

# Package ‘AgiMicroRna’

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|            |                                  |
|------------|----------------------------------|
| basicLimma | <i>Linear models Using limma</i> |
|------------|----------------------------------|

---

## Description

Differential expression analysis using the linear model features implemented in the limma package. A linear model is fitted to each miRNA gene so that the fold change between different experimental conditions and their standard errors can be estimated. Empirical Bayes methods are applied to obtain moderated statistics

## Usage

```
basicLimma(eset, design, CM, verbose = FALSE)
```

## Arguments

|         |                                                              |
|---------|--------------------------------------------------------------|
| eset    | ExpressionSet containing the processed log-expression values |
| design  | design matrix                                                |
| CM      | contrast matrix                                              |
| verbose | logical, if TRUE prints out output                           |

## Details

In our data example (see the target file in Table 1 in vignette), we have used a paired design (by subject) to assess the differential expression between two treatments B and C vs a control treatment A. That is, we want to obtain the microRNAs that are differentially expressed between conditions A vs B and A vs C. The linear model that we are going to fit to every miRNA is defined by equation:  $y = \text{Treatment} + \text{Subject} + \text{error term}$ . This model is going to estimate the treatment effect and then, the comparison between the different treatments are done in terms of contrasts between the estimates of the treatment effects. To fit the model, we need first to define a design matrix. The design matrix

is an incidence matrix that relates each array/sample/file to its given experimental conditions, in our case, relates each file to one of the three treatments and with its particular subject. If treatment is a factor variable, we can define the design matrix using `model.matrix(~ -1 + treatment + subject)`. Then the linear model can be fitted using `fit=lmFit(eset,design)`. This will get the treatment estimates for each microRNA in the `eset` object:

```
treatmentA treatmentB treatmentC subject2 hsa-miR-152 7.5721 7.656 7.566 -0.1157 hsa-miR-15a* 0.9265 1.066 1.211 -0.2242 hsa-miR-337-5p 6.2448 7.298 7.084 -0.4489
```

We can define the contrasts of interest using a contrast matrix as in `CM=cbind(MSC\_AvsMSC\_B=c(1,-1,0), MSC\_AvsMSC\_C=c(1,0,-1))`

And then, we can estimate those contrasts using `fit2=contrasts.fit(fit,CM)`. Finally, we can obtain moderated statistics using `fit2=eBayes(fit2)`.

The function 'basicLimma' implemented in AgiMicroRna produces the last `fit2` object, that has in `fit2$coeff` the M values, in `fit2$t` the moderated-t statistic of the contrasts, and in `fit2$p.value` the corresponding p value of each particular contrasts. Be aware that these p values must be corrected by multiple testing.

```
MSC\_AvsMSC\_B MSC\_AvsMSC\_C hsa-miR-152 0.67567761 0.977326746 hsa-miR-15a* 0.68019442 0.413657270 hsa-miR-337-5p 0.03737814 0.075248741
```

See `limmaUsersGuide()` for a complete description of the limma package.

## Value

An `MArrayLM` object of the package limma

## Author(s)

Pedro Lopez-Romero

## References

Smyth, G. K. (2005). Limma: linear models for microarray data. In: 'Bioinformatics and Computational Biology Solutions using R and Bioconductor'. R. Gentleman, V. Carey, S. Dudoit, R. Irizarry, W. Huber (eds), Springer, New York, pages 397–420.

Smyth, G. K. (2004). Linear models and empirical Bayes methods for assessing differential expression in microarray experiments. *Statistical Applications in Genetics and Molecular Biology*, Vol. 3, No. 1, Article 3. <http://www.bepress.com/sagmb/vol3/iss1/art3>

## See Also

An 'RGList' example containing processed data is in `ddPROC` and an overview of how the processed data is produced is given in `filterMicroRna`. The `ExpressionSet` object can be generated using `esetMicroRna`

## Examples

```
## Not run:
data(targets.micro)
data(ddPROC)
esetPROC=esetMicroRna(ddPROC,targets.micro,makePLOT=FALSE,verbose=FALSE)

levels.treatment=levels(factor(targets.micro$Treatment))
treatment=factor(as.character(targets.micro$Treatment),
  levels=levels.treatment)
```

```

levels.subject=levels(factor(targets.micro$Subject))
subject=factor(as.character(targets.micro$Subject),
               levels=levels.subject)

design=model.matrix(~ -1 + treatment + subject )

CM=cbind(MSC_AvsMSC_B=c(1,-1,0,0),
         MSC_AvsMSC_C=c(1,0,-1,0))

fit2=basicLimma(esetPROC,design,CM,verbose=TRUE)

names(fit2)
head(fit2$coeff)
head(fit2$p.value)
plot(fit2$Amean,fit2$coeff[,1],xlab="A",ylab="M")
abline(h=0)
abline(h=c(-1,1),col="red")
plot(fit2$coeff[,1],fit2$p.value[,1], xlab="M",ylab="p value")

## End(Not run)

```

---

boxplotMicroRna

*Boxplot*


---

## Description

It creates a Boxplot using the matrix columns as input

## Usage

```
boxplotMicroRna(object, maintitle, colorfill, xlab, ylab)
```

## Arguments

|           |                                                                    |
|-----------|--------------------------------------------------------------------|
| object    | A matrix containing by columns the expression arrays in log2 scale |
| maintitle | character to indicate the title of the graph                       |
| colorfill | color to fill the boxplot                                          |
| xlab      | title for the x axe                                                |
| ylab      | title for the y axe                                                |

## Author(s)

Pedro Lopez-Romero

## Examples

```
data(dd.micro)
MMM=log2(dd.micro$meanS)
      boxplotMicroRna(MMM,
      maintitle="log2 Mean Signal",
      colorfill="green",
      xlab="Samples",
      ylab="expression")
```

---

|         |                                                                   |
|---------|-------------------------------------------------------------------|
| cvArray | <i>Coefficient of variation of replicated probes within array</i> |
|---------|-------------------------------------------------------------------|

---

## Description

Identifies replicated features at probe and at gene level and computes the coefficient of variation of the array

## Usage

```
cvArray(ddDUP, foreground = c("MeanSignal", "ProcessedSignal"), targets, verbose=FALSE)
```

## Arguments

|            |                                                                               |
|------------|-------------------------------------------------------------------------------|
| ddDUP      | uRNAList, containing the output from readMicroRnaAFE                          |
| foreground | Specifies the signal used, only "MeanSignal" or "ProcessedSignal" can be used |
| targets    | data.frame with the target structure                                          |
| verbose    | logical, if TRUE prints out output                                            |

## Details

In the Agilent microRNA platforms the features are replicated at a probe level and normally, a single microRNA is interrogated by either two or four sets of replicated probes. The replication of the probes allows computing the coefficient of variation (CV) for each array as a measure of the reproducibility of the array. The CV is computed for every set of replicated probes and the CV median is reported as the array CV. A lower array CV indicates a better array reproducibility.

