Using refGenome package

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1 refGenome package

The refGenome package provides functionality for managing of genome annotation data, especially for Ensembl and UCSC data.

2 Object types inside refGenome package

The central classes inside this package are refGenome derived (S4) classes. The class contains two slots: ev (environment) and basedir (character). All annotation data is kept in data.frames inside the ev slot. Saving and loading refGenome derived objects works on the complete content of the environment. This mechanism also avoids generation of copies and allows addition of new data inside of member functions. The basedir slot keeps a path on a hard-disc which is intended as location where data files and object versions can be kept.

The package contains three derived class lineages refGenome, refExons and refJunctions. For each lineage there are classes for Ensembl and UCSC defined, e.g. ensemblGenome and ucscGenome. The exon classes focus on annotated exon positions and the junction classes focus on adjacent exons.

2.1 Creation of empty refGenome objects

Empty objects of refGenome derived classes can be created with ensembleGenome() or ucscGenome(). After creation of an empty object the first step usually is to set the basedir address:

- > library(refGenome)
- > beg <- ensemblGenome()</pre>
- > basedir(beg) <- system.file("extdata", package="refGenome")

The "basedir" folder is intended to contain all data which is associated with the current annotation set, e.g. downloaded gtf files, saved object data, saved SQLite versions of the data and potenially sequence information. In order to fill an empty object, annotation data has to be imported from external files.

2.2 Importing annotation data

The basic importing mechanism for refGenome objects is to import a "gtf" file. Therefore, the "gtf" files have to be downloaded. The download source and mechanism is explained for ensemblGenome and ucscGenome separately. There are specialized mechanisms in order to provide additional information either from within the gtf file (ensembl) or via other external files (ucsc).

2.3 Saving and loading data

The data content of refGenome objects can be saved and re-loaded in several ways. One way is the saveGenome method where the content is written into a compressed ".RData" file. One alternative is to write the content into a SQLite database via writeDB.

3 Ensembl Genomes

The ensemblGenome class is specialized for managing annotation data for ensemble Genomes.

3.1 Download and import data

For ensemblGenome objects, gtf files can be downloaded from Ensemble servers. Therefore, go to

```
http://www.ensembl.org/info/data/ftp/index.html
```

and choose a file from the "Gene sets" column. They are labeled "GTF". For example Version 62 of human genomic annotation can be downloaded from

```
\label{lem:condition} $$ ftp://ftp.ensembl.org/pub/release-80/gtf/homo_sapiens/Homo_sapiens. $$ GRCh38.80.gtf.gz $$
```

A copy of the obtained file should then be placed in the the "basedir" directory. With the appropriate setting of basedir, annotation data can be imported with:

```
2 GL000213.1 protein_coding
                                      CDS 138767 139287
  3 GL000213.1 protein_coding start_codon 139285 139287
  4 GL000213.1 protein_coding
                                    exon 134276 134390
  5 GL000213.1 protein_coding
                                      CDS 134276 134390
  6 GL000213.1 protein_coding
                                      exon 133943 134116
  score strand frame
                         protein_id transcript_name
                                <NA> BX072566.1-201
                   0 ENSP00000329990 BX072566.1-201
3
                   0
                                <NA>
                                     BX072566.1-201
4
                                <NA>
                                     BX072566.1-201
5
                   1 ENSP00000329990
                                     BX072566.1-201
                                <NA>
                                     BX072566.1-201
         gene_id
                   transcript_id gene_name exon_number
1 ENSG00000237375 ENST00000327822 BX072566.1
2 ENSG00000237375 ENST00000327822 BX072566.1
3 ENSG00000237375 ENST00000327822 BX072566.1
                                                       1
4 ENSG00000237375 ENST00000327822 BX072566.1
                                                       2
                                                       2
5 ENSG00000237375 ENST00000327822 BX072566.1
6 ENSG00000237375 ENST00000327822 BX072566.1
                                                       3
```

The top lines of the contained table are shown when the object is printed.

