affy

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AffyBatch-class Class AffyBatch

Description

This is a class representation for Affymetrix GeneChip probe level data. The main component are the intensities from multiple arrays of the same CDF type. It extends eSet.

Objects from the Class

Objects can be created using the function read.affybatch or the wrapper ReadAffy.

Slots

- **cdfName:** Object of class character representing the name of CDF file associated with the arrays in the AffyBatch.
- nrow: Object of class integer representing the physical number of rows in the arrays.
- **ncol:** Object of class integer representing the physical number of columns in the arrays.
- **assayData:** Object of class AssayData containing the raw data, which will be at minimum a matrix of intensity values. This slot can also hold a matrix of standard errors if the 'sd' argument is set to TRUE in the call to ReadAffy.
- **phenoData:** Object of class AnnotatedDataFrame containing phenotypic data for the samples.
- **annotation** A character string identifying the annotation that may be used for the ExpressionSet instance.

AffyBatch-class

featureData Object of class AnnotatedDataFrame containing feature-level (e.g., probesetlevel) information.

experimentData: Object of class "MIAME" containing experiment-level information. **notes:** Object of class "character" Vector of explanatory text

Extends

Class "eSet", directly.

Methods

cdfName signature(object = "AffyBatch"): **Obtains the cdfName** slot.

pm<- signature (object = "AffyBatch"): replaces the perfect match intensities

pm signature(object = "AffyBatch"): extracts the pm intensities.

mm<- signature (object = "AffyBatch"): replaces the mismatch intensities.

mm signature(object = "AffyBatch"): extracts the mm intensities.

- probes signature(object = "AffyBatch", which): extract the perfect match or mismatch probe intensities. Uses which can be "pm" and "mm".
- **exprs** signature(object = "AffyBatch"): extracts the expression matrix.
- exprs<- signature(object = "AffyBatch", value = "matrix"): replaces the expression matrix.
- se.exprs signature(object = "AffyBatch"): extracts the matrix of standard errors of
 expression values, if available.
- se.exprs<- signature(object = "AffyBatch", value = "matrix"): replaces the matrix of standard errors of expression values.
- [<- signature(x = "AffyBatch"): replaces subsets.</pre>

[signature(x = "AffyBatch"): subsets by array.

- boxplot signature(x = "AffyBatch"): creates a boxplots of log base 2 intensities
 (pm, mm or both). Defaults to both.
- computeExprSet signature(x = "AffyBatch", summary.method = "character"):
 For each probe set computes an expression value using summary.method.
- featureNames signature(object = "AffyBatch"): return the probe set names also referred to as the Affymetrix IDs. Notice that one can not assign featureNames. You must do this by changing the cdfenvs.
- geneNames signature(object="AffyBatch'"): deprecated, use featureNames
- getCdfInfo signature(object = "AffyBatch"): retrieve the environment that defines
 the location of probes by probe set.

image signature(x = "AffyBatch"): creates an image for each sample.

indexProbes signature(object = "AffyBatch", which = "character"): returns a list with locations of the probes in each probe set. The affyID corresponding to the probe set to retrieve can be specified in an optional parameter genenames. By default, all the affyIDs are retrieved. The names of the elements in the list returned are the affyIDs. which can be "pm", "mm", or "both". If "both" then perfect match locations are given followed by mismatch locations.

signature(object = "AffyBatch", which = "missing")(i.e., calling indexProbes
without a "which" argument) is the same as setting "which" to "pm".

