Hervé Pagès

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1 Introduction

In the context of an RNA-seq experiment, encoding the overlaps between the aligned reads and the transcripts can be used for detecting those overlaps that are "compatible" with the splicing of the transcript.

Various tools are provided in the *IRanges* and *GenomicRanges* packages for working with *overlap encodings*. In this vignette, we illustrate the use of these tools on the single-end and paired-end reads of an RNA-seq experiment.

2 Load reads from a BAM file

2.1 Load single-end reads from a BAM file

BAM file untreated1_chr4.bam (located in the *pasillaBamSubset* data package) contains single-end reads from the "Pasilla" experiment and aligned against the dm3 genome (see ?untreated1_chr4 in the *pasillaBamSubset* package for more information about those reads):

- > library(pasillaBamSubset)
- > untreated1_chr4()
- [1] "/home/biocbuild/bbs-2.13-bioc/R/library/pasillaBamSubset/extdata/untreated1_chr4.bam"

We use the readGAlignments function defined in the *GenomicRanges* package to load the reads into a *GAlignments* object. It's probably a good idea to get rid of the PCR or optical duplicates (flag bit 0x400 in the SAM format, see the SAM Spec ¹ for the details), as well as reads not passing quality controls (flag bit 0x200 in the SAM format). We do this by creating a *ScanBamParam* object that we pass to readGAlignments (see ?ScanBamParam in the *Rsamtools* package for the details). Note that we also use use.names=TRUE in order to load the *query names* (aka *query template names*, see QNAME field in the SAM Spec) from the BAM file (readGAlignments will use them to set the names of the returned object):

- > library(GenomicRanges)
- > library(Rsamtools)
- > flag0 <- scanBamFlag(isDuplicate=FALSE, isNotPassingQualityControls=FALSE)
- > param0 <- ScanBamParam(flag=flag0)</pre>

¹http://samtools.sourceforge.net/

> U1.GAL <- readGAlignments(untreated1_chr4(), use.names=TRUE, param=param0)
> head(U1.GAL)

GAlignments with 6 alignments and 0 metadata columns:

	seqnames	${\tt strand}$	cigar	qwidth	start	end	width	ngap
	<rle></rle>	<rle></rle>	<character></character>	<pre><integer></integer></pre>	<pre><integer></integer></pre>	<integer></integer>	<pre><integer></integer></pre>	<integer></integer>
SRR031729.3941844	chr4	_	75M	75	892	966	75	0
SRR031728.3674563	chr4	-	75M	75	919	993	75	0
SRR031729.8532600	chr4	+	75M	75	924	998	75	0
SRR031729.2779333	chr4	+	75M	75	936	1010	75	0
SRR031728.2826481	chr4	+	75M	75	949	1023	75	0
SRR031728.2919098	chr4	_	75M	75	967	1041	75	0
seqlengths:								
chr2L chr2H	R chr31	L chi	c3R chr4	\mathtt{chrM}	chrX o	chrYHet		

chr2L chr2R chr3L chr3R chr4 chrM chrX chrYHet 23011544 21146708 24543557 27905053 1351857 19517 22422827 347038

Because the aligner used to align those reads can report more than 1 alignment per *original query* (i.e. per read stored in the input file, typically a FASTQ file), we shouldn't expect the names of U1.GAL to be unique:

```
> U1.GAL_names_is_dup <- duplicated(names(U1.GAL))
> table(U1.GAL_names_is_dup)
```

```
U1.GAL_names_is_dup
FALSE TRUE
190770 13585
```

Storing the query names in a factor will be useful as we will see later in this document:

```
> U1.uqnames <- unique(names(U1.GAL))
> U1.GAL_qnames <- factor(names(U1.GAL), levels=U1.uqnames)</pre>
```

Note that we explicitly provide the levels of the factor to enforce their order. Otherwise factor() would put them in lexicographic order which is not advisable because it depends on the locale in use.

Another object that will be useful to keep near at hand is the mapping between each *query name* and its first occurence in U1.GAL_qnames:

```
> U1.GAL_dup2ung <- match(U1.GAL_qnames, U1.GAL_qnames)
```

Our reads can have up to 2 gaps (a gap corresponds to an N operation in the CIGAR):

> head(unique(cigar(U1.GAL)))

```
[1] "75M" "35M6727N40M" "22M6727N53M" "13M6727N62M" "26M292N49M" "62M21227N13M"
```

> table(ngap(U1.GAL))

```
0 1 2
184039 20169 147
```

Also, the following table indicates that indels were not allowed/supported during the alignment process (no I or D CIGAR operations):

> colSums(cigarOpTable(cigar(U1.GAL)))

M	I	D	N	S	H	P	=	X
15326625	0	0 216	82582	0	0	0	0	0

2.2 Load paired-end reads from a BAM file

BAM file untreated3_chr4.bam (located in the pasillaBamSubset data package) contains paired-end reads from the "Pasilla" experiment and aligned against the dm3 genome (see ?untreated3_chr4 in the pasillaBamSubset package for more information about those reads). We use the readGAlignmentPairs function to load them into a GAlignmentPairs object:

- > U3.galp <- readGAlignmentPairs(untreated3_chr4(), use.names=TRUE, param=param0)
- > head(U3.galp)

GAlignmentPairs with 6 alignment pairs and 0 metadata columns:

```
seqnames strand :
                                         ranges --
                     <Rle> <Rle> :
                                      <IRanges> --
                                                       <IRanges>
SRR031715.1138209
                      chr4
                                +: [169,
                                           205] -- [ 326, 362]
 SRR031714.756385
                      chr4
                                + : [943,
                                           979] -- [1086, 1122]
                                           980] -- [1119, 1155]
SRR031714.2355189
                      chr4
                                + : [944,
SRR031714.5054563
                                + : [946, 982] -- [ 986, 1022]
                      chr4
SRR031715.1722593
                                + : [966, 1002] -- [1108, 1144]
                      chr4
                                + : [966, 1002] -- [1114, 1150]
SRR031715.2202469
                      chr4
```

seqlengths:

chr2L chr2R chr3L chr3R chr4 chrM chrX chrYHet 23011544 21146708 24543557 27905053 1351857 19517 22422827 347038

The show method for *GAlignmentPairs* objects displays two ranges columns, one for the *first* alignment in the pair (the left column), and one for the *last* alignment in the pair (the right column). The strand column corresponds to the strand of the *first* alignment.

> head(first(U3.galp))

GAlignments with 6 alignments and 0 metadata columns:

_		_							
	\$	seqnames	strand	cigar	qwidth	start	end	width	ngap
		<rle></rle>	<rle></rle>	<character></character>	<pre><integer></integer></pre>	<pre><integer></integer></pre>	<pre><integer></integer></pre>	<pre><integer></integer></pre>	<integer></integer>
RR031715.113	38209	chr4	+	37M	37	169	205	37	0
SRR031714.75	6385	chr4	+	37M	37	943	979	37	0
RR031714.235	55189	chr4	+	37M	37	944	980	37	0
RR031714.505	54563	chr4	+	37M	37	946	982	37	0
RR031715.172	22593	chr4	+	37M	37	966	1002	37	0
RR031715.220	2469	chr4	+	37M	37	966	1002	37	0
eqlengths:									
chr2L	chr2R	chr3L	chi	c3R chr4	chrM	chrX o	chrYHet		
23011544 211	146708	24543557	279050	053 1351857	19517	22422827	347038		
	SRR031714.75 RR031714.235 RR031714.505 RR031715.172 RR031715.220 eqlengths: chr2L	RR031715.1138209 SRR031714.756385 RR031714.2355189 RR031714.5054563 RR031715.1722593 RR031715.2202469 eqlengths: chr2L chr2R	RR031715.1138209 chr4 SRR031714.756385 chr4 RR031714.2355189 chr4 RR031714.5054563 chr4 RR031715.1722593 chr4 RR031715.2202469 chr4 eqlengths: chr2L chr2R chr3L	RR031715.1138209	<pre></pre>	<pre></pre>	<pre></pre>	<pre></pre>	<pre></pre>

> head(last(U3.galp))

GAlignments with 6 alignments and 0 metadata columns:

	seqnames	${\tt strand}$	cigar	qwidth	start	end	width	ngap
	<rle></rle>	<rle></rle>	<character></character>	<pre><integer></integer></pre>	<pre><integer></integer></pre>	<pre><integer></integer></pre>	<integer></integer>	<integer></integer>
SRR031715.1138209	chr4	_	37M	37	326	362	37	0
SRR031714.756385	chr4	_	37M	37	1086	1122	37	0
SRR031714.2355189	chr4	_	37M	37	1119	1155	37	0
SRR031714.5054563	chr4	_	37M	37	986	1022	37	0
SRR031715.1722593	chr4	_	37M	37	1108	1144	37	0
SRR031715.2202469	chr4	_	37M	37	1114	1150	37	0

```
seqlengths:
```

```
chr2L chr2R chr3L chr3R chr4 chrM chrX chrYHet
23011544 21146708 24543557 27905053 1351857 19517 22422827 347038
```

According to the SAM format specifications, the aligner is expected to mark each alignment pair as *proper* or not (flag bit 0x2 in the SAM format). The SAM Spec only says that a pair is *proper* if the *first* and *last* alignments in the pair are "properly aligned according to the aligner". So the exact criteria used for setting this flag is left to the aligner.

We use isProperPair to extract this flag from the GAlignmentPairs object:

```
> table(isProperPair(U3.galp))
```

```
FALSE TRUE 29518 45828
```

Even though we could do *overlap encodings* with the full object, we keep only the *proper* pairs for our downstream analysis:

```
> U3.GALP <- U3.galp[isProperPair(U3.galp)]</pre>
```

Because the aligner used to align those reads can report more than 1 alignment per *original query template* (i.e. per pair of sequences stored in the input files, typically 1 FASTQ file for the *first* ends and 1 FASTQ file for the *last* ends), we shouldn't expect the names of U3.GALP to be unique:

```
> U3.GALP_names_is_dup <- duplicated(names(U3.GALP))
> table(U3.GALP_names_is_dup)

U3.GALP_names_is_dup
FALSE TRUE
43659 2169
```

Storing the *query template names* in a factor will be useful:

```
> U3.ugnames <- unique(names(U3.GALP))</pre>
```

```
> U3.GALP_qnames <- factor(names(U3.GALP), levels=U3.uqnames)
```

as well as having the mapping between each query template name and its first occurence in U3.GALP_qnames:

> U3.GALP_dup2unq <- match(U3.GALP_qnames, U3.GALP_qnames)

Our reads can have up to 1 gap per end:

```
> head(unique(cigar(first(U3.GALP))))
```

```
[1] "37M" "6M58N31M" "25M56N12M" "19M62N18M" "29M222N8M" "9M222N28M"
```

> head(unique(cigar(last(U3.GALP))))

```
[1] "37M" "12M58N25M" "19M58N18M" "27M2339N10M" "29M2339N8M" "9M222N28M"
```

> table(ngap(first(U3.GALP)), ngap(last(U3.GALP)))

```
0 1
0 44510 596
1 637 85
```

Like for our single-end reads, the following tables indicate that indels were not allowed/supported during the alignment process:

> colSums(cigarOpTable(cigar(first(U3.GALP))))

M	I	D	N	S	H	P	=	Х			
1695636	0	0	673919	0	0	0	0	0			
<pre>> colSums(cigarOpTable(cigar(last(U3.GALP))))</pre>											
M	I	D	N	S	H	P	=	Х			
1695636	0	0	630395	0	0	0	0	0			

3 Validate the alignments produced by the aligner

In this section we show how to validate the alignments produced by the aligner by comparing the *original query sequences* (aka "true" or "real" query sequences, or query sequences **before** alignment) with the *reference query sequences* (i.e. the query sequences **after** alignment).

Note that even though this step is not strictly required for computing the *overlap encodings*, some of the concepts and string-based computations described in this section are slightly related to some ideas introduced later in this document.

