Package 'GEOmetadb'

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Type Package
Title A compilation of metadata from NCBI GEO
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Depends GEOquery,RSQLite
Suggests knitr, rmarkdown, dplyr, tm, wordcloud
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biocViews Infrastructure
Description The NCBI Gene Expression Omnibus (GEO) represents the largest public repository of microarray data. However, finding data of interest can be challenging using current tools. GEOmetadb is an attempt to make access to the metadata associated with samples, platforms, and datasets much more feasible. This is accomplished by parsing all the NCBI GEO metadata into a SQLite database that can be stored and queried locally. GEOmetadb is simply a thin wrapper around the SQLite database along with associated documentation. Finally, the SQLite database is updated regularly as new data is added to GEO and can be downloaded at will for the most up-to-date metadata. GEOmetadb paper: http://bioinformatics.oxfordjournals.org/cgi/content/short/24/23/2798 .
<pre>URL http://gbnci.abcc.ncifcrf.gov/geo/</pre>
VignetteBuilder knitr
License Artistic-2.0
NeedsCompilation no
R topics documented:
GEOmetadb-package 2 columnDescriptions 2 geoConvert 3 getBiocPlatformMap 4 getSQLiteFile 5

6

Index

2 columnDescriptions

GEOmetadb-package

Query NCBI GEO metadata from a local SQLite database

Description

The NCBI Gene Expression Omnibus (GEO) represents the largest public repository of microarray data. However, finding data of interest can be challenging using current tools. GEOmetadb is an attempt to make access to the metadata associated with samples, platforms, and datasets much more feasible. This is accomplished by parsing all the NCBI GEO metadata into a SQLite database that can be stored and queried locally. GEOmetadb is simply a thin wrapper around the SQLite database along with associated documentation. Finally, the SQLite database is updated regularly as new data is added to GEO and can be downloaded at will for the most up-to-date metadata.

Details

Package: GEOmetadb Type: Package Version: 1.1.5 Date: 2008-09-09 License: Artistic-2.0

Author(s)

Jack Zhu and Sean Davis

Maintainer: Jack Zhu <zhujack@mail.nih.gov>

References

http://meltzerlab.nci.nih.gov/apps/geo, http://gbnci.abcc.ncifcrf.gov/geo/

Examples

```
if(file.exists('GEOmetadb.sqlite')) {
    a <- columnDescriptions()[1:5,]
    b <- geoConvert('GPL97','GSM')
} else {
    print("use getSQLiteFile() to get a copy of the GEOmetadb SQLite file and then rerun the example")
}</pre>
```

columnDescriptions

Get column descriptions for the GEOmetadb database

geoConvert 3

Description

Searching the GEOmetadb database requires a bit of knowledge about the structure of the database and column descriptions. This function returns those column descriptions for all columns in all tables in the database.

Usage

```
columnDescriptions(sqlite_db_name='GEOmetadb.sqlite')
```

Arguments

sqlite_db_name The filename of the GEOmetadb sqlite database file

Value

A three-column data.frame including TableName, FieldName, and Description.

Author(s)

Sean Davis <sdavis2@mail.nih.gov>

References

http://meltzerlab.nci.nih.gov/apps/geo

Examples

```
if(file.exists('GEOmetadb.sqlite')) {
  columnDescriptions()[1:5,]
} else {
  print("You will need to usethe getSQLiteFile() function to get a copy
  of the SQLite database file before this example will work")
}
```

geoConvert

Cross-reference between GEO data types

Description

A common task is to find all the GEO entities of one type associated with another GEO entity (eg., find all GEO samples associated with GEO platform 'GPL96'). This function provides a very fast mapping between entity types to facilitate queries of this type.

Usage

```
geoConvert(in\_list,\ out\_type = c("gse",\ "gpl",\ "gsm",\ "gds",\ "smatrix"),\ sqlite\_db\_name = "GEOmetallist",\ sqlite\_db\_name = "GEOmetall
```

Arguments

```
in_list Character vector of GEO entities to convert from.

out_type Character vector of GEO entity types to which to convert.

sqlite_db_name The filename of the GEOmetadb sqlite database file
```

Value

A list of data.frames.

Author(s)

Jack Zhu <zhujack@mail.nih.gov>

References

http://meltzerlab.nci.nih.gov/apps/geo, http://gbnci.abcc.ncifcrf.gov/geo/

Examples

```
if(file.exists("GEOmetadb.sqlite")) {
  geoConvert('GPL96',out_type='GSM')
} else {
  print("Run getSQLiteFile() to get a copy of the GEOmetadb SQLite file
and then rerun the example")
}
```

getBiocPlatformMap

Get mappings between GPL and Bioconductor microarry annotation packages

Description

Query the gpl table and get GPL information of a given list of Bioconductor microarry annotation packages. Note currently the GEOmetadb does not contains all the mappings, but we are trying to construct a relative complete list.

Usage

```
getBiocPlatformMap(con, bioc='all')
```

Arguments

con Connection to the GEOmetadb.sqlite database

bioc Character vector of Biocondoctor microarry annotation packages, e.g. c('hgu133plus2','hgu95av2').

'all' returns all mappings.

Value

A six-column data.frame including GPL title, GPL accession, bioc_package, manufacturer, organism, data_row_count.

Author(s)

Jack Zhu <zhujack@mail.nih.gov>, Sean Davis <sdavis2@mail.nih.gov>

References

http://meltzerlab.nci.nih.gov/apps/geo

getSQLiteFile 5

Examples

```
if(file.exists('GEOmetadb.sqlite')) {
    con <- dbConnect(SQLite(), "GEOmetadb.sqlite")
getBiocPlatformMap(con)[1:5,]
getBiocPlatformMap(con, bioc=c('hgu133a','hgu95av2'))
dbDisconnect(con)
} else {
    print("You will need to usethe getSQLiteFile() function to get a copy
of the SQLite database file before this example will work")
}</pre>
```

getSQLiteFile

Download and unzip the most recent GEOmetadb SQLite file

Description

This function is the standard method for downloading and unzipping the most recent GEOmetadb SQLite file from the server.

Usage

```
getSQLiteFile(destdir = getwd(), destfile = "GEOmetadb.sqlite.gz")
```

Arguments

destdir The destination directory of the downloaded file

destfile The filename of the downloaded file. This filename should end in ".gz" as the

unzipping assumes that is the case

Value

Prints some diagnostic information to the screen.

Returns the local filename for use later.

Author(s)

Sean Davis <sdavis2@mail.nih.gov>

References

http://meltzerlab.nci.nih.gov/apps/geo, http://gbnci.abcc.ncifcrf.gov/geo/

Examples

```
## Not run: geometadbfile <- getSQLiteFile()</pre>
```

Index

```
*Topic IO
    geoConvert, 3
    getSQLiteFile, 5
*Topic database
    columnDescriptions, 2
    geoConvert, 3
    getBiocPlatformMap, 4
    getSQLiteFile, 5
*Topic package
    GEOmetadb-package, 2
\verb|columnDescriptions|, 2
geoConvert, 3
{\tt GEOmetadb}\,({\tt GEOmetadb-package}),\,2
GEOmetadb-package, 2
getBiocPlatformMap, 4
getSQLiteFile, 5
```