How to use MeSH-related Packages

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April 17, 2014

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1 Introduction

This document provides the way to use MeSH-related packages; MeSH.db, MeSH.AOR.db, MeSH.PCR.db, org.MeSH.XXX.db-type packages, MeSHDbi, and meshr packages. MeSH (Medical Subject Headings) is the NLM (U. S. National Library of Medicine) controlled vocabulary used to manually index articles for MEDLINE/PubMed [1] and is a collection of a comprehensive life science vocabulary. MeSH contains more than 25,000 clinical and



Figure 1: MeSH Term

biological terms. The amount of MeSH term is about twice as large as that of GO (Gene Ontology)[2] and its categories are also wider. MeSH in 2014 proposed its 19 categories and MeSH.db provides 16 of them, which are actually assigned to some MeSH terms. Each category is expressed as single capital alphabet as abbreviation defined by NLM. Therefore MeSH is an expected to be much detailed and exhaustive gene annotation tool. Some software or databases using MeSH are now proposed [3, 4, 5, 6].

This vignette introduces R/Bioconductor packages for handling MeSH in R. Original MeSH data is accessible by NLM FTP site (http://www.nlm.nih.gov/mesh/filelist.html). The data are downloadable as plain-text format (ASCII MeSH; d2014.bin / q2014.bin). These files were pre-processed by our data-processing pipeline (figure 2) and corresponding information is summarized as a table in SQLite3 file and packed into MeSH.db, MeSH.AOR.db, and MeSH.PCR.db.

Abbreviation	Category
A	Anatomy
В	Organisms
С	Diseases
D	Chemicals and Drugs
E	Analytical, Diagnostic and Therapeutic Techniques and Equipment
F	Psychiatry and Psychology
G	Phenomena and Processes
Н	Disciplines and Occupations
I	Anthropology, Education, Sociology and Social Phenomena
J	Technology and Food and Beverages
K	Humanities
L	Information Science
M	Persons
N	Health Care
V	Publication Type
Z	Geographical Locations

1.1 About MeSH

MeSH.db provides the corresponding table which contains MeSH ID, MeSH term, MeSH category, synonym, qualifier ID, and qualifier term. Qualifier term means more rough annotation (subheadings) than MeSH. MeSH has hierarchical structure like GO. Such structure is provided as MeSH.AOR.db (AOR: ancestor-offspring Relationships) and MeSH.PCR.db (PCR: parent-child Relationships as corresponding table.

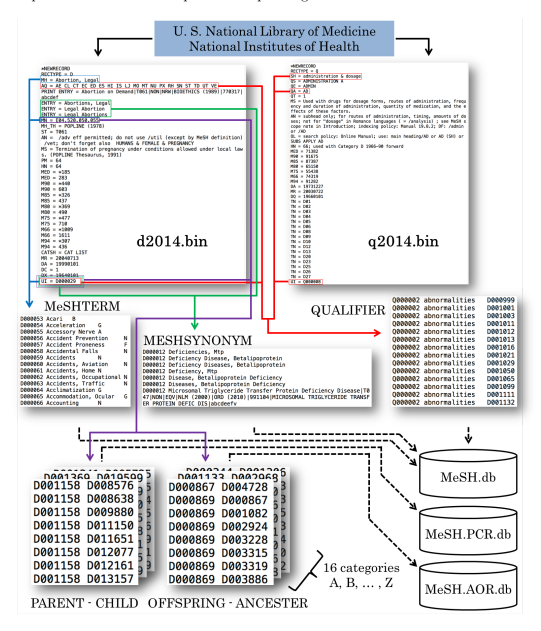


Figure 2: Data pre-process for MeSH.db

1.2 The correspondence between MeSH ID and NCBI Entrez Gene ID

org.MeSH.XXX.db (XXX is an abbreviation of species name such as Hsa: Homo sapiens) packages provide the correspondence between Entrez Gene IDs and NLM MeSH IDs. Such correspondence in wide variety of organisms are summarized as each org.MeSH.XXX.db by three way of methods, Gendoo[4], gene2pubmed, and RBBH (reciprocal BLAST best Hit).

