Package: qsmooth (via r-universe)

June 30, 2024

Title Smooth quantile normalization

Version 1.21.0

Imports SummarizedExperiment, utils, sva, stats, methods, graphics, Hmisc
Depends R (>= 4.0)
Suggests bodymapRat, quantro, knitr, rmarkdown, BiocStyle, testthat
VignetteBuilder knitr
RoxygenNote 7.1.1
Encoding UTF-8
License GPL-3
Description Smooth quantile normalization is a generalization of quantile normalization, which is average of the two types of assumptions about the data generation process: quantile normalization and quantile normalization between groups.
biocViews Normalization, Preprocessing, MultipleComparison, Microarray, Sequencing, RNASeq, BatchEffect
Repository https://bioc.r-universe.dev
RemoteUrl https://github.com/bioc/qsmooth
RemoteRef HEAD
RemoteSha cbbedd71a186bb1c547e170fad5d7e2c877eb187
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Description

This function applies a generalization of quantile normalization called smoothed quantile normalization. This function defines the qsmooth class and constructor.

Usage

```
qsmooth(object, group_factor, batch = NULL, norm_factors = NULL, window = 0.05)
```

Arguments

object	an object which is a matrix or data.frame with observations (e.g. probes or genes) on the rows and samples as the columns. Alternatively, a user can provide a SummarizedExperiment object and the assay(object, "counts") will be used as input for the qsmooth normalization.
group_factor	a group level continuous or categorial covariate associated with each sample or column in the object. The order of the group_factor must match the order of the columns in object.
batch	(Optional) batch covariate (multiple batches are not allowed). If batch covariate is provided, Combat() from sva is used prior to qsmooth normalization to remove batch effects. See Combat() for more details.
norm_factors	optional normalization scaling factors.
window	window size for running median which is a fraction of the number of rows in object. Default is 0.05.

Details

Quantile normalization is one of the most widely used normalization tools for data analysis in genomics. Although it was originally developed for gene expression microarrays it is now used across many different high-throughput applications including RNAseq and ChIPseq. The methodology relies on the assumption that observed changes in the empirical distribution of samples are due to unwanted variability. Because the data is transformed to remove these differences it has the potential to remove interesting biologically driven global variation. Therefore, applying quantile normalization, or other global normalization methods that rely on similar assumptions, may not be an appropriate depending on the type and source of variation.

This function computes a weight at every quantile that compares the variability between groups relative to within groups. In one extreme quantile normalization is applied and in the other extreme quantile normalization within each biological condition is applied. The weight shrinks the group-level quantile normalized data towards the overall reference quantiles if variability between groups is sufficiently smaller than the variability within groups. See the vignette for more details.

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Value

A object of the class qsmooth that contains a numeric vector of the qsmooth weights in the qsmoothWeights slot and a matrix of normalized values after applying smoothed quantile normalization in the qsmoothData slot.

Examples

qsmooth-class

the qsmooth class

Description

Objects of this class store all the values needed information to work with a qsmooth object

Value

qsmoothWeights returns the qsmooth weights and qsmoothData returns the qsmooth normalized data

Slots

```
qsmoothWeights qsmooth weights
qsmoothData qsmooth normalized data
```

Examples

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qsmoothData

Generic function that returns the qsmooth normalized data

Description

Given a qsmooth object, this function returns the qsmooth normalized data Accessors for the 'qsmoothData' slot of a qsmooth object.

Usage

```
qsmoothData(object)
## S4 method for signature 'qsmooth'
qsmoothData(object)
```

Arguments

object

an object of class qsmooth.

Value

The normalized data after applying smoothed quantile normalization.

Examples

qsmoothGC

qsmoothGC

Description

This function applies smoothed quantile normalization separately for groups of features that are binned according to their GC-content.

Usage

```
qsmoothGC(object, group_factor, gc, nGroups = 50, round = TRUE, ...)
```

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Arguments

object an object which is a matrix or data. frame with observations (e.g. probes or genes) on the rows and samples as the columns. Alternatively, a user can provide a SummarizedExperiment object and the assay(object, "counts") will be used as input for the qsmooth normalization. group_factor a group level continuous or categorial covariate associated with each sample or column in the object. The order of the group_factor must match the order of the columns in object. GC-content of the features, ordered according to the features in object. gc nGroups The number of equally-sized bins used to group the GC-content values. Groups are created using Hmisc::cut2. Should normalized values be rounded to integers? round

(Optional) Additional arguments passed to qsmooth.

Value

A matrix of normalized counts.

References

Van den Berge K., Chou H., Roux de Bézieux H., Street K., Risso D., Ngai J., Dudoit S. Normalization benchmark of ATAC-seq datasets shows the importance of accounting for GC-content effects. https://www.biorxiv.org/content/10.1101/2021.01.26.428252v2

Examples

qsmoothPlotWeights

Plot weights from qsmooth *function*.

Description

This function plots a scatterplot showing the qsmoothWeights along the y-axis and the quantiles on the x-axis.

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Usage

```
qsmoothPlotWeights(
  object,
  xLab = "quantiles",
  yLab = "weights",
  mainLab = "qsmooth weights")
```

Arguments

object a qsmooth object from qsmooth
xLab label for x-axis. Default is "quantiles"
yLab label for y-axis. Default is "weights"
mainLab title of plot. Default is "qsmooth weights"

Value

A scatterplot will be created showing the qsmoothWeights along the y-axis and the quantiles on the x-axis.

Examples

qsmoothWeights

Generic function that returns the qsmooth weights

Description

Given a qsmooth object, this function returns the qsmooth weights Accessors for the 'qsmoothWeights' slot of a qsmooth object.

Usage

```
qsmoothWeights(object)

## S4 method for signature 'qsmooth'
qsmoothWeights(object)
```

Arguments

object an object of class qsmooth.

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Value

The weights calculated for each feature after applying smoothed quantile normalization.

Examples

qstats

qstats

Description

This function is a helper function that computes quantile statistics for the function qsmooth.

Usage

```
qstats(object, group_factor, window = 0.05)
```

Arguments

object an object which is a data frame or matrix with observations (e.g. probes or

genes) on the rows and samples as the columns.

group_factor a group level continuous or categorial covariate associated with each sample or

column in the object. The order of the group_factor must match the order of

the columns in object.

window window size for running median which is a fraction of the number of rows in

object. Default is 0.05.

Value

A list of quantile statistics including

Q sample quantiles Qref reference quantile

Qhat linear model fit at each quantile

SST total sum of squares
SSB between sum of squares
SSE within sum of squares

roughWeights SSE/SST

smoothWeights smoothed weights computed using a running median with a given window size.

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Examples

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