# Package: pathfindR.data (via r-universe)

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Title Data Package for patninds
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Maintainer Ege Ulgen <egeulgen@gmail.com></egeulgen@gmail.com>
<b>Description</b> This is a data-only package, containing data needed to run the CRAN package 'pathfindR', a package for enrichment analysis utilizing active subnetworks. This package contains protein-protein interaction network data, data related to gene sets and example input/output data.
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biocarta\_descriptions BioCarta Pathways - Descriptions

## **Description**

A named vector containing the descriptions for each human BioCarta pathway. *Generated on 27 Apr 2024*.

## Usage

biocarta\_descriptions

## **Format**

named vector containing 292 character values, the descriptions for the given pathways.

biocarta\_genes

BioCarta Pathways - Gene Sets

## **Description**

A list containing the genes involved in each human BioCarta pathway. Each element is a vector of gene symbols located in the given pathway. *Generated on 27 Apr 2024*.

## Usage

biocarta\_genes

#### **Format**

list containing 292 vectors of gene symbols. Each vector corresponds to a gene set.

cell\_markers\_descriptions

Human Cell Markers - Descriptions

#### **Description**

A named vector containing descriptions of different cell types from different tissues in human. Names of the vectors are Cell Ontology IDs (if available) of the cell types in the following format: "tissue type, cancer type, cell name" For more information, refer to the article: Hu C, Li T, Xu Y, Zhang X, Li F, Bai J, et al. CellMarker 2.0: an updated database of manually curated cell markers in human/mouse and web tools based on scRNA-seq data. Nucleic Acids Res. 2022 Oct 27;gkac947. *Generated on 27 Apr 2024*.

#### Usage

cell\_markers\_descriptions

#### **Format**

named vector containing 1986 character values, the descriptions for the given human cell types.

cell\_markers\_gsets

Human Cell Markers - Gene Sets

## **Description**

A list containing the sets of genes that are cell markers of different cell types from different tissues in human. Each element is a vector of cell marker gene symbols for the given cell type. Names correspond to the Cell Ontology ID (if available) of the cell type. For more information, refer to the article: Hu C, Li T, Xu Y, Zhang X, Li F, Bai J, et al. CellMarker 2.0: an updated database of manually curated cell markers in human/mouse and web tools based on scRNA-seq data. Nucleic Acids Res. 2022 Oct 27;gkac947. *Generated on 27 Apr 2024*.

#### Usage

cell\_markers\_gsets

#### **Format**

list containing 1986 vectors. Each vector corresponds to a cell marker gene set for a given human cell type.

example\_active\_snws

Example Active Subnetworks

#### **Description**

A list of vectors containing genes for each active subnetwork that passed the filtering step. *Generated on 27 Apr 2024*.

#### Usage

example\_active\_snws

#### **Format**

list containing 150 vectors. Each vector is the set of genes for the given active subnetwork.

example\_comparison\_output

Second Example Output for the pathfindR Enrichment Workflow (H.sapiens. - Rheumatoid Arthritis data)

## **Description**

The data frame containing the results of pathfindR's active-subnetwork-oriented enrichment work-flow performed on the rheumatoid arthritis dataset GSE84074 https://www.ncbi.nlm.nih.gov/geo/query/acc.cgi?acc=GSE84074. Analysis via run\_pathfindR was performed using the default settings. *Generated on 27 Apr 2024*.

