# Practical: Ranges

## Martin Morgan (mtmorgan@fhcrc.org)

## 27-28 February, 2014

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# 1 Working with ranges

Start by loading the *GenomicRanges* package and defining the plotRanges helper function

Ranges describe both features of interest (e.g., genes, exons, promoters) and reads aligned to the genome. *Bioconductor* has very powerful facilities for working with ranges, some of which are summarized in Table<sup>1</sup>. These are implemented in the *GenomicRanges* package; see [1] for a more comprehensive conceptual orientation.

**The GRanges class** Instances of *GRanges* are used to specify genomic coordinates. Suppose we wish to represent two *D. melanogaster* genes. The first is located on the positive strand of chromosome 3R, from position 19967117 to 19973212. The second is on the minus strand of the X chromosome, with 'left-most' base at 18962306, and right-most base at 18962925. The coordinates are *1-based* (i.e., the first nucleotide on a chromosome is numbered 1, rather than 0), *left-most* (i.e., reads on the minus strand are defined to 'start' at the left-most coordinate, rather than the 5' coordinate), and *closed* (the start and end coordinates are included in the range; a range with identical start and end coordinates has width 1, a 0-width range is represented by the special construct where the end coordinate is one less than the start coordinate). A complete definition of these genes as *GRanges* is:

Table 1: Selected	Bioconductor	packages f	or representing	g and mani	pulating range	es, strings,	and other	data s	struc-
tures.									

Package	Description
IRanges	Defines important classes (e.g., IRanges, Rle) and methods (e.g.,
	findOverlaps, countOverlaps) for representing and manipulating ranges of
	consecutive values. Also introduces DataFrame, SimpleList and other classes
	tailored to representing very large data.
GenomicRanges	Range-based classes tailored to sequence representation (e.g., GRanges,
	GRangesList), with information about strand and sequence name.
GenomicFeatures	Foundation for manipulating data bases of genomic ranges, e.g., representing
	coordinates and organization of exons and transcripts of known genes.

The components of a *GRanges* object are defined as vectors, e.g., of seqnames, much as one would define a *data.frame*. The start and end coordinates are grouped into an *IRanges* instance. The optional seqlengths argument specifies the maximum size of each sequence, in this case the lengths of chromosomes 3R and X in the 'dm2' build of the *D. melanogaster* genome. This data is displayed as

genes

```
## GRanges with 2 ranges and 0 metadata columns:
## seqnames ranges strand
## <Rle> <IRanges> <Rle>
## [1] chr3R [19967117, 19973212] +
## [2] chrX [18962306, 18962925] -
## ----
## seqlengths:
## chr3R chrX
## 27905053 22422827
```

The GRanges class has many useful methods defined on it. Consult the help page

?GRanges

and package vignettes

vignette(package="GenomicRanges")

for a comprehensive introduction. A *GRanges* instance can be subset, with accessors for getting and updating information.

```
genes<sup>[2]</sup>
## GRanges with 1 range and 0 metadata columns:
   seqnames ranges strand
<Rle> <IRanges> <Rle>
##
##
    [1] chrX [18962306, 18962925]
##
## ----
## seqlengths:
##
     chr3R
                  chrX
##
      27905053 22422827
strand(genes)
## factor-Rle of length 2 with 2 runs
## Lengths: 1 1
## Values : + -
## Levels(3): + - *
width(genes)
## [1] 6096 620
length(genes)
## [1] 2
```



Figure 1: Ranges

nar	nes(genes) <- c("FBgn00	39155", "FB	gn0085359")	)
gen	nes # now with names			
##	GRanges with 2 ranges	and 0 metada	ata columna	5:
##	seqnames		ranges	strand
##	<rle></rle>		<iranges></iranges>	<rle></rle>
##	FBgn0039155 chr3R	[19967117,	19973212]	+
##	FBgn0085359 chrX	[18962306,	18962925]	-
##				
##	seqlengths:			
##	chr3R chrX			
##	27905053 22422827			

strand returns the strand information in a compact representation called a *run-length encoding*. The 'names' could have been specified when the instance was constructed; once named, the *GRanges* instance can be subset by name like a regular vector.