3.1 Validate the single-end alignments

3.1.1 Load the original query sequences

To load the *original query sequences*, we reload the BAM file but now we explicitly request the SEQ field by using what="seq" in our call to ScanBamParam. To further validate the alignments produced by the aligner, we also need to load the NM tag which is a predefined tag described in the SAM Spec as the "Edit distance to the reference, including ambiguous bases but excluding clipping" (note that tags are optional fields i.e. not all BAM files have them):

```
> param1 <- ScanBamParam(flag=flag0, what="seq", tag="NM")
> U1.GAL <- readGAlignments(untreated1_chr4(), use.names=TRUE, param=param1)
> U1.GAL_qseq <- mcols(U1.GAL)$seq
> names(U1.GAL_qseq) <- names(U1.GAL)</pre>
> head(U1.GAL_qseq)
  A DNAStringSet instance of length 6
    width seq
                                                                                 names
[1]
       75 CTGTGGTGACCACACACACAGAATGGTTCGGGC...GGGTTCCCTGCCCCTTTCCTGGCTAGGTTGTCC SRR031729.3941844
      75 TCGGGCCCAATTAGAGGGTTCCCTGCCCCTTTC...TTGTCCGCTAGCTCATTTCCTGGGCTGTTGTTG SRR031728.3674563
[2]
      75 CCCAATTAGAGGATTCTCTGCCCCTTTCCTGGC...CGGTAGCTCATTTCCCGGGATGTTGTTGTCC SRR031729.8532600
[4]
      75 GTTCTCTGCCCCTTTCCTGGCTAGGTTGTCCGC...TCCCGAGATGTTGTTGTTGTCCCGGGACCCACCT SRR031729.2779333
[5]
       75 TTCCTGGCTAGGTTGTCCGCTAGCTCATTTCCC...TTGTGTCCCGGGACACCCTTATTGTGAGTTTG SRR031728.2826481
[6]
      75 GCTAGCTCATTTCCCGGGAGGTTGTTGTCCC...CTTATTGTGAGTTTGTTGACAGCTCCAAGTTTG SRR031728.2919098
```

Because the BAM format imposes that the read sequence is "reverse complemented" when the read is aligned to the minus strand, we "reverse complement" it again to get the *original query sequences*:

```
> U1.GAL_oqseq <- U1.GAL_qseq
> U1.GAL_is_on_minus <- as.logical(strand(U1.GAL) == "-")</pre>
> U1.GAL_oqseq[U1.GAL_is_on_minus] <- reverseComplement(U1.GAL_oqseq[U1.GAL_is_on_minus])
> head(U1.GAL_oqseq)
  A DNAStringSet instance of length 6
    width seq
                                                                                 names
[1]
       75 GGACAACCTAGCCAGGAAAGGGGCAGGGAACCC...GCCCGAACCATTCTGTGGTGTTGGTCACCACAG SRR031729.3941844
      75 CAACAACAGCCCAGGAAATGAGCTAGCGGACAA...GAAAGGGGCAGGGAACCCTCTAATTGGGCCCGA SRR031728.3674563
[2]
      75 CCCAATTAGAGGATTCTCTGCCCCTTTCCTGGC...CGGTAGCTCATTTCCCGGGATGTTGTTGTTCCC SRR031729.8532600
[3]
      75 GTTCTCTGCCCCTTTCCTGGCTAGGTTGTCCGC...TCCCGAGATGTTGTTGTTGTCCCGGGACCCACCT SRR031729.2779333
[4]
[5]
       75 TTCCTGGCTAGGTTGTCCGCTAGCTCATTTCCC...TTGTGTCCCGGGACACCCTTATTGTGAGTTTG SRR031728.2826481
      75 CAAACTTGGAGCTGTCAACAAACTCACAATAAG...GGGACACAACCACCTCCCGGGAAATGAGCTAGC SRR031728.2919098
[6]
```

Note that sequences with the same *query name* correspond to the same *original query* and therefore must be the same. Let's do a quick sanity check:

```
> stopifnot(all(U1.GAL_oqseq == U1.GAL_oqseq[U1.GAL_dup2unq]))
```

Finally, let's reduce U1.GAL_oqseq to one original query sequence per unique query name:

```
> U1.oqseq <- U1.GAL_oqseq[!U1.GAL_names_is_dup]
```

If we had access to the FASTQ file used as input to the aligner, U1.oqseq would be the subset of this file made of the sequences with at least 1 alignment reported in BAM file untreated1_chr4.bam.

3.1.2 Compute the reference query sequences

> library(BSgenome.Dmelanogaster.UCSC.dm3)

The *reference query sequences* can easily be computed by extracting the nucleotides mapped to each read from the reference genome. This of course requires that we have access to the reference genome used by the aligner. In Bioconductor, the full genome sequence for the dm3 assembly is stored in the *BSgenome.Dmelanogaster.UCSC.dm3* data package ²:

```
> Dmelanogaster
Fly genome
| organism: Drosophila melanogaster (Fly)
| provider: UCSC
| provider version: dm3
| release date: Apr. 2006
| release name: BDGP Release 5
| single sequences (see '?seqnames'):
    chr2L
               chr2R
                           chr3L
                                       chr3R
                                                   chr4
                                                              chrX
                                                                          chrU
                                                                                     chrM
               chr2RHet
    chr2LHet
                           chr3LHet
                                       chr3RHet
                                                  chrXHet
                                                              chrYHet
                                                                          chrUextra
| multiple sequences (see '?mseqnames'):
    upstream1000 upstream2000 upstream5000
| (use the '$' or '[[' operator to access a given sequence)
   Let's start by converting our GAlignments object U1.GAL into a GRangesList object:
> U1.grl <- grglist(U1.GAL, order.as.in.query=TRUE)
   To extract the portions of the reference genome corresponding to the ranges in U1.grl, we can use the extract-
TranscriptsFromGenome function defined in the GenomicFeatures package:
> library(GenomicFeatures)
> U1.GAL_rqseq <- extractTranscriptsFromGenome(Dmelanogaster, U1.grl)
> head(U1.GAL_rqseq)
  A DNAStringSet instance of length 6
    width seq
                                                                                    names
       75 GGACAACCTAGCCAGGAAAGGGGCAGAGAACCC...GCCCGAACCATTCTGTGGTGTTGGTCACCACAG SRR031729.3941844
[1]
       75 CAACAACATCCCGGGAAATGAGCTAGCGGACAA...GAAAGGGGCAGAGAACCCTCTAATTGGGCCCGA SRR031728.3674563
```

3.1.3 Compare the original query sequences with the reference query sequences

[3]

[5]

[6]

We can use the neditAt function defined in the *Biostrings* package to compute the edit distance between 2 strings. Because the aligned reads have no indels, we should only see mismatches (typically a small number) during that comparison so we don't need to call neditAt with with.indels=TRUE. And because calling neditAt in a loop is inefficient, we only do this comparison for the first 500 sequences in U1.GAL_oqseq:

75 CCCAATTAGAGGGTTCTCTGCCCCTTTCCTGGC...CGCTAGCTCATTTCCCGGGATGTTGTTGTTCCC SRR031729.8532600

75 GTTCTCTGCCCCTTTCCTGGCTAGGTTGTCCGC...TCCCGGGATGTTGTTGTGTCCCGGGACCCACCT SRR031729.2779333 75 TTCCTGGCTAGGTTGTCCGCTAGCTCATTTCCC...TTGTGTCCCGGGACCCACCTTATTGTGAGTTTG SRR031728.2826481

75 CAAACTTGGAGCTGTCAACAACTCACAATAAG...GGGACACAACAACATCCCGGGAAATGAGCTAGC SRR031728.2919098

```
> U1.GAL_nedit500 <- sapply(1:500, function(i) neditAt(U1.GAL_oqseq[[i]], U1.GAL_rqseq[[i]]))
> table(U1.GAL_nedit500)
```

²See http://bioconductor.org/packages/release/data/annotation/ for the full list of annotation packages available in the current release of Bioconductor.

```
U1.GAL_nedit500

0 1 2 3 4 5

273 101 97 22 6 1
```

Yes, the first 500 sequences in U1.GAL_oqseq are "close" to the first 500 sequences in U1.GAL_rqseq.

Now let's compare the edit distance reported by neditAt with the edit distance reported by the aligner (NM tag). Because the latter excludes the N CIGAR operations, it should actually be the same as the former. We confirm this for the 500 edit distances computed in U1.GAL_nedit500:

```
> U1.GAL_NM <- mcols(U1.GAL)$NM
> stopifnot(all(U1.GAL_NM[1:500] == U1.GAL_nedit500))
```

Note that the maximum observed number of mismatches tells us how many mismatches per read were tolerated by the aligner:

```
> table(U1.GAL_NM)
U1.GAL_NM
     0      1      2      3      4      5      6
135911     44033     15684     5172      2233      948      374
```

3.2 Validate the paired-end alignments

3.2.1 Load the original query sequences

width seq

37 GTCTCCAGCAGAGCAGATGGAGCAACGGCCTATAGAG

To load the *original query sequences*, we reload the BAM file and request the SEQ field (and also the NM tag). Since we've removed the improper pairs from our current U3.GALP object, we need to do this again but now we do it at load time which is equivalent to doing it afterward (i.e. not only do we have the guarantee to end up with the same elements in U3.GALP, but also to have them in the same order):

```
> flag2 <- scanBamFlag(isDuplicate=FALSE,
                       isNotPassingQualityControls=FALSE,
                       isProperPair=TRUE)
> param2 <- ScanBamParam(flag=flag2, what="seq", tag="NM")
> U3.GALP <- readGAlignmentPairs(untreated3_chr4(), use.names=TRUE, param=param2)
  Let's extract the first and last sequences from U3.GALP:
> U3.GALP_gseq1 <- mcols(first(U3.GALP))$seq</pre>
> U3.GALP_qseq2 <- mcols(last(U3.GALP))$seq</pre>
> names(U3.GALP_qseq1) <- names(U3.GALP_qseq2) <- names(U3.GALP)
> head(U3.GALP_qseq1)
  A DNAStringSet instance of length 6
    width seq
                                                                                   names
       37 CCGTTTCTGAAGGAGATGGCTCATGGAGTACCTGCCT
[1]
                                                                                   SRR031715.1138209
[2]
       37 GCCCCTTTCCTGGCTAGGTTGTCCGCTAGCTCATTTC
                                                                                   SRR031714.756385
[3]
       37 CCTTTCCTGGCTAGGTTGTCCGCTAGCTCATTTCCCG
                                                                                   SRR031714.5054563
       37 CGCTAGCTCATTTCCCGGGCTGTTGTTGTCCCGGG
[4]
                                                                                   SRR031715.1722593
[5]
       37 CGCTAGCTCATTTCCCGAGATGTTGTTGTCCCGGG
                                                                                   SRR031715.2202469
       37 GCTTTGCTGAGCGCCTTTATGGCTGCTTGACTATCAG
                                                                                   SRR031714.3544437
> head(U3.GALP_qseq2)
  A DNAStringSet instance of length 6
```

names

SRR031715.1138209

[2]	37 AGCTTTGCTGAGCGCCTTTATGGCTGCTTGACTATCA	SRR031714.756385
[3]	37 TGTTGTTGTCCCGGGACCCACCTTATTGTGAGTTT	SRR031714.5054563
[4]	37 GCTGCTTGACTATCAGACAGTATAGCAATGTCCTTGC	SRR031715.1722593
[5]	37 TGACTATCAGACAGTATAGCAATGTCCTTGCCATGAT	SRR031715.2202469
[6]	37 GTCCGCTAGCTCATTTCCCGGGATGTTTTTGTGTCCC	SRR031714.3544437

To obtain the original query sequences we "reverse complement" the sequences that are aligned to the minus strand:

```
> U3.GALP_oqseq1 <- U3.GALP_qseq1
> U3.GALP_first_is_on_minus <- as.logical(strand(first(U3.GALP)) == "-")
> U3.GALP_oqseq1[U3.GALP_first_is_on_minus] <- reverseComplement(U3.GALP_oqseq1[U3.GALP_first_is_on_minus])
> U3.GALP_oqseq2 <- U3.GALP_qseq2
> U3.GALP_last_is_on_minus <- as.logical(strand(last(U3.GALP)) == "-")
> U3.GALP_oqseq2[U3.GALP_last_is_on_minus] <- reverseComplement(U3.GALP_oqseq2[U3.GALP_last_is_on_minus])</pre>
```

Note that sequence pairs with the same *query template name* correspond to the same *original query pairs* and therefore should be the same. Let's do a quick sanity check:

```
> stopifnot(all(U3.GALP_oqseq1 == U3.GALP_oqseq1[U3.GALP_dup2unq]))
> stopifnot(all(U3.GALP_oqseq2 == U3.GALP_oqseq2[U3.GALP_dup2unq]))
```

Finally, let's reduce U3.GALP_oqseq1 and U3.GALP_oqseq2 to one *original query sequence* per unique *query template* name:

```
> U3.oqseq1 <- U3.GALP_oqseq1[!U3.GALP_names_is_dup]
> U3.oqseq2 <- U3.GALP_oqseq2[!U3.GALP_names_is_dup]</pre>
```

If we had access to the 2 FASTQ files used as input to the aligner, U3.oqseq1 and U3.oqseq2 would be the subsets of those files made of the sequence pairs with at least 1 alignment pair reported in BAM file untreated3_chr4.bam.