## Value

It prints out the results of the replication for the NON CONTROL FEATURES at a probe and gene level.

## Author(s)

Pedro Lopez-Romero

## Examples

```
## Not run:
data(dd.micro)
data(targets.micro)

cvArray(dd.micro, "MeanSignal", targets.micro, verbose=TRUE)

graphics.off()

## End(Not run)
```

---

|          |                                |
|----------|--------------------------------|
| dd.micro | <i>data example (uRNAList)</i> |
|----------|--------------------------------|

---

## Description

Data, extracted from scanned images using Agilent Feature Extraction Software, are stored in a `uRNAList` object.

## Usage

```
data(dd.micro)
```

## Details

A data example is provided. The data example includes 3 experimental conditions with two replicates.

For these data, chips were scanned using the Agilent G2567AA Microarray Scanner System (Agilent Technologies) Image analysis and data collection were carried out using the Agilent Feature Extraction 9.1.3.1. (AFE).

Data, collected with the Agilent Feature Extraction Software, are stored in a `uRNAList` object with the following components:

```
uRNAList$TGS matrix, 'gTotalGeneSignal'
uRNAList$TPS matrix, 'gTotalProbeSignal'
uRNAList$meanS matrix, 'gMeanSignal'
uRNAList$procS matrix, 'gProcessedSignal'
uRNAList$targets data.frame, 'FileName'
uRNAList$genes$ProbeName vector of characters, 'AGilent Probe Name'
uRNAList$genes$GeneName vector of characters, 'microRNA Name'
uRNAList$genes$ControlType vector of integers, '0' = Feature, '1' = Positive control, '-1' =
  Negative control
uRNAList$other$IsGeneDetected matrix, FLAG to classify signal if 'IsGeneDetected=1' or
  'not=0'
uRNAList$other$IsSaturated matrix, FLAG to classify signal if 'IsSaturated = 1' or 'not=0'
uRNAList$other$IsFeatPopnOL matrix, FLAG to classify signal if 'IsFeatPopnOL = 0' or
  'not=1'
```

**uRNAList\$other\$gIsFeatNonUnifOL** matrix, FLAG to classify signal if 'gIsFeatNonUnifOL = 0' or 'not=1'

**uRNAList\$other\$gBGMedianSignal** matrix, gBGMedianSignal

**uRNAList\$other\$gBGUsed** matrix, gBGUsed

#### Author(s)

Pedro Lopez-Romero

#### See Also

readMicroRnaAFE.Rd

---

ddPROC

*Processed miRNA data (uRNAList)*


---

#### Description

Filtered and Normalized miRNA data stored in a uRNAList object.

#### Usage

```
data(ddPROC)
```

#### Details

ddPROC is originated after the processing of the dd.micro raw data.

#### Author(s)

Pedro Lopez-Romero

#### See Also

An overview of how ddPROC is obtained is given in filterMicroRna

---

dim.uRNAList

*Retrieve the Dimensions of an uRNAList Object*


---

#### Description

Retrieve the number of rows (genes) and columns (arrays) for an uRNAList object.

#### Usage

```
## S3 method for class 'uRNAList'
dim(x)
## S3 method for class 'uRNAList'
length(x)
```

**Arguments**

x                      an object of class uRNAList

**Details**

This function and this file, has been borrowed from the files created by Gordon Smyth for the limma package.

**Value**

Numeric vector of length 2. The first element is the number of rows (genes) and the second is the number of columns (arrays).

**Author(s)**

Pedro Lopez-Romero

---

|                   |                                                           |
|-------------------|-----------------------------------------------------------|
| dimnames.uRNAList | <i>Retrieve the Dimension Names of an uRNAList Object</i> |
|-------------------|-----------------------------------------------------------|

---

**Description**

Retrieve the dimension names of a microarray data object.

**Usage**

```
## S3 method for class 'uRNAList'
dimnames(x)
## S3 replacement method for class 'uRNAList'
dimnames(x) <- value
```

**Arguments**

x                      an object of class uRNAList  
value                  a possible value for dimnames(x)

**Details**

The dimension names of a microarray object are the same as those of the most important matrix component of that object. A consequence is that rownames and colnames will work as expected. This function and this file, has been borrowed from the files created by Gordon Smyth for the limma package.

**Value**

Either NULL or a list of length 2.

**Author(s)**

Pedro Lopez-Romero



---

|              |                                             |
|--------------|---------------------------------------------|
| esetMicroRna | <i>ExpressionSet object from a uRNAList</i> |
|--------------|---------------------------------------------|

---

## Description

It creates an 'ExpressionSet' object from a 'uRNAList' with unique probe names. Typically, the 'uRNAList' object contains the Total Gene Processed data.

## Usage

```
esetMicroRna(uRNAList, targets, makePLOT=FALSE, verbose=FALSE)
```

## Arguments

|          |                                                                                                                                       |
|----------|---------------------------------------------------------------------------------------------------------------------------------------|
| uRNAList | An uRNAList containing normally the processed data                                                                                    |
| targets  | data.frame with the targets structure                                                                                                 |
| makePLOT | logical, if TRUE it makes a 'heatmap' with the 100 greater variance genes, a 'hierarchical cluster' with all the genes and a pca plot |
| verbose  | logical, if TRUE prints out output                                                                                                    |

## Details

It creates an ExpressionSet object from a uRNAList. Usually this function is applied to an uRNAList object containing the Total Gene Processed data.

