# Saying Hello to the Bioconductor Ranges Infrastructure

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#### Package

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## 1 Introduction

The primary purpose of the *HelloRanges* package, from the pedagogical perspective, is to map *bedtools* "recipes" to R scripts. The package was born out of the recognition that *Bioconductor* lacks a cookbook that explains how to achieve common, specific tasks using the Ranges infrastructure. And that writing a compiler is more fun (and hopefully more useful) than writing documentation. The goal is to enable those who already use R/Bioconductor for modeling and plotting to unify their workflow by integrating their upstream processing.

*HelloRanges* provides an *R* function corresponding to each *bedtools* command. The output is an *R* language object, and we can print the object to copy and integrate the code into an R script. Ideally, the code is first integrated (by copy/paste) into an R script. Unlike bedtools, the result of evaluation is an R object that is suitable for further analysis. There is no automatic output to files, because once we are in *R*, we want to stay there. Assuming that the reader is interested in learning Bioconductor, we encourage the reader to inspect the code prior to evaluation by printing the language object. The generated code is meant to be correct, readable, conformant to best practices and performant (in that order). While the code is much more verbose than the corresponding *bedtools* call, we argue that the explicit, low-level Ranges API has the advantage of being self-documenting and more flexible. And, of course, it directly integrates with the rest of *Bioconductor*.

Obviously, performing I/O with each operation will have a negative impact on performance, so it is recommended to import the data once, and perform subsequent operations on in-memory data structures. If memory is exhausted, consider distributing computations.

For the sake of comparison, this tutorial closely follows that of *bedtools* itself (http:// quinlanlab.org/tutorials/bedtools/bedtools.html). We will analyze the data from Maurano et al [1] assessment of Dnasel hypersensitivy across a range of fetal tissues (20 samples). The *bedtools* tutorial mostly consists of arbitrary range operations on the annotation tracks. Near the end, we will compare samples from the Maurano study in terms of their mutual overlap.

# 2 Data

The data are provided via the *HelloRangesData* package, which we load presently:

```
> library(HelloRanges)
```

```
> library(HelloRangesData)
```

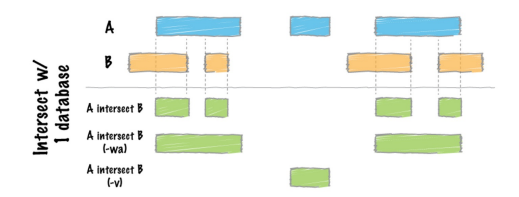
To have convenient paths to the data, we switch our working directory to the one with the data files:

```
> oldwd <- setwd(system.file("extdata", package="HelloRangesData"))</pre>
```

In our working directory are 20 BED files from the Dnasel study, as well as BED files representing the CpG islands ('cpg.bed'), Refseq exons ('exons.bed'), disease-associated SNPs ('gwas.bed'), and functional annotations output by chromHMM given ENCODE human embrionic stem cell ChIP-seq data ('hesc.chromHmm.bed'). There is also a 'hg19.genome' file indicating the chromosome lengths of the hg19 genome build.

One of the advantages of *R*, compared to the shell, is its unified package management system. *R* packages can contain data, and even completed analyses, in addition to libraries of functions. *Bioconductor* provides many annotations and sample datasets through packages. Packages make obtaining data easy, and they also help with reproducibility and provenance through versioning. Thus, they are more convenient and safer compared to downloading from live URLs. Some packages provide APIs to download data from repositories, and we can use them to programmatically generate data packages in a reproducible way.

For example, the TxDb packages provide transcript annotations for an individual reference genome. *Bioconductor* provides pre-built TxDb packages for the commonly used genome builds, and it is easy to generate a custom TxDb package using annotations from Biomart, UCSC, or just a GFF file. The *rtracklayer* package supports automated retrieval of any dataset from the UCSC table browser, without any pointing-and-clicking involved. We will demonstrate some of these tools later in the tutorial.



# 3 Overlap and Intersection

One of the most useful ways to compare two tracks is to determine which ranges overlap, and where they intersect (see the above image from the *bedtools* tutorial).

By default, *bedtools* outputs the region of intersection for each overlap between the two tracks. We compile the code for intersecting the CpG islands and the exons.

```
> code <- bedtools_intersect("-a cpg.bed -b exons.bed -g hg19.genome")
> code
{
    genome <- import("hg19.genome")
    gr_a <- import("cpg.bed", genome = genome)
    gr_b <- import("exons.bed", genome = genome)
    pairs <- find0verlapPairs(gr_a, gr_b, ignore.strand = TRUE)
    ans <- pintersect(pairs, ignore.strand = TRUE)
    ans
}</pre>
```

This code should be integrated into an R script that implements a larger workflow. For the purposes of this tutorial, we will call eval on the language object to yield the result:

> ans <- eval(code)								
> mcols(ans)	> mcols(ans)\$hit <- NULL							
> ans								
<b>60</b>					-			
GRanges object	ct with 45	500 ranges	and I	metadat	a column:			
sec	qnames	rar	nges st	rand	name			
	<rle></rle>	<irang< td=""><td>ges&gt; &lt;</td><td>Rle&gt;  </td><td><character></character></td><td></td></irang<>	ges> <	Rle>	<character></character>			
[1]	chr1	29321-29	9370	*	CpG:_116			
[2]	chr1	135125-135	5563	*	CpG:_30			
[3]	chr1	327791-328	3229	*	CpG:_29			
[4]	chr1	327791-328	3229	*	CpG:_29			
[5]	chr1	327791-328	3229	*	CpG:_29			
[45496]	chrY 592	13949-59214	4117	*	CpG:_36			
[45497]	chrY 592	13949-59214	4117	*	CpG:_36			
[45498]	chrY 592	13949-59214	4117	*	CpG:_36			
[45499]	chrY 592	13949-59214	4117	*	CpG:_36			
[45500]	chrY 592	13949-59214	4117	*	CpG:_36			
seqinfo: 93	3 sequence	s from hg19	9 genom	ie				

The result is an instance of *GRanges*, the central data structure in *Bioconductor* for genomic data. A *GRanges* takes the form of a table and resembles a BED file, with a column for the chromosome, start, end, strand. We will see *GRanges* a lot, along with its cousin, *GRangesList*, which stores what *bedtools* calls "split" ranges.

