

# Package ‘TSEA’

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**Type** Package

**Title** Tissue-Specific Enrichment Analysis

**Version** 1.0

**Date** 2018-08-21

**Author** Guangsheng Pei

**Maintainer** Guangsheng Pei <peiguangsheng@gmail.com>

**Imports** pheatmap, RColorBrewer

**Description** Tissue-specific enrichment analysis to assess lists of candidate genes or RNA-Seq expression profiles.

**License** GPL (>= 2)

**NeedsCompilation** no

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## Description

Tissue-specific enrichment analysis to assess lists of candidate genes and tissue-specific expression  
decode analysis for RNA-seq data to decode RNA expression matrices tissue heterogeneity.

## Details

Since disease and physiological condition are often associated with a specific tissue, understanding the tissue-specific genes (TSG) expression patterns will substantially reduce false discoveries in biomedical research. However, due to cell complexity in human system, heterogeneous tissues are frequently collected. Making it difficult to distinguish gene expression variability and mislead result interpretation. Here, we present TSEA, an R package that conducts Tissue-Specific Enrichment Analysis (TSEA) using two built-in reference panels: the Genotype-Tissue Expression (GTEx) data and the ENCyclopedia Of DNA Elements (ENCODE) data. We implemented two major functions in TSEA to assess lists of candidate genes or expression matrices.

The DESCRIPTION file:

```
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Type:         Package
Title:        Tissue-Specific Enrichment Analysis
Version:      1.0
Date:         2018-08-21
Author:       Guangsheng Pei
Maintainer:   Guangsheng Pei <peiguangsheng@gmail.com>
Imports:      pheatmap, RColorBrewer
Description:  Tissue-specific enrichment analysis to assess lists of candidate genes or RNA-Seq expression profiles.
License:      GPL (>= 2)
```

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tsea.plot	Tissue-specific enrichment analysis result heatmap plot
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## Author(s)

Guangsheng Pei  
 Maintainer: Guangsheng Pei

## References

Pei G., Dai Y., Zhao Z. Jia P. (2018) Tissue-Specific Enrichment Analysis (TSEA) to decode tissue heterogeneity. Bioinformatics, In submission.