Gendoo is the web-application based on text-mining of PubMed. Co-occurrence relations in PubMed document are exhaustively retrieved and much relevant correspondence are filtered by some information science techniques.

gene2pubmed is the correspondence between Entrez Gene IDs and NLM PubMed IDs. These relationship is manually assigned by NCBI curator teams. We also summarized the relationship between MeSH Terms and PubMed IDs from licensed-PubMed, then merged as Gene IDs - MeSH IDs correspondence.

For some minor species including non-model organisms, which have no sufficient databases for annotation, we defined 15 well-annotated organisms and 100 minor-organisms, then conducted RBBH between all possible combinations using BLASTP search.

Method Way of corresponding Entrez Gene IDs and MeSH IDs	
Gendoo	Text-mining
gene2pubmed	Manual curation by NCBI teams
RBBH	sequence homology with BLASTP search (E-value $< 10^{-50}$)



Figure 3: 120 organisms for org.MeSH.XXX.db and those source databases $\,$

1.3 Database interface package for MeSH-related packages

We also implemented a database interface (DBI) package named MeSHDbi. This package is important because of two reasons. First reason is a unification of DBI functions for MeSH-related packages. MeSH.db, MeSH.AOR.db, MeSH.PCR.db, and org.MeSH.XXX.db packages inherit the MeSHDb-class defined by MeSHDbi and behavior of these packages is uniformly designed. Second reason is supporting construction of user's original org.MeSH.XXX.db package. Due to the rapid development of DNA sequence technology, wide variety of genome sequences are more and more determined and the correspondence of Gene IDs and MeSH IDs may be designed by many databases [3, 4, 5, 6]. Therefore, we prepared the function to create org.MeSH.XXX.db package for a situation in which users can retrieved the relationship between Gene IDs and MeSH IDs by some means.

1.4 MeSH term enrichment analysis

To analyze MeSH-related packages with omics data, we implement *meshr* package, which is for conducting enrichment analysis using MeSH data. This package internally imports *MeSH.db*, *MeSH.AOR.db*, *MeSH.PCR.db* and *org.MeSH.XXX.db*, then conducts enrichment analysis to detect highly enriched MeSH terms in gene sets of interesting species.

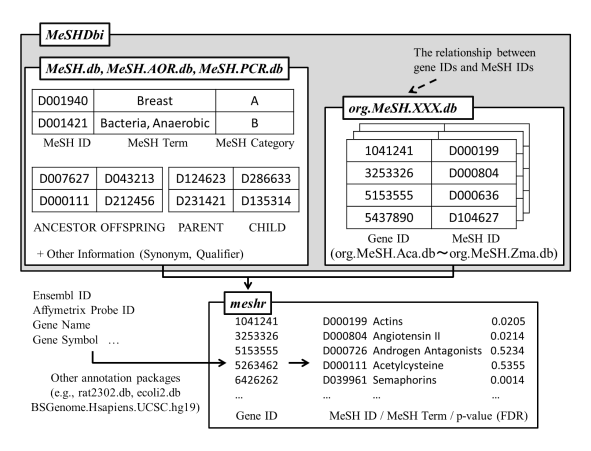


Figure 4: The relationship of meshr and other MeSH-related packages

2 Exercise

2.1 Access MeSH Term

2.1.1 columns, keytypes, keys, and select

In our packages, all data are extracted by only 4 functions defined by AnnotationDbi; **keytypes**, **columns**, **keys** and **select**. In this section, we demonstrate how to use these functions by using MeSH.db.

At first, install and load the MeSH.db.

> library(MeSH.db)

ls function shows all objects in this package. MeSH.db object is generated. This is also package's name and all MeSH-related packages provide the object named as package name (e.g., MeSH.db, org.MeSH.Mmu.db).

```
> ls("package:MeSH.db")
[1] "MeSH.db"

> MeSH.db

[1] "##### class ####"
[1] "MeSHDb"
attr(,"package")
[1] "MeSHDbi"
[1] "##### connection #####"

<SQLiteConnection: DBI CON (31073, 0)>
[1] "#### package name ####"
[1] "MeSH.db"
```

Here, we use columns, keytypes, keys and select against MeSH.db.

columns returns the rows which we can retrieve in MeSH.db.

```
> columns(MeSH.db)
```

