# Package 'NanoporeRNASeq'

July 4, 2024

Type Package

Title Nanopore RNA-Seq Example data

Version 1.14.0

Description The NanoporeRNASeq package contains long read RNA-Seq data generated using Oxford Nanopore Sequencing. The data consists of 6 samples from two human cell lines (K562 and MCF7) that were generated by the SG-NEx project. Each of these cell lines has three replicates, with 1 direct RNA sequencing data and 2 cDNA sequencing data. Reads are aligned to chromosome 22 (Grch38) and stored as bam files. The original data is from the SG-NEx project.

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**Encoding UTF-8** 

LazyData true

**Depends** R(>=4.0.0), ExperimentHub (>=1.15.3)

**Suggests** knitr, bambu, ggbio, BSgenome.Hsapiens.NCBI.GRCh38, circlize, ComplexHeatmap, apeglm, rlang, rmarkdown, GenomicAlignments, Rsamtools

Enhances parallel

**biocViews** ExperimentHub, ExperimentData, RNASeqData, Genome, SequencingData

bugReports https://github.com/GoekeLab/NanoporeRNASeq/issues

URL https://github.com/GoekeLab/NanoporeRNASeq

RoxygenNote 7.1.1

VignetteBuilder knitr

git\_url https://git.bioconductor.org/packages/NanoporeRNASeq

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# Description

Annotation GRangesList prepared from bambu for Granges of human genome (Grch38) chromosome 22 (1:25409234)

# Usage

data("HsChr22BambuAnnotation")

#### **Format**

SummarizedExperiment

NanoporeRNASeq

NanoporeRNASeq package with long-read RNA sequencing data

# **Description**

The NanoporeRNASeq package contains long-read RNA-Seq data generated using Oxford Nanopore Sequencing. The data consists of 6 samples from two human cell lines (K562 and MCF7). Each of these cell lines has three replicates, with 1 direct RNA sequencing data and 2 cDNA sequencing data. Reads are aligned to chromosome 22 (Grch38) and stored as bam files. The original data is from the SG-NEx project. Please see the package vignette for examples and use cases.

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# **Details**

Data objects include:

- SGNexSamples sample information of the bam files
- HsChr22BambuAnnotation annotation GRangesList

For detailed information on usage, see the package vignette, by typing vignette("NanoporeRNASeq"), or the workflow linked to on the first page of the vignette.

#### Author(s)

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SGNexSamples

SG-Nex samples from Nanopore RNA-Seq

# Description

Sample information description for K562 and MCF7 samples from SG-Nex

# Usage

```
data("SGNexSamples")
```

#### **Format**

DataFrame

#### **Details**

SGNexSamples is a DataFrame containing the following information:

- sample\_id sample names of the bam files
- Platform sequencing platform
- cellLine cell line used
- protocol sequencing protocols
- cancer\_type- cancer type of the cell line

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