## 3.1 Sequence information

Consider the simplest invocation of *bedtools intersect*:

```
> code <- bedtools_intersect("-a cpg.bed -b exons.bed")
> code
{
    genome <- Seqinfo(genome = NA_character_)
    gr_a <- import("cpg.bed", genome = genome)
    gr_b <- import("exons.bed", genome = genome)
    pairs <- findOverlapPairs(gr_a, gr_b, ignore.strand = TRUE)
    ans <- pintersect(pairs, ignore.strand = TRUE)
    ans
}</pre>
```

The first line creates an object representing the structure of the genome build:

```
> genome <- eval(code[[2L]])
> genome
Seqinfo object with no sequences:
```

It is an empty object, because the genome identifier is unspecified (NA\_character\_). Having unspecified genome bounds is dangerous and leads to accidents involving incompatible genome builds. Besides error checking, the bounds are useful when computing coverage and finding the gaps (see below). Luckily, *bedtools* lets us specify the genome as an argument:

```
> bedtools_intersect("-a cpg.bed -b exons.bed -g hg19.genome")
{
    genome <- import("hg19.genome")
    gr_a <- import("cpg.bed", genome = genome)
    gr_b <- import("exons.bed", genome = genome)
    pairs <- find0verlapPairs(gr_a, gr_b, ignore.strand = TRUE)
    ans <- pintersect(pairs, ignore.strand = TRUE)
    ans
}</pre>
```

We now have a populated genome, since the tutorial had provided the 'hg19.genome' file. However, in general, information on the genome build should be centralized, not sitting somewhere in a file. *Bioconductor* provides *GenomeInfoDb* as its central source of sequence information. We can hook into that by passing the genome identifier instead of a file name:

```
> bedtools_intersect("-a cpg.bed -b exons.bed -g hg19")
{
    genome <- Seqinfo(genome = "hg19")
    gr_a <- import("cpg.bed", genome = genome)
    gr_b <- import("exons.bed", genome = genome)
    pairs <- find0verlapPairs(gr_a, gr_b, ignore.strand = TRUE)
    ans <- pintersect(pairs, ignore.strand = TRUE)
    ans
}</pre>
```

## 3.2 Annotations

The next step is the import of the CpG islands and exons using the import function from *rtracklayer*, which can load pretty much any kind of genomic data into the appropriate type of *Bioconductor* object. In this case, we are loading the data as a *GRanges*:

```
> gr_a
```

G

Ranges o	object with	n 28691	ranges	and	1 meta	data	column:
	seqnames		rar	nges	strand		name
	<rle></rle>		<irang< td=""><td>ges&gt;</td><td><rle></rle></td><td>  &lt;</td><td>character&gt;</td></irang<>	ges>	<rle></rle>	<	character>
[1]	] chr1	:	28736-29	9810	*	1	CpG:_116
[2]	] chr1	13	5125-135	5563	*	1	CpG:_30
[3]	] chr1	32	7791-328	3229	*	1	CpG:_29
[4]	] chr1	43	7152-438	3164	*	1	CpG:_84
[5]	] chr1	449	9274-450	9544	*	1	CpG:_99
						•	
[28687]	] chrY	276101	16-27613	1088	*	1	CpG:_76
[28688]	] chrY	285555	36-28555	5932	*	1	CpG:_32
[28689]	] chrY	287733	16-28773	3544	*	1	CpG:_25

[28690] chrY 59213795-59214183 \* | CpG:\_36 [28691] chrY 59349267-59349574 \* | CpG:\_29 -----seqinfo: 93 sequences from hg19 genome

The *rtracklayer* package can also download data directly from the UCSC table browser. For example, we could get the CpG islands directly:

```
> ucsc <- browserSession()
> genome(ucsc) <- "hg19"
> cpgs <- ucsc[["CpG Islands"]]</pre>
```

Gene annotations, including exon coordinates, should also be stored more formally than in a file, and Bioconductor provides them through its TxDb family of packages:

> library(TxDb.Hsapiens.UCSC.hg19.knownGene)

> exons <- exons(TxDb.Hsapiens.UCSC.hg19.knownGene)</pre>

#### 3.3 Finding Overlaps

The next step is to find all of the overlaps. The workhorse function is **findOverlaps** from the *IRanges* package. Here, we use the variant **findOverlapPairs**, a convenience for creating a *Pairs* object that matches up the overlapping ranges:

> pairs

Pairs	objec <sup>.</sup>	t with	45500	pairs	and	0	metadata	colu	imns:
				fi	rst				second
				<grange< td=""><td>es&gt;</td><td></td><td></td><td>&lt;</td><td>GRanges&gt;</td></grange<>	es>			<	GRanges>
	[1]	cl	hr1:28	736-298	310		chr1:	29321	-29370:-
	[2]	chr	1:13512	25-1355	563		chr1:13	4773-	139696:-
	[3]	chr	1:32779	91-3282	229		chr1:32	4439-	328581:+
	[4]	chr	1:32779	91-3282	229		chr1:32	4439-	328581:+
	[5]	chr	1:32779	91-3282	229		chr1:32	7036-	328581:+
[454	496] cl	rY:592	213795	-59214	183 (	chr	Y:592139	49-59	214117:+
[454	197] cl	rY:592	213795	-59214	183 (	chr	Y:592139	49-59	214117:+
[454	498] cl	rY:592	213795	-59214	183 (	chr	Y:592139	49-59	214117:+
[454	499] cl	rY:592	213795	-59214	183 (	chr	Y:592139	49-59	214117:+
[455	500] cl	nrY:592	213795	-592142	183 (	chr	Y:592139	49-59	214117:+

Although the ranges are displayed as succinct strings, they data are still represented as *GRanges* objects.

Users of *bedtools* will be familiar with *Pairs* as the analog of the BEDPE file format. We can use *rtracklayer* to export *Pairs* to BEDPE:

> export(pairs, "pairs.bedpe")

A key parameter to findOverlapPairs is ignore.strand=TRUE. By default, all operations on *GRanges* take strand into account when determining whether two ranges overlap, and deciding on the orientation of a range. This is surprising to many novice users, particularly to

those with *bedtools* experience. Most functions take the ignore.strand argument to control this behavior. To avoid confusion, the code generated by *HelloRanges* is always explicit about how it is treating strand. Users are encouraged to follow the same practice.

#### 3.4 Computing intersections

The final step is to find the actual intersecting region between the member of each overlapping pair. We do this with the **pintersect** function, which is the "parallel" or "pairwise" version of the default **intersect** function. If we had just called **intersect**(gr\_a, gr\_b) instead, the entire set of ranges would have been treated as a set, and overlapping ranges in gr\_a and gr\_b would have been merged (this is rarely desirable and requires an extra merge step in *bedtools*).

Notice again the importance of ignore.strand=TRUE. Without that, ranges on opposite strands would have zero intersection.

