# Using the GEOquery package

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## 1 Overview of GEO

The NCBI Gene Expression Omnibus (GEO) serves as a public repository for a wide range of high-throughput experimental data. These data include single and dual channel microarraybased experiments measuring mRNA, genomic DNA, and protein abundance, as well as non-array techniques such as serial analysis of gene expression (SAGE), mass spectrometry proteomic data, and high-throughput sequencing data.

At the most basic level of organization of GEO, there are four basic entity types. The first three (Sample, Platform, and Series) are supplied by users; the fourth, the dataset, is compiled and curated by GEO staff from the user-submitted data.<sup>1</sup>

#### 1.1 Platforms

A Platform record describes the list of elements on the array (e.g., cDNAs, oligonucleotide probesets, ORFs, antibodies) or the list of elements that may be detected and quantified in that experiment (e.g., SAGE tags, peptides). Each Platform record is assigned a unique and stable GEO accession number (GPLxxx). A Platform may reference many Samples that have been submitted by multiple submitters.

#### 1.2 Samples

A Sample record describes the conditions under which an individual Sample was handled, the manipulations it underwent, and the abundance measurement of each element derived from it. Each Sample record is assigned a unique and stable GEO accession number (GSMxxx). A Sample entity must reference only one Platform and may be included in multiple Series.

#### 1.3 Series

A Series record defines a set of related Samples considered to be part of a group, how the Samples are related, and if and how they are ordered. A Series provides a focal point and description of the experiment as a whole. Series records may also contain tables describing extracted data, summary conclusions, or analyses. Each Series record is assigned a unique and stable GEO accession number (GSExxx). Series records are available in a couple of formats which are handled by GEOquery independently. The smaller and new GSEMatrix files are quite fast to parse; a simple flag is used by GEOquery to choose to use GSEMatrix files (see below).

#### 1.4 Datasets

GEO DataSets (GDSxxx) are curated sets of GEO Sample data. A GDS record represents a collection of biologically and statistically comparable GEO Samples and forms the basis

<sup>&</sup>lt;sup>1</sup>See http://www.ncbi.nih.gov/geo for more information

of GEO's suite of data display and analysis tools. Samples within a GDS refer to the same Platform, that is, they share a common set of probe elements. Value measurements for each Sample within a GDS are assumed to be calculated in an equivalent manner, that is, considerations such as background processing and normalization are consistent across the dataset. Information reflecting experimental design is provided through GDS subsets.

## 2 Getting Started using GEOquery

Getting data from GEO is really quite easy. There is only one command that is needed, getGEO. This one function interprets its input to determine how to get the data from GEO and then parse the data into useful R data structures. Usage is quite simple:

```
> library(GEOquery)
```

This loads the GEOquery library.

```
> gds <- getGEO("GDS10")</pre>
```

```
File stored at:
/tmp/RtmpnFTpz5/GDS10.soft
```

Now, gds contains the R data structure (of class GDS) that represents the GDS1 entry from GEO. You'll note that the filename used to store the download was output to the screen (but not saved anywhere) for later use to a call to getGEO(filename=...).

We can do the same with any other GEO accession, such as GSM3, a GEO sample.

```
> gsm <- getGEO("GSM3")</pre>
```

File stored at: /tmp/RtmpnFTpz5/GSM3.soft

## **3** GEOquery Data Structures

The GEOquery data structures really come in two forms. The first, comprising GDS, GPL, and GSM all behave similarly and accessors have similar effects on each. The fourth GEOquery data structure, GSE is a composite data type made up of a combination of GSM and GPL objects. I will explain the first three together first.

#### 3.1 The GDS, GSM, and GPL classes

Each of these classes is comprised of a metadata header (taken nearly verbatim from the SOFT format header) and a GEODataTable. The GEODataTable has two simple parts, a Columns part which describes the column headers on the Table part. There is also a *show* method for each class. For example, using the gsm from above:

> Meta(gsm)

\$channel\_count
[1] "1"

\$contact\_address
[1] "6 Center Drive"

\$contact\_city
[1] "Bethesda"

\$contact\_country
[1] "USA"

\$contact\_department
[1] "LCDB"

\$contact\_email
[1] "oliver@helix.nih.gov"

\$contact\_fax
[1] "301-496-5239"

\$contact\_institute
[1] "NIDDK, NIH"

\$contact\_name
[1] "Brian,,Oliver"

\$contact\_phone
[1] "301-496-5495"

\$contact\_state
[1] "MD"

\$contact\_web\_link
[1] "http://www.niddk.nih.gov/intram/people/boliver.htm"