- [1] "MESHID" "MESHTERM" "CATEGORY" "SYNONYM" "QUALIFIERID"
- [6] "QUALIFIER"

keytypes returns the rows which can be used as the optional parameter in **keys** and **select** functions against MeSH.db.

```
> keytypes(MeSH.db)
```

- [1] "MESHID" "MESHTERM" "CATEGORY" "SYNONYM" "QUALIFIERID"
- [6] "QUALIFIER"

keys function returns the value of keytype.

```
> k <- keys(MeSH.db, keytype = "MESHID")
> length(k)
[1] 24416
> head(k)
```

[1] "D000001" "D000002" "D000003" "D000004" "D000005" "D000006"

select function returns rows in particular columns, which are having user-specified keys. This function provides the data as a dataframe. Now, we will retrieve the rows in which MESHID is equivalent to MESHTERM.

```
> select(MeSH.db, keys = k[1:10], columns = c("MESHID", "MESHTERM"),
+ keytype = "MESHID")
```

MESHTERM	MESHID	
Calcimycin	D000001	1
Temefos	D000002	141
Abattoirs	D000003	219
${\tt Abbreviations} \ {\tt as} \ {\tt Topic}$	D000004	246
Abdomen	D000005	255
Abdomen, Acute	D000006	271
Abdominal Injuries	D000007	382
Abdominal Neoplasms	D000008	493
Abdominal Muscles	D000009	619
Abducens Nerve	D000010	700

2.1.2 Annotation of Leukemia

Next, we will retrieve some information about *Leukemia* by our packages.

select function retrieves rows in which MESHTERM is "Leukemia" in the MeSH.db table.

```
> LEU <- select(MeSH.db, keys = "Leukemia", columns = c("MESHID",
      "MESHTERM", "CATEGORY", "SYNONYM"), keytype = "MESHTERM")
> LEU
    MESHID MESHTERM CATEGORY
   D007938 Leukemia
38 D007938 Leukemia
75 D007938 Leukemia
                            C
112 D007938 Leukemia
                            С
149 D007938 Leukemia
                            C
                                                 SYNONYM
                                         Leucocythaemias
1
38 Leucocythaemia|T191|NON|EQV|NLM (2012)|110224|abcdef
75
                                          Leucocythemias
```

112 Leucocythemia|T191|NON|EQV|NLM (2012)|110224|abcdef

select function shows that MESHID of Leukemia is D007938 and Leukemia is categorized as C (Diseases). Leukemia has some synonyms like Leucocythaemias, Leucocythaemias, Leucocythaemias and Leukemias.

Leukemias

As mentioned above, MeSH has hierarchical structures. MeSH.AOR.db and MeSH.PCR.db packages provide such hierarchical information of MeSH. For example, MeSH.AOR.db enable us to examine the top terms of Leukemia.

D009370 has found above Leukemia.

149

These MeSH IDs can be translated to MeSH Term.

```
> select(MeSH.db, keys = ANC[1, 1], columns = c("MESHTERM"), keytype = "MESHID")
```

MESHTERM

1 Neoplasms by Histologic Type

In this way, we can specify that Leukemia is categorized as one of NeoplasmsbyHistologicType. Once keytype-parameter set to opposite direction (OFFSPRING to ANCESTOR), other MeSH IDs in lower hierarchies also can be retrieved.

```
> OFF <- select(MeSH.AOR.db, keys = "D007938", columns = c("ANCESTOR",
      "OFFSPRING"), keytype = "ANCESTOR")
> OFF
  ANCESTOR OFFSPRING
1 D007938
             D007942
2 D007938
             D007943
3 D007938
             D007945
4 D007938
             D007946
5 D007938
             D007951
6 D007938
             D007952
7 D007938
             D007953
8 D007938
             D016582
9 D007938
             D016583
> select(MeSH.db, keys = OFF[, 2], columns = c("MESHTERM"), keytype = "MESHID")
                        MESHTERM
1
          Leukemia, Experimental
505
            Leukemia, Hairy Cell
              Leukemia, Lymphoid
764
1023
             Leukemia, Mast-Cell
1208
               Leukemia, Myeloid
2022
           Leukemia, Plasma Cell
2281 Leukemia, Radiation-Induced
2466
                Leukemia, Feline
2571
        Enzootic Bovine Leukosis
There are a lot of MeSH terms, which means Leukemia has many lower hierarchies.
```