## Value

An ExpressionSet object

## Author(s)

Pedro Lopez-Romero

## See Also

An 'uRNAList' example containing processed data is in ddPROC and an overview of how the processed data is produced is given in filterMicroRna

---

|                |                        |
|----------------|------------------------|
| filterMicroRna | <i>Filtering Genes</i> |
|----------------|------------------------|

---

## Description

Filter genes out according to their Quality Flag

## Usage

```
filterMicroRna(ddNORM,
  dd,
  control,
  IsGeneDetected,
  wellaboveNEG,
  limIsGeneDetected,
  limNEG,
  makePLOT,
  targets,
  verbose,
  writeout)
```

## Arguments

|                   |                                                                                                                                                                         |
|-------------------|-------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| ddNORM            | uRNAList with the Total Gene Signal in log2 scale to be FILTERED out according to a Quality FLAG                                                                        |
| dd                | uRNAList, containing the output from readMicroRnaAFE                                                                                                                    |
| control           | logical, if TRUE it removes controls                                                                                                                                    |
| IsGeneDetected    | logical, if TRUE it filters genes according to gIsGeneDetected Flag. Flag = 1, then gene is detected                                                                    |
| wellaboveNEG      | logical, if TRUE it filter genes whose expression is not above a limit value defined by the expression of negative controls. Limit= Mean(negative) + 1.5 x sd(negative) |
| limIsGeneDetected | for a given feature xi accros samples, is the minimum in at least one experimental condition with a IsGeneDetected-FLAG = 1 (Is Detected)                               |
| limNEG            | for a given feature xi accros samples, is the minimum in at least one experimental condition with intensity > Limit established for negative controls (Mean + 1.5 x SD) |
| makePLOT          | logical, if TRUE makes QC plots with the remaining signals                                                                                                              |
| targets           | data.frame with the targets structure                                                                                                                                   |
| verbose           | logical, if TRUE prints out output                                                                                                                                      |
| writeout          | logical, if TRUE writes out output files                                                                                                                                |

## Details

Agilent Feature Extraction software provides a flag for each spot that identifies different quantification errors of the signal. Quantification flags were used to filter out signals that did not reach a minimum established criterion of quality.

## Value

The function returns a uRNAList containing the FILTERED data. In order to allow the tracking of those microRNAs that may have been filtered out from the original raw data, the following files are given:

NOCtrl\_exprs.txt: Log2 Normalized Total Gene Signals for the Non Control Genes  
 NOCtrl\_FlagIsGeneDetected.txt: IsGeneDetected Flag for the Non Control Genes, 1 = detected  
 IsNOTGeneDetected.txt: Genes that not are not detected according to IsGeneDetected Flag

**Author(s)**

Pedro Lopez-Romero

**References**Agilent Feature Extraction Reference Guide <http://www.Agilent.com>**Examples**

```

data(dd.micro,verbose=FALSE)
data(targets.micro,verbose=FALSE)
ddTGS=tgsMicroRna(dd.micro,half=TRUE,makePLOT=FALSE,verbose=FALSE)
ddNORM=tgsNormalization(ddTGS,'quantile',
                        makePLOTpre=FALSE,makePLOTpost=TRUE,targets.micro,verbose=FALSE)
ddPROC=filterMicroRna(ddNORM,
                      dd.micro,
                      control=TRUE,
                      IsGeneDetected=TRUE,
                      wellaboveNEG=FALSE,
                      limIsGeneDetected=50,
                      limNEG=25,
                      makePLOT=FALSE,
                      targets.micro,
                      verbose=FALSE,
                      writeout=FALSE)

```

getDecideTests

*Differential expression analysis an multiplicity of the tests***Description**

It Uses the decideTests function of the 'limma' package to classify the list of genes as up, down or not significant after correcting by the multiplicity of the tests.

**Usage**

```
getDecideTests(fit2, DEmethod, MTestmethod, PVcut,verbose=FALSE)
```

**Arguments**

|             |                                                                                                         |
|-------------|---------------------------------------------------------------------------------------------------------|
| fit2        | MArrayLM object                                                                                         |
| DEmethod    | method for decideTests, only 'separate' or 'nestedF' are implemented. see decideTests in limma package. |
| MTestmethod | method for multiple test, choices are 'none','BH', 'BY', ... see p.adjust                               |
| PVcut       | p value threshold to declare significant features                                                       |
| verbose     | logical, if TRUE prints out output                                                                      |

**Value**

A 'TestResults' object of the 'limma' package It prints out the number of UP and DOWN genes for every contrasts according to the p value limit specified

**Author(s)**

Pedro Lopez-Romero

**References**

Smyth, G. K. (2005). Limma: linear models for microarray data. In: 'Bioinformatics and Computational Biology Solutions using R and Bioconductor'. R. Gentleman, V. Carey, S. Dudoit, R. Irizarry, W. Huber (eds), Springer, New York, pages 397–420.

**See Also**

An overview of miRNA differential expression analysis is given in `basicLimma`

**Examples**

```
## Not run:
DE=getDecideTests(fit2,
  DEmethod="separate",
  MTestmethod="BH",
  PVcut=0.10,
  verbose=TRUE)

## End(Not run)
```

---

HeatMapMicroRna

*HeatMap*

---

**Description**

Creates a HeatMap graph using the 'heatmap.2' function

**Usage**

```
HeatMapMicroRna(object, size, maintitle)
```

**Arguments**

|                        |                                                               |
|------------------------|---------------------------------------------------------------|
| <code>object</code>    | A expression Matrix                                           |
| <code>size</code>      | number of highest variance genes to be considered in the plot |
| <code>maintitle</code> | title of the plot                                             |

**Author(s)**

Pedro Lopez-Romero

**See Also**

`heatmap.2`

**Examples**

```
data(ddPROC)
  HeatMapMicroRna(ddPROC$TGS,
    size=100,
    maintitle="100 High Var genes")
```

---

|                  |                                |
|------------------|--------------------------------|
| hierclusMicroRna | <i>Hierarchical clustering</i> |
|------------------|--------------------------------|

---

**Description**

Hierarchical cluster of samples using the 'hclust' function

**Usage**

```
hierclusMicroRna(object, GErep, methdis, methclu, sel, size)
```

**Arguments**

|         |                                                                                             |
|---------|---------------------------------------------------------------------------------------------|
| object  | An expression Matrix                                                                        |
| GErep   | Numerical vector that relates each sample with its experimental condition                   |
| methdis | the distance measure to be used. Options are 'euclidean' and 'pearson'. see 'dist' function |
| methclu | the agglomeration method to be used by the 'hclust' function                                |
| sel     | logical, if TRUE selects the 'size' highest variance genes for the plot                     |
| size    | selects the 'size' highest variance genes for the plot if 'sel=TRUE'                        |

**Author(s)**

Pedro Lopez-Romero

**See Also**

hclust, dist

**Examples**

```
data(targets.micro)
data(ddPROC)

hierclusMicroRna(ddPROC$TGS, GErep,
  methdis="euclidean",
    methclu="complete",
  sel=FALSE, 100)
```

mvaBASIC

*MVA plot***Description**

For each array, the M value is computed for every spot as the difference between the spot intensity in the array and the averaged intensity for that feature over the whole set of arrays. It does not make a distinction between the different kind of features in the array as the mvaMicroRna() does.