As the GRanges function suggests, the *GRanges* class extends the *IRanges* class by adding information about seqnames, strand, and other information particularly relevant to representing ranges that are on genomes. The *IRanges* class and related data structures (e.g., *RangedData*) are meant as a more general description of ranges defined in an arbitrary space. Many methods implemented on the *GRanges* class are 'aware' of the consequences of genomic location, for instance treating ranges on the minus strand differently (reflecting the 5' orientation imposed by DNA) from ranges on the plus strand.

**Operations on ranges** The *GRanges* class has many useful methods. We use *IRanges* to illustrate these operations to avoid complexities associated with strand and seqnames, but the operations are comparable on *GRanges*. We begin with a simple set of ranges:

These and some common operations are illustrated in the upper panel of Figure<sup>1</sup> and summarized in Table<sup>2</sup>.

Methods on ranges can be grouped as follows:

**Intra-range** methods act on each range independently. These include flank, narrow, reflect, resize, restrict, and shift, among others. An illustration is shift, which translates each range by the amount specified by the shift argument. Positive values shift to the right, negative to the left; shift can be a vector, with each element of the vector shifting the corresponding element of the *IRanges* instance. Here we shift all ranges to the right by 5, with the result illustrated in the middle panel of Figure~1.

shi	ift(	ir, 5	5)			
##	IRar	nges	oſ	ler	ıgth	7
##		stai	ct	end	widt	h
##	[1]	1	12	20		9
##	[2]	1	14	16		3
##	[3]	1	17	17		1
##	[4]	1	19	23		5
##	[5]	4	27	31		5
##	[6]	4	28	32		5
##	[7]	2	29	33		5

**Inter-range** methods act on the collection of ranges as a whole. These include disjoin, reduce, gaps, and range. An illustration is reduce, which reduces overlapping ranges into a single range, as illustrated in the lower panel of Figure 1.

```
reduce(ir)
## IRanges of length 2
## start end width
## [1] 7 18 12
## [2] 22 28 7
```

coverage is an inter-range operation that calculates how many ranges overlap individual positions. Rather than returning ranges, coverage returns a compressed representation (run-length encoding)

```
cvg <- coverage(ir)
cvg
## integer-Rle of length 28 with 12 runs
## Lengths: 6 2 4 1 2 3 3 1 1 3 1 1
## Values : 0 1 2 1 2 1 0 1 2 3 2 1
## plot(as.integer(cvg), type="s", xlab="Coordinate", ylab="Depth of coverage")</pre>
```

The run-length encoding can be interpreted as 'a run of length 6 of nucleotides covered by 0 ranges, followed by a run of length 2 of nucleotides covered by 1 range...'.

Between methods act on two (or sometimes more) *IRanges* instances. These include intersect, setdiff, union, pintersect, psetdiff, and punion.

The countOverlaps and findOverlaps functions operate on two sets of ranges. countOverlaps takes its first argument (the query) and determines how many of the ranges in the second argument (the subject) each overlaps. The result is an integer vector with one element for each member of query. findOverlaps performs a similar operation but returns a more general matrix-like structure that identifies each pair of query / subject overlaps. Both arguments allow some flexibility in the definition of 'overlap'.

