# Package 'gDNAinRNAseqData'

June 12, 2025

Title RNA-seq data with different levels of gDNA contamination
Version 1.9.0
<b>Description</b> Provides access to BAM files generated from RNA-seq data produced with different levels of gDNA contamination. It currently allows one to download a subset of the data published by Li et al., BMC Genomics, 23:554, 2022. This subset of data is formed by BAM files with about 100,000 alignments with three different levels of gDNA contamination.
<b>Depends</b> R (>= $4.3$ )
Imports RCurl, XML, ExperimentHub, BiocGenerics, Rsamtools
Suggests BiocStyle, knitr, rmarkdown
VignetteBuilder knitr
License Artistic-2.0
NeedsCompilation no
Encoding UTF-8
biocViews ExperimentHub, ExperimentData, ExpressionData, SequencingData, RNASeqData, Homo_sapiens_Data
<pre>URL https://github.com/functionalgenomics/gDNAinRNAseqData</pre>
BugReports https://github.com/functionalgenomics/gDNAinRNAseqData
RoxygenNote 7.2.3
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gDNAinRNAseqData-package

RNA-seq data with different levels of gDNA contamination

## **Description**

This package provides access to RNA-seq BAM files containing different levels of genomic DNA (gDNA) contamination.

## Usage

```
LiYu22subsetBAMfiles(path = tempdir(), offline = FALSE)
LiYu22phenoData(bamfiles)
```

#### **Arguments**

path (Default='tempdir()') Filesystem path where to store the BAM files.

offline (Default='FALSE') If there is no internet connection, but the data has been pre-

viously downloaded, setting 'offline=TRUE' allows one to retrive the data from

the ExperimentHub cache.

bamfiles full filesystem paths to where the BAM files were downloaded with 'LiYu22subsetBAMfiles()'.

## Details

Currently, this package allows one to download a subset of the data published in:

Li, X., Zhang, P., and Yu. Y. Gene expressed at low levels raise false discovery rates in RNA samples contaminated with genomic DNA. BMC Genomics, 23:554, 2022.

The subset of the data accessible through this package corresponds to BAM files containing about 100,000 alignments sampled uniformly at random for the RNA-seq experiments produced from total RNA libraries mixed with different concentrations of gDNA, concretely 0% (no contamination), 1% and 10%; see Fig. 2 from Li et al. (2022).

#### Value

'LiYu22subsetBAMfiles()' returns a string character vector of filesystem paths to the downloaded BAM files.

'LiYu22phenoData()' returns a 'data.frame' object with the gDNA contamination levels for the BAM files specified in the 'bamfiles' parameter, according to the publication by Li et al. (2022).

### **Functions**

- LiYu22subsetBAMfiles(): downloads the BAM files from the RNA-seq data through the ExperimentHub, and returns the path in the filesystem where the BAM files are stored.
- LiYu22phenoData(): retrieves phenotypic data from the BAM files downloaded with 'LiYu22subsetBAMfiles()'.

#### References

Li, X., Zhang, P., and Yu. Y. Gene expressed at low levels raise false discovery rates in RNA samples contaminated with genomic DNA. BMC Genomics, 23:554, 2022.

## **Examples**

```
## for LiYu2subsetBAMfiles()
bamfiles <- LiYu22subsetBAMfiles()
bamfiles

## for LiYu22phenoData()
bamfiles <- LiYu22subsetBAMfiles()
LiYu22phenoData(bamfiles)</pre>
```

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