Using the *GEOmetadb* package

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1 Overview of *GEOmetadb*

The NCBI Gene Expression Omnibus (GEO) represents the largest repository of microarray data in existence. One difficulty in dealing with GEO is finding the microarray data that is of interest. As part of the NCBI Entrez search system, GEO can be searched online via web pages or using NCBI Eutils. However, the web search is not as full-featured as it could be, particularly for programmatic access. NCBI Eutils offers another option for finding data within the vast stores of GEO, but it is cumbersome to use, often requiring multiple

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complicated Eutils calls to get at the relevant information. We have found it **absolutely critical** to have ready access not just to the microarray data, but to the metadata describing the microarray experiments. To this end we have created *GEOmetadb*.

1.1 What is *GEOmetadb*?

The *GEOmetadb* is an attempt to make querying the metadata describing microarray experiments, platforms, and datasets both easier and more powerful. At the heart of *GEOmetadb* is a SQLite database that stores nearly all the metadata associated with all GEO data types including GEO samples (GSM), GEO platforms (GPL), GEO data series (GSE), and curated GEO datasets (GDS), as well as the relationships between these data types. This database is generated by our server by parsing all the records in GEO and needs to be downloaded via a simple helper function to the user's local machine before *GEOmetadb* is useful. Once this is done, the entire GEO database is accessible with simple SQL-based queries. With the GEOmetadb database, queries that are simply not possible using NCBI tools or web pages are often quite simple. The relationships between the tables in the GEOmetadb SQLite database can be seen in figure 1.

1.2 Conversion capabilities

A very typical problem for large-scale consumers of GEO data is to determine the relationships between various GEO accession types. As examples, consider the following questions:

- What samples are associated with GEO platform "GPL96", which represents the Affymetrix hgu133a array?
- What GEO Series were performed using "GPL96"?
- What samples are in my favorite three GEO Series records?
- How many samples are associated with the ten most popular GEO platforms?

Because these types of questions are common, *GEOmetadb* contains the function geoConvert that addresses these questions directly and efficiently.

1.3 What *GEOmetadb* is not

We have faithfully parsed and maintained in GEO when creating *GEOmetadb*. This means that limitations inherent to GEO are also inherent in *GEOmetadb*. We have made no attempt to curate, semantically recode, or otherwise "clean up" GEO; to do so would require significant resources, which we do not have.

GEOmetadb does not contain any microarray data. For access to microarray data from within R/Bioconductor, please look at the *GEOquery* package. In fact, we would expect that many users will find that the combination of *GEOmetadb* and *GEOquery* is quite powerful.

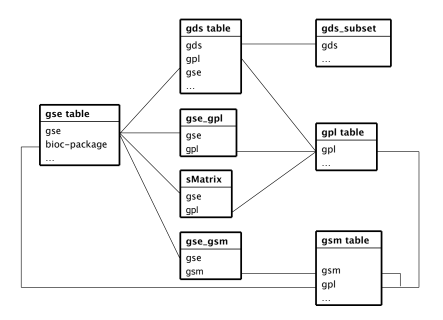


Figure 1: A graphical representation (sometimes called an *Entity-Relationship Diagram*) of the relationships between the tables in the GEOmetadb SQLite database

2 Getting Started

Once *GEOmetadb* is installed (see the Bioconductor website for full installation instructions), we are ready to begin.

2.1 Getting the *GEOmetadb* database

This package does not come with a pre-installed version of the database. This has the advantage that the user will get the most up-to-date version of the database to start; the database can be re-downloaded using the same command as often as desired. First, load the library.

> library(GEOmetadb)

The download and uncompress steps are done automatically with a single command, getSQLiteFile.

```
> getSQLiteFile()
```

The default storage location is in the current working directory and the default filename is "GEOmetadb.sqlite"; it is best to leave the name unchanged unless there is a pressing reason to change it.

Since this SQLite file is of key importance in *GEOmetadb*, it is perhaps of some interest to know some details about the file itself.

> file.info('GEOmetadb.sqlite')

```
size isdir mode

GEOmetadb.sqlite 2091419648 FALSE 644

mtime

GEOmetadb.sqlite 2012-10-01 20:40:10

ctime

GEOmetadb.sqlite 2012-10-01 20:40:10

atime uid gid uname

GEOmetadb.sqlite 2012-10-01 20:40:10 691 692 biocbuild

grname

GEOmetadb.sqlite compbio
```

Now, the SQLite file is available for connection. The standard DBI functionality as implemented in RSQLite function dbConnect makes the connection to the database. The dbDisconnect function disconnects the connection.

```
> con <- dbConnect(SQLite(),'GEOmetadb.sqlite')
> dbDisconnect(con)
```

[1] TRUE

The variable con is an *RSQLite* connection object.