And here is our result:

> ans

GRanges	object	with	45500	ranges	and	1 metac	lata	column:
	seqn	ames		rar	iges	strand		name
	<	Rle>		<irang< td=""><td>es&gt;</td><td><rle></rle></td><td>  &lt;</td><td>character&gt;</td></irang<>	es>	<rle></rle>	<	character>
[1	1]	chr1	2	29321-29	370	*		CpG:_116
[2	2]	chr1	135	5125-135	563	*		CpG:_30
[3	3]	chr1	327	791-328	8229	*		CpG:_29
[4	4]	chr1	327	791-328	8229	*		CpG:_29
[5	5]	chr1	327	791-328	8229	*		CpG:_29
[45496	5]	chrY !	5921394	49-59214	117	*		CpG:_36
[45497	7]	chrY !	5921394	49-59214	117	*		CpG:_36
[45498	3]	chrY !	5921394	49-59214	117	*		CpG:_36
[45499	9]	chrY !	5921394	49-59214	117	*		CpG:_36
[45500	9]	chrY !	5921394	49-59214	117	*		CpG:_36
seqint	fo: 93	seque	nces fr	om hg19	) ger	nome		

Again, a *GRanges* object. The hit column indicates whether the pair overlapped at all (as opposed to one range being of zero width). It's useless in this case.

## 3.5 Keeping the original features

To keep the original form of the overlapping features, the generated code simply neglects to call **pintersect** and ends up with the **pairs** object introduced previously:

```
> bedtools_intersect("-a cpg.bed -b exons.bed -g hg19 -wa -wb")
{
    genome <- Seqinfo(genome = "hg19")
    gr_a <- import("cpg.bed", genome = genome)
    gr_b <- import("exons.bed", genome = genome)
    pairs <- find0verlapPairs(gr_a, gr_b, ignore.strand = TRUE)</pre>
```

}

```
ans <- pairs
ans
```

## 3.6 Computing the amount of overlap

To compute the width of the overlapping regions, we query the initial result for its width and store as an annotation on the pairs:

```
> bedtools_intersect("-a cpg.bed -b exons.bed -g hg19 -wo")
{
    genome <- Seqinfo(genome = "hg19")
    gr_a <- import("cpg.bed", genome = genome)
    gr_b <- import("exons.bed", genome = genome)
    pairs <- findOverlapPairs(gr_a, gr_b, ignore.strand = TRUE)
    ans <- pairs
    mcols(ans)$overlap_width <- width(pintersect(ans, ignore.strand = TRUE))
    ans
}</pre>
```

This code reveals that *GRanges*, along with every other vector-like object in the Ranges infrastructure, is capable of storing tabular annotations, accessible via mcols. We actually saw this before with the "name" column on the Cpg Islands. Here, we use it to store the overlap width.

## 3.7 Counting the number of overlaps

A common query, particularly in RNA-seq analysis, is how many ranges in the subject overlap each query range. The countOverlaps function serves this particular purpose:

```
> bedtools_intersect("-a cpg.bed -b exons.bed -g hg19 -c")
{
    genome <- Seqinfo(genome = "hg19")
    gr_a <- import("cpg.bed", genome = genome)
    gr_b <- import("exons.bed", genome = genome)
    ans <- gr_a
    mcols(ans)$overlap_count <- countOverlaps(gr_a, gr_b, ignore.strand = TRUE)
    ans
}</pre>
```

## 3.8 Excluding queries with overlaps

We might instead want to exclude all query ranges that overlap any subject range, i.e., any CpG island that overlaps an exon. The subsetByOverlaps function is tasked with restricting by overlap. By passing invert=TRUE, we exclude ranges with overlaps.

```
> bedtools_intersect("-a cpg.bed -b exons.bed -g hg19 -v")
{
    genome <- Seqinfo(genome = "hg19")
    gr_a <- import("cpg.bed", genome = genome)
    gr_b <- import("exons.bed", genome = genome)
    subsetByOverlaps(gr_a, gr_b, invert = TRUE, ignore.strand = TRUE)
}</pre>
```

## 3.9 Restricting by fraction of overlap

The *bedtools* suite has deep support for restricting overlaps by the fraction of the query/subject range that is overlapped. This is not directly supported by the *Bioconductor* infrastructure, but we can filter post-hoc:

```
> bedtools_intersect("-a cpg.bed -b exons.bed -g hg19 -f 0.5 -wo")
{
    genome <- Seqinfo(genome = "hg19")
    gr_a <- import("cpg.bed", genome = genome)
    gr_b <- import("exons.bed", genome = genome)
    pairs <- find0verlapPairs(gr_a, gr_b, ignore.strand = TRUE)
    olap <- pintersect(pairs, ignore.strand = TRUE)
    keep <- width(olap)/width(first(pairs)) >= 0.5
    pairs <- pairs[keep]
    ans <- pairs
    mcols(ans)$overlap_width <- width(olap)[keep]
    ans
}</pre>
```

## 3.10 Performance

Comparing the performance of *bedtools* and *IRanges* is like comparing apples and oranges. The typical *Bioconductor* workflow imports the data once, paying an upfront cost, and then operates efficiently on in-memory data structures. The BED parser is implemented in R code and will not compete with the parsing performance of special purpose C code. The intersect operation itself is also slower than *bedtools*, but it's still reasonably close for being mostly implemented in R.

```
> a <- import("exons.bed")
> b <- import("hesc.chromHmm.bed")
> system.time(pintersect(findOverlapPairs(a, b, ignore.strand=TRUE),
+ ignore.strand=TRUE))
user system elapsed
0.628 0.016 0.645
```

## 3.11 Multiple subjects

Often, we are interested in intersections with mutiple annotation tracks, or multiple samples. Note that the command line parser used by *helloRanges* requires that the filenames be comma-separated, instead of space-separated. This is probably more readable anyway.

