# Package 'MeSHDbi'

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Title DBI to construct MeSH-related package from sqlite file Description The package is unified implementation of MeSH.db, MeSH.AOR.db, and MeSH.PCR.db and also is interface to construct Gene-MeSH package (MeSH.XXX.eg.db). loadMeSHDbiPkg import sqlite file and generate MeSH.XXX.eg.db. Version 1.24.0 Author Koki Tsuyuzaki Maintainer Koki Tsuyuzaki <k.t.the-answer@hotmail.co.jp> **Depends** R (>= 3.0.1), BiocGenerics (>= 0.15.10) Imports methods, AnnotationDbi (>= 1.31.19), RSQLite, Biobase Suggests RUnit License Artistic-2.0 biocViews Annotation, AnnotationData, Infrastructure git\_url https://git.bioconductor.org/packages/MeSHDbi git\_branch RELEASE\_3\_11 git\_last\_commit aa52d99 git\_last\_commit\_date 2020-04-27 Date/Publication 2020-10-16

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listDatabases

# Description

This function returns the scientific name of species used in the package.

#### Usage

listDatabases(x)

#### Arguments

х

MeSHDb object such as MeSH.Mmu.eg.db

#### Author(s)

Koki Tsuyuzaki

# Examples

# library("MeSH.Mmu.eg.db")

# listDatabases(MeSH.Mmu.eg.db)

| makeGeneMeSHPackage | Making MeSHDb packages from corresponding table as single data |
|---------------------|--|
|                     | frame.   |

# Description

makeGeneMeSHPackage is a method that generates a package that will load an appropriate MeSHDb object that will in turn point to existing annotation packages.

# Usage

#### Arguments

| pkgname    | What is the desired package name.  |
|------------|--|
| data       | Data frame contains GENEID (e.g., 100036770), MESHID (e.g.D000465), CAT-EGORY (e.g., G), SOURCEID (pubmed id), and SOURCEDB (e.g., gendoo) |
| metadata   | Data frame contains metadata of the package  |
| organism   | The name of the organism this package represents   |
| version    | What is the version number for this package?   |
| maintainer | Who is the package maintainer? (must include email to be valid)  |
| author     | Who is the creator of this package?  |
| destDir    | A path where the package source should be assembled.   |
| license    | What is the license (and it's version)   |

#### Details

The purpose of this method is to create a special package that will depend on existing annotation packages and which will load a special MeSHDb object that will allow proper dispatch of special select methods. These methods will allow the user to easily query across multiple annotation resources via information contained by the MeSHDb object. Because the end result will be a package that treats all the data mapped together as a single source, the user is encouraged to take extra care to ensure that the different packages used are from the same build etc.

#### Value

A special package to load an MeSHDb object.

#### Author(s)

Koki Tsuyuzaki

#### See Also

MeSHDb

#### Examples

```
## makeGeneMeSHPackage enable users to construct
## user's own custom MeSH package
```

```
## this is test data which means the relationship between
## Entrez gene IDs of Pseudomonas aeruginosa PA01
## and its MeSH IDs.
data(PA01)
head(PA01)
```

```
# We are also needed to prepare meta data as follows.
data(metaPA01)
metaPA01
```

```
## sets up a temporary directory for this example
## (users won't need to do this step)
destination <- tempfile()
dir.create(destination)</pre>
```

MeSHDb-class MeSHDb objects

#### Description

MeSHDb is the simple class for providing the relationship between Entrez gene IDs and MeSH IDs. It provides the database connection and easily accessible with columns, keytypes, keys and select. Some users may use additional functions such as dbconn, dbfile, dbschema, dbInfo, and species for much complex data acquisition.

columns shows which kinds of data can be returned for the MeSHDb object.

keytypes allows the user to discover which keytypes can be passed in to select or keys and the keytype argument.

keys returns keys for the database contained in the MeSHDb object. This method is already documented in the keys manual page but is mentioned again here because it's usage with select is so intimate. By default it will return the primary keys for the database, but if used with the keytype argument, it will return the keys from that keytype.

select will retrieve the data as a data.frame based on parameters for selected keys, columns, and keytype arguments.

dbconn returns the connection with database in the package.

dbfile returns the absolute path sqlite file is saved.

dbschema returns the database schema.

dbInfo returns the many meta information about the package.

species returns the species name.