#### Usage

example\_comparison\_output

#### **Format**

A data frame with 38 rows and 9 columns:

**ID** ID of the enriched term

Term\_Description Description of the enriched term

Fold\_Enrichment Fold enrichment value for the enriched term

**occurrence** the number of iterations that the given term was found to enriched over all iterations **support** the median support (proportion of active subnetworks leading to enrichment within an iteration) over all iterations

**lowest\_p** the lowest adjusted-p value of the given term over all iterations

highest\_p the highest adjusted-p value of the given term over all iterations

**Up\_regulated** the up-regulated genes in the input involved in the given term, comma-separated **Down\_regulated** the down-regulated genes in the input involved in the given term, comma-separated

#### See Also

example\_pathfindR\_input for the RA differentially-expressed genes data frame example\_pathfindR\_output for the RA example pathfindR enrichment output example\_pathfindR\_output\_clustered for the RA example pathfindR clustering output example\_experiment\_matrix for the RA differentially-expressed genes expression matrix run\_pathfindR for details on the pathfindR enrichment analysis

example\_custom\_genesets\_result

Custom Gene Set Enrichment Results

#### **Description**

A data frame consisting of pathfindR enrichment analysis results on the example TF target genes data (target gene sets of CREB and MYC). *Generated on 27 Apr 2024*.

## Usage

```
example_custom_genesets_result
```

#### **Format**

data frame containing 2 rows and 9 columns. Each row is a gene set (the TF target gene sets).

example\_experiment\_matrix

Example Experiment Matrix for pathfindR - Enriched Term Scoring

#### **Description**

A matrix containing the  $log_2$ -transformed and quantile-normalized expression values of the differentially-expressed genes for 18 rheumatoid arthritis (RA) patients and 15 healthy subjects. The matrix contains expression values of 572 significantly differentially-expressed genes (see example\_pathfindR\_input) with adj.P.Val <= 0.05. Generated on 28 Sep 2019.

#### Usage

```
example_experiment_matrix
```

#### **Format**

A matrix with 572 rows and 33 columns.

#### Source

https://www.ncbi.nlm.nih.gov/geo/query/acc.cgi?acc=GSE15573

#### See Also

example\_pathfindR\_input for the RA differentially-expressed genes data frame example\_pathfindR\_output for the RA example pathfindR enrichment output score\_terms for details on calculating agglomerated scores of enriched terms

example\_mmu\_input

Example Input for Mus musculus - Myeloma Analysis

## **Description**

A dataset containing the differentially-expressed genes and adjusted p-values for the GEO dataset GSE99393. The RNA microarray experiment was perform to detail the global program of gene expression underlying polarization of myeloma-associated macrophages by CSF1R antibody treatment. The samples were 6 murine bone marrow derived macrophages co-cultured with myeloma cells (myeloma-associated macrophages), 3 of which were treated with CSF1R antibody (treatment group) and the rest were treated with control IgG antibody (control group). In this dataset, differentially-expressed genes with llogFCl >= 2 and FDR < 0.05 are presented. *Generated on 1 Nov 2019*.

## Usage

example\_mmu\_input

## **Format**

A data frame with 45 rows and 2 variables:

Gene\_Symbol MGI gene symbols of the differentially-expressed genes

FDR adjusted p values, via the Benjamini & Hochberg (1995) method

#### Source

```
https://www.ncbi.nlm.nih.gov/geo/query/acc.cgi?acc=GSE99393
```

## See Also

example\_mmu\_output for the example mmu enrichment output. run\_pathfindR for details on the pathfindR enrichment analysis.

example\_mmu\_output

example\_mmu\_output

Example Output for Mus musculus - Myeloma Analysis

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#### **Description**

A dataset containing the results of pathfindR's active-subnetwork-oriented enrichment workflow performed on the Mus musculus myeloma differential expression dataset example\_mmu\_input. *Generated on 27 Apr 2024.* 

## Usage

example\_mmu\_output

#### **Format**

A data frame with 34 rows and 9 columns:

**ID** ID of the enriched term

**Term\_Description** Description of the enriched term

Fold Enrichment Fold enrichment value for the enriched term

occurrence the number of iterations that the given term was found to enriched over all iterations support the median support (proportion of active subnetworks leading to enrichment within an iteration) over all iterations

**lowest\_p** the lowest adjusted-p value of the given term over all iterations

**highest\_p** the highest adjusted-p value of the given term over all iterations

**Up\_regulated** the up-regulated genes in the input involved in the given term, comma-separated **Down\_regulated** the down-regulated genes in the input involved in the given term, comma-separated