4 UCSC Genomes

Downloading of annotation data for UCSC genomes is a bit more complicated than for Ensemble Genomes because additional data must be downloaded in separate files. The Homepage for UCSC browser can be found under:

```
http://genome.ucsc.edu/
```

In order to import UCSC annotation data into refGenome objects files containing the data have to be downloaded from the USCS Table Browser which can be found under:

```
http://genome.ucsc.edu/cgi-bin/hgTables
```

or by following the "Table Browser" link in the left panel on the homepage. On the Table Browser:

- Select genome, assembly and track (UCSC genes)
- Choose table (knownGene)
- Choose output format (GTF -gene transfer format for knownGene table)
- Insert a name for the output file
- Download the file (get output)

The basic table to be imported is "knownGene". The knownGene table has to be downloaded in GTF format (otherwise the read.gtf function will complain about "wrong number of columns").

In order to extend the available information additionally the tables "kgXref", "knownToEnsembl" and "knownIsoforms" can be downloaded and imported. These tables come in plain "csv" format. Select "all fields from selected table" as output format.

Do not use "add custom tracks" or modify the tables elsewhere tracks because the importing functions will check for apropriate number of columns.

After downloading, all tables should be placed into a separate folder which we from now on call "basedir".ucscGenome objects keep a basedir as standard location for all writing and reading procedures.

```
> uc <- ucscGenome()
> basedir(uc) <- "/my/ucsc/basedir"
> read.gtf(uc, "ucsc_knownGene.gtf")
> addXref(uc, "kgXref.csv")
> addEnsembl(uc, "knownToEnsembl.csv")
> addIsoforms(uc, "ucsc_knownisoforms.csv")
```

4.1 Load stored data

Once, annotation data is imported and stored, ucscGenome objects can be restored with the loadGenome function which is shown below on example data:

```
> ucfile <- system.file("extdata", "hs.ucsc.small.RData", package="refGenome")
> uc <- loadGenome(ucfile)
> ensfile <- system.file("extdata", "hs.ensembl.62.small.RData", package="refGenome")
> ens <- loadGenome(ensfile)</pre>
```

5 Extracting data subsets

There are specialized functions for extracting data for multiple purposes.

5.1 Extracting data for sets of seqid's

For preparation of seqid based extraction, the contained seqid's can be tabled:

```
> tableSeqids(ens)
1 GL000213.1
111 24
```

Extraction of subsets based on seqid can be done with extractSeqids. The sequence id's for extraction are specified as regular expression:

```
> en1 <- extractSeqids(ens, "^1$")
> en1
```

```
Object of class 'ensemblGenome' with 111 rows and 15 columns.
   id seqid start end feature score strand frame
         1 11869 12227 exon
34
         1 11872 12227
                          exon
41 3
         1 11874 12227
                        exon
28 4
         1 12010 12057
                        exon
29 5
         1 12179 12227
                          exon
                           CDS
         1 12190 12227
          gene_id
                    transcript_id
                                          source
25 ENSG00000223972 ENST00000456328
                                      pseudogene
34 ENSG00000249291 ENST00000515242 protein_coding
41 ENSG00000253101 ENST00000518655 protein_coding
28 ENSG00000223972 ENST00000450305
                                      pseudogene
29 ENSG00000223972 ENST00000450305
                                      pseudogene
35 ENSG00000249291 ENST00000515242 protein_coding
   gene_name transcript_name exon_number
                DDX11L1-002
25
     DDX11L1
34 AL627309.2 AL627309.2-201
                                       1
    DDX11L11
              DDX11L11-201
                                       1
28
     DDX11L1
                 DDX11L1-001
                                       1
     DDX11L1
29
                 DDX11L1-001
                                       2
35 AL627309.2 AL627309.2-201
                  transcript_biotype
25
                processed_transcript
34
             nonsense_mediated_decay
41
             nonsense_mediated_decay
28 transcribed_unprocessed_pseudogene
29 transcribed_unprocessed_pseudogene
35
             nonsense_mediated_decay
```

It looks cumbersome for single chromosomes but allows extraction of complex patterns.

