# Package 'raerdata'

December 10, 2024

```
Version 1.5.0

Description raerdata is an ExperimentHub package that provides a collection of files useful for demostrating functionality in the raer package. Datasets include 10x genomics scRNA-seq, bulk RNA-seq, and paired whole-genome and RNA-seq data. Additionally databases of human and mouse RNA editing sites are provided.
```

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**Imports** ExperimentHub, Rsamtools, BiocGenerics, rtracklayer, SingleCellExperiment

**Suggests** rmarkdown, knitr, BiocStyle, testthat (>= 3.0.0)

**Title** A collection of datasets for use with raer package

VignetteBuilder knitr

**Roxygen** list(markdown = TRUE)

**RoxygenNote** 7.2.3 **Encoding** UTF-8

**biocViews** SingleCellData, SequencingData, RNASeqData, ExperimentHub, PackageTypeData, ExpressionData

BugReports https://github.com/rnabioco/raerdata/issues

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## **Description**

Study GSE99249 examined RNA editing in WT, ADAR1KO, and ADAR1-p150 HEK293T cells treated with and without interferon beta.

## Usage

GSE99249()

#### **Details**

GSE99249() will download BAM and BAM index files from 6 RNA-seq libraries. 3 libraries are ADAR1 knockout cells treated with interferon beta and 3 libraries are wild type cells treated with interferon beta. The BAM files contain alignments from chromosome 18.

#### Value

A list containing:

- bams A BamFileList object, indicating the BAM file paths and BAI indexes.
- fasta A path to a fasta file from chr18 of hg38
- snps a GRanges object containing known SNPs from the REDIportal database (hg38)

## **Examples**

GSE99249()

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NA12878

Whole genome and RNA sequencing data from NA12878 cell line

## **Description**

Whole genome and RNA sequencing data from NA12878 cell line

#### **Usage**

NA12878()

#### **Details**

Will download BAM and BAM index files from whole genome and RNA sequencing of the NA12878 cell line, The data is from the first megabase of chromosome 4. Additionally a fasta file and a database of known SNPs will be downloaded.

#### Value

A list containing:

- bams A BamFileList object, indicating the BAM file paths and BAI indexes.
- fasta A path to a fasta file containing the genome sequence of the first megabase of chr4 (hg38)
- snps a GRanges object containing SNPs from the first megabase of chr4

#### **Examples**

NA12878()

pbmc\_10x

single cell RNA sequencing data from human PBMCs

#### Description

A 10x Genomics 3' single cell RNA-seq library from 10k PBMCs. The BAM file contains alignments from chr16. A SingleCellExperiment is also provided with pre-processed gene expression data, a UMAP projection and cell type annotations.

## Usage

pbmc\_10x()

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#### **Details**

pbmc\_10x() will download a BAM, BAM index file, REDIportal RNA editing sites, and a Single-CellExperiment object from the ExperimentHub.

#### Value

A list containing:

- bam a BamFile object indicating the BAM and BAI file paths. Contains alignments from only chr16 (hg38).
- sites a GRanges object containing known RNA editing sites from the REDIportal database (hg38).
- sce a <u>SingleCellExperiment</u> object containing gene expression data, a UMAP projection and cell type annotations.

#### See Also

https://www.10xgenomics.com/resources/datasets/10k-human-pbmcs-3-v3-1-chromium-x-with-intronic-rea

### **Examples**

pbmc\_10x()

raerdata

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## **Description**

A collection of datasets and databases to demonstrate RNA-editing analysis approaches using the raer package.

#### Details

atlases a collection of RNA editing databases

NA12878 Whole genome and RNA sequencing data from the NA12878 cell line

GSE99249 RNA sequencing data from a study that examined RNA editing in WT, ADAR1KO, and ADAR1-p150 HEK293T cells treated with and without interferon beta.

pbmc\_10x single cell RNA sequencing data from human PBMCs from 10x Genomics

rediportal\_full\_mm10

rediportal\_full\_mm10 Databases of known RNA editing sites

#### **Description**

Databases of known RNA editing sites

## Usage

```
rediportal_full_mm10()
rediportal_coords_mm10()
rediportal_full_hg38()
rediportal_coords_hg38()
gabay_sites_mm10()
gabay_sites_hg38()
```

#### **Details**

rediportal\_full\_hg38() will download the human REDIportal database for hg38 which has been converted into a GRanges object. The GRanges is supplemented with additional columns of information provided by the REDIportal database, including gene location, repeat type, dbSNP annotation, and potential for amino-acid recoding.

rediportal\_coords\_hg38() will download the human REDIportal database for hg38 which has been converted into a GRanges object, which only contains the coordinates of the editing site.

rediportal\_full\_mm10() will download the mouse REDIportal database for mm10 which has been converted into a GRanges object. The GRanges is supplemented with additional columns of information provided by the REDIportal database, including gene location, repeat type, dbSNP annotation, and potential for amino-acid recoding.

rediportal\_coords\_mm10()will download the mouse REDIportal database for mm10 which has been converted into a GRanges object, which only contains the coordinates of the editing site.

gabay\_sites\_hg38() will download high-confidence human CDS editing sites (hg38).

gabay\_sites\_mm10() will download high-confidence mouse CDS editing sites (lifted-over from hg38 to mm10).

#### Value

A GRanges object.

#### **Examples**

```
gabay_sites_hg38()
```

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