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1 Introduction

Nearly all sequence junctions formed by a rearrangement occur in non-coding regions of the genome. With whole genome sequencing, we can identify the location of the noncoding sequence junctions. When these junctions occur within a gene (usually intronic), we can evaluate whether the new transcript produced by the rearrangement is in-frame. This vignette provides an overview, extending the rearrangement analyses in the svrearrange package.

Our analysis to identify in-frame fusions begin with a RearrangementList for ovarian cell line CGOV11T_1 created by and available from the svrearrange package.

2 Analysis pipeline

```
suppressPackageStartupMessages(library(svrearrange))
##load_all("svrearrange")
##suppressPackageStartupMessages(library(svfusions))
extdata <- system.file("extdata", package="svrearrange")
rfile <- file.path(extdata, "CGOV11T_1.bam.rds")
rlist <- readRDS(rfile)
rlist
## An object of class 'RearrangementList'
## number rearrangement objects: 68
## Use '[[i]]' to return a single Rearrangement object'</pre>
```

The above rlist contains 68 rearrangements. We begin by selecting only the rearrangements in which both ends of a sequence junction are within 5kb of a known transcript.

```
build <- "hg19"
near.coding <- seqJunctionNearTx(rlist, build)
rlist <- rlist[ near.coding ]
length(rlist)
## [1] 26</pre>
```

As discussed in the vignette for the svrearrange package, each candidate rearrangement has two possible 5-prime to 3-prime orientations that can be inferred by the orientation of the rearranged read pairs (RRPs) and split reads (SRs). We use the function fiveTo3List to order each rearrangement in the rlist object by these two orientations. In doing so, the length of the rlist object doubles.

```
rlist2 <- fiveTo3List(rlist, build)
length(rlist2)
## [1] 52</pre>
```

Having established the 5-prime to 3-prime orientations in the rlist2 object, we make the definition of the sequence junctions more precise.

```
jxns <- seqJunctions_Rlist(rlist2)
jxns
## GRanges object with 52 ranges and 2 metadata columns:
## seqnames ranges strand | rid 3p</pre>
```

```
<Rle>
                            <IRanges> <Rle> | <character>
                                                                 <GRanges>
##
           chr3 [ 58908808, 58908808]
                                         - | 10546-10582 chr3:59949878:-
    5p
##
    5p
           chr3 [ 59949878, 59949878]
                                           + | 10582-10546 chr3:58908808:+
##
    5p
           chr3 [171752966, 171752966]
                                         - | 13607-44076 chr15:99550846:-
##
          chr15 [ 99550785, 99550785]
                                          - | 44076-13607 chr3:171753007:-
    5p
##
           chr3 [187389262, 187389262]
                                           - | 14161-14186 chr3:187988635:-
    5p
##
                                  ... ... .
                                                      . . .
##
    5p
           chr2 [213914447, 213914447]
                                         + |
                                                9181-9136 chr2:213286255:+
##
          chr2 [218936235, 218936235]
                                         - | 9258-33360 chr11:62154698:+
    5p
##
    5p
          chr11 [ 62154698, 62154698]
                                          - | 33360-9258 chr2:218936235:+
##
          chr2 [219352749, 219352749]
                                          + | 9281-33369 chr11:62362005:-
    5p
##
          chr11 [ 62362005, 62362005]
                                          + | 33369-9281 chr2:219352749:-
    5p
##
    seqinfo: 23 sequences from an unspecified genome
```