**Usage**

```
mvaBASIC(object, colorfill, maintitle)
```

**Arguments**

|           |                                    |
|-----------|------------------------------------|
| object    | An expression matrix in log2 scale |
| colorfill | color of the plot                  |
| maintitle | title of the plot                  |

**Author(s)**

Pedro Lopez-Romero

**Examples**

```
data(dd.micro)
op=par(mfrow=c(1,1),ask=TRUE)
mvaBASIC(log2(dd.micro$meanS),
colorfill="red",
          maintitle=" log2 Mean Signal")
par(op)
```

mvaMicroRna

*MA plot***Description**

For each array, the M value is computed for every spot as the difference between the spot intensity in the array and the median intensity for that feature over the whole set of arrays. Every kind of feature is identified with different color (microRNA genes, positive controls, etc ...) The input must be an uRNAList object created by the user, in such a way that the uRNAList\$meanS field contains the expression matrix that we want to use in log2 scale (see example below) The gProcessedSignal computed by the Agilent Feature Extraction software normally contains negative values, so a small constant has to be added to the signals before log transformation.

**Usage**

```
mvaMicroRna(uRNAList, maintitle, verbose=FALSE)
```

**Arguments**

|           |                                                                                                                                      |
|-----------|--------------------------------------------------------------------------------------------------------------------------------------|
| uRNAList  | A uRNAList object. It uses the expression matrix stored in the uRNAList\$meanS slot. Input expression matrix should be in log2 scale |
| maintitle | character to indicate the title of the graph                                                                                         |
| verbose   | logical, if TRUE it prints details                                                                                                   |

**Author(s)**

Pedro Lopez-Romero

**Examples**

```
data(dd.micro)
op=par(mfrow=c(1,1),ask=TRUE)

MMM=dd.micro$procS      ## gProcessedSignal

min=min(MMM)           ## transforming gProcessedSignal to positive values
for(i in 1:dim(MMM)[2]){ ## before log2 transformation
  MMM[,i]=MMM[,i]+(abs(min)+ 5)
}

ddaux=dd.micro
ddaux$meanS=log2(MMM)
mvaMicroRna(ddaux,maintitle="ProcessedSignal",verbose=FALSE)
rm(ddaux)
par(op)
```

---

|                 |                 |
|-----------------|-----------------|
| PCAplotMicroRna | <i>PCA plot</i> |
|-----------------|-----------------|

---

**Description**

It is a wrapper for the 'plotPCA' of the 'affycoretools' package

**Usage**

```
PCAplotMicroRna(eset, targets)
```

**Arguments**

|         |                                      |
|---------|--------------------------------------|
| eset    | An Expression Set object             |
| targets | data.frame with the target structure |

**Author(s)**

Pedro Lopez-Romero

**Examples**

```
data(targets.micro)
data(ddPROC)
esetPROC=esetMicroRna(ddPROC,targets.micro,makePLOT=FALSE,verbose=FALSE)

PCAplotMicroRna(esetPROC,targets.micro)
```

---

`plotDensityMicroRna`      *Density Plots of Intensity Signals*

---

**Description**

Creates a density plot with the arrays intensities

**Usage**

```
plotDensityMicroRna(object, maintitle)
```

**Arguments**

|                        |                                     |
|------------------------|-------------------------------------|
| <code>object</code>    | An expression matrix, in log2 scale |
| <code>maintitle</code> | title of the plot                   |

**Author(s)**

Pedro Lopez-Romero

**Examples**

```
data(dd.micro)
plotDensity(log2(dd.micro$meanS), maintitle="log2 Mean Signal")
```

---

`pvalHistogram`      *Histogram of the p values*

---

**Description**

Creates an histogram of the pvalues. For multiple contrasts, creates an histogram for every t.test pvalue (separate) or a single histogram for the F.test pvalue (nestedF). A uniform histogram will indicate no differential expression in the data set, whereas a right skewed histogram, will indicate some significant differential expression

**Usage**

```
pvalHistogram(fit2, DE, PVcut, DEmethod, MTestmethod, CM, verbose=FALSE)
```

**Arguments**

|                          |                                                                      |
|--------------------------|----------------------------------------------------------------------|
| <code>fit2</code>        | MArrayLM object                                                      |
| <code>DE</code>          | TestResults object                                                   |
| <code>PVcut</code>       | limit p value to declare significant features                        |
| <code>DEmethod</code>    | method for decideTests, only 'separate' or 'nestedF' are implemented |
| <code>MTestmethod</code> | method for multiple test                                             |
| <code>CM</code>          | contrast matrix                                                      |
| <code>verbose</code>     | logical, if TRUE prints out output                                   |



**Author(s)**

Pedro Lopez-Romero

**See Also**

An overview of miRNA differential expression analysis is given in `basicLimma`. An example of how to get the 'TestResults' object is in `getDecideTests`.

**Examples**

```
## Not run:
pvalHistogram(fit2, DE, PVcut=0.10,
              DEmethod="separate", MTestmethod="BH", CM)

## End(Not run)
```

---

qcPlots

*Plots for Quality Assessment*


---

**Description**

It creates BoxPlots, Density Plots, MA plots, RLE plots and hierarchical clustering plots with the sample data set.

**Usage**

```
qcPlots(dd,
        offset,
        MeanSignal=TRUE,
        ProcessedSignal=FALSE,
        TotalProbeSignal=FALSE,
        TotalGeneSignal=FALSE,
        BGMedianSignal=FALSE,
        BGUsed=FALSE,
        targets)
```

**Arguments**

|                  |                                                                 |
|------------------|-----------------------------------------------------------------|
| dd               | A uRNAList object containing the output from readMicroRnaAFE    |
| offset           | numeric value to add to the intensities before log transforming |
| MeanSignal       | logical, if TRUE "gMeanSignal" is used                          |
| ProcessedSignal  | logical, if TRUE "gProcessedSignal" is used                     |
| TotalProbeSignal | logical, if TRUE "gTotalProbeSignal" is used                    |
| TotalGeneSignal  | logical, if TRUE "gTotalGeneSignal" is used                     |
| BGMedianSignal   | logical, if TRUE "gBGMedianSignal" is used                      |
| BGUsed           | logical, if TRUE "gBGUsed" is used                              |
| targets          | data.frame with the target structure                            |

## Details

The signals loaded from the AFE data files can be used for the quality assesment using the graphical utilities included in the `qcPlots` function. For the `gMeanSignal`, the `BoxPlots`, `Density Plots`, `MA plots`, `RLE plots` and `hierachical clustering plots` are done. For the `gProcessedSignal` the same plots are done, except the `hierachical clustering`. For the `gTotalProbeSignal` and the `gTotalGeneSignal` only the `BoxPlots` and `Density Plots` are done, and finally, for the `Background` signals only the `Boxplots` are done.

