Package 'miRcomp'

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- Title Tools to assess and compare miRNA expression estimatation methods
- **Description** Based on a large miRNA dilution study, this package provides tools to read in the raw amplification data and use these data to assess the performance of methods that estimate expression from the amplification curves.
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Imports utils, methods, graphics, KernSmooth, stats

VignetteBuilder knitr

Suggests BiocStyle, knitr, rmarkdown, RUnit, BiocGenerics, shiny

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biocViews Software, qPCR, Preprocessing, QualityControl

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accuracy

Assess the accuracy of the expression estimates

Description

This function assesses the signal detect slope (a measure of accuracy) for each feature.

Usage

```
accuracy(object1, qcThreshold1, object2=NULL, qcThreshold2=NULL,
commonFeatures=TRUE, bins=3, label1=NULL, label2=NULL)
```

Arguments

object1	a list containing two elements: ct (the expression estimates) and qc (quality scores)
qcThreshold1	a numeric threshold corresponding to object1\$qc below which values are con- sidered low quality.
object2	an optional second list of the same format as object1, used to compare two methods.
qcThreshold2	a numeric threshold corresponding to object2\$qc below which values are con- sidered low quality.
commonFeatures	if TRUE and object2 is non-NULL, only high quality non-NA features in com- mon between both objects are used.
bins	the number of bins to divide the data into.
label1	optional label corresponding to object 1 to be used in plotting.
label2	optional label corresponding to object 2 to be used in plotting.

Value

A plot of signal detect slopes stratified by difference in pure sample expression is produced, and a summary (median and MAD) of the signal detect slopes in each bin is returned.

completeFeatures

Author(s)

Matthew N. McCall

Examples

completeFeatures Determine the Number of Completely Observed Features

Description

This function determines the number of features that are good quality and non-NA across all samples using a given quality threshold.

Usage

```
completeFeatures(object1, qcThreshold1, object2=NULL, qcThreshold2=NULL,
label1=NULL, label2=NULL)
```

Arguments

object1	a list containing two elements: ct (the expression estiamtes) and qc (quality scores)
qcThreshold1	a numeric threshold corresponding to object1\$qc below which values are con- sidered low quality.
object2	an optional second list of the same format as object1, used to compare two methods.
qcThreshold2	a numeric threshold corresponding to object2\$qc below which values are con- sidered low quality.
label1	optional label corresponding to object 1 to be used in plotting.
label2	optional label corresponding to object 2 to be used in plotting.

Value

The function generates a table of the number of complete, partial, and absent features.

Author(s)

Matthew N. McCall

Examples

expressionComp

Compare expression estimates between two methods

Description

This function allows one to easily compare the expression estimates generated by two different algorithms. Feature / sample combinations that differ above a given threshold can be easily highlighted.

Usage

```
expressionComp(object1, qcThreshold1,
```

```
object2, qcThreshold2,
label1=NULL, label2=NULL,
diffThreshold=2,
plotType=c("scatterplot","MAplot"))
```

Arguments

object1	a list containing two elements: ct (the expression estimates) and qc (quality scores)
qcThreshold1	a numeric threshold corresponding to object1\$qc below which values are con- sidered low quality.
object2	a list containing two elements: ct (the expression estimates) and qc (quality scores)
qcThreshold2	a numeric threshold corresponding to object2\$qc below which values are con- sidered low quality.
label1	an optional label for plotting on the x-axis
label2	an optional label for plotting on the y-axis
diffThreshold	feature / sample combinations for which the expression estimates differ by more than this value are returned by the function and highlighted in the plot.
plotType	scatterplot or MA-plot

Value

This function plots the expression estimates produced by two different methods. Feature / sample combinations for which the expression estimates differ by more than diffThreshold are returned by the function and highlighted in the plot.

