# affycomp

October 5, 2010

SD

SD Assessment Functions

# Description

These functions create assessments, figures, and tables for expression standard errors

# Usage

### **Arguments**

a list of lists with the necessary components to create the Figure. See details.

method.names a character vector with the names of the expression measures methodologies being compared.

main title of the Figure.

# **Details**

This uses the dilution data. The exprsets need to have standard error estimates in the assayDataElement (exprset, Read the vignette for more details. The functions work similarly to those assessing expression measures.

All these files need the result of assessDD

### Value

Depends on the call.

### Author(s)

#### **Examples**

```
data(rma.sd.assessment) ##this was produced with affycomp.assess
data(lw.sd.assessment) ##this one too
affycomp.compfig7(list(rma.sd.assessment,lw.sd.assessment))
affycomp.figure7(rma.sd.assessment)
```

```
affycomp.compfigs.auxiliary

Auxiliary functions to create comparitive Figures
```

### **Description**

These functions are auxiliary function to affycompPlot. These Figures are used to compare expression measures. They take lists with components created by the assessDilution and assessSpikeIn functions.

#### Usage

```
affycomp.compfig2(l, method.names = as.character(1:length(l)),
                  add.legend = TRUE, main = "Figure 2")
affycomp.compfig3(l, method.names = as.character(1:length(l)),
                  main = "Figure 3")
affycomp.compfig4a(l, method.names = as.character(1:length(l)),
                   add.legend = TRUE, main = "Figure 4a")
affycomp.compfig4b(1, method.names = as.character(1:length(1)),
                   add.legend = TRUE, main = "Figure 4b")
affycomp.compfig4c(l, method.names = as.character(1:length(l)),
                   add.legend = TRUE, rotate=TRUE, main = "Figure 4c")
affycomp.compfig5a(1, method.names = as.character(1:length(1)),
                  add.legend = TRUE, main = "Figure 5a", maxfp=100)
affycomp.compfiq5b(1, method.names = as.character(1:length(1)),
                  add.legend = TRUE, main = "Figure 5b", maxfp=100)
affycomp.compfig5cde(1, method.names = as.character(1:length(1)),
                  add.legend = TRUE, main = "Figure 5c", maxfp=100,
                  type=c("low", "med", "high"))
affycomp.compfig5c(l, method.names = as.character(1:length(l)),
                  add.legend = TRUE, main = "Figure 5c", maxfp=100)
affycomp.compfig5d(1, method.names = as.character(1:length(1)),
                  add.legend = TRUE, main = "Figure 5d", maxfp=100)
affycomp.compfig5e(l, method.names = as.character(1:length(l)),
                  add.legend = TRUE, main = "Figure 5e", maxfp=100)
```

# **Arguments**

a list of lists with the necessary components to create the Figure. See details.

method.names a character vector with the names of the expression measures methodologies being compared.

add.legend logical. If TRUE a legend is added.

main title of the Figure.

rotate in the case of compfig4c one can eiher show the actual local slopes or the bias (local slope minus 1).

maxfp range of the false positives in ROC will be from 0 to maxfp

compfig5cdef is the engine for 5c, 5d, and 5e. type tells is which of these 4 to

run.

### **Details**

type

These are similar to the functions defined in affycomp.figures.auxiliary. Main difference is that here you send lists with the result of the assessment functions as components.

#### Value

Figures are produced.

### Author(s)

Rafael A. Irizarry

### **Examples**

```
data(rma.assessment)
data(mas5.assessment)
affycomp.compfig2(list(rma.assessment$Dilution, mas5.assessment$Dilution))
affycomp.compfig3(list(rma.assessment$Dilution, mas5.assessment$Dilution))
affycomp.compfig4a(list(rma.assessment$Signal, mas5.assessment$Signal))
affycomp.compfig4b(list(rma.assessment$Dilution, mas5.assessment$Dilution))
affycomp.compfig5a(list(rma.assessment$FC, mas5.assessment$FC))
affycomp.compfig5b(list(rma.assessment$FC, mas5.assessment$FC))
```

```
affycomp.figures.auxiliary

Auxiliary functions to create Figures
```

### **Description**

These functions are auxiliary function to affycompPlot. These Figures are used to assess an expression measure. They take components created by the assessDilution and assessSpikeIn functions.