3.2.2 Compute the reference query sequences

Because our reads are paired-end, we extract separately the ranges corresponding to their *first* ends (aka *first* segments in BAM jargon) and those corresponding to their *last* ends (aka *last* segments in BAM jargon):

```
> U3.grl_first <- grglist(first(U3.GALP), order.as.in.query=TRUE)
> U3.grl_last <- grglist(last(U3.GALP, invert.strand=TRUE), order.as.in.query=TRUE)</pre>
```

Then we extract the portions of the reference genome corresponding to the ranges in *GRangesList* objects U3.grl_first and U3.grl_last:

```
> U3.GALP_rqseq1 <- extractTranscriptsFromGenome(Dmelanogaster, U3.grl_first)
> U3.GALP_rqseq2 <- extractTranscriptsFromGenome(Dmelanogaster, U3.grl_last)</pre>
```

3.2.3 Compare the original query sequences with the reference query sequences

Because the aligned reads have no indels, we should only see mismatches (typically a small number) during that comparison so we don't need to call neditAt with with.indels=TRUE. Let's do this comparison for the first 500 sequences in U3.GALP_oqseq1 and in reverseComplement(U3.GALP_oqseq2):

```
> U3.GALP_first_nedit500 <- sapply(1:500, function(i)
+    neditAt(U3.GALP_oqseq1[[i]], U3.GALP_rqseq1[[i]])
+ )
> table(U3.GALP_first_nedit500)

U3.GALP_first_nedit500
0 1 2
337 135 28
```

Yes, the first 500 sequences in U3.GALP_oqseq1 and in reverseComplement(U3.GALP_oqseq2) are "close" to the first 500 sequences in U3.GALP_rqseq1 and in U3.GALP_rqseq2, respectively.

Now let's compare the edit distance reported by neditAt with the edit distance reported by the aligner (NM tag). Because the latter excludes the N CIGAR operations, it should actually be the same as the former. We confirm this for the 500 edit distances computed in U3.GALP_first_nedit500 and U3.GALP_last_nedit500:

```
> U3.GALP_first_NM <- mcols(first(U3.GALP))$NM
> stopifnot(all(U3.GALP_first_NM[1:500] == U3.GALP_first_nedit500))
> U3.GALP_last_NM <- mcols(last(U3.GALP))$NM
> stopifnot(all(U3.GALP_last_NM[1:500] == U3.GALP_last_nedit500))
```

Note that the following table tells us how many mismatches per read were tolerated by the aligner:

```
> table(U3.GALP_first_NM, U3.GALP_last_NM)
```

Up to 2 mismatches per end.

3.3 Conclusion

In addition to validate the alignments produced by the aligner, the validation described in this section is also an efficient and accurate way to make sure that the reference genome we've picked up is the same as the reference genome used by the aligner, at least for the regions covered by the reads.

In other words, if it's known that the 2 reference genomes are different, then this validation could still be performed, and, if successful, would indicate that the 2 genomes are probably substitutable for most analysis happening downstream of the BAM file.

4 Find all the overlaps between the reads and transcripts

4.1 Load the transcripts from a TranscriptDb object

In order to compute overlaps between reads and transcripts, we need access to the genomic positions of a set of known transcripts and their exons. It is essential that the reference genome of this set of transcripts and exons be **exactly** the same as the reference genome used to align the reads.

We could use the makeTranscriptDbFromUCSC function defined in the *GenomicFeatures* package to make a *TranscriptDb* object containing the dm3 transcripts and their exons retrieved from the UCSC Genome Browser³. The Bioconductor project however provides a few annotation packages containing *TranscriptDb* objects for the most commonly studied organisms (those data packages are sometimes called the *TxDb* packages). One of them is the *TxDb.Dmelanogaster.-UCSC.dm3.ensGene* package. It contains a *TranscriptDb* object that was made by pointing the makeTranscriptDbFromUCSC function to the dm3 genome and *Ensembl Genes* track ⁴. We can use it here:

³http://genome.ucsc.edu/cgi-bin/hgGateway

⁴See http://genome.ucsc.edu/cgi-bin/hgTrackUi?hgsid=276880911&g=ensGene for a description of this track.

```
> library(TxDb.Dmelanogaster.UCSC.dm3.ensGene)
> TxDb.Dmelanogaster.UCSC.dm3.ensGene
TranscriptDb object:
| Db type: TranscriptDb
| Supporting package: GenomicFeatures
| Data source: UCSC
| Genome: dm3
| Organism: Drosophila melanogaster
| UCSC Table: ensGene
Resource URL: http://genome.ucsc.edu/
| Type of Gene ID: Ensembl gene ID
| Full dataset: yes
| miRBase build ID: NA
| transcript_nrow: 29173
| exon_nrow: 76920
| cds_nrow: 62135
| Db created by: GenomicFeatures package from Bioconductor
| Creation time: 2013-09-19 00:24:08 -0700 (Thu, 19 Sep 2013)
| GenomicFeatures version at creation time: 1.13.40
RSQLite version at creation time: 0.11.4
| DBSCHEMAVERSION: 1.0
> txdb <- TxDb.Dmelanogaster.UCSC.dm3.ensGene
  We extract the exons grouped by transcript in a GRangesList object:
> exbytx <- exonsBy(txdb, by="tx", use.names=TRUE)
> length(exbytx) # nb of transcripts
[1] 29173
```

We check that all the exons in any given transcript belong to the same chromosome and strand. Knowing that our set of transcripts is free of this sort of trans-splicing events typically allows some significant simplifications during the downstream analysis ⁵. A quick and easy way to check this is to take advantage of the fact that seqnames and strand return *RleList* objects. So we can extract the number of Rle runs for each transcript and make sure it's always 1:

```
> table(elementLengths(runLength(seqnames(exbytx))))
    1
29173
> table(elementLengths(runLength(strand(exbytx))))
    1
29173
```

Therefore the strand of any given transcript is unambiguously defined and can be extracted with:

> exbytx_strand <- unlist(runValue(strand(exbytx)), use.names=FALSE)

We will also need the mapping between the transcripts and their gene. We start by using transcripts to extract this information from our *TranscriptDb* object txdb, and then we construct a named factor that represents the mapping:

```
> tx <- transcripts(txdb, columns=c("tx_name", "gene_id"))
> head(tx)
```

⁵Dealing with trans-splicing events is not covered in this document.

```
GRanges with 6 ranges and 2 metadata columns:
                       ranges strand |
      seqnames
                                                            gene_id
                                           tx_name
                    <IRanges> <Rle> | <character> <CharacterList>
         <Rle>
  [1]
         chr2L [ 7529, 9484]
                                  + | FBtr0300689
                                                       FBgn0031208
  [2]
         chr2L [ 7529, 9484]
                                   + | FBtr0300690
                                                        FBgn0031208
         chr2L [ 7529, 9484]
                                                        FBgn0031208
  [3]
                                  + | FBtr0330654
  [4]
         chr2L [21952, 24237]
                                                        FBgn0263584
                                 + | FBtr0309810
  [5]
         chr2L [66584, 71390]
                                 + | FBtr0306539
                                                        FBgn0067779
         chr2L [67043, 71081]
                                                        FBgn0067779
  [6]
                                   + | FBtr0306536
  seqlengths:
       chr2L
                 chr2R
                           chr3L
                                     chr3R
                                                 chr4 ... chr3RHet
                                                                      chrXHet chrYHet chrUextra
    23011544 21146708 24543557 27905053
                                             1351857 ...
                                                           2517507
                                                                      204112
                                                                                 347038 29004656
> df <- mcols(tx)</pre>
> exbytx2gene <- as.character(df$gene_id)</pre>
> exbytx2gene <- factor(exbytx2gene, levels=unique(exbytx2gene))
> names(exbytx2gene) <- df$tx_name</pre>
> exbytx2gene <- exbytx2gene[names(exbytx)]</pre>
> head(exbytx2gene)
FBtr0300689 FBtr0300690 FBtr0330654 FBtr0309810 FBtr0306539 FBtr0306536
FBgn0031208 FBgn0031208 FBgn0031208 FBgn0263584 FBgn0067779 FBgn0067779
15682 Levels: FBgn0031208 FBgn0263584 FBgn0067779 FBgn0031213 FBgn0031214 FBgn0031216 ... FBgn0085792
> nlevels(exbytx2gene) # nb of genes
[1] 15682
```

4.2 Single-end overlaps

4.2.1 Find the single-end overlaps

We are ready to compute the overlaps with the findOverlaps function. Note that the strand of the queries produced by the RNA-seq experiment is typically unknown so we use ignore.strand=TRUE:

```
> U1.0V00 <- findOverlaps(U1.GAL, exbytx, ignore.strand=TRUE)
```

U1.0V00 is a *Hits* object that contains 1 element per overlap. Its length gives the number of overlaps:

- > length(U1.0V00)
- [1] 563552

4.2.2 Tabulate the single-end overlaps

We will repeatedly use the 2 following little helper functions to "tabulate" the overlaps in a given *Hits* object (e.g. U1.0V00), i.e. to count the number of overlaps for each element in the query or for each element in the subject:

```
> nhitPerQuery <- function(x) tabulate(queryHits(x), nbins=queryLength(x))
> nhitPerSubject <- function(x) tabulate(subjectHits(x), nbins=subjectLength(x))</pre>
```

Number of transcripts for each alignment in U1.GAL:

```
> U1.GAL_ntx <- nhitPerQuery(U1.0V00)
```

- > mcols(U1.GAL)\$ntx <- U1.GAL_ntx
- > head(U1.GAL)

GAlignments with 6 alignments and 3 metadata col	Alignments wi	nments and 3 metada	ta columns:
--	---------------	---------------------	-------------

	seqnames	strand	cigar	qwidth	start	end	width	ngap	
	<rle></rle>	<rle></rle>	<character></character>	<pre><integer></integer></pre>	<pre><integer></integer></pre>	<pre><integer></integer></pre>	<integer></integer>	<integer></integer>	
SRR031729.3941844	chr4	-	75M	75	892	966	75	0	
SRR031728.3674563	chr4	_	75M	75	919	993	75	0	1
SRR031729.8532600	chr4	+	75M	75	924	998	75	0	
SRR031729.2779333	chr4	+	75M	75	936	1010	75	0	
SRR031728.2826481	chr4	+	75M	75	949	1023	75	0	1
SRR031728.2919098	chr4	-	75M	75	967	1041	75	0	1

seq

<DNAStringSet>

SRR031729.3941844 CTGTGGTGACCAACACCACAGAATGGTTCGGGCCCAATTAGAGGGTTCCCTGCCCCTTTCCTGGCTAGGTTGTCC
SRR031728.3674563 TCGGGCCCAATTAGAGGGTTCCCTGCCCCTTTCCTGGCTAGGTTGTCC
SRR031729.8532600 CCCAATTAGAGGATTCTCTGCCCCTTTCCTGGCTAGGTTGTCCGGTAGCTCATTTCCCGGGATGTTGTTGTCC
SRR031729.2779333 GTTCTCTGCCCCTTTCCTGGCTAGGTTGTCCCGTAGCTCATTTCCCGAGATGTTGTTGTCCCGGGACCCACCT
SRR031728.2826481 TTCCTGGCTAGGTTGTCCGCTAGCTCATTTCCCGGGACCCACCTTATTGTGAGTTTG
SRR031728.2919098 GCTAGCTCATTTCCCGGGAGGTTGTTCTGTGCCCCAAGTTTG

	141.1	IIUA
	<integer></integer>	<integer></integer>
SRR031729.3941844	1	0
SRR031728.3674563	3	0
SRR031729.8532600	2	0
SRR031729.2779333	1	0
SRR031728.2826481	2	0
SRR031728.2919098	1	0

seqlengths:

chr2L chr2R chr3L chr3R chr4 chrM chrX chrYHet 23011544 21146708 24543557 27905053 1351857 19517 22422827 347038

ntx

> table(U1.GAL_ntx)

U1.GAL_ntx

10 Λ 2 3 4 5 6 7 8 9 11 12 47076 9493 26146 82427 5291 14530 8158 610 1952 2099 492 4945

 $> mean(U1.GAL_ntx >= 1)$

[1] 0.7696362

76% of the alignments in U1.GAL have an overlap with at least 1 transcript in exbytx.

