Working with the ShortRead Package

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In this document we show how to use the *Genominator* package with the *ShortRead* package and conduct a simple differential expression analysis.

1 Importing data

- > require(Genominator)
 > require(ShortRead)
- > require(yeastRNASeq)

The data we will use are described in the vignette of the yeastRNASeq and was originally published in [1]. But to summarize, we have data from two different yeasts, a wild-type strain ("wt") and a mutant strain ("mut"). Part of the RNA degradation pathway is knocked out in the mutant. For each strain we have two lanes worth of data – it is the exact same sample and library preparation sequenced in both lanes. Only 500,000 raw reads from each lane is part of the yeastRNASeq package and they have been aligned with Bowtie, keeping only unique hits with up to two mismatches. Since not all reads align, we will be working with a up to 500,000 reads per lane.

The data is available as a list of AlignedRead class objects:

```
> data(yeastAligned)
> yeastAligned[[1]]
class: AlignedRead
length: 423318 reads; width: 26 cycles
chromosome: Scchr05 Scchr15 ... Scchr08 Scchr13
position: 541317 885627 ... 488228 667296
strand: - + ... - +
alignQuality: NumericQuality
alignData varLabels: similar mismatch
> sapply(yeastAligned, length)
mut_1_f mut_2_f wt_1_f
                         wt_2_f
423318 420848 410349 430264
  In the package we also have the Bowtie output files, which we will use for illustration:
> list.files(file.path(system.file(package = "yeastRNASeq"), "reads"),
             pattern = "bowtie")
[1] "mut_1_f.bowtie.gz" "mut_2_f.bowtie.gz" "wt_1_f.bowtie.gz"
[4] "wt_2_f.bowtie.gz"
```

The Bowtie output files are compressed using gzip, ShortRead can handle this.

Before importing a set of aligned reads using *Genominator* one usually has to make some decisions regarding chromosome names as well as names of the columns in the resulting database.

Note the chromosome names are a bit special: Scchr01. In our experience it is quite common for different annotation sources to use different shorthands for the chromosomes (for yeast we have at least seen chr1, chr01, Scchr01, chr1). This is a bit of a pain and is very hard to automate. In *Genominator* we expect the user to explicitly convert the chromosome names to a common representation and we furthermore require chromosome names to be integers. For the sake of importing alignment files this is accomplished by using the chrMap argument that is a simple character vector which the chromosome names are matched up against. The first element in the vector gets assigned the integer 1 and so on. If there are chromosomes in the alignment file that does not appear in the chrMap vector, the corresponding reads are not imported. This is often useful, but can also lead to loss of data.

We construct the chrMap object by

```
> chrMap <- paste("Scchr", sprintf("%02d", 1:16), sep = "")
> unique(chromosome((yeastAligned[[1]])))

[1] Scchr05 Scchr15 Scchr13 Scchr16 Scchr01 Scchr14 Scchr12 Scchr03 Scchr07
[10] Scchr11 Scmito Scchr10 Scchr08 Scchr04 Scchr09 Scchr06 Scchr02
17 Levels: Scchr01 Scchr02 Scchr03 Scchr04 Scchr05 Scchr06 Scchr07 ... Scmito
```

We see that by using this version of chrMap we will drop reads aligning to the mitochondrial chromosome.

The other decision we need to make is how to map filenames to names of the columns of the resulting

The other decision we need to make is how to map filenames to names of the columns of the resulting database. Note that whatever choice we make is more or less permanent. Usually we prefer short, descriptive names. It makes a lot of sense to construct these names programmatic from the filenames:

The way this is all specific is through a named vector of filenames, the names of the vector corresponds to column names in the resulting database. If two (or more) file entries have the same name, they will be joined into one column.