\$`contact\_zip/postal\_code` [1] "20892" \$data\_row\_count [1] "3456" \$description [1] "Testis dissected from adult (12-24 hours post-eclosion) Drosophila melanogaster of [2] "Keywords = gonad, male, sex" \$geo\_accession [1] "GSM3" \$last\_update\_date [1] "May 27 2005" \$molecule\_ch1 [1] "total RNA" \$organism\_ch1 [1] "Drosophila melanogaster" \$platform\_id [1] "GPL5" \$series\_id [1] "GSE462" \$source\_name\_ch1 [1] "y w[67c1]/Y testis" \$status [1] "Public on Oct 18 2000" \$submission\_date [1] "Oct 18 2000" \$supplementary\_file [1] "NONE" \$title

[1] "testis a" \$type [1] "RNA" > Table(gsm)[1:5, ] ID\_REF SIGNAL\_RAW BKD\_FORM NORM\_FORM BKD\_RAW NORM\_VALUE CONST 1 1 138392.65 no 101113.7775 395070.1312 39542 no 2 2 100973.49 no 101113.7775 395070.1312 39542 no 3 118994.03 no 101113.7775 395070.1312 39542 3 no 4 no 101113.7775 395070.1312 39542 4 108126.05 yes 5 293362.11 no 101113.7775 395070.1312 39542 5 no VALUE 1 76820.87249 2 39401.7125 3 57422.25249 4 46554.2725 5 231790.3324 > Columns(gsm) Column Description 1 ID\_REF 2 SIGNAL\_RAW raw signal BKD\_FORM З 4 NORM\_FORM BKD\_RAW raw background as taken in four quarters of microarray 5 6 NORM\_VALUE normalization value 7 constant value CONST 8 VALUE

The GPL behaves exactly as the GSM class. However, the GDS has a bit more information associated with the *Columns* method:

```
> Columns(gds)
```

	sample	tissue	strain	disease.state
1	GSM582	spleen	NOD	diabetic
2	GSM589	spleen	NOD	diabetic
3	GSM583	spleen	Idd3	diabetic-resistant
4	GSM590	spleen	Idd3	diabetic-resistant
5	GSM584	spleen	Idd5	diabetic-resistant
6	GSM591	spleen	Idd5	diabetic-resistant

7	GSM585	spleen	Idd3	+Idd5	diabeti	c-res	istant
8	GSM592	spleen	Idd3	+Idd5	diabeti	c-res:	istant
9	GSM586	spleen		Idd9	diabeti	c-res:	istant
10	GSM593	spleen		Idd9	diabeti	c-res:	istant
11	GSM587	spleen	B10	.H2g7	1	nondia	abetic
12	GSM594	spleen	B10	.H2g7	1	nondia	abetic
13	GSM588	spleen	B10.H2g7	Idd3	1	nondia	abetic
14	GSM595	spleen	B10.H2g7	Idd3	1	nondia	abetic
15	GSM596	thymus	-	NOD		dia	abetic
16	GSM603	thymus		NOD		dia	abetic
17	GSM597	thymus		Idd3	diabeti	c-res:	istant
18	GSM604	thymus		Idd3	diabeti	c-res:	istant
19	GSM598	thymus		Idd5	diabeti	c-res:	istant
20	GSM605	thymus		Idd5	diabeti	c-res:	istant
21	GSM599	thymus	Idd3	+Idd5	diabeti	c-res:	istant
22	GSM606	thymus	Idd3	+Idd5	diabeti	c-res:	istant
23	GSM600	thymus		Idd9	diabeti	c-res:	istant
24	GSM607	thymus		Idd9	diabeti	c-res:	istant
25	GSM601	thymus	B10	.H2g7	1	nondia	abetic
26	GSM608	thymus	B10	.H2g7	1	nondia	abetic
27	GSM602	thymus	B10.H2g7	Idd3	1	nondia	abetic
28	GSM609	thymus	B10.H2g7	Idd3	1	nondia	abetic
						desc	ription
1		Valı	ue for GS	M582:	NOD_S1;	<pre>src:</pre>	Spleen
2		Valı	ue for GS	M589:	NOD_S2;	<pre>src:</pre>	Spleen
3		Value	e for GSM	1583:	Idd3_S1;	<pre>src:</pre>	Spleen
4		Value	e for GSM	1590:	Idd3_S2;	<pre>src:</pre>	Spleen
5		Value	e for GSM	1584:	Idd5_S1;	<pre>src:</pre>	Spleen
6		Value	e for GSM	1591:	Idd5_S2;	<pre>src:</pre>	Spleen
7		Value 1	or GSM58	5: Id	d3+5_S1;	<pre>src:</pre>	Spleen
8		Value 1	for GSM59	2: Id	d3+5_S2;	<pre>src:</pre>	Spleen
9		Value	e for GSM	1586:	Idd9_S1;	<pre>src:</pre>	Spleen
10		Value	e for GSM	1593:	Idd9_S2;	<pre>src:</pre>	Spleen
11	Va	alue for	GSM587:	B10.	H2g7_S1;	<pre>src:</pre>	Spleen
12	Va	alue for	GSM594:	B10.	H2g7_S2;	<pre>src:</pre>	Spleen
13	Value 1	for GSM5	588: B10.	H2g7	Idd3_S1;	<pre>src:</pre>	Spleen
14	Value 1	for GSM5	595: B10.	H2g7	Idd3_S2;	<pre>src:</pre>	Spleen
15		Valı	ue for GS	M596:	NOD_T1;	<pre>src:</pre>	Thymus
16		Valı	ue for GS	M603:	NOD_T2;	<pre>src:</pre>	Thymus
17		Value	e for GSM	1597:	Idd3_T1;	<pre>src:</pre>	Thymus
18		Value	e for GSM	604:	Idd3_T2;	<pre>src:</pre>	Thymus
19		Value	e for GSM	1598:	Idd5_T1;	<pre>src:</pre>	Thymus

```
20
            Value for GSM605: Idd5_T2; src: Thymus
21
          Value for GSM599: Idd3+5_T1; src: Thymus
22
          Value for GSM606: Idd3+5_T2; src: Thymus
23
            Value for GSM600: Idd9_T1; src: Thymus
24
            Value for GSM607: Idd9_T2; src: Thymus
25
        Value for GSM601: B10.H2g7_T1; src: Thymus
26
        Value for GSM608: B10.H2g7_T2; src: Thymus
27 Value for GSM602: B10.H2g7 Idd3_T1; src: Thymus
28 Value for GSM609: B10.H2g7 Idd3_T2; src: Thymus
```

