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.¹

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).

¹See http://www.ncbi.nih.gov/geo for more information

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 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.

```
> # If you have network access, the more typical way to do this
```

```
> # would be to use this:
```

```
> # gds <- getGEO("GDS507")
```

```
> gds <- getGEO(filename=system.file("extdata/GDS507.soft.gz",package="GEOquery"))</pre>
```

Now, gds contains the R data structure (of class GDS) that represents the GDS507 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.

```
> # If you have network access, the more typical way to do this
> # would be to use this:
> # gds <- getGEO("GSM11805")
> gsm <- getGEO(filename=system.file("extdata/GSM11805.txt.gz",package="GEOquery"))</pre>
```

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:

```
> # Look at gsm metadata:
> Meta(gsm)
$channel_count
[1] "1"
$comment
[1] "Raw data provided as supplementary file"
$contact_address
[1] "715 Albany Street, E613B"
$contact_city
[1] "Boston"
$contact_country
[1] "USA"
$contact_department
[1] "Genetics and Genomics"
$contact_email
[1] "mlenburg@bu.edu"
$contact_fax
[1] "617-414-1646"
$contact_institute
[1] "Boston University School of Medicine"
$contact_name
[1] "Marc, E., Lenburg"
$contact_phone
[1] "617-414-1375"
$contact_state
```

```
[1] "MA"
$contact_web_link
[1] "http://gg.bu.edu"
$`contact_zip/postal_code`
[1] "02130"
$data_row_count
[1] "22283"
$description
[1] "Age = 70; Gender = Female; Right Kidney; Adjacent Tumor Type = clear cell; Adjacent
[2] "Keywords = kidney"
[3] "Keywords = renal"
[4] "Keywords = RCC"
[5] "Keywords = carcinoma"
[6] "Keywords = cancer"
[7] "Lot batch = 2004638"
$geo_accession
[1] "GSM11805"
$last_update_date
[1] "May 28 2005"
$molecule_ch1
[1] "total RNA"
$organism_ch1
[1] "Homo sapiens"
$platform_id
[1] "GPL96"
$series_id
[1] "GSE781"
$source_name_ch1
[1] "Trizol isolation of total RNA from normal tissue adjacent to Renal Cell Carcinoma"
$status
```

```
[1] "Public on Nov 25 2003"
$submission_date
[1] "Oct 20 2003"
$supplementary_file
[1] "ftp://ftp.ncbi.nih.gov/pub/geo/DATA/supplementary/samples/GSM11nnn/GSM11805/GSM1180
$title
[1] "NO35 Normal Human Kidney U133A"
$type
[1] "RNA"
> # Look at data associated with the GSM:
> # but restrict to only first 5 rows, for brevity
> Table(gsm)[1:5,]
          ID_REF VALUE ABS_CALL
1 AFFX-BioB-5_at 953.9
                               Ρ
2 AFFX-BioB-M_at 2982.8
                               Ρ
3 AFFX-BioB-3_at 1657.9
                               Ρ
4 AFFX-BioC-5_at 2652.7
                               Ρ
5 AFFX-BioC-3_at 2019.5
                               Ρ
> # Look at Column descriptions:
> Columns(gsm)
    Column
    ID_REF
1
2
    VALUE
3 ABS_CALL
                                                                   Description
1
                          MAS 5.0 Statistical Algorithm (mean scaled to 500)
2
3 MAS 5.0 Absent, Marginal, Present call with Alpha1 = 0.05, Alpha2 = 0.065
   The GPL behaves exactly as the GSM class. However, the GDS has a bit more informa-
```

tion associated with the Columns method:

> Columns(gds)

sample disease.state individual 1 GSM11815 RCC 035

2	GSM11832	RCC	023
3	GSM12069	RCC	001
4	GSM12083	RCC	005
5	GSM12101	RCC	011
6	GSM12106	RCC	032
7	GSM12274	RCC	2
8	GSM12299	RCC	3
9	GSM12412	RCC	4
10	GSM11810	normal	035
11	GSM11827	normal	023
12	GSM12078	normal	001
13	GSM12099	normal	005
14	GSM12269	normal	1
15	GSM12287	normal	2
16	GSM12301	normal	3
17	GSM12448	normal	4