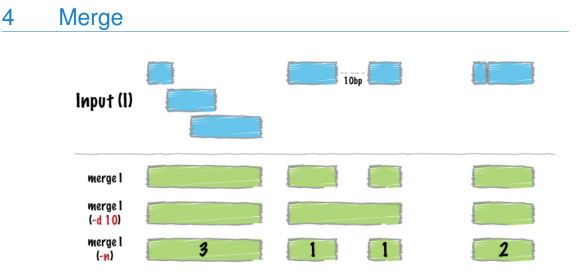
```
> code <- bedtools_intersect(</pre>
+
      paste("-a exons.bed",
              "-b cpg.bed,gwas.bed,hesc.chromHmm.bed -wa -wb -g hg19",
+
+
             "-names cpg,gwas,chromhmm"))
> ans <- eval(code)</pre>
> code
{
    genome <- Seqinfo(genome = "hg19")</pre>
    gr_a <- import("exons.bed", genome = genome)</pre>
    b <- c("cpg.bed", "gwas.bed", "hesc.chromHmm.bed")</pre>
    names(b) <- c("cpg", "gwas", "chromhmm")</pre>
    bl <- List(lapply(b, import, genome = genome))</pre>
    gr_b <- stack(bl, "b")</pre>
    pairs <- findOverlapPairs(gr_a, gr_b, ignore.strand = TRUE)</pre>
    ans <- pairs
    ans
}
```

Inspecting the code, we see that we need to loop over the database files and then **stack** them into a single *GRanges* grouped by the column "b":

```
> second(ans)
```

> second(ans)							
GRanges object w	ith 541210 ranges and	2 metadata columns	5:				
seqnar	mes ranges	strand   b	name				
<r]< td=""><td>le&gt; <iranges></iranges></td><td><rle>   <rle></rle></rle></td><td><character></character></td></r]<>	le> <iranges></iranges>	<rle>   <rle></rle></rle>	<character></character>				
[1] cł	hr1 11538-11937	*   chromhmm	11_Weak_Txn				
[2] cł	hr1 11938-12137	*   chromhmm	14_Repetitive/CNV				
[3] cł	hr1 12138-14137	*   chromhmm	11_Weak_Txn				
[4] cł	hr1 12138-14137	*   chromhmm	11_Weak_Txn				
[5] cł	hr1 12138-14137	*   chromhmm	11_Weak_Txn				
[541206] cł	hrY 59213795-59214183	*   cpg	CpG:_36				
[541207] cł	hrY 59213795-59214183	*   cpg	CpG:_36				
[541208] cł	hrY 59213795-59214183	*   cpg	CpG:_36				
[541209] cł	hrY 59213795-59214183	*   cpg	CpG:_36				
[541210] cł	hrY 59213795-59214183	*   cpg	CpG:_36				
seqinfo: 298 se	equences (2 circular)	from hg19 genome					

The "b" column is an *Rle* object, a run-length encoded form of an ordinary R vector, in this case a factor. Since the data are sorted into groups, this encoding is more efficient than a dense representation. The "seqnames" and "strand" columns also benefit from run-length encoding. Not only can we fit more data into memory, many operations become faster.



There are many ways to summarize interval data. In the Ranges infrastructure, we call some of them range (min start to max end), reduce (bedtools merge), disjoin (involved in bedtools multiinter) and coverage (counting the number of ranges overlapping each position, bedtools genomecov). We are presently concerned with reduce, which combines overlapping and adjacent ranges into a single range. The corresponding bedtools merge command requires the data to be sorted; however, reduce does not have this constraint.

```
> bedtools_merge("-i exons.bed")
{
   genome <- Seqinfo(genome = NA_character_)
   gr_a <- import("exons.bed", genome = genome)
   ans <- reduce(gr_a, ignore.strand = TRUE)
   ans</pre>
```

```
}
```

# 4.1 Aggregation

As with any reduction, we often want to simultaneously aggregate associated variables and report the summaries in the result.

We count the number of ranges overlapping each merged range:

```
> code <- bedtools_merge("-i exons.bed -c 1 -o count")
> code
{
    genome <- Seqinfo(genome = NA_character_)
    gr_a <- import("exons.bed", genome = genome)
    ans <- reduce(gr_a, ignore.strand = TRUE, with.revmap = TRUE)
    mcols(ans) <- aggregate(gr_a, mcols(ans)$revmap, seqnames.count = lengths(seqnames),
        drop = FALSE)
    ans
}</pre>
```

The key to aggregation with reduce is the with.revmap=TRUE argument. That yields a "revmap" column on the result. It is an *IntegerList* holding the subscripts corresponding to each group. We pass it to aggregate to indicate the grouping. The named arguments to aggregate, in this case seqnames.count, are effectively evaluated with respect to each group (although they are actually evaluated only once).

This yields the result:

```
> eval(code)
```

GRanges objec	ct with 2	29241 ranges and	2 metadata co	olumns:
Se	eqnames	ranges	strand	grouping
	<rle></rle>	<iranges></iranges>	<rle>   <mar< td=""><td>yToManyGrouping&gt;</td></mar<></rle>	yToManyGrouping>
[1]	chr1	11874-12227	*	1
[2]	chr1	12613-12721	*	2
[3]	chr1	13221-14829	*	3,4
[4]	chr1	14970-15038	*	5
[5]	chr1	15796-15947	*	6
[229237]	chrY 5	59337091-59337236	*	459744,459745
[229238]	chrY 5	59337949-59338150	*	459746,459747
[229239]	chrY 5	59338754-59338859	*	459748,459749
[229240]	chrY 5	59340194-59340278	*	459750
[229241]	chrY 5	59342487-59343488	*	459751,459752
Se	eqnames.c	count		
	<inte< td=""><td>eger&gt;</td><td></td><td></td></inte<>	eger>		
[1]		1		
[2]		1		
[3]		2		
[4]		1		
[5]		1		
[229237]		2		
[229238]		2		
[229239]		2		
[229240]		1		
[229241]		2		
seqinfo: 49	) sequenc	ces from an unspec	cified genome;	no seqlengths

We see that the grouping has been preserved on the object, in case we wish to aggregate further through joins.

Counting the overlaps by counting the "seqnames" is a little circuitous. Instead, we could have just counted the elements in each group:

> identical(lengths(ans\$grouping), ans\$seqnames.count)

[1] TRUE

Note that this counting is very fast, because the "revmap" *IntegerList* is not actually a list, but a partitioned vector, and the partitioning already encodes the counts. This is an example of where the flexibility and efficient in-memory representations of *Bioconductor* are particularly effective.