#### 3.2 The GSE class

The GSE is the most confusing of the GEO entities. A GSE entry can represent an arbitrary number of samples run on an arbitrary number of platforms. The GSE has a metadata section, just like the other classes. However, it doesn't have a GEODataTable. Instead, it contains two lists, accessible using GPLList and GSMList, that are each lists of GPL and GSM objects. To show an example:

```
> gse <- getGEO("GSE462", GSEMatrix = FALSE)
File stored at:
/tmp/RtmpnFTpz5/GSE462.soft
Parsing....
^{PLATFORM} = GPL5
^{SAMPLE} = GSM3
^{SAMPLE} = GSM4
^{SAMPLE} = GSM5
^{SAMPLE} = GSM6
^{SAMPLE} = GSM7
^SAMPLE = GSM8
^{SAMPLE} = GSM9
> Meta(gse)
$contact_address
[1] "6 Center Drive"
$contact_city
[1] "Bethesda"
$contact_country
[1] "USA"
```

\$contact\_department

[1] "LCDB" \$contact\_email [1] "oliver@helix.nih.gov" \$contact\_fax [1] "301-496-5239" \$contact\_institute [1] "NIDDK, NIH" \$contact\_name [1] "Brian,,Oliver" \$contact\_phone [1] "301-496-5495" \$contact\_state [1] "MD" \$contact\_web\_link [1] "http://www.niddk.nih.gov/intram/people/boliver.htm" \$`contact\_zip/postal\_code` [1] "20892" \$contributor [1] "Justen,,Andrews" "Gerard,G,Bouffard" "Chris,,Cheadle" [4] "Jining,,LÃij" "Kevin,G,Becker" "Brian,,Oliver" \$geo\_accession [1] "GSE462" \$last\_update\_date [1] "Oct 28 2005" \$platform\_id [1] "GPL5" \$pubmed\_id [1] "11116097"

