame with results.

find.de.genes.welch

Find differentially expressed genes using Welch (t-test w/ unequal variance) test

#### **Description**

Find differentially expressed genes using Welch (t-test w/ unequal variance) test

### Usage

```
find.de.genes.welch(expr, sample.mapping, states, out.file = NULL,
   state.background = NULL, adjust.method = "BH", cutoff = 0.05)
```

### **Arguments**

expr expression matrix.

sample.mapping Sample - condition mapping.

states Conditions to be considered as case.

out.file File to write output. If NULL (default) not used.

state.background

Condition to be considered as control.

adjust.method Multiple hypothesis testing correction method. Defaults to BH.

cutoff Adjust p-value cutoff. Defaults to 0.05.

get.fdr.matrix

#### Value

Data frame with results

get.data.set

Get expression data set

## Description

Get expression data set

## Usage

```
get.data.set(geo.id, output.dir, is.annotation = F)
```

## Arguments

geo.id GEO id.

output.dir Output directory to write / look for files.

#### Value

data set

get.fdr.matrix

Get fdr matrix

## Description

Calculates FDRs from z-scores for each gene in each sample.

## Usage

```
get.fdr.matrix(z, adjust.method, out.file = NULL)
```

#### **Arguments**

z Data frame containing z-scores (probes vs samples).

adjust.method P-value correction method (see p.adjust).
out.file Output file for writing z score matrix.

### Value

Data frame containing FDR values.

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```
get.peeps.from.z.matrix
```

Get peeps from z-score matrix

#### **Description**

Returns peeps for each sample in a given z-score matrix.

#### Usage

```
get.peeps.from.z.matrix(z, cutoff, convert.to.pvalues)
```

## **Arguments**

z Matrix containing z-scores (genes vs samples), tab separated.

 $\hbox{cutoff} \qquad \hbox{Threshold for deciding peeps, either a $z$-score or adjusted $p$-value (if convert.to.pvalues=$T$).} \\ \hbox{convert.to.pvalues}$ 

Flag to convert z-scores to p-values. If TRUE, the z-scores are converted to P-values which are then corrected for multiple hypothesis testing.

#### Value

Data frame containing sample name and geneid of genes in the peeps

#### **Examples**

```
\verb|peeps <- get.peeps.from.z.matrix(z, cutoff=0.05, convert.to.pvalues=T)|\\
```

get.z.matrix

Get z matrix

## **Description**

Returns z-score matrix for a given GEO data set. The z-scores are calculated for each gene in each sample using the mean and sd over provided control samples.

#### Usage

```
get.z.matrix(expr, sample.mapping, states.control = NULL,
    states.case = NULL, method = "mean", out.file = NULL)
```

#### **Arguments**

expr Expression matrix (genes vs samples), tab separated.

sample.mapping Sample - condition mapping.

states.control Label of control (background) samples. If NULL sample.mapping is assumed

to include the following types: "case" "control".

states.case Label of case (disease) samples.

method Method to calculate the z score, defaults to mean and sd, use median for med

and mad.

out.file Output file for writing z score matrix.

get.z.score

#### Value

Data frame containing z-scores

#### **Examples**

```
gds.data = fetch.expression.data("GDS4966", do.log2=F, probe.conversion="Gene ID")
expr = gds.data$expr
sample.mapping = gds.data$sample.mapping
z = get.z.matrix(expr, sample.mapping,
states.control = c("healthy donor"),
states.case = c("tuberculosis", "latent tuberculosis infection"))
```

get.z.score

Get z score

#### **Description**

Calculates z-score for each gene in each sample using the mean and sd over provided control samples.

## Usage

```
get.z.score(expr, samples.background, method = "mean",
   samples.to.exclude = NULL)
```

#### Arguments

expr Expression matrix (probes vs samples). samples.background

Names of the background (control) samples.

method Method to calculate the z score, defaults to mean and sd, use median for med

and mad.

samples.to.exclude

Names of the samples to be excluded from background (for CV).

#### Value

Data frame containing z-scores

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