## Package 'GEOquery'

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Type Package Title Get data from NCBI Gene Expression Omnibus (GEO) Version 2.36.0 Date 2015-09-27 Author Sean Davis <sdavis2@mail.nih.gov> Maintainer Sean Davis <sdavis2@mail.nih.gov> BugReports https://github.com/seandavi/GEOquery/issues/new

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Suggests limma, knitr, rmarkdown, RUnit, BiocGenerics

VignetteBuilder knitr

URL https://github.com/seandavi/GEOquery

biocViews Microarray, DataImport, OneChannel, TwoChannel, SAGE

**Description** The NCBI Gene Expression Omnibus (GEO) is a public repository of microarray data. Given the rich and varied nature of this resource, it is only natural to want to apply Bio-Conductor tools to these data. GEOquery is the bridge between GEO and BioConductor.

License GPL-2

NeedsCompilation no

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```
Converting
```

Convert a GDS data structure to a BioConductor data structure

## Description

Functions to take a GDS data structure from getGEO and coerce it to limma MALists or ExpressionSets.

## Usage

```
GDS2MA(GDS,do.log2=FALSE,GPL=NULL,AnnotGPL=TRUE,getGPL=TRUE)
GDS2eSet(GDS,do.log2=FALSE,GPL=NULL,AnnotGPL=TRUE,getGPL=TRUE)
```

## Arguments

GDS	The GDS datastructure returned by getGEO
do.log2	Boolean, should the data in the GDS be log2 transformed before inserting into the new data structure
GPL	Either a GPL data structure (from a call to getGEO) or NULL. If NULL, this will cause a call to getGEO to produce a GPL. The gene information from the GPL is then used to construct the genes slot of the resulting limma MAList object or the featureData slot of the ExpressionSet instance.
AnnotGPL	In general, the annotation GPL files will be available for GDS records, so the default is to use these files over the user-submitted GPL files
getGPL	A boolean defaulting to TRUE as to whether or not to download and include GPL information when converting to ExpressionSet or MAList. You may want to set this to FALSE if you know that you are going to annotate your featureData using Bioconductor tools rather than relying on information provided through NCBI GEO. Download times can also be greatly reduced by specifying FALSE.

## Details

This function just rearranges one data structure into another. For GDS, it also deals appropriately with making the "targets" list item for the limma data structure and the phenoData slot of ExpressionSets.

#### Value

GDS2MA	A limma MAList
GDS2eSet	An ExpressionSet object

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#### GDS-class

#### Author(s)

Sean Davis

## References

See the limma and ExpressionSet help in the appropriate packages

## Examples

```
## Not run: gds505 <- getGEO('GDS505')
## Not run: MA <- GDS2MA(gds505)
## Not run: eset <- GDS2eSet(gds505)</pre>
```

GDS-class

Class "GDS"

## Description

A class describing a GEO GDS entity

### **Objects from the Class**

Objects can be created by calls of the form new("GDS", ...)

#### Slots

gpl: Object of class "GPL"

dataTable: Object of class "GEODataTable" containing the data table for the GDS

header: Object of class "list" containing the metadata for the GDS; can be accessed via the Meta accessor

## Extends

Class "GEOData", directly.

## Methods

No methods defined with class "GDS" in the signature, but methods applying to GEOData are also applicable here.

## Author(s)

Sean Davis

#### See Also

GEOData-class

Generic functions Generic functions for GEOquery

#### Description

The main documentation is in the Class documentation

#### Author(s)

Sean Davis

#### See Also

GEOData-class

GEOData-cla	ss Class	"GEOData"

#### Description

A virtual class for holding GEO samples, platforms, and datasets

#### **Objects from the Class**

Objects can be created by calls of the form new("GEOData", ...).

## Slots

header: Object of class "list" containing metadata

#### Methods