Note that countOverlaps can be used directly on U1.GAL and exbytx for computing U1.GAL_ntx:

- > U1.GAL_ntx_again <- countOverlaps(U1.GAL, exbytx, ignore.strand=TRUE)
- > stopifnot(identical(unname(U1.GAL_ntx_again), U1.GAL_ntx))

Because U1.GAL can (and actually does) contain more than 1 alignment per *original query* (aka read), we also count the number of transcripts for each read:

- > U1.OV10 <- remapHits(U1.OV00, query.map=U1.GAL_qnames)
- > U1.uqnames_ntx <- nhitPerQuery(U1.0V10)</pre>
- > names(U1.uqnames_ntx) <- U1.uqnames</pre>
- > table(U1.uqnames_ntx)

U1.uqnames_ntx

12 2 3 5 7 4 6 8 10 11 1 39503 9298 18394 82346 5278 14536 9208 610 2930 488 2099 4944 1136

```
> mean(U1.uqnames_ntx >= 1)
```

[1] 0.7929287

78.4% of the reads have an overlap with at least 1 transcript in exbytx. Number of reads for each transcript:

- > U1.exbytx_nOV10 <- nhitPerSubject(U1.OV10)</pre>
- > names(U1.exbytx_nOV10) <- names(exbytx)</pre>
- $> mean(U1.exbytx_nOV10 >= 50)$

[1] 0.009015185

Only 0.869% of the transcripts in exbytx have an overlap with at least 50 reads. Top 10 transcripts:

> head(sort(U1.exbytx_nOV10, decreasing=TRUE), n=10)

FBtr0308296 FBtr0089175 FBtr0089176 FBtr0112904 FBtr0289951 FBtr0089243 FBtr0333672 FBtr0089186
40654 40529 40529 11735 11661 11656 10087 10084
FBtr0089187 FBtr0089172
10084 6749

4.3 Paired-end overlaps

4.3.1 Find the paired-end overlaps

Like with our single-end overlaps, we call findOverlaps with ignore.strand=TRUE:

> U3.0V00 <- findOverlaps(U3.GALP, exbytx, ignore.strand=TRUE)

Like U1.0V00, U3.0V00 is a Hits object. Its length gives the number of paired-end overlaps:

- > length(U3.0V00)
- [1] 113827

4.3.2 Tabulate the paired-end overlaps

Number of transcripts for each alignment pair in U3.GALP:

- > U3.GALP_ntx <- nhitPerQuery(U3.0V00)
- > mcols(U3.GALP)\$ntx <- U3.GALP_ntx
- > head(U3.GALP)

GAlignmentPairs with 6 alignment pairs and 1 metadata column:

	seqnames	strand	:	1	ranges		1	ranges		ntx
	<rle></rle>	<rle></rle>	:	<ira< td=""><td>anges></td><td></td><td><ir:< td=""><td>anges></td><td> <integ< td=""><td>ger></td></integ<></td></ir:<></td></ira<>	anges>		<ir:< td=""><td>anges></td><td> <integ< td=""><td>ger></td></integ<></td></ir:<>	anges>	<integ< td=""><td>ger></td></integ<>	ger>
SRR031715.1138209	chr4	+	:	[169,	205]		[326,	362]		0
SRR031714.756385	chr4	+	:	[943,	979]		[1086,	1122]	1	0
SRR031714.5054563	chr4	+	:	[946,	982]		[986,	1022]	1	0
SRR031715.1722593	chr4	+	:	[966,	1002]		[1108,	1144]		0
SRR031715.2202469	chr4	+	:	[966,	1002]		[1114,	1150]		0
SRR031714.3544437	chr4	_	:	[1087,	1123]		[963,	999]		0
seqlengths:										
chr?I chr?I	2 chr31	chr	٠QI	2 61	nr/l	cl	n r M	chrY	chrVHot	

chr2L chr2R chr3L chr3R chr4 chrM chrX chrYHet 23011544 21146708 24543557 27905053 1351857 19517 22422827 347038

```
U3.GALP_ntx
                        3
                                     5
                                                         8
                                                                      10
                                                                            11
                                                                                   12
    0
                                                                           803
12950 2080 5854 17025 1078 3083 2021
                                                  70
                                                       338
                                                              370
                                                                                   97
                                                                      59
> mean(U3.GALP_ntx >= 1)
[1] 0.7174217
   71% of the alignment pairs in U3.GALP have an overlap with at least 1 transcript in exbytx.
   Note that countOverlaps can be used directly on U3.GALP and exbytx for computing U3.GALP_ntx:
> U3.GALP_ntx_again <- countOverlaps(U3.GALP, exbytx, ignore.strand=TRUE)
> stopifnot(identical(unname(U3.GALP_ntx_again), U3.GALP_ntx))
   Because U3.GALP can (and actually does) contain more than 1 alignment pair per original query template, we also
count the number of transcripts for each template:
> U3.0V10 <- remapHits(U3.0V00, query.map=U3.GALP_qnames)</pre>
> U3.uqnames_ntx <- nhitPerQuery(U3.0V10)</pre>
> names(U3.uqnames_ntx) <- U3.uqnames
> table(U3.uqnames_ntx)
U3.uqnames_ntx
                 2
                        3
                              4
                                     5
                                            6
                                                   7
                                                         8
                                                                      10
                                                                            11
                                                                                   12
11851 2061 4289 17025 1193
                                 3084 2271
                                                  70
                                                       486
                                                              370
                                                                      59
                                                                           803
                                                                                   97
> mean(U3.uqnames_ntx >= 1)
[1] 0.7285554
   72.3% of the templates have an overlap with at least 1 transcript in exbytx.
   Number of templates for each transcript:
> U3.exbytx_nOV10 <- nhitPerSubject(U3.OV10)</pre>
> names(U3.exbytx_nOV10) <- names(exbytx)</pre>
> mean(U3.exbytx_nOV10 >= 50)
[1] 0.00712988
   Only 0.756% of the transcripts in exbytx have an overlap with at least 50 templates.
   Top 10 transcripts:
> head(sort(U3.exbytx_nOV10, decreasing=TRUE), n=10)
FBtr0308296 FBtr0089175 FBtr0089176 FBtr0112904 FBtr0089243 FBtr0289951 FBtr0333672 FBtr0089186
                     7573
                                  7572
                                               2750
                                                             2732
                                                                          2732
                                                                                       2260
                                                                                                     2260
       7574
FBtr0089187 FBtr0310542
       2260
                     1698
```

5 Encode the overlaps between the reads and transcripts

5.1 Single-end encodings

> table(U3.GALP_ntx)

The *overlap encodings* are strand sensitive so we will compute them twice, once for the "original alignments" (i.e. the alignments of the *original queries*), and once again for the "flipped alignments" (i.e. the alignments of the "flipped *original queries*"). We extract the ranges of the "original" and "flipped" alignments in 2 *GRangesList* objects with:

```
> U1.grlf <- flipQuery(U1.grl) # flipped
and encode their overlaps with the transcripts:
> U1.ovencA <- encodeOverlaps(U1.grl, exbytx, hits=U1.0V00)
> U1.ovencB <- encodeOverlaps(U1.grlf, exbytx, hits=U1.0V00)</pre>
```

U1.ovencA and U1.ovencB are 2 OverlapsEncodings objects of the same length as Hits object U1.0V00. For each hit in U1.0V00, we have 2 corresponding encodings, one in U1.ovencA and one in U1.ovencB, but only one of them encodes a hit between alignment ranges and exon ranges that are on the same strand. We use the selectEncoding-WithCompatibleStrand function to merge them into a single OverlapsEncodings of the same length. For each hit in U1.0V00, this selects the encoding corresponding to alignment ranges and exon ranges with compatible strand:

 ${\tt OverlapEncodings} \ {\tt of} \ {\tt length} \ 563552$

	${\tt Loffset}$	${\tt Roffset}$	${\tt encoding}$	${\tt flippedQuery}$
[1]	0	3	1:i:	TRUE
[2]	4	0	1:k:	FALSE
[3]	4	0	1:k:	TRUE
[4]	4	0	1:k:	TRUE
[5]	4	0	1:k:	TRUE
[6]	4	0	1:k:	TRUE
[7]	4	0	1:k:	TRUE
[8]	4	0	1:i:	TRUE
[9]	4	0	1:i:	TRUE
[563544]	23	0	1:i:	FALSE
[563545]	24	0	1:i:	FALSE
[563546]	24	0	1:i:	FALSE
[563547]	23	0	1:i:	FALSE
[563548]	22	0	1:i:	TRUE
[563549]	23	0	1:i:	TRUE
[563550]	24	0	1:i:	TRUE
[563551]	24	0	1:i:	TRUE
[563552]	23	0	1:i:	TRUE

As a convenience, the 2 above calls to encodeOverlaps + merging step can be replaced by a single call to encodeOverlaps on U1.grl (or U1.grlf) with flip.query.if.wrong.strand=TRUE:

```
> U1.ovenc_again <- encodeOverlaps(U1.grl, exbytx, hits=U1.0V00, flip.query.if.wrong.strand=TRUE)
> stopifnot(identical(U1.ovenc_again, U1.ovenc))
```

Unique encodings in U1.ovenc:

```
> U1.unique_encodings <- levels(U1.ovenc)</pre>
```

[1] 120

> head(U1.unique_encodings)

```
[1] "1:c:" "1:e:" "1:f:" "1:h:" "1:i:" "1:j:"
```

> length(U1.unique_encodings)

Encodings are sort of cryptic but utilities are provided to extract specific meaning from them. Use of these utilities is covered later in this document.