MeSH.PCR.db provides the directly lower (or upper) terms.

```
> library("MeSH.PCR.db")
> CHI <- select(MeSH.PCR.db, keys = LEU[1, 1], columns = c("PARENT",
      "CHILD"), keytype = "PARENT")
> head(CHI)
  PARENT
            CHILD
1 D007938 D001353
2 D007938 D001752
3 D007938 D004915
4 D007938 D007939
5 D007938 D007940
6 D007938 D007941
```

 $Leukemia \ {\bf has\ a\ lot\ of\ subtypes\ like}\ Avian Leukosis, Blast Crisis, Leukemia, Erythroblastic, Acute and so on.$

2.1.3 Other functions

Some optional functions for much complex data acquisition are also provided. In this section, users may need some basic SQL knowledge (see also RSQLite).

dbInfo returns the information of the package. dbfile returns the directory where sqlite file is stored. dbschema returns the schema of database. dbconn returns the connection constructed by RSQLite.

> dbInfo(MeSH.db)

```
NAME
                                                         VALUE
       SOURCEDATE
                                                   19-Nov-2013
1
2
       SOURCENAME
                                    Medical Subject Headings
        SOURCEURL http://www.nlm.nih.gov/mesh/filelist.html
3
         DBSCHEMA
                                                       MeSH.db
5 DBSCHEMAVERSION
                                                           1.0
6
          package
                                                       MeSH.db
7
          Db type
                                                       MeSH.Db
```

- > dbfile(MeSH.db)
- [1] "/Library/Frameworks/R.framework/Versions/3.1/Resources/library/MeSH.db/extdata/MeSH.db
- > dbschema(MeSH.db)
- [1] "CREATE TABLE DATA (\n MESHID CHAR(7) NOT NULL, \n MESHTERM VARCHAR(100) NOT NULL, \n
- [2] "CREATE TABLE METADATA (\n NAME VARCHAR(80),\n VALUE VARCHAR(255)\n)"
- > dbconn(MeSH.db)

```
<SQLiteConnection: DBI CON (31073, 0)>
```

dbschema shows the data is stored as a table named "DATA" in the sqlite database and the table has six columns; MESHID, MESHTERM, CATEGORY, SYNONYM, QUALIFIERID, and QUALIFIER. Therefore, we can retrieve data by much complex SQL query like below;

```
> library("RSQLite")
```

- > SQL1 <- paste("SELECT MESHTERM, QUALIFIERID, QUALIFIER FROM DATA",
- + "WHERE MESHID = 'D000001'", "AND QUALIFIERID = 'Q000494'")
- > dbGetQuery(dbconn(MeSH.db), SQL1)

	MESHTERM	QUALIFIERID	QUALIFIER
1	Calcimycin	Q000494	pharmacology
2	Calcimycin	Q000494	pharmacology
3	Calcimycin	Q000494	pharmacology
4	Calcimycin	Q000494	pharmacology
5	Calcimycin	Q000494	pharmacology

```
> SQL2 <- paste("SELECT ANCESTOR, OFFSPRING FROM DATA", "WHERE OFFSPRING = 'D0000002'",
      "OR OFFSPRING = 'D0000003'", "OR OFFSPRING = 'D0000004'", "OR ANCESTOR = 'D009275'")
> dbGetQuery(dbconn(MeSH.AOR.db), SQL2)
 ANCESTOR OFFSPRING
1 D063086 D000002
2 D008462 D000003
3 D009275 D000004
4 D009275 D000851
5 D009275 D004850
> SQL3 <- paste("SELECT PARENT, CHILD FROM DATA", "WHERE PARENT = 'D0000005'",
      "AND NOT CHILD = 'D004312'")
> dbGetQuery(dbconn(MeSH.PCR.db), SQL3)
    PARENT CHILD
1 D000005 D006119
2 D000005 D007264
3 D000005 D008643
4 D000005 D008646
5 D000005 D009852
6 D000005 D010529
7 D000005 D010537
8 D000005 D012187
9 D000005 D014472
10 D000005 D034841
11 D000005 D034861
```

