Package ‘shinyMethyl’

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Title Interactive visualization for Illumina methylation arrays

Version 1.10.0

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Imports RColorBrewer

Suggests shinyMethylData, minfiData, BiocStyle, RUnit, digest, knitr

Description Interactive tool for visualizing Illumina methylation array data. Both the 450k and EPIC array are supported.

Url https://github.com/Jfortin1/shinyMethyl

VignetteBuilder knitr

License Artistic-2.0

biocViews DNAMethylation, Microarray, TwoChannel, Preprocessing, QualityControl

NeedsCompilation no

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shinyMethyl-package

Interactive visualization for Illumina methylation arrays

Description

Interactive tool for visualizing Illumina methylation array data (450k and EPIC arrays).

Author(s)

Jean-Philippe Fortin <jfortin@jhsph.edu>

runShinyMethyl

Run the interactive shinyMethyl session

Description

Function to run the interactive shinyMethyl session from a shinyMethylSet.

Usage

runShinyMethyl(shinyMethylSet1, shinyMethylSet2 = NULL)

Arguments

shinyMethylSet1
shinyMethylSet that must be extracted from an RGChannelSet

shinyMethylSet2
optional shinyMethylSet that must be extracted from an GenomicRatioSet

Value

No value returned. Instead the shinyMethyl interactive session is launched.

Author(s)

Jean-Philippe Fortin

See Also

shinyMethylSet
Examples

## Not run:
if (require(minfiData)){
    baseDir <- system.file("extdata", package = "minfiData")
    targets <- read.450k.sheet(baseDir)
    targets$Sample_Plate <- substr(targets$Slide,1,7)
    RGSet <- read.450k.exp(base = baseDir, targets = targets)

    summarized.data <- shinySummarize(RGSet)
    runShinyMethyl(summarized.data)
}
## End(Not run)

shinyMethylSet-class

shinyMethylSet instances

Description

This class holds summarized data from Illumina methylation microarrays for interactive visualization purpose.

Usage

## Constructor

shinyMethylSet(sampleNames = new("character"),
               phenotype = new("data.frame"),
               mQuantiles = new(vector("list",5)),
               betaQuantiles = new(vector("list",5)),
               methQuantiles = new(vector("list",5)),
               unmethQuantiles = new(vector("list",5)),
               cnQuantiles = new(vector("list",5)),
               greenControls = new(vector("list",12)),
               redControls = new(vector("list",12)),
               pca = new("list"),
               originObject = new("character"),
               array = new("character"))

## Data extraction / Accessors

## S4 method for signature 'shinyMethylSet'
getM(object)
## S4 method for signature 'shinyMethylSet'
getMeth(object)
## S4 method for signature 'shinyMethylSet'
getUnmeth(object)
## S4 method for signature 'shinyMethylSet'
getBeta(object)
## S4 method for signature 'shinyMethylSet'
getM(object)
## S4 method for signature 'shinyMethylSet'
getCN(object)
## S4 method for signature 'shinyMethylSet'
pData(object)
## S4 method for signature 'shinyMethylSet'
sampleNames(object)

### Arguments

- **object**: A shinyMethylSet.
- **sampleNames**: A character vector
- **phenotype**: A data.frame object.
- **methQuantiles**: A list of 5 matrices containing several quantiles for the methylation values (between zero and infinity) separated by probe type and autosomal/sex probes. Each row is a quantile and each column is a sample.
- **unmethQuantiles**: Similar to Meth Quantiles
- **betaQuantiles**: Similar to Meth but for Beta-values
- **mQuantiles**: Similar to Meth but for M-values
- **cnQuantiles**: Similar to Meth but for Copy Number
- **green Controls**: A list containing the matrices of different raw control probes intensities in the green channel
- **red Controls**: Similar to green Controls but for the red channel
- **pca**: List containing the PCA scores for the 20,000 most variable CpGs and the percentages of variance explained
- **originObject**: Name of the minfi object from which the data were extracted
- **array**: Name of the Illumina array platform: 450k or EPIC

### Details

These class is a representation of a Meth matrix and a Unmeth matrix linked to a pData data frame. A creation Method slot is present to indicate from which object type the shinyMethylSet has been created: either a RGChannelSet or GenomicRatioSet

A MethylSet stores meth and Unmeth.

### Constructor

Instances are constructed using the shinyMethylSet function with the arguments outlined above.

### Accessors

- In the following code, object is a shinyMethylSet.
- `getMeth(object), getUnmeth(object)` Get the Meth or Unmeth matrix.
- `getBeta(object)` get Beta, see details.
- `getM(object)` get M-values, see details.
- `getCN(object)` get copy number values which are defined as the sum of the methylation and unmethylation channel.
- `sampleNames(object)` Get the sampleNames
- `combine(object1, object2)` combine shinyMethylSet objects
Author(s)
Jean-Philippe Fortin <jfortin@jhsph.edu>

See Also
Objects of this class are typically created by using the function shinySummarize on a RGChannelSet or GenomicRatioSet.

Examples
showClass("shinyMethylSet")

shinySummarize-methods
Summarizing a large 450K experiment into a shinyMethylSet

Description
Summarizing methylation data from a RGChannelSet or GenomicRatioSet into a shinyMethylSet needed to launch the interactive interface of shinyMethyl. That works for both the 450k and EPIC arrays.

Usage
## S4 method for signature 'RGChannelSet'
shinySummarize(object)
## S4 method for signature 'GenomicRatioSet'
shinySummarize(object)

Arguments
object Either a RGChannelSet or a GenomicRatioSet.

Details
Either a RGChannelSet is transformed into a shinyMethylSet, or a GenomicRatioSet is transformed to a shinyMethylSet

Author(s)
Jean-Philippe Fortin <jfortin@jhsph.edu>

See Also
shinyMethylSet for the output object and RGChannelSet or GenomicRatioSet for the input object.

Examples
if (require(minfiData)){
  library(minfiData)
  summarized.data <- shinySummarize(RGsetEx)
}
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