#### Usage

```
affycomp.figure1(l,main="Figure 1",xlim=NULL,ylim=NULL)
affycomp.figure1b(l,main="Figure 1b",xlim=NULL,ylim=NULL,cex=0.85,all=FALSE)
affycomp.figure2(l,main="Figure 2")
affycomp.figure2b(l,main="Figure 2b")
affycomp.figure3(l, main = "Figure 3")
affycomp.figure4a(l, main = "Figure 4a",equal.lims=FALSE)
affycomp.figure4b(l, main = "Figure 4b")
affycomp.figure5a(l, main = "Figure 5a",maxfp=100)
affycomp.figure5b(l, main = "Figure 5b",maxfp=100)
affycomp.figure5c(l, main = "Figure 5c",maxfp=100)
affycomp.figure5d(l, main = "Figure 5d",maxfp=100)
affycomp.figure5e(l, main = "Figure 5d",maxfp=100)
affycomp.figure5e(l, main = "Figure 5e",maxfp=100)
affycomp.figure6a(l, main = "Figure 6a",xlim = NULL, ylim = NULL)
affycomp.figure6b(l, main = "Figure 6b",xlim = NULL, ylim = NULL)
```

#### **Arguments**

1	A list with the necessary components to create the Figure. See details.		
main	Title for the Figure.		
maxfp	range of the false positives in ROC will be from 0 to maxfp		
xlim	x-axis limits.		
ylim	y-axis limits.		
cex	size of numbers in figure 1b.		
all	logical. If TRUE all spikeins are shown. Otherwise, only those resulting in smaller, realistic, fold changes are shown.		
equal.lims	logical. If TRUE the limits of x-axis and y-axis will have same range.		
rotate	in the case of compfig4c one can eiher show the actual local slopes or the bias (local slope minus 1).		

#### **Details**

Read the vignette for more details on what each Figure is. You can read assessSpikeIn and assessDilution to see which assessments are needed.

### Value

Figures are produced.

#### Author(s)

Rafael A. Irizarry

### **Examples**

```
data(rma.assessment)
affycomp.figure1(rma.assessment$MA)
affycomp.figure2(rma.assessment$Dilution)
```

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```
affycomp.figure3(rma.assessment$Dilution)
affycomp.figure4a(rma.assessment$Signal)
affycomp.figure4b(rma.assessment$Dilution)
affycomp.figure5a(rma.assessment$FC)
affycomp.figure5b(rma.assessment$FC2)
affycomp.figure6a(rma.assessment$FC)
affycomp.figure6b(rma.assessment$FC)
```

affycompTable

Expression Assessment Table

### **Description**

These functions takes as an argument the output of the assessment functions.

# Usage

```
affycompTable(...,Table=NULL, assessment.list=NULL, method.names=NULL)

tableAll(...,assessment.list=NULL, method.names=NULL)

tableDilution(l, method.names=NULL)

tableFC(l, method.names=NULL)

tableFC2(l, method.names=NULL)

tableSignal(l, method.names=NULL)

tableLS(l, method.names=NULL)

tableSpikeInSD(l, method.names=NULL)

tableMA2(l, method.names=NULL)

tableOverallSNR(...,assessment.list=NULL, method.names=NULL, ngenes=12626, rank=TRUE)
```

## **Arguments**

6 affycompPlot

#### **Details**

Read the vignette for more details on what the entries of the table are. affycompTable has a few entries per graph. tableAll has more entries. Once an assessment is used this function knows what to do. You can call any of the assessment functions described in assessSpikeIn, assessDilution, assessSD, assessLS, assessMA2, and assessSpikeInSD.

Note tableRanks and tableOverallSNR work on the results from assessSpikeIn2.

#### Value

A matrix. One column per each method and one row for each comparison. tableOverallSNR is an exception. Where rows represnt methods.

### Author(s)

Rafael A. Irizarry

### **Examples**

```
data(rma.assessment) ##this was produced with affycomp.assess
data(mas5.assessment) ##this one too
tmp <- affycompTable(mas5.assessment,rma.assessment)
format(tmp,digit=2)</pre>
```

affycompPlot

Assessment Plots

### Description

Function that makes assessment plot

# Usage

affycompPlot 7

#### **Arguments**

