

Package ‘brgedata’

December 10, 2024

Title Exposures, Gene Expression and Methylation data for illustration purposes

Version 1.28.0

Description This package contains several sets of omics data including Gene Expression (ExpressionSet), Methylation (GenomicRatioSet), Proteome and Exposome (ExposomeSet). This data is used in vignettes and examples at MEAL, MultiDataSet and omicRexposome.

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Encoding UTF-8

LazyData true

RoxygenNote 6.0.1

VignetteBuilder knitr

Depends R (>= 3.4), Biobase

Imports SummarizedExperiment

Suggests minfi, MultiAssayExperiment, knitr, rmarkdown, rexposome, BiocStyle

biocViews ExperimentData,Homo_sapiens_Data,MicroarrayData,MethylationArrayData

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Description

brgedata

brge_expo	ExposomeSet <i>for testing purposes</i>
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Description

ExposomeSet with full set of 15 exposures grouped in 3 families for 110 individuals.

Usage

```
data("brge_expo")
```

Format

An object of class ExposomeSet of dimension 15 x 110 x 6.

Value

An ExposomeSet object.

brge_gexp	ExpressionSet <i>for testing purposes</i>
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Description

ExpressionSet with full set of 67528 features from Affymetrix HTA 2.0 for 100 simulated samples.

Usage

```
data("brge_gexp")
```

Format

An object of class ExpressionSet with 67528 rows and 100 columns.

Value

An ExpressionSet object.

Examples

```
data("brge_gexp")
dim(brge_gexp)
sampleNames(brge_gexp)
```

brge_methy

GenomicRatioSet *for testing purposes*

Description

GenomicRatioSet with 476946 features from Illumina 450k methylation array and 115 simulated samples.

Usage

```
data("brge_methy")
```

Format

An object of class GenomicRatioSet with 476946 rows and 20 columns.

Value

An GenomicRatioSet object.

Examples

```
data("brge_methy")
dim(brge_methy)
sampleNames(brge_methy)
```

brge_prot

ExpressionSet *for testing purposes*

Description

ExpressionSet with full set of 47 proteins for 90 simulated samples.

Usage

```
data("brge_prot")
```

Format

An object of class ExpressionSet with 47 rows and 90 columns.

Value

An ExpressionSet object.

Examples

```
data("brge_prot")  
dim(brge_prot)  
sampleNames(brge_prot)
```

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