Making and Utilizing TranscriptDb Objects

Marc Carlson Patrick Aboyoun Hervé Pagès Seth Falcon Martin Morgan

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1 Introduction

The *GenomicFeatures* package retrieves and manages transcript-related features from the UCSC Genome Bioinformatics¹ and BioMart² data resources. The package is useful for ChIP-chip, ChIP-seq, and RNA-seq analyses.

> library("GenomicFeatures")

2 Transcript Metadata

2.1 TranscriptDb Objects

The GenomicFeatures package uses TranscriptDb objects to store transcript metadata. This class maps the 5' and 3' untranslated regions (UTRs), protein coding sequences (CDSs) and exons for a set of mRNA transcripts to their associated genome. TranscriptDb objects also have accessors functions to allow such features to be retrieved individually or grouped together in a way that reflects the underlying biology.

All *TranscriptDb* objects are backed by a SQLite database that manages genomic locations and the relationships between pre-processed mRNA transcripts, exons, protein coding sequences, and their related gene identifiers.

2.2 Creating New *TranscriptDb* Objects

The GenomicFeatures package provides functions to create TranscriptDb objects based on data downloaded from UCSC Genome Bioinformatics or BioMart. The following subsections demonstrate the use of these functions.

¹http://genome.ucsc.edu/

²http://www.biomart.org/

There is also support for creating TranscriptDb objects from custom data sources using makeTranscriptDb; see the help page for this function for details.

2.2.1 Using makeTranscriptDbFromUCSC

The function makeTranscriptDbFromUCSC downloads UCSC Genome Bioinformatics transcript tables (e.g. "knownGene", "refGene", "ensGene") for a genome build (e.g. "mm9", "hg19"). Use the supportedUCSCtables utility function to get the list of supported tables.

> supportedUCSCtables()[1:4,]

		${\tt track}$	subtrack
knownGene	UCSC	Genes	<na></na>
knownGeneOld3	Old UCSC	Genes	<na></na>
${\tt wgEncodeGencodeManualV3}$	Gencode	Genes	Genecode Manual
wgEncodeGencodeAutoV3	Gencode	Genes	Genecode Auto

> mm9KG <- makeTranscriptDbFromUCSC(genome = "mm9", tablename = "knownGene")

The function makeTranscriptDbFromUCSC also takes an important argument called circ_seqs to label which chromosomes are circular. The argument is a character vector of strings that correspond to the circular chromosomes (as labeled by the source). To discover what the source calls their chromosomes, use the getChromInfoFromUCSC function to list them. By default, there is a supplied character vector that will attempt to label all the mitochondrial chromosomes as circular by matching to them. This is the DEFAULT_CIRC_SEQS vector. It contains strings that usually correspond to mitochondrial chromosomes. Once the database has been generated with the circular chromosomes tagged in this way, all subsequent analysis of these chromosomes will be able to consider their circularity for analysis. So it is important for the user to make sure that they pass in the correct strings to the circ_seqs argument to ensure that the correct sequences are tagged as circular by the database.

> head(getChromInfoFromUCSC("hg19"))

chrom length 1 chr1 249250621 2 chr2 243199373

```
3 chr3 198022430
4 chr4 191154276
5 chr5 180915260
6 chr6 171115067
```

2.2.2 Using makeTranscriptDbFromBiomart

Retrieve data from BioMart by specifying the mart and the data set to the makeTranscriptDbFromBiomart function (not all BioMart data sets are currently supported):

As with the makeTranscriptDbFromUCSC function, the makeTranscriptDbFromBiomart function also has a circ_seqs argument that will default to using the contents of the DEFAULT_CIRC_SEQS vector. And just like those UCSC sources, there is also a helper function called getChromInfoFromBiomart that can show what the different chromosomes are called for a given source.

Using the makeTranscriptDbFromBiomart makeTranscriptDbFromUCSC functions can take a while and may also require some bandwidth as these methods have to download and then assemble a database from their respective sources. It is not expected that most users will want to do this step every time. Instead, we suggest that you save your annotation objects and label them with an appropriate time stamp so as to facilitate reproducible research.

2.3 Saving and Loading a *TranscriptDb* Object

Once a TranscriptDb object has been created, it can be saved to avoid the time and bandwidth costs of recreating it and to make it possible to reproduce results with identical genomic feature data at a later date. Since TranscriptDb objects are backed by a SQLite database, the save format is a SQLite database file (which could be accessed from programs other than Rif desired). Note that it is not possible to serialize a TranscriptDb object using R's save function.

```
> saveFeatures(mm9KG, file="fileName.sqlite")
```

A TranscriptDb object can be initialized from a file using loadFeatures.