We now import the alignment files using importFromAlignedReads, which uses the readAligned function from *ShortRead* to parse the files.

```
chr location strand mut_1 mut_2 wt_1 wt_2
    1
            3888
                        1
                               1
                                     NΑ
                                           NΑ
                                                  NΑ
1
2
    1
            3970
                              NA
                                     NA
                                             1
                                                  NA
3
    1
            3988
                              NA
                                                 NA
                        1
                                      1
                                           NA
4
    1
            4126
                       -1
                              NA
                                     NΑ
                                           NΑ
                                                   1
5
    1
            4242
                        1
                               1
                                     NA
                                           NA
                                                  NA
    1
            4296
                       -1
                               1
                                     NA
                                           NΑ
                                                  NA
```

This will create a database my.db in the current directory. The reads are automatically associated to the genomic location corresponding to the 5' end of the read – this is slightly different from some other programs that uses the lefternmost location of the read (the difference is in reads mapping to the reverse strand).

2 Annotation

There are at least two easy ways to retrieve annotation using Bioconductor: the *biomaRt* package which accesses Ensembl and the *rtracklayer* package which accesses the UCSC genome browser. There are oftentimes species-specific databases (for yeast we have at least SGD) which is a third source of annotation.

Using biomaRt

We will use biomaRt to retrieve information from the Ensembl database. We will illustrate a few pitfalls by comparing two different, but similar looking queries. More information on using biomaRt can be found it its excellent vignette.

The following code was run in January 2010.

```
ensembl_gene_id chromosome_name start_position end_position strand
1 YHR055C VIII 214535 214720 -1
2 YPR161C XVI 864445 866418 -1
gene_biotype
```

1 protein_coding

2 protein_coding

> head(ensembl.transcript, n = 2)

```
ensembl_gene_id ensembl_transcript_id ensembl_exon_id chromosome_name

1 YHR055C YHR055C YHR055C.1 VIII

2 YPR161C YPR161C YPR161C.1 XVI

start_position end_position strand gene_biotype exon_chrom_start
```

```
1
          214535
                        214720
                                   -1 protein_coding
                                                                 214535
2
          864445
                                   -1 protein_coding
                                                                 864445
                        866418
  exon_chrom_end rank
          214720
                     1
1
          866418
                     1
> dim(ensembl.gene)
[1] 7124
            6
> dim(ensembl.transcript)
[1] 7547
           11
> subset(ensembl.gene, ensembl_gene_id == "YPR098C")
  ensembl_gene_id chromosome_name start_position end_position strand
                                            728945
          YPR098C
                               XVI
                                                         729526
                                                                     -1
    gene_biotype
7 protein_coding
> subset(ensembl.transcript, ensembl_gene_id == "YPR098C")
  ensembl_gene_id ensembl_transcript_id ensembl_exon_id chromosome_name
7
          YPR098C
                                 YPR098C
                                                YPR098C.1
                                                                       XVI
          YPR098C
                                 YPR098C
                                                YPR098C.2
                                                                       XVI
8
  start_position end_position strand
                                         gene_biotype exon_chrom_start
7
          728945
                        729526
                                   -1 protein_coding
                                                                 729480
          728945
                        729526
                                   -1 protein_coding
                                                                 728945
8
  exon_chrom_end rank
7
          729526
                     1
8
          729383
> length(unique(ensembl.transcript$ensembl_transcript_id))
```

[1] 7124

Note that altering the query a bit leads to a quite different number of rows being retrieved. In this case it is because each row in one query correspond to a gene, whereas each row in the other query correspon to an exon of a transcript.

A key observation here is that the columns start_position and end_position contains the start and end position of the gene which is different from the start and end position of the exon when we look at a gene consiting of two exons. Furthermore, it is not even clear from the output of the first query that genes with multiple exons exists. In this example, there is no difference between the transcript id and the gene id, because (at least according to this annotation), there are no genes in yeast that produces multiple transcripts.

In this case, we would argue that the right object to use is yeastAnno.transcripts. Below we will post-process this object for use with *Genominator*.