5.2 Paired-end encodings

Let's encode the overlaps in U3.0V00:

- > U3.grl <- grglist(U3.GALP, order.as.in.query=TRUE)
 > U3.ovenc <- encodeOverlaps(U3.grl, exbytx, hits=U3.OVOO, flip.query.if.wrong.strand=TRUE)</pre>
- > U3.ovenc

OverlapEncodings of length 113827

```
Loffset Roffset encoding flippedQuery
[1]
                4
                        0 1--1:i--k:
                                               TRUE
[2]
                4
                        0 1--1:i--i:
                                               TRUE
[3]
                4
                        0 1--1:i--k:
                                              FALSE
[4]
                4
                        0 1--1:i--k:
                                              FALSE
[5]
                4
                                               TRUE
                        0 1--1:a--c:
[6]
                4
                        0 1--1:i--m:
                                              FALSE
               3
[7]
                        1 1--1:i--i:
                                              FALSE
[8]
               3
                        1 1--1:i--i:
                                              FALSE
[9]
               2
                        2 1--1:i--i:
                                               TRUE
             . . .
                                  . . .
                                                . . .
. . .
[113819]
              23
                        0 1--1:i--i:
                                              FALSE
                        0 1--1:i--i:
                                              FALSE
[113820]
              24
              24
                        0 1--1:i--i:
[113821]
                                              FALSE
[113822]
              23
                        0 1--1:i--i:
                                              FALSE
              22
                                               TRUE
[113823]
                        0 1--1:i--i:
              23
[113824]
                        0 1--1:i--i:
                                               TRUE
              24
                        0 1--1:i--i:
                                               TRUE
[113825]
[113826]
              24
                        0 1--1:i--i:
                                               TRUE
[113827]
              23
                        0 1--1:i--i:
                                               TRUE
```

Unique encodings in U3.ovenc:

```
> U3.unique_encodings <- levels(U3.ovenc)
```

- > length(U3.unique_encodings)
- [1] 123
- > head(U3.unique_encodings)

```
[1] "1--1:a--c:" "1--1:a--i:" "1--1:a--j:" "1--1:a--k:" "1--1:b--i:" "1--1:b--k:"
```

- > U3.ovenc_table <- table(encoding(U3.ovenc))</pre>
- > tail(sort(U3.ovenc_table))

```
1--1:i--m: 1--1:i--k: 1--1:c--i: 1--2:i--jm:a--af: 2--1:jm--m:af--i: 852 1485 1714 2480 2700 1--1:i--i: 100084
```

6 "Compatible" overlaps

We are interested in a particular type of overlap where the read overlaps the transcript in a "compatible" way, that is, in a way compatible with the splicing of the transcript. The isCompatibleWithSplicing function can be used on an OverlapEncodings object to detect this type of overlap. Note that isCompatibleWithSplicing can also be used on a character vector or factor.

6.1 "Compatible" single-end overlaps

6.1.1 "Compatible" single-end encodings

U1.ovenc contains 7 unique encodings "compatible" with the splicing of the transcript:

> sort(U1.ovenc_table[isCompatibleWithSplicing(U1.unique_encodings)])

```
2:jm:ag: 2:gm:af: 3:jmm:agm:aaf: 1:j: 1:f: 2:jm:af: 32 79 488 1538 1555 72929 1:i: 455176
```

Encodings "1:i:" (455176 occurences in U1.ovenc), "2:jm:af:" (72929 occurences in U1.ovenc), and "3:jmm:agm:aaf:" (488 occurences in U1.ovenc), correspond to the following overlaps:

```
• "1:i:"
```

```
- read (no gap):
                             00000000
   - transcript:
                         >>>>>>>>>
• "2:jm:af:"
   - read (1 gap):
                             00000---000
   - transcript:
                         >>>>>>>
                                     >>>>>>>
• "3:jmm:agm:aaf:"
   - read (2 gaps):
                               00---0
   - transcript:
                         >>>>>>
                                    >>>>
                                           >>>>>
```

For clarity, only the exons involved in the overlap are represented. The transcript can of course have more upstream and downstream exons, which is denoted by the ... on the left side (5' end) and right side (3' end) of each drawing. Note that the exons represented in the 2nd and 3rd drawings are consecutive and adjacent in the processed transcript.

Encodings "1:f:" and "1:j:" are variations of the situation described by encoding "1:i:". For "1:f:", the first aligned base of the read (or "flipped" read) is aligned with the first base of the exon. For "1:j:", the last aligned base of the read (or "flipped" read) is aligned with the last base of the exon:

```
• "1:f:"
      - read (no gap):
                              00000000
      - transcript:
                              >>>>>>>>>
   • "1:j:"
       - read (no gap):
                                    0000000
      - transcript:
                              >>>>>>>>>
                        . . .
> U1.0V00_is_comp <- isCompatibleWithSplicing(U1.ovenc)
> table(U1.0V00_is_comp) # 531797 "compatible" overlaps
U1.0V00_is_comp
 FALSE TRUE
 31755 531797
```

Finally, let's extract the "compatible" overlaps from U1.0V00:

> U1.compOV00 <- U1.OV00[U1.OV00_is_comp]

Note that high-level convenience wrapper findCompatibleOverlaps can be used for computing the "compatible" overlaps directly between a *GAlignments* object (containing reads) and a *GRangesList* object (containing transcripts):

- > U1.compOVOO_again <- findCompatibleOverlaps(U1.GAL, exbytx)
- > stopifnot(identical(U1.comp0V00_again, U1.comp0V00))

6.1.2 Tabulate the "compatible" single-end overlaps

Number of "compatible" transcripts for each alignment in U1.GAL:

- > U1.GAL_ncomptx <- nhitPerQuery(U1.compOV00)</pre>
- > mcols(U1.GAL)\$ncomptx <- U1.GAL_ncomptx
- > head(U1.GAL)

GAlignments with 6 alignments and 4 metadata columns:

	seqnames	strand	cigar	qwidth	start	end	width	ngap
	<rle></rle>	<rle></rle>	<character></character>	<pre><integer></integer></pre>	<integer></integer>	<pre><integer></integer></pre>	<pre><integer></integer></pre>	<integer> </integer>
SRR031729.3941844	chr4	-	75M	75	892	966	75	0
SRR031728.3674563	chr4	-	75M	75	919	993	75	0
SRR031729.8532600	chr4	+	75M	75	924	998	75	0
SRR031729.2779333	chr4	+	75M	75	936	1010	75	0
SRR031728.2826481	chr4	+	75M	75	949	1023	75	0
SRR031728.2919098	chr4	_	75M	75	967	1041	75	0

seq

<DNAStringSet>

SRR031729.3941844 CTGTGGTGACCAACACCACAGAATGGTTCGGGCCCAATTAGAGGGTTCCCTGCCCCTTTCCTGGCTAGGTTGTCC
SRR031728.3674563 TCGGGCCCAATTAGAGGGTTCCCTGCCCCTTTCCTGGCTAGGTTGTTC
SRR031729.8532600 CCCAATTAGAGGATTCTCTGCCCCTTTCCTGGCTAGGTTGTCCGGTAGCTCATTTCCCGGGATGTTGTTGTCC
SRR031729.2779333 GTTCTCTGCCCCTTTCCTGGCTAGGTTGTCCGCTAGCTCATTTCCCGAGATGTTGTTGTCCCGGGACCCACCT
SRR031728.2826481 TTCCTGGCTAGGTTGTCCGCTAGCTCATTTCCCGGGGCCCACCTTATTGTGAGTTTG
SRR031728.2919098 GCTAGCTCATTTCCCGGGAGGTTGTTCTCCCGGGACCCACCTTATTGTGACACCTCCAAGTTTG

	1411	HUX	ncompex
	<integer></integer>	<pre><integer></integer></pre>	<integer></integer>
SRR031729.3941844	1	0	0
SRR031728.3674563	3	0	0
SRR031729.8532600	2	0	0
SRR031729.2779333	1	0	0
SRR031728.2826481	2	0	0
SRR031728.2919098	1	0	0

seqlengths:

chr2L chr2R chr3L chr3R chr4 chrM chrX chrYHet 23011544 21146708 24543557 27905053 1351857 19517 22422827 347038

> table(U1.GAL_ncomptx)

U1.GAL_ncomptx

2 3 4 5 6 7 8 10 11 12 51101 9848 33697 72987 5034 14021 7516 581 1789 530 2015 4389 847

> mean(U1.GAL_ncomptx >= 1)

[1] 0.7499401

75% of the alignments in U1.GAL are "compatible" with at least 1 transcript in exbytx.

Note that high-level convenience wrapper countCompatibleOverlaps can be used directly on U1.GAL and exbytx for computing U1.GAL_ncomptx:

- > U1.GAL_ncomptx_again <- countCompatibleOverlaps(U1.GAL, exbytx)
- > stopifnot(identical(U1.GAL_ncomptx_again, U1.GAL_ncomptx))

Number of "compatible" transcripts for each read:

- > U1.compOV10 <- remapHits(U1.compOV00, query.map=U1.GAL_qnames)
- > U1.uqnames_ncomptx <- nhitPerQuery(U1.compOV10)</pre>
- > names(U1.uqnames_ncomptx) <- U1.uqnames</pre>
- > table(U1.uqnames_ncomptx)

U1.uqnames_ncomptx

```
0 1 2 3 4 5 6 7 8 9 10 11 12
42886 9711 26075 72989 5413 14044 8584 581 2706 2015 530 4389 847
```

> mean(U1.ugnames_ncomptx >= 1)

[1] 0.7751953

77.5% of the reads are "compatible" with at least 1 transcript in exbytx. Number of "compatible" reads for each transcript:

- > U1.exbytx_ncompOV10 <- nhitPerSubject(U1.compOV10)</pre>
- > names(U1.exbytx_ncompOV10) <- names(exbytx)</pre>
- > mean(U1.exbytx_ncompOV10 >= 50)

[1] 0.008706681

Only 0.87% of the transcripts in exbytx are "compatible" with at least 50 reads. Top 10 transcripts:

> head(sort(U1.exbytx_ncompOV10, decreasing=TRUE), n=10)

```
FBtr0308296 FBtr0089175 FBtr0089176 FBtr0089243 FBtr0289951 FBtr0112904 FBtr0089186 FBtr0089187 40309 40158 33490 11365 11332 11284 10018 9627 FBtr0333672 FBtr0089172 9568 6599
```

Note that this "top 10" is slightly different from the "top 10" we obtained earlier when we counted **all** the overlaps.

6.2 "Compatible" paired-end overlaps

6.2.1 "Compatible" paired-end encodings

U3.ovenc contains 13 unique paired-end encodings "compatible" with the splicing of the transcript:

> sort(U3.ovenc_table[isCompatibleWithSplicing(U3.unique_encodings)])

Paired-end encodings "1--1:i--i:" (100084 occurences in U3.ovenc), "2--1:jm--m:af--i:" (2700 occurences in U3.ovenc), "1--2:i--jm:a--af:" (2480 occurences in U3.ovenc), "1--1:i--m:a--i:" (287 occurences in U3.ovenc), and "2--2:jm--mm:af--jm:aa--af:" (153 occurences in U3.ovenc), correspond to the following paired-end overlaps: • "1--1:i--i:" - paired-end read (no gap on the first end, no gap on the last end): 0000 0000 - transcript: >>>>>>>> • "2--1:jm--m:af--i:" - paired-end read (1 gap on the first end, no gap on the last end): 000---0 0000 ... >>>>>> - transcript: • "1--2:i--jm:a--af:" - paired-end read (no gap on the first end, 1 gap on the last end): 0000 00---00 - transcript: ... >>>>>>> ... • "1--1:i--m:a--i:" - paired-end read (no gap on the first end, no gap on the last end): 0000 0000 - transcript: ... >>>>>>> • "2--2:jm--mm:af--jm:aa--af:" - paired-end read (1 gap on the first end, 1 gap on the last end): 000---0 - transcript: ... >>>>> >>>>> Note: switch use of "first" and "last" above if the read was "flipped". > U3.0V00_is_comp <- isCompatibleWithSplicing(U3.ovenc)</pre> > table(U3.0V00_is_comp) # 106835 "compatible" paired-end overlaps U3.0V00_is_comp FALSE TRUE 6992 106835 Finally, let's extract the "compatible" paired-end overlaps from U3.0V00: > U3.compOV00 <- U3.OV00[U3.OV00_is_comp] Note that, like with our single-end reads, high-level convenience wrapper findCompatibleOverlaps can be used for computing the "compatible" paired-end overlaps directly between a GAlignmentPairs object (containing paired-end reads) and a GRangesList object (containing transcripts): > U3.compOVOO_again <- findCompatibleOverlaps(U3.GALP, exbytx)