#### 4.2 Merging close features

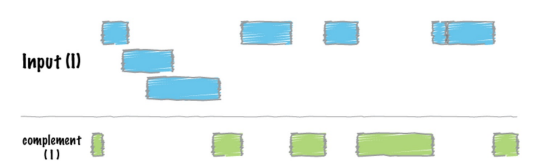
By default, features are merged if they are overlapping or adjacent, i.e., the min.gapwidth (the minimum gap allowed to not be merged) is 1. To merge features that are up to, say, 1000bp away, we need to pass min.gapwidth=1001:

```
> bedtools_merge("-i exons.bed -d 1000")
{
    genome <- Seqinfo(genome = NA_character_)
    gr_a <- import("exons.bed", genome = genome)
    ans <- reduce(gr_a, ignore.strand = TRUE, min.gapwidth = 1001L)
    ans
}</pre>
```

Here is another example showing how to merge multiple columns at once:

```
> bedtools_merge("-i exons.bed -d 90 -c 1,4 -o count,collapse")
{
    genome <- Seqinfo(genome = NA_character_)
    gr_a <- import("exons.bed", genome = genome)
    ans <- reduce(gr_a, ignore.strand = TRUE, with.revmap = TRUE,
        min.gapwidth = 91L)
    mcols(ans) <- aggregate(gr_a, mcols(ans)$revmap, seqnames.count = lengths(seqnames),
        name.collapse = unstrsplit(name, ","), drop = FALSE)
    ans
}</pre>
```

## 5 Finding the Gaps

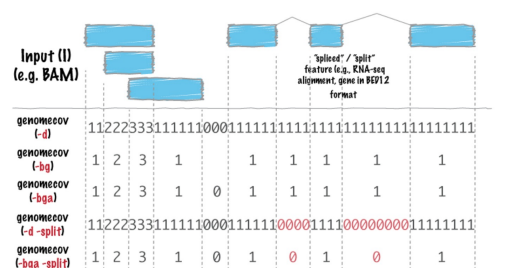


The *bedtools complement* tool finds the gaps in the sequence, i.e., the regions of sequence the track does not cover. This is where having the sequence bounds is critical.

```
> bedtools_complement("-i exons.bed -g hg19.genome")
{
    genome <- import("hg19.genome")
    gr_a <- import("exons.bed", genome = genome)
    ans <- setdiff(as(seqinfo(gr_a), "GRanges"), unstrand(gr_a))
    ans
}</pre>
```

The call to setdiff is a set operation, along with intersect and union. Set operations behave a bit surprisingly with respect to strand. The "unstranded" features, those with "\*" for their strand, are considered to be in a separate space from the stranded features. If we pass ignore.strand=TRUE, both arguments are unstranded and the result is unstranded (strand information is discarded). This makes sense, because there is no obvious way to merge a stranded and unstranded feature. Since we are looking for the gaps, we do not care about the strand, so discarding the strand is acceptable. Best practice is to make this explicit by calling unstrand instead of assuming the reader understands the behavior of ignore.strand=TRUE.

# 6 Computing Genomic Coverage



One of the useful ways to summarize ranges, particularly alignments, is to count how many times a position is overlapped by a range. This is called the coverage. Unlike *bedtools genomecov*, we do not require the data to be sorted.

### 6.1 Coverage vector

To compute the coverage, we just call coverage. For consistency with *bedtools*, which drops zero runs with the "-bg" option, we convert the coverage vector to a *GRanges* and subset:

```
> bedtools_genomecov("-i exons.bed -g hg19.genome -bg")
```

```
{
    genome <- import("hg19.genome")
    gr_a <- import("exons.bed", genome = genome)
    cov <- coverage(gr_a)
    ans <- GRanges(cov)
    ans <- subset(ans, score > 0)
    ans
}
```

### 6.2 Coverage histogram

The default behavior of *genomecov* is to compute a histogram showing the number and fraction of positions covered at each level. It does this for the individual chromosome, and the entire genome. While computing the coverage vector is as simple as calling coverage, forming the histogram is a bit complicated. This is a bit esoteric, but it lets us demonstrate how to aggregate data in R:

```
> code <- bedtools_genomecov("-i exons.bed -g hg19.genome")
> ans <- eval(code)
> code
{
    genome <- import("hg19.genome")
    gr_a <- import("exons.bed", genome = genome)
    cov <- coverage(gr_a)
    tablist <- List(lapply(cov, table))
    mcols(tablist)$len <- lengths(cov, use.names = FALSE)
    covhist <- stack(tablist, "seqnames", "count", "coverage")
    margin <- aggregate(covhist, ~coverage, count = sum(NumericList(count)))[-1L]
    margin <- DataFrame(seqnames = Rle("genome"), margin, len = sum(as.numeric(lengths(cov))))
    covhist <- rbind(covhist, margin)
    ans <- within(covhist, fraction <- count/len)
    ans
```

```
}
```

The cov object is an *RleList*, with one *Rle* per sequence (chromosome).

```
> cov
RleList of length 93
$chr1
integer-Rle of length 249250621 with 46230 runs
  Lengths: 11873
                 354
                        385
                             109
                                    499 ...
                                              64 2722
                                                         677 1868 37276
  Values :
                    1
                          0
                                1
                                     0 ...
                                               2
                                                     0
                                                        1
                                                                 2
              0
                                                                      0
$chr10
integer-Rle of length 135534747 with 18962 runs
  Lengths: 92827 1227
                             111
                                    78 ... 1936 1120
                        500
                                                        3293
                                                               260 36289
  Values :
              0
                 1
                          0
                               1
                                     0 ...
                                               0
                                                     8
                                                           0
                                                                 8
                                                                      0
```

```
$chr11
```

integer-Rle of length 135006516 with 25138 runs 4387 93 58 227 ... Lengths: 126986 104 22067 864 630961 Values : 0 1 0 ... 1 1 0 1 0 0 \$chr11\_gl000202\_random integer-Rle of length 40103 with 1 run Lengths: 40103 Values : 0 \$chr12 integer-Rle of length 133851895 with 25290 runs Lengths: 87983 34 239 136 177 ... 2463 271 1428 30 39473 0 ... Values : 1 0 1 0 1 1 0 0 0 <88 more elements>

We tabulate each coverage vector individually, then stack the tables into an initial histogram. Then, we aggregate over the entire genome and combine the genome histogram with the perchromosome histogram. The call to NumericList is only to avoid integer overflow. Finally, we compute the fraction covered and end up with:

> ans

DataFrame with 653 rows and 5 columns							
	seqnames	coverage	count	len	fraction		
	<rle></rle>	<factor></factor>	<numeric></numeric>	<numeric></numeric>	<numeric></numeric>		
1	1	0	241996316	249250621	0.97089554		
2	1	1	4276763	249250621	0.01715848		
3	1	2	1475526	249250621	0.00591985		
4	1	3	710135	249250621	0.00284908		
5	1	4	388193	249250621	0.00155744		
649	genome	73	135	3137161264	4.30325e-08		
650	genome	74	263	3137161264	8.38338e-08		
651	genome	75	1921	3137161264	6.12337e-07		
652	genome	76	705	3137161264	2.24725e-07		
653	genome	77	2103	3137161264	6.70351e-07		

This takes 3 minutes for bedtools, but closer to 3 seconds for us, probably because it is working too hard to conserve memory.

# 7 Combining operations

## 7.1 Chaining

Most real-world workflows involve multiple operations, chained together. The R objects produced *HelloRanges* can be passed directly to existing R functions, and *HelloRanges* defines an ordinary R function corresponding to each bedtools operation. The arguments of the

function correspond to *bedtools* arguments, except they can be R objects, like *GRanges*, in addition to filenames. These functions with ordinary R semantics are prefixed by  $R_{-}$ , so the analog to <u>bedtools\_intersect</u> is <u>R\_bedtools\_intersect</u>.