\$sample\_id [1] "GSM10" "GSM3" "GSM4" "GSM5" "GSM6" "GSM7" "GSM8" "GSM9" \$status [1] "Public on Jul 16 2003" \$submission\_date [1] "Jun 25 2003" \$summary [1] "Identification and annotation of all the genes in the sequenced Drosophila genome i \$title [1] "Analysis of transcription in the Drosophila melanogaster testis" \$type [1] "other" > names(GSMList(gse)) [1] "GSM10" "GSM3" "GSM4" "GSM5" "GSM6" "GSM7" "GSM8" "GSM9" > GSMList(gse)[[1]] An object of class "GSM" channel\_count [1] "1" contact\_address [1] "6 Center Drive" contact\_city [1] "Bethesda" contact\_country [1] "USA" contact\_department [1] "LCDB" contact\_email [1] "oliver@helix.nih.gov" contact\_fax [1] "301-496-5239" contact\_institute [1] "NIDDK, NIH" contact\_name [1] "Brian,,Oliver"

```
contact_phone
[1] "301-496-5495"
contact_state
[1] "MD"
contact_web_link
[1] "http://www.niddk.nih.gov/intram/people/boliver.htm"
contact_zip/postal_code
[1] "20892"
data_row_count
[1] "3456"
description
[1] "Whole adult male minus (12-24 hours post-eclosion) Drosophila melanogaster of the g
geo_accession
[1] "GSM10"
last_update_date
[1] "Mar 09 2006"
molecule_ch1
[1] "total RNA"
organism_ch1
[1] "Drosophila melanogaster"
platform_id
[1] "GPL5"
series_id
[1] "GSE462"
source_name_ch1
[1] "y w[67c1] female"
status
[1] "Public on Oct 18 2000"
submission_date
[1] "Oct 18 2000"
title
[1] "female b"
type
[1] "RNA"
An object of class "GEODataTable"
***** Column Descriptions *****
      Column
                     Description
1
      ID_REF
2 SIGNAL_RAW
                      raw signal
   BKD_FORM
З
4 NORM_FORM
5
     BKD_RAW
                  raw background
```

```
6 NORM_VALUE normalization value
7
       CONST
                  constant value
8
       VALUE
***** Data Table *****
  ID_REF SIGNAL_RAW BKD_FORM NORM_FORM BKD_RAW NORM_VALUE CONST
                                                                       VALUE
            4486.49
                            0
                                       0 3379.579
                                                    23337.54 39542 55845.45
1
       1
2
       2
                            0
                                       0 3379.579
                                                    23337.54 39542 41058.05
            3482.51
                                                    23337.54 39542 45916.78
3
            3812.39
                            0
                                       0 3379.579
       З
4
            3257.56
                                       0 3379.579
                                                    23337.54 39542 37744.81
       4
                            1
5
                            0
                                       0 3379.579
                                                    23337.54 39542 69843.97
       5
            5436.91
3450 more rows ...
> names(GPLList(gse))
[1] "GPL5"
```

See below for an additional, preferred method of obtaining GSE information.

## 4 Converting to BioConductor ExpressionSets and limma MALists

GEO datasets are (unlike some of the other GEO entities), quite similar to the *limma* data structure *MAList* and to the *Biobase* data structure *ExpressionSet*. Therefore, there are two functions, GDS2MA and GDS2eSet that accomplish that task.

#### 4.1 Getting GSE Series Matrix files as an ExpressionSet

GEO Series are collections of related experiments. In addition to being available as SOFT format files, which are quite large, NCBI GEO has prepared a simpler format file based on tab-delimited text. The getGEO function can handle this format and will parse very large GSEs quite quickly. The data structure returned from this parsing is a list of ExpressionSets. As an example, we download and parse GSE2553.

```
> gse2553 <- getGEO("GSE2553", GSEMatrix = TRUE)
```

Found 1 file(s)
GSE2553\_series\_matrix.txt.gz
File stored at:
/tmp/RtmpnFTpz5/GPL1977.soft

> show(gse2553)

```
$GSE2553_series_matrix.txt.gz
ExpressionSet (storageMode: lockedEnvironment)
assayData: 12600 features, 181 samples
  element names: exprs
phenoData
  sampleNames: GSM48681, GSM48682, ..., GSM48861 (181 total)
  varLabels and varMetadata description:
    title: NA
    geo_accession: NA
    . . . : . . .
    data_row_count: NA
    (27 total)
featureData
  featureNames: 1, 2, ..., 12600 (12600 total)
  fvarLabels and fvarMetadata description:
    ID: NA
    PenAt: NA
    . . . : . . .
    Chimeric_Cluster_IDs: NA
    (13 total)
  additional fvarMetadata: Column, Description
experimentData: use 'experimentData(object)'
Annotation: GPL1977
> show(pData(phenoData(gse2553[[1]]))[1:5, c(1, 6, 8)])
                                                                   title type
GSM48681
                              Patient sample ST18, Dermatofibrosarcoma
                                                                          RNA
GSM48682
                                    Patient sample ST410, Ewing Sarcoma
                                                                          RNA
GSM48683
                                     Patient sample ST130, Sarcoma, NOS
                                                                          RNA
GSM48684 Patient sample ST293, Malignant Peripheral Nerve Sheath Tumor
                                                                          RNA
GSM48685
                                      Patient sample ST367, Liposarcoma
                                                                          RNA
                                  source_name_ch1
GSM48681
                             Dermatofibrosarcoma
GSM48682
                                    Ewing Sarcoma
GSM48683
                                     Sarcoma, NOS
GSM48684 Malignant Peripheral Nerve Sheath Tumor
GSM48685
                                      Liposarcoma
```