2.2 A word about SQL

The Structured Query Language, or SQL, is a very powerful and standard way of working with relational data. GEO is composed of several data types, all of which are related to each other; in fact, NCBI uses a relational SQL database for metadata storage and querying. SQL databases and SQL itself are designed specifically to work efficiently with just such data. While the goal of many programming projects and programmers is to hide the details of SQL from the user, we are of the opinion that such efforts may be counterproductive, particularly with complex data and the need for *ad hoc* queries, both of which are characteristics with GEO metadata. We have taken the view that exposing the power of SQL will enable users to maximally utilize the vast data repository that is GEO. We understand that many users are not accustomed to working with SQL and, therefore, have devoted a large section of the vignette to working examples. Our goal is not to teach SQL, so a quick tutorial of SQL is likely to be beneficial to those who have not used it before. Many such tutorials are available online and can be completed in 30 minutes or less.

3 Examples

3.1 Interacting with the database

The functionality covered in this section is covered in much more detail in the DBI and RSQLite package documentation. We cover enough here only to be useful.

Again, we connect to the database.

```
> con <- dbConnect(SQLite(),'GEOmetadb.sqlite')</pre>
```

The dbListTables function lists all the tables in the SQLite database handled by the connection object con.

```
> geo_tables <- dbListTables(con)
> geo_tables
```

[1]	"gds"	"gds_subset"
[3]	"geoConvert"	"geodb_column_desc"
[5]	"gpl"	"gse"
[7]	"gse_gpl"	"gse_gsm"
[9]	"gsm"	"metaInfo"
[11]	"sMatrix"	

There is also the dbListFields function that can list database fields associated with a table.

```
> dbListFields(con,'gse')
```

[1]	"ID"	"title"
[3]	"gse"	"status"
[5]	"submission_date"	"last_update_date"
[7]	"pubmed_id"	"summary"
[9]	"type"	"contributor"
[11]	"web_link"	"overall_design"
[13]	"repeats"	"repeats_sample_list"
[15]	"variable"	"variable_description"
[17]	"contact"	"supplementary_file"

Sometimes it is useful to get the actual SQL schema associated with a table. As an example of doing this and using an *RSQLite* shortcut function, sqliteQuickSQL, we can get the table schema for the *gpl* table.

> sqliteQuickSQL(con,'PRAGMA TABLE_INFO(gpl)')

	cid	name	type	notnull	dflt_value	pk
1	0	ID	REAL	0	<na></na>	0
2	1	title	TEXT	0	<na></na>	0
3	2	gpl	TEXT	0	<na></na>	0
4	3	status	TEXT	0	<na></na>	0
5	4	submission_date	TEXT	0	<na></na>	0
6	5	last_update_date	TEXT	0	<na></na>	0
7	6	technology	TEXT	0	<na></na>	0
8	7	distribution	TEXT	0	<na></na>	0
9	8	organism	TEXT	0	<na></na>	0
10	9	manufacturer	TEXT	0	<na></na>	0
11	10	manufacture_protocol	TEXT	0	<na></na>	0
12	11	coating	TEXT	0	<na></na>	0
13	12	catalog_number	TEXT	0	<na></na>	0
14	13	support	TEXT	0	<na></na>	0
15	14	description	TEXT	0	<na></na>	0

16	15	web_link	TEXT	0	<na></na>	0
17	16	contact	TEXT	0	<na></na>	0
18	17	data_row_count	REAL	0	<na></na>	0
19	18	<pre>supplementary_file</pre>	TEXT	0	<na></na>	0
20	19	<pre>bioc_package</pre>	TEXT	0	<na></na>	0