## Author(s)

Pedro Lopez-Romero

## References

Boldstad B.M., Collin F., Brettschneider J., Simpson, K., Cope L., Irizarry R. A., Speed T. P. Quality Assesment of Affymetrix GeneChip Data. In *Bioinformatics and Computational Biology Solutions Using R and Bioconductor*. (eds.) Gentleman R., Carey V. J., Huber W., Irizarry R. A., Dudoit S. (2005). Springer.

## See Also

`boxplotMicroRna`, `plotDensityMicroRna`, `RleMicroRna`, `mvaMicroRna` and `hierclusMicroRna`

## Examples

```
## Not run:
data(dd.micro)
qcPlots(dd.micro,offset=5,
        MeanSignal=TRUE,
        ProcessedSignal=TRUE,
        TotalProbeSignal=TRUE,
        TotalGeneSignal=TRUE,
        BGMedianSignal=TRUE,
        BGUsed=TRUE,
        targets.micro)
graphics.off()

## End(Not run)
```

---

readMicroRnaAFE

*Read Agilent Feature Extraction txt data files*

---

## Description

Read the data files generated by the Agilent Feature Extraction image analysis software

## Usage

```
readMicroRnaAFE(targets,verbose=FALSE)
```

## Arguments

|         |                                                                                                |
|---------|------------------------------------------------------------------------------------------------|
| targets | A data frame that specifies experimental conditions under which each sample has been obtained. |
| verbose | logical, if TRUE prints out output                                                             |

## Details

The function reads the \*.txt files generated by the AFE Software using the 'read.maimages' function of 'limma' package.

Data, collected with the Agilent Feature Extraction Software, are stored in a uRNAList object with the following components:

```
- dd.micro\TGS 'gTotalGeneSignal' - dd.micro\TPS 'gTotalProbeSignal' - dd.micro\meanS 'gMeanSignal'
- dd.micro\procS 'gProcessedSignal' - dd.micro\targets 'targets' - dd.micro\genes\ProbeName
'Probe Name' - dd.micro\genes\GeneName 'microRNA Name' - dd.micro\genes\ControlType
'FLAG to specify the sort of feature' - dd.micro\other\IsGeneDetected 'FLAG IsGeneDetected'
- dd.micro\other\IsSaturated 'FLAG IsSaturated' - dd.micro\other\IsFeatNonUnifOL 'FLAG
IsFeatNonUnifOL' - dd.micro\other\IsFeatPopnOL 'FLAG IsFeatPopnOL' - dd.micro\other\gBGMedianSignal
'gBGMedianSignal' - dd.micro\other\gBGUsed 'gBGUsed'
```

## Value

A uRNAList containing the following elements:

```
uRNAList\TGS    matrix, 'gTotalGeneSignal'
uRNAList\TPS    matrix, 'gTotalProbeSignal'
uRNAList\meanS
                matrix, 'gMeanSignal'
uRNAList\procS
                matrix, 'gProcessedSignal'
uRNAList\targets
                data.frame, 'FileName'
uRNAList\genes\ProbeName
                character, 'AGilent Probe Name'
uRNAList\genes\GeneName
                character, 'microRNA Name'
uRNAList\genes\ControlType
                integer, '0' = Feature, '1' = Positive control, '-1' = Negative control
uRNAList\other\IsGeneDetected
                matrix, FLAG to classify signal if 'IsGeneDetected=1' or 'not=0'
uRNAList\other\IsSaturated
                matrix, FLAG to classify signal if 'IsSaturated = 1' or 'not=0'
uRNAList\other\IsFeatPopnOL
                matrix, FLAG to classify signal if 'IsFeatPopnOL = 0' or 'not=1'
uRNAList\other\IsFeatNonUnifOL
                matrix, FLAG to classify signal if 'IsFeatNonUnifOL = 0' or 'not=1'
uRNAList\other\gBGMedianSignal
                matrix, gBGMedianSignal
uRNAList\other\gBGUsed
                matrix, gBGUsed
```

**Author(s)**

Pedro Lopez-Romero

**References**

Agilent Feature Extraction Reference Guide <http://www.Agilent.com>

Smyth, G. K. (2005). Limma: linear models for microarray data. In: 'Bioinformatics and Computational Biology Solutions using R and Bioconductor'. R. Gentleman, V. Carey, S. Dudoit, R. Irizarry, W. Huber (eds), Springer, New York, pages 397–420.

**See Also**

A data example can be found in `dd.micro` See also `readTargets` to see how to build the target file and the example given in `targets.micro`

**Examples**

```
## Not run:
data(targets.micro)
dd.micro = readMicroRnaAFE(targets.micro)

## End(Not run)
```

---

`readTargets`

*read the target file*

---

**Description**

The target file is a txt file created by the user where every input file (array, sample) is attached to a experimental condition

**Usage**

```
readTargets(infile, verbose=FALSE)
```

**Arguments**

|                      |                                                           |
|----------------------|-----------------------------------------------------------|
| <code>infile</code>  | name of the target file, for instance 'targets.micro.txt' |
| <code>verbose</code> | logical, if TRUE prints out output                        |

**Details**

In the 'target' file (see Table 1 in vignette) we specify the experimental conditions under which the data have been generated. The target file **MUST** contain the following mandatory columns: -FileName : Name of the array data file -Treatment : Treatment effect -GRep : Treatment effect in numeric code, from '1' to 'n', being 'n' the number of the levels of the treatment effect

Other explanatory variables specifying the experimental conditions might be also included.

**Value**

A 'data.frame' containing by the columns specified in the input file targets.txt. This 'targets.txt' file must be created by the user.