Using rtracklayer

Here we will use the rtracklayer package to access the UCSC genome browser. UCSC sometimes have different tables for a specific genome. We will take a closer look at the SGD table (based on the name we presume that it is supposed to package information from SGD (= Saccharomyces Genome Database)) and the ENS table (which we assume is an attempt to package information from Ensembl)

The following code was run in January 2010

```
> require(rtracklayer)
> session <- browserSession()</pre>
> genome(session) <- "sacCer2"</pre>
> ucsc.sgdGene <- getTable(ucscTableQuery(session, "sgdGene"))
> ucsc.ensGene <- getTable(ucscTableQuery(session, "ensGene"))
  We will also examine the output from these two tables
> data(yeastAnno.sources)
> ucsc.sgdGene <- yeastAnno.sources$ucsc.sgdGene
> ucsc.ensGene <- yeastAnno.sources$ucsc.ensGene
> head(ucsc.sgdGene, n = 2)
        name chrom strand txStart txEnd cdsStart cdsEnd exonCount exonStarts
                    + 130801 131986 130801 131986
1 73 YAL012W chrI
                                                                       130801.
2 585 YAL069W chrI
                              334
                                      649
                                               334
                                                      649
                                                                          334.
  exonEnds proteinID
1 131986,
              P31373
      649,
                n/a
> head(ucsc.ensGene, n = 2)
        name chrom strand txStart txEnd cdsStart cdsEnd exonCount exonStarts
  bin
1 73 YAL012W chrI
                        + 130801 131986
                                            130801 131986
                                                                       130801,
2 585 YAL069W chrI
                              334
                                      649
                                               334
                                                      649
                                                                  1
                                                                          334,
  exonEnds score name2 cdsStartStat cdsEndStat exonFrames
              O YALO12W
1 131986,
                                cmpl
                                            cmpl
      649,
              O YALO69W
                                 cmpl
                                            cmpl
                                                         0,
> dim(ucsc.sgdGene)
[1] 6717
          12
> dim(ucsc.ensGene)
[1] 7124
          16
> subset(ucsc.sgdGene, name == "YPR098C")
           name chrom strand txStart txEnd cdsStart cdsEnd exonCount
5756 590 YPR098C chrXVI
                            - 728944 729526
                                              728944 729526
         exonStarts
                          exonEnds proteinID
5756 728944,729479, 729383,729526,
> subset(ucsc.ensGene, name == "YPR098C")
            name chrom strand txStart txEnd cdsStart cdsEnd exonCount
6104 590 YPR098C chrXVI
                            - 728944 729526
                                               728944 729526
         exonStarts
                          exonEnds score name2 cdsStartStat cdsEndStat
6104 728944,729479, 729383,729526,
                                    0 YPR098C
                                                         cmpl
                                                                    cmpl
     exonFrames
6104
          2,0,
> subset(ucsc.sgdGene, name == "YER102W")
          name chrom strand txStart txEnd cdsStart cdsEnd exonCount
                          + 362728 363698
373 587 YER102W chrV
                                              363095 363698
        exonStarts
                         exonEnds proteinID
373 362728,363095, 363088,363698,
                                    P05754
```

```
> subset(ucsc.ensGene, name == "YER102W")
bin name chrom strand txStart txEnd cdsStart cdsEnd exonCount
424 587 YER102W chrV + 363095 363698 363095 363698 1
    exonStarts exonEnds score name2 cdsStartStat cdsEndStat exonFrames
424 363095, 363698, 0 YER102W cmpl cmpl 0,
```

From this we see that the two tables have quite a different number of genes in them, that the two tables look very similar on the two exon gene considered in the previous section (although one table has the additional information of a protein ID), but that the two tables differ on at least one gene (actually, there are 757 entries that have same value in the name column but different values in the exonStarts column).