3.2 Writing SQL queries and getting results

Select 5 records from the gse table and show the first 7 columns.

```
> rs <- dbGetQuery(con,'select * from gse limit 5')</pre>
> rs[,1:7]
  ID
   1
1
2
  2
3
  3
4
  4
5
  5
                                                       title
1
                                       NHGRI_Melanoma_class
2
                                     Cerebellar development
3
              Renal Cell Carcinoma Differential Expression
4
      Diurnal and Circadian-Regulated Genes in Arabidopsis
5 Global profile of germline gene expression in C. elegans
                      status submission_date
   gse
1 GSE1 Public on Jan 22 2001
                                   2001-01-22
2 GSE2 Public on Apr 26 2001
                                   2001-04-19
3 GSE3 Public on Jul 19 2001
                                   2001-07-19
4 GSE4 Public on Jul 20 2001
                                   2001-07-20
5 GSE5 Public on Jul 24 2001
                                   2001-07-24
  last_update_date pubmed_id
1
        2005-05-29
                    10952317
2
        2005-05-29
                          NA
З
        2005-05-29
                    11691851
4
        2005-05-29
                   11158533
5
        2005-07-18
                    11030340
```

Get the GEO series accession and title from GEO series that were submitted by "Sean Davis". The "

	a so	
1	gse GSE2553	
2	GSE4406	
2 3	GSE4400 GSE5357	
4	GSE7376	
5	GSE7882	
6	GSE8486	
7	GSE9328	
8	GSE14543	
9	GSE15621	
	GSE16087	
11		
12	GSE16091	
13	GSE16102	
14	GSE18544	
15	GSE19063	
16	GSE20016	
17	GSE25164	
18	GSE22520	
	GSE25127	
20	GSE29428	
1		
2		Gene expression profiling of CD4+ T-cells an
3 4		Detection of novel ampl
4 5		Detection of novel ampl Gene Expression and Comparative Genomic Hybridization of Duct
5 6		Whole genome DNAse hype
0 7		whole Peneme Punes The
8		A molecul
9		Acute Lymphocytic Leu
10		Gene expressi
11		Gene express
12		Gene expression pr
13		Gene expression profile
14		ession Profiling of a Mouse Xenograft Model of âĂIJTriple-NegativeâĂŦ Breast
15		Genome-wide map of PAX3-FKHR
		of Human Brain Metastases of Breast Cancer Reveal the Association between HK
17	•	
18		Mouse Models of Alveolar/Embryonal Rhabdo
19		Ewing Sarcoma c
20	A r	nethyl-deviator epigenotype of estrogen-receptor-positive breast carcinoma is

As another example, GEOmetadb can find all samples on GPL96 (Affymetrix hgu133a) that have .CEL files available for download.

```
> rs <- dbGetQuery(con,paste("select gsm,supplementary_file",
+ "from gsm where gpl='GPL96'",
+ "and supplementary_file like '%CEL.gz'"))
> dim(rs)
[1] 21975 2
```

But why limit to only GPL96? Why not look for all Affymetrix arrays that have .CEL files? And list those with their associated GPL information, as well as the Bioconductor annotation package name?

```
> rs[1:5,]
```

	<pre>bioc_package</pre>	gpl	gsm
1	hu6800	GPL80	GSM575
2	hu6800	GPL80	GSM576
3	hu6800	GPL80	GSM577
4	hu6800	GPL80	GSM578
5	hu6800	GPL80	GSM579

```
supplementary_file
```

```
1 ftp://ftp.ncbi.nih.gov/pub/geo/DATA/supplementary/samples/GSMnnn/GSM575/GSM575.cel.gz
2 ftp://ftp.ncbi.nih.gov/pub/geo/DATA/supplementary/samples/GSMnnn/GSM576/GSM576.cel.gz
3 ftp://ftp.ncbi.nih.gov/pub/geo/DATA/supplementary/samples/GSMnnn/GSM577/GSM577.cel.gz
4 ftp://ftp.ncbi.nih.gov/pub/geo/DATA/supplementary/samples/GSMnnn/GSM578/GSM578.cel.gz
5 ftp://ftp.ncbi.nih.gov/pub/geo/DATA/supplementary/samples/GSMnnn/GSM579/GSM579.cel.gz
```

Of course, we can combine programming and data access. A simple sapply example shows how to query each of the tables for number of records.

```
> getTableCounts <- function(tableName,conn) {
+ sql <- sprintf("select count(*) from %s",tableName)
+ return(dbGetQuery(conn,sql)[1,1])
+ }
> do.call(rbind,sapply(geo_tables,getTableCounts,con,simplify=FALSE))
```

	[,1]
gds	2721
gds_subset	15275
geoConvert	3565064
<pre>geodb_column_desc</pre>	104
gpl	10722
gse	32661
gse_gpl	38459
gse_gsm	894949
gsm	765310
metaInfo	2
sMatrix	39186

3.3 Conversion of GEO entity types

Large-scale consumers of GEO data might want to convert GEO entity type from one to others, e.g. finding all GSM and GSE associated with 'GPL96'. Function goeConvert does the conversion with a very fast mapping between entity types.