**See Also**

<https://github.com/bsml320/TSEA/blob/master/README.md>

**Examples**

```
#Download the built-in data from https://github.com/bsml320/TSEA/tree/master/data
#load("data/GTEX_t_score.rda")
#load("data/ENCODE_z_score.rda")
#library(pheatmap)

#=====
#Example 1: Lists of candidate genes for single sample.
#load("data/GWAS_gene.rda")
#query_gene_list = GWAS_gene

#TSEA in GTEX panel
#tsea_t = tsea.analysis(query_gene_list, GTEX_t_score, ratio = 0.05,
#p.adjust.method = "bonferroni")
#write.csv(tsea_t,"1.1.GWAS_TSEA_in_GTEX_panel.csv")
#pdf ("1.2.GWAS_TSEA_in_GTEX_panel.pdf", 5, 9, onefile = FALSE)
#tsea.plot(tsea_t, threshold = 0.05)
#dev.off()
#tsea_t_summary = tsea.summary(tsea_t)
#write.csv(tsea_t_summary,"1.3.GWAS_summary_in_GTEX_panel.csv")

#TSEA in ENCODE panel
#tsea_z = tsea.analysis(query_gene_list, ENCODE_z_score, ratio = 0.05,
#p.adjust.method = "bonferroni")
#write.csv(tsea_z,"1.4.GWAS_TSEA_in_ENCODE_panel.csv")
#pdf ("1.5.GWAS_TSEA_in_ENCODE_panel.pdf", 5, 9, onefile = FALSE)
#tsea.plot(tsea_z, threshold = 0.05)
#dev.off()
#tsea_z_summary = tsea.summary(tsea_z)
#write.csv(tsea_z_summary,"1.6.GWAS_summary_in_ENCODE_panel.csv")

#=====
#Example 2: Lists of candidate genes for multiple samples.
#load("data/GWAS_gene_multiple.rda")
#query_gene_list = GWAS_gene_multiple

#TSEA in GTEX panel
#tsea_t_multi = tsea.analysis.multiple(query_gene_list,
#GTEX_t_score, ratio = 0.05, p.adjust.method = "BH")
#write.csv(tsea_t_multi,"2.1.GWAS_multi_TSEA_in_GTEX_panel.csv")
#pdf ("2.2.GWAS_multi_TSEA_in_GTEX_panel.pdf", 6, 6, onefile = FALSE)
#tsea.plot(tsea_t_multi, threshold = 0.05)
#dev.off()
#tsea_t_multi_summary = tsea.summary(tsea_t_multi)
#write.csv(tsea_t_multi_summary,"2.3.GWAS_multi_summary_in_GTEX_panel.csv")

#TSEA in ENCODE panel
#tsea_z_multi = tsea.analysis.multiple(query_gene_list,
#ENCODE_z_score, ratio = 0.05, p.adjust.method = "BH")
#write.csv(tsea_z_multi,"2.4.GWAS_multi_TSEA_in_ENCODE_panel.csv")
#pdf ("2.5.GWAS_multi_TSEA_in_ENCODE_panel.pdf", 7, 7, onefile = FALSE)
#tsea.plot(tsea_z_multi, threshold = 0.05)
```

```

#dev.off()
#tsea_z_multi_summary = tsea.summary(tsea_z_multi)
#write.csv(tsea_z_multi_summary, "2.6.GWAS_multi_summary_in_ENCODE_panel.csv")

#####
#Example 3: RNA expression profiles TSEA in ENCODE panel.
#load("data/query_GTEEx.rda")
#query_matrix = query_GTEEx

#load("data/correction_factor.rda")
#RNA expression profiles z-score normalization
#query_mat_zscore_nor = tsea.expression.normalization(query_matrix,
#correction_factor, normalization = "z-score")

#RNA expression profiles TSEA in ENCODE panel
#tseaed_in_ENCODE = tsea.expression.decode(query_mat_zscore_nor,
#ENCODE_z_score, ratio = 0.05, p.adjust.method = "BH")
#write.csv(tseaed_in_ENCODE, "3.1.RNAseq_TSEA_in_ENCODE_panel.csv")
#pdf ("3.2.RNAseq_TSEA_in_ENCODE_panel.pdf", 10, 9, onefile = FALSE)
#tsea.plot(tseaed_in_ENCODE, threshold = 0.05)
#dev.off()
#tseaed_in_ENCODE_summary = tsea.summary(tseaed_in_ENCODE)
#write.csv(tseaed_in_ENCODE_summary, "3.3.RNAseq_summary_in_ENCODE_panel.csv")

#####
#Example 4: RNA expression profiles TSEA in GTEEx panel.
#load("data/query_ENCODE.rda")
#query_matrix = query_ENCODE

#RNA expression profiles abundance normalization
#query_mat_abundance_nor = tsea.expression.normalization(query_matrix,
#correction_factor, normalization = "abundance")

#RNA expression profiles TSEA in GTEEx panel
#tseaed_in_GTEEx = tsea.expression.decode(query_mat_abundance_nor,
#GTEEx_t_score, ratio = 0.05, p.adjust.method = "BH")
#write.csv(tseaed_in_GTEEx, "4.1.RNAseq_TSEA_in_GTEEx_panel.csv")
#pdf ("4.2.RNAseq_TSEA_in_GTEEx_panel.pdf", 10, 9, onefile = FALSE)
#tsea.plot(tseaed_in_GTEEx, threshold = 0.05)
#dev.off()
#tseaed_in_GTEEx_summary = tsea.summary(tseaed_in_GTEEx)
#write.csv(tseaed_in_GTEEx_summary, "4.3.RNAseq_summary_in_GTEEx_panel.csv")

```

---

tsea.analysis

*Tissue-specific enrichment analysis for query gene list*


---

## Description

Tissue-specific enrichment analysis by Fisher's Exact Test for given gene list.

## Usage

```

tsea.analysis(query_gene_list, score, ratio = 0.05,
p.adjust.method = "BH")

```

**Arguments**

<code>query_gene_list</code>	a gene symbol list object.
<code>score</code>	a gene tissue-specific score matrix, c("GTEx_t_score" or "ENCODE_z_score"), can be loaded by <code>data(GTEx)</code> or <code>data(ENCODE)</code> , the default value is recommended "GTEx_t_score".
<code>ratio</code>	the threshold to define tissue-specific genes (with top t-score or z-score), the default value is 0.05.
<code>p.adjust.method</code>	<code>p.adjust.method</code> , c("holm", "hochberg", "hommel", "bonferroni", "BH", "BY", "fdr", "none")

**Details**

Tissue-specific enrichment analysis by Fisher's Exact Test for given gene list.