Some comments on annotation

As we see here, there are different sources of annotation that differ, even for a relatively simple and well-studied species as *S. cerevisiae*. We cannot give any recommendation as to what annotation source to use, that depends on the biological question and possibly other factors.

However, some effort ought to be spend on making sure that the used annotation matches up with the genome used. Even for yeast there are several different genomes. And while they differ in only a few substitutions and insertion/deletions, the insertion/deletions can easily lead to "off-by-a-little" errors.

Finally we note that for this specific example, none of the queries above display the SGD classification of genes into "verified", "dubious", and "uncharacterized", a classification that is often important when analyzing ones results. This information is obtainable directly from SGD and perhaps from a better use of the annotation tools above.

Post-processing the annotation

In the following we will work with the ensembl.transcript object, which we now post-process for use with *Genominator*.

In Genominator an annotation object is a data-frame with columns chr, start, end, strand where chr is an integer (and should match up with whatever was used in the import step earlier) and strand has values in $\{-1,1,0\}$ with 0 indicating that there is no strand information (and hence 0 matches both 1 and -1).

```
> yAnno <- yeastAnno.sources$ensembl.transcript
> yAnno$chr <- match(yAnno$chr, c(as.character(as.roman(1:16)), "MT", "2-micron"))
> yAnno$start <- yAnno$start_position
> yAnno$end <- yAnno$end_position
> rownames(yAnno) <- yAnno$ensembl_exon_id
> yAnno.simple <- yAnno[yAnno$chr %in% 1:16, c("chr", "start", "end", "strand")]
> head(yAnno.simple, n = 2)
          chr start
                        end strand
YHR055C.1
            8 214535 214720
YPR161C.1 16 864445 866418
                                 _1
> head(yAnno, n = 2)
          ensembl_gene_id ensembl_transcript_id ensembl_exon_id chromosome_name
YHR055C.1
                  YHR055C
                                         YHR055C
                                                       YHR055C.1
                                                                             VIII
YPR161C.1
                  YPR161C
                                         YPR161C
                                                       YPR161C.1
                                                                              XVI
                                                gene_biotype exon_chrom_start
          start_position end_position strand
YHR055C.1
                  214535
                               214720
                                           -1 protein_coding
                                                                        214535
YPR161C.1
                  864445
                               866418
                                           -1 protein_coding
                                                                        864445
          exon_chrom_end rank chr
                                  start
                                             end
YHR055C.1
                  214720
                            1
                                8 214535 214720
YPR161C.1
                               16 864445 866418
                  866418
                            1
```

(note that we remove all annotation on the mitochondria and the plasmid).

A useful function is validAnnotation that checks whether the produced data.frame satisfy the annotation assumptions

> validAnnotation(yAnno)

[1] TRUE

NME1

Q0010

182

0

170

3 Gene level counts

It is easy to obtain region level counts for a given annotation object:

```
> geneCounts.1 <- summarizeByAnnotation(eData, yAnno, ignoreStrand = TRUE)
> head(geneCounts.1)
```

```
mut_1 mut_2 wt_1 wt_2
YHR055C.1
               0
                      0
                            0
YPR161C.1
              38
                     39
                           35
                                 34
YOL138C.1
              31
                     34
                           40
                                 27
YDR395W.1
              54
                     54
                           47
                                 46
YGR129W.1
              28
                     26
                            4
                                  5
                    177
                          150
YPR165W.1
             184
```

This produces a matrix containing the read counts per gene. Such a matrix is ready for analysis by various packages as well as the functionality in *Genominator*.

We use ignoreStrand = TRUE because the experimental assay does not keep strand of origin, so we count reads on either strand, inside each atomic region.

