# An Introduction to GSEABase

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24 September, 2007

```
> ## FIXME: <adjMat> adjacency matrix -- color w. +/- 1
> ## FIXME: limma topTable --> GeneColorSet
> ## w. verbose=TRUE
> library(GSEABase)
> library(hgu95av2.db)
> library(GO.db)
```

### 1 GeneSet

[5] "31444\_s\_at" "31445\_at"

A GeneSet stores a set of related gene identifiers. Important components of the gene set are a vector of identifiers, general descriptive information about the set, and information about how the gene set was constructed. To construct a gene set, use GeneSet. For example, to create a gene set from the identifiers in a subset of the sample ExpressionSet in the Biobase package (see Biobase documentation for a description of expression sets) use

```
> data(sample.ExpressionSet) # from Biobase
> egs <- GeneSet(sample.ExpressionSet[201:250,], setName="Sample")
> egs
setName: Sample
geneIds: 31440_at, 31441_at, ..., 31489_at (total: 50)
geneIdType: Annotation (hgu95av2)
collectionType: ExpressionSet
details: use 'details(object)'

Each gene set may have a name. The gene set contains 50 identifiers ('genes')
from the expression set. These are accessible using geneIds, e.g.,
> head(geneIds(egs))
[1] "31440_at" "31441_at" "31442_at" "31443_at"
```

The gene set records that the identifiers are probeset names from the annotation package hgu95av2, and that the source of the gene set was an expression set. Additional details are available:

```
> details(egs)
setName: Sample
geneIds: 31440_at, 31441_at, ..., 31489_at (total: 50)
geneIdType: Annotation (hgu95av2)
collectionType: ExpressionSet
setIdentifier: malbec1:24416:Mon Oct 30 18:47:14 2017:1
description: Smoking-Cancer Experiment
  (longDescription available)
organism: Homo sapiens
pubMedIds:
urls: www.lab.not.exist
contributor: Pierre Fermat
setVersion: 0.0.1
creationDate:
The set identifier, set version, and creation date provide mechanisms for carefully
```

curating gene sets. Additional information is automatically copied from the expression set used to create egs.

```
## FIXME: GeneSet(AnnotationIdentifier("hgu95av2")) --> non-empty
## FIXME: GeneSet(AnnotationIdentifier("hgu95av2"),
## collectionType=GOCollection()) filters on GOCollection (or KEGG)
```

View additional methods for creating gene sets with:

> showMethods("GeneSet", inherited=FALSE)

```
Function: GeneSet (package GSEABase)
type="BroadCollection"
type="ExpressionSet"
type="GOCollection"
type="GeneIdentifierType"
type="character"
type="missing"
```

These are described on the GeneSet help page.

The identifier type of gene sets created from expression sets is Annotation-Identifier. Additional predefined identifiers are available:

> names(slot(getClass("GeneIdentifierType"), "subclasses"))

```
[1] "NullIdentifier"
                            "EnzymeIdentifier"
[3] "GenenameIdentifier"
                            "RefseqIdentifier"
[5] "SymbolIdentifier"
                            "UnigeneIdentifier"
                            "ENSEMBLIdentifier"
[7] "UniprotIdentifier"
[9] "AnnotationIdentifier" "EntrezIdentifier"
[11] "GOAllFrameIdentifier" "KEGGFrameIdentifier"
```

It is possible to map between identifier types (provided the corresponding map in the annotation package exists):

```
> mapIdentifiers(egs, EntrezIdentifier())
```

```
setName: Sample
geneIds: 6932, 643332, ..., 4287 (total: 31)
geneIdType: EntrezId (hgu95av2)
collectionType: ExpressionSet
details: use 'details(object)'
```

mapIdentifiers consults the gene set to determine that annotation (probeset) identifiers are to be converted to Entrez IDs using the hgu95av2ENTREZID environment in the hgu95av2.db (or hgu95av2) package. mapIdentifiers can automatically determine many of the common maps; it is also possible to provide as a third argument an environment containing an arbitrary map. Use the verbose argument of mapIdentifiers to be informed when the map between identifier types is not 1:1.

A gene set can be created with different types of identifier, e.g., to create a gene set with Entrez IDs, use

```
> library(annotate) # getEG
> eids <- unique(getEG(geneIds(egs), "hgu95av2"))
> eids <- eids[!is.na(eids)]
> GeneSet(EntrezIdentifier(), geneIds=as.character(eids))
setName: NA
geneIds: 6932, 643332, ..., 4287 (total: 31)
geneIdType: EntrezId
collectionType: Null
details: use 'details(object)'
```

The collectionType of a gene set provides additional information about a gene set. Available collection types are

> names(slot(getClass("CollectionType"), "subclasses"))

```
[1] "NullCollection" "ExpressionSetCollection"
[3] "ComputedCollection" "CollectionIdType"
[5] "BroadCollection" "KEGGCollection"
[7] "OMIMCollection" "PMIDCollection"
[9] "ChrCollection" "ChrlocCollection"
[11] "MapCollection" "PfamCollection"
[13] "PrositeCollection" "GOCollection"
[15] "OBOCollection"
```

Collection types are most important when creating gene sets. For instance, the *GOCollection* class allows creation of gene sets based on gene ontology (GO) terms. The following command creates a gene set from two terms, including all genes with a particular evidence code.