#### 4.2 Converting GDS to an ExpressionSet

Taking our gds object from above, we can simply do:

```
> eset <- GDS2eSet(gds, do.log2 = TRUE)</pre>
```

```
File stored at:
/tmp/RtmpnFTpz5/GPL24.annot
```

Now, eset is an *ExpressionSet* that contains the same information as in the GEO dataset, including the sample information, which we can see here:

```
> eset
```

```
ExpressionSet (storageMode: lockedEnvironment)
assayData: 39114 features, 28 samples
  element names: exprs
phenoData
  sampleNames: GSM582, GSM589, ..., GSM609
                                             (28 total)
  varLabels and varMetadata description:
    sample: NA
    tissue: NA
    . . . : . . .
    description: NA
    (5 total)
featureData
  featureNames: 1, 2, ..., 39114 (39114 total)
  fvarLabels and fvarMetadata description:
    ID: ID from Platform data table
    Gene.title: Entrez Gene name
    . . . : . . .
    GO.Component.1: Gene Ontology Component identifier
    (21 total)
  additional fvarMetadata: Column
experimentData: use 'experimentData(object)'
  pubMedIds: 11827943
Annotation:
> pData(eset)
       sample tissue
                                        disease.state
                           strain
GSM582 GSM582 spleen
                               NOD
                                              diabetic
GSM589 GSM589 spleen
                               NOD
                                              diabetic
GSM583 GSM583 spleen
                             Idd3 diabetic-resistant
GSM590 GSM590 spleen
                              Idd3 diabetic-resistant
GSM584 GSM584 spleen
                              Idd5 diabetic-resistant
GSM591 GSM591 spleen
                              Idd5 diabetic-resistant
GSM585 GSM585 spleen
                         Idd3+Idd5 diabetic-resistant
GSM592 GSM592 spleen
                         Idd3+Idd5 diabetic-resistant
GSM586 GSM586 spleen
                              Idd9 diabetic-resistant
```

GSM593	GSM593	spleen		Idd9	diabeti	c-res	istant
GSM587	GSM587 s	spleen	B10	.H2g7	1	nondia	abetic
GSM594	GSM594 s	spleen	B10	.H2g7	]	nondia	abetic
GSM588	GSM588 s	spleen	B10.H2g7	Idd3	1	nondia	abetic
GSM595	GSM595	spleen	B10.H2g7	Idd3	1	nondia	abetic
GSM596	GSM596	thymus		NOD		dia	abetic
GSM603	GSM603	thymus		NOD		dia	abetic
GSM597	GSM597	thymus		Idd3	diabeti	c-res	istant
GSM604	GSM604 t	thymus		Idd3	diabeti	c-res	istant
GSM598	GSM598 t	thymus		Idd5	diabeti	c-res	istant
GSM605	GSM605 t	thymus		Idd5	diabeti	c-res	istant
GSM599	GSM599 t	thymus	Idd3-	+Idd5	diabeti	c-res	istant
GSM606	GSM606 t	thymus	Idd3-	+Idd5	diabeti	c-res	istant
GSM600	GSM600	thymus		Idd9	diabeti	c-res	istant
GSM607	GSM607	thymus		Idd9	diabeti	c-res	istant
GSM601	GSM601	thymus	B10	.H2g7	]	nondia	abetic
GSM608	GSM608	thymus	B10	.H2g7	1	nondia	abetic
GSM602	GSM602	thymus	B10.H2g7	Idd3	1	nondia	abetic
GSM609	GSM609	thymus	B10.H2g7	Idd3	1	nondia	abetic
						desci	ription
GSM582		Valı	ie for GSN	4582:	NOD_S1;	<pre>src:</pre>	${\tt Spleen}$
GSM589		Valı	le for GSN	1589:	NOD_S2;	<pre>src:</pre>	${\tt Spleen}$
GSM583		Value	e for GSM	583: 1	Idd3_S1;	<pre>src:</pre>	Spleen
GSM590		Value	e for GSMS	590: 1	Idd3_S2;	<pre>src:</pre>	${\tt Spleen}$
GSM584		Value	e for GSM	584: 1	Idd5_S1;	<pre>src:</pre>	Spleen
GSM591		Value	e for GSM	591: 1	Idd5_S2;	<pre>src:</pre>	Spleen
GSM585		Value 1	for GSM588	5: Ido	13+5_S1;	<pre>src:</pre>	Spleen
GSM592	1	Value 1	for GSM592	2: Ido	13+5_S2;	<pre>src:</pre>	Spleen
GSM586		Value	e for GSM	586: 1	Idd9_S1;	<pre>src:</pre>	Spleen
GSM593		Value	e for GSMS	593: 1	Idd9_S2;	<pre>src:</pre>	Spleen
GSM587	Val	lue foi	GSM587:	B10.H	H2g7_S1;	<pre>src:</pre>	Spleen
GSM594	Val	lue foi	GSM594:	B10.H	H2g7_S2;	<pre>src:</pre>	Spleen
GSM588	Value fo	or GSM5	588: B10.H	12g7 ]	Idd3_S1;	<pre>src:</pre>	Spleen
GSM595	Value fo	or GSM5	595: B10.H	12g7 ]	Idd3_S2;	<pre>src:</pre>	Spleen
GSM596		Valı	ie for GSN	1596:	NOD_T1;	<pre>src:</pre>	Thymus
GSM603		Valı	ie for GSN	1603:	NOD_T2;	<pre>src:</pre>	Thymus
GSM597		Value	e for GSM	597: 1	Idd3_T1;	<pre>src:</pre>	Thymus
GSM604		Value	e for GSM6	504: I	Idd3_T2;	<pre>src:</pre>	Thymus
GSM598		Value	e for GSM	598: 1	Idd5_T1;	<pre>src:</pre>	Thymus
GSM605		Value	e for GSM6	305: I	Idd5_T2;	<pre>src:</pre>	Thymus
GSM599	1	Value 1	for GSM599	9: Ido	d3+5_T1;	<pre>src:</pre>	Thymus
GSM606	,	Value 1	for GSM606	5: Ido	13+5_T2;	<pre>src:</pre>	Thymus

```
GSM600Value for GSM600: Idd9_T1; src: ThymusGSM607Value for GSM607: Idd9_T2; src: ThymusGSM601Value for GSM601: B10.H2g7_T1; src: ThymusGSM608Value for GSM608: B10.H2g7_T2; src: ThymusGSM602Value for GSM602: B10.H2g7 Idd3_T1; src: ThymusGSM609Value for GSM609: B10.H2g7 Idd3_T2; src: Thymus
```

