

Package ‘DFD’

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Type Package

Title Extract Drugs from Differential Expression Data from LINCS Database

Version 0.1.0

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Description Get Drug information from given differential expression profile. The package search for the bioactive compounds from reference databases such as LINCS containing the genome-wide gene expression signature (GES) from tens of thousands of drug and genetic perturbations (Subramanian et al. (2017) <[DOI:10.1016/j.cell.2017.10.049](https://doi.org/10.1016/j.cell.2017.10.049)>).

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Encoding UTF-8

biocViews

Imports stringr, gprofiler2, signatureSearch, signatureSearchData

URL <https://github.com/MohmedSoudy/DFD>

BugReports <https://github.com/MohmedSoudy/DFD/issues>

RoxygenNote 7.2.3

NeedsCompilation no

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Repository CRAN

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convert_id	<i>Convert Gene Symbols to ENTREZ IDs</i>
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Description

The function is used to convert gene symbols to entrez ids and map the genes to human orthologs

Usage

```
convert_id(gene_symbols)
```

Arguments

gene_symbols gene symbols

Value

IDs that are converted from gene symbols to ENTREZ gene ids

Author(s)

Mohamed Soudy <Mohmedsoudy2009@gmail.com>

Examples

```
convert_id(c("TP53", "A2M"))
```

filter_drugs	<i>Re-rank drugs based on the number of targets</i>
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Description

The function is used to re-rank drugs based on their targets

Usage

```
filter_drugs(drug_frame)
```

Arguments

drug_frame drugs data frame returned by ‘get_drugs’ function

Value

re-ranked drug data frame based on their number of targets

Author(s)

Mohamed Soudy <Mohmedsoudy2009@gmail.com>

`get_drugs`

Get Drugs associated with the differential expression profile

Description

The function is used to get list of drugs that are associated with differential expression profile

Usage

```
get_drugs(up_regulated, down_regulated)
```

Arguments

`up_regulated` up-regulated genes returned by ‘prepare_ids’ function

`down_regulated` down-regulated genes returned by ‘prepare_ids’ function

Value

significant drugs data frame that contains list of drugs with their targets

Author(s)

Mohamed Soudy <Mohmedsoudy2009@gmail.com>

`prepare_ids`

Prepare IDs for CMAP Search

Description

The function is used to prepare the ids for the CMAP search

Usage

```
prepare_ids(up_regulated, down_regulated)
```

Arguments

`up_regulated` up regulated gene symbols

`down_regulated` down regulated gene symbols

Value

list containing up_regulated and down_regulated genes ENTREZ gene ids

Author(s)

Mohamed Soudy <Mohmedsoudy2009@gmail.com>

read_id

Read Gene Symbols from CSV file into up and down regulated lists

Description

The function is used to read symbols from a CSV file. The gene symbols should be in the first column

Usage

```
read_id(csv_path)
```

Arguments

csv_path	absolute path of CSV file containing gene symbols and sign
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Value

list containing up_regulated and down_regulated genes symbols

Author(s)

Mohamed Soudy <Mohmedsoudy2009@gmail.com>

run_pipeline

Run the main pipeline for getting drugs from differentail expression profile

Description

The function is used to run the main pipeline by extracting the drug list given differential expressed genes

Usage

```
run_pipeline(degs_path, output_path = NULL)
```

Arguments

degs_path	path to csv file containing degs see example file at https://raw.githubusercontent.com/MohmedSoudy/datas
	expression.csv
output_path	absolute path to output directory

run_pipeline

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Value

significant drug data frame after the re-ranking step

Author(s)

Mohamed Soudy <Mohmedsoudy2009@gmail.com>

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