3 Retrieving Transcript, Exon, and Coding Sequence Ranges

3.1 Loading some sample genomic feature data

Here was are loading a previously created *TranscriptDb* object based on UCSC known gene data. This database only contains a small subset of the possible annotations for human and is only included to demonstrate and test the functionality of the *GenomicFeatures* package.

```
> samplefile <- system.file("extdata", "UCSC_knownGene_sample.sqlite",
                            package="GenomicFeatures")
> txdb <- loadFeatures(samplefile)
> txdb
TranscriptDb object:
| Db type: TranscriptDb
| Data source: UCSC
| Genome: hg18
| UCSC Table: knownGene
| Type of Gene ID: Entrez Gene ID
| Full dataset: no
| transcript_nrow: 135
| exon_nrow: 544
| cds_nrow: 324
| Db created by: GenomicFeatures package from Bioconductor
| Creation time: 2010-09-21 16:27:22 -0700 (Tue, 21 Sep 2010)
| GenomicFeatures version at creation time: 1.1.12
| RSQLite version at creation time: 0.9-2
| DBSCHEMAVERSION: 1.0
```

3.2 Working with Basic Features

The most basic operations on a TranscriptDb object retrieve the genomic coordinates or ranges for exons, transcripts or coding sequences. The functions transcripts, exons, and cds return the coordinate information as a GRanges object.

For example, all transcripts present in a TranscriptDb object can be obtained as follows:

```
> GR <- transcripts(txdb)
> GR[1:3]
```

GRanges with 3 ranges and 2 elementMetadata values

	seqnames]	ranges	strand		tx_id	tx_name
	<rle></rle>	<ira< td=""><td>anges></td><td><rle></rle></td><td></td><td><integer></integer></td><td><character></character></td></ira<>	anges>	<rle></rle>		<integer></integer>	<character></character>
[1]	chr1	[1116,	4121]	+		1	uc001aaa.2
[2]	chr1	[1116,	4272]	+		2	uc009vip.1
[3]	chr1	[4269,	6628]	_	1	3	uc009vis.1

seqlengths

```
        chr1
        chr1_random
        chr10
        ...
        chrX_random
        chrY

        247249719
        1663265
        135374737
        ...
        1719168
        57772954
```

The transcripts function returns a *GRanges* class object. You can learn a lot more about the manipulation of these objects by reading the *GenomicRanges* introductory vignette. The show method for a *GRanges* object will display the ranges, seqnames (a chromosome or a contig), and strand on the left side and then present related metadata on the right side. At the bottom, the seqlengths display all the possible seqnames along with the length of each sequence.

In addition, the **transcripts** function can also be used to retrieve a subset of the transcripts available such as those on the +-strand of chromosome 1.

```
> GR <- transcripts(txdb, vals <- list(tx_chrom = "chr1", tx_strand = "+"))
> length(GR)

[1] 2
> unique(strand(GR))

[1] +
Levels: + - *
```

The exons and cds functions can also be used in a similar fashion to retrive genomic coordinates for exons and coding sequences.

3.3 Working with Grouped Features

Often one is interested in how particular genomic features relate to each other, and not just their location. For example, it might be of interest to

group transcripts by gene or to group exons by transcript. Such groupings are supported by the transcriptsBy, exonsBy, and cdsBy functions.

The following call can be used to group transcripts by genes:

```
> GRList <- transcriptsBy(txdb, by = "gene")
> length(GRList)
[1] 51
> names(GRList)[10:13]
[1] "23192" "245938" "255403" "26751"
> GRList[11:12]
GRangesList of length 2
$245938
GRanges with 1 range and 2 elementMetadata values
                     ranges strand |
    segnames
       <Rle>
                  <IRanges>
                              <Rle> | <integer> <character>
[1]
       chr20 [16351, 25296]
                                                  uc002wcw.1
$255403
GRanges with 1 range and 2 elementMetadata values
    segnames
                       ranges strand |
                                            tx_id
                                                      tx_name
       <Rle>
                    <IRanges>
                               <Rle> | <integer> <character>
[1]
        chr4 [43227, 146490]
                                   + |
                                              10 uc003fzt.2
seqlengths
                 chr1_random
                                      chr10 ...
                                                   chrX_random
                                                                         chrY
          chr1
     247249719
                                  135374737 ...
                                                       1719168
                                                                     57772954
                     1663265
```

The transcriptsBy function returns a *GRangesList* class object. As with *GRanges* objects, you can learn more about these objects by reading the *GenomicRanges* introductory vignette. The show method for a *GRangesList* object will display as a list of *GRanges* objects. And, at the bottom the seqlengths will be displayed once for the entire list.