**Author(s)**

Pedro Lopez-Romero

**References**

Smyth, G. K. (2005). Limma: linear models for microarray data. In: 'Bioinformatics and Computational Biology Solutions using R and Bioconductor'. R. Gentleman, V. Carey, S. Dudoit, R. Irizarry, W. Huber (eds), Springer, New York, pages 397–420.

**See Also**

An example of a target file can be found in `targets.micro`

---

|             |                                |
|-------------|--------------------------------|
| RleMicroRna | <i>Relative Log Expression</i> |
|-------------|--------------------------------|

---

**Description**

RLE: Relative Log Expression

**Usage**

```
RleMicroRna(object, maintitle, colorfill)
```

**Arguments**

|                        |                      |
|------------------------|----------------------|
| <code>object</code>    | An expression matrix |
| <code>maintitle</code> | title of the plot    |
| <code>colorfill</code> | color of the plot    |

**Details**

Each Boxplot corresponds to a sample and displays the Relative Log Expression computed for every spot in the array as the difference between the spot intensity and the median intensity for the same feature accros all the arrays. Since majority of the spots are expected not to be differentially expressed, the plot should show boxplots centered around zero and all of them having the approximately the same dispersion. An array showing greater dispersion than the other, or being not centered at zero could have quality problems.

**Author(s)**

Pedro Lopez-Romero

**References**

Boldstad B.M., Collin F., Brettschneider J., Simpson, K., Cope L., Irizarry R. A., Speed T. P. Quality Assesement of Affymetrix GeneChip Data. In Bioinformatics and Computational Biology Solutions Using R and Bioconductor. (eds.) Gentleman R., Carey V. J., Huber W., Irizarry R. A., Dudoit S. (2005). Springer.

## Examples

```
data(dd.micro)
      RleMicroRna(log2(dd.micro$meanS),
maintitle="log2 Mean Signal RLE",
colorfill="orange")
```

---

rmaMicroRna

*Getting the Total Gene Signal by RMA algorithm*


---

## Description

The function creates an `uRNAList` containing the `TotalGeneSignal` computed by the RMA algorithm. This signal can be used for the statistical analysis.

## Usage

```
rmaMicroRna(dd, normalize, background)
```

## Arguments

|                         |                                                                                                                                             |
|-------------------------|---------------------------------------------------------------------------------------------------------------------------------------------|
| <code>dd</code>         | <code>uRNAList</code> , containing the output from <code>readMicroRnaAFE</code>                                                             |
| <code>normalize</code>  | logical, if TRUE the signal is normalized between arrays using the 'quantile' method                                                        |
| <code>background</code> | logical, if TRUE the signal is background corrected by fitting a normal + exponential convolution model to a vector of observed intensities |

## Details

The function creates an `uRNAList` output that contains in the `uRNAList$TGS`, `uRNAList$TPS`, `uRNAList$meanS` & `uRNAList$procS` slots the Total Gene Signal (TGS) computed by the RMA algorithm. The function uses the robust multiarray average (RMA) method from the 'affy' package. RMA obtains an estimate of the expression measure for each gene using all the replicated probes for that gene. First, RMA obtains a background corrected intensity by fitting a normal + exponential convolution model to a vector of observed intensities. The normal part represents the background and the exponential part represents the signal intensities. Then the arrays are normalized using 'quantile' normalization. Finally, for each probe set that interrogates the same microRNA, RMA fits a linear model to the background-corrected, normalized and log2 transformed probe intensities. This model produces an estimate of the gene signal taking into account the probe effect. The model parameters estimates are obtained by median polish. The estimates of the gene expression signals are referred as RMA estimates. Normally, each microRNA is interrogated by 16 probes either using 2 different probes, each of them replicated 8 times, or using 4 different probes replicated 4 times. First, function 'rmaMicroRna' obtains a background corrected signal using the 'rma.background.correct' function of the package 'preprocessCore', then the signal is normalized between arrays using the 'limma' function 'normalizeBetweenArrays' with the 'quantile' method. Then, the median of the replicated probes is obtained, leading to either 2 or 4 different measures for each gene. These measures correspond to different probes for the same genes that are summarized into a single RMA linear model described above.

## Value

`uRNAList` containing the Total Gene Signal computed by the RMA algorithm in log 2 scale.

**Author(s)**

Pedro Lopez-Romero

**References**

- Irizarry, R., Hobbs, B., Collin, F., Beazer-Barclay, Y., Antonellis, K., Scherf, U., Speed, T. (2003) Exploration, normalization, and summaries of high density oligonucleotide array probe level data. *Biostatistics*. 4, 249-264
- Gautier, L., Cope, L., Bolstad, B. M., and Irizarry, R. A. (2004). *affy*—analysis of Affymetrix GeneChip data at the probe level. *Bioinformatics* 20, 3, 307-315.
- Bolstad B. M. (). *preprocessCore*: A collection of pre-processing functions. R package version 1.4.0
- Smyth, G. K. (2005). *Limma*: linear models for microarray data. In: 'Bioinformatics and Computational Biology Solutions using R and Bioconductor'. R. Gentleman, V. Carey, S. Dudoit, R. Irizarry, W. Huber (eds), Springer, New York, pages 397 - 420

**Examples**

```
data(dd.micro)
ddTGS.rma=rmaMicroRna(dd.micro, normalize=TRUE, background=TRUE)
dim(ddTGS.rma)
RleMicroRna(ddTGS.rma$TGS, "RLE TGS.rma", "blue")
```

---

significantMicroRna      *Summarize Differential Expression Results*

---

**Description**

The function summarizes the results from the differential expression analysis using the different objects that are obtained after 'limma' has been used, such as the 'MArrayLM' object with the statistics and the 'TestResults' object highlighting the significant features.

**Usage**

```
significantMicroRna(eset, ddset, targets, fit2,
CM, DE, DEmethod, MTestmethod, PVcut, Mcut, verbose=FALSE)
```

**Arguments**

|             |                                                                          |
|-------------|--------------------------------------------------------------------------|
| eset        | ExpressionSet containing the Total Gene processed data                   |
| ddset       | An uRNAList object containing the Total Gene processed data              |
| targets     | data.frame with the target structure                                     |
| fit2        | MArrayLM object from eBayes 'limma' function                             |
| CM          | Contrast matrix                                                          |
| DE          | TestResults object                                                       |
| DEmethod    | method used in decideTests, only 'separate' or 'nestedF' are implemented |
| MTestmethod | method for multiple test                                                 |
| PVcut       | p value threshold to declare significant features                        |
| Mcut        | M value threshold to select within significant features                  |
| verbose     | logical, if TRUE prints out output                                       |

## Details

A list containing the genes with their statistics is generated. The significant genes above the PVcut p values are also given in a html file that links the selected miRNAs to the miRBase <http://microrna.sanger.ac.uk/>. A MA plots indicating the differentially expressed genes are also displayed.

