An Overview of the S4Vectors package

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1 Introduction

The *S4Vectors* package provides a framework for representing vector-like and list-like objects as S4 objects. It defines two central virtual classes, *Vector* and *List*, and a set of generic functions that extend the semantic of ordinary vectors and lists in *R*. Package developers can easily implement vector-like or list-like objects as *Vector* and/or *List* derivatives. A few low-level *Vector* and *List* derivatives are implemented in the *S4Vectors* package itself e.g. *Hits*, *Rle*, and *DataFrame*). Many more are implemented in the *IRanges* and *GenomicRanges* infrastructure packages, and in many other Bioconductor packages.

In this vignette, we will rely on simple, illustrative example datasets, rather than large, realworld data, so that each data structure and algorithm can be explained in an intuitive, graphical manner. We expect that packages that apply S4Vectors to a particular problem domain will provide vignettes with relevant, realistic examples.

The *S4Vectors* package is available at bioconductor.org and can be downloaded via **BiocMan** ager::install:

```
> if (!require("BiocManager"))
+ install.packages("BiocManager")
> BiocManager::install("S4Vectors")
```

```
> library(S4Vectors)
```

2 Vector-like and list-like objects

In the context of the *S4Vectors* package, a vector-like object is an ordered finite collection of elements. All vector-like objects have three main properties: (1) a notion of length or number of elements, (2) the ability to extract elements to create new vector-like objects, and (3) the ability to be concatenated with one or more vector-like objects to form larger vector-like objects. The main functions for these three operations are **length**, [, and c. Supporting these operations provide a great deal of power and many vector-like object manipulations can be constructed using them.

Some vector-like objects can also have a list-like semantic, which means that individual elements can be extracted with [[.

In *S4Vectors* and many other Bioconductor packages, vector-like and list-like objects derive from the *Vector* and *List* virtual classes, respectively. Note that *List* is a subclass of *Vector*.

The following subsections describe each in turn.

2.1 Vector-like objects

As a first example of vector-like objects, we'll look at *Rle* objects. In *R*, atomic sequences are typically stored in atomic vectors. But there are times when these object become too large to manage in memory. When there are lots of consecutive repeats in the sequence, the data can be compressed and managed in memory through a run-length encoding where a data value is paired with a run length. For example, the sequence $\{1, 1, 1, 2, 3, 3\}$ can be represented as values = $\{1, 2, 3\}$, run lengths = $\{3, 1, 2\}$.

The *Rle* class defined in the *S4Vectors* package is used to represent a run-length encoded (compressed) sequence of *logical*, *integer*, *numeric*, *complex*, *character*, *raw*, or *factor* values. Note that the *Rle* class extends the *Vector* virtual class:

```
> showClass("Rle")
```

Class "Rle" [package "S4Vectors"]

Slots:

Name: values lengths elementMetadata Class: vector_OR_factor integer_OR_LLint DataFrame_OR_NULL

Name: metadata Class: list

```
Extends:
Class "Vector", directly
Class "Annotated", by class "Vector", distance 2
Class "vector_OR_Vector", by class "Vector", distance 2
```

One way to construct *Rle* objects is through the *Rle* constructor function:

```
> set.seed(0)
> lambda <- c(rep(0.001, 4500), seq(0.001, 10, length=500),
+ seq(10, 0.001, length=500))
> xVector <- rpois(1e7, lambda)</pre>
```

```
> yVector <- rpois(1e7, lambda[c(251:length(lambda), 1:250)])
> xRle <- Rle(xVector)
> yRle <- Rle(yVector)
Rle objects are vector-like objects:
> length(xRle)
[1] 10000000
> xRle[1]
integer-Rle of length 1 with 1 run
Lengths: 1
Values : 0
```

```
> zRle <- c(xRle, yRle)</pre>
```

2.1.1 Subsetting a vector-like object

As with ordinary R atomic vectors, it is often necessary to subset one sequence from another. When this subsetting does not duplicate or reorder the elements being extracted, the result is called a *subsequence*. In general, the [function can be used to construct a new sequence or extract a subsequence, but its interface is often inconvenient and not amenable to optimization. To compensate for this, the *S4Vectors* package supports seven additional functions for sequence extraction:

- 1. window Extracts a subsequence over a specified region.
- 2. subset Extracts the subsequence specified by a logical vector.
- 3. head Extracts a consecutive subsequence containing the first n elements.
- 4. tail Extracts a consecutive subsequence containing the last n elements.
- 5. rev Creates a new sequence with the elements in the reverse order.
- 6. rep Creates a new sequence by repeating sequence elements.