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limitOfDetection

Author(s)

Matthew N. McCall

Examples

```
data(lifetech)
data(qpcRdefault)
tmp <- expressionComp(object1=lifetech, qcThreshold1=1.25,
object2=qpcRdefault,qcThreshold2=0.99,plotType="scatter")</pre>
```

lifetech

The processed data generated using the LifeTech software.

Description

These data are also used to demonstrate the functionality of the miRcomp package.

Usage

data(lifetech)

Format

A list with two elements.

ct a miRNA x sample matrix of expression estimates

qc a miRNA x sample matrix of quality scores (here AmpScores)

Examples

data(lifetech)

limitOfDetection Assess the limit of detection of a given method

Description

This function assesses the limit of detection in one of two ways: (1) the distribution of expression estimates stratified by the proportion of poor quality values within replicates, (2) the average vs expected expression for the two most diluted sample types.

Usage

limitOfDetection(object, qcThreshold, plotType=c("boxplot","scatterplot","MAplot"))

Arguments

object	a list containing two elements: ct (the expression estiamtes) and qc (quality scores)
qcThreshold	a numeric threshold corresponding to object1\$qc below which values are con- sidered low quality.
plotType	the desired output type – boxplot is option (1); scatterplot is option (2), an MA-plot is option (3).

Value

This function assesses the limit of detection in several ways. If plotType is boxplot, then boxplots of expression estimates stratified by the proportion of poor quality values within replicates is displayed. The function also outputs a list with the values plotted in each box of the boxplot. If plotType is scatterplot, then the average within replicate expression vs expected expression (based on pure sample expression) is displayed for the 0.1/0.1 dilution and 0.01/0.01 dilution. If plotType is MAplot, then the difference in expression (average within replicate expression - expected expression) is displayed for the 0.1/0.1 dilution. For both plotTypes, scatterplot and MAplot, the function outputs a matrix containing estimates of the limit of detection for four different tolerances. Specifically, the two columns correspond to the two dilutions (0.1/0.1 and 0.01/0.01) and rows correspond to the median difference between the observed and expected values. The values in the matrix are the expected expression values such that the median absolute difference of all larger expected expression values is approximately equal to the given tolerance.

Author(s)

Matthew N. McCall

Examples

```
data(lifetech)
tmp <- limitOfDetection(object=lifetech,qcThreshold=1.25)
data(qpcRdefault)
limitOfDetection(object=qpcRdefault,qcThreshold=0.99,plotType="scatter")</pre>
```

miRcompShinyApp Launch a Shiny app to use the miRcomp package interactively

Description

This function launches the miRcomp Shiny app locally.

Usage

miRcompShinyApp()

Author(s)

Matthew N. McCall and Lauren Kemperman

precision

Examples

miRcompShinyApp()

precision

Assess the precision of the expression estimates

Description

This function assesses the within-replicate precision for each feature.

Usage

Arguments

object1	a list containing two elements: ct (the expression estiamtes) and qc (quality scores)
qcThreshold1	a numeric threshold corresponding to object1\$qc below which values are con- sidered low quality.
object2	an optional second list of the same format as object1, used to compare two methods.
qcThreshold2	a numeric threshold corresponding to object2\$qc below which values are con- sidered low quality.
commonFeatures	if TRUE and object2 is non-NULL, only high quality non-NA features in com- mon between both objects are used.
statistic	whether to compute the standard deviation (sd) or coefficient of variation (cv).
scale	optional scaling of the values. This can help with visualizing the distributions.
bins	the number of bins to divide the data into.
label1	optional label corresponding to object 1 to be used in plotting.
label2	optional label corresponding to object 2 to be used in plotting.

Value

A boxplot of either the standard deviation or coefficient of variation stratified by expression is produced. The values plotted in each box of the boxplot are returned.

Author(s)

Matthew N. McCall

qpcRb5

Examples

qpcRb4	The processed data generated using the 4 parameter sigmoidal method
	from the qpcR software.

Description

These data are also used to demonstrate the functionality of the miRcomp package.