Value for GSM11815: C035 Renal Clear Cell Carcinoma U133B; src: Trizol iso 1 2 Value for GSM11832: C023 Renal Clear Cell Carcinoma U133B; src: Trizol iso Value for GSM12069: C001 Renal Clear Cell Carcinoma U133B; src: Trizol iso З 4 Value for GSM12083: C005 Renal Clear Cell Carcinoma U133B; src: Trizol iso Value for GSM12101: C011 Renal Clear Cell Carcinoma U133B; src: Trizol iso 5 6 Value for GSM12106: C032 Renal Clear Cell Carcinoma U133B; src: Trizol iso 7 Value for GSM12274: C2 Renal Clear Cell Carcinoma U133B; src: Trizol iso 8 Value for GSM12299: C3 Renal Clear Cell Carcinoma U133B; src: Trizol iso 9 Value for GSM12412: C4 Renal Clear Cell Carcinoma U133B; src: Trizol iso 10 Value for GSM11810: NO35 Normal Human Kidney U133B; src: Trizol isolation of tot 11 Value for GSM11827: NO23 Normal Human Kidney U133B; src: Trizol isolation of tot 12 Value for GSM12078: NO01 Normal Human Kidney U133B; src: Trizol isolation of tot 13 Value for GSM12099: NO05 Normal Human Kidney U133B; src: Trizol isolation of tot 14 Value for GSM12269: N1 Normal Human Kidney U133B; src: Trizol isolation of tot 15 Value for GSM12287: N2 Renal Clear Cell Carcinoma U133B; src: Trizol isolation of tot 16 Value for GSM12301: N3 Renal Clear Cell Carcinoma U133B; src: Trizol isolation of tot 17 Value for GSM12448: N4 Renal Clear Cell Carcinoma U133B; src: Trizol isolation of tot

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:

```
> # Again, with good network access, one would do:
> # gse <- getGEO("GSE781",GSEMatrix=FALSE)</pre>
> gse <- getGEO(filename=system.file("extdata/GSE781_family.soft.gz",package="GEOquery</pre>
Parsing....
> Meta(gse)
$contact_address
[1] "715 Albany Street, E613B"
$contact_city
[1] "Boston"
$contact_country
[1] "USA"
$contact_department
[1] "Genetics and Genomics"
$contact_email
[1] "mlenburg@bu.edu"
$contact_fax
[1] "617-414-1646"
$contact_institute
[1] "Boston University School of Medicine"
$contact_name
[1] "Marc, E., Lenburg"
$contact_phone
[1] "617-414-1375"
$contact_state
[1] "MA"
$contact_web_link
[1] "http://gg.bu.edu"
$`contact_zip/postal_code`
[1] "02130"
```

\$contributor [1] "Marc, E, Lenburg" "Louis,S,Liou" "Norman, P, Gerry" [4] "Garrett,M,Frampton" "Herbert, T, Cohen" "Michael, F, Christman" \$email [1] "geo@ncbi.nlm.nih.gov" \$geo_accession [1] "GSE781" \$institute [1] "NCBI NLM NIH" \$last_update_date [1] "May 29 2005" \$name [1] "Gene Expression Omnibus (GEO)" \$platform_id [1] "GPL96" "GPL97" \$pubmed_id [1] "14641932" \$sample_id [1] "GSM11805" "GSM11810" "GSM11814" "GSM11815" "GSM11823" "GSM11827" [7] "GSM11830" "GSM11832" "GSM12067" "GSM12069" "GSM12075" "GSM12078" [13] "GSM12079" "GSM12083" "GSM12098" "GSM12099" "GSM12100" "GSM12101" [19] "GSM12105" "GSM12106" "GSM12268" "GSM12269" "GSM12270" "GSM12274" [25] "GSM12283" "GSM12287" "GSM12298" "GSM12299" "GSM12300" "GSM12301" [31] "GSM12399" "GSM12412" "GSM12444" "GSM12448" \$status [1] "Public on Nov 25 2003" \$submission_date [1] "Oct 24 2003" \$summary