## See Also

example\_mmu\_input for the example mmu input. run\_pathfindR for details on the pathfindR enrichment workflow.

example\_pathfindR\_input

Example Input for the pathfindR Enrichment Workflow - Rheumatoid Arthritis (H.sapiens)

## Description

A dataset containing the differentially-expressed genes along with the associated  $log_2$ (fold-change) values and FDR adjusted p-values for the GEO dataset GSE15573. This microarray dataset aimed to characterize gene expression profiles in the peripheral blood mononuclear cells of 18 rheumatoid arthritis (RA) patients versus 15 healthy subjects. Differentially-expressed genes with adj.P.Val < 0.05 are presented in this data frame. *Generated on 1 Nov 2019*.

#### Usage

```
example_pathfindR_input
```

#### **Format**

A data frame with 572 rows and 3 variables:

Gene.symbol HGNC gene symbols of the differentially-expressed genes

logFC log<sub>2</sub>(fold-change) values

adj.P.Val adjusted p values, via the Benjamini & Hochberg (1995) method

#### Source

```
https://www.ncbi.nlm.nih.gov/geo/query/acc.cgi?acc=GSE15573
```

#### See Also

example\_pathfindR\_output for the RA example pathfindR enrichment output example\_pathfindR\_output\_clustered for the RA example pathfindR clustering output example\_experiment\_matrix for the RA differentially-expressed genes expression matrix run\_pathfindR for details on the pathfindR enrichment analysis

example\_pathfindR\_output

 $\label{lem:example output for the pathfind Renrichment Workflow - Rheumatoid \\ Arthritis$ 

## **Description**

The data frame containing the results of pathfindR's active-subnetwork-oriented enrichment work-flow performed on the rheumatoid arthritis differential-expression data frame example\_pathfindR\_input. Analysis via run\_pathfindR was performed using the default settings. *Generated on 27 Apr 2024*.

#### Usage

```
example_pathfindR_output
```

## **Format**

A data frame with 121 rows and 9 columns:

**ID** ID of the enriched term

Term\_Description Description of the enriched term

Fold\_Enrichment Fold enrichment value for the enriched term

occurrence the number of iterations that the given term was found to enriched over all iterationssupport the median support (proportion of active subnetworks leading to enrichment within an iteration) over all iterations

**lowest\_p** the lowest adjusted-p value of the given term over all iterations

highest\_p the highest adjusted-p value of the given term over all iterations

Up\_regulated the up-regulated genes in the input involved in the given term, comma-separated

**Down\_regulated** the down-regulated genes in the input involved in the given term, comma-separated

#### See Also

example\_pathfindR\_input for the RA differentially-expressed genes data frame example\_pathfindR\_output\_clustered for the RA example pathfindR clustering outputs example\_experiment\_matrix for the RA differentially-expressed genes expression matrix run\_pathfindR for details on the pathfindR enrichment analysis

example\_pathfindR\_output\_clustered

Example Output for the pathfindR Clustering Workflow - Rheumatoid Arthritis

#### **Description**

A dataset containing the results of pathfindR's clustering and partitioning workflow performed on the rheumatoid arthritis enrichment results example\_pathfindR\_output. The clustering and partitioning function cluster\_enriched\_terms was used with the default settings (i.e. hierarchical clustering was performed and the agglomeration method was "average"). *Generated on 27 Apr* 2024.

