Package 'shinyMethylData'

September 19, 2024

Version 1.24.0	
Title Example dataset of input data for shinyMethyl	
Description Extracted data from 369 TCGA Head and Neck Cancer DNA methylation samples. The extracted data serve as an example dataset for the package shinyMethyl. Original samples are from 450k methylation arrays, and were obtained from The Cancer Genome Atlas (TCGA). 310 samples are from tumor, 50 are matched normals and 9 are technical replicates of a control cell line.	
Maintainer Jean-Philippe Fortin <jfortin@jhsph.edu></jfortin@jhsph.edu>	
License Artistic-2.0	
Depends R (>= $3.0.0$)	
LazyData yes	
biocViews Genome, CancerData	
Url https://github.com/Jfortin1/shinyMethylData	
Author Jean-Philippe Fortin [cre, aut], Kasper Daniel Hansen [aut]	
git_url https://git.bioconductor.org/packages/shinyMethylData	
git_branch RELEASE_3_19	
git_last_commit 1c95a44	
git_last_commit_date 2024-04-30	
Repository Bioconductor 3.19	
Date/Publication 2024-09-19	
Contents	
• •	2
Index	4

2 summary.tcga.raw

summary.tcga.norm

Example dataset of input data for shinyMethyl

Description

Extracted data from 369 TCGA Head and Neck Cancer DNA methylation samples. The extracted data serve as an example dataset for the package shinyMethyl. Original samples are from 450k methylation arrays, and were obtained from The Cancer Genome Atlas (TCGA). 310 samples are from tumor, 50 are matched normals and 9 are technical replicates of a control cell line.

Usage

```
data(summary.tcga.norm)
```

Format

A list containing the necessary information to launch a shinyMethyl session. See the links below for more details on the data.

References

The Cancer Genome Atltas (TCGA) Head and Neck Cancer dataset: http://cancergenome.nih.gov/cancersselected/headandne

See Also

These data objects were created by See shinySummarize for details on how to perform the data extraction. See runShinyMethyl for how to launch a shinyMethyl session.

Examples

```
data(summary.tcga.norm)
## Not run:
runShinyMethyl(summary.tcga.norm)
## End(Not run)
```

summary.tcga.raw

Example dataset of input data for shinyMethyl

Description

Extracted data from 369 TCGA Head and Neck Cancer DNA methylation samples. The extracted data serve as an example dataset for the package shinyMethyl. Original samples are from 450k methylation arrays, and were obtained from The Cancer Genome Atlas (TCGA). 310 samples are from tumor, 50 are matched normals and 9 are technical replicates of a control cell line.

summary.tcga.raw 3

Usage

```
data(summary.tcga.raw)
```

Format

A list containing the necessary information to launch a shinyMethyl session. See the links below for more details on the data.

References

The Cancer Genome Atltas (TCGA) Head and Neck Cancer dataset: http://cancergenome.nih.gov/cancersselected/headandne

See Also

See shinySummarize for details on how to perform the data extraction. See runShinyMethyl for how to launch a shinyMethyl session.

Examples

```
data(summary.tcga.raw)
## Not run:
runShinyMethyl(summary.tcga.raw)
## End(Not run)
```

Index

```
* datasets

summary.tcga.norm, 2

summary.tcga.raw, 2

runShinyMethyl, 2, 3

shinySummarize, 2, 3

summary.tcga.norm, 2

summary.tcga.raw, 2
```