

LRBase.Bta.eg.db

August 24, 2021

LRBase.Bta.eg.db

Annotation package that provides correspondence between Ligand-Receptor genes as Entrez Gene ID

Description

This data represents a collection of annotation packages that can be used as a single object named as package name. This object can be used with the standard four accessor method for all AnnotationDbi objects. Namely: `columns`, `keytypes`, `keys` and `select`. Users are encouraged to read the vignette from the LRBaseDbi package for more details.

Usage

```
LRBase.Bta.eg.db
```

Value

LRBase.XXX.eg.db-type package

Author(s)

Koki Tsuyuzaki

Examples

```
LRBase.Bta.eg.db
cls <- columns(LRBase.Bta.eg.db)
cls
kts <- keytypes(LRBase.Bta.eg.db)
kt <- kts[2]
kts
ks <- head(keys(LRBase.Bta.eg.db, keytype=kt))
ks
res <- select(LRBase.Bta.eg.db, keys=ks, columns=cls, keytype=kt)
head(res)

species(LRBase.Bta.eg.db)
dbInfo(LRBase.Bta.eg.db)
dbfile(LRBase.Bta.eg.db)
dbschema(LRBase.Bta.eg.db)
```

```
dbconn(LRBase.Bta.eg.db)
lrPackageName(LRBase.Bta.eg.db)
lrNomenclature(LRBase.Bta.eg.db)
lrListDatabases(LRBase.Bta.eg.db)
lrVersion(LRBase.Bta.eg.db)
```

Index

LRBase.Bta.eg.db, [1](#)