Note that the resulting matrix has one row for each row in the annotation object. In this annotation object, rows correspond to exons. We can get gene level counts either by using a tapply or directly from Genominator by

```
> geneCounts.2 <- summarizeByAnnotation(eData, yAnno, ignoreStrand = TRUE,
                                          groupBy = "ensembl_gene_id")
> head(geneCounts.2)
         mut_1 mut_2 wt_1 wt_2
15S_rRNA
             0
                    0
                         0
                              0
                    0
21S_rRNA
             0
                              0
             7
                             12
HRA1
                   14
LSR1
           389
                  402
                        50
                              60
```

Note that – because of the way reads are represented in the database when we use importFromAligne–dReads – that a read is counting as part of a gene if the first sequenced base maps within the region defined by the gene. This may create some concern at gene boundaries.

There are further complications. In yeast, it is very common for two genes to overlap each other on opposite strands. In other organisms, a gene may have several transcripts. For this purpose, *Genominator* supports the computation of various gene models.

Union-intersection (UI) genes were introduced by []. The UI representation of a gene is the set of bases that are annotated as being part of every transcript of the gene and that are not part of any transcript of any other gene.

We may transform our annotation object into a UI representation by

7

6

```
end strand ensembl_gene_id
  chr
       start
       99699
    1
              99869
                           1
                                         HRA1
1
2
    1 147596 151168
                          -1
                                      YALO01C
3
      143709 147533
                                      YALOO2W
                           1
4
      142176 142368
                                      YAL003W
                                      YAL003W
5
    1 142471 143162
                           1
                                      YAL005C
    1 139505 140761
                          -1
```

In this step we loose all of the additional columns of the yAnno object.

4 Statistical Analysis

We can see how "good" the replicates are by assessing whether it fits the Poisson model of constant gene expression across lanes with variable sequencing effort.

```
> groups <- gsub("_[0-9]_f", "", colnames(geneCounts))
> groups
> plot(regionGoodnessOfFit(geneCounts, groups), chisq = TRUE)
```

5 Working with Priming Weights

In a recent publication [3] we describe how the use of random priming for Illumina RNA-Seq impacts the nucleotide content of the reads and we describe a method for alleviating this bias.

The method associates a weight with each read and instead of counting the number of reads in a given genomic interval, the weights in the interval are summed. Because of this, the use of weights happen at the data import step.

Since our example data was generated using random priming, we illustrate the methodology. The first step is to compute the weights using the function computePrimingWeights and an AlignedRead object. Next, the weights are associated with each read using addPrimingWeights. Once reads have an associated weight, the importFromAlignedReads function uses these. Because of the need to compute the weights, for now, it is not possible to have importFromAlignedReads work directly on filenames.

We start with the yeastAligned object which was simply a list of *AlignedRead*, generated using an lapply on a vector of filenames.

In this case, we have around 410.000-430.000 reads per sample, with each read having a length of only 26 bases. In order to compute the priming weights we need to assess the k-mer distribution at the end of the reads. In [3], the end is based on reads having at least 35 bases, so we need to modify this. We also makes the weights a bit shorter (as described in Hansen et al., this does not change the effect much)

```
> weightsList <- lapply(yeastAligned, computePrimingWeights,
                         unbiasedIndex = 20:21, weightsLength = 6L)
> sapply(weightsList, summary)
        mut_1_f mut_2_f
                         wt_1_f
                                  wt_2_f
Min.
         0.1378
                 0.1415
                          0.1725
                                  0.2048
1st Qu.
         0.7466
                 0.7424
                          0.7592
                                  0.7580
Median
         1.1710
                 1.1590
                          1.1760
                                  1.1670
         1.6160
                 1.6160
                          1.6160
Mean
                                  1.6030
         1.9050 1.9220
                         1.9350
3rd Qu.
                                  1.9220
Max.
        26.6700 24.4000 18.4300 22.6700
```