Adding mcols and metadata The *GRanges* class (actually, most of the data structures defined or extending those in the *IRanges* package) has two additional very useful data components. The mcols function allows information on each range to be stored and manipulated (e.g., subset) along with the *GRanges* instance. The element metadata is represented as a *DataFrame*, defined in *IRanges* and acting like a standard *R data.frame* but with the ability to hold more complicated data structures as columns (and with element metadata of its own, providing an enhanced alternative to the *Biobase* class *AnnotatedDataFrame*).

metadata allows addition of information to the entire object. The information is in the form of a list; any data can be provided.

metadata(genes) <- list(CreatedBy="A. User", Date=date())</pre>

Category	Function	Description
Accessors	start, end, width	Get or set the starts, ends and widths
	names	Get or set the names
	mcols, metadata	Get or set metadata on elements or object
	length	Number of ranges in the vector
	range	Range formed from min start and max end
Ordering	<, <=, >, >=, ==, !=	Compare ranges, ordering by start then width
	sort, order, rank	Sort by the ordering
	duplicated	Find ranges with multiple instances
	unique	Find unique instances, removing duplicates
Arithmetic	r + x, r - x, r * x	Shrink or expand ranges $r$ by number $x$
	shift	Move the ranges by specified amount
	resize	Change width, anchoring on start, end or mid
	distance	Separation between ranges (closest endpoints)
	restrict	Clamp ranges to within some start and end
	flank	Generate adjacent regions on start or end
Set operations	reduce	Merge overlapping and adjacent ranges
	intersect, union, setdiff	Set operations on reduced ranges
	pintersect, punion, psetdiff	Parallel set operations, on each x[i], y[i]
	gaps, pgap	Find regions not covered by reduced ranges
	disjoin	Ranges formed from union of endpoints
Overlaps	findOverlaps	Find all overlaps for each $x$ in $y$
	countOverlaps	Count overlaps of each x range in y
	nearest	Find nearest neighbors (closest endpoints)
	precede, follow	Find nearest y that x precedes or follows
	x %in% y	Find ranges in $\mathbf{x}$ that overlap range in $\mathbf{y}$
Coverage	coverage	Count ranges covering each position
Extraction	r[i]	Get or set by logical or numeric index
	r[[i]]	Get integer sequence from start[i] to end[i]
	subsetByOverlaps	Subset $x$ for those that overlap in $y$
	head, tail, rev, rep	Conventional R semantics
Split, combine	split	Split ranges by a factor into a RangesList
	c	Concatenate two or more range objects

Table 2: Common operations on IRanges, GRanges and GRangesList.

**The GRangesList class** The GRanges class is extremely useful for representing simple ranges. Some nextgeneration sequence data and genomic features are more hierarchically structured. A gene may be represented by several exons within it. An aligned read may be represented by discontinuous ranges of alignment to a reference. The *GRangesList* class represents this type of information. It is a list-like data structure, which each element of the list itself a *GRanges* instance. The ENSEMBL genes identified earlier can be represented as a *GRangesList*.

##	GRangesL	ist of ]	Length 6:				
##	\$84929						
##	GRanges	with 10	ranges and 2	2 metadata	columns:		
##	S	eqnames		ranges	strand	exon_id	exon_name
##		<rle></rle>		<iranges></iranges>	<rle>  </rle>	<integer></integer>	<character></character>
##	[1]	chr9	[133777825,	133779710]	-	132272	<na></na>
##	[2]	chr9	[133780621,	133780800]	-	132273	<na></na>
##	[3]	chr9	[133787179,	133787275]	-	132274	<na></na>
##	[4]	chr9	[133799131,	133799267]	-	132275	<na></na>
##	[5]	chr9	[133799624,	133799783]	-	132276	<na></na>
##	[6]	chr9	[133804954,	133805433]	-	132277	<na></na>
##	[7]	chr9	[133806160,	133806183]	-	132278	<na></na>

