Package 'NanoporeRNASeq'

December 19, 2024

Type Package

Title Nanopore RNA-Seq Example data

Version 1.16.0

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

 ${\bf URL} \ {\tt https://github.com/GoekeLab/NanoporeRNASeq}$

RoxygenNote 7.1.1 **VignetteBuilder** knitr

git_url https://git.bioconductor.org/packages/NanoporeRNASeq

 $\textbf{git_branch} \ \ RELEASE_3_20$

git_last_commit 8b705c0

git_last_commit_date 2024-10-29

Repository Bioconductor 3.20

Date/Publication 2024-12-19

Author Jonathan Goeke [aut],

Ying Chen [cre], Yuk Kei Wan [aut]

Maintainer Ying Chen <chen_ying@gis.a-star.edu.sg>

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BambuAnnotation of the first half of Human Sapiens Chr22

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.

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)

Ying Chen, Yuk Kei Wan, Jonathan Göke

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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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