5.2 Extracting primary assembly data

Usually the interesting part of the annotation data is the the primary assembly (where alternative haplotypes are excluded). Therefore functions which return the proper terms are supplied:

```
> ensPrimAssembly()
[1] "^([0-9]{1,2})$|^[XY]|MT$"
> ucPrimAssembly()
[1] "^chr[0-9XYM]{1,2}$"

Extraction of primary assembly seqid's i is done by:
> enpa<-extractSeqids(ens,ensPrimAssembly())
> tableSeqids(enpa)
```

```
1
111
> ucpa<-extractSeqids(uc,ucPrimAssembly())</pre>
> tableSeqids(ucpa)
chr1
   6
```

Extract features 5.3

Subsets defined by features can allso be tabled and extracted:

> tableFeatures(enpa)

41

42

```
CDS
           exon start_codon stop_codon
  8
             98
                           3
```

> enpf<-extractFeature(enpa, "exon")</pre>

```
> enpf
Object of class 'ensemblGenome' with 98 rows and 15 columns.
   id seqid start
                    end feature score strand frame
          1 11869 12227
                           exon
34 2
          1 11872 12227
                           exon
41 3
          1 11874 12227
                           exon
28 4
          1 12010 12057
                           exon
29 5
          1 12179 12227
                           exon
          1 12595 12721
                           exon
                     transcript_id
           gene_id
                                           source
25 ENSG00000223972 ENST00000456328
                                       pseudogene
34 ENSG00000249291 ENST00000515242 protein_coding
41 ENSG00000253101 ENST00000518655 protein_coding
28 ENSG00000223972 ENST00000450305
                                       pseudogene
29 ENSG00000223972 ENST00000450305
                                       pseudogene
42 ENSG00000253101 ENST00000518655 protein_coding
    gene_name transcript_name exon_number
                  DDX11L1-002
      DDX11L1
                                        1
34 AL627309.2 AL627309.2-201
                                        1
    DDX11L11
                 DDX11L11-201
                                        1
                  DDX11L1-001
28
      DDX11L1
                                        1
29
      DDX11L1
                  DDX11L1-001
                                        2
     DDX11L11
                 DDX11L11-201
42
                   transcript_biotype
25
                 processed_transcript
34
```

nonsense_mediated_decay

nonsense_mediated_decay

nonsense_mediated_decay

28 transcribed_unprocessed_pseudogene 29 transcribed_unprocessed_pseudogene

5.4 Extract data for single genes and transcripts

There are some functions which extract objects that contain data for single genes (or transcripts). These functions provide a closer insight into specific regeions.

Objects which contain data for vectors of gene-names can be extracted with

```
> dxe <- extractByGeneName(enpa, "DDX11L1")
> dxu <- extractByGeneName(ucpa, "DDX11L1")</pre>
```

When gene-names did not match in the gtf-table of the object, a message including all names of not matching gene-names will be printed. When no gene-name matches, a message will be printed and the function returns NULL, which can be tested for later on.

Additionally subsets can also be extacted based on gene-id

```
> dxe <- extractByGeneId(enpa, "ENSG00000223972")
> dxu <- extractByGeneId(ucpa, "ENSG00000223972")</pre>
```

[1] "ENSG00000223972"

From these extracts we can view the contained transcripts with the tableTranscript.id function:

```
> tableTranscript.id(enpa)
```

```
ENST00000408384 ENST00000417324 ENST00000423562
                              8
ENST00000430492 ENST00000438504 ENST00000450305
                             12
ENST00000456328 ENST00000461467 ENST00000469289
              3
                              2
ENST00000473358 ENST00000488147 ENST00000515242
              3
                             11
ENST00000518655 ENST00000537342 ENST00000538476
ENST00000541675
> tableTranscript.id(ucpa)
uc001aaa.3 uc010nxr.1
         3
                    3
```

