# Package 'AffyRNADegradation'

## October 8, 2015

Type Package
<b>Title</b> Analyze and correct probe positional bias in microarray data due to RNA degradation
<b>Version</b> 1.14.0
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<b>Depends</b> R ( $>= 2.9.0$ ), methods, affy
Suggests AmpAffyExample
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<b>Description</b> The package helps with the assessment and correction of RNA degradation effects in Affymetrix 3' expression arrays. The parameter d gives a robust and accurate measure of RNA integrity. The correction removes the probe positional bias, and thus improves comparability of samples that are affected by RNA degradation.
License GPL-2
Collate AllClasses.R decayFunction.R AffyRNADegradation.R probeInfo.R tongs.R
<b>biocViews</b> GeneExpression, Microarray, OneChannel, Preprocessing, QualityControl
NeedsCompilation no
R topics documented:
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AffyDegradationBatch-class

Class AffyDegradationBatch

#### **Description**

This class represents Affymetrix GeneChip probe level data that has been analysed and corrected for the probe location bias.

#### **Objects from the Class**

Objects can be created using the function RNADegradation.

#### **Slots**

location.type: Object of class character describing the type of probe position used for the analysis (probe index or probe location).

afbatch: Object of class AffyBatch containing corrected probe level data.

stats: Object of class matrix containg various statistical parameters from the analysis.

means.pm: Object of class matrix containing the average PM probe intensites for probe locations of expressed genes.

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#### Methods

**afbatch** signature(x = "AffyDegradationBatch"): returns as AffyBatch object containing corrected probe level data.

**d** signature(x = "AffyDegradationBatch"): returns a vector with a measure of RNA degradation for each sample

plotDx signature(x = "AffyDegradationBatch"): plots the probe location bias for all samples
 of the experiment.

#### Author(s)

Mario Fasold

#### See Also

AffyBatch

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#### **Examples**

```
if (require(AmpAffyExample)) {
    # Get example data
    data(AmpData)

    affy.deg <- RNADegradation(AmpData[,4])

    ## Plot degradation
    plotDx(affy.deg)

    ## Get degradation measure
    d(affy.deg)

## Get AffyBatch object with corrected probe intensities
    afbatch(affy.deg)
}</pre>
```

 ${\tt AffyRNADegradation}$ 

AffyRNADegradation: analyze and correct RNA degradation effects

#### **Description**

The AffyRNADegradation package helps in the assessment of RNA degradation effects in Affymetrix 3' expression arrays. The parameter d gives a robust and accurate measure of RNA integrity. The correction removes the probe positional bias, and thus improves comparability of samples that are affected by RNA degradation.

#### **Details**

Package: AffyRNADegradation

Type: Package
Version: 0.1.0
Date: 2011-10-13
License: GPL >=2

The RNADegradation function performs both analysis and correction of RNA degradation effects, returning an object of type AffyDegradationBatch. The class provides accessor functions to obtain the degradation parameter d and a AffyBatch object containing corrected probe intensities. A more detailed look on the RNA degradation effects can be gained through the tongs plot, the Dx plot and further statistics.

#### Author(s)

Mario Fasold

GetTongs

#### **Examples**

```
if (require(AmpAffyExample)) {
    # Load example data (AmpData affybatch)
    data(AmpData)

## Compute and correct degradation for a single chip (for speed)
    affy.deg <- RNADegradation(AmpData[,4])

## Show degradation parameter d
    d(affy.deg)

## Replace data with corrected data for further analysis
    AmpData <- afbatch(affy.deg)
}</pre>
```

GetTongs

Generate and visualize the tongs plot

#### **Description**

The tongs plot and the related degradation hook show the relationship between expression level and probe location bias. They are required for the correct estimation of RNA degradation effects.

#### Usage

```
GetTongs(affyData, chip.idx = 1)
PlotTongs(tongs)
PlotDegradationHooks(affyData, ...)
```

#### **Arguments**

```
affyData an AffyBatch object.

chip.idx index of the sample to compute the tongs for.

tongs the tongs plot data table.

... optional graphical parameters.
```

#### Value

Tongs a table containing Tongs plot values ordered by expression level.

#### Author(s)

Mario Fasold

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#### **Examples**

```
if (require(AmpAffyExample)) {
    # Get example data
    data(AmpData)

    tongs <- GetTongs(AmpData, chip.idx = 4)
    PlotTongs(tongs)

PlotDegradationHooks(AmpData[,c(3,4)])
}</pre>
```

RNADegradation

Compute degradation parameters and correct probe intensities

#### **Description**

Given an AffyBatch object, the function RNADegradation computes the probe positional bias and several statistical parameters, including a robust and accurate measure of RNA degradation. Probe intensities are corrected for the bias for each sample within the AffyBatch object.

#### Usage

#### **Arguments**

```
affyData an AffyBatch object containing the probe level microarray data.

location.type index-based probe alignment (x=k) if "index", or location-based alignment (x=L) if "absolute".

location.file.dir directory containing the probe location file(s).

plot.images if TRUE plots a set of debug images.
```

#### Value

An AffyDegradationBatch object.

#### Author(s)

Mario Fasold

#### See Also

AffyDegradationBatch

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### Examples

```
if (require(AmpAffyExample)) {
    # Get example data
    data(AmpData)

affy.deg <- RNADegradation(AmpData[,4])
}</pre>
```

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