AnnotationDbi: Introduction To Bioconductor Annotation Packages

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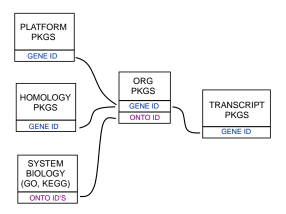


Figure 1: Annotation Packages: the big picture

Bioconductor provides extensive annotation resources. These can be gene centric, or genome centric. Annotations can be provided in packages curated by Bioconductor, or obtained from web-based resources. This vignette is primarily concerned with describing the annotation resources that are available as packages. More advanced users who wish to learn about how to make new annotation packages should see the vignette titled "Creating select Interfaces for custom Annotation resources" from the AnnotationForge package.

Gene centric Annotation Dbi packages include:

• Organism level: e.g. org.Mm.eg.db.

- Platform level: e.g. hgu133plus2.db, hgu133plus2.probes, hgu133plus2.cdf.
- Homology level: e.g. hom.Dm.inp.db.
- System-biology level: GO.db

Genome centric GenomicFeatures packages include

- Transcriptome level: e.g. TxDb.Hsapiens.UCSC.hg19.knownGene
- Generic genome features: Can generate via GenomicFeatures

One web-based resource accesses biomart, via the biomaRt package:

 Query web-based 'biomart' resource for genes, sequence, SNPs, and etc.

The most popular annotation packages have been modified so that they can make use of a new set of methods to more easily access their contents. These four methods are named: columns, keytypes, keys and select. And they are described in this vignette. They can currently be used with all chip, organism, and *TranscriptDb* packages along with the popular GO.db package.

For the older less popular packages, there are still conventient ways to retrieve the data. The *How to use bimaps from the ".db" annotation packages* vignette in the *AnnotationDbi* package is a key reference for learnign about how to use bimap objects.

Finally, all of the '.db' (and most other *Bioconductor* annotation packages) are updated every 6 months corresponding to each release of *Bioconductor*. Exceptions are made for packages where the actual resources that the packages are based on have not themselves been updated.

0.1 AnnotationDb objects and the select method

As previously mentioned, a new set of methods have been added that allow a simpler way of extracting identifier based annotations. All the annotation packages that support these new methods expose an object named exactly the same as the package itself. These objects are collectively called *AnntoationDb* objects for the class that they all inherit from. The more specific classes (the ones that you will actually see in the wild) have names like *OrgDb*, *ChipDb* or *TranscriptDb* objects. These names correspond to the kind of package (and underlying schema) being represented. The methods that can be applied to all of these objects are columns, keys, keytypes and select.

0.2 ChipDb objects and the select method

An extremely common kind of Annotation package is the so called platform based or chip based package type. This package is intended to make the manufacturer labels for a series of probes or probesets to a wide range of gene-based features. A package of this kind will load an ChipDb object. Below is a set of examples to show how you might use the standard 4 methods to interact with an object of this type.

First we need to load the package:

R> library(hgu95av2.db)

If we list the contents of this package, we can see that one of the many things loaded is an object named after the package "hgu95av2.db":

R> ls("package:hgu95av2.db")

[1]	"hgu95av2"	"hgu95av2.db"
[3]	"hgu95av2ACCNUM"	"hgu95av2ALIAS2PR0BE"
[5]	"hgu95av2CHR"	"hgu95av2CHRLENGTHS"
[7]	"hgu95av2CHRLOC"	"hgu95av2CHRLOCEND"
[9]	J	· ·
	"hgu95av2ENSEMBL"	"hgu95av2ENSEMBL2PR0BE"
[11]	"hgu95av2ENTREZID"	"hgu95av2ENZYME"
[13]	"hgu95av2ENZYME2PR0BE"	"hgu95av2GENENAME"
[15]	"hgu95av2G0"	"hgu95av2G02ALLPR0BES"
[17]	"hgu95av2G02PR0BE"	"hgu95av2MAP"
[19]	"hgu95av2MAPCOUNTS"	"hgu95av20MIM"
[21]	"hgu95av2ORGANISM"	"hgu95av20RGPKG"
[23]	"hgu95av2PATH"	"hgu95av2PATH2PR0BE"
[25]	"hgu95av2PFAM"	"hgu95av2PMID"
[27]	"hgu95av2PMID2PROBE"	"hgu95av2PROSITE"
[29]	"hgu95av2REFSEQ"	"hgu95av2SYMBOL"
[31]	"hgu95av2UNIGENE"	"hgu95av2UNIPROT"
[33]	"hgu95av2_dbInfo"	"hgu95av2_dbconn"
[35]	"hgu95av2_dbfile"	"hgu95av2_dbschema"