Covert 'GPL96' to other possible types in the GEOmetadb.sqlite.

```
> conversion <- geoConvert('GPL96')</pre>
```

Check what GEO types and how many entities in each type in the conversion.

```
> lapply(conversion, dim)
$gse
[1] 932
          2
$gsm
[1] 31484
              2
$gds
[1] 325
          2
$sMatrix
[1] 983
          2
> conversion$gse[1:5,]
  from_acc
             to_acc
1
     GPL96 GSE1000
2
     GPL96 GSE10024
3
     GPL96 GSE10043
4
     GPL96 GSE10072
5
     GPL96 GSE10089
```

```
> conversion$gsm[1:5,]
  from_acc
              to_acc
     GPL96 GSM100386
1
2
     GPL96 GSM100454
3
     GPL96 GSM100455
4
     GPL96 GSM100456
5
     GPL96 GSM100457
> conversion$gds[1:5,]
  from_acc to_acc
1
     GPL96 GDS1023
2
     GPL96 GDS1036
3
     GPL96 GDS1050
4
     GPL96 GDS1062
5
     GPL96 GDS1063
> conversion$sMatrix[1:5,]
  from_acc
                                   to_acc
1
     GPL96 GSE1000_series_matrix.txt.gz
2
     GPL96 GSE10024_series_matrix.txt.gz
     GPL96 GSE10043_series_matrix.txt.gz
3
4
     GPL96 GSE10072_series_matrix.txt.gz
     GPL96 GSE10089_series_matrix.txt.gz
5
```

3.4 Mappings between GPL and Bioconductor microarry annotation packages

The function getBiocPlatformMap is to 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.

Get GPL information of 'hgu133a' and 'hgu95av2':

```
> getBiocPlatformMap(con, bioc=c('hgu133a', 'hgu95av2'))
                                            title
                                                     gpl
1 [HG-U133A] Affymetrix Human Genome U133A Array GPL96
2
    [HG_U95A] Affymetrix Human Genome U95A Array GPL91
  bioc_package manufacturer
                                 organism
                 Affymetrix Homo sapiens
1
       hgu133a
2
      hgu95av2
                 Affymetrix Homo sapiens
  data_row_count
           22283
1
2
           12626
```

3.5 More advanced queries

Now, for something a bit more complicated, we would like to find all the human breast cancer-related Affymetrix gene expression GEO series.

```
> sql <- paste("SELECT DISTINCT gse.title,gse.gse",</pre>
+
                "FROM",
                II
                   gsm JOIN gse_gsm ON gsm.gsm=gse_gsm.gsm",
+
                II.
                   JOIN gse ON gse_gsm.gse=gse.gse",
+
                11
                   JOIN gse_gpl ON gse_gpl.gse=gse.gse",
+
                II.
                   JOIN gpl ON gse_gpl.gpl=gpl.gpl",
+
                "WHERE",
+
                   gsm.molecule_ch1 like '%total RNA%' AND",
+
                   gse.title LIKE '%breast cancer%' AND",
+
                   gpl.organism LIKE '%Homo sapiens%'",sep=" ")
                II
+
> rs <- dbGetQuery(con,sql)</pre>
> dim(rs)
[1] 391
          2
> print(rs[1:5,],right=FALSE)
  title
1 Breast Cancer aCGH
2 Human mammary epithelium and breast cancer
3 Cell-type specific responses to chemotherapeutics in breast cancer
4 E2 and SERM Treatment of MCF-7 Breast Cancer Cells
5 Estradiol Treated Breast Cancer Cells Expressing Mutant Estrogen Receptors
  gse
1 GSE3281
2 GSE53
3 GSE763
4 GSE848
5 GSE1045
   Finally, it is probably a good idea to close the connection, please see DBI for detail.
```

```
> dbDisconnect(con)
```

[1] TRUE

If you want to remove old GEOmetadb.sqlite file before retrieve a new version from the server, execute the following codes:

```
> file.remove('GEOmetadb.sqlite')
```

[1] TRUE