Data for interesting transcripts can be extracted by extractTranscript:

```
> extractTranscript(ens, "ENST00000456328")
```

```
Object of class 'ensemblGenome' with 3 rows and 15 columns.
    transcript_id id seqid start end feature score strand
1 ENST00000456328 1 1 11869 12227 exon . +
2 ENST00000456328 9 1 12613 12721 exon . +
3 ENST00000456328 14 1 13221 14409 exon . +
```

```
gene_id
                           source gene_name
     . ENSG00000223972 pseudogene
                                    DDX11L1
      . ENSG00000223972 pseudogene
                                    DDX11L1
      . ENSG00000223972 pseudogene
                                    DDX11L1
 transcript_name exon_number
                               transcript_biotype
     DDX11L1-002
                           1 processed_transcript
1
2
     DDX11L1-002
                           2 processed_transcript
3
     DDX11L1-002
                           3 processed_transcript
> extractTranscript(uc, "uc010nxr.1")
Object of class 'ucscGenome' with 3 rows and 14 columns.
  transcript_id id segid start
                                end feature score strand
    uc010nxr.1 4 chr1 11874 12227
                                       exon
    uc010nxr.1 5 chr1 12646 12697
                                       exon
                                                0
    uc010nxr.1 6 chr1 13221 14409
                                       exon
 frame
          gene_id
                          source gene_name
                                                   ensembl
      . uc010nxr.1 hg19_knownGene DDX11L1 ENST00000456328
1
      . uc010nxr.1 hg19_knownGene DDX11L1 ENST00000456328
      . uc010nxr.1 hg19_knownGene DDX11L1 ENST00000456328
  clusterId
         1
1
2
         1
3
         1
```

6 Accumulate data for whole genes

The function getGenePositions accumulates position data for whole genes. Genes are grouped by gene_name. For both, ensemblGenome and ucscGenome the gene_name column is not present after the standard gtf-import. For ucscGenome, addXref must be used. Respective warnings are thrown.

```
> gpe <- getGenePositions(ens)</pre>
> gpe
            gene_id gene_name
                                    seqid start
  2 ENSG00000223972
                      DDX11L1
                                        1 11869 14409
  7 ENSG00000249291 AL627309.2
                                        1 11872 14412
8 8 ENSG00000253101 DDX11L11
                                       1 11874 14409
  3 ENSG00000227232
                        WASH7P
                                        1 14363 29806
  6 ENSG00000243485 MIR1302-10
                                        1 29554 31109
  1 ENSG00000221311 MIR1302-10
                                        1 30366
                                                  30503
5
  5 ENSG00000237613 FAM138A
                                        1 34554 36081
  4 ENSG00000237375 BX072566.1 GL000213.1 108007 139339
  strand start_codon stop_codon
2
                 NA
                            NA
7
              12190
                            NA
              13548
                         13817
3
                 NΑ
                            NΑ
6
                 NA
                            NΑ
```

There is a slight difference between both results: The last column is gene_id for *ensemblGenome* and clusterID for *ucscGenome*. This is due to different information which is available for each.

7 Exon and splice-junction based views (only for Ensembl genomes)

7.1 Extract exon based table

[refExons.refGenome] Extracting tables. [refExons.refGenome] Adding 'CDS'.

[refExons.refGenome] Adding 'start_codon'.