We can look at this object to learn more about it:

R> hgu95av2.db

ChipDb object:

| DBSCHEMAVERSION: 2.1

- | Db type: ChipDb
- | Supporting package: AnnotationDbi
- | DBSCHEMA: HUMANCHIP_DB | ORGANISM: Homo sapiens
- | SPECIES: Human
- | MANUFACTURER: Affymetrix
- | CHIPNAME: Human Genome U95 Set
- | MANUFACTURERURL: http://www.affymetrix.com/support/technical/byproduct.affx?product
- | EGSOURCEDATE: 2013-Sep12 | EGSOURCENAME: Entrez Gene
- | EGSOURCEURL: ftp://ftp.ncbi.nlm.nih.gov/gene/DATA
- | CENTRALID: ENTREZID
- | TAXID: 9606
- | GOSOURCENAME: Gene Ontology
- | GOSOURCEURL: ftp://ftp.geneontology.org/pub/go/godatabase/archive/latest-lite/
- | GOSOURCEDATE: 20130907 | GOEGSOURCEDATE: 2013-Sep12 | GOEGSOURCENAME: Entrez Gene
- | GOEGSOURCEURL: ftp://ftp.ncbi.nlm.nih.gov/gene/DATA
- | KEGGSOURCENAME: KEGG GENOME
- | KEGGSOURCEURL: ftp://ftp.genome.jp/pub/kegg/genomes
- | KEGGSOURCEDATE: 2011-Mar15
- | GPSOURCENAME: UCSC Genome Bioinformatics (Homo sapiens) | GPSOURCEURL: ftp://hgdownload.cse.ucsc.edu/goldenPath/hg19
- | GPSOURCEDATE: 2010-Mar22 | UPSOURCENAME: Uniprot
- | UPSOURCEURL: http://www.UniProt.org/ | UPSOURCEDATE: Mon Sep 16 12:05:19 2013
- | ENSOURCEDATE: 2013-Sep3 | ENSOURCENAME: Ensembl
- | ENSOURCEURL: ftp://ftp.ensembl.org/pub/current_fasta

If we want to know what kinds of data are retriveable via select, then we should use the columns method like this:

R> columns(hgu95av2.db)

[1]	"PROBEID"	"ENTREZID"	"PFAM"	"IPI"
[5]	"PROSITE"	"ACCNUM"	"ALIAS"	"CHR"
[9]	"CHRLOC"	"CHRLOCEND"	"ENZYME"	"MAP"

[13]	"PATH"	"PMID"	"REFSEQ"	"SYMBOL"
[17]	"UNIGENE"	"ENSEMBL"	"ENSEMBLPROT"	"ENSEMBLTRANS"
[21]	"GENENAME"	"UNIPROT"	"GO"	"EVIDENCE"
[25]	"ONTOLOGY"	"GOALL"	"EVIDENCEALL"	"ONTOLOGYALL"
[29]	"OMIM"	"UCSCKG"		

If we are further curious to know more about those values for columns, we can consult the help pages. Asking about any of these values will pull up a manual page describing the different fields and what they mean.

R> help("SYMBOL")

If we are curious about what kinds of fields we could potentiall use as keys to query the database, we can use the **keytypes** method. In a perfect world, this method will return values very similar to what was returned by **columns**, but in reality, some kinds of values make poor keys and so this list is often shorter.

R> keytypes(hgu95av2.db)

[1]	"ENTREZID"	"PFAM"	"IPI"	"PROSITE"
[5]	"ACCNUM"	"ALIAS"	"CHR"	"CHRLOC"
[9]	"CHRLOCEND"	"ENZYME"	"MAP"	"PATH"
[13]	"PMID"	"REFSEQ"	"SYMBOL"	"UNIGENE"
[17]	"ENSEMBL"	"ENSEMBLPROT"	"ENSEMBLTRANS"	"GENENAME"
[21]	"UNIPROT"	"GO"	"EVIDENCE"	"ONTOLOGY"
[25]	"GOALL"	"EVIDENCEALL"	"ONTOLOGYALL"	"PROBEID"
[29]	"OMIM"	"UCSCKG"		