```
lists produced by the assessment functions (one for each method) separated by
                  commas.
assessment.list
                  Alternatively, one can also send a list of lists produced by one of the assessment
                  functions.
method.names A character vector with the names of the epxression measure methodology.
figure 1. xlim x-axis lim used for the plots in Figure 1.
figure1.ylim y-axis lim used for the plots in Figure 1.
figure1b.xlim
                  x-axis lim used for the plots in Figure 1b.
figure1b.ylim
                  y-axis lim used for the plots in Figure 1b.
figure6a.xlim
                  x-axis lim used for the plots in Figure 6a.
figure6a.ylim
                  y-axis lim used for the plots in Figure 6a.
figure6b.xlim
                  x-axis lim used for the plots in Figure 6b.
figure6b.ylim
                  y-axis lim used for the plots in Figure 6b.
1
                  list with assessment lists as components.
                  a dummy variable to know what function call to create.
what
```

### **Details**

Read the vignette for more details on what each Figure is. Once an assessment is used this function knows what to do. You can call any of the assessment functions described in assessSpikeIn, assessDilution and assessSD.

affycomp.figures, affycomp.figure.calls, affycomp.compfigs.calls are auxiliary functions.

### Value

Figures are produced.

#### Author(s)

Rafael A. Irizarry

### **Examples**

```
data(rma.assessment)
data(mas5.assessment)
affycompPlot(rma.assessment, mas5.assessment)
affycompPlot(rma.assessment$FC)
affycompPlot(rma.assessment$Dilution, mas5.assessment$Dilution)
affycompPlot(rma.assessment$Dilution, mas5.assessment$Dilution)
affycompPlot(rma.assessment$Signal, mas5.assessment$Signal)
affycompPlot(rma.assessment$Dilution, mas5.assessment$Dilution)
affycompPlot(rma.assessment$Dilution, mas5.assessment$Dilution)
```

8 assess Dilution

assessAll	
assessati	

Assessment functions

### **Description**

Assessment functions. Takes a couple of ExpressionSet, one for spike in another for the dilution and returns a list with necessary information to create assessment plots.

# Usage

```
assessAll(d,s,method.name=NULL,verbose=TRUE)
affycomp(d,s,method.name=NULL,verbose=TRUE,return.it=TRUE)
```

### Arguments

А	An Evnrassion Sat (	containing th	e expression meas	sures for the Gene Logic's

dilution experiment.

s An ExpressionSet containing expression measures for the Affymetrix's

spike-in experiment.

method.name Name of expression measure being assessed.

verbose verbosity flag.

return.it if TRUE returns assessment lists.

### **Details**

 ${\tt assessAll\ performs\ assessments\ for\ Figures\ 1-6}.\ It\ is\ a\ wrapper\ for\ assessDilution\ and\ assessSpikeIn.}$ 

affycomp is a wrapper that does it all... including the plotting and creation of table.

### Value

Lists with the necessary information to make the Figures.

# Author(s)

Rafael A. Irizarry

assessDilution

Assessment functions for Dilution Data

### **Description**

Assessment function. Takes an ExpressionSet and returns a list with necessary information to create assessment plots.

# Usage

```
assessDilution(exprset,method.name=NULL)
```

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### **Arguments**

exprset An ExpressionSet containing expression measures for Affymetrix's spike-

in experiment.

method.name Name of expression measure being assessed.

#### **Details**

 ${\tt assessDilution}\ performs\ the\ assessment\ for\ the\ plots\ related\ to\ Dilution\ (Figures\ 2,\ 3,\ 4b)$ 

### Value

Lists with the necessary information to make the Figures.

### Author(s)

Rafael A. Irizarry

assessSD

SD Assessment functions

### **Description**

Assessment function for standard deviation estimates. Takes a dilution data ExpressionSet and returns a list with necessary information to create assessment plot.

# Usage