Exon based view on annotation data can be obtained with ensemblexons which returns an object of class ensemblexons. Basically ensemblexons calls extractFeature for feature type "exon". Information about presence of cds start or end and start-codon or stop-codon is added.

```
[refExons.refGenome] Adding 'stop_codon'.
[refExons.refGenome] Finished.
[refExons.refGenome] Extracting tables.
[refExons.refGenome] Adding 'CDS'.
[refExons.refGenome] Adding 'start_codon'.
[refExons.refGenome] Adding 'stop_codon'.
[refExons.refGenome] Finished.
> enex
Object of class 'ensemblExons' with 109 rows and 18 columns.
  id seqid start end score strand frame
                                                  gene_id
                         . + . ENSG00000223972
53 1
         1 11869 12227
74 2
                                       . ENSG00000249291
         1 11872 12227
         1 11874 12227
                                       . ENSG00000253101
47 4
         1 12010 12057
                                        . ENSG00000223972
48 5
         1 12179 12227
                                        . ENSG00000223972
         1 12595 12721
                                        . ENSG00000253101
    transcript_id
                          source gene_name
53 ENST00000456328
                      pseudogene
                                    DDX11L1
```

```
74 ENST00000515242 protein_coding AL627309.2
77 ENST00000518655 protein_coding
                                     DDX11L11
47 ENST00000450305
                        pseudogene
                                      DDX11L1
48 ENST00000450305
                        pseudogene
                                      DDX11L1
78 ENST00000518655 protein_coding
                                     DDX11L11
   transcript_name exon_number
53
       DDX11L1-002
74
    AL627309.2-201
77
      DDX11L11-201
                              1
47
       DDX11L1-001
                              1
48
       DDX11L1-001
                              2
78
      DDX11L11-201
                    transcript_biotype cds_start cds_end
53
                 processed_transcript
                                               NA
                                                       NA
74
              nonsense_mediated_decay
                                              318
                                                        0
77
              nonsense_mediated_decay
                                               NA
                                                       NA
47 transcribed_unprocessed_pseudogene
                                               NΑ
                                                       NΑ
                                                       NA
48 transcribed_unprocessed_pseudogene
                                               NA
              nonsense_mediated_decay
                                               NΑ
                                                       NA
   start_codon stop_codon
53
            NΑ
74
           318
                        NΑ
77
                        NA
            NA
47
            NA
                        NA
48
            NA
                        NΑ
78
            NΑ
                        NΑ
```

7.2 Extract splice-junction based views from ensemblExons

From ensemblexons information about adjacency of exons (which defines annotated splice-sites) can be obtained by putting exons with equal transcript_id and subsequent exon_number side by side.

The start and end positions of adjacent exons are renamed to lstart, lend and rstart and rend. The "l" prefix refers to the exon with lower start and end coordinates (i.e. left, lower exon_number). The "r" prefix refers to the exons with higher start and end coordinates (i.e. right, higher exon_number).

Setting coding=TRUE will restrict the result to exons for which source and gene_biotype equal "protein_coding".

```
> jens <- getSpliceTable(ens)</pre>
```

[getSpliceTable.refGenome] Finished.

> jens

```
5 5 GL000213.1 121073 121143 126648 126718 ENSG00000237375
6 6 GL000213.1 126648 126718 129228 129365 ENSG00000237375
   gene_name strand
                    transcript_id lexid rexid
1 BX072566.1
                 - ENST00000327822
                                      112
2 BX072566.1
                 - ENST00000327822
                                      115
                                            117
3 BX072566.1
                 - ENST00000327822
                                     117
                                            119
4 BX072566.1
                  - ENST00000327822
                                     119
                                            121
5 BX072566.1
                  - ENST00000327822
                                     121
                                            123
6 BX072566.1
                  - ENST00000327822
                                     123
                                            125
       transcript_biotype
1 nonsense_mediated_decay
2 nonsense_mediated_decay
3 nonsense_mediated_decay
4 nonsense_mediated_decay
5 nonsense_mediated_decay
6 nonsense_mediated_decay
> juc <- getSpliceTable(uc)</pre>
[getSpliceTable.refGenome] Finished.
> juc
Object of class 'ucscJunctions' with 4 rows and 12 columns.
                                        gene_id gene_name
  id segid 1start lend rstart rend
 1 chr1 11874 12227 12613 12721 uc001aaa.3
                                                 DDX11L1
2 2 chr1 12613 12721 13221 14409 uc001aaa.3
                                                  DDX11L1
3 3 chr1 11874 12227 12646 12697 uc010nxr.1
                                                  DDX11L1
4 4 chr1 12646 12697 13221 14409 uc010nxr.1
                                                 DDX11I.1
  strand transcript_id lexid rexid
1
       +
            uc001aaa.3
                         1
2
            uc001aaa.3
                           2
                                 3
3
       +
            uc010nxr.1
                           4
                                 5
                                 6
            uc010nxr.1
                           5
```

This generally leads to repeated occurrence of start and and positons when a splice-junction is contained in multiple transcripts. Additionally a handful splice-sites with multiple gene-id's are present.