#### 4.3 Converting GDS to an MAList

No annotation information (called platform information by GEO) was retrieved from because *ExpressionSet* does not contain slots for gene information, typically. However, it is easy to obtain this information. First, we need to know what platform this GDS used. Then, another call to getGEO will get us what we need.

```
> Meta(gds)$platform
```

```
[1] "GPL24"
```

```
> gpl <- getGEO("GPL5")
```

```
File stored at:
/tmp/RtmpnFTpz5/GPL5.soft
```

So, gpl now contains the information for GPL5 from GEO. Unlike *ExpressionSet*, the limma *MAList* does store gene annotation information, so we can use our newly created gpl of class *GPL* in a call to GDS2MA like so:

```
> MA <- GDS2MA(gds, GPL = gpl)
> MA
```

```
An object of class "MAList"
$M
```

	GSM582	GSM589	GSM583	GSM590	GSM584	GSM591	GSM585	GSM592	GSM586	GSM593
[1,]	101	54	111	55	87	30	99	43	105	56
[2,]	26	23	30	27	19	22	32	19	24	25
[3,]	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA
[4,]	233	162	252	178	214	144	238	147	250	166
[5,]	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA
	GSM587	GSM594	GSM588	GSM595	GSM596	GSM603	GSM597	GSM604	GSM598	GSM605
[1,]	GSM587 43	GSM594 14	GSM588 112	GSM595 43	GSM596 97	GSM603 36	GSM597 117	GSM604 40	GSM598 125	GSM605 45
[1,] [2,]	GSM587 43 14	GSM594 14 49	GSM588 112 32	GSM595 43 29	GSM596 97 31	GSM603 36 22	GSM597 117 26	GSM604 40 26	GSM598 125 35	GSM605 45 26
[1,] [2,] [3,]	GSM587 43 14 NA	GSM594 14 49 7	GSM588 112 32 NA	GSM595 43 29 4	GSM596 97 31 10	GSM603 36 22 22	GSM597 117 26 NA	GSM604 40 26 15	GSM598 125 35 NA	GSM605 45 26 23
[1,] [2,] [3,] [4,]	GSM587 43 14 NA 86	GSM594 14 49 7 22	GSM588 112 32 NA 236	GSM595 43 29 4 139	GSM596 97 31 10 216	GSM603 36 22 22 112	GSM597 117 26 NA 241	GSM604 40 26 15 130	GSM598 125 35 NA 270	GSM605 45 26 23 144