When multiple contrasts are done, the method for the selection of the significant genes can be either 'separated' or 'nestedF'. See decideTests in package limma *limma* for a detailed description on these two methods. When 'separated' is used a list with all the genes that have been analyzed in limma is given. The list includes the following columns:

PROBE - Probe name (one of the probes interrogating the gene) GENE - miRNA name PROBE chr\\_coord - Agilent chromosomal location M - Fold change A - Mean of the intensity for that miRNA t - moderated t-statistic pval - p value of the t-statistic adj.pval - p value adjusted by 'MTest-method' fdr.pval - p value adjusted by fdr

Some times, the user can be set 'MTestmethod = none', in this case, it might be interesting to still see the fdr value, despite of the fact that the user has decided not apply any multiple testing correction.

If the 'nestedF' is used, then two lists are provided for each contrasts. A first containing the selected significant genes, and a second list containing the rest of the genes that have been analyzed. The columns given in this case is:

PROBE - Probe name (one of the probes interrogating the gene) GENE - miRNA name PROBE chr\\_coord - Agilent chromosomal location M - Fold change A - Mean of the intensity for that miRNA t - moderated t-statistic t pval - p value of the t-statistic F - F statistic (null hypothesis:  $C_i = C_j$ , for all contrasts  $i, j$ ) adj.F.pval - F p value adjusted by 'MTestmethod' fdr.F.pval - F p value adjusted by fdr

The html files, both for the 'separated' and 'nestedF' method, includes only the selected as significant genes.

## Author(s)

Pedro Lopez-Romero

## References

Smyth, G. K. (2005). Limma: linear models for microarray data. In: 'Bioinformatics and Computational Biology Solutions using R and Bioconductor'. R. Gentleman, V. Carey, S. Dudoit, R. Irizarry, W. Huber (eds), Springer, New York, pages 397–420.

miRBase: the home of microRNA data <http://microrna.sanger.ac.uk/>

## See Also

A 'uRNAList' example containing processed data is in ddPROC and an overview of how the processed data is produced is given in filterMicroRna. The ExpressionSet object can be generated using esetMicroRna An overview of miRNA differential expression analysis is given in basicLimma An example of how to get the 'TestResults' object is in getDecideTests

## Examples

```
data(targets.micro)
data(ddPROC)
esetPROC=esetMicroRna(ddPROC, targets.micro, makePLOT=FALSE)
```



```

levels.treatment=levels(factor(targets.micro$Treatment))
treatment=factor(as.character(targets.micro$Treatment),
  levels=levels.treatment)

levels.subject=levels(factor(targets.micro$Subject))
subject=factor(as.character(targets.micro$Subject),
  levels=levels.subject)

design=model.matrix(~ -1 + treatment + subject )

CM=cbind(MSC_AvsMSC_B=c(1,-1,0,0),
  MSC_AvsMSC_C=c(1,0,-1,0))

fit2=basicLimma(esetPROC,design,CM,verbose=TRUE)

DE=getDecideTests(fit2,
  DEmethod="separate",
  MTestmethod="BH",
  PVcut=0.10)

significantMicroRna(esetPROC,
  ddPROC,
  targets.micro,
  fit2,
  CM,
  DE,
  DEmethod="separate",
  MTestmethod="BH",
  PVcut=0.10,
  Mcut=0,
  verbose=TRUE)

```

---

summary.uRNAList

*Summaries of Microarray Data Objects*


---

## Description

Briefly summarize microarray data objects.

## Usage

```
## S3 method for class 'uRNAList'
summary(object, ...)
```

## Arguments

|        |                              |
|--------|------------------------------|
| object | an object of class uRNAList  |
| ...    | other arguments are not used |

## Details

The data objects are summarized as if they were lists, i.e., brief information about the length and type of the components is given. This function and this file, has been borrowed from the files created by Gordon Smyth for the limma package.

**Value**

A table.

**Author(s)**

Pedro Lopez-Romero

---

targets.micro

*Example of target file*

---

**Description**

Example of target file

**Usage**

```
data(targets.micro)
```

**Format**

A data frame with 4 observations on the following 5 variables.

FileName names of the Files Ast.txt Bst.txt Aunst.txt Bunst.txt

Treatment Assigns level for Treatment Effect to each File (mandatory)

GErep a numeric vector tha numerates the FACTOR of the Treatment Effect (mandatory)

Subject Assigns level for Subject Effect to each File

**Details**

It is a tab-delimited text format file. The target file is created by the user with the intention of carrying out a differential expression analysis in future steps using 'limma'. Here is where the factors that are going to be included in the linear model that is fitted to each gen are specified. The targets file assigns each data file to a particular experimental conditions. First column 'FileName' is mandatory and includes the image data files names. Second column 'Treatment' is also mandatory and includes the treament effect. Third column 'GErep' is also mandatory, and includes the treatment effect in a numeric code, from 1 to n, being n the number of Treatment effect levels.

**Author(s)**

Pedro Lopez-Romero

**References**

Gordon K. Smyth, M. Ritchie, N. Thorne, J. Wettenhall (2007). limma: Linear Models for Microarray Data User's Guide.

**See Also**

readTargets

---

tgsMicroRna

---

*Getting the Total Gene Signal*

---

## Description

The function creates an `uRNAList` containing the `TotalGeneSignal` computed by the Agilent Feature Extraction software. This signal can be used for the statistical analysis after a possible normalization step.