If we want to extract some sample keys of a particular type, we can use the keys method.

```
R> head(keys(hgu95av2.db, keytype="SYMBOL"))
```

```
[1] "A1BG" "A2M" "A2MP1" "NAT1" "NAT2" "NATP"
```

And finally, if we have some keys, we can use **select** to extract them. By simply using appropriate argument values with select we can specify what keys we want to look up values for (keys), what we want returned back (columns) and the type of keys that we are passing in (keytype)

```
R> #1st get some example keys
R> k <- head(keys(hgu95av2.db,keytype="PROBEID"))</pre>
R> # then call select
R> select(hgu95av2.db, keys=k, columns=c("SYMBOL", "GENENAME"), keytype="PROBEID")
    PROBEID
             SYMBOL
    1000 at
1
              MAPK3
2
    1001_at
               TIE1
3 1002_f_at CYP2C19
4 1003_s_at
              CXCR5
    1004_at
5
              CXCR5
6
    1005_at
              DUSP1
                                                           GENENAME
1
                                mitogen-activated protein kinase 3
2 tyrosine kinase with immunoglobulin-like and EGF-like domains 1
3
           cytochrome P450, family 2, subfamily C, polypeptide 19
4
                                chemokine (C-X-C motif) receptor 5
5
                                chemokine (C-X-C motif) receptor 5
6
                                    dual specificity phosphatase 1
```

And as you can see, when you call the code above, select will try to return a data.frame with all the things you asked for matched up to each other.

0.3 OrgDb objects and the select method

An organism level package (an 'org' package) uses a central gene identifier (e.g. Entrez Gene id) and contains mappings between this identifier and other kinds of identifiers (e.g. GenBank or Uniprot accession number, RefSeq id, etc.). The name of an org package is always of the form org.<Ab>.<id>.db (e.g. org.Sc.sgd.db) where <Ab> is a 2-letter abbreviation of the organism (e.g. Sc for Saccharomyces cerevisiae) and <id> is an abbreviation (in lower-case) describing the type of central identifier (e.g. sgd for gene identifiers assigned by the Saccharomyces Genome Database, or eg for Entrez Gene ids).

Just as the chip packages load a ChipDb object, the org packages will load a OrgDb object. The following exercise should acquaint you with the use of these methods in the context of an organism package.

Exercise 1

Display the OrgDb object for the org.Hs.eg.db package.

Use the columns method to discover which sorts of annotations can be extracted from it. Is this the same as the result from the keytypes method? Use the keytypes method to find out.

Finally, use the keys method to extract UNIPROT identifiers and then pass those keys in to the select method in such a way that you extract the gene symbol and KEGG pathway information for each. Use the help system as needed to learn which values to pass in to columns in order to achieve this.

Solution:

```
R> library(org.Hs.eg.db)
R> columns(org.Hs.eg.db)
```

```
"PFAM"
                                     "IPI"
 [1] "ENTREZID"
                                                      "PROSITE"
 [5] "ACCNUM"
                     "ALIAS"
                                     "CHR"
                                                      "CHRLOC"
                                     "MAP"
 [9] "CHRLOCEND"
                     "ENZYME"
                                                      "PATH"
[13] "PMID"
                     "REFSEQ"
                                     "SYMBOL"
                                                      "UNIGENE"
[17] "ENSEMBL"
                     "ENSEMBLPROT"
                                     "ENSEMBLTRANS" "GENENAME"
[21] "UNIPROT"
                     "GO"
                                     "EVIDENCE"
                                                      "ONTOLOGY"
[25] "GOALL"
                     "EVIDENCEALL"
                                     "ONTOLOGYALL"
                                                      "MIMO"
[29] "UCSCKG"
```

R> help("SYMBOL") ## for explanation of these columns and keytypes values

```
R> keytypes(org.Hs.eg.db)
```

```
[1] "ENTREZID"
                      "PFAM"
                                      "IPI"
                                                      "PROSITE"
 [5] "ACCNUM"
                      "ALIAS"
                                      "CHR"
                                                      "CHRLOC"
 [9] "CHRLOCEND"
                      "ENZYME"
                                      "MAP"
                                                      "PATH"
[13] "PMID"
                      "REFSEQ"
                                      "SYMBOL"
                                                      "UNIGENE"
[17] "ENSEMBL"
                      "ENSEMBLPROT"
                                      "ENSEMBLTRANS" "GENENAME"
[21] "UNIPROT"
                      "GO"
                                      "EVIDENCE"
                                                      "ONTOLOGY"
[25] "GOALL"
                      "EVIDENCEALL"
                                      "ONTOLOGYALL"
                                                      "MIMO"
[29] "UCSCKG"
R> uniKeys <- head(keys(org.Hs.eg.db, keytype="UNIPROT"))</pre>
```

```
R> cols <- c("SYMBOL", "PATH")
```