The following code illustrates how these functions are used on an *Rle* vector:

```
> xSnippet <- window(xRle, 4751, 4760)
> xSnippet
integer-Rle of length 10 with 9 runs
Lengths: 1 1 1 1 1 1 1 1 2
Values : 4 6 5 4 6 2 6 7 5
> head(xSnippet)
integer-Rle of length 6 with 6 runs
Lengths: 1 1 1 1 1 1
Values : 4 6 5 4 6 2
> tail(xSnippet)
integer-Rle of length 6 with 5 runs
Lengths: 1 1 1 1 2
Values : 6 2 6 7 5
```

```
> rev(xSnippet)
integer-Rle of length 10 with 9 runs
Lengths: 2 1 1 1 1 1 1 1 1
Values : 5 7 6 2 6 4 5 6 4
> rep(xSnippet, 2)
integer-Rle of length 20 with 18 runs
Lengths: 1 1 1 1 1 1 1 1 2 1 1 1 1 1 1 1 1 2
Values : 4 6 5 4 6 2 6 7 5 4 6 5 4 6 2 6 7 5
> subset(xSnippet, xSnippet >= 5L)
integer-Rle of length 7 with 5 runs
Lengths: 1 1 2 1 2
Values : 6 5 6 7 5
```

2.1.2 Concatenating vector-like objects

The *S4Vectors* package uses two generic functions, **c** and **append**, for concatenating two *Vector* derivatives. The methods for *Vector* objects follow the definition that these two functions are given the *base* package.

```
> c(xSnippet, rev(xSnippet))
```

```
integer-Rle of length 20 with 17 runs
Lengths: 1 1 1 1 1 1 1 4 1 1 1 1 1 1 1 1
Values : 4 6 5 4 6 2 6 7 5 7 6 2 6 4 5 6 4
```

```
> append(xSnippet, xSnippet, after=3)
```

```
integer-Rle of length 20 with 18 runs
Lengths: 1 1 1 1 1 1 1 1 1 1 2 1 1 1 1 2
Values : 4 6 5 4 6 5 4 6 2 6 7 5 4 6 2 6 7 5
```

2.1.3 Looping over subsequences of vector-like objects

In *R*, for looping can be an expensive operation. To compensate for this, the *S4Vectors* package provides aggregate and shiftApply methods (shiftApply is a new generic function defined in *S4Vectors*) to perform calculations over subsequences of vector-like objects.

The aggregate function combines sequence extraction functionality of the window function with looping capabilities of the sapply function. For example, here is some code to compute medians across a moving window of width 3 using the function aggregate:

```
> xSnippet
integer-Rle of length 10 with 9 runs
Lengths: 1 1 1 1 1 1 1 1 2
Values : 4 6 5 4 6 2 6 7 5
> aggregate(xSnippet, start=1:8, width=3, FUN=median)
[1] 5 5 5 4 6 6 6 5
```

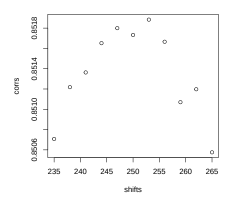


Figure 1: Correlation between xRle and yRle for various shifts.

The shiftApply function is a looping operation involving two vector-like objects whose elements are lined up via a positional shift operation. For example, the elements of xRle and yRle were simulated from Poisson distributions with the mean of element i from yRle being equivalent to the mean of element i + 250 from xRle. If we did not know the size of the shift, we could estimate it by finding the shift that maximizes the correlation between xRle and yRle.

```
> cor(xRle, yRle)
[1] 0.5739224
> shifts <- seq(235, 265, by=3)
> corrs <- shiftApply(shifts, yRle, xRle, FUN=cor)</pre>
```

```
> plot(shifts, corrs)
```

The result is shown in Fig. 1.