[1] "Each total RNA sample is hybridized to two different arrays: Affymetrix U133A (GP

```
[2] ""
 [3] "For most of the normal tissue samples there is a renal clear cell carcinoma sample
 [4] ""
 [5] "For most of the renal clear cell carcinoma samples there is a corresponding adjace
 [6] "Keywords = kidney"
 [7] "Keywords = renal"
 [8] "Keywords = RCC"
 [9] "Keywords = carcinoma"
[10] "Keywords = cancer"
[11] "Keywords: parallel sample"
$supplementary_file
[1] "ftp://ftp.ncbi.nih.gov/pub/geo/DATA/supplementary/series/GSE781/GSE781_RAW.tar"
$title
[1] "Normal and Renal Cell Carcinoma Kidney Tissue, Human"
$type
[1] "Expression profiling by array"
$web_link
[1] "http://www.ncbi.nlm.nih.gov/projects/geo"
> # names of all the GSM objects contained in the GSE
> names(GSMList(gse))
 [1] "GSM11805" "GSM11810" "GSM11814" "GSM11815" "GSM11823" "GSM11827"
 [7] "GSM11830" "GSM11832" "GSM12067" "GSM12069" "GSM12075" "GSM12078"
[13] "GSM12079" "GSM12083" "GSM12098" "GSM12099" "GSM12100" "GSM12101"
[19] "GSM12105" "GSM12106" "GSM12268" "GSM12269" "GSM12270" "GSM12274"
[25] "GSM12283" "GSM12287" "GSM12298" "GSM12299" "GSM12300" "GSM12301"
[31] "GSM12399" "GSM12412" "GSM12444" "GSM12448"
> # and get the first GSM object on the list
> GSMList(gse)[[1]]
An object of class "GSM"
channel_count
[1] "1"
comment
[1] "Raw data provided as supplementary file"
contact_address
[1] "715 Albany Street, E613B"
```

```
contact_city
[1] "Boston"
contact_country
[1] "USA"
contact_department
[1] "Genetics and Genomics"
contact_email
[1] "mlenburg@bu.edu"
contact_fax
[1] "617-414-1646"
contact_institute
[1] "Boston University School of Medicine"
contact_name
[1] "Marc, E., Lenburg"
contact_phone
[1] "617-414-1375"
contact_state
[1] "MA"
contact_web_link
[1] "http://gg.bu.edu"
contact_zip/postal_code
[1] "02130"
data_row_count
[1] "22283"
description
[1] "Age = 70; Gender = Female; Right Kidney; Adjacent Tumor Type = clear cell; Adjacent
[2] "Keywords = kidney"
[3] "Keywords = renal"
[4] "Keywords = RCC"
[5] "Keywords = carcinoma"
[6] "Keywords = cancer"
[7] "Lot batch = 2004638"
geo_accession
[1] "GSM11805"
last_update_date
[1] "May 28 2005"
molecule_ch1
[1] "total RNA"
organism_ch1
[1] "Homo sapiens"
platform_id
[1] "GPL96"
```

```
series_id
[1] "GSE781"
source_name_ch1
[1] "Trizol isolation of total RNA from normal tissue adjacent to Renal Cell Carcinoma"
status
[1] "Public on Nov 25 2003"
submission_date
[1] "Oct 20 2003"
supplementary_file
[1] "ftp://ftp.ncbi.nih.gov/pub/geo/DATA/supplementary/samples/GSM11nnn/GSM11805/GSM1180
title
[1] "N035 Normal Human Kidney U133A"
type
[1] "RNA"
An object of class "GEODataTable"
***** Column Descriptions *****
    Column
1
    ID_REF
2
    VALUE
3 ABS_CALL
                                                                  Description
1
2
                          MAS 5.0 Statistical Algorithm (mean scaled to 500)
3 MAS 5.0 Absent, Marginal, Present call with Alpha1 = 0.05, Alpha2 = 0.065
***** Data Table *****
          ID_REF VALUE ABS_CALL
1 AFFX-BioB-5_at 953.9
                               Ρ
2 AFFX-BioB-M_at 2982.8
                               Ρ
                               Ρ
3 AFFX-BioB-3_at 1657.9
4 AFFX-BioC-5_at 2652.7
                               Ρ
5 AFFX-BioC-3_at 2019.5
                               Ρ
22278 more rows ...
> # and the names of the GPLs represented
> names(GPLList(gse))
```

[1] "GPL96" "GPL97"