6.2.2 Tabulate the "compatible" paired-end overlaps

Number of "compatible" transcripts for each alignment pair in U3.GALP:

> stopifnot(identical(U3.compOVOO_again, U3.compOVOO))

```
> U3.GALP_ncomptx <- nhitPerQuery(U3.comp0V00)
> mcols(U3.GALP)$ncomptx <- U3.GALP_ncomptx
> head(U3.GALP)
```

GAlignmentPairs with 6 alignment pairs and 2 metadata columns:

	seqnames	strand	:]	ranges		ranges	ı	ntx	ncomptx
	<rle></rle>	<rle></rle>	:		<ira< td=""><td>anges></td><td> <if< td=""><td>langes></td><td>-</td><td><integer></integer></td><td><integer></integer></td></if<></td></ira<>	anges>	 <if< td=""><td>langes></td><td>-</td><td><integer></integer></td><td><integer></integer></td></if<>	langes>	-	<integer></integer>	<integer></integer>
SRR031715.1138209	chr4	+	:		169,	205]	 [326,	362]	-	0	0
SRR031714.756385	chr4	+	:		943,	979]	 [1086,	1122]	-	0	0
SRR031714.5054563	chr4	+	:		946,	982]	 [986,	1022]	-	0	0
SRR031715.1722593	chr4	+	:		966,	1002]	 [1108,	1144]	-	0	0
SRR031715.2202469	chr4	+	:	[966,	1002]	 [1114,	1150]		0	0
SRR031714.3544437	chr4	-	:	[:	1087,	1123]	 [963,	999]	-	0	0

seqlengths:

chr2L chr2R chr3L chr3R chr4 chrM chrX chrYHet 23011544 21146708 24543557 27905053 1351857 19517 22422827 347038

> table(U3.GALP_ncomptx)

U3.GALP_ncomptx

0 1 2 3 4 5 6 7 8 9 10 11 12 13898 2028 8091 14334 1099 2950 1865 84 296 330 88 699 66

> mean(U3.GALP_ncomptx >= 1)

[1] 0.6967356

69.7% of the alignment pairs in U3.GALP are "compatible" with at least 1 transcript in exbytx.

Note that high-level convenience wrapper countCompatibleOverlaps can be used directly on U3.GALP and exbytx for computing U3.GALP_ncomptx:

- > U3.GALP_ncomptx_again <- countCompatibleOverlaps(U3.GALP, exbytx)
- > stopifnot(identical(U3.GALP_ncomptx_again, U3.GALP_ncomptx))

Number of "compatible" transcripts for each template:

- > U3.compOV10 <- remapHits(U3.compOV00, query.map=U3.GALP_qnames)
- > U3.uqnames_ncomptx <- nhitPerQuery(U3.compOV10)</pre>
- > names(U3.uqnames_ncomptx) <- U3.uqnames</pre>
- > table(U3.uqnames_ncomptx)

U3.uqnames_ncomptx

0 1 2 3 4 5 6 7 8 9 10 11 12 12783 2026 6531 14334 1210 2950 2114 84 444 330 88 699 66

> mean(U3.uqnames_ncomptx >= 1)

[1] 0.7072081

70.7% of the templates are "compatible" with at least 1 transcript in exbytx. Number of "compatible" templates for each transcript:

- > U3.exbytx_ncompOV10 <- nhitPerSubject(U3.compOV10)</pre>
- > names(U3.exbytx_ncompOV10) <- names(exbytx)</pre>
- > mean(U3.exbytx_ncompOV10 >= 50)

[1] 0.007061324

Only 0.7% of the transcripts in exbytx are "compatible" with at least 50 templates. Top 10 transcripts:

```
> head(sort(U3.exbytx_ncompOV10, decreasing=TRUE), n=10)
```

```
FBtr0308296 FBtr0089175 FBtr0089176 FBtr0289951 FBtr0089243 FBtr0112904 FBtr0089187 FBtr0089186 7425 7419 5227 2686 2684 2640 2257 2250 FBtr0333672 FBtr0310542 2206 1649
```

Note that this "top 10" is slightly different from the "top 10" we obtained earlier when we counted **all** the paired-end overlaps.

7 Project the alignments on the transcriptome

7.1 Project the single-end alignments on the transcriptome

The extractQueryStartInTranscript function computes for each overlap the position of the *query start* in the transcript:

```
> U1.0V00_qstart <- extractQueryStartInTranscript(U1.grl, exbytx,
                                                      hits=U1.0V00, ovenc=U1.ovenc)
> head(subset(U1.0V00_qstart, U1.0V00_is_comp))
   \verb|startInTranscript| firstSpannedExonRank| startInFirstSpannedExon
1
                  100
                                           1
                 4229
                                           5
                                                                   137
8
9
                 4229
                                           5
                                                                   137
                                           5
10
                 4207
                                                                   115
11
                 4207
                                           5
                                                                   115
                 4187
12
                                                                    95
```

U1.0V00_qstart is a data frame with 1 row per overlap and 3 columns:

- 1. startInTranscript: the 1-based start position of the read with respect to the transcript. Position 1 always corresponds to the first base on the 5' end of the transcript sequence.
- 2. firstSpannedExonRank: the rank of the first exon spanned by the read, that is, the rank of the exon found at position startInTranscript in the transcript.
- 3. startInFirstSpannedExon: the 1-based start position of the read with respect to the first exon spanned by the read.

Having this information allows us for example to compare the read and transcript nucleotide sequences for each "compatible" overlap. If we use the *reference query sequence* instead of the *original query sequence* for this comparison, then it should match **exactly** the sequence found at the *query start* in the transcript.

Let's start by using the extractTranscriptsFromGenome to extract the transcript sequences (aka transcriptome) from the dm3 reference genome:

```
> txseq <- extractTranscriptsFromGenome(Dmelanogaster, exbytx)
```

For each "compatible" overlap, the read sequence in U1.GAL_rqseq must be an *exact* substring of the transcript sequence in exbytx_seq:

Because of this relationship between the *reference query sequence* and the transcript sequence of a "compatible" overlap, and because of the relationship between the *original query sequences* and the *reference query sequences*, then the edit distance reported in the NM tag is actually the edit distance between the *original query* and the transcript of a "compatible" overlap.

7.2 Project the paired-end alignments on the transcriptome

For a paired-end read, the query start is the start of its "left end".

```
> U3.0V00_Lqstart <- extractQueryStartInTranscript(U3.grl, exbytx,
                                                      hits=U3.0V00, ovenc=U3.ovenc)
> head(subset(U3.0V00_Lgstart, U3.0V00_is_comp))
   startInTranscript firstSpannedExonRank startInFirstSpannedExon
2
                4118
                                          5
7
                                          4
                 3940
                                                                  31
8
                 3940
                                          4
                                                                  31
9
                                          3
                3692
                                                                 320
10
                 3692
                                          3
                                                                 320
                                          3
                 3690
                                                                 318
11
```

Note that extractQueryStartInTranscript can be called with for.query.right.end=TRUE if we want this information for the "right ends" of the reads:

```
> U3.0V00_Rqstart <- extractQueryStartInTranscript(U3.grl, exbytx,
+
                                                     hits=U3.0V00, ovenc=U3.ovenc,
                                                     for.query.right.end=TRUE)
> head(subset(U3.0V00_Rqstart, U3.0V00_is_comp))
   startInTranscript firstSpannedExonRank startInFirstSpannedExon
2
                4267
                                          5
                                                                 175
7
                3948
                                          4
                                                                  39
8
                3948
                                          4
                                                                  39
                                          3
9
                3849
                                                                 477
10
                3849
                                          3
                                                                 477
```

Like with single-end reads, having this information allows us for example to compare the read and transcript nucleotide sequences for each "compatible" overlap. If we use the *reference query sequence* instead of the *original query sequence* for this comparison, then it should match **exactly** the sequences of the "left" and "right" ends of the read in the transcript. Let's assign the "left and right reference query sequences" to each overlap:

```
> U3.0V00_Lrqseq <- U3.GALP_rqseq1[queryHits(U3.0V00)]
> U3.0V00_Rrqseq <- U3.GALP_rqseq2[queryHits(U3.0V00)]</pre>
```

For the single-end reads, the sequence associated with a "flipped query" just needed to be "reverse complemented". For paired-end reads, we also need to swap the 2 sequences in the pair:

```
> flip_idx <- which(flippedQuery(U3.ovenc))
> tmp <- U3.0V00_Lrqseq[flip_idx]
> U3.0V00_Lrqseq[flip_idx] <- reverseComplement(U3.0V00_Rrqseq[flip_idx])
> U3.0V00_Rrqseq[flip_idx] <- reverseComplement(tmp)</pre>
```

Let's assign the transcript sequence to each overlap:

```
> U3.0V00_txseq <- txseq[subjectHits(U3.0V00)]</pre>
```

For each "compatible" overlap, we expect the "left and right reference query sequences" of the read to be *exact* substrings of the transcript sequence. Let's check the "left reference query sequences":

```
> stopifnot(all(
      U3.0V00_Lrqseq[U3.0V00_is_comp] ==
+
+
          narrow(U3.0V00_txseq[U3.0V00_is_comp],
+
                 start=U3.0V00_Lqstart$startInTranscript[U3.0V00_is_comp],
                 width=width(U3.0V00_Lrqseq)[U3.0V00_is_comp])
+ ))
   and the "right reference query sequences":
> stopifnot(all(
      U3.0V00_Rrqseq[U3.0V00_is_comp] ==
          narrow(U3.0V00_txseq[U3.0V00_is_comp],
+
                 start=U3.0V00_Rqstart$startInTranscript[U3.0V00_is_comp],
                 width=width(U3.0V00_Rrqseq)[U3.0V00_is_comp])
+ ))
```

8 Align the reads to the transcriptome

Aligning the reads to the reference genome is not the most efficient nor accurate way to count the number of "compatible" overlaps per *original query*. Supporting junction reads (i.e. reads that align with at least 1 gap) introduces a significant computational cost during the alignment process. Then, as we've seen in the previous sections, each alignment produced by the aligner needs to be broken into a set of ranges (based on its CIGAR) and those ranges compared to the ranges of the exons grouped by transcript.

A more straightforward and accurate approach is to align the reads directly to the transcriptome, and without allowing the typical gap that the aligner needs to introduce when aligning a junction read to the reference genome. With this approach, a "hit" between a read and a transcript is necessarily compatible with the splicing of the transcript. In case of a "hit", we'll say that the read and the transcript are "string-based compatible" (to differentiate from our previous notion of "compatible" overlaps that we will call "encoding-based compatible" from now on, unless the context is clear).

8.1 Align the single-end reads to the transcriptome

8.1.1 Find the "hits"

The single-end reads are in U1.oqseq, the transcriptome is in exbytx_seq.

Since indels were not allowed/supported during the alignment of the reads to the reference genome, we don't need to allow/support them either for aligning the reads to the transcriptome. Also since our goal is to find (and count) "compatible" overlaps between reads and transcripts, we don't need to keep track of the details of the alignments between the reads and the transcripts. Finally, since BAM file untreated1_chr4.bam is not the full output of the aligner but the subset obtained by keeping only the alignments located on chr4, we don't need to align U1.oqseq to the full transcriptome, but only to the subset of exbytx_seq made of the transcripts located on chr4.