R> select(org.Hs.eg.db, keys=uniKeys, columns=cols, keytype="UNIPROT")

```
UNIPROT SYMBOL PATH
1 PO4217 A1BG <NA>
```

```
2
    P01023
              A2M 04610
3
    F5H5R8
             NAT1 00232
4
    F5H5R8
             NAT1 00983
5
    F5H5R8
             NAT1 01100
6
    P18440
             NAT1 00232
7
    P18440
             NAT1 00983
8
    P18440
             NAT1 01100
9
    Q400J6
             NAT1 00232
10
   Q400J6
             NAT1 00983
    Q400J6
             NAT1 01100
11
12
   A4Z6T7
             NAT2 00232
   A4Z6T7
13
             NAT2 00983
14 A4Z6T7
             NAT2 01100
```

So how could you use select to annotate your results? This next exercise should hlep you to understand how that should generally work.

Exercise 2

Please run the following code snippet (which will load a fake data result that I have provided for the purposes of illustration):

```
R> load(system.file("extdata", "resultTable.Rda", package="AnnotationDbi"))
R> head(resultTable)
```

```
logFC LR.statistic
                                                  PValue
                                                                  FDR
             logConc
                                   378.0732 3.269307e-84 2.613484e-80
100418920 -9.639471 -4.679498
100419779 -10.638865 -4.264830
                                   291.1028 2.859424e-65 1.142912e-61
100271867 -11.448981 -4.009603
                                   222.3653 2.757135e-50 7.346846e-47
100287169 -11.026699 -3.486593
                                   206.7771 6.934967e-47 1.385953e-43
100287735 -11.036862 3.064980
                                   204.1235 2.630432e-46 4.205535e-43
100421986 -12.276297 -4.695736
                                   190.5368 2.427556e-43 3.234314e-40
```

The rownames of this table happen to provide entrez gene identifiers for each row (for human). Find the gene symbol and gene name for each of the rows in resultTable and then use the merge method to attach those annotations to it.

Solution:

R> resultTable <- merge(resultTable, annots, by.x=0, by.y="ENTREZID") R> head(resultTable) logFC LR.statistic Row.names logConc **PValue FDR** 1 100127888 -10.57050 182.8937 1.131473e-41 1.130624e-38 2.758937 2 100131223 -12.37808 -4.654318 179.2331 7.126423e-41 6.329847e-38 3 100271381 -12.06340 3.511937 188.4824 6.817155e-43 7.785191e-40 222.3653 2.757135e-50 7.346846e-47 4 100271867 -11.44898 -4.009603 5 100287169 -11.02670 -3.486593 206.7771 6.934967e-47 1.385953e-43 6 100287735 -11.03686 3.064980 204.1235 2.630432e-46 4.205535e-43 SYMBOL **GENENAME** 1 LOC100127888 uncharacterized LOC100127888

```
1 LOC100127888 uncharacterized LOC100127888
2 LOC100131223 ADP-ribosylation factor-like 8B pseudogene
3 RPS28P8 ribosomal protein S28 pseudogene 8
4 MPVQTL1 Mean platelet volume QTL1
```

5 <NA> <NA> <NA> 6 TTTY13B testis-specific transcript, Y-linked 13B

0.4 Using select with GO.db

When you load the GO.db package, a *GODb* object is also loaded. This allows you to use the columns, keys, keytypes and select methods on the contents of the GO ontology. So if for example, you had a few GO IDs and wanted to know more about it, you could do it like this:

A cellular process that results in the biosynthesis of constituent macromolecu 2 Interacting selectively and non-covalently with any protein or protein complex (a c

0.5 Using select with TranscriptDb packages

A *TranscriptDb* package (a 'TxDb' package) connects a set of genomic coordinates to various transcript oriented features. The package can also contain Identifiers to features such as genes and transcripts, and the internal

schema describes the relationships between these different elements. All TranscriptDb containing packages follow a specific naming scheme that tells where the data came from as well as which build of the genome it comes from.

Exercise 3

Display the TranscriptDb object for the TxDb.Hsapiens.UCSC.hg19.knownGene package.

As before, use the columns and keytypes methods to discover which sorts of annotations can be extracted from it.