```
Accession signature(object = "GEOData"): returns the GEO acccession for the current object
Columns signature(object = "GEOData"): returns the column descriptions for the current
object
Meta signature(object = "GEOData"): returns the metadata for the current object
Table signature(object = "GEOData"): returns the "Table" for the current object
dataTable signature(object = "GEOData"): returns the dataTable (column info and data) for
the current object
show signature(object = "GEOData"): a convenience method for showing a GEO object
```

#### Author(s)

Sean Davis

#### GEODataTable-class

#### See Also

GDS-class, GPL-class, GSM-class, GEODataTable-class,

GEODataTable-class Class "GEODataTable"

#### Description

Contains the column descriptions and data for the datatable part of a GEO object

#### **Objects from the Class**

Objects can be created by calls of the form new("GEODataTable", ...).

### Slots

columns: Object of class "data.frame" containing information about the columns in the datatable table: Object of class "data.frame" containing the actual data

#### Methods

**Columns** signature(object = "GEODataTable"): get the column portion of the GEODataTable **Table** signature(object = "GEODataTable"): get the table portion of the GEODataTable **show** signature(object = "GEODataTable"): convenience show method

## Author(s)

Sean Davis

getGEO

Get a GEO object from NCBI or file

#### Description

This function is the main user-level function in the GEOquery package. It directs the download (if no filename is specified) and parsing of a GEO SOFT format file into an R data structure specifically designed to make access to each of the important parts of the GEO SOFT format easily accessible.

#### Usage

```
getGEO(GEO = NULL, filename = NULL, destdir = tempdir(), GSElimits=NULL,
GSEMatrix=TRUE,AnnotGPL=FALSE,getGPL=TRUE)
```

#### Arguments

GEO	A character string representing a GEO object for download and parsing. (eg., 'GDS505','GSE2','GSM2','GPL96')
filename	The filename of a previously downloaded GEO SOFT format file or its gzipped representation (in which case the filename must end in .gz). Either one of GEO or filename may be specified, not both. GEO series matrix files are also handled. Note that since a single file is being parsed, the return value is not a list of esets, but a single eset when GSE matrix files are parsed.
destdir	The destination directory for any downloads. Defaults to the architecture-dependent tempdir. You may want to specify a different directory if you want to save the file for later use. Doing so is a good idea if you have a slow connection, as some of the GEO files are HUGE!
GSElimits	This argument can be used to load only a contiguous subset of the GSMs from a GSE. It should be specified as a vector of length 2 specifying the start and end (inclusive) GSMs to load. This could be useful for splitting up large GSEs into more manageable parts, for example.
GSEMatrix	A boolean telling GEOquery whether or not to use GSE Series Matrix files from GEO. The parsing of these files can be many orders-of-magnitude faster than parsing the GSE SOFT format files. Defaults to TRUE, meaning that the SOFT format parsing will not occur; set to FALSE if you for some reason need other columns from the GSE records.
AnnotGPL	A boolean defaulting to FALSE as to whether or not to use the Annotation GPL information. These files are nice to use because they contain up-to-date information remapped from Entrez Gene on a regular basis. However, they do not exist for all GPLs; in general, they are only available for GPLs referenced by a GDS
getGPL	A boolean defaulting to TRUE as to whether or not to download and include GPL information when getting a GSEMatrix file. You may want to set this to FALSE if you know that you are going to annotate your featureData using Bioconductor tools rather than relying on information provided through NCBI GEO. Download times can also be greatly reduced by specifying FALSE.

#### Details

getGEO functions to download and parse information available from NCBI GEO (http://www.ncbi.nlm.nih.gov/geo). Here are some details about what is avaible from GEO. All entity types are handled by getGEO and essentially any information in the GEO SOFT format is reflected in the resulting data structure.

From the GEO website:

The Gene Expression Omnibus (GEO) from NCBI 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), and mass spectrometry proteomic data. At the most basic level of organization of GEO, there are three entity types that may be supplied by users: Platforms, Samples, and Series. Additionally, there is a curated entity called a GEO dataset.

#### getGEO

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.

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.

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).

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.

#### Value

An object of the appropriate class (GDS, GPL, GSM, or GSE) is returned. If the GSEMatrix option is used, then a list of ExpressionSet objects is returned, one for each SeriesMatrix file associated with the GSE accession. If the filename argument is used in combination with a GSEMatrix file, then the return value is a single ExpressionSet.

#### Warning

Some of the files that are downloaded, particularly those associated with GSE entries from GEO are absolutely ENORMOUS and parsing them can take quite some time and memory. So, particularly when working with large GSE entries, expect that you may need a good chunk of memory and that coffee may be involved when parsing....

#### Author(s)

Sean Davis

#### See Also

getGEOfile

## Examples

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