12 D000005 D054048

2.2 org.MeSH.XXX.db-type packages

2.2.1 Annotation of 120 organisms

As well as MeSH.db, MeSH.AOR.db, and MeSH.PCR.db, org.MeSH.XXX.db-type packages also use 4 functions (**keytypes**, **columns**, **keys** and **select**) to extract data.

```
> library("org.MeSH.Hsa.db")
> columns(org.MeSH.Hsa.db)
[1] "GENEID"
                   "MESHID"
                                   "MESHCATEGORY" "SOURCEID"
                                                                  "SOURCEDB"
> keytypes(org.MeSH.Hsa.db)
[1] "GENEID"
                   "MESHID"
                                   "MESHCATEGORY" "SOURCEID"
                                                                  "SOURCEDB"
> key_HSA <- keys(org.MeSH.Hsa.db, keytype = "MESHID")</pre>
> select(MeSH.db, keys = key_HSA[1:10], columns = c("MESHID", "MESHTERM"),
      keytype = "MESHID")
      MESHID
                                   MESHTERM
1
     D000001
                                Calcimycin
141 D000002
                                    Temefos
219 D000005
                                    Abdomen
                            Abdomen, Acute
235 D000006
346 D000007
                        Abdominal Injuries
457 D000008
                       Abdominal Neoplasms
583 D000009
                         Abdominal Muscles
664 D000010
                            Abducens Nerve
1139 D000011 Abelson murine leukemia virus
1178 D000012
                      Abetalipoproteinemia
```

Moreover, these packages have other additional functions like **species**, **nomenclature**, **listDatabases**. In each org.MeSH.XXX.db, **species** function returns the common name and **nomenclature** returns the scientific name.

```
> library("org.MeSH.Aca.db")
> library("org.MeSH.Atu.K84.db")
> library("org.MeSH.Bsu.168.db")
> library("org.MeSH.Syn.db")
> species(org.MeSH.Hsa.db)

[1] "Human"
> species(org.MeSH.Aca.db)

[1] "Lizard"
> species(org.MeSH.Atu.K84.db)
```

```
[1] NA
> species(org.MeSH.Bsu.168.db)
[1] NA
> species(org.MeSH.Syn.db)
[1] "Cyanobacteria"
> nomenclature(org.MeSH.Hsa.db)
[1] "Homo sapiens"
> nomenclature(org.MeSH.Aca.db)
[1] "Anolis carolinensis"
> nomenclature(org.MeSH.Atu.K84.db)
[1] "Agrobacterium Tumefacienes K84"
> nomenclature(org.MeSH.Bsu.168.db)
[1] "Bacillus subtilis subsp. spizizenii str. 168"
> nomenclature(org.MeSH.Syn.db)
[1] "Synechocystis"
   listDatabases function returns the source of data (figure 3). In regard to RBBH, name
of organisms is returned. These values are important when users specify the database for
MeSH Term enrichment analysis (see the section 2.4).
> listDatabases(org.MeSH.Hsa.db)
     SOURCEDB
       gendoo
1
2 gene2pubmed
```

> listDatabases(org.MeSH.Aca.db)

SOURCEDB

Arabidopsis thaliana

Bacillus subtilis subsp. spizizenii str. 168

Bos taurus

Caenorhabditis elegans

Danio rerio

Drosophila melanogaster

```
Escherichia coli str K-12 substr. MG1655
7
8
                                   Gallus gallus
9
                                    Homo sapiens
                                    Mus musculus
10
11
                               Rattus norvegicus
                 Saccharomyces cerevisiae S288c
12
13
                Schizosaccharomyces pombe 972h-
14
                                      Sus scrofa
15
                                  Xenopus laevis
16
                                     gene2pubmed
```

> listDatabases(org.MeSH.Atu.K84.db)

