Package 'DFD'

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Title Extract Drugs from Differential Expression Data from LINCS

Type Package

Database
Version 0.3.0
Maintainer Mohamed Soudy < Mohmed Soudy 2009@gmail.com>
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>).
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BugReports https://github.com/MohmedSoudy/DFD/issues RoxygenNote 7.2.3 NeedsCompilation no Author Mohamed Soudy [aut, cre] Repository CRAN Date/Publication 2025-02-11 13:50:11 UTC Contents
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convert_id

Convert Gene Symbols to ENTREZ IDs

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 < Mohmedsoudy 2009@gmail.com>

Examples

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

filter_drugs

Re-rank drugs based on the number of targets

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

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Author(s)

Mohamed Soudy < Mohmedsoudy 2009@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 < Mohmedsoudy 2009@gmail.com>

get_pert_by_type

Filter the drug based on the tested cell-line

Description

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

Usage

```
get_pert_by_type(perts, perts_type = c("cancer", "Normal"), high_targets = TRUE)
```

Arguments

perts drugs data frame returned by 'get_drugs' function

perts_type type of drug, whether it's from a normal or cancerous cell line high_targets Boolean paramter to rank their drugs per the number of targets plot_activities

Value

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

Author(s)

Mohamed Soudy < Mohmedsoudy 2009@gmail.com>

parse_perts

Parse the perturbagens to find out their frequency of occurrence.

Description

The function is used to parse drugs based on their targets

Usage

```
parse_perts(perts)
```

Arguments

perts

drugs data frame returned by 'get_drugs' function

Value

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

Author(s)

Mohamed Soudy < Mohmedsoudy 2009@gmail.com>

plot_activities

Visualise the number of perturbagens and the top activities

Description

The function is used to provide a handy visualization of the top activities

Usage

```
plot_activities(perts, top = 10, directorypath = NULL)
```

Arguments

perts drugs data frame returned by 'get_drugs' function or 'get_pert_by_type' func-

tion

top Number of activities to be visualised (N) The default value is set to 10.

directorypath path to save the output figure

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Value

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

Author(s)

Mohamed Soudy < Mohmedsoudy 2009@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 < Mohmedsoudy 2009@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

run_pipeline

Value

list containing up_regulated and down_regulated genes symbols

Author(s)

Mohamed Soudy < Mohmedsoudy 2009@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

Value

significant drug data frame after the re-ranking step

Author(s)

Mohamed Soudy < Mohmedsoudy 2009@gmail.com>

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