2.1.4 More on *Rle* objects

When there are lots of consecutive repeats, the memory savings through an RLE can be quite dramatic. For example, the <u>xRle</u> object occupies less than one third of the space of the original <u>xVector</u> object, while storing the same information:

```
> as.vector(object.size(xRle) / object.size(xVector))
```

```
[1] 0.3020726
```

```
> identical(as.vector(xRle), xVector)
```

[1] TRUE

The functions **runValue** and **runLength** extract the run values and run lengths from an *Rle* object respectively:

```
> head(runValue(xRle))
```

[1] 0 1 0 1 0 1

> head(runLength(xRle)) [1] 780 1 208 1 1599

The *Rle* class supports many of the basic methods associated with R atomic vectors including the Ops, Math, Math2, Summary, and Complex group generics. Here is a example of manipulating *Rle* objects using methods from the Ops group:

1

> xRle > 0 logical-Rle of length 10000000 with 197127 runs Lengths: 780 1 208 1 1599 ... 1 91 1 927 Values : FALSE TRUE FALSE TRUE FALSE ... TRUE FALSE TRUE FALSE > xRle + yRle integer-Rle of length 10000000 with 1957707 runs Lengths: 780 1 208 1 13 1 413 ... 5 1 91 1 507 1 419 Values : 0 1 0 1 0 1 0 ... 1 0 0 1 0 1 0 > xRle > 0 | yRle > 0logical-Rle of length 10000000 with 210711 runs 1 Lengths: 780 1 208 13 ... 1 507 1 419 Values : FALSE TRUE FALSE TRUE FALSE ... TRUE FALSE TRUE FALSE

Here are some from the Summary group:

> range(xRle) [1] 0 26 > sum(xRle > 0 | yRle > 0)

[1] 2105185

And here is one from the Math group:

```
> log1p(xRle)
numeric-Rle of length 10000000 with 1510219 runs
Lengths: 780 1 208 ... 91 1 927
Values : 0.000000 0.693147 0.000000 ... 0.000000 0.693147 0.000000
```

As with atomic vectors, the cor and shiftApply functions operate on *Rle* objects:

For more information on the methods supported by the *Rle* class, consult the Rle man page.

2.2 List-like objects

Just as with ordinary *R* list objects, List-derived objects support [[for element extraction, c for concatenating, and lapply/sapply for looping. lapply and sapply are familiar to many *R* users since they are the standard functions for looping over the elements of an *R* list object.

In addition, the *S4Vectors* package introduces the <u>endoapply</u> function to perform an endomorphism equivalent to <u>lapply</u>, i.e. it returns a *List* derivative of the same class as the input rather than a *list* object.

An example of *List* derivative is the *DataFrame* class:

```
> showClass("DataFrame")
Virtual Class "DataFrame" [package "S4Vectors"]
Slots:
Name:
             elementType elementMetadata
                                                    metadata
Class:
               character DataFrame_OR_NULL
                                                        list
Extends:
Class "RectangularData", directly
Class "List", directly
Class "DataFrame_OR_NULL", directly
Class "Vector", by class "List", distance 2
Class "list_OR_List", by class "List", distance 2
Class "Annotated", by class "List", distance 3
Class "vector_OR_Vector", by class "List", distance 3
```

```
Known Subclasses: "DFrame"
```

One way to construct DataFrame objects is through the DataFrame constructor function:

```
> df <- DataFrame(x=xRle, y=yRle)</pre>
> sapply(df, class)
    х
         y
"Rle" "Rle"
> sapply(df, summary)
                х
                           у
Min.
        0.000000 0.000000
1st Qu. 0.0000000 0.0000000
Median
        0.000000 0.000000
Mean
        0.9090338 0.9096009
3rd Qu. 0.0000000 0.0000000
Max.
        26.000000 27.0000000
> sapply(as.data.frame(df), summary)
                х
                           v
Min.
        0.000000 0.000000
1st Qu. 0.0000000 0.0000000
Median 0.0000000 0.0000000
```

