# Package 'yeastNagalakshmi'

October 10, 2024

### Description

The yeast genome data was retrieved from the sequence read archive, aligned with bwa, and converted to BAM format with samtools.

#### **Details**

Package: yeastNagalakshmi

Type: Package Version: 0.99.0

biocViews: ExperimentData, yeast

License: Artistic-2.0

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The package contains three files in extdata sub-directory. Two of them are pertained to RNA sequencing data in BAM format, and one is a TranscriptDb object of yeast from transcript annotations available at the UCSC Genome Browser.

#### Author(s)

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Maintainer: Biocore Team c/o BioC user list <bioconductor@stat.math.ethz.ch>

#### References

Nagalakshmi et. al., *The transcriptional landscape of the yeast genome defined by RNA sequencing*, Science, 320:1344:1349, June 2008.

#### **Examples**

```
y <- system.file("extdata", package="yeastNagalakshmi")
dir(y)</pre>
```

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