With those simplifications in mind, we write the following function that we will use to find the "hits" between the reads and the transcriptome:

```
subject_hits <- rep.int(seq_len(length(x)), elementLengths(x))</pre>
          new("Hits", queryHits=query_hits, subjectHits=subject_hits,
                        queryLength=pattern_length, subjectLength=length(x))
      }
+
      .isHitInTranscriptBounds <- function(hits, qseq, txseq)</pre>
          sapply(seq_len(length(hits)),
               function(i) {
                   pattern <- qseq[[queryHits(hits)[i]]]</pre>
                   subject <- txseq[[subjectHits(hits)[i]]]</pre>
                   v <- matchPattern(pattern, subject,</pre>
                                       max.mismatch=max.mismatch, fixed=FALSE)
+
                   any(1L <= start(v) & end(v) <= length(subject))</pre>
               })
+
      }
+
      if (!is.null(which.txseq)) {
          txseq0 <- txseq
+
          txseq <- txseq[which.txseq]</pre>
      }
      names(qseq) <- NULL</pre>
      other <- alphabetFrequency(qseq, baseOnly=TRUE)[ , "other"]</pre>
+
      is_clean <- other == OL # "clean" means "no IUPAC ambiguity code"
      ## Find hits for "clean" original queries.
      qseq0 <- qseq[is_clean]</pre>
      pdict0 <- PDict(qseq0, max.mismatch=max.mismatch)</pre>
      m0 <- vwhichPDict(pdict0, txseq,</pre>
                          max.mismatch=max.mismatch, fixed="pattern")
      hits0 <- .asHits(m0, length(qseq0))</pre>
+
      hits0@queryLength <- length(qseq)</pre>
      hitsO@queryHits <- which(is_clean)[hitsO@queryHits]
      ## Find hits for non "clean" original queries.
      qseq1 <- qseq[!is_clean]</pre>
      m1 <- vwhichPDict(qseq1, txseq,</pre>
                          max.mismatch=max.mismatch, fixed=FALSE)
      hits1 <- .asHits(m1, length(qseq1))</pre>
      hits1@queryLength <- length(qseq)
      hits1@queryHits <- which(!is_clean)[hits1@queryHits]
      ## Combine the hits.
      query_hits <- c(queryHits(hits0), queryHits(hits1))</pre>
      subject_hits <- c(subjectHits(hits0), subjectHits(hits1))</pre>
+
      if (!is.null(which.txseq)) {
          ## Remap the hits.
          txseq <- txseq0
           subject_hits <- which.txseq[subject_hits]</pre>
          hits0@subjectLength <- length(txseq)</pre>
      }
```

```
## Order the hits.
+ oo <- IRanges:::orderIntegerPairs(query_hits, subject_hits)
+ hitsO@queryHits <- query_hits[oo]
+ hitsO@subjectHits <- subject_hits[oo]
+
+ if (max.mismatch != OL) {
+ ## Keep only "in bounds" hits.
+ is_in_bounds <- .isHitInTranscriptBounds(hitsO, qseq, txseq)
+ hitsO <- hitsO[is_in_bounds]
+ }
+ hitsO
+ }</pre>
```

Let's compute the index of the transcripts in exbytx_seq located on chr4 (findSequenceHits will restrict the search to those transcripts):

```
> chr4tx <- transcripts(txdb, vals=list(tx_chrom="chr4"))
> chr4txnames <- mcols(chr4tx)$tx_name
> which.txseq <- match(chr4txnames, names(txseq))</pre>
```

We know that the aligner tolerated up to 6 mismatches per read. The 3 following commands find the "hits" for each *original query*, then find the "hits" for each "flipped *original query*", and finally merge all the "hits" (note that the 3 commands take about 1 hour to complete on a modern laptop):

8.1.2 Tabulate the "hits"

Number of "string-based compatible" transcripts for each read:

```
> U1.ugnames_nsbcomptx <- nhitPerQuery(U1.sbcompHITS)</pre>
> names(U1.uqnames_nsbcomptx) <- U1.uqnames</pre>
> table(U1.uqnames_nsbcomptx)
U1.uqnames_nsbcomptx
                       3
                             4
                                   5
                                          6
                                                7
                                                      8
                                                                  10
                                                                        11
                                                                               12
40555 10080 25299 74609 5207 14265 8643
                                              610 3410 2056
                                                                 534 4588
                                                                              914
> mean(U1.uqnames_nsbcomptx >= 1)
[1] 0.7874142
```

77.7% of the reads are "string-based compatible" with at least 1 transcript in exbytx. Number of "string-based compatible" reads for each transcript:

```
> U1.exbytx_nsbcompHITS <- nhitPerSubject(U1.sbcompHITS)
> names(U1.exbytx_nsbcompHITS) <- names(exbytx)
> mean(U1.exbytx_nsbcompHITS >= 50)
```

[1] 0.008809516

Only 0.865% of the transcripts in exbytx are "string-based compatible" with at least 50 reads. Top 10 transcripts:

```
> head(sort(U1.exbytx_nsbcompHITS, decreasing=TRUE), n=10)
```

```
FBtr0308296 FBtr0089175 FBtr0089176 FBtr0089243 FBtr0289951 FBtr0112904 FBtr0089186 FBtr0333672 40548 40389 34275 11605 11579 11548 10059 9742 FBtr0089187 FBtr0089172 9666 6704
```

8.1.3 A closer look at the "hits"

```
[WORK IN PROGRESS, might be removed or replaced soon...]

Any "encoding-based compatible" overlap is of course "string-based compatible":
```

```
> stopifnot(length(setdiff(U1.compOV10, U1.sbcompHITS)) == 0)
```

but the reverse is not true:

> length(setdiff(U1.sbcompHITS, U1.compOV10))

[1] 13549

8.2 Align the paired-end reads to the transcriptome

[COMING SOON...]

9 "Almost compatible" overlaps

In many aspects, "compatible" overlaps can be seen as perfect. We are now insterested in a less perfect type of overlap where the read overlaps the transcript in a way that *would* be "compatible" if 1 or more exons were removed from the transcript. In that case we say that the overlap is "almost compatible" with the transcript. The isCompatibleWithSkippedExons function can be used on an *OverlapEncodings* object to detect this type of overlap. Note that isCompatibleWithSkippedExons can also be used on a character vector of factor.

9.1 "Almost compatible" single-end overlaps

9.1.1 "Almost compatible" single-end encodings

U1.ovenc contains 7 unique encodings "almost compatible" with the splicing of the transcript:

> sort(U1.ovenc_table[isCompatibleWithSkippedExons(U1.unique_encodings)])

```
      2:jm:am:am:am:am:af:
      2:jm:am:am:am:af:
      2:jm:am:am:am:af:

      1
      1
      4
      7

      3:jmm:agm:aam:aaf:
      3:jmm:agm:aam:aaf:
      2:jm:am:am:af:
      2:jm:am:am:af:

      9
      21
      144
      1015
```

Encodings "2:jm:am:af:" (1015 occurences in U1.ovenc), "2:jm:am:am:af:" (144 occurences in U1.ovenc), and "3:jmm:agm:aam:aaf:" (21 occurences in U1.ovenc), correspond to the following overlaps:

```
• "2:jm:am:af:"
```

```
- read (1 gap): 00000-----000
- transcript: ... >>>>> >>> ...
```

• "2:jm:am:am:af:"

```
- read (1 gap): 00000------000
- transcript: ... >>>>> >>> ...
```

• "3:jmm:agm:aam:aaf:"

```
- read (2 gaps):
                                  00---00
      - transcript:
                             >>>>>
                                      >>>>
                                                   >>>>>>
                       . . .
> U1.0V00_is_acomp <- isCompatibleWithSkippedExons(U1.ovenc)</pre>
> table(U1.0V00_is_acomp) # 1202 "almost compatible" overlaps
U1.0V00_is_acomp
FALSE
        TRUE
```

Finally, let's extract the "almost compatible" overlaps from U1.0V00:

> U1.acomp0V00 <- U1.0V00[U1.0V00_is_acomp]</pre>

9.1.2 Tabulate the "almost compatible" single-end overlaps

Number of "almost compatible" transcripts for each alignment in U1.GAL:

- > U1.GAL_nacomptx <- nhitPerQuery(U1.acomp0V00)</pre>
- > mcols(U1.GAL)\$nacomptx <- U1.GAL_nacomptx</pre>
- > head(U1.GAL)

562350

1202

GAlignments with 6 alignments and 5 metadata columns:

	seqnames	strand	cigar	qwidth	start	end	width	ngap	
	<rle></rle>	<rle></rle>	<character></character>	<integer></integer>	<integer></integer>	<pre><integer></integer></pre>	<pre><integer></integer></pre>	<integer></integer>	
SRR031729.3941844	chr4	-	75M	75	892	966	75	0	
SRR031728.3674563	chr4	-	75M	75	919	993	75	0	
SRR031729.8532600	chr4	+	75M	75	924	998	75	0	
SRR031729.2779333	chr4	+	75M	75	936	1010	75	0	
SRR031728.2826481	chr4	+	75M	75	949	1023	75	0	
SRR031728.2919098	chr4	-	75M	75	967	1041	75	0	

<DNAStringSet>

seq

SRR031729.3941844 CTGTGGTGACCACACACACAGAATGGTTCGGGCCCAATTAGAGGGTTCCCTGCCCCTTTCCTGGCTAGGTTGTCC SRR031728.3674563 TCGGGCCCAATTAGAGGGTTCCCTGCCCCTTTCCTGGCTAGGTTGTCCGCTAGCTCATTTCCTGGGCTGTTGTTG SRR031729.8532600 CCCAATTAGAGGATTCTCTGCCCCTTTCCTGGCTAGGTTGTCCGGTAGCTCATTTCCCGGGATGTTGTTGTTGTCC SRR031729.2779333 GTTCTCTGCCCCTTTCCTGGCTAGGTTGTCCGCTAGCTCATTTCCCGAGATGTTGTTGTTGTCCCGGGACCCACCT SRR031728.2826481 TTCCTGGCTAGGTTGTCCGCTAGCTCATTTCCCGGGCTGTTGTTGTCCCGGGACACACCTTATTGTGAGTTTG SRR031728.2919098 GCTAGCTCATTTCCCGGGAGGTTGTTGTGTCCCGGGACCCACCTTATTGTGAGTTTGTTGACAGCTCCAAGTTTG

	NM	ntx	ncomptx	${\tt nacomptx}$
	<pre><integer></integer></pre>	<pre><integer></integer></pre>	<pre><integer></integer></pre>	<integer></integer>
SRR031729.3941844	1	0	0	0
SRR031728.3674563	3	0	0	0
SRR031729.8532600	2	0	0	0
SRR031729.2779333	1	0	0	0
SRR031728.2826481	2	0	0	0
SRR031728.2919098	1	0	0	0

seqlengths:

chr2L chr2R chr3L chr3R chr4 chrMchrX chrYHet 23011544 21146708 24543557 27905053 1351857 19517 22422827 347038

> table(U1.GAL_nacomptx)

U1.GAL_nacomptx

0	1	2	3	4	5	6	7	8	9	10	11	12
203800	283	101	107	19	24	2	3	1	3	4	4	4

- > mean(U1.GAL_nacomptx >= 1)
- [1] 0.002715862

Only 0.27% of the alignments in U1.GAL are "almost compatible" with at least 1 transcript in exbytx. Number of "almost compatible" alignments for each transcript:

- > U1.exbytx_nacomp0V00 <- nhitPerSubject(U1.acomp0V00)</pre>
- > names(U1.exbytx_nacomp0V00) <- names(exbytx)</pre>
- > table(U1.exbytx_nacompOV00)

U1.exbytx_nacomp0V00

0	1	2	3	4	5	6	7	8	9	10	12	13	14	17	18
29039	50	8	15	12	2	3	7	5	7	3	2	1	1	1	2
20	21	32	34	44	55	59	77	170							
1	3	2	1	3	2	1	1	1							

> mean(U1.exbytx_nacompOV00 >= 50)

Γ17 0.0001713914

Only 0.017% of the transcripts in exbytx are "almost compatible" with at least 50 alignments in U1.GAL. Finally note that the "query start in transcript" values returned by extractQueryStartInTranscript are also defined for "almost compatible" overlaps:

> head(subset(U1.0V00_qstart, U1.0V00_is_acomp))

	startInTranscript	${\tt firstSpannedExonRank}$	startInFirstSpannedExon
144233	133	1	133
144234	133	1	133
144247	151	1	151
144248	151	1	151
146615	757	7	39
146616	689	8	39