In addition, we require the 3-prime genomic region of the junction to lie strictly within a transcript, while the 5-prime genomic region forming the junction can occur either in the promoter or within the 5-prime transcript. Operationally, we define the promoter as the genomic region 5kb upstream of the transcription start site of the 5-prime gene. We refer to the remaining junctions as coding_jxns. As the coding junctions are named by the rearrangement ids, we can use these names to subset the rlist2 object, excluding those rearrangements that do not meet the above criteria.

```
coding_jxns <- codingJunctions(jxns, build)</pre>
coding_jxns
## GRanges object with 44 ranges and 3 metadata columns:
##
                                         ranges strand |
                segnames
                                                                rid
                                     <IRanges> <Rle> | <character>
##
                   <Rle>
                 chr3 [ 58908808, 58908808]
##
    10546 - 10582
                                                   - | 10546-10582
    10582-10546 chr3 [ 59949878, 59949878]
##
                                                   + | 10582-10546
    13607-44076 chr3 [171752966, 171752966]
                                                   - | 13607-44076
##
                 chr3 [187389262, 187389262]
##
    14161-14186
                                                    - | 14161-14186
    16502-16504 chr5 [ 59150971, 59150971]
                                                   + | 16502-16504
##
##
            . . .
                    . . .
                                                  . . . .
                 chr2 [213914447, 213914447]
##
     9181-9136
                                                          9181-9136
                 chr2 [218936235, 218936235]
##
     9258-33360
                                                         9258-33360
##
     33360-9258
                   chr11 [ 62154698, 62154698]
                                                    - | 33360-9258
     9281-33369 chr2 [219352749, 219352749]
                                                    + | 9281-33369
                   chr11 [ 62362005, 62362005]
##
     33369-9281
                                                    + | 33369-9281
##
                              3р
                                                  tx_name
##
                       <GRanges>
                                               <character>
##
    10546-10582 chr3:59949878:-
                                                NM_198463
##
    10582-10546 chr3:58908808:+
                                   NM_001166243, NM_002012
##
    13607-44076 chr15:99550846:-
                                        promoter_NM_022763
##
    14161-14186 chr3:187988635:-
                                        promoter_NM_001048
##
    16502-16504 chr5:59646153:+ NM_001104631,NM_001165899
##
            . . .
                                   NM_016260, NM_001079526
##
      9181-9136 chr2:213286255:+
##
     9258-33360 chr11:62154698:+
                                                NM_198483
     33360-9258 chr2:218936235:+ NM_001083926,NM_025080
##
     9281-33369 chr11:62362005:-
                                                 NM_020935
```

```
## 33369-9281 chr2:219352749:- promoter_NM_022830
## -----
## seqinfo: 23 sequences from hg19 genome
rlist2 <- rlist2[ names(coding_jxns) ]</pre>
```

Next, we use the fuseCDS_Rlist function to extract for each rearrangement the full CDS of the fused sequence in the somatic genome (fusions), the partial CDS of the 5-prime (tum.5p) and 3-prime (tum.3p) transcripts that are involved in the fusion, and the full CDS of the 5-prime (ref.5p) and 3-prime transcripts in the reference genome. To speed up computation in the remaining portions of this vignette, we restrict the RearrangementList object to a rearrangement involving the fusion of the tyrosine kinase IKAROS to a known driver ERBB4. Note, we use a single bracket [when subsetting the RearrangementList so that the resulting object is still an instance of RearrangementList (albeit a length-one list).

```
ikaros.erbb4 <- "9136-9181"
coding_jxns <- coding_jxns[ikaros.erbb4]
cds.list <- fuseCDS_Rlist(rlist2[ikaros.erbb4], coding_jxns)
cds.list
## List of length 6
## names(6): fusions tum.5p tum.3p ref.5p ref.3p coding_junctions</pre>
```

For each fusion, we translate the CDS in each of the three possible reading frames and partition the amino acid sequence of the fusion protein into the sequences derived from the 5-prime and 3-prime transcripts. In particular, by partioning the amino acid sequence of the fusion into its 5-prime and 3-prime parts, we can compare the 5-prime partition to the amino acid sequence of the 5-prime reference protein and the 3-prime partition to the amino acid sequence of the 3-prime reference protein. The function translateCDS translates the CDS in the 3 possible reading frames.

```
proteins <- translateCDS(cds.list)
proteins
## List of length 5
## names(5): fusion tum5p tum3p ref5p ref3p</pre>
```