We will continue with a separate set of weights for each lane.

```
> yeastAligned2 <- mapply(addPrimingWeights, yeastAligned, weightsList)
> alignData(yeastAligned2[[1]])
```

```
An object of class 'AlignedDataFrame'
  readName: 1 2 ... 423318 (423318 total)
  varLabels: similar mismatch weights
  varMetadata: labelDescription
> head(alignData(yeastAligned2[[1]])$weights)
[1] 0.3824834 0.3925501 0.7834928 0.8106667 1.0075188 1.1889401
   Now the weights have been added the AlignedRead objects. Once this has happened, importFromA-
lignedReads will use them.
> eData2 <- importFromAlignedReads(yeastAligned2, chrMap = chrMap,</pre>
                                    dbFilename = "my.db", tablename = "weights")
   We can now easily get the re-weighted gene level counts as usual.
> reweightedCounts <- summarizeByAnnotation(eData2, yAnno, ignoreStrand = TRUE,
                                             groupBy = "ensembl_gene_id")
> head(reweightedCounts)
           mut_1_f
                      mut_2_f
                                  wt_1_f
                                            wt_2_f
           0.00000
                                0.000000
15S_rRNA
                     0.000000
                                          0.000000
21S_rRNA
           0.00000
                     0.000000
                                0.000000 0.000000
HRA1
                     9.707854 2.012202 12.249379
           5.04497
LSR1
         333.17849 367.034724 47.275727 50.420965
NME1
         184.64176 174.321823
                               5.525353
                                          6.791041
Q0010
           0.00000
                     0.000000 0.000000 0.000000
```

SessionInfo

- R version 3.2.0 (2015-04-16), x86_64-unknown-linux-gnu
- Locale: LC_CTYPE=en_US.UTF-8, LC_NUMERIC=C, LC_TIME=en_US.UTF-8, LC_COLLATE=C, LC_MONETARY=en_US.UTF-8, LC_MESSAGES=en_US.UTF-8, LC_PAPER=en_US.UTF-8, LC_NAME=C, LC_ADDRESS=C, LC_TELEPHONE=C, LC_MEASUREMENT=en_US.UTF-8, LC_IDENTIFICATION=C
- Base packages: base, datasets, grDevices, graphics, grid, methods, parallel, stats, stats4, utils
- Other packages: BiocGenerics 0.14.0, BiocParallel 1.2.0, Biostrings 2.36.0, DBI 0.3.1, GenomeGraphs 1.28.0, GenomeInfoDb 1.4.0, GenomicAlignments 1.4.0, GenomicRanges 1.20.0, Genominator 1.22.0, IRanges 2.2.0, RSQLite 1.0.0, Rsamtools 1.20.0, S4Vectors 0.6.0, ShortRead 1.26.0, XVector 0.8.0, biomaRt 2.24.0, yeastRNASeq 0.5.1
- Loaded via a namespace (and not attached): AnnotationDbi 1.30.0, Biobase 2.28.0, RColorBrewer 1.1-2, RCurl 1.95-4.5, XML 3.98-1.1, bitops 1.0-6, futile.logger 1.4, futile.options 1.0.0, hwriter 1.3.2, lambda.r 1.1.7, lattice 0.20-31, latticeExtra 0.6-26, tools 3.2.0, zlibbioc 1.14.0

References

- [1] Lee, A., Hansen, K.D., Bullard, J., Dudoit, S. and Sherlock, G. (2008) Novel Low Abundance and Transient RNAs in Yeast Revealed by Tiling Microarrays and Ultra High-Throughput Sequencing Are Not Conserved Across Closely Related Yeast Species. *PLoS Genet*, 4e1000299.
- [2] Bullard, J.H., Purdom, E.A., Hansen, K.D., and Dudoit, S. (2010) Evaluation of statistical methods for normalization and differential expression in mRNA-Seq experiments. *BMC Bioinformatics*, **11**, 94.
- [3] Hansen, K.D., Brenner, S.E. and Dudoit, S. (2010) Biases in Illumina transcriptome sequencing caused by random hexamer priming. *Nucleic Acids Res*, doi: 10.1093/nar/gkq224