Consider a use case similar to the one mentioned in the *bedtools* tutorial: find the regions of the CpG islands that are not covered by exons. We could do this directly with **bed** tools\_subtract, but let us instead compute the coverage of the exons, find the regions of zero coverage, and intersect those with the CpG islands.

First, we generate the code for the coverage operation (and ideally copy it to a script). The result of evaluating that code is a *GRanges*, which we subset for the regions with zero score.

```
> code <- bedtools_genomecov("-i exons.bed -g hg19.genome -bga")</pre>
> gr0 <- subset(eval(code), score == 0L) # compare to: awk '$4==0'</pre>
> gr0
GRanges object with 229334 ranges and 1 metadata column:
           segnames
                                ranges strand |
                                                     score
              <Rle>
                            <IRanges> <Rle> | <integer>
       [1]
               chr1
                               1-11873
                                             * |
                                                         0
       [2]
               chr1
                          12228-12612
                                             * |
                                                         0
       [3]
               chr1
                          12722-13220
                                                         0
                                             * |
       [4]
               chr1
                          14830-14969
                                             * |
                                                         0
               chr1
                           15039-15795
                                                         0
       [5]
                                             * |
                . . .
       . . .
                                    . . .
                                           ... .
                                                        . . .
  [229330]
               chrY 59337237-59337948
                                             * |
                                                         0
               chrY 59338151-59338753
                                                         0
  [229331]
                                             * |
               chrY 59338860-59340193
                                                         0
  [229332]
                                             * |
                                                         0
  [229333]
               chrY 59340279-59342486
                                             * |
  [229334]
               chrY 59343489-59373566
                                             * |
                                                         0
  - - - - - - -
  seqinfo: 93 sequences from an unspecified genome
```

Next, we pass gro directly to the R analog of *intersect*, R\_bedtools\_intersect:

```
> code <- R_bedtools_intersect("cpg.bed", gr0)
> code
{
    genome <- Seqinfo(genome = NA_character_)
    gr_a <- import("cpg.bed", genome = genome)
    gr_b <- gr0
    pairs <- find0verlapPairs(gr_a, gr_b, ignore.strand = TRUE)
    ans <- pintersect(pairs, ignore.strand = TRUE)
    ans
}</pre>
```

The generated code already refers to gr0 explicitly, so it is easy to copy this into the script.

To generalize, the chaining workflow is:

- 1. Generate code for first operation,
- 2. Integrate and evaluate the code,
- 3. Interactively inspect the result of evaluation,

- 4. Perform intermediate operations, while inspecting results,
- 5. Call R analog to generate second stage code.

Generating and integrating R code is the best way to learn, and the best way to produce a readable, flexible and performant script. However, there are probably those who are tempted to evaluate the code directly, as we have done in this vignette. Further, there are those who wish to chain these operations together with the so-called "pipe" operator, because it would come so tantalizing close to the syntax of the shell. Thus, we created a third family of functions, prefixed by do., which provide the same interface as the R\_ family, except they evaluate the generated code:

```
> do_bedtools_genomecov("exons.bed", g="hg19.genome", bga=TRUE) %>%
```

```
+ subset(score > OL) %>%
```

+ do\_bedtools\_intersect("cpg.bed", .)

#### 7.2 Coalescence

In the previous section, we chained together independent operations. Having access to the underlying code gives us the flexibility to merge operations so that they are faster than the sum of their parts. We call this coalescence.

Consider a use case cited by the *bedtools* tutorial: compute the distribution of coverage over all exons. To integrate better with this tutorial, we adapt that to finding the distribution of exon coverage over all CpG islands.

We could mimic the example by computing the coverage complete histogram and extracting only the margin:

```
> bedtools_coverage("-a cpg.bed -b exons.bed -hist -g hg19.genome")
{
    genome <- import("hg19.genome")</pre>
    gr_a <- import("cpg.bed", genome = genome)</pre>
    gr_b <- import("exons.bed", genome = genome)</pre>
    cov <- unname(coverage(gr_b)[gr_a])</pre>
    tab <- t(table(cov))</pre>
    tab <- cbind(tab, all = rowSums(tab))</pre>
    covhist <- DataFrame(as.table(tab))</pre>
    colnames(covhist) <- c("coverage", "a", "count")</pre>
    len <- c(lengths(cov, use.names = FALSE), sum(lengths(cov)))</pre>
    covhist$len <- rep(len, each = nrow(tab))</pre>
    covhist <- subset(covhist, count > 0L)
    covhist$fraction <- with(covhist, count/len)</pre>
    ans <- gr_a
    covhistList <- split(covhist, ~a)[, -2L]</pre>
    mcols(ans)$coverage <- head(covhistList, -1L)</pre>
    metadata(ans)$coverage <- covhistList$all</pre>
    ans
```

}

The code is quite complex, because the Ranges infrastructure does not attempt to generate high-level summaries of the data. The rationale, which is validated in this case, is that the desired summary depends on the specific question, and the number of questions is effectively infinite. In this case, we only care about the margin, i.e., metadata(ans)\$coverage.

Thus, we can simplify the code. We begin with the same lines:

> genome <- import("hg19.genome") > gr\_a <- import("cpg.bed", genome = genome) > gr\_b <- import("exons.bed", genome = genome) > cov <- unname(coverage(gr\_b)[gr\_a])</pre>

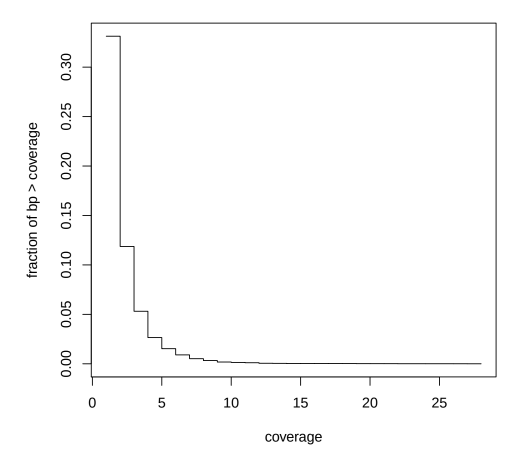
And summarize all of the coverage at once:

```
> all_cov <- unlist(cov)
> df <- as.data.frame(table(coverage=all_cov))
> df$fraction <- df$Freq / length(all_cov)</pre>
```

This is much faster, because we are only computing one table, not 30,000, and the table method for *Rle* is very efficient.