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.

```
> # Note that GSEMatrix=TRUE is the default
> gse2553 <- getGEO('GSE2553',GSEMatrix=TRUE)</pre>
> show(gse2553)
$GSE2553_series_matrix.txt.gz
ExpressionSet (storageMode: lockedEnvironment)
assayData: 12600 features, 181 samples
  element names: exprs
protocolData: none
phenoData
 sampleNames: GSM48681 GSM48682 ... GSM48861 (181 total)
 varLabels: title geo_accession ... data_row_count (30 total)
 varMetadata: labelDescription
featureData
 featureNames: 1 2 ... 12600 (12600 total)
 fvarLabels: ID PenAt ... Chimeric_Cluster_IDs (13 total)
 fvarMetadata: Column Description labelDescription
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
                                                                         R.NA
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>
```

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
```

GSM12106 GSM12106

GSM12274 GSM12274

```
ExpressionSet (storageMode: lockedEnvironment)
assayData: 22645 features, 17 samples
  element names: exprs
protocolData: none
phenoData
  sampleNames: GSM11815 GSM11832 ... GSM12448 (17 total)
  varLabels: sample disease.state individual description
  varMetadata: labelDescription
featureData
  featureNames: 200000_s_at 200001_at ... AFFX-TrpnX-M_at (22645 total)
  fvarLabels: ID Gene title ... GO:Component ID (21 total)
  fvarMetadata: Column labelDescription
experimentData: use 'experimentData(object)'
  pubMedIds: 14641932
Annotation:
> pData(eset)
           sample disease.state individual
                            RCC
GSM11815 GSM11815
                                        035
GSM11832 GSM11832
                            RCC
                                        023
GSM12069 GSM12069
                            RCC
                                        001
GSM12083 GSM12083
                            RCC
                                        005
GSM12101 GSM12101
                            RCC
                                        011
```

RCC

RCC

032

2

GSM12299	GSM12299	RCC	3
GSM12412	GSM12412	RCC	4
GSM11810	GSM11810	normal	035
GSM11827	GSM11827	normal	023
GSM12078	GSM12078	normal	001
GSM12099	GSM12099	normal	005
GSM12269	GSM12269	normal	1
GSM12287	GSM12287	normal	2
GSM12301	GSM12301	normal	3
GSM12448	GSM12448	normal	4

Value for GSM11815: CO35 Renal Clear Cell Carcinoma U133B; src: Triz GSM11815 GSM11832 Value for GSM11832: C023 Renal Clear Cell Carcinoma U133B; src: Triz GSM12069 Value for GSM12069: C001 Renal Clear Cell Carcinoma U133B; src: Triz GSM12083 Value for GSM12083: C005 Renal Clear Cell Carcinoma U133B; src: Triz Value for GSM12101: C011 Renal Clear Cell Carcinoma U133B; src: Triz GSM12101 GSM12106 Value for GSM12106: CO32 Renal Clear Cell Carcinoma U133B; src: Triz GSM12274 Value for GSM12274: C2 Renal Clear Cell Carcinoma U133B; src: Triz GSM12299 Value for GSM12299: C3 Renal Clear Cell Carcinoma U133B; src: Triz GSM12412 Value for GSM12412: C4 Renal Clear Cell Carcinoma U133B; src: Triz GSM11810 Value for GSM11810: NO35 Normal Human Kidney U133B; src: Trizol isolation GSM11827 Value for GSM11827: NO23 Normal Human Kidney U133B; src: Trizol isolation GSM12078 Value for GSM12078: NO01 Normal Human Kidney U133B; src: Trizol isolation GSM12099 Value for GSM12099: N005 Normal Human Kidney U133B; src: Trizol isolation GSM12269 Value for GSM12269: N1 Normal Human Kidney U133B; src: Trizol isolation GSM12287 Value for GSM12287: N2 Renal Clear Cell Carcinoma U133B; src: Trizol isolation GSM12301 Value for GSM12301: N3 Renal Clear Cell Carcinoma U133B; src: Trizol isolation GSM12448 Value for GSM12448: N4 Renal Clear Cell Carcinoma U133B; src: Trizol isolation

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.