	GSM599	GSM606	GSM600	GSM607	GSM601	GSM60	8 GSM602	GSM6	609			
[1,]	99	1	109	38	87	1	8 72		16			
[2,]	18	13	25	32	28	4	0 14		41			
[3,]	NA	29	9	25	11	4	0 NA		22			
[4,]	239	148	211	139	208	1	6 174		15			
[5,]	NA	NA	NA	NA	NA	N	A NA		NA			
3910	9 more 1	rows	•									
\$A												
NULL												
\$tar	gets											
sa	mple tis	ssue sti	rain	disea	ase.sta	te						
1 GS	M582 spl	leen	NOD		diabet	ic						
2 GS	M589 spl	leen	NOD		diabet	ic						
3 GS	M583 spl	leen 1	Idd3 di	abetic-	resista	nt						
4 GS	M590 spl	leen 1	Idd3 di	abetic-	resista	nt						
5 GS	M584 spl	leen 1	Idd5 di	abetic-	resista	nt						
	-			des	criptio	n						
1 V	alue foi	r GSM582	2: NOD_	S1; src	: Splee	n						
2 V	alue foi	r GSM589	9: NOD_	S2; src	: Splee	n						
3 Va	lue for	GSM583	: Idd3_	S1; src	: Splee	n						
4 Va	lue for	GSM590	: Idd3_	S2; src	: Splee	n						
5 Va	lue for	GSM584	: Idd5_	S1; src	: Splee	n						
23 m	ore rows	5			-							
\$gen	es											
ID	GB_AC	CC BSCC	_ID	CLONE_II	SUB.A	RRAY D	UPLICATE	ROW	COLUMN	PCR_QC	SPOT_ID	)
1 1	AI94454	49 bs03g	g07 FBgi	n0033989	9	1	a	1	1	passed		
2 2	AI94469	95 bs040	c11 FBg	n003282:	1	1	a	1	2	passed		
3 3	AI94474	41 bs041	h01 FBg	n0034374	1	1	a	1	3	passed		
4 4	AI94480	01 bs051	f04 FBg	n003942:	1	1	a	1	4	failed		
55	AI94504	43 bs080	c11 FBg	n004537(	C	1	a	1	5	passed		
			0							1		
1												g
2												
3												
4 gi	4505995	5 ref NH	P_00269	7.1 PPPI	M1B  pr	otein <sup>.</sup>	phosphat	ase 1	.B (form	nerly 20	C), magn	lesium-dep
5		-			Ŧ					5	. 0	1
E_	VAL SPOT	r_QC										
1 2e	-08 44	4364										
2 <	NA> 16	6957										

3 <NA> 17896 4 1e-25 16363 5 <NA> 83502 39109 more rows ... \$notes [[1]] [1] "able\_begin" \$channel\_count [1] "1" \$description [1] "Examination of spleen and thymus of type 1 diabetes nonobese diabetic (NOD) mouse, \$feature\_count [1] "39114" \$order [1] "none" \$platform [1] "GPL24" \$platform\_organism [1] "Mus musculus" \$platform\_technology\_type [1] "in situ oligonucleotide" \$pubmed\_id [1] "11827943" \$reference\_series [1] "GSE11" \$sample\_count [1] "28" \$sample\_organism [1] "Mus musculus"

```
$sample_type
[1] "RNA"
$title
[1] "Type 1 diabetes gene expression profiling"
$type
[1] "gene expression array-based"
$update_date
[1] "Jul 15 2003"
$value_type
[1] "count"
```

Now, MA is of class *MAList* and contains not only the data, but the sample information and gene information associated with GDS1.

#### 4.4 Converting GSE to an ExpressionSet

First, make sure that using the method described above in the section "Getting GSE Series Matrix files as an ExpressionSet" for using GSE Series Matrix files is not sufficient for the task, as it is much faster and simpler. If it is not (i.e., other columns from each GSM are needed), then this method will be needed.

Converting a GSE object to an ExpressionSet object currently takes a bit of R data manipulation due to the varied data that can be stored in a GSE and the underlying GSM and GPL objects. However, using a simple example will hopefully be illustrative of the technique.

First, we need to make sure that all of the *GSMs* are from the same platform:

\$GSM5 [1] "GPL5"

\$GSM6 [1] "GPL5"

### \$GSM7

[1] "GPL5"

#### \$GSM8

[1] "GPL5"

#### \$GSM9

[1] "GPL5"

Indeed, they all used GPL5 as their platform (which we could have determined by looking at the GPLList for gse, which shows only one GPL for this particular GSE.). So, now we would like to know what column represents the data that we would like to extract. Looking at the first few rows of the Table of a single GSM will likely give us an idea (and by the way, GEO uses a convention that the column that contains the single "measurement" for each array is called the "VALUE" column, which we could use if we don't know what other column is most relevant).