SOURCEDB

```
1
                            Arabidopsis thaliana
2
   Bacillus subtilis subsp. spizizenii str. 168
3
                                      Bos taurus
4
                          Caenorhabditis elegans
5
                                     Danio rerio
6
                         Drosophila melanogaster
7
       Escherichia coli str K-12 substr. MG1655
8
                                   Gallus gallus
9
                                    Homo sapiens
10
                                    Mus musculus
                               Rattus norvegicus
11
12
                 Saccharomyces cerevisiae S288c
13
                Schizosaccharomyces pombe 972h-
14
                                      Sus scrofa
                                  Xenopus laevis
15
```

> listDatabases(org.MeSH.Bsu.168.db)

SOURCEDB

- 1 gene2pubmed
- > listDatabases(org.MeSH.Syn.db)

SOURCEDB

1 gendoo

2.3 MeSHDbi

2.3.1 User's custom org.MeSH.XXX.db package

Although most of users may not be conscious of this package, MeSHDbi is important for our MeSH packages. This package regulates class definition of MeSH object (MeSHDb-class). Besides, this package constructs user's original org.MeSH.XXX.db package. **makeGeneMeSHPackage** easily constructs such package.

```
> library("MeSHDbi")
> example("makeGeneMeSHPackage")
mGMSHP> ## makeGeneMeSHPackage enable users to construct
mGMSHP> ## user's own custom MeSH package
mGMSHP>
mGMSHP> ## this is test data which means the relationship between
mGMSHP> ## Entrez gene IDs of Pseudomonas aeruginosa PAO1
mGMSHP> ## and its MeSH IDs.
mGMSHP> data(PAO1)
mGMSHP> head(PAO1)
  GENEID MESHID MESHCATEGORY SOURCEID
1 877657 D000265
                            D
                                937805
2 877657 D000265
                            D 937805
3 877657 D001412
                            B 937805
4 877657 D001412
                           B 937805
5 877657 D001426
                           D 937805
6 877657 D001483
                           G 937805
                                      SOURCEDB
1 Bacillus subtilis subsp. spizizenii str. 168
2 Bacillus subtilis subsp. spizizenii str. 168
3 Bacillus subtilis subsp. spizizenii str. 168
4 Bacillus subtilis subsp. spizizenii str. 168
5 Bacillus subtilis subsp. spizizenii str. 168
6 Bacillus subtilis subsp. spizizenii str. 168
mGMSHP> # We are also needed to prepare meta data as follows.
mGMSHP> data(metaPAO1)
mGMSHP> metaPA01
             NAME.
                                                  VALUE.
1
       SOURCEDATE
                                           31-July-2013
2
       SOURCENAME
                                                 BLASTP
3
        SOURCEURL ftp://ftp.ncbi.nlm.nih.gov/gene/DATA/
         DBSCHEMA
                                   org.Pae.PAO1.MeSH.db
5 DBSCHEMAVERSION
6
         ORGANISM
                            Pseudomonas aeruginosa PAO1
```

```
7
          SPECIES
8
          package
                                          AnnotationDbi
          Db type
                                                BLASTDb
mGMSHP> ## sets up a temporary directory for this example
mGMSHP> ## (users won't need to do this step)
mGMSHP> destination <- tempfile()
mGMSHP> dir.create(destination)
mGMSHP> ## makes an Organism package for human called Homo.sapiens
mGMSHP> makeGeneMeSHPackage(pkgname = "org.MeSH.Pae.db",
mGMSHP+
                                                 data = PAO1,
mGMSHP+
                  metadata = metaPAO1,
mGMSHP+
                                                 organism = "Pseudomonas aeruginosa PAO1",
mGMSHP+
                                                 version = "1.0.0",
mGMSHP+
                                                 maintainer = "Koki Tsuyuzaki <k.t.the-answe
mGMSHP+
                                                 author = "Koki Tsuyuzaki",
                                                 destDir = destination,
mGMSHP+
mGMSHP+
                                                 license="Artistic-2.0")
Creating package in /var/folders/k0/tk8gl4bj2_v2mbjx80ydsznw0000gn/T//RtmpQXcFsU/file796166
[1] TRUE
```

2.4 meshr

2.4.1 MeSH enrichment analysis

The *meshr* package is designed to conduct an enrichment analysis for MeSH. The idea behind this package is analogous to GO enrichment analysis, where sets of genes is analyzed to extract common annotated biological properties. The usage of *meshr* closely follows that of the Bioconductor *GOstats* package. Thus, users who are familiar with *GOstats* may easily handle *meshr*.