```
> #get the platform from the GDS metadata
> Meta(gds)$platform
```

```
[1] "GPL97"
```

```
> #So use this information in a call to getGEO
> gpl <- getGEO(filename=system.file("extdata/GPL97.annot.gz",package="GEOquery"))</pre>
```

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)</pre>
> MA
An object of class "MAList"
$M
     GSM11815 GSM11832 GSM12069 GSM12083 GSM12101 GSM12106 GSM12274 GSM12299
[1,]
       4254.0
                 5298.2
                           4026.5
                                     3498.4
                                               3566.4
                                                         4903.1
                                                                   6372.6
                                                                            4829.1
[2,]
      17996.2
                12010.7
                          10283.5
                                     2534.7
                                                        13354.0
                                                                   8563.8
                                                                           17247.6
                                              11048.4
[3,]
      41678.8
                39116.9
                          38758.9
                                    32847.7
                                              39633.9
                                                        43511.2
                                                                 46856.7
                                                                           47032.4
[4,]
      65390.9
                34806.2
                          31257.2
                                    28308.5
                                             67447.5
                                                        56989.9
                                                                 57972.5
                                                                           57570.5
[5,]
      19030.1
                15813.6
                          16355.7
                                     9579.7
                                              14273.5
                                                        17217.0
                                                                 19116.9
                                                                           17487.6
     GSM12412 GSM11810 GSM11827 GSM12078 GSM12099 GSM12269 GSM12287 GSM12301
                                     3729.9
[1,]
       5205.8
                 2756.8
                           3932.0
                                               3223.4
                                                         3640.5
                                                                   4886.3
                                                                            4070.2
[2,]
      16018.5
                 6077.0
                          15703.8
                                    10138.5
                                             11614.4
                                                         8460.5
                                                                 10282.6
                                                                           11844.3
[3,]
      22152.2
                26660.7
                          26373.6
                                    23809.6
                                              24749.3
                                                                           22733.7
                                                        21936.8
                                                                 31462.8
[4,]
      29062.2
                35140.9
                          23629.3
                                    22100.5
                                              21651.0
                                                        18550.7
                                                                 23496.5
                                                                           21315.4
[5,]
      14671.6
                17733.1
                          18022.4
                                    17957.4
                                              15958.0
                                                        15799.8
                                                                 16685.8
                                                                           18817.3
     GSM12448
[1,]
       3482.1
[2,]
       9741.6
[3,]
      25395.5
[4,]
      28631.4
[5,]
      17421.1
22640 more rows ...
$A
NULL
$targets
    sample disease.state individual
1 GSM11815
                      RCC
                                   035
2 GSM11832
                       RCC
                                   023
3 GSM12069
                      RCC
                                   001
4 GSM12083
                       RCC
                                   005
5 GSM12101
                      RCC
                                   011
1 Value for GSM11815: C035 Renal Clear Cell Carcinoma U133B; src: Trizol isolation of to
```

2 Value for GSM11815: C035 Renal Clear Cell Carcinoma U133B; src: Irizol isolation of to 2 Value for GSM11832: C023 Renal Clear Cell Carcinoma U133B; src: Trizol isolation of to 3 Value for GSM12069: C001 Renal Clear Cell Carcinoma U133B; src: Trizol isolation of to 4 Value for GSM12083: C005 Renal Clear Cell Carcinoma U133B; src: Trizol isolation of to 5 Value for GSM12101: C011 Renal Clear Cell Carcinoma U133B; src: Trizol isolation of to 12 more rows ...

\$genes

ID Gene title 1 200000_s_at PRP8 pre-mRNA processing factor 8 homolog (S. cerevisiae) 2 200001_at calpain, small subunit 1 200002_at 3 ribosomal protein L35 4 200003_s_at ribosomal protein L28 5 200004_at eukaryotic translation initiation factor 4 gamma, 2 Gene symbol Gene ID UniGene title UniGene symbol UniGene ID 1 PRPF8 10594 2 CAPNS1 826 З RPL35 11224 4 RPL28 6158 5 EIF4G2 1982 1 Homo sapiens PRP8 pre-mRNA processing factor 8 homolog (S. cerevisiae 2 Homo sapiens calpain, small subunit 1 (CAPNS1), transcript 3 Homo sapiens ribosomal protein L3 4 Homo sapiens ribosomal protein L2 5 Homo sapiens eukaryotic translation initiation factor 4 gamma, 2 (EIF4G2), transcript GI GenBank Accession Platform_CLONEID Platform_ORF Platform_SPOTID 1 91208425 NM_006445 <NA> <NA> <NA> 2 51599152 NM_001749 <NA> <NA> <NA> 3 78190471 NM_007209 <NA> <NA> <NA> 4 34486095 NM_000991 <NA> <NA> <NA> 5 111494227 NM_001418 <NA> <NA> <NA> Chromosome location 1 17p13.3 2 19q13.12 3 9q34.1 4 19q13.4 5 11p15 Chromosome annotation 1 Chromosome 17, NC_000017.9 (1500673..1534926, complement) 2 Chromosome 19, NC_000019.8 (41322758..41333095) 3 Chromosome 9, NC_000009.10 (126659979..126664061, complement) 4 Chromosome 19, NC_000019.8 (60589112..60595265) 5 Chromosome 11, NC_000011.8 (10775169..10787158, complement)