#### > Table(GSMList(gse)[[1]])[1:5, ]

	ID_REF	SIGNAL_RAW	BKD_FORM	NORM_FORM	BKD_RAW	NORM_VALUE	CONST	VALUE
1	1	4486.49	0	0	3379.579	23337.54	39542	55845.45
2	2	3482.51	0	0	3379.579	23337.54	39542	41058.05
3	3	3812.39	0	0	3379.579	23337.54	39542	45916.78
4	4	3257.56	1	0	3379.579	23337.54	39542	37744.81
5	5	5436.91	0	0	3379.579	23337.54	39542	69843.97

> Columns(GSMList(gse)[[1]])[1:5, ]

Column Description 1 ID\_REF 2 SIGNAL\_RAW raw signal 3 BKD\_FORM 4 NORM\_FORM

5 BKD\_RAW raw background

We will indeed use the "VALUE" column. We then want to make a matrix of these values like so:

```
> probesets <- Table(GPLList(gse)[[1]])$ID
> data.matrix <- do.call("cbind", lapply(GSMList(gse), function(x) {
        tab <- Table(x)
        mymatch <- match(probesets, tab$ID_REF)
        return(tab$VALUE[mymatch])
+ }))
> data.matrix <- apply(data.matrix, 2, function(x) {
        as.numeric(as.character(x))
        + })
> data.matrix <- log2(data.matrix)
> data.matrix[1:5, ]
```

GSM10GSM3GSM4GSM5GSM6GSM7GSM8GSM9[1,]15.7691516.2292116.1300015.6503417.0921415.4585316.0947415.23515[2,]15.3253815.26597NaN15.2040616.4759614.8577615.1488514.89007[3,]15.4867315.8093214.1625915.1804816.2123515.0609415.3824214.96986[4,]15.2039915.5066313.4158215.0593916.1859314.7986114.8046015.01923[5,]16.0918517.8224618.3827016.2457016.6096415.9001116.0096215.88859

Note that we do a "match" to make sure that the values and the platform information are in the same order. Finally, to make the *ExpressionSet* object:

```
> require(Biobase)
> rownames(data.matrix) <- probesets</pre>
> colnames(data.matrix) <- names(GSMList(gse))</pre>
> pdata <- data.frame(samples = names(GSMList(gse)))</pre>
> rownames(pdata) <- names(GSMList(gse))</pre>
> pheno <- as(pdata, "AnnotatedDataFrame")</pre>
> eset2 <- new("ExpressionSet", exprs = data.matrix, phenoData = pheno)</pre>
> eset2
ExpressionSet (storageMode: lockedEnvironment)
assayData: 3455 features, 8 samples
  element names: exprs
phenoData
  sampleNames: GSM10, GSM3, ..., GSM9 (8 total)
  varLabels and varMetadata description:
    samples: NA
featureData
  featureNames: 1, 2, ..., 3455 (3455 total)
  fvarLabels and fvarMetadata description: none
experimentData: use 'experimentData(object)'
Annotation:
```

So, using a combination of lapply on the GSMList, one can extract as many columns of interest as necessary to build the data structure of choice. Because the GSM data from the GEO website are fully downloaded and included in the GSE object, one can extract foreground and background as well as quality for two-channel arrays, for example. Getting array annotation is also a bit more complicated, but by replacing "platform" in the lapply call to get platform information for each array, one can get other information associated with each array. Future work with this package will likely focus on better tools for manipulating GSE data.

### 5 Accessing Raw Data from GEO

NCBI GEO accepts (but has not always required) raw data such as .CEL files, .CDF files, images, etc. Sometimes, it is useful to get quick access to such data. A single function, getGEOSuppFiles, can take as an argument a GEO accession and will download all the raw data associate with that accession. By default, the function will create a directory in the current working directory to store the raw data for the chosen GEO accession. Combining a simple sapply statement or other loop structure with getGEOSuppFiles makes for a very simple way to get gobs of raw data quickly and easily without needing to know the specifics of GEO raw data URLs.

## 6 Conclusion

The GEOquery package provides a bridge to the vast array resources contained in the NCBI GEO repositories. By maintaining the full richness of the GEO data rather than focusing on getting only the "numbers", it is possible to integrate GEO data into current Bioconductor data structures and to perform analyses on that data quite quickly and easily. These tools will hopefully open GEO data more fully to the array community at large.

## 7 sessionInfo

- R version 2.8.0 (2008-10-20), x86\_64-unknown-linux-gnu
- Locale: LC\_CTYPE=en\_US;LC\_NUMERIC=C;LC\_TIME=en\_US;LC\_COLLATE=en\_US;LC\_MONETARY=C;LC\_M
- Base packages: base, datasets, graphics, grDevices, methods, stats, tools, utils
- Other packages: Biobase 2.2.0, GEOquery 2.6.0, limma 2.16.0, RCurl 0.91-0