We now have a simple *data.frame* that we can plot as an inverted ECDF:

> plot((1-cumsum(fraction)) ~ as.integer(coverage), df, type="s", + ylab = "fraction of bp > coverage", xlab="coverage")



# 8 Jaccard Statistic

In order to compare the Dnasel hypersenstivity across tissues, we will employ the *bedtools jaccard* statistic, a measure of similarity between two tracks. It is defined as the total width of their intersection over the total width of their union.

We might expect, for example, that the similarity within a tissue is higher than that between two tissues, and this is indeed the case:

```
> code <- bedtools_jaccard(
+    paste("-a fHeart-DS16621.hotspot.twopass.fdr0.05.merge.bed",
+          "-b fHeart-DS15839.hotspot.twopass.fdr0.05.merge.bed"))
> heart_heart <- eval(code)
> code <- bedtools_jaccard(
+        paste("-a fHeart-DS16621.hotspot.twopass.fdr0.05.merge.bed",
+          "-b fSkin_fibro_bicep_R-DS19745.hg19.hotspot.twopass.fdr0.05.merge.bed"))
> heart_skin <- eval(code)</pre>
```

The generated code makes the statistic self-documenting:

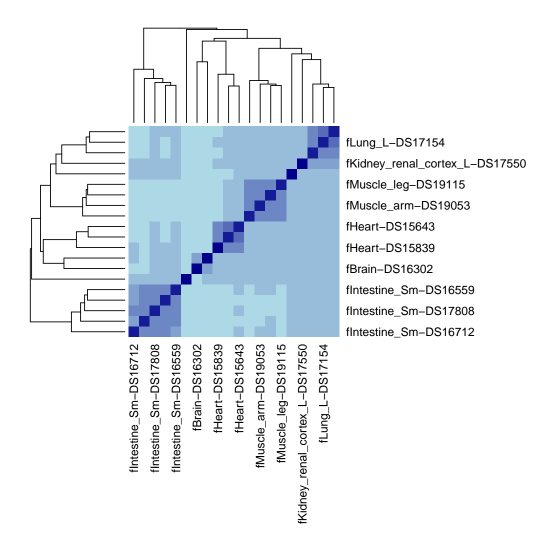
```
> code
{
    genome <- NA_character_
    gr_a <- import("fHeart-DS16621.hotspot.twopass.fdr0.05.merge.bed",
        genome = genome)
    gr_b <- import("fSkin_fibro_bicep_R-DS19745.hg19.hotspot.twopass.fdr0.05.merge.bed",
        genome = genome)
    intersects <- intersect(gr_a, gr_b, ignore.strand = TRUE)
    intersection <- sum(width(intersects))
    union <- sum(width(union(gr_a, gr_b, ignore.strand = TRUE)))
    ans <- DataFrame(intersection, union, jaccard = intersection/union,
        n_intersections = length(intersects))
    ans
}</pre>
```

We can compute the statistic over all pairs of samples using functionality included with R, through the *parallel* package. There is no need to learn yet another syntax, such as that of the *parallel* UNIX utility. Nor do we need to download a custom python script, and repeatedly call perl and awk.

```
> files <- Sys.glob("*.merge.bed")
> names(files) <- sub("\\..*", "", files)
> ncores <- if (.Platform$0S.type == "windows") 1L else 4L
> ans <- outer(files, files,
+ function(a, b) mcmapply(do_bedtools_jaccard, a, b,
+ mc.cores=ncores))
> jaccard <- apply(ans, 1:2, function(x) x[[1]]$jaccard)</pre>
```

Since we are already in R, it is easy to create a simple plot:

```
> palette <- colorRampPalette(c("lightblue", "darkblue"))(9)
> heatmap(jaccard, col=palette, margin=c(14, 14))
```



## 9 Exercises

These were adapted from the *bedtools* tutorial. Try to complete these exercises using *Bioconductor* directly.

- 1. Create a *GRanges* containing the non-exonic regions of the genome.
- 2. Compute the average distance from the GWAS SNPs to the closest exon (Hint: ?bed tools\_closest and ?distanceToNearest).
- Compute the exon coverage in 500kb windows across the genome (Hint: ?bedtools\_makewindows and ?tileGenome).
- 4. How many exons are completely overlapped by an enhancer (from 'hesc.chromHmm.bed') (Hint: ¿%within%)?
- 5. What fraction of the disease-associated SNPs are exonic (Hint: (Hint: ¿%over%))?

- Create intervals representing the canonical 2bp splice sites on either side of each exon (bonus: exclude splice sites at the first and last exons) (Hint: ?bedtools\_flank, ?in tronsByTranscript).
- 7. Which hESC ChromHMM state represents the most number of base pairs in the genome? (Hint: ?xtabs).

#### 9.1 Answers

Below, we give the *bedtools*-style answer first, followed by the essential call against the *Bioconductor* API.

First, we load the files into R objects for convenience:

```
> genome <- import("hg19.genome")
> exons <- import("exons.bed", genome=genome)
> gwas <- import("gwas.bed", genome=genome)
> hesc.chromHmm <- import("hesc.chromHmm.bed", genome=genome)</pre>
```

Here are the numbered answers:

```
1. > bedtools_complement("-i exons.bed -g hg19.genome")
  {
      genome <- import("hg19.genome")</pre>
      gr_a <- import("exons.bed", genome = genome)</pre>
      ans <- setdiff(as(seqinfo(gr_a), "GRanges"), unstrand(gr_a))</pre>
      ans
  }
  > ## or without HelloRanges:
  > setdiff(as(seqinfo(exons), "GRanges"), unstrand(exons))
  GRanges object with 229334 ranges and 0 metadata columns:
              seqnames
                                   ranges strand
                               <IRanges> <Rle>
                 <Rle>
          [1]
                  chr1
                                 1-11873
                                               *
          [2]
                  chr1
                             12228-12612
          [3]
                  chr1
                             12722-13220
                  chr1
                             14830-14969
          [4]
          [5]
                 chr1
                             15039-15795
                                               *
          . . .
                  . . .
                                      . . .
                                             . . .
     [229330]
                 chrY 59337237-59337948
                                              *
     [229331]
                 chrY 59338151-59338753
                 chrY 59338860-59340193
     [229332]
     [229333]
                  chrY 59340279-59342486
                 chrY 59343489-59373566
     [229334]
                                               *
     seqinfo: 93 sequences from hg19 genome
```