RNA binding///RNA splicing factor activity, transesterificatio 1 2 calcium ion binding///calcium-dependent cysteine-type endopeptida 3 mRNA binding///protein binding///struc RNA binding///protein binding///structural constituent of ribosome///struc 4 5 protein binding///protein binding///translation initiation factor activity///translati 1 RNA splicing///nuclear mRNA splicing, via spliceosome///nuclear mRNA splicing, via spl 2 З 4 5 RNA metabolic process///cell cycle arrest/// GO:Component nuclear speck///nucleus///snRNP U5///spliceosome 1 cytoplasm///plasma membrane 2 3 cytosol///cytosolic large ribosomal subunit///intracellular///nucleolus///ribosome cytosol///cytosolic large ribosomal subunit///intracellular///ribosome 4 eukaryotic translation initiation factor 4F complex 5 GO:Function GD:0003723///GD:0031202///GD:0005515 1 GD:0005509///GD:0004198///GD:0005515 2 GD:0003729///GD:0005515///GD:0003735 3 4 GD:0003723///GD:0005515///GD:0003735///GD:0003735 5 GD:0005515///GD:0005515///GD:0003743///GD:0003743 GO:Process 1 GD:0008380///GD:0000398///GD:0000398///GD:0050896///GD:0007601 2 GD:0008284 3 GD:0006414 GD:0006412///GD:0006414 4 GD:0016070///GD:0007050///GD:0008219///GD:0006446 5 GO:Component GD:0016607///GD:0005634///GD:0005682///GD:0005681 1 GD:0005737///GD:0005886 2 3 GD:0005829///GD:0022625///GD:0005622///GD:0005730///GD:0005840 GD:0005829///GD:0022625///GD:0005622///GD:0005840 4 5 GD:0016281 22640 more rows ... \$notes \$channel_count [1] "1" \$dataset_id

[1] "GDS507" "GDS507" "GDS507" "GDS507" "GDS507" "GDS507" "GDS507"

[9] "GDS507" "GDS507" "GDS507"

\$description

[1] "Investigation into mechanisms of renal clear cell carcinogenesis (RCC). Comparison

- [2] "RCC"
- [3] "normal"
- [4] "035"
- [5] "023"
- [6] "001"
- [7] "005"
- [8] "011"
- [9] "032"
- [10] "1"
- [11] "2"
- [12] "3"
- [13] "4"

```
$email
```

[1] "geo@ncbi.nlm.nih.gov"

\$feature_count [1] "22645"

\$institute

[1] "NCBI NLM NIH"

\$name

[1] "Gene Expression Omnibus (GEO)"

\$order

[1] "none"

\$platform
[1] "GPL97"

\$platform_organism
[1] "Homo sapiens"

```
$platform_technology_type
[1] "in situ oligonucleotide"
```

\$pubmed_id [1] "14641932" \$ref [1] "Nucleic Acids Res. 2005 Jan 1;33 Database Issue:D562-6" \$reference_series [1] "GSE781" \$sample_count [1] "17" \$sample_id [1] "GSM11815,GSM11832,GSM12069,GSM12083,GSM12101,GSM12106,GSM12274,GSM12299,GSM12412" [2] "GSM11810, GSM11827, GSM12078, GSM12099, GSM12269, GSM12287, GSM12301, GSM12448" [3] "GSM11810,GSM11815" [4] "GSM11827,GSM11832" [5] "GSM12069,GSM12078" [6] "GSM12083,GSM12099" [7] "GSM12101" [8] "GSM12106" [9] "GSM12269" [10] "GSM12274,GSM12287" [11] "GSM12299,GSM12301" [12] "GSM12412,GSM12448" \$sample_organism [1] "Homo sapiens" \$sample_type [1] "RNA" \$title [1] "Renal clear cell carcinoma (HG-U133B)" \$type [1] "gene expression array-based" "disease state" [3] "disease state" "individual" [5] "individual" "individual" [7] "individual" "individual" [9] "individual" "individual"

```
[11] "individual" "individual"
```

```
[13] "individual"
$update_date
[1] "Mar 04 2004"
$value_type
[1] "count"
$web_link
```

[1] "http://www.ncbi.nlm.nih.gov/projects/geo"

Now, MA is of class *MAList* and contains not only the data, but the sample information and gene information associated with GDS507.