## Usage

```
tgsMicroRna(dd, offset, half, makePLOT=FALSE, verbose=FALSE)
```

## Arguments

|                       |                                                                                 |
|-----------------------|---------------------------------------------------------------------------------|
| <code>dd</code>       | <code>uRNAList</code> , containing the output from <code>readMicroRnaAFE</code> |
| <code>offset</code>   | integer. To use this option set <code>half = FALSE</code>                       |
| <code>half</code>     | logical, if <code>TRUE</code> half option is used                               |
| <code>makePLOT</code> | logical, if <code>TRUE</code> QC plots with the Total Gene Signal are displayed |
| <code>verbose</code>  | logical, if <code>TRUE</code> prints out some summary results                   |

## Details

The function creates a `uRNAList` object that contains in the `uRNAList$TGS`, `uRNAList$TPS`, `uRNAList$meanS` & `uRNAList$procS` the Total Gene Signal (TGS) as computed by the Agilent Feature Extraction algorithms. This TGS is not in  $\log_2$  scale. All the replicated genes have the same estimated TGS, and the function simply picks one gene from each set of replicated genes. To maintain the format of the `uRNAList`, every selected gene retains a probe name attached to them. This probe name is not meaningful any more, since the signal corresponds to the total gene signal and not to the probe signal. The TGS processed by AFE contains some negative values. To get signals with positive values we can either add a positive small constant to all the signals (`offset`) or we can select the 'half' option, which set to 0.5 all the values that are smaller than 0.5. To use the `offset` option we have to set `half=FALSE`, otherwise the half method is used by default. The `offset` option, adds to each signal the quantity  $(\text{abs}(\min(\text{ddTGS}\$TGS)) + \text{offset})$ , where `ddTGS$TGS` is the matrix that contains the `TotalGeneSignal`.

## Value

`uRNAList` containing the `TotalGeneSignal` computed by the Agilent Feature Extraction software. Optionally, it can generate a boxplot, a density plot and a MA plot with the Total Gene Signal.

## Author(s)

Pedro Lopez-Romero

## References

Agilent Feature Extraction Reference Guide <http://www.Agilent.com>

Examples

```
data(dd.micro)
data(targets.micro)
ddTGS=tgsgMicroRna(dd.micro,half=TRUE,makePLOT=FALSE,verbose=FALSE)
```

---

|                   |                                     |
|-------------------|-------------------------------------|
| tgsgNormalization | <i>Normalization Between Arrays</i> |
|-------------------|-------------------------------------|

---

Description

Normalization between arrays of the Total Gene Signal. The function is a wrapper of the 'limma' 'normalizeBetweenArrays' with ('none','quantile','scale') methods

Usage

```
tgsgNormalization(ddTGS, NORMmethod = "quantile", makePLOTpre = FALSE, makePLOTpost = FALSE, targets
```

Arguments

|              |                                                                                                   |
|--------------|---------------------------------------------------------------------------------------------------|
| ddTGS        | uRNAList, containing the output from tgsgMicroRna                                                 |
| NORMmethod   | character specifying the normalization method, 'none','quantile','scale'. The default is quantile |
| makePLOTpre  | logical, if TRUE QC plots with the Raw Total Gene Signal are displayed                            |
| makePLOTpost | logical, if TRUE QC plots with the Normalized Total Gene Signal are displayed                     |
| targets      | data.frame with the target structure                                                              |
| verbose      | logical, if TRUE prints out output                                                                |

Value

A uRNAList object containing the Normalized Total Gene Signal in log 2 scale

Author(s)

Pedro Lopez-Romero

References

Smyth, G. K. (2005). Limma: linear models for microarray data. In: 'Bioinformatics and Computational Biology Solutions Using R and Bioconductor'. R. Gentleman, V. Carey, S. Dudoit, R. Irizarry, W. Huber (eds), Springer, New York, pages 397 - 420

Smyth, G. K., and Speed, T. P. (2003). Normalization of cDNA microarray data. Methods 31, 265-273.

## Examples

```
## Not run:
data(dd.micro)
data(targets.micro)
ddTGS=tgsMicroRna(dd.micro, half=TRUE, makePLOT=FALSE, verbose=FALSE)

ddNORM=tgsNormalization(ddTGS, 'quantile',
                        makePLOTpre=FALSE, makePLOTpost=TRUE, targets.micro, verbose=TRUE)
graphics.off()

## End(Not run)
```

---

uRNAList-class

uRNAList - class

---

## Description

A list-based class (similar to the RGList class in limma package) for the storing of Agilent chips microRNA data uRNAList objects are created by read.agiMicroRna

## uRNAList Components

uRNAList objects are created by new("uRNAList", Newagi) where Newagi is a list. with the following components:

**uRNAList** matrix, 'gTotalGeneSignal'

**uRNAList** matrix, 'gTotalProbeSignal'

**uRNAList** matrix, 'gMeanSignal'

**uRNAList** matrix, 'gProcessedSignal'

**uRNAList** data.frame, 'FileName'

**uRNAList** vector of characters, 'AGilent Probe Name'

**uRNAList** vector of characters, 'microRNA Name'

**uRNAList** vector of integers, '0' = Feature, '1' = Positive control, '-1' = Negative control

**uRNAList** matrix, FLAG to classify signal if 'IsGeneDetected=1' or 'not=0'

**uRNAList** matrix, FLAG to classify signal if 'IsSaturated = 1' or 'not=0'

**uRNAList** matrix, FLAG to classify signal if 'IsFeatPopnOL = 0' or 'not=1'

**uRNAList** matrix, FLAG to classify signal if 'IsFeatNonUnifOL = 0' or 'not=1'

**uRNAList** matrix, gBGMedianSignal

**uRNAList** matrix, gBGUsed

## Author(s)

Pedro Lopez-Romero

**Examples**

```
## Not run:
  data(dd.micro)

## End(Not run)
```

---

|           |                                                                                   |
|-----------|-----------------------------------------------------------------------------------|
| writeEset | <i>Writes the expression data matrix of an ExpressionSet object in a txt file</i> |
|-----------|-----------------------------------------------------------------------------------|

---

**Description**

Writes the expression data matrix of an ExpressionSet object in a file.

**Usage**

```
writeEset(eset, ddPROC, targets, verbose=FALSE)
```

**Arguments**

|         |                                                              |
|---------|--------------------------------------------------------------|
| eset    | An Expression object, normally containing the processed data |
| ddPROC  | An RGList object, normally containing the processed data     |
| targets | data.frame with the targets structure                        |
| verbose | logical, if TRUE prints out output                           |

**Details**

Writes the expression data matrix of an ExpressionSet object in a file.

**Author(s)**

Pedro Lopez-Romero

**See Also**

An 'RGList' example containing processed data is in ddPROC and an overview of how the processed data is produced is given in filterMicroRna. The ExpressionSet object can be generated using esetMicroRna

**Examples**

```
## Not run:
data(ddPROC)
data(targets.micro)
esetPROC=esetMicroRna(ddPROC, targets.micro, makePLOT=TRUE, verbose=FALSE)
writeEset(esetPROC, ddPROC, targets.micro, verbose=TRUE)

## End(Not run)
```

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