2. > bedtools\_closest("-a gwas.bed -b exons.bed -d")

```
{
       genome <- Seqinfo(genome = NA_character_)</pre>
       gr_a <- import("gwas.bed", genome = genome)</pre>
       gr_b <- import("exons.bed", genome = genome)</pre>
       hits <- nearest(gr_a, gr_b, ignore.strand = TRUE, select = "all")</pre>
       ans <- pair(gr_a, gr_b, hits, all.x = TRUE)</pre>
       mcols(ans)$distance <- distance(ans)</pre>
       ans
  }
  > ## or
  > distanceToNearest(gwas, exons)
  Hits object with 17674 hits and 1 metadata column:
             queryHits subjectHits | distance
             <integer> <integer> | <integer>
         [1]
                   1
                              235 |
                                          1319
                    2
                               249 |
         [2]
                                              0
                    3
         [3]
                              426
                                              0
                   4
         [4]
                              1163 |
                                           2385
                   5
         [5]
                             1163 |
                                          2894
         . . .
                  . . .
                               ... .
                                            . . .
     [17670] 17676
                            455399 |
                                            371
     [17671] 17677
                            455399 |
                                            371
     [17672]
             17678
                            455464 |
                                          5898
     [17673]
                 17679
                            455674 |
                                         324706
     [17674]
                 17680
                            456097 |
                                         135628
     - - - - - - - -
     queryLength: 17680 / subjectLength: 459752
3. > code <- bedtools_makewindows("-g hg19.genome -w 500000")
  > code
  {
       genome <- import("hg19.genome")</pre>
       ans <- tile(as(genome, "GRanges"), width = 500000L)</pre>
       ans
  }
  > windows <- unlist(eval(code))</pre>
  > R_bedtools_intersect(windows, exons, c=TRUE)
  {
       genome <- Seqinfo(genome = NA_character_)</pre>
       gr_a <- windows
      gr_b <- exons
      ans <- gr_a
      mcols(ans)$overlap_count <- count0verlaps(gr_a, gr_b, ignore.strand = TRUE)</pre>
       ans
  }
  > ## or
  > str(countOverlaps(tileGenome(seqinfo(exons), tilewidth=500000),
```

```
exons))
   +
   int [1:6275] 37 197 477 445 209 96 83 271 9 12 ...
4. > bedtools_intersect(
         paste("-a exons.bed -b <\"grep Enhancer hesc.chromHmm.bed\"",</pre>
  +
   +
               "-f 1.0 -wa -u"))
   {
       genome <- Seqinfo(genome = NA_character_)</pre>
       gr_a <- import("exons.bed", genome = genome)</pre>
       gr_b <- import(BEDFile(pipe("\"grep Enhancer hesc.chromHmm.bed\"")),</pre>
           genome = genome)
       hits <- findOverlaps(gr_a, gr_b, ignore.strand = TRUE, type = "within")
       gr_a[countQueryHits(hits) > 0L]
   }
  > quote(length(ans))
  length(ans)
  > ## or
  > sum(exons %within%
         subset(hesc.chromHmm, grepl("Enhancer", name)))
   +
   [1] 13746
5. > bedtools_intersect("-a gwas.bed -b exons.bed -u")
   {
       genome <- Seqinfo(genome = NA_character_)</pre>
       gr_a <- import("gwas.bed", genome = genome)</pre>
       gr_b <- import("exons.bed", genome = genome)</pre>
       subsetByOverlaps(gr_a, gr_b, ignore.strand = TRUE)
   }
  > quote(length(gr_a)/length(ans))
  length(gr_a)/length(ans)
  > ## or
  > mean(gwas %over% exons)
   [1] 0.09191176
6. > bedtools_flank("-l 2 -r 2 -i exons.bed -g hg19.genome")
   {
       genome <- import("hg19.genome")</pre>
       gr_a <- import("exons.bed", genome = genome)</pre>
       left <- flank(gr_a, 2, ignore.strand = TRUE)</pre>
       right <- flank(gr_a, 2, start = FALSE, ignore.strand = TRUE)
       ans <- zipup(Pairs(left, right))</pre>
       ans
```

```
}
```

```
> ## or, bonus:
  > txid <- sub("_exon.*", "", exons$name)</pre>
  > tx <- split(exons, txid)</pre>
  > bounds <- range(tx)</pre>
  > transpliced <- lengths(bounds) > 1
  > introns <- unlist(psetdiff(unlist(bounds[!transpliced]),</pre>
                                tx[!transpliced]))
  +
  > Pairs(resize(introns, 2L), resize(introns, 2L, fix="end"))
  Pairs object with 386125 pairs and 0 metadata columns:
                                   first
                                                              second
                               <GRanges>
                                                           <GRanges>
          [1]
                 chr12:9220777-9220778:- chr12:9220436-9220437:-
                 chr12:9221334-9221335:- chr12:9220821-9220822:-
          [2]
                 chr12:9222339-9222340:- chr12:9221439-9221440:-
          [3]
          [4]
                chr12:9223082-9223083:- chr12:9222410-9222411:-
          [5]
                 chr12:9224953-9224954:- chr12:9223175-9223176:-
          . . .
                                      . . .
                                                                  . . .
     [386121] chr9:123626325-123626326:- chr9:123625028-123625029:-
     [386122] chr9:123627986-123627987:- chr9:123626395-123626396:-
     [386123] chr9:123628304-123628305:- chr9:123628109-123628110:-
     [386124] chr9:123629146-123629147:- chr9:123628375-123628376:-
     [386125] chr9:123631084-123631085:- chr9:123629244-123629245:-
  > ## better way to get introns:
  > library(TxDb.Hsapiens.UCSC.hg19.knownGene)
  > txdb <- TxDb.Hsapiens.UCSC.hg19.knownGene</pre>
  > introns <- unlist(intronsByTranscript(txdb))</pre>
7. > system.time(names(which.max(xtabs(width ~ name,
                                       hesc.chromHmm))))
     user system elapsed
    0.092 0.000 0.092
  > ## or
  > names(which.max(sum(with(hesc.chromHmm,
                              splitAsList(width, name)))))
   +
   [1] "13_Heterochrom/lo"
  > ## or
  > df <- aggregate(hesc.chromHmm, ~ name, totalWidth=sum(width))</pre>
  > df$name[which.max(df$totalWidth)]
   [1] 13_Heterochrom/lo
  15 Levels: 10_Txn_Elongation 11_Weak_Txn 12_Repressed ... 9_Txn_Transition
```

# References

 Matthew T Maurano, Richard Humbert, Eric Rynes, Robert E Thurman, Eric Haugen, Hao Wang, Alex P Reynolds, Richard Sandstrom, Hongzhu Qu, Jennifer Brody, et al. Systematic localization of common disease-associated variation in regulatory dna. *Science*, 337(6099):1190–1195, 2012.