**Value**

nothing

**Note**

nothing

**Author(s)**

Guangsheng Pei

**References**

Pei G., Dai Y., Zhao Z. Jia P. (2018) Tissue-Specific Enrichment Analysis (TSEA) to decode tissue heterogeneity. Bioinformatics, In submission.

**See Also**

<https://github.com/bsml320/TSEA/>

**Examples**

```
#load("data/GWAS_gene.rda")
#query_gene_list = GWAS_gene
#tsea_t = tsea.analysis(query_gene_list, GTEx_t_score, 0.05,
#p.adjust.method = "bonferroni")
```

---

```
tsea.analysis.multiple
```

*Tissue-specific enrichment analysis for multi query gene lists*

---

## Description

Tissue-specific enrichment analysis by Fisher's Exact Test for multiple gene list.

## Usage

```
tsea.analysis.multiple(query_gene_list, score, ratio = 0.05,
p.adjust.method = "BH")
```

## Arguments

<code>query_gene_list</code>	a 0~1 gene~sample table object, row should be gene symbol, column should be sample name. In the table, gene labeled with 1 indicated it is target gene for a given sample, while 0 indicated it is not target in a given sample.
<code>score</code>	a gene tissue-specific score matrix, c("GTEx_t_score" or "ENCODE_z_score"), can be loaded by data(GTEx) or data(ENCODE), the default value is recommended "GTEx_t_score".
<code>ratio</code>	the threshold to define tissue-specific genes (with top t-score or z-score), the default value is 0.05.
<code>p.adjust.method</code>	p.adjust.method, c("holm", "hochberg", "hommel", "bonferroni", "BH", "BY", "fdr", "none")

## Details

Tissue-specific enrichment analysis by Fisher's Exact Test for multiple gene list.

## Value

nothing

## Note

nothing

## Author(s)

Guangsheng Pei

## References

Pei G., Dai Y., Zhao Z. Jia P. (2018) Tissue-Specific Enrichment Analysis (TSEA) to decode tissue heterogeneity. Bioinformatics, In submission.

## See Also

<https://github.com/bsml320/TSEA/>

**Examples**

```
#load("data/GWAS_gene_multiple.rda")
#query_gene_list = GWAS_gene_multiple
#tsea_t_multi = tsea.analysis.multiple(query_gene_list,
#GTEx_t_score, 0.05, p.adjust.method = "BH")
```

---

```
tsea.expression.decode
```

*Tissue-specific enrichment analysis for RNA-Seq expression profiles*

---

**Description**

Tissue-specific enrichment analysis to decode whether a given RNA-seq sample (RPKM) with potential confounding effects based on expression profiles.

**Usage**

```
tsea.expression.decode(query_mat_normalized_score, score,
ratio = 0.05, p.adjust.method = "BH")
```

**Arguments**

<code>query_mat_normalized_score</code>	a normalized RNA-seq RPKM object, which produced by "tsea.expression.normalization".
<code>score</code>	a gene tissue-specific score matrix, c("GTEx_t_score" or "ENCODE_z_score"), can be loaded by data(GTEx) or data(ENCODE), the default value is recommended "GTEx_t_score".
<code>ratio</code>	the threshold to define tissue-specific genes (with top t-score or z-score), the default value is 0.05.
<code>p.adjust.method</code>	p.adjust.method, c("holm", "hochberg", "hommel", "bonferroni", "BH", "BY", "fdr", "none")

**Details**

Tissue-specific enrichment analysis for RNA-Seq expression profiles.

**Value**

nothing

**Note**

nothing

**Author(s)**

Guangsheng Pei

## References

Pei G., Dai Y., Zhao Z. Jia P. (2018) Tissue-Specific Enrichment Analysis (TSEA) to decode tissue heterogeneity. Bioinformatics, In submission.

## See Also

<https://github.com/bsml320/TSEA/>

## Examples

```
#load("data/query_GTEEx.rda")
#query_matrix = query_GTEEx
#load("data/correction_factor.rda")
#RNA expression profiles z-score normalization
#query_mat_zscore_nor = tsea.expression.normalization(query_matrix,
#correction_factor, normalization = "z-score")

#RNA expression profiles TSEA in ENCODE panel
#tseaed_in_ENCODE = tsea.expression.decode(query_mat_zscore_nor,
#ENCODE_z_score, 0.05, p.adjust.method = "BH")
```

---

```
tsea.expression.normalization
```

*RNA-Seq expression profiles normalization*

---

## Description

To avoid the data bias and adapt better data heterogeneity, before `tsea.expression.decode()` analysis, the raw discrete RPKM value have to normalized to continuous variable meet the normal distribution before t-test.

