Package: LRBaseDbi (via r-universe)

July 27, 2024

Title DBI to construct LRBase-related package
Description Interface to construct LRBase package (LRBase.XXX.eg.db).
Version 2.15.0
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Depends R (>= $3.5.0$)
Imports methods, stats, utils, AnnotationDbi, RSQLite, DBI, Biobase
Suggests testthat, BiocStyle, AnnotationHub
VignetteBuilder utils
biocViews Infrastructure
License Artistic-2.0
Repository https://bioc.r-universe.dev
RemoteUrl https://github.com/bioc/LRBaseDbi
RemoteRef HEAD
RemoteSha b60c019e05035354a71db04ceda4b325b87d2e33
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Description

LRBaseDb is the simple class for providing the relationship between Entrez gene IDs and LRBase 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, species, lrNomenclature, lrListDatabases, and lrVersion for much complex data acquisition.

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

1rNomenclature returns the scientific name.

lrListDatabases returns the list of databases to correspond Gene ID of ligend gene and Gene ID of receptor gene.

1rVersion returns the version of LRBaseDb.

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)
lrNomenclature(x)
lrListDatabases(x)
lrVersion(x)
```

Arguments

```
{\bf x} the LRBaseDb object converted by LRBaseDbi::LRBaseDb. object same as {\bf x}
```

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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 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.

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, species, and lrNomenclature each return a character vector or possible values. select, dbschema, lrListDatabases, and lrVersion each return a data.frame. dbconn returns database connection.

Author(s)

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See Also

dbConnect

Examples

```
# library("LRBaseDbi")
# library("AnnotationHub")

# # Data retrieval from AnnotationHub
# ah <- AnnotationHub()
# dbfile <- query(ah, c("LRBaseDb", "Sus scrofa", "v001"))[[1]]

# # Constructor
# LRBase.Ssc.eg.db <- LRBaseDbi::LRBaseDb(dbfile)

# # show
# LRBase.Ssc.eg.db

# # dbconn
# dbconn(LRBase.Ssc.eg.db)

# # dbfile
# dbfile(LRBase.Ssc.eg.db)

# # dbschema
# dbschema
# dbschema(LRBase.Ssc.eg.db)</pre>
```

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```
# # dbInfo
# dbInfo(LRBase.Ssc.eg.db)
# # species
# species(LRBase.Ssc.eg.db)
# # 1rNomenclature
# lrNomenclature(LRBase.Ssc.eg.db)
# # lrListDatabases
# lrListDatabases(LRBase.Ssc.eg.db)
# # lrVersion
# lrVersion(LRBase.Ssc.eg.db)
# # columns
# cols <- columns(LRBase.Ssc.eg.db)</pre>
# # keytypes
# kts <- keytypes(LRBase.Ssc.eg.db)</pre>
# # keys
# ks <- keys(LRBase.Ssc.eg.db, keytype="GENEID_L")[seq(10)]</pre>
# # select
# out <- select(LRBase.Ssc.eg.db,</pre>
   columns=cols,
#
     keys=ks,
     keytype="GENEID_L")
```

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