The unifyJunc therefore calculates nGenes which represents the multiplicity of each gene-id at each splice-site and then selects a gene-id for which nGenes is maximal.

unifyJuncs adds a uid column to the basic gtf table which identifies each seqid, left-end, right-start combination uniquely. unifyJuncs also adds a new ujs table inside the contained environment.

 ${\tt getUnifiedJuncs}$ takes the result of unifyJuncs and adds gene_name and strand information.

```
> ujens <- unifyJuncs(jens)
> ujuc <- unifyJuncs(juc)
> jeg <- getGenePositions(jens)
> jug <- getGenePositions(juc)
> ujens
```

```
Object of class 'ensemblJunctions' with 51 rows and 12 columns.
  id seqid lstart lend rstart rend nSites
                                                     gene_id
            12010 12057
                        12179 12227
                                           1 ENSG00000223972
            11874 12227
                         12595 12721
                                           1 ENSG00000253101
3
                         12613 12721
                                           3 ENSG00000223972
            11869 12227
            12613 12697
                         12975 13052
                                           1 ENSG00000223972
5
  5
         1
            12613 12721
                        13221 14409
                                           1 ENSG00000223972
6
            12613 12721 13225 14412
  6
         1
                                           1 ENSG00000249291
  strand fexid cnNmd
                      gene_name
1
            41
                   1
                        DDX11L1
2
            64
                   0
                       DDX11L11
3
            42
                   2
                        DDX11L1
4
            43
                   1
                        DDX11L1
5
            47
                   1
                        DDX11L1
6
            63
                   0 AL627309.2
> jug
        gene_id gene_name seqid start
                                         end strand
  1 uc001aaa.3
                  DDX11L1 chr1 11874 14409
  start_codon stop_codon
```

The result tables of unifyJuncs and getGenePositions are stored inside the internal environment of ensemblJunctions. From there, the results can easily be reproduced without recalculation. The tables are automatically included in saveGenome and load.ensembl.juncs mechanisms.

8 Overlapping

The overlap function is used to supply annotation for genomic ranges. The function takes two data.frame's which contain query (qry) and reference (ref) ranges respectively. Each dataset will be identified by it's id.

The routine assumes that query and reference tables are ascending sorted by column 'start'. Otherwise the result will be incorrect (i.e. missing hits). The function assumes that there is no overlap between reference ranges. It will otherwise return only one, possibly arbitrary, hit per query range.

The function returns a data.frame. For each query range, there will be one row.

	overlap	leftDiff	rightDiff	queryid	refid
0	no	0	5	1	0
1	1	2	3	2	1
2	n	1	2	3	3
3	b	2	2	4	4
4	r	2	5	5	4
5	no	5	0	6	0

The query and reference record are identified by "queryid" and "refid". The type of overlap is encoded in the "overlap" column. The overlap encodings are explained as follows:

- no. The query range does not overlap with any reference ranges.
- 1 The query range overhangs the matching reference range on the left (lower coordinate) side.
- **n** The query range is completely contained within a reference range. There is no overhang.
- **b** The query range overhangs the matching reference range on both sides.
- r The query range overhangs the matching reference range on the right (higher coordinate) side.

The added "leftDiff" and "rightDiff" columns contain the distance between the query and reference range boundaries: leftDiff is the difference between the left (lower coordinate) margins and rightDiff is the difference between the right (higher coordinate) margins.

8.1 Overlapping for splice-junctions

The overlapJuncs function is specialized for annotation of splice events in BAM alignment data. The function takes two arguments: a data.frame providing query data and a refJunctions object providing annotation data.

8.1.1 Query data

Query data for overlapJuncs are BAM alignment gap-sites. These gap-sites are defined by one or multiplie gapped alignments of RNA-seq reads to a reference genome. Due to the splicing process, a variable fraction of the reads align in two or more fractions to the reference genome. Two adjacent alignment fractions define the position of a gap-site.

