HowTo: Build and use chromosomal information

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1 Overview

The annotate package provides a class that can be used to model chromosomal information about a species, using one of the metadata packages provided by Bioconductor. This class contains information about the organism and its chromosomes and provides a standardized interface to the information in the metadata packages for other software to quickly extract necessary chromosomal information. An example of using `chromLocation` objects in other software can be found with the `alongChrom` function of the `geneplotter` package in Bioconductor.

2 The chromLocation class

The `chromLocation` class is used to provide a structure for chromosomal data of a particular organism. In this section, we will discuss the various slots of the class and the methods for interacting with them. Before this though, we will create an object of class `chromLocation` for demonstration purposes later. The helper function `buildChromLocation` is used, and it takes as an argument the name of a Bioconductor metadata package, which is itself used to extract the data. For this vignette, we will be using the `hgu95av2` package.

```r
> library("annotate")
> z <- buildChromLocation("hgu95av2")
> z
```

Instance of a chromLocation class with the following fields:

- Organism: Homo sapiens
- Data source: hgu95av2
- Number of chromosomes for this organism: 25
- Chromosomes of this organism and their lengths in base pairs:
  1 : 246127941
  2 : 243615958
  3 : 199344050
  4 : 191731959
Once we have an object of the chromLocation class, we can now access its various slots to get the information contained within it. There are six slots in this class:

- **organism:** This lists the organism that this object is describing.
- **dataSource:** Where this data was acquired from.
- **chromLocs:** A list with an element for every unique chromosome name, where each element contains a named vector where the names are probe IDs and the values describe the location of that probe on the chromosome. Negative values indicate that the location is on the antisense strand.
- **probesToChrom:** A hash table which will translate a probe ID to the chromosome it belongs to.
- **chromInfo:** A numerical vector representing each chromosome, where the names are the names of the chromosomes and the values are the lengths of those chromosomes.
- **geneSymbols:** An environment that maps a probe ID to the appropriate gene symbol.

There is a basic 'get' type method for each of these slots, all with the same name as the respective slot. In the following example, we will demonstrate these basic methods. For the probesToChrom and geneSymbols methods, the return value is an environment which maps a probe ID to other values, we will be using the probe ID '32972_at', which was selected at random for these examples. We
are showing only part of the \texttt{chromLocs} method's output as it is quite long in its entirety.

\begin{verbatim}
> organism(z)
[1] "Homo sapiens"
> dataSource(z)
[1] "hgu95av2"

> names(chromLocs(z))

[1] "1" "10" "11" "12" "13" "14" "15" "16" "16_random" "17" "17_random" "18" "19" "2" "20" "21" "22" "3" "4" "4_random" "5" "6" "6_cox_hap1" "6_qbl_hap2" "7" "8" "9" "X" "Y" "3_random" "5_h2_hap1" "19_random" "2_random" "8_random" "22_random" "6_random" "X_random" "1_random"

> chromLocs(z)[["Y"]]

32930_f_at  31911_at  35930_at  32991_f_at  266_s_at  35885_at  35929_s_at  15145847  14324840  9914563  -6793959  -19611913  13322553  9914563  38182_at  40097_at  31534_at  40030_at  41214_at  38355_at  32864_at  20213723  21146998  2863545  7202013  2769622  13526170  -2714896  37583_at  629_at  39168_at  34215_at  31415_at  40342_at  -20326690  57739639  -2414454  1670485  -18390255  18756722  -23684986  40342_at  1185_at  40436_g_at  36553_s_at  31412_at  31412_at  40342_at  25389451  1415508  -1465044  -1482031  -22627290  23045931  -14607046  32677_at  41138_at  40435_at  36554_at  35073_at  34753_at  35447_s_at  14677491  2619227  -1465044  -1482031  505078  57623412  1674347  34172_s_at  33593_at  33593_at  33593_at  41108_at  34477_at  34477_at  1670485  -24600763  26177651  -24601329  -161425  -13918783  34777_at  31411_at  31411_at  31411_at  33665_s_at  33665_s_at  33665_s_at  31601_s_at  -13869656  23539797  25173538  -25586439  1361570  1347700  1347700  31601_s_at  31601_s_at  31601_s_at  31601_s_at  31601_s_at  31601_s_at  31601_s_at  22106177  -22435611  -22459154  22082645  22106186

> get("32972_at", probesToChrom(z))

[1] "X"

> chromInfo(z)
\end{verbatim}
Another method which can be used to access information about the particular *chromLocation* object is the `nChrom` method, which will list how many chromosomes this organism has:

```r
> nChrom(z)
[1] 25
```

## 3 Summary

The *chromLocation* class has a simple design, but can be powerful if one wants to store the chromosomal data contained in a Bioconductor package into a single object. These objects can be created once and then passed around to multiple functions, which can cut down on computation time to access the desired information from the package. These objects allow access to basic but also important information, and provide a standard interface for writers of other software to access this information.