```
assessSD(exprset, method.name=NULL, logx=FALSE)
```

### **Arguments**

exprset An ExpressionSet containing expression measures for Affymetrix's spike-

in experiment.

method.name Name of expression measure being assessed.

logx Logical. If TRUE expression is logged for plot. See details.

# **Details**

assessD does the assessment for Figure 7. This requires the ExpressionSet to have standard error estimates for the expression measure. Some expression (such as dChip) will have SEs for the original scale. Others, like RMA will have them for the log scale. For original scales, making logx=TRUE is recommended.

### Value

Lists with the necessary information to make the Figures.

### Author(s)

10 assessSpikeIn

assessSpikeIn Assessment functions for Spike In Data
--

# **Description**

These functions are assessment functions. Each takes an ExpressionSet and returns a list with necessary information to create assessment plots.

# Usage

```
assessSpikeIn(s,method.name=NULL,verbose=TRUE)
assessMA(exprset,method.name=NULL)
assessSignal(exprset,method.name=NULL)
assessFC(exprset,method.name=NULL)
assessFC2(exprset,method.name=NULL)
```

### **Arguments**

s An ExpressionSet containing expression measures for Affymetrix's spike-

in experiment.

exprset An ExpressionSet containing expression measures for Affymetrix's spike-

in experiment.

method.name Name of expression measure being assessed.

verbose logical. If TRUE show messages.

#### **Details**

assessMA performs the assessment for the MA-plot (Figure 1), assessSignal performs the assessment for signal detection plot (Figure 4a), assessFC performs assessments used by fold-change related plots (Figures 5a, 6a, 6b). assessFC2 is for the ROC for genes with nominal fold changes of 2 (Figure 5b). assessSpikeIn is a wrapper for all these and returns a list of lists.

#### Value

Lists with the necessary information to make the Figures.

# Author(s)

assessSpikeIn2

assessSpikeIn2	New Assessment functions for Spike In Data
----------------	--

# Description

These functions are assessment functions. Each takes an ExpressionSet and returns a list with necessary information to create assessment plots.

# Usage

```
assessSpikeIn2(s, method.name=NULL, verbose=TRUE)
assessSpikeInSD(exprset, method.name=NULL, span=1/3)
assessLS(exprset, method.name=NULL)
assessMA2(exprset, method.name=NULL)
```

# Arguments

S	An ExpressionSet containing expression measures for Affymetrix's spike-in experiment.
exprset	An ExpressionSet containing expression measures for Affymetrix's spike-in experiment.

method.name Name of expression measure being assessed.

verbose logical. If TRUE show messages.

span span used in call to loess.

### **Details**

assessMA2 performs the assessment for the second MA-plot (Figure 1b), assessLS performs the assessment for signal detection plot (Figure 4c), assessMA2 also performs assessments used by fold-change related plots (Figures 5c-f). assessSpikeInSD is for the standard deviation assessment of Figure 2b. assessSpikeIn2 is a wrapper for all these and returns a list of lists.

# Value

Lists with the necessary information to make the Figures.

### Author(s)

12 exprset.log

dilution.phenodata Phenotypic Information for Dilution Study

# **Description**

This object is of class phenoData with necessary information for the assessemnts.

#### **Usage**

```
data(dilution.phenodata)
```

#### **Format**

An object of class phenoData

#### **Source**

Two sources of cRNA A (human liver tissue) and B (Central Nervous System cell line) have been hybridized to human array (HGU95Av2) in a range of proportions and dilutions. This object described these.

For more information see Irizarry, R.A., et al. (2001) http://www.biostat.jhsph.edu/~ririzarr/papers/index.html

exprset.log

Take log base 2 of Expression

# Description

Take log base 2 of the expression matrix in an ExpressionSet

# Usage

```
exprset.log(exprset)
```

### **Arguments**

exprset

ExpressionSet

### **Details**

This functions takes log base 2 of the expression matrix in an ExpressionSet. Negatives are converted to the smallest non-negative entry.

# Value

An ExpressionSet

# Author(s)

```
hgu133a.spikein.phenodata
```

phenotypic information for HGU133A spike in study

# Description

This objact is of class phenoData with necessary information for the assessemnts.

# Usage