The position of the alignment fractions are named **lstart** and **lend** for the left (i.e. lower genomic coordinates) alignment interval and **rstart** and **rend** for the right (i.e. higher genomic coordinates) alignment interval.

The two genomic intervals of a gap-site implicitly define a third interval: The gap in between (alignment gap).

The biological meaning of gap-sites is that alignment gaps correspond to introns and the left and right align regions correspond to adjacent exon boundaries.

Due to the fact that gapped alignments do not cover whole exons, the inner boundaries of gap-sites are biological meaningful and the outer boundaries of gap-sites usually are technical artifacts. Therefore the overlapJuncs function searches and rates overlaps by inner gap boundaries (lend, rstart) whereas the outer boundaries solely define whether any overlap with annotated splice-sites is present.

The query data.frame defines one gap-site in each row. Therefore the table is expected to contain the following columns:

- id: Consecutive (unique) integral values.
- **seqid**: Character values for reference sequence itentifiers (e.g. 'chr1' or '1').
- lstart: Position of first nucleotide of left alignment region
- lend: Position of last nucleotide of left alignment region.
- rstart: Position of first nucleotide of right alignment region.
- rend: Position of last nucleotide of right alignment region.

All genomic positions are 1-based (i.e. first position of sequence has position 1).

```
> # + + + + + + + + + + + + + + + + + #
 # A) Example query data
 #++++++#
                            1
                                    2
                                            3
                                                           5
                                                                   6
                                                                           7 ##
 qry <- data.frame(id = 1:7, seqid = "1",</pre>
             lstart = c(10100L, 11800L, 12220L, 12220L, 12220L, 32000L, 40000L),
                      c(10100L, 12000L, 12225L, 12227L, 12227L, 32100L, 40100L),
             rstart = c(10200L, 12200L, 12057L, 12613L, 12650L, 32200L, 40200L),
                      c(10300L, 12250L, 12179L, 12620L, 12700L, 32300L, 40300L))
> ##
                            1
                                    2
                                            3
                                                           5
                                                                           7 ##
```

8.1.2 Reference data

The reference data for the overlapJuncs function is can be obtained from a refGenome object with the getSpliceJuncs function.

[getSpliceTable.refGenome] Finished.

8.1.3 Overlapping results

Junction based annotation of annotation gap-sites is done by the overlapJuncs function.

```
> res <- overlapJuncs(qry, junc)</pre>
```

The overlapJuncs function returns a data.frame. As an overlap result is returned for every query record, the output and the query data.frame contain the same number of rows.

The first two columns (qid and refid) provide the query id (the id from the qry data.frame) and a refid (the id for the identified optimal overlap). Records without overlap can be identified withe a refid value of 0.

The validity of the overlap is defined by the distance of the inner boundaries between the query and the reference site. The distance for the left alignment region is given in ldiff and the distance for the right alignment region is given in rdiff. The sum of the absolute ldiff and rdiff values is given by sod (sum of distances). The optimal hit is defined as the one with the lowest sod value. An exact hit will have sod=0.

The result table provides gene_id, transcript_id, gene_name and strand.

9 Workflows

9.1 Establish a standard set of refGenome objects for Ensembl

The following example assumes, that a downloaded *GTF* file has been downloaded and unzipped into a target location (endir):

```
> library(refGenome)
> endir <- "/.../refGenomes/hsEns76"</pre>
> # Read GTF
> en76 <- ensemblGenome()</pre>
> basedir(en76) <- endir
> read.gtf(en76, "Homo_sapiens.GRCh38.76.gtf")
> saveGenome(en76, "en76.RData")
> # Extract primary assembly
> enpa76 <- extractSeqids(en76, ensPrimAssembly())</pre>
> saveGenome(enpa76, "enpa76.RData")
> # Extract Exons
> enex76 <- refExons(enpa76)
> saveGenome(enex76, "enex76.RData")
> # Extract Junctions
> enjc76 <- getSpliceTable(enpa76)</pre>
> saveGenome(enjc76, "enjc76.RData")
> # Extract data.frame
```