This creates a gene set by consulting an object in the GO.db package. A gene set created from an expression set, and with collection type GOCollection() consults the appropriate environment in the annotation package associated with the expression set.

Gene sets encoded in XML following the schema and conventions of the Broad Institute can be read into  ${\sf R}$  as follows:

```
> f1 <- system.file("extdata", "Broad1.xml", package="GSEABase")
> bgs <- GeneSet(BroadCollection(), urls=f1)
> bgs

setName: chr16q24
geneIds: GALNS, C16ORF44, ..., TRAPPC2L (total: 129)
geneIdType: Symbol
collectionType: Broad
  bcCategory: c1 (Positional)
  bcSubCategory: NA
details: use 'details(object)'
```

The example above uses a local file, but the source for the gene set might also be a web address accessible via the http:// protocol. The file name is added to the url of the gene set. The set name and category of the Broad collection indicate that the gene set contains gene symbols from band 24 of the q arm of chromosome 16. The probe sets in chip hgu95av2 corresponding to these symbols can be determined by mapping identifiers

```
> bgs1 <- mapIdentifiers(bgs, AnnotationIdentifier("hgu95av2"))
> bgs1

setName: chr16q24
geneIds: 32100_r_at, 32101_at, ..., 35807_at (total: 36)
geneIdType: Annotation (hgu95av2)
collectionType: Broad
  bcCategory: c1 (Positional)
```

```
bcSubCategory: NA
details: use 'details(object)'
   Subsetting creates sets with just the symbols identified. Subsetting can use
indices or symbol names.
> bgs[1:5]
setName: chr16q24
geneIds: GALNS, C160RF44, ..., L0C646365 (total: 5)
geneIdType: Symbol
collectionType: Broad
  bcCategory: c1 (Positional)
  bcSubCategory: NA
details: use 'details(object)'
> bgs[c("GALNS", "LOC646365")]
setName: chr16q24
geneIds: GALNS, LOC646365 (total: 2)
geneIdType: Symbol
collectionType: Broad
  bcCategory: c1 (Positional)
  bcSubCategory: NA
details: use 'details(object)'
  Logical operations provide a convenient way to identify genes with particular
properties. For instance, the intersection
> egs & bgs1
setName: (Sample & chr16q24)
geneIds: (total: 0)
geneIdType: Annotation (hgu95av2)
collectionType: Computed
details: use 'details(object)'
is empty (note that the identifiers in the two sets were of the same type), indi-
cating that none of the identifiers in egs are on 16q24. Additional operations
on sets include union (performed with |) and difference (setdiff).
   Methods exist to directly subset expression sets using gene sets
> sample.ExpressionSet[bgs,]
ExpressionSet (storageMode: lockedEnvironment)
assayData: 1 features, 26 samples
  element names: exprs, se.exprs
protocolData: none
phenoData
```

sampleNames: A B ... Z (26 total)

varLabels: sex type score
varMetadata: labelDescription

featureData: none

experimentData: use 'experimentData(object)'

Annotation: hgu95av2

Remember that sample.ExpressionSet contains just 500 probe sets, so the small size of the subset is not surprising. Note also that subsetting required mapping of symbol identifiers in bgs to *AnnotationIdentifiers*; this map used the annotation information in the expression set, and is not necessarily 1:1.

## $2 \quad Gene Color Set$

A GeneColorSet is a gene set with "coloring" to indicate how features of genes and phenotypes are associated. The following sample data describes how changes in expression levels of several genes (with Entrez and Symbol names) influence cisplatin resistance phenotype.

#### > tbl

	Entrez.ID	<pre>Gene.Symbol</pre>	Expression.level	Phenotype.response
1	1244	ABCC2	Increase	Resistant
2	538	ATP7A	Increase	Resistant
3	540	ATP7B	Increase	Resistant
4	9961	MVP	Increase	Resistant
5	7507	XPA	Increase	Resistant
6	2067	ERCC1	Increase	Resistant
7	672	BRCA1	Increase	Resistant
8	3725	JUN	Increase	Resistant
9	2730	GCLM	Increase	Resistant

Note that three different aspects of data influence coloring: the phenotype under consideration (cisplatin resistance), whether expression responses refer to increasing or decreasing levels of gene expression, and whether the phenotypic response represents greater resistance or sensitivity to cisplatin. Here is the resulting gene color set:

```
setName: A color set
geneIds: 1244, 538, ..., 2730 (total: 9)
geneIdType: EntrezId
collectionType: Null
phenotype: Cisplatin resistance
geneColor: Increase, Increase, ..., Increase
  levels: Increase
phenotypeColor: Resistant, Resistant, ..., Resistant
  levels: Resistant
details: use 'details(object)'
```

Gene color sets can be used in the same way as gene sets, e.g., for subsetting expression sets (provided the map between identifiers is 1:1, so that coloring corresponding to identifiers can be determined). The coloring method allows access to the coloring information with a data frame interface; phenotype, geneColor and phenotypeColor provide additional accessors.