```
data(hgu133a.spikein.phenodata)
```

### **Format**

An object of class phenoData

#### Source

This comes from an experiments where 16 different cRNA fragments have been added to the hybridization mixture of the GeneChip arrays at different pM concentrations. For more information see Irizarry, R.A., et al. (2001) http://www.biostat.jhsph.edu/~ririzarr/papers/index.html

lw.sd.assessment

An example of the result of an SD assessment

### **Description**

The Dilution files were processed with the dChip package (using PM-only) and then the function assessD was applied.

# Usage

```
data(lw.sd.assessment)
```

### **Format**

A list.

14 readin

mas5.assessment An ex

An example of the result of the assessments

## **Description**

The Dilution and both (HGU95 and HGU133) Spike-in cel files were processed with MAS 5.0 software and then the functions assessAll and assessSpikeIn2 were applied.

# Usage

```
data(rma.assessment)
data(rma.assessment.133)
data(rma.assessment2)
data(rma.assessment2.133)
```

#### **Format**

A list of list.

readin

Read Expression Date Sets

### **Description**

Reads a comma-delimited file containing the expression values of the dilution and spike-in data sets and creates a ExpressionSet

# Usage

```
read.dilution(filename)
read.spikein(filename, cdfName=c("hgu95a","hgu133a"), remove.xhyb=TRUE)
read.newspikein(filename)
```

### **Arguments**

filename character containing the filename to be read.

cdfName are we reading data from the hgu95a or hgu133a spike-in experiment?

remove.xhyb logical. If TRUE possible cross hybridizers are removed from the HGU133A

spikein. See remove.hgu133a.xhyb.

# **Details**

The file to be read must be comma-delimited with the first row containing the cel filenames (case sensitive). The first column must be the Affymetrix gene identifiers. read.dilution will put things in the right place.

read.newspikein is a wrapper to read results from the hgu133a spikein experiment.

### Value

An ExpressionSet.

remove.hgu133a.xhyb

### Author(s)

Rafael A. Irizarry

```
remove.hgu133a.xhyb

*Remove crosshybridizers*
```

### **Description**

This functions removes possible cross hybridizers from Affymetrix HGU133A spike-in experiment

### Usage

```
remove.hgu133a.xhyb(s, bp = c("200", "150", "100"))
```

# Arguments

```
an ExpressionSet containing the HGU133A spike-in
```

bp number of base pair matches needed to define a possible cross hybridizer

### **Details**

Some details are contained in the help file for hgu133a.spikein.xhyb

### Value

An ExpressionSet with probeset removed

# See Also

```
hgu133a.spikein.xhyb
```

rma.assessment

An example of the result of an assessment

# Description

The Dilution and Spike-in cel files were processed with the affy version 1.0 package rma add-on function and then the functions assessAll and assessSpikeIn2 were applied.

# Usage

```
data(rma.assessment)
data(rma.assessment.133)
data(rma.assessment2)
data(rma.assessment2.133)
```

### **Format**

A list of list.

16 spikein.phenodata

rma.sd.assessment An example of the result of an SD assessment

# Description

The Dilution files were processed with the affy version 1.0 package rma add-on function and then the function assessSD was applied.

### Usage

```
data(rma.sd.assessment)
```

#### **Format**

A list.

spikein.phenodata phenotypic information for spike in study

# Description

This objact is of class phenoData with necessary information for the assessemnts.

# Usage

```
data(spikein.phenodata)
```

### **Format**

An object of class phenoData

### Source

This comes from an experiments where 16 different cRNA fragments have been added to the hybridization mixture of the GeneChip arrays at different pM concentrations. For more information see Irizarry, R.A., et al. (2001) http://www.biostat.jhsph.edu/~ririzarr/papers/index.html

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hgu133a.spikein.xhyb

Cross hybridizers

# Description

Probe Sets likely to crosshybridize to spiked-in probesets in the Affymetrix HGU133A spike in.

This objact is list. Each component of the list contains probeset names of possible crosshybridizers. The sequences of each spiked-in clone were collected and blasted against all HG-U133A target sequences. Target sequences are the ~600bp regions from which probes were selected. Thresholds of 100, 150 and 200bp were used and define the three components of the list.

# Usage

```
data(hgu133a.spikein.xhyb)
```

### **Format**

A list

#### **Source**

Simon Cawley <simon\_cawley@affymetrix.com>

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