The meshr package accepts selected and universal genes as input, and returns significantly overrepresented MeSH terms. It is used in conjunction with MeSH.db package and one of the annotation packages, (e.g., org.MeSH.Hsa.db). This section serves as a quick guide to the meshr, while illustrating entire process to perform a MeSH enrichment analysis.

Here, we use the example data set taken from the Bioconductor package *cummeRbund*. The example data are located in library/cummeRbund/extdata/. This RNA-Seq data were taken from three samples, "iPS", "hESC", and "Fibroblasts". We first created two objects of gene sets, i.e., selected and universal genes, by comparing significantly regulated genes between iPS and hESC under the significance level of 0.05, then mapped the Gene Symbols to Entrez Gene IDs through the *org.Hs.eg.db* package. Pre-processed Gene IDs are easily accessible by **data** function.

```
> library("meshr")
> data(geneid.cummeRbund)
> data(sig.geneid.cummeRbund)
```

Finally 303 universal genes and 104 selected genes are detected and subsequently used for the MeSH enrichment analysis.

```
> dim(geneid.cummeRbund)[1]
[1] 303
> dim(sig.geneid.cummeRbund)[1]
[1] 104
```

We proceed to uncover a characteristic of MeSH terms that the set of identified genes share each other via the *meshr* package. We first load the required packages.

```
> library("fdrtool")
> library("org.MeSH.Hsa.db")
```

We create a parameter instance by specifying the objects of selected and universal genes, the name of the annotation package, the category of MeSH, the database of correspondence between Gene IDs and MeSH IDs (see also ${\bf listDatabases}$ in the section 2.2.1), p-value cutoff, and the choice of a multiple-testing correction method. In this first example, we use org.MeSH.Hsa.db because the above RNA-seq data is extracted from human cells. We

choose D (Drugs and chemicals) category, gendoo database, p-value cutoff 0.05, and no multiple-testing adjustment. For more details on description of all the arguments, readers are referred to the **MeSHHyperGParams-class** help page.

```
> meshParams <- new("MeSHHyperGParams", geneIds = sig.geneid.cummeRbund[,
+ 2], universeGeneIds = geneid.cummeRbund[, 2], annotation = "org.MeSH.Hsa.db",
+ category = "D", database = "gendoo", pvalueCutoff = 0.05,
+ pAdjust = "none")</pre>
```

The **meshHyperGTest** function carries out a hypergeometric test and returns an instance of class **MeSHHyperGResult**.

> meshR <- meshHyperGTest(meshParams)</pre>

Simply typing the **MeSHHyperGResult** class gives a brief description of the analysis, including the choice of a MeSH category, the annotation data used, and a total number of identified overrepresented MeSH terms.

> meshR

```
MeSH enrichment analysis for category Chemicals and Drugs
Annotation package used: org.MeSH.Hsa.db
The correspondance is retrived from: gendoo
Number of MeSH terms identified: 128
```

Full details of the result is obtained by calling the **summary** function on the **MeSH-HyperGResult** instance. This presents significantly enriched MeSH ID, MeSH term, and their associated p-values.

> head(summary(meshR))

MESHID		MESHTERM	PVALUE
D011509		Proteoglycans	0.0000123822
D051744		Protein Kinase C-epsilon	0.0001575585
D040262	Receptors,	Vascular Endothelial Growth Factor	0.0006466433
D017077		Culture Media, Conditioned	0.0008313715
D051380		GRB2 Adaptor Protein	0.0032728303
D009218		Mvosins	0.0034614243

Switching to test another MeSH category and another database can be easily done. For example, to choose the category as G (Phenomena and Processes) and the database as gene2pubmed, we can do the following.

```
> category(meshParams) <- "G"
> database(meshParams) <- "gene2pubmed"
> meshR <- meshHyperGTest(meshParams)
> summary(meshR)

MESHID MESHTERM PVALUE
1 D005819 Genetic Markers 0.03966931
```

3 Setup

```
This vignette was built on:
```

```
> sessionInfo()
```

R version 3.1.0 (2014-04-10)

Platform: x86_64-apple-darwin10.8.0 (64-bit)

locale:

[1] ja_JP.UTF-8/ja_JP.UTF-8/ja_JP.UTF-8/ja_JP.UTF-8

attached base packages:

[1] grid parallel stats graphics grDevices utils datasets

[8] methods base

other attached packages:

[1] meshr_0.99.0 org.Hs.eg.db_2.14.0

[3] cummeRbund_2.6.0 Gviz_1.8.0

[5] rtracklayer_1.24.0 GenomicRanges_1.16.0 [7] IRanges_1.22.2 fastcluster_1.1.13 ggplot2_0.9.3.1 [9] reshape2_1.2.2

[11] Category_2.30.0 GO.db_2.14.0

[13] Matrix_1.1-3 AnnotationDbi_1.26.0

[15] GenomeInfoDb_1.0.1 Biobase_2.24.0 [17] BiocGenerics_0.10.0 fdrtool_1.2.11

[19] org.MeSH.Syn.db_0.99.0 org.MeSH.Bsu.168.db_0.99.0

[21] org.MeSH.Atu.K84.db_0.99.0 org.MeSH.Aca.db_0.99.0

[23] org.MeSH.Hsa.db_0.99.0 RSQLite_0.11.4 [25] DBI_0.2-7 MeSH.PCR.db_0.99.0 [27] MeSH.AOR.db_0.99.0 MeSH.db_0.99.0

[29] MeSHDbi_1.0.0

loaded via a namespace (and not attached):

	±		
[1]	annotate_1.42.0	BatchJobs_1.2	BBmisc_1.5
[4]	BiocParallel_0.6.0	biomaRt_2.20.0	Biostrings_2.32.0
[7]	biovizBase_1.12.0	bitops_1.0-6	brew_1.0-6
[10]	BSgenome_1.32.0	cluster_1.15.2	codetools_0.2-8
[13]	colorspace_1.2-4	dichromat_2.0-0	digest_0.6.4
[16]	fail_1.2	foreach_1.4.2	Formula_1.1-1
[19]	genefilter_1.46.0	GenomicAlignments_1.0.0	<pre>GenomicFeatures_1.16.0</pre>
[22]	graph_1.42.0	GSEABase_1.26.0	gtable_0.1.2
[25]	Hmisc_3.14-4	iterators_1.0.7	labeling_0.2
[28]	lattice_0.20-29	latticeExtra_0.6-26	MASS_7.3-31
[31]	matrixStats_0.8.14	munsell_0.4.2	plyr_1.8.1
[34]	proto_0.3-10	R.methodsS3_1.6.1	RBGL_1.40.0

[37] RColorBrewer_1.0-5	Rcpp_0.11.1	RCurl_1.95-4.1
[40] Rsamtools_1.16.0	scales_0.2.3	sendmailR_1.1-2
[43] splines_3.1.0	stats4_3.1.0	stringr_0.6.2
[46] survival_2.37-7	tools_3.1.0	VariantAnnotation_1.10.0
[49] XML_3.98-1.1	xtable_1.7-3	XVector_0.4.0
[52] zlibbioc_1.10.0		

References

- [1] S. J. Nelson and et al. The MeSH translation maintenance system: structure, interface design, and implementation. *Stud. Health Technol. Inform.*, 107: 67-69, 2004.
- [2] M. Ashburner and et al. Gene ontology: tool for the unification of biology. The Gene Ontology Consortium. *Nat. Genet.*, 25(1): 25-29, 2000.
- [3] T. Nakazato and et al. BioCompass: a novel functional inference tool that utilizes MeSH hierarchy to analyze groups of genes. *In Silico Biol.*, 8(1): 53-61, 2007.
- [4] T. Nakazato and et al. Nucleic Acids Res. Gendoo: functional profiling of gene and disease features using MeSH vocabulary., 37: W166-W169, 2009.
- [5] D. J. Saurin and et al. GeneMeSH: a web-based microarray analysis tool for relating differentially expressed genes to MeSH terms. *BMC Bioinformatics*, 11: 166, 2010.
- [6] M. A. Sartor and et al. Metab2MeSH: annotating compounds with medical subject headings. *Bioinformatics*, 28(10): 1408-1410, 2012.