##	[8]	chr9	[133813923]	, 13381403	5] –	13	2279	<na></na>	
##	[9]	chr9	[133813923]	, 133814239	9] –	13	2280	<na></na>	
##	[10]	chr9	[133814390]	, 13381445	5] -	13	2281	<na></na>	
##									
##	\$8140								
##	GRanges	with 10	ranges and	2 metadata	a columns	:			
##		seqnames		ranges	strand	exon_id	exon_name		
##	[1]	chr16	[87863629,	87866631]	-	215168	<na></na>		
##	[2]	chr16	[87868020,	87868197]	-	215169	<na></na>		
##	[3]	chr16	[87870104,	87870253]	-	215170	<na></na>		
##	[4]	chr16	[87871451,	87871547]	-	215171	<na></na>		
##	[5]	chr16	[87872320,	87872423]	-	215172	<na></na>		
##	[6]	chr16	[87873308,	87873431]	-	215173	<na></na>		
##	[7]	chr16	[87874035,	87874079]	-	215174	<na></na>		
##	[8]	chr16	[87874656,	87874761]	-	215175	<na></na>		
##	[9]	chr16	[87885330,	87885455]	-	215176	<na></na>		
##	[10]	chr16	[87902491,	87903100]	-	215177	<na></na>		
##									
##	• • •								
##	<4 more	e elements	5>						
##									
##	seqleng	ths:							
##			chr1		chr2		chrUn_gl	000249	
##		249	9250621	24	43199373			38502	

The *GRangesList* object has methods one would expect for lists (e.g., length, sub-setting). Many of the methods introduced for working with *IRanges* are also available, with the method applied element-wise.

### 1.1 Selecting gene sequences

### Exercise 1

This exercise uses annotation packages to go from gene identifiers to coding sequences.

- a. Map from an informal gene SYMBOL, e.g., BRCA1, to ENTREZID gene identifiers using the org.Hs.eg.db package and the select function, use the TxDb.Hsapiens.UCSC.hg19.knownGene package and a second map to go from ENTREZID to TXNAME.
- b. Extract the coding sequence grouped by transcript using the TxDb.Hsapiens.UCSC.hg19.knownGene package and cdsBy function; select just those transcripts we are interested in.
- c. Retrieve the nucleotide sequence from the BSgenome.Hsapiens.UCSC.hg19 package using the function extractTranscriptsFromGenome.
- d. Verify that the coding sequences are all multiples of 3, and translate from nucleotide to amino acid sequence.

# **Solution:** Map from gene SYMBOL to ENTREZID, and from ENTREZID to TXNAME and extract the releveant coding sequence, grouped by transcript

```
library(org.Hs.eg.db)
egid <- select(org.Hs.eg.db, "BRCA1", "ENTREZID", "SYMBOL")$ENTREZID
library(TxDb.Hsapiens.UCSC.hg19.knownGene)
txdb <- TxDb.Hsapiens.UCSC.hg19.knownGene
egToTx <- select(txdb, egid, "TXNAME", "GENEID")
## Warning: 'select' resulted in 1:many mapping between keys and return rows
cdsByTx <- cdsBy(txdb, "tx", use.names=TRUE)[egToTx$TXNAME]</pre>
```

### Retrieve the sequence

```
library(BSgenome.Hsapiens.UCSC.hg19)
txx <- extractTranscriptsFromGenome(Hsapiens, cdsByTx)
## Warning: 'extractTranscriptsFromGenome' is deprecated.
## Use 'extractTranscriptSeqs' instead.
## Warning: 'extractTranscripts' is deprecated.
## Use 'extractTranscriptSeqs' instead.
## See help("Deprecated")</pre>
```