#### 3 GeneSetCollection

A GeneSetCollection is a collection of gene sets. Sets in the collection must have distinct setNames, but can be a mix of GeneSet and GeneColorSet. Two convenient ways to create a gene set collection are by specifying a source of identifiers (e.g., an ExpressionSet or AnnotationIdentifier) and how the identifiers are to be induced into sets (e.g., by consulting the GO or KEGG ontologies):

```
> gsc <- GeneSetCollection(sample.ExpressionSet[201:250,], setType=GOCollection())</pre>
> gsc
GeneSetCollection
  names: GO:0000122, GO:0000186, ..., GO:1990380 (297 total)
  unique identifiers: 31440_at, 31468_f_at, ..., 31480_f_at (32 total)
  types in collection:
    geneIdType: AnnotationIdentifier (1 total)
    collectionType: GOCollection (1 total)
> gsc[["GO:0005737"]]
setName: GO:0005737
geneIds: 31450_s_at, 31451_at, ..., 31489_at (total: 6)
geneIdType: Annotation (hgu95av2)
collectionType: GO
  ids: GO:0005737 (1 total)
  evidenceCode: EXP IDA IPI IMP IGI IEP ISS ISO ISA ISM IGC IBA IBD IKR IRD RCA TAS NAS IC N
  ontology: CC MF BP
details: use 'details(object)'
```

In this example, the annotation identifiers of the sample expression set are organized into gene sets based on their presence in GO pathways. Providing arguments such as evidenceCode to *GOCollection* act to select just those pathways satisfying the GO collection constraint:

```
> GeneSetCollection(sample.ExpressionSet[201:300,],
                     setType=GOCollection(evidenceCode="IMP"))
GeneSetCollection
  names: GO:0000122, GO:0000226, ..., GO:1902282 (102 total)
  unique identifiers: 31520_at, 31489_at, ..., 31442_at (22 total)
  types in collection:
    geneIdType: AnnotationIdentifier (1 total)
    collectionType: GOCollection (1 total)
Sets in the collection are named after the GO terms, and can be accessed by
numeric index or name.
   A file or url containing several gene sets defined by Broad XML can be used
to create a gene set collection, e.g.,
    ## FIXME: BroadCollection default to paste("c", 1:4, sep="")
    ## FIXME: GeneSetCollection(BroadCollection(), urls=f1); filters on bcCategory
> fl <- system.file("extdata", "Broad.xml", package="GSEABase")</pre>
> gss <- getBroadSets(f1)</pre>
> gss
{\tt GeneSetCollection}
 names: chr5q23, chr16q24 (2 total)
  unique identifiers: ZNF474, CCDC100, ..., TRAPPC2L (215 total)
  types in collection:
    geneIdType: SymbolIdentifier (1 total)
    collectionType: BroadCollection (1 total)
> names(gss)
[1] "chr5q23"
               "chr16q24"
   Identifiers within a gene set collection can be mapped to a common type
(provided maps are available) with, for example,
> gsc <- mapIdentifiers(gsc, EntrezIdentifier())</pre>
> gsc
GeneSetCollection
 names: GO:0000122, GO:0000186, ..., GO:1990380 (297 total)
  unique identifiers: 6932, 2911, ..., 4111 (27 total)
  types in collection:
    geneIdType: EntrezIdentifier (1 total)
```

collectionType: GOCollection (1 total)

```
> gsc[["GO:0005737"]]
setName: G0:0005737
geneIds: 6014, 392, ..., 4287 (total: 6)
geneIdType: EntrezId (hgu95av2)
collectionType: GO
  ids: GO:0005737 (1 total)
  evidenceCode: EXP IDA IPI IMP IGI IEP ISS ISO ISA ISM IGC IBA IBD IKR IRD RCA TAS NAS IC N
  ontology: CC MF BP
details: use 'details(object)'
   A convenient way to visualize a GeneSetCollection is with the Reporting-
Tools package.
> ## 'interesting' gene sets
> idx <- sapply(gsc, function(x) length(geneIds(x))) > 2
> library(ReportingTools)
> gscReport <- HTMLReport(</pre>
      shortName="gsc_example",
      title="GSEABase Vignette GeneSetCollection",
      basePath=tempdir())
> publish(gsc[idx], gscReport, annotation.db="org.Hs.eg")
> url <- finish(gscReport)</pre>
The report can be viewed with
> browseURL(url)
  This concludes a brief tour of gene sets and gene set collections available in
```

the GSEABase package.