Usage

data(qpcRb4)

Format

A list with two elements.

ct a miRNA x sample matrix of expression estimates

qc a miRNA x sample matrix of quality scores (here R-squared values)

Examples

data(qpcRb4)

qpcRb5	The processed data generated using the 5 parameter sigmodial method
	from the qpcR software.

Description

These data are also used to demonstrate the functionality of the miRcomp package.

Usage

```
data(qpcRb5)
```

Format

A list with two elements.

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qpcR15

- ct a miRNA x sample matrix of expression estimates
- qc a miRNA x sample matrix of quality scores (here R-squared values)

Examples

data(qpcRb5)

qpcRdefault	The processed data generated using the default method (4 parameter
	log-logistic) implemented in the $qpcR$ software package.

Description

These data are also used to demonstrate the functionality of the miRcomp package.

Usage

data(qpcRdefault)

Format

A list with two elements.

- ct a miRNA x sample matrix of expression estimates
- qc a miRNA x sample matrix of quality scores (here R-squared values)

Examples

data(qpcRdefault)

qpcR15	The processed data generated using the 5 parameter log-logistic
	method from the qpcR software.

Description

These data are also used to demonstrate the functionality of the miRcomp package.

Usage

```
data(qpcR15)
```

Format

A list with two elements.

- ct a miRNA x sample matrix of expression estimates
- qc a miRNA x sample matrix of quality scores (here R-squared values)

Examples

data(qpcR15)

qpcRlinexp	The processed data generated using the linear-exponential method im-
	plemented in the $qpcR$ software package.

Description

These data are also used to demonstrate the functionality of the miRcomp package.

Usage

data(qpcRlinexp)

Format

A list with two elements.

- ct a miRNA x sample matrix of expression estimates
- qc a miRNA x sample matrix of quality scores (here R-squared values)

Examples

data(qpcRlinexp)

qualityAssessment Quality Assessment for MicroRNA Expression Estimates

Description

This function produces several figures to examine quality scores.

Usage

titrationResponse

Arguments

object1	a list containing two elements: ct (the expression estiamtes) and qc (quality scores)
object2	an optional second list of the same format as object1, used to compare two methods.
cloglog1	if TRUE, the -log(-log(object1\$qc)) is plotted. This is useful to visualize certain quality scores such as R-squared.
cloglog2	if TRUE, the -log(-log(object2\$qc)) is plotted. This is useful to visualize certain quality scores such as R-squared.
na.rm	if TRUE, quality scores corresponding to NA expression estimates are removed from analysis. This is only used if plotType=="boxplot".
plotType	type of plot to return. Options are a scatterplot or a boxplot.
label1	optional label corresponding to object 1 to be used in plotting.
label2	optional label corresponding to object 2 to be used in plotting.

Value

The function generates a plot of the desired type.

Author(s)

Matthew N. McCall

Examples

```
data(lifetech)
qualityAssessment(lifetech, plotType="boxplot")
```

data(qpcRdefault)
qualityAssessment(object1=lifetech,object2=qpcRdefault,cloglog2=TRUE)

titrationResponse Assess monotone signal across titration

Description

This function determines

Usage

Arguments

object1	a list containing two elements: ct (the expression estiamtes) and qc (quality scores)
qcThreshold1	a numeric threshold corresponding to object1\$qc below which values are con- sidered low quality.
object2	an optional second list of the same format as object1, used to compare two methods.
qcThreshold2	a numeric threshold corresponding to object2\$qc below which values are con- sidered low quality.
commonFeatures	if TRUE and object2 is non-NULL, only high quality non-NA features in com- mon between both objects are used.
label1	optional label corresponding to object 1 to be used in plotting.
label2	optional label corresponding to object 2 to be used in plotting.

Value

A table listing the number of features showing a monotone titration response and a figure showing the proportion of features showing a monotone titration response vs the difference in pure sample expression.

Author(s)

Matthew N. McCall

Examples

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