```
getGEOfile
```

#### Description

This function simply downloads a SOFT format file associated with the GEO accession number given.

#### Usage

```
getGEOfile(GEO, destdir = tempdir(), AnnotGPL = FALSE, amount = c("full", "brief", "quick", "data"))
```

#### Arguments

GEO	Character string, the GEO accession for download (eg., GDS84, GPL96, GSE2553, or GSM10)
destdir	Directory in which to store the resulting downloaded file. Defaults to tempdir()
AnnotGPL	A boolean defaulting to FALSE as to whether or not to use the Annotation GPL information. These files are nice to use because they contain up-to-date information remapped from Entrez Gene on a regular basis. However, they do not exist for all GPLs; in general, they are only available for GPLs referenced by a GDS
amount	Amount of information to pull from GEO. Only applies to GSE, GPL, or GSM. See details

## Details

This function downloads GEO SOFT files based on accession number. It does not do any parsing. The first two arguments should be fairly self-explanatory, but the last is based on the input to the acc.cgi url at the geo website. In the default "full" mode, the entire SOFT format file is downloaded. Both "brief" and "quick" offer shortened versions of the files, good for "peeking" at the file before a big download on a slow connection. Finally, "data" downloads only the data table part of the SOFT file and is good for downloading a simple EXCEL-like file for use with other programs (a convenience).

#### Value

Invisibly returns the full path of the downloaded file.

#### Author(s)

Sean Davis

#### References

http://www.ncbi.nlm.nih.gov/geo/query/acc.cgi

## getGEOSuppFiles

#### See Also

getGE0

## Examples

# myfile <- getGEOfile('GDS10')</pre>

getGEOSuppFiles Get Supplemental Files from GEO

#### Description

NCBI GEO allows supplemental files to be attached to GEO Series (GSE), GEO platforms (GPL), and GEO samples (GSM). This function "knows" how to get these files based on the GEO accession. No parsing of the downloaded files is attempted, since the file format is not generally knowable by the computer.

#### Usage

```
getGEOSuppFiles(GEO, makeDirectory = TRUE, baseDir = getwd())
```

#### Arguments

GEO	A GEO accession number such as GPL1073 or GSM1137
makeDirectory	Should a "subdirectory" for the downloaded files be created? Default is TRUE. If FALSE, the files will be downloaded directly into the baseDir.
baseDir	The base directory for the downloads. Default is the current working directory.

#### Details

Again, just a note that the files are simply downloaded.

#### Value

A data frame is returned invisibly with rownames representing the full path of the resulting downloaded files and the records in the data.frame the output of file.info for each downloaded file.

#### Author(s)

Sean Davis <sdavis2@mail.nih.gov>

#### Examples

# a <- getGEOSuppFiles('GSM1137')
# a</pre>

getGSEDataTables

## Description

In some cases, instead of individual sample records (GSM) containing information regarding sample phenotypes, the GEO Series contains that information in an attached data table. And example is given by GSE3494 where there are two data tables with important information contained within them. Using getGEO with the standard parameters downloads the GSEMatrix file which, unfortunately, does not contain the information in the data tables. This function simply downloads the "header" information from the GSE record and parses out the data tables into R data.frames.

#### Usage