9.2 "Almost compatible" paired-end overlaps

9.2.1 "Almost compatible" paired-end encodings

U3. ovenc contains 5 unique paired-end encodings "almost compatible" with the splicing of the transcript:

> sort(U3.ovenc_table[isCompatibleWithSkippedExons(U3.unique_encodings)])

```
2--1:jm--m:am--m:af--i: 1--2:i--jm:a--am:a--af: 5
2--2:jm--mm:am--mm:af--jm:aa--af: 1--2:i--jm:a--am:a--af: 5
2--1:jm--m:am--m:af--i: 73
```

Paired-end encodings "2--1:jm--m:am--m:af--i:" (73 occurences in U3.ovenc), "1--2:i--jm:a--am:a--af:" (53 occurences in U3.ovenc), and "2--2:jm--mm:am--mm:af--jm:aa--af:" (9 occurences in U3.ovenc), correspond to the following paired-end overlaps:

- "2--1:jm--m:af--i:"
 - paired-end read (1 gap on the first end, no gap on the last end): 000----- 0000
 - transcript: ... >>>> >>>> ...

```
• "1--2:i--jm:a--am:a--af:"
      - paired-end read (no gap on the first end, 1 gap on the
                               0000 00-----00
        last end):
                                           >>> >>>>>
       - transcript:
                              >>>>>>
   • "2--2:jm--mm:am--mm:af--jm:aa--af:"
      - paired-end read (1 gap on the first end, 1 gap on the
        last end):
                                  0----000 00---00
      - transcript:
                              >>>> >>>>>
   Note: switch use of "first" and "last" above if the read was "flipped".
> U3.0V00_is_acomp <- isCompatibleWithSkippedExons(U3.ovenc)
> table(U3.0V00_is_acomp) # 141 "almost compatible" paired-end overlaps
U3.0V00_is_acomp
        TRUE
 FALSE
113686
         141
   Finally, let's extract the "almost compatible" paired-end overlaps from U3.0V00:
```

> U3.acompOV00 <- U3.OV00[U3.OV00_is_acomp]

9.2.2 Tabulate the "almost compatible" paired-end overlaps

Number of "almost compatible" transcripts for each alignment pair in U3.GALP:

```
> U3.GALP_nacomptx <- nhitPerQuery(U3.acompOV00)
> mcols(U3.GALP)$nacomptx <- U3.GALP_nacomptx</pre>
> head(U3.GALP)
```

GAlignmentPairs with 6 alignment pairs and 3 metadata columns:

	seqnames	strand	:	rang	ges		ranges		ntx	ncomptx	nacomptx
	<rle></rle>	<rle></rle>	:	<irange< td=""><td>es></td><td><ir< td=""><td>anges></td><td></td><td><integer></integer></td><td><integer></integer></td><td><integer></integer></td></ir<></td></irange<>	es>	<ir< td=""><td>anges></td><td></td><td><integer></integer></td><td><integer></integer></td><td><integer></integer></td></ir<>	anges>		<integer></integer>	<integer></integer>	<integer></integer>
SRR031715.1138209	chr4	+	:	[169, 20)5]	[326,	362]	-	0	0	0
SRR031714.756385	chr4	+	:	[943, 97	'9]	[1086,	1122]	-	0	0	0
SRR031714.5054563	chr4	+	:	[946, 98	32]	[986,	1022]	-	0	0	0
SRR031715.1722593	chr4	+	:	[966, 100	2]	[1108,	1144]	-	0	0	0
SRR031715.2202469	chr4	+	:	[966, 100	2]	[1114,	1150]	-	0	0	0
SRR031714.3544437	chr4	_	:	[1087, 112	23]	[963,	999]	-	0	0	0

seqlengths:

```
chr2L
            chr2R
                    chr3L
                              chr3R
                                        chr4
                                                chrM
                                                          chrX chrYHet
23011544 21146708 24543557 27905053 1351857
                                                19517 22422827
                                                                347038
```

> table(U3.GALP_nacomptx)

U3.GALP_nacomptx

```
0
           1
                  2
                        3
                                           11
45734
         74
                 4
                       13
                                      1
                                             1
                               1
```

> mean(U3.GALP_nacomptx >= 1)

[1] 0.002051148

Only 0.2% of the alignment pairs in U3.GALP are "almost compatible" with at least 1 transcript in exbytx. Number of "almost compatible" alignment pairs for each transcript:

- > U3.exbytx_nacomp0V00 <- nhitPerSubject(U3.acomp0V00)</pre>
- > names(U3.exbytx_nacompOVOO) <- names(exbytx)</pre>
- > table(U3.exbytx_nacomp0V00)

U3.exbytx_nacomp0V00

0	1	5	8	12	13	66
29143	22	4	1	1	1	1

> mean(U3.exbytx_nacompOV00 >= 50)

[1] 3.427827e-05

Only 0.0034% of the transcripts in exbytx are "almost compatible" with at least 50 alignment pairs in U3.GALP. Finally note that the "query start in transcript" values returned by extractQueryStartInTranscript are also defined for "almost compatible" paired-end overlaps:

> head(subset(U3.0V00_Lqstart, U3.0V00_is_acomp))

	startInTranscript	firstSpannedExonRank	startInFirstSpannedExon
27617	1549	12	45
27629	1562	12	58
27641	1562	12	58
27692	1567	12	63
27814	1549	12	45
42870	659	4	101

> head(subset(U3.0V00_Rqstart, U3.0V00_is_acomp))

	startInTranscript	${\tt firstSpannedExonRank}$	${\tt startInFirstSpannedExon}$
27617	2135	14	115
27629	2141	14	121
27641	2135	14	115
27692	2048	14	28
27814	2136	14	116
42870	866	6	19

10 Detect novel splice junctions

10.1 By looking at single-end overlaps

An alignment in U1.GAL with "almost compatible" overlaps but no "compatible" overlaps suggests the presence of one or more transcripts that are not in our annotations.

First we extract the index of those alignments (nsj here stands for "novel splice junction"):

- > $U1.GAL_is_nsj$ <- $U1.GAL_nacomptx$!= OL & $U1.GAL_ncomptx$ == OL
- > head(which(U1.GAL_is_nsj))
- [1] 57972 57974 58321 67251 67266 67267

We make this an index into U1.0V00:

> U1.0V00_is_nsj <- queryHits(U1.0V00) %in% which(U1.GAL_is_nsj)</pre>

We intersect with U1.0V00_is_acomp and then subset U1.0V00 to keep only the overlaps that suggest novel splicing:

- > U1.0V00_is_nsj <- U1.0V00_is_nsj & U1.0V00_is_acomp</pre>
- > U1.nsj0V00 <- U1.0V00[U1.0V00_is_nsj]</pre>

For each overlap in U1.nsj0V00, we extract the ranks of the skipped exons (we use a list for this as there might be more than 1 skipped exon per overlap):

```
> U1.nsj0V00_skippedex <- extractSkippedExonRanks(U1.ovenc)[U1.0V00_is_nsj]
> names(U1.nsj0V00_skippedex) <- queryHits(U1.nsj0V00)</pre>
> table(elementLengths(U1.nsj0V00_skippedex))
  1 2
          3
              4
                   5
234 116 7
              1
   Finally, we split U1.nsj0V00_skippedex by transcript names:
> f <- factor(names(exbytx)[subjectHits(U1.nsj0V00)], levels=names(exbytx))</pre>
> U1.exbytx_skippedex <- split(U1.nsj0V00_skippedex, f)
   U1.exbytx_skippedex is a named list of named lists of integer vectors. The first level of names (outer names) are
transcript names and the second level of names (inner names) are alignment indices into U1.GAL:
> head(names(U1.exbytx_skippedex)) # transcript names
[1] "FBtr0300689" "FBtr0300690" "FBtr0330654" "FBtr0309810" "FBtr0306539" "FBtr0306536"
   Transcript FBtr0089124 receives 7 hits. All of them skip exons 9 and 10:
> U1.exbytx_skippedex$FBtr0089124
$ 104549
[1] 9 10
$`104550`
[1] 9 10
$`104553`
[1] 9 10
$`104557`
[1] 9 10
$`104560`
[1] 9 10
$`104572`
[1] 9 10
$ 104577
[1] 9 10
   Transcript FBtr0089147 receives 4 hits. Two of them skip exon 2, one of them skips exons 2 to 6, and one of them
skips exon 10:
> U1.exbytx_skippedex$FBtr0089147
$`72828`
[1] 10
$`74018`
```

[1] 2 3 4 5 6

```
$`74664`
[1] 2
$`74670`
[1] 2
```

A few words about the interpretation of U1.exbytx_skippedex: Because of how we've conducted this analysis, the aligments reported in U1.exbytx_skippedex are guaranteed to not have any "compatible" overlaps with other known transcripts. All we can say, for example in the case of transcript FBtr0089124, is that the 7 reported hits that skip exons 9 and 10 show evidence of one or more unknown transcripts with a splice junction that corresponds to the gap between exons 8 and 11. But without further analysis, we can't make any assumption about the exons structure of those unknown transcripts. In particular, we cannot assume the existence of an unknown transcript made of the same exons as transcript FBtr0089124 minus exons 9 and 10!

By looking at paired-end overlaps

[COMING SOON...]

11 sessionInfo()

```
> sessionInfo()
R version 3.0.2 (2013-09-25)
Platform: x86_64-unknown-linux-gnu (64-bit)
locale:
 [1] LC_CTYPE=en_US.UTF-8
                                LC_NUMERIC=C
                                                            LC_TIME=en_US.UTF-8
 [4] LC_COLLATE=C
                                LC_MONETARY=en_US.UTF-8
                                                            LC_MESSAGES=en_US.UTF-8
 [7] LC_PAPER=en_US.UTF-8
                                LC_NAME=C
                                                            LC_ADDRESS=C
[10] LC_TELEPHONE=C
                                LC_MEASUREMENT=en_US.UTF-8 LC_IDENTIFICATION=C
attached base packages:
[1] parallel stats
                        graphics grDevices utils
                                                       datasets methods
                                                                            base
other attached packages:
 [1] BSgenome.Dmelanogaster.UCSC.dm3_1.3.19
                                                 BSgenome.Scerevisiae.UCSC.sacCer2_1.3.19
 [3] KEGGgraph_1.20.0
                                                 graph_1.40.0
 [5] XML_3.98-1.1
                                                 KEGG.db_2.10.1
 [7] RSQLite_0.11.4
                                                 DBI_0.2-7
 [9] BSgenome. Hsapiens. UCSC. hg19_1.3.19
                                                 BSgenome_1.30.0
[11] TxDb.Hsapiens.UCSC.hg19.knownGene_2.10.1
                                                 edgeR_3.4.0
[13] limma_3.18.0
                                                 DESeq_1.14.0
[15] lattice_0.20-24
                                                 locfit_1.5-9.1
[17] AnnotationHub_1.2.0
                                                 TxDb.Dmelanogaster.UCSC.dm3.ensGene_2.10.1
[19] GenomicFeatures_1.14.0
                                                 AnnotationDbi_1.24.0
[21] Biobase_2.22.0
                                                 pasillaBamSubset_0.0.8
[23] Rsamtools_1.14.1
                                                 Biostrings_2.30.0
[25] GenomicRanges_1.14.1
                                                 XVector_0.2.0
[27] IRanges_1.20.0
                                                 BiocGenerics_0.8.0
loaded via a namespace (and not attached):
 [1] BiocInstaller_1.12.0 BiocStyle_1.0.0
                                                RColorBrewer_1.0-5
                                                                     RCurl_1.95-4.1
 [5] annotate_1.40.0
                          biomaRt_2.18.0
                                                                      genefilter_1.44.0
                                                bitops_1.0-6
```

[9] geneplotter_1.40.0 grid_3.0.2 rjson_0.2.13 rtracklayer_1.22.0 [13] splines_3.0.2 stats4_3.0.2 survival_2.37-4 tools_3.0.2 [17] xtable_1.7-1 zlibbioc_1.8.0