Use the keys method to extract just a few gene identifiers and then pass those keys in to the select method in such a way that you extract the transcript ids and transcript starts for each.

Solution:

```
R> library(TxDb.Hsapiens.UCSC.hg19.knownGene)
R> txdb <- TxDb. Hsapiens. UCSC. hg19. knownGene
R> txdb
TranscriptDb object:
| Db type: TranscriptDb
| Supporting package: GenomicFeatures
| Data source: UCSC
| Genome: hg19
| Organism: Homo sapiens
| UCSC Table: knownGene
Resource URL: http://genome.ucsc.edu/
| Type of Gene ID: Entrez Gene ID
| Full dataset: yes
| miRBase build ID: GRCh37
| transcript_nrow: 82960
| exon_nrow: 289969
| cds_nrow: 237533
| Db created by: GenomicFeatures package from Bioconductor
| Creation time: 2013-09-19 00:15:15 -0700 (Thu, 19 Sep 2013)
| GenomicFeatures version at creation time: 1.13.40
RSQLite version at creation time: 0.11.4
| DBSCHEMAVERSION: 1.0
R> columns(txdb)
```

```
[1] "CDSID"
                   "CDSNAME"
                                 "CDSCHROM"
                                               "CDSSTRAND"
                                                            "CDSSTART"
 [6] "CDSEND"
                   "EXONID"
                                 "EXONNAME"
                                               "EXONCHROM"
                                                            "EXONSTRAND"
                   "EXONEND"
[11] "EXONSTART"
                                 "GENEID"
                                               "TXID"
                                                            "EXONRANK"
[16] "TXNAME"
                   "TXCHROM"
                                 "TXSTRAND"
                                                            "TXEND"
                                               "TXSTART"
R> keytypes(txdb)
[1] "GENEID"
                "TXID"
                           "TXNAME"
                                       "EXONID"
                                                   "EXONNAME" "CDSID"
[7] "CDSNAME"
R> keys <- head(keys(txdb, keytype="GENEID"))</pre>
R> cols <- c("TXID", "TXSTART")</pre>
R> select(txdb, keys=keys, columns=cols, keytype="GENEID")
      GENEID TXID
                      TXSTART
           1 70455
1
                     58858172
2
           1 70456 58859832
          10 31944 18248755
3
4
         100 72132
                     43248163
5
        1000 65378
                     25530930
6
        1000 65379
                     25530930
7
       10000 7895 243651535
8
       10000 7896 243663021
9
       10000 7897 243663021
10 100008586 75890
                     49217763
```

As is widely known, in addition to providing access via the select method, TranscriptDb objects also provide access via the more familiar transcripts, exons, cds, transcriptsBy, exonsBy and cdsBy methods. For those who do not yet know about these other methods, more can be learned by seeing the vignette called: $Making\ and\ Utilizing\ TranscriptDb\ Objects$ in the $GenomicFeatures\ package$.

The version number of R and packages loaded for generating the vignette were:

```
R version 3.0.2 (2013-09-25)
Platform: i386-w64-mingw32/i386 (32-bit)
```

R.>

locale:

- [1] LC_COLLATE=C
- [2] LC_CTYPE=English_United States.1252
- [3] LC_MONETARY=English_United States.1252
- [4] LC_NUMERIC=C
- [5] LC_TIME=English_United States.1252

attached base packages:

- [1] parallel stats graphics grDevices utils datasets
- [7] methods base

other attached packages:

- [1] TxDb.Hsapiens.UCSC.hg19.knownGene_2.10.1
- [2] GenomicFeatures_1.14.0
- [3] GenomicRanges_1.14.0
- [4] XVector_0.2.0
- [5] IRanges_1.20.0
- [6] GO.db_2.10.1
- [7] hgu95av2.db_2.10.1
- [8] AnnotationForge_1.4.0
- [9] org.Hs.eg.db_2.10.1
- [10] RSQLite_0.11.4
- [11] DBI_0.2-7
- [12] AnnotationDbi_1.24.0
- [13] Biobase_2.22.0
- [14] BiocGenerics_0.8.0

loaded via a namespace (and not attached):

- [1] BSgenome_1.30.0 Biostrings_2.30.0 RCurl_1.95-4.1
- [4] Rsamtools_1.14.0 XML_3.98-1.1 biomaRt_2.18.0
- [7] bitops_1.0-6 rtracklayer_1.22.0 stats4_3.0.2
- [10] tools_3.0.2 zlibbioc_1.8.0