```
getGSEDataTables(GSE)
```

#### Arguments

GSE The GSE identifier, such as "GSE3494".

#### Value

A list of data.frames.

#### Author(s)

Sean Davis <sdavis2@mail.nih.gov>

#### References

http://www.ncbi.nlm.nih.gov/geo

#### See Also

getGE0

## Examples

dfl = getGSEDataTables("GSE3494")
lapply(dfl,head)

GPL-class

## Description

Contains a full GEO Platform entity

## **Objects from the Class**

Objects can be created by calls of the form new("GPL", ...).

## Slots

dataTable: Object of class "GEODataTable"

header: Object of class "list" containing metadata associated with the GPL

## Extends

Class "GEOData", directly.

#### Methods

No methods defined with class "GPL" in the signature, but methods applicable to GEOData are also applicable here.

### Author(s)

Sean Davis

## See Also

GEOData-class

GSE-class Class "GSE"

## Description

Contains a GEO Series entity

## **Objects from the Class**

Objects can be created by calls of the form new("GSE", ...).

header: Object of class "list" containing metadata for the series

gsms: Object of class "list" containing a list of items of class GSM associated with the series gpls: Object of class "list" containing a list of items of class GPL associate with the series

#### Methods

**GPLList** signature(object = "GSE"): returns a list, each item of the list being a GPL object **GSMList** signature(object = "GSE"): returns a list, each item of the list being a GSM object **Meta** signature(object = "GSE"): returns a list, the metadata associated with the GSE

#### Author(s)

Sean Davis

### See Also

GPL-class,GSM-class

GSM-class

Class "GSM"

#### Description

A class containing a GEO Sample entity

## **Objects from the Class**

Objects can be created by calls of the form new("GSM", ...).

#### Slots

dataTable: Object of class "GEODataTable"
header: Object of class "list" containing the metadata associated with the sample

### Extends

```
Class "GEOData", directly.
```

#### Methods

No methods defined with class "GSM" in the signature, but methods that apply to the GEOData also apply here.

#### Author(s)

Sean Davis

## gunzip

## See Also

GEOData-class

gunzip

Gunzip a file

## Description

gunzip a file

## Usage

```
gunzip(filename, destname = gsub("[.]gz$", "", filename), overwrite = FALSE, remove = TRUE, BFR.SIZE =
```

## Arguments

filename	The filename to be unzipped
destname	The destination file
overwrite	Boolean indicating whether or not to overwrite a destfile of the same name
remove	Boolean indicating whether or not to remove the original file after completion
BFR.SIZE	The size of the read buffer

## Details

This function was stripped out of R.utils due to breaking some stuff on the bioconductor build machine.

## Value

Invisibly, the number of bytes read.

## Author(s)

Original author: Henrik Bengtsson

## See Also

gzfile

parseGE0

## Description

Workhorse GEO parsers.

## Usage

```
parseGEO(fname, GSElimits, destdir=tempdir(), AnnotGPL=FALSE, getGPL=TRUE)
parseGPL(fname)
parseGDS(fname)
parseGSE(fname, GSElimits)
parseGSM(fname)
```

## Arguments

fname	The filename of a SOFT format file. If the filename ends in .gz, a gzfile() connection is used to read the file directly.
GSElimits	Used to limit the number of GSMs parsed into the GSE object; useful for memory management for large GSEs.
destdir	The destination directory into which files will be saved (to be used for caching)
AnnotGPL	Fetch the annotation GPL if available
getGPL	Fetch the GPL associated with a GSEMatrix entity (should remain TRUE for all normal use cases)

#### Details

These are probably not useful to the end-user. Use getGEO to access these functions. parseGEO simply delegates to the appropriate specific parser. There should be no reason to use the parseGPL, parseGDS, parseGSE, or parseGSM functions directly.

## Value

parseGEO returns an object of the associated type. For example, if it is passed the text from a GDS entry, a GDS object is returned.

#### Author(s)

Sean Davis

## See Also

getGE0

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