For each of these three functions, there is a limited set of options that can be passed into the by argument to allow grouping. For the transcriptsBy function, you can group by gene, exon or cds, whereas for the exonsBy and cdsBy functions can only be grouped by transcript (tx) or gene.

So as a further example, to extract all the exons for each transcript you can call:

```
> GRList <- exonsBy(txdb, by = "tx")
```

> length(GRList)

[1] 135

> names(GRList)[10:13]

[1] "10" "11" "12" "13"

> GRList[[12]]

GRanges with 4 ranges and 3 elementMetadata values

exon_rank	exon_name	exon_id	strand	ranges	seqnames	
<integer></integer>	<character></character>	<integer></integer>	<rle> </rle>	Ranges	<rle></rle>	
1	NA	50	+	[43227, 43385]	chr4	[1]
2	NA	51	+	[49323, 49449]	chr4	[2]
3	NA	52	+	[49951, 50046]	chr4	[3]
4	NA	56	+	[57732, 58380]	chr4	[4]

seqlengths

chr1	chr1_random	chr10	chrX_random	chrY
247249719	1663265	135374737	1719168	57772954

As you can see, the *GRangesList* objects returned from each function contain locations and identifiers grouped into a list like object according to the type of feature specified in the by argument. The object returned can then be used by functions like findOverlaps to contextualize alignments from high-throughput sequencing.

The identifiers used to label the *GRanges* objects depend upon the data source used to create the *TranscriptDb* object. So the list identifiers will not always be Entrez Gene IDs, as they were in the first example. Furthermore, some data sources do not provide a unique identifier for all features. In this situation, the group label will be a synthetic ID created by *GenomicFeatures* to keep the relations between features consistent in the database this was the case in the 2nd example. Even though the results will sometimes have to come back to you as synthetic IDs, you can still always retrieve the original IDs. In the following example, we will find the original transcript names stored in the database for each ID by calling the transcripts function.

```
> tx_ids <- names(GRList)
> vals <- list(tx_id=tx_ids)</pre>
> txs <- transcripts(txdb, vals, columns = c("tx_id", "tx_name"))
> head(values(txs))
DataFrame with 6 rows and 2 columns
      tx_id
                tx_name
  <integer> <character>
1
          1 uc001aaa.2
2
             uc009vip.1
3
          3
             uc009vis.1
4
         39 uc001ifj.1
5
             uc001ifi.1
         37
6
         38
             uc009xhe.1
```

Finally, the order of the results in a *GRangesList* object can vary with the way in which things were grouped. In most cases the grouped elements of the *GRangesList* object will be listed in the order that they occurred along the chromosome. However, when exons or CDS are grouped by transcript, they will instead be grouped according to their position along the transcript itself. This is important because alternative splicing can mean that the order along the transcript can be different from that along the chromosome.

3.4 Prespecifed grouping functions

The intronsByTranscript, fiveUTRsByTranscript and threeUTRsByTranscript are convenience functions that provide behavior equivalent to the grouping functions, but in prespecified form. These functions return a *GRangesList* object grouped by transcript for introns, 5' UTR's, and 3' UTR's, respectively.

```
> length(intronsByTranscript(txdb))
[1] 135
> length(fiveUTRsByTranscript(txdb))
[1] 61
> length(threeUTRsByTranscript(txdb))
[1] 58
```

3.5 Convenience functions for computing overlap

The transcriptsByOverlaps, exonsByOverlaps and cdsByOverlaps functions return a *GRangesList* object containing data about transcripts, exons, or coding sequences that overlap genomic coordinates specified by a *GRanges* object. So for example, lets just mock up some fake data:

```
> gr <- GRanges(
+ seqnames = rep("chr5",4),
+ ranges = IRanges(start = c(244620, 244670, 245804, 247502),
+ end = c(244652, 244702, 245836, 247534)),
+ strand = rep("+", 4))</pre>
```

