Package 'Autoseed'

January 20, 2025

Description For researchers to quickly and comprehensively acquire disease genes, so as to under-

The data is integrated from three public databases. The three databases are 'eDGAR', 'DrugBank'

Title Retrieve Disease-Related Genes from Public Sources

we developed this program to acquire disease-related genes.

stand the mechanism of disease,

Type Package

Version 0.1.0

and 'MalaCards'. The 'eDGAR' is a comprehensive database, containing data on the relationship between disease and genes. 'DrugBank' contains information on 13443 drugs and 5157 targets. 'MalaCards' integrates human disease information, including disease-related genes.
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AutoSeed

Get disease-related genes from eDGAR, DrugBank and MalaCards

Description

Get disease-related genes from eDGAR, DrugBank and MalaCards

Usage

```
AutoSeed(search)
```

Arguments

search

Name of the disease

Value

result\$edgar: Containing Disease Name, OMIM ID and Genesymbol (Data comes from the eDGAR) result\$malacards: Containing genes related to the disease (Data comes from the MalaCards) result\$drugbank: Containing genes related to the disease (Data comes from the DrugBank)

Examples

```
result = AutoSeed("diabetes")
```

drugbank

Relationship between gene and disease

Description

The data was collected in the drugbank. We got 27728 relationships between various disease and their related genes.

Usage

```
data("drugbank")
```

Format

A data frame with 27728 observations on the following 2 variables.

Details

There are two columns in the data. The first column is genes and the second column is diseases.

drugbank_disease_gene

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Source

https://www.drugbank.ca/

Examples

```
data(drugbank)
```

drugbank_disease_gene Get disease-related genes from DrugBank.

Description

Get disease-related genes from DrugBank.

Usage

```
drugbank_disease_gene(search)
```

Arguments

search

Name of the disease, character

Value

The genes related to the disease, list

Examples

```
result = drugbank_disease_gene("diabetes")
```

edgar

Relationship between the gene and disease in edgar

Description

The function "edgar_disease_gene()" will generate the relationship between gene and disease depend on this dataset.

Usage

```
data("edgar")
```

Format

A data frame with 1038340 observations on the following 2 variables.

4 edgar_disease_gene

Details

We got a total of 1038340 diseases and genes. There are two columns in the data. The first column is genes and the second column is diseases.

Source

http://edgar.biocomp.unibo.it/gene_disease_db/

Examples

data(edgar)

edgar_disease_gene

Get disease-related genes from eDGAR

Description

Get disease-related genes from eDGAR

Usage

```
edgar_disease_gene(disease)
```

Arguments

disease

Name of the disease, character

Value

a vector containing genesymbol related to the disease

Examples

```
result = edgar_disease_gene("diabetes")
```

mala 5

mala

The relationship between gene and disease

Description

The data was collected in the Malacards. We got 241306 relationships between various disease and their related genes.

Usage

```
data("mala")
```

Format

A data frame with 241306 observations on the following 2 variables.

Details

We got a total of 241306 diseases and genes. There are two columns in the data. The first column is disease and the second column is gene.

Source

```
<a href="https://doi.org/10.1093/nar/gkw1012">https://doi.org/10.1093/nar/gkw1012</a>
```

Examples

```
data(mala)
```

```
malacards_disease_gene
```

Get disease-related genes from MalaCards

Description

Get disease-related genes from MalaCards

Usage

```
malacards_disease_gene(disease)
```

Arguments

disease

Name of the disease

Value

The genes related to the disease, character vector

Examples

```
result = malacards_disease_gene("diabetes")
```

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