An overview of yeastRNASeq

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Modified: 15 January 2010. Compiled: October 19, 2016

```
> require(yeastRNASeq)
```

This package contains data from

```
> x <- citation(package = "yeastRNASeq")[[1]]</pre>
> print(x, bibtex = FALSE)
  Albert Lee, Kasper D. Hansen, James Bullard, Sandrine
  Dudoit, Gavin Sherlock (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 4(12): e1000299.
  doi:10.1371/journal.pgen.1000299
A BibTeX entry for LaTeX users is
  @Article{,
    title = {Novel low abundance and transient RNAs in yeast revealed by tiling mi
    author = {Albert Lee and Kasper D. Hansen and James Bullard and Sandrine Dudoi
    year = \{2008\},
    journal = {PLoS Genet},
    volume = \{4\},
    number = \{12\},
    pages = \{e1000299\},\
    url = {http://dx.doi.org/10.1371/journal.pgen.1000299},
    doi = {10.1371/journal.pgen.1000299},
    note = {R package version 0.12.0},
  }
```

which describes some experiments in S. cerevisiae comparing various mutant strains to a wild-type strain. A full BibTex entry can be obtained by

> citation("yeastRNASeq")

The subset of the data which this package contains is more specifically data from a wild-type and a single mutant yeast. For each condition (mutant, wild-type) there is two lanes worth of data, each lane containing a sample of 500,000 raw (unaligned) reads from each of 2 lanes each. Each of the four lanes have been aligned against the yeast genome using Bowtie.

The raw reads are contained in 4 FASTQ files and the Bowtie alignment are contained in 4 Bowtie output files. There are 500,000 reads in each of the FASTQ files and fewer reads in each of the Bowtie files. The filenames are

```
> list.files(file.path(system.file(package = "yeastRNASeq"), "reads"))
```

```
[1] "mut_1_f.bowtie.gz" "mut_1_f.fastq.gz" "mut_2_f.bowtie.gz"
[4] "mut_2_f.fastq.gz" "wt_1_f.bowtie.gz" "wt_1_f.fastq.gz"
[7] "wt_2_f.bowtie.gz" "wt_2_f.fastq.gz"
```

The reads were aligned to the yeast genome obtained from ftp://ftpmips.gsf.de/yeast/sequences (which was the basis for the Bowtie index available at the Bowtie website at the time of alignment).

These files are ready to be parsed by the tools in the ShortRead package. As an example we read the alignment files by

The constructed object aligned is a list with 4 elements. Each element correspond to a lane and is an object of class AlignedRead.

The output from this operation has already been stored as an R object and is accessible by

```
> data(yeastAligned)
> yeastAligned[["mut_1_f"]]
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
```

The percent of aligned reads is

> round(sapply(aligned, length) / 500000, 2)
mut_1_f mut_2_f wt_1_f wt_2_f

0.85 0.84 0.82 0.86

There are two additional objects available in the package, purely for illustrative purposes (do not use them for analysis). The object yeastAnno is annotation obtained from Ensembl using biomaRt and is a data.frame of annotation:

```
> data(yeastAnno)
> dim(yeastAnno)
[1] 7124
            6
> head(yeastAnno, n = 2)
  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
```

> table(yeastAnno\$gene_biotype)

| ncRNA | protein_coding | pseudogene | rRNA |
|-------|----------------|------------|------|
| 9 | 6698 | 21 | 14 |
| snRNA | snoRNA | tRNA | |
| 6 | 77 | 299 | |

The other object is called geneLevelData and is a matrix of counts per gene.

Such a matrix may be constructed from yeastAligned and yeastAnno using either the functionality in the IRanges and ShortRead packages or by using the functionality of the Genominator package (which also contains a vignette describing a simplified differential analysis of this dataset).

Note that the data does not contain any biological replicates. In the original publication this was addressed by also analyzing a set of tiling microarrays.