The amino acid sequence obtained by translating the CDS involved in the 5-prime and 3-prime transcripts are represented as AAStringSet objects. Since a single gene can have multiple transcripts, we translate every possible combination of 5-prime and 3-prime transcripts in each of the 3 frames. Here, we show the amino acid sequences for the two possible combinations of transcripts involving IKAROS and ERBB4:

```
proteins$fusion[[1]]
## $frame1
##
    A AAStringSet instance of length 2
       width seq
                                                             names
        363 MKPATGLWVWVSLLVAAGTVQP...RSQDRYEFSSHIVRGEHTFH* NM_001042599-NM_0...
        363 MKPATGLWVWVSLLVAAGTVQP...RSQDRYEFSSHIVRGEHTFH* NM_001042599-NM_0...
## [2]
##
## $frame2
##
    A AAStringSet instance of length 2
##
      width seq
                                                             names
## [1]
        362 *SRRQDFGSG*AFSWRRGPSSP...TEARTVMSFHHTLFEGSTHST NM_001042599-NM_0...
        362 *SRRQDFGSG*AFSWRRGPSSP...TEARTVMSFHHTLFEGSTHST NM_001042599-NM_0...
```

```
##
## $frame3
## A AAStringSet instance of length 2
## width seq names
## [1] 362 EAGDRTLGLGEPSRGGGDRPAQ...QKPGPL*VFITHCSRGAHIPL NM_001042599-NM_0...
## [2] 362 EAGDRTLGLGEPSRGGGDRPAQ...QKPGPL*VFITHCSRGAHIPL NM_001042599-NM_0...
```

We can also list the amino acid sequences derived from the 5-prime gene and the 3-prime gene:

```
proteins$tum5p[[1]]
## $frame1
## A AAStringSet instance of length 2
    width seq
## [1] 27 MKPATGLWVWVSLLVAAGTVQPSDSQS
                                                          NM_001042599-NM_0...
## [2] 27 MKPATGLWVWVSLLVAAGTVQPSDSQS
                                                          NM_001042599-NM_0...
##
## $frame2
## A AAStringSet instance of length 2
     width seq
                                                          names
## [1] 26 *SRRQDFGSG*AFSWRRGPSSPAILS
                                                          NM_001042599-NM_0...
## [2]
         26 *SRRODFGSG*AFSWRRGPSSPAILS
                                                          NM_001042599-NM_0...
## $frame3
## A AAStringSet instance of length 2
     width seq
                                                          names
## [1] 26 EAGDRTLGLGEPSRGGGDRPAQRFSV
                                                          NM_001042599-NM_0...
                                                          NM_001042599-NM_0...
## [2]
       26 EAGDRTLGLGEPSRGGGDRPAQRFSV
proteins$tum3p[[1]]
## $frame1
## A AAStringSet instance of length 2
     width seq
## [1] 335 WVNLTSATTVDEATSSAVHWRS...TEARTVMSFHHTLFEGSTHST NM_016260
## [2] 335 WVNLTSATTVDEATSSAVHWRS...TEARTVMSFHHTLFEGSTHST NM_001079526
## $frame2
   A AAStringSet instance of length 2
     width seq
                                                          names
## [1] 334 G*TSOVOLLWTKLOAAOFTGGA...LOKPGPL*VFITHCSRGAHIP NM_016260
## [2] 334 G*TSQVQLLWTKLQAAQFTGGA...LQKPGPL*VFITHCSRGAHIP NM_001079526
##
## $frame3
## A AAStringSet instance of length 2
    width seq
## [1] 334 GKPHKCNYCGRSYKQRSSLEEH...YRSQDRYEFSSHIVRGEHTFH NM_016260
## [2] 334 GKPHKCNYCGRSYKQRSSLEEH...YRSQDRYEFSSHIVRGEHTFH NM_001079526
```

The function partitionAASequence reorganizes the list of amino acid sequences by frame to facilitate downstream computation.

```
partition.fusion <- partitionAASequence(proteins)
partition.fusion
## List of length 3
## names(3): frame1 frame2 frame3</pre>
```

For a fusion to be in-frame, we require that the amino acid sequence derived from the 5- and 3-prime transcripts to be subsequences of the full amino acid sequence of the reference genome that was translated in the same frame. In addition, we require that there be no premature stop codons. The following code accomplishes these two tasks, with the ref.frames object below containing the full amino acid sequence of the transcripts associated with IKAROS and ERBB4.

```
nostop.list <- noPrematureStop(partition.fusion)</pre>
nostop.list
## $frame1
## LogicalList of length 1
## [["9136-9181"]] TRUE TRUE
## $frame2
## LogicalList of length 1
## [["9136-9181"]] FALSE FALSE
##
## $frame3
## LogicalList of length 1
## [["9136-9181"]] FALSE FALSE
ref.frames <- organizeReferenceByFrame(proteins)</pre>
inframe.list <- inFrameList(fusion.frames=partition.fusion,</pre>
                             ref.frames=ref.frames)
inframe.list
## $frame1
## LogicalList of length 1
## [["9136-9181"]] TRUE TRUE
## $frame2
## LogicalList of length 1
## [["9136-9181"]] FALSE FALSE
## $frame3
## LogicalList of length 1
## [["9136-9181"]] FALSE FALSE
```