#### Translate to amino acid sequence

all	(wid	l <mark>th</mark> (txx)	) %% 3 == 0) # sanity check
##	[1]	TRUE	
tra	nsla	te(txx)	# amino acid sequence
##	A	AAStrin	ngSet instance of length 20
##		width	seq
##	[1]	760	MDLSALRVEEVQNVINAMQKILECPICLELIKEPVSTKMCEAPVVTREWVLDSVALYQCQELDTYLIPQIPHSHY*
##	[2]	1793	MSLQESTRFSQLVEELLKIICAFQLDTGLEYANSYNFAMCEAPVVTREWVLDSVALYQCQELDTYLIPQIPHSHY*
##	[3]	174	MDAEFVCERTLKYFLGIAGGKWVVSYFWVTQSIKERKMMCEAPVVTREWVLDSVALYQCQELDTYLIPQIPHSHY*
##	[4]	700	MDLSALRVEEVQNVINAMQKILECPICLELIKEPVSTKCCYGPFTNMPTGCPPNCGCAARCLDRGQWLPCNWADV*
##	[5]	1817	MLKLLNQKKGPSQCPLCKNDITKRSLQESTRFSQLVEEMCEAPVVTREWVLDSVALYQCQELDTYLIPQIPHSHY*
##			••••
##	[16]	1365	MDLSALRVEEVQNVINAMQKILECPICLELIKEPVSTKSESQGVGLSDKELVSDDEERGTGLEENNQEEQSMDSNL
##	[17]	1365	MDLSALRVEEVQNVINAMQKILECPICLELIKEPVSTKSESQGVGLSDKELVSDDEERGTGLEENNQEEQSMDSNL
##	[18]	1318	MLKLLNQKKGPSQCPLCKNDITKRSLQESTRFSQLVEESESQGVGLSDKELVSDDEERGTGLEENNQEEQSMDSNL
##	[19]	1339	MDLSALRVEEVQNVINAMQKILECPICLELIKEPVSTKSESQGVGLSDKELVSDDEERGTGLEENNQEEQSMDSNL
##	[20]	1069	$\texttt{MNVEKAEFCNKSKQPGLARSQHNRWAGSKETCNDRRTP\ldotsSESQGVGLSDKELVSDDEERGTGLEENNQEEQSMDSNL}$

### 1.2 Summarizing overlaps

### Exercise 2

A basic operation in RNA-seq and other work flows is to count the number of times aligned reads overlap features of interest.

- a. Load the RNAseqData.HNRNPC.bam.chr14 experiment data package and get the paths to the BAM files it contains.
- b. Load the 'transcript db' package that contains the coordinates of each exon of the UCSC 'known genes' track of hg19.
- c. Extract the exon coordinates grouped by gene; the result is an GRangesList object that we will discuss more latter.
- d. Use the summarizeOverlaps function with the exon coordinates and BAM files to generate a count of the number of reads overlapping each gene. Visit the help page ?summarizeOverlaps to read about the counting strategy used.
- e. The counts can be extracted from the return value of summarizeOverlaps using the function assay. This is standard R matrix. How many reads overlapped regions of interest in each sample? How many genes had non-zero counts?

Solution: Point to BAM files

library(RNAseqData.HNRNPC.bam.chr14)
fls <- RNAseqData.HNRNPC.bam.chr14\_BAMFILES</pre>

Get the gene model; this could also come from, e.g., a GFF or GTF file.

```
library(BiocParallel)
library(TxDb.Hsapiens.UCSC.hg19.knownGene)
ex <- exonsBy(TxDb.Hsapiens.UCSC.hg19.knownGene, "gene")</pre>
```

Summarize the number of reads overlapping each region of interest

```
counts <- summarizeOverlaps(ex, fls)
colSums(assay(counts))
## ERR127306 ERR127307 ERR127308 ERR127309 ERR127302 ERR127303 ERR127304 ERR127305
## 340669 373302 371666 331540 313817 331160 331639 329672
sum(rowSums(assay(counts)) != 0)
## [1] 528
```

## References

[1] Michael Lawrence, Wolfgang Huber, Hervé Pagès, Patrick Aboyoun, Marc Carlson, Robert Gentleman, Martin<sup>T</sup>. Morgan, and Vincent<sup>J</sup>. Carey. Software for computing and annotating genomic ranges. *PLoS Comput Biol*, 9(8):e1003118, 08 2013. URL: http://dx.doi.org/10.1371%2Fjournal.pcbi.1003118, doi: 10.1371/journal.pcbi.1003118.