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 GSMand 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:

```
> gsmplatforms <- lapply(GSMList(gse),function(x) {Meta(x)$platform})
> gsmplatforms
$GSM11805
[1] "GPL96"
$GSM11814
[1] "GPL96"
$GSM11815
[1] "GPL97"
$GSM11823
[1] "GPL96"
```

\$GSM12105 [1] "GPL96"

\$GSM12101 [1] "GPL97"

\$GSM12100 [1] "GPL96"

\$GSM12099 [1] "GPL97"

\$GSM12098 [1] "GPL96"

\$GSM12083 [1] "GPL97"

\$GSM12079 [1] "GPL96"

\$GSM12078 [1] "GPL97"

\$GSM12075 [1] "GPL96"

\$GSM12069 [1] "GPL97"

\$GSM12067 [1] "GPL96"

\$GSM11832 [1] "GPL97"

\$GSM11830 [1] "GPL96"

\$GSM11827 [1] "GPL97"

\$GSM12444 [1] "GPL96"

\$GSM12412 [1] "GPL97"

\$GSM12399 [1] "GPL96"

\$GSM12301 [1] "GPL97"

\$GSM12300 [1] "GPL96"

\$GSM12299 [1] "GPL97"

\$GSM12298 [1] "GPL96"

\$GSM12287 [1] "GPL97"

\$GSM12283 [1] "GPL96"

\$GSM12274 [1] "GPL97"

\$GSM12270 [1] "GPL96"

\$GSM12269 [1] "GPL97"

\$GSM12268 [1] "GPL96"

\$GSM12106 [1] "GPL97" \$GSM12448 [1] "GPL97"

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
                  VALUE ABS_CALL
1 AFFX-BioB-5_at
                 953.9
                                Ρ
2 AFFX-BioB-M_at 2982.8
                                Ρ
3 AFFX-BioB-3_at 1657.9
                                Ρ
4 AFFX-BioC-5_at 2652.7
                                Ρ
5 AFFX-BioC-3_at 2019.5
                                Ρ
> # and get the column descriptions
> Columns(GSMList(gse)[[1]])[1:5,]
       Column
1
       ID_REF
2
        VALUE
З
     ABS_CALL
NA
         <NA>
NA.1
         <NA>
                                                                       Description
1
2
                              MAS 5.0 Statistical Algorithm (mean scaled to 500)
3
     MAS 5.0 Absent, Marginal, Present call with Alpha1 = 0.05, Alpha2 = 0.065
NA
                                                                              <NA>
NA.1
                                                                              <NA>
```

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

```
> # get the probeset ordering
> probesets <- Table(GPLList(gse)[[1]])$ID
> # make the data matrix from the VALUE columns from each GSM
> # being careful to match the order of the probesets in the platform
```