Each element of the nostop.list and inframe.list lists are an object of class LogicalList, each evaluated from a different reading frame. The LogicalList objects are named by the rearrangement identifier. The helper function inFrameNoStop combines these to lists to create a single list that is TRUE for rearrangements that are in-frame and have no premature stop codons.

```
inframe.nostop <- inFrameNoStop(nostop.list, inframe.list)
inframe.nostop
## $frame1
## LogicalList of length 1
## [["9136-9181"]] TRUE TRUE</pre>
```

```
##
## $frame2
## LogicalList of length 1
## [["9136-9181"]] FALSE FALSE
##
## $frame3
## LogicalList of length 1
## [["9136-9181"]] FALSE FALSE
```

Finally, validFusions returns a list object containing only the fusions that are in-frame and have no premature stops. For each such fusion, the amino acid sequence, CDS of the fusion, partial CDS of the 5-prime and 3-prime transcripts, and the full CDS of the reference transcripts are available.

The above data can be summarized in tabular format using the fusionTable2 function:

```
tab <- fusionTable2(valid.fusions)</pre>
head(tab)
## DataFrame with 2 rows and 15 columns
    gene.5prime gene.3prime
                               tx.5prime
                                           tx.3prime junction.5prime
##
    <character> <character> <character> <character>
                                                         <character>
## 1
          ERBB4
                    IKZF2 NM_001042599
                                           NM_016260 chr2:213286255
## 2
                     IKZF2 NM_001042599 NM_001079526 chr2:213286255
          ERBB4
    junction.3prime exons.5prime exons.3prime strand.5prime strand.3prime
        <character> <character> <character> <character>
##
                                                             <character>
## 1 chr2:213914447
                     exons: 1-1 exons: 5-7
## 2 chr2:213914447 exons: 1-1 exons: 5-7
     rearrangement.id aa.5prime.start aa.5prime.end aa.3prime.start
                                       <integer>
##
         <character>
                        <numeric>
                                                      <numeric>
## 1
           9136-9181
                                  1
                                               27
                                                            192
           9136-9181
## 2
                                   1
                                               27
                                                              166
##
    aa.3prime.end
        <integer>
##
## 1
              527
## 2
              501
```

Note, this table provides both the genomic coordinates of the fusion (junction.5prime, junction.3prime), but also the portion of the amino acid sequence that was retained by the fusion (aa.5prime.start, aa.5prime.end, aa.3prime.start, aa.3prime.end). In the following code chunk, we construct amino acid ranges for the genes involved in the fusion and the protein domains for these genes in the reference genome.

```
tumor_aa_ranges <- aa_granges(tab)
extdata2 <- system.file("extdata", package="svfusions")</pre>
```

```
up <- readRDS(file.path(extdata2, "uniprot.rds"))</pre>
  up2 <- uniprotFeatures(up, tab, strwrap.width=20)</pre>
  domain_aa_ranges <- GRanges(up2$hugo, IRanges(up2$start, up2$end),</pre>
                                    chromosome=up2$seqnames,
                                    description=up2$description,
                                    short.desc=up2$short.desc,
                                    aa_len=up2$aa_len)
  domains <- subsetByOverlaps(domain_aa_ranges, tumor_aa_ranges, type="within")</pre>
  domains
## GRanges object with 4 ranges and 4 metadata columns:
##
                          ranges strand | chromosome description short.desc
          segnames
             <Rle> <IRanges> <Rle> | <factor> <character> <character>
    [1] IKZF2 [168, 190] * | chr2 C2H2-type 3 C2H2-type 3 [2] IKZF2 [196, 219] * | chr2 C2H2-type 4 C2H2-type 4 [3] IKZF2 [471, 493] * | chr2 C2H2-type 5 C2H2-type 5 [4] IKZF2 [499, 523] * | chr2 C2H2-type 6 C2H2-type 6
##
##
##
##
              aa_len
##
##
          <integer>
##
     [1]
                 526
     [2]
                 526
##
     [3]
                  526
##
##
     [4]
                 526
##
    seqinfo: 2 sequences from an unspecified genome; no seqlengths
```