#### Usage

```
columns(x)
keytypes(x)
keys(x, keytype, ...)
select(x, keys, columns, keytype, ...)
dbconn(x)
dbfile(x)
dbschema(x, file = "", show.indices = FALSE)
dbInfo(x)
species(object)
```

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#### Arguments

| x            | the MeSHDb object. But in practice this will mean an object derived from an MeSHDb object such as a MeSH.Hsa.eg.db, MeSH.Mmu.eg.db or many other MeSH.XXX.eg.db (XXX means abbreviation of species name).                        |
|--------------|--|
| object       | same as x  |
| keys         | the keys to select records for from the database. All possible keys are returned by using the keys method.   |
| columns      | the columns or kinds of things that can be retrieved from the database. As with keys, all possible columns are returned by using the columns method.   |
| keytype      | the keytype that matches the keys used. For the select methods, this is used to indicate the kind of ID being used with the keys argument. For the keys method this is used to indicate which kind of keys are desired from keys |
|              | other arguments.   |
| file         | The file argument must be a connection, or a character string naming the file to print to (see the file argument of the cat function for the details).   |
| show.indices | The CREATE INDEX statements are not shown by default. Use show.indices=TRUE to get them.   |

## Value

keys, columns, keytypes, dbfile, dbInfo, and species each return a character vector or possible values. select and dbschema each return a data.frame. dbconn returns database connection.

#### Author(s)

Koki Tsuyuzaki

#### See Also

dbConnect

#### Examples

```
# # load a package that creates an MeSHDb object
# library(MeSH.Mmu.eg.db)
# MeSH.Mmu.eg.db
# ## then the methods can be used on this object.
# cls <- columns(MeSH.Mmu.eg.db)</pre>
# cls
# kts <- keytypes(MeSH.Mmu.eg.db)</pre>
# kt <- kts[2]
# kts
# ks <- head(keys(MeSH.Mmu.eg.db, keytype=kts[2]))</pre>
# ks
# res <- select(MeSH.Mmu.eg.db, keys=ks, columns=cls, keytype=kt)</pre>
# head(res)
# dbconn(MeSH.Mmu.eg.db)
# dbfile(MeSH.Mmu.eg.db)
# dbschema(MeSH.Mmu.eg.db)
```

```
# dbInfo(MeSH.Mmu.eg.db)
# species(MeSH.Mmu.eg.db)
```

meshVersion

#### Description

This function returns the version of MeSH used in the package.

#### Usage

meshVersion(x)

#### Arguments

```
х
```

MeSHDb object such as MeSH.Mmu.eg.db

#### Author(s)

Koki Tsuyuzaki

#### Examples

# library("MeSH.Mmu.eg.db")

```
# meshVersion(MeSH.Mmu.eg.db)
```

metaPA01

Metadata to construct user's original MeSHDb package

### Description

Meta data to construct user's custom MeSHDb

#### Usage

```
data(metaPA01)
```

#### Details

- SOURCEDATE: The date the source data is retrived
- SOURCENAME: Type of source data
- SOURCEURL: The URL of source data
- DBSCHEMA: Database schema
- DBSCHEMAVERSION: The version of database schema
- ORGANISM: The scientific name
- SPECIES: The common name of the species
- package: The pacakge name
- Db type: The type of name (or class name)

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#### nomenclature

#### Examples

```
data(metaPA01)
head(metaPA01)
```

nomenclature A function

```
A function to return the scientific name of package
```

#### Description

This function returns the scientific name of species used in the package.

#### Usage

nomenclature(x)

#### Arguments

#### х

MeSHDb object such as MeSH.Mmu.eg.db

#### Author(s)

Koki Tsuyuzaki

#### Examples

# library("MeSH.Mmu.eg.db")

# nomenclature(MeSH.Mmu.eg.db)

packageName A function to return the name of package

#### Description

This function returns the name of package

#### Usage

packageName(x)

#### Arguments

х

MeSHDb object such as MeSH.Mmu.eg.db

### Author(s)

Koki Tsuyuzaki

## Examples

- # library("MeSH.Mmu.eg.db")
- # packageName(MeSH.Mmu.eg.db)

#### PA01

# Description

Correspondance between Entrez gene IDs of Pseudomonas aeruiginosa PAO1 and MeSH IDs is provided as demo data. This is to demonstrate how to construct user's original MeSHDb package. The data is based on reciprocal BLASTP best hit (E-value < 200) against Bacillus subtilis subsp. spizizenii str. 168.

#### Usage

data(PAO1)

### Details

- 1st Column: Entrez gene ID
- 2nd Column: MeSH ID
- 3rd Column: MeSH Category
- 4th Column: Source ID. In this case, this is the Entrez gend ID of Bacillus subtilis subsp. spizizenii str. 168
- 5th Column: Source Database. In this case, species name.

#### Examples

data(PAO1) head(PAO1)

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