

Extending *GenomicRanges*

Michael Lawrence, Bioconductor Team

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1 Introduction

The goal of *GenomicRanges* is to provide general containers for genomic data. The central class, at least from the user perspective, is *GRanges*, which formalizes the notion of ranges, while allowing for arbitrary “metadata columns” to be attached to it. These columns offer the same flexibility as the venerable *data.frame* and permit users to adapt *GRanges* to a wide variety of *ad hoc* use-cases.

The more we encounter a particular problem, the better we understand it. We eventually develop a systematic approach for solving the most frequently encountered problems, and every systematic approach deserves a systematic implementation. For example, we might want to formally store genetic variants, with information on alleles and read depths. The metadata columns, which were so useful during prototyping, are inappropriate for extending the formal semantics of our data structure: for the sake of data integrity, we need to ensure that the columns are always present and that they meet certain constraints.

We might also find that our prototype does not scale well to the increased data volume that often occurs when we advance past the prototype stage. *GRanges* is meant mostly for prototyping and stores its data in memory as simple R data structures. We may require something more specialized when the data are large; for example, we might store the data as a Tabix-indexed file, or in a database.

The *GenomicRanges* package does not directly solve either of these problems, because there are no general solutions. However, it is adaptable to specialized use cases.

2 The *GenomicRanges* abstraction

Unbeknownst to many, most of the *GRanges* implementation is provided by methods on the *GenomicRanges* class, the virtual parent class of *GRanges*. *GenomicRanges* methods provide everything except for the actual data storage and retrieval, which *GRanges* implements directly using slots. For example, the ranges are retrieved like this:

```
> library(GenomicRanges)
> selectMethod(ranges, "GRanges")

Method Definition:

function (x, use.names = TRUE, use.mcols = FALSE, ...)
{
  .local <- function (x, use.names = TRUE, use.mcols = FALSE)
  {
    if (!isTRUEorFALSE(use.names))
      stop("'use.names' must be TRUE or FALSE")
    if (!isTRUEorFALSE(use.mcols))
      stop("'use.mcols' must be TRUE or FALSE")
    ans <- x@ranges
    if (!use.names)
      names(ans) <- NULL
    if (use.mcols)
      mcols(ans) <- mcols(x)
    ans
  }
  .local(x, use.names, use.mcols, ...)
}
<environment: namespace:GenomicRanges>

Signatures:
      x
target  "GRanges"
defined "GRanges"
```

An alternative implementation is *DelegatingGenomicRanges*, which stores all of its data in a delegate *GenomicRanges* object:

```
> selectMethod(ranges, "DelegatingGenomicRanges")

Method Definition:

function (x, use.names = TRUE, use.mcols = FALSE, ...)
ranges(x@delegate, ...)
<environment: namespace:GenomicRanges>

Signatures:
      x
target  "DelegatingGenomicRanges"
defined "DelegatingGenomicRanges"
```

This abstraction enables us to pursue more efficient implementations for particular tasks. One example is *GNCList*, which is indexed for fast range queries, we expose here:

```
> getSlots("GNCList")["granges"]  
  
granges  
"GRanges"
```

The *MutableRanges* package in svn provides other, untested examples.

3 Formalizing `mcols`: Extra column slots

An orthogonal problem to data storage is adding semantics by the formalization of metadata columns, and we solve it using the “extra column slot” mechanism. Whenever *GenomicRanges* needs to operate on its metadata columns, it also delegates to the internal `extraColumnSlotNames` generic, methods of which should return a character vector, naming the slots in the *GenomicRanges* subclass that correspond to columns (i.e., they have one value per range). It extracts the slot values and manipulates them as it would a metadata column – except they are now formal slots, with formal types.

An example is the *VRanges* class in *VariantAnnotation*. It stores information on the variants by adding these column slots:

```
> GenomicRanges::extraColumnSlotNames(VariantAnnotation::VRanges())  
  
[1] "ref"          "alt"          "totalDepth"  
[4] "refDepth"     "altDepth"     "sampleNames"  
[7] "softFilterMatrix"
```

Mostly for historical reasons, *VRanges* extends *GRanges*. However, since the data storage mechanism and the set of extra column slots are orthogonal, it is probably best practice to take a composition approach by extending *DelegatingGenomicRanges*.