#### Usage

example\_pathfindR\_output\_clustered

#### Format

A data frame with 121 rows and 11 columns:

**ID** ID of the enriched term

**Term\_Description** Description of the enriched term

Fold\_Enrichment Fold enrichment value for the enriched term

occurrence the number of iterations that the given term was found to enriched over all iterations

**support** the median support (proportion of active subnetworks leading to enrichment within an iteration) over all iterationss

**lowest\_p** the lowest adjusted-p value of the given term over all iterations

highest\_p the highest adjusted-p value of the given term over all iterations

Up\_regulated the up-regulated genes in the input involved in the given term, comma-separated

Down\_regulated the down-regulated genes in the input involved in the given term, comma-separated

**Cluster** the cluster to which the enriched term is assigned

Status whether the enriched term is the "Representative" term in its cluster or only a "Member"

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## See Also

example\_pathfindR\_input for the RA differentially-expressed genes data frame example\_experiment\_matrix for the RA differentially-expressed genes expression matrix run\_pathfindR for details on the pathfindR enrichment analysis example\_pathfindR\_output for the RA example pathfindR enrichment output cluster\_enriched\_terms for details on clustering methods

go\_all\_genes

Gene Ontology - All Gene Ontology Gene Sets

#### **Description**

A list containing the genes involved in each GO ontology term. Each element is a vector of gene symbols located in the given gene set. *Generated on 27 Apr 2024*.

#### **Usage**

go\_all\_genes

#### **Format**

list containing 15450 vectors of gene symbols. Each vector corresponds to a GO gene set.

kegg\_descriptions

KEGG Pathways - Descriptions

## **Description**

A named vector containing the descriptions for each Homo sapiens KEGG pathway. Names of the vector correspond to the KEGG ID of the pathway. Pathways that did not contain any genes were discarded. *Generated on 27 Apr 2024*.

#### Usage

kegg\_descriptions

#### **Format**

named vector containing 358 character values, the descriptions for the given pathways.

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kegg\_genes

KEGG Pathways - Gene Sets

#### **Description**

A list containing the genes involved in each Homo sapiens KEGG pathway. Each element is a vector of gene symbols located in the given pathway. Names correspond to the KEGG ID of the pathway. Pathways that did not contain any genes were discarded. *Generated on 27 Apr 2024*.

#### **Usage**

kegg\_genes

#### **Format**

list containing 358 vectors of gene symbols. Each vector corresponds to a pathway.

## **Description**

A named vector containing the descriptions for each Mus musculus KEGG pathway. Names of the vector correspond to the KEGG ID of the pathway. Pathways that did not contain any genes were discarded. *Generated on 27 Apr 2024*.

## Usage

mmu\_kegg\_descriptions

#### **Format**

named vector containing 355 character values, the descriptions for the given pathways.

mmu\_kegg\_genes

Mus Musculus KEGG Pathways - Gene Sets

## **Description**

A list containing the genes involved in each Mus musculus KEGG pathway. Each element is a vector of gene symbols located in the given pathway. Names correspond to the KEGG ID of the pathway. Pathways that did not contain any genes were discarded. *Generated on 27 Apr 2024*.

## Usage

mmu\_kegg\_genes

#### **Format**

list containing 355 vectors of gene symbols. Each vector corresponds to a pathway.

pathfindR.data\_updates

Table of Data for pathfindR

## Description

Data frame containing all the data for pathfindR along with descriptions and last update dates.

#### Usage

pathfindR.data\_updates

## **Format**

A data frame with 30 rows and 6 columns:

Category Category of the data

Name Name of the data

**Description** Description of the data

Source of the data

Version Version of the data (if applicable)

Last Update Last update date

reactome\_descriptions 13

reactome\_descriptions Reactome Pathways - Descriptions

## **Description**

A named vector containing the descriptions for each human Reactome pathway. Names of the vector correspond to the Reactome ID of the pathway. *Generated on 27 Apr 2024*.

## Usage

reactome\_descriptions

#### **Format**

named vector containing 2681 character values, the descriptions for the given pathways.

reactome\_genes

Reactome Pathways - Gene Sets

## Description

A list containing the genes involved in each human Reactome pathway. Each element is a vector of gene symbols located in the given pathway. Names correspond to the Reactome ID of the pathway. *Generated on 27 Apr 2024*.

## Usage

reactome\_genes

#### **Format**

list containing 2681 vectors of gene symbols. Each vector corresponds to a pathway.

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