## Usage

```
tsea.expression.normalization(query_mat,
correction_factor, normalization = "abundance")
```

## Arguments

<code>query_mat</code>	a RNA-seq RPKM object, row name should be gene symbol, and column name should be sample name.
<code>correction_factor</code>	<code>correction_factor</code> , a gene table object contain genes average expression level and standard variance in GTEEx database, can be loaded by <code>data(correction_factor)</code> .
<code>normalization</code>	normalization methods, c("z-score", "abundance")

## Details

As RNA-Seq samples are often heterogeneous, before in-depth analysis, it is necessary to decode tissue heterogeneity to avoid samples with confounding effects. However, the raw discrete RPKM value have to normalized to continuous variable meet the normal distribution before t-test.



**Value**

nothing

**Note**

nothing

**Author(s)**

Guangsheng Pei

**References**

Pei G., Dai Y., Zhao Z. Jia P. (2018) Tissue-Specific Enrichment Analysis (TSEA) to decode tissue heterogeneity. Bioinformatics, In submission.

**See Also**

<https://github.com/bsml320/TSEA/>

**Examples**

```
#load("data/query_GTEEx.rda")
#query_matrix = query_GTEEx
#load("data/correction_factor.rda")
#RNA expression profiles z-score normalization
#query_mat_zscore_nor = tsea.expression.normalization(query_matrix,
#correction_factor, normalization = "z-score")

#RNA expression profiles TSEA in ENCODE panel
#tseaed_in_ENCODE = tsea.expression.decode(query_mat_zscore_nor,
#ENCODE_z_score, 0.05, p.adjust.method = "BH")
```

tsea.plot

*Tissue-specific enrichment analysis result heatmap plot***Description**

Heat map plot for tissue-specific enrichment analysis result.

**Usage**

```
tsea.plot(tsea_result, threshold = 0.05)
```

**Arguments**

tsea_result	the result of tissue-specific enrichment analysis, which produced by "tsea.analysis", "tsea.analysis.multiple" or "tsea.expression.decode".
threshold	the p-value threshold to define if the gene list or RNA-seq data enriched in a given tissue, p-value greater than threshold will not be labeled in the plot. The default value is 0.05.

**Details**

Heat map plot for tissue-specific enrichment analysis result

**Value**

nothing

**Note**

nothing

**Author(s)**

Guangsheng Pei

**References**

Pei G., Dai Y., Zhao Z. Jia P. (2018) Tissue-Specific Enrichment Analysis (TSEA) to decode tissue heterogeneity. Bioinformatics, In submission.

**See Also**

<https://github.com/bsml320/TSEA/>

**Examples**

```
#load("data/GWAS_gene_multiple.rda")
#query_gene_list = GWAS_gene_multiple

#TSEA in GTEx panel
#tsea_t_multi = tsea.analysis.multiple(query_gene_list,
#GTEx_t_score, 0.05, p.adjust.method = "BH")
#tsea.plot(tsea_t_multi, 0.05)
```

---

tsea.summary

*Tissue-specific enrichment analysis result summary*


---

**Description**

Tissue-specific enrichment analysis result summary (list the top 3 most enriched tissues) from the given gene list or RNA-seq expression profiles.

**Usage**

```
tsea.summary(tsea_result)
```

**Arguments**

tsea\_result the result of tissue-specific enrichment analysis, which produced by "tsea.analysis", "tsea.analysis.multiple" or "tsea.expression.decode".

**Details**

Tissue-specific enrichment analysis result summary

**Value**

nothing

**Note**

nothing

**Author(s)**

Guangsheng Pei

**References**

Pei G., Dai Y., Zhao Z. Jia P. (2018) Tissue-Specific Enrichment Analysis (TSEA) to decode tissue heterogeneity. Bioinformatics, In submission.

**See Also**

<https://github.com/bsml320/TSEA/>

**Examples**

```
#load("data/query_GTEEx.rda")
#query_matrix = query_GTEEx
#load("data/correction_factor.rda")
#RNA expression profiles z-score normalization
#query_mat_zscore_nor = tsea.expression.normalization(query_matrix,
#correction_factor, normalization = "z-score")

#RNA expression profiles TSEA in ENCODE panel
#tseaed_in_ENCODE = tsea.expression.decode(query_mat_zscore_nor,
#ENCODE_z_score, 0.05, p.adjust.method = "BH")
#tseaed_in_ENCODE_summary = tsea.summary(tseaed_in_ENCODE)
```

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