Mean 0.9090338 0.9096009 3rd Qu. 0.0000000 0.0000000 26.000000 27.0000000 Max. > endoapply(df, `+`, 0.5) DataFrame with 10000000 rows and 2 columns х У <Rle> <Rle> 1 0.5 0.5 2 0.5 0.5 3 0.5 0.5 4 0.5 0.5 5 0.5 0.5 9999996 0.5 0.5 0.5 0.5 9999997 9999998 0.5 0.5 9999999 0.5 0.5 10000000 0.5 0.5

For more information on DataFrame objects, consult the DataFrame man page.

See the "An Overview of the *IRanges* package" vignette in the *IRanges* package for many more examples of *List* derivatives.

3 Vector Annotations

Often when one has a collection of objects, there is a need to attach metadata that describes the collection in some way. Two kinds of metadata can be attached to a *Vector* object:

- Metadata about the object as a whole: this metadata is accessed via the metadata accessor and is represented as an ordinary *list*;
- 2. Metadata about the individual elements of the object: this metadata is accessed via the mcols accessor (mcols stands for metadata columns) and is represented as a DataFrame object. This DataFrame object can be thought of as the result of binding together one or several vector-like objects (the metadata columns) of the same length as the Vector object. Each row of the DataFrame object annotates the corresponding element of the Vector object.

4 Session Information

Here is the output of sessionInfo() on the system on which this document was compiled:

```
R version 4.3.1 (2023-06-16)
Platform: x86_64-pc-linux-gnu (64-bit)
Running under: Ubuntu 22.04.3 LTS
Matrix products: default
BLAS: /home/biocbuild/bbs-3.17-bioc/R/lib/libRblas.so
LAPACK: /usr/lib/x86_64-linux-gnu/lapack/liblapack.so.3.10.0
```

locale: [1] LC_CTYPE=en_US.UTF-8 LC_NUMERIC=C [3] LC_TIME=en_GB LC_COLLATE=C [5] LC_MONETARY=en_US.UTF-8 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 time zone: America/New_York tzcode source: system (glibc) attached base packages: [1] stats4 stats graphics grDevices utils datasets [7] methods base other attached packages: [1] graph_1.78.0 ShortRead_1.58.0 [3] GenomicAlignments_1.36.0 SummarizedExperiment_1.30.2 [5] Biobase_2.60.0 MatrixGenerics_1.12.3 [7] matrixStats_1.0.0 Rsamtools_2.16.0 [9] GenomicRanges_1.52.0 Biostrings_2.68.1 [11] GenomeInfoDb_1.36.3 XVector_0.40.0 [13] BiocParallel_1.34.2 IRanges_2.34.1 [15] S4Vectors_0.38.2 BiocGenerics_0.46.0 [17] Matrix_1.6-1.1 BiocStyle_2.28.1 loaded via a namespace (and not attached): [1] sass_0.4.7 bitops_1.0-7 [3] jpeg_0.1-10 lattice_0.21-8 [5] digest_0.6.33 evaluate_0.21 [7] grid_4.3.1 RColorBrewer_1.1-3 fastmap_1.1.1 [9] bookdown_0.35 [11] jsonlite_1.8.7 BiocManager_1.30.22 [13] codetools_0.2-19 jquerylib_0.1.4 [15] abind_1.4-5 cli_3.6.1 [17] rlang_1.1.1 crayon_1.5.2 [19] cachem_1.0.8 DelayedArray_0.26.7 [21] yaml_2.3.7 S4Arrays_1.0.6 [23] tools_4.3.1 parallel_4.3.1 [25] deldir_1.0-9 interp_1.1-4 [27] GenomeInfoDbData_1.2.10 hwriter_1.3.2.1 [29] R6_2.5.1 png_0.1-8 [31] zlibbioc_1.46.0 bslib_0.5.1 xfun_0.40 [33] Rcpp_1.0.11 [35] knitr_1.44 latticeExtra_0.6-30 [37] htmltools_0.5.6 rmarkdown_2.25 [39] compiler_4.3.1 RCurl_1.98-1.12