```
> # with those in the GSMs
> 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,]
```

	GSM11805	GSM11810	GSM11814 GSM1181	GSM11823	GSM11827	GSM11830
[1,]	10.926963	NA	11.105254 NA	11.275019	NA	11.438636
[2,]	5.749534	NA	7.908092 NA	7.093814	NA	7.514122
[3,]	7.066089	NA	7.750205 NA 12.479755 NA	7.244126	NA	7.962896
[4,]	12.660353	NA	12.479755 NA	12.215897	NA	11.458355
[5,]	6.195741	NA	6.061776 NA	6.565293	NA	6.583459
			GSM12069 GSM12075			
[1,]	NA 1	11.424376	NA 11.222799 NA 6.407693	5 NA	11.469845	NA
[2,]	NA	7.901470	NA 6.407693	B NA	5.165912	NA
[3,]	NA	7.337176	NA 6.569850	S NA	7.477354	NA
[4,]	NA 1	11.397568	NA 12.529870) NA	12.240046	NA
[5,]	NA	6.877744	NA 6.652486	S NA	3.981853	NA
	GSM12098	GSM12099	GSM12100 GSM1210:	GSM12105	GSM12106	GSM12268
			10.835971 NA			
[2,]	6.556123	NA	8.207014 NA	6.816344	NA	6.563768
[3,]	7.708739	NA	7.428779 NA	7.754888	NA	7.126188
[4,]	12.336534	NA	11.762839 NA	11.237509	NA	12.412490
[5,]	5.501439	NA	6.247928 NA	6.017922	NA	6.525129
	GSM12269	GSM12270	GSM12274 GSM12283	GSM12287	GSM12298	GSM12299
			NA 11.181028			
[2,]	NA	7.353147	NA 5.770829) NA	6.912889	NA
[3,]	NA	8.742815	NA 7.339850) NA	7.602142	NA
			NA 12.678380			
[5,]	NA	6.683696	NA 5.918863	B NA	5.837943	NA
	GSM12300	GSM12301	GSM12399 GSM12412	GSM12444	GSM12448	
[1,]	11.078151	NA	11.535178 NA	11.105450	NA	
[2,]	4.812498	NA	7.471675 NA	7.488644	NA	
[3,]	7.383704	NA	7.471675 NA 7.432959 NA	7.381110	NA	
[4,]	12.090939	NA	11.421802 NA 5.419539 NA	12.172834	NA	
[5,]	6.281698	NA	5.419539 NA	5.469235	NA	

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)
> # go through the necessary steps to make a compliant ExpressionSet
> 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)
> eset2
ExpressionSet (storageMode: lockedEnvironment)
assayData: 22283 features, 34 samples
  element names: exprs
protocolData: none
phenoData
  sampleNames: GSM11805 GSM11810 ... GSM12448 (34 total)
  varLabels: samples
  varMetadata: labelDescription
featureData: 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.

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 Use Cases

GEOquery can be quite powerful for gathering a lot of data quickly. A few examples can be useful to show how this might be done for data mining purposes.

6.1 Getting all Series Records for a Given Platform

For data mining purposes, it is sometimes useful to be able to pull all the GSE records for a given platform. GEOquery makes this very easy, but a little bit of knowledge of the GPL record is necessary to get started. The GPL record contains both the GSE and GSM accessions that reference it. Some code is useful to illustrate the point:

```
> gp197 <- getGEO('GPL97')
> Meta(gp197)$title
[1] "[HG-U133B] Affymetrix Human Genome U133B Array"
> head(Meta(gp197)$series_id)
[1] "GSE362" "GSE473" "GSE620" "GSE674" "GSE781" "GSE907"
> length(Meta(gp197)$series_id)
[1] 141
> head(Meta(gp197)$sample_id)
[1] "GSM3922" "GSM3924" "GSM3926" "GSM3928" "GSM3930" "GSM3932"
> length(Meta(gp197)$sample_id)
```

[1] 6081

The code above loads the GPL97 record into R. The Meta method extracts a list of header information from the GPL record. The "title" gives the human name of the platform. The "series_id" gives a vector of series ids. Note that there are more than 120 series associated with this platform and more than 5100 samples. Code like the following could be used to download all the samples or series. I show only the first 5 samples as an example:

```
> gsmids <- Meta(gp197)$sample_id
> gsmlist <- sapply(gsmids[1:5],getGEO)
> names(gsmlist)
[1] "GSM3922" "GSM3924" "GSM3926" "GSM3928" "GSM3930"
```

7 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.

8 sessionInfo

- R version 3.0.2 (2013-09-25), x86_64-unknown-linux-gnu
- Locale: LC_CTYPE=en_US.UTF-8, LC_NUMERIC=C, LC_TIME=en_US.UTF-8, LC_COLLATE=C, LC_MONETARY=en_US.UTF-8, LC_MESSAGES=en_US.UTF-8, LC_PAPER=en_US.UTF-8, LC_NAME=C, LC_ADDRESS=C, LC_TELEPHONE=C, LC_MEASUREMENT=en_US.UTF-8, LC_IDENTIFICATION=C
- Base packages: base, datasets, grDevices, graphics, methods, parallel, stats, utils
- Other packages: Biobase 2.22.0, BiocGenerics 0.8.0, GEOquery 2.28.0, limma 3.18.0
- Loaded via a namespace (and not attached): RCurl 1.95-4.1, XML 3.98-1.1, tools 3.0.2