Then we can call the convenience function to see what transcripts overlap with our ranges.

```
> transcriptsByOverlaps(txdb, gr)
```

```
GRanges with 1 range and 2 elementMetadata values
seqnames ranges strand | tx_id tx_name
<Rle> <IRanges> <Rle> | <integer> <character>
[1] chr5 [244626, 248468] + | 14 uc003jal.1
```

seqlengths

```
        chr1
        chr1_random
        chr10
        ...
        chrX_random
        chrY

        247249719
        1663265
        135374737
        ...
        1719168
        57772954
```

The convenience functions can be a great shortcut, but because they have to make assumptions about how the results are compared and represented, they are ultimately not as flexible as just using the basic and grouping accessors in combination with findOverlaps.

4 A typical example treating gr as RNA-seq data

Let's suppose that you have run an experiment. After mapping all your reads to a genome and collapsing them into a set of ranges, you want to find out which genomic features a particular range overlaps with. What would be the usual way to proceed?

For this example, let's also assume that you are only interested in mapping the ranges that overlap with exons (not introns). From our TranscriptDb object, we want to recover the annotations for all of the relevant

exons, but grouped according to their transcripts. Therefore, we want to use exonsby and group them by transcripts.

```
> annotGr <- exonsBy(txdb, "tx")</pre>
```

Then we need to use the findOverlaps method to learn which of our data ranges, gr, will overlap with the in exons that we have grouped by transcripts.

> OL <- findOverlaps(query = annotGr, subject = gr)

Finally, once we have called findOverlaps we can subset out the annotations that meet our criteria. The queryHits method will allow us to retrieve only the parts of the query that overlapped from our original findOverlaps call. Once we have subsetted out annotations in this way, the length of the resulting GRangesList object is also the number of transcripts that overlap with our data.

```
> tdata <- annotGr[unique(queryHits(OL)),]
> tdata
```

 ${\tt GRangesList}$ of length 1

\$14

GRanges with 2 ranges and 3 elementMetadata values

	seqnames	ranges	strand		exon_id	exon_name	exon_rank
	<rle></rle>	Ranges	<rle></rle>		<pre><integer></integer></pre>	<character></character>	<integer></integer>
[1]	chr5	[244626, 245552]	+		75	NA	1
[2]	chr5	[247823, 248468]	+	Ī	76	NA	2

seqlengths

```
chr1 chr1_random chr10 ... chrX_random chrY
247249719 1663265 135374737 ... 1719168 57772954
```

> length(tdata)

[1] 1

By using findOverlaps along with the different accessors in this way, it is possible to connect any data that has been represented as a *GRanges* object with the annotations stored in a *TranscriptDb* object. Calling find-Overlaps along with the appropriate *GRanges* object not only allows users

to quickly determine what has overlapped, but also controls what criteria are used for determining whether an overlap has occurred. This can be done by passing in an alternate type parameter to findOverlaps. In addition, because the basic accessors allow for the users to retrieve data grouped in different ways, the user has control over which parts of a transcript or gene are included in the overlap. For a more complete example of how you could approach RNA-seq analysis, including explanation on methods that will help to tally up the counts etc. please see the *GenomicRanges* Use Cases vignette.

5 Session Information

```
R version 2.13.2 (2011-09-30)
```

Platform: x86_64-unknown-linux-gnu (64-bit)

locale:

```
[1] LC_CTYPE=en_US.UTF-8 LC_NUMERIC=C
[3] LC_TIME=en_US.UTF-8 LC_COLLATE=C
```

[5] LC_MONETARY=C LC_MESSAGES=en_US.UTF-8

[7] LC_PAPER=en_US.UTF-8 LC_NAME=C
[9] LC_ADDRESS=C LC_TELEPHONE=C

[11] LC_MEASUREMENT=en_US.UTF-8 LC_IDENTIFICATION=C

attached base packages:

[1] stats graphics grDevices utils datasets methods base

other attached packages:

[1] GenomicFeatures_1.4.5 GenomicRanges_1.4.8 IRanges_1.10.6

loaded via a namespace (and not attached):

```
[1] BSgenome_1.20.1 Biostrings_2.20.4 DBI_0.2-5 RCurl_1.6-10
```

[5] RSQLite_0.10.0 XML_3.4-3 biomaRt_2.8.1 rtracklayer_1.12.5

[9] tools_2.13.2