- intensity signature(object = "AffyBatch"): extract the exprs slot, i.e. the intensities.
- **length** signature (x = "AffyBatch"): returns the number of samples.
- pmindex signature(object = "AffyBatch"): return the location of perfect matches in the intensity matrix.
- mmindex signature(object = "AffyBatch"): return the location of the mismatch intensities.
- **dim** signature (x = "AffyBatch"): Row and column dimensions.
- ncol signature(x = "AffyBatch"): An accessor function for ncol.
- **nrow** signature (x = "AffyBatch"): an accessor function for nrow.
- normalize signature(object = "AffyBatch"): a method to normalize. The method
 accepts an argument method. The default methods is specified in package options (see the
 main vignette).
- **normalize.methods** signature (object = "AffyBatch"): returns the normalization methods defined for this class. See normalize.
- probeNames signature(object = "AffyBatch"): returns the probe set associated with
 each row of the intensity matrix.
- probeset signature(object = "AffyBatch", genenames=NULL, locations=NULL):
 Extracts ProbeSet objects related to the probe sets given in genenames. If an alternative
 set of locations defining pms and mms a list with those locations should be passed via the
 locations argument.
- bg.correct signature(object = "AffyBatch", method="character") applies background correction methods defined by method.

Note

This class is better described in the vignette.

See Also

related methods merge.AffyBatch, pairs.AffyBatch, and eSet

Examples

```
if (require(affydata)) {
    ## load example
    data(Dilution)
    ## nice print
    print(Dilution)
    pm(Dilution)[1:5,]
    mm(Dilution)[1:5,]
    ## get indexes for the PM probes for the affyID "1900_at"
```

affy-options

```
mypmindex <- pmindex(Dilution,"1900_at")
## same operation using the primitive
mypmindex <- indexProbes(Dilution, which="pm", genenames="1900_at")[[1]]
## get the probe intensities from the index
intensity(Dilution)[mypmindex, ]
description(Dilution) ##we can also use the methods of eSet
sampleNames(Dilution)
abstract(Dilution)
}</pre>
```

affy-options Options for the affy package

Description

Description of the options for the affy package.

Note

The affy package options are contained in the Bioconductor options. The options are:

- use.widgets: a logical used to decide on the default of widget use.
- compress.cel: a logical
- compress.cdf: a logical
- probes.loc: a list. Each element of the list is it self a list with two elements *what* and *where*. When looking for the informations about the locations of the probes on the array, the elements in the list will be looked at one after the other. The first one for which *what* and *where* lead to the matching locations information is used. The element *what* can be one of *package, environment* or *file*. The element *where* depends on the corresponding element *what*.
 - if *package*: location for the package (like it would be for the argument lib.loc for the function library.)
 - if *environment*: an environment to look for the information (like the argument env for the function get).
 - if *file*: a character with the path in which a CDF file can be found.

Examples

```
## get the options
opt <- getOption("BioC")
affy.opt <- opt$affy
## list their names
names(affy.opt)
## set the option compress.cel
affy.opt$compress.cel <- TRUE
options(BioC=opt)
```

AffyRNAdeg

Description

Uses ordered probes in probeset to detect possible RNA degradation. Plots and statistics used for evaluation.

Usage

Arguments

abatch	An object of class AffyBatch-class.	
log.it	A logical argument: If log.it=T, then probe data is log2 tranformed	
rna.deg.obj	Output from AffyRNAdeg	
signif.digits		
	Number of significant digits to show.	
transform	Possible choices are "shift.scale", "shift.only", and "neither". "Shift" vertically staggers the plots for individual chips, to make the display easier to read. "Scale" normalizes so that standard deviation is equal to 1.	
cols	A vector of colors for plot, length = number of chips	
	further arguments for plot function.	

Details

Within each probeset, probes are numbered directionally from the 5' end to the 3' end. Probe intensities are averaged by probe number, across all genes. If log.it=FALSE and transform="Neither", then plotAffyRNAdeg simply shows these means for each chip. Shifted and scaled versions of the plot can make it easier to see.

Value

AffyRNAdeg returns a list with the following components:

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oun		· • •			\sim	~

	names of samples, derived from affy batch object	
means.by.number		
	average intensity by probe position	
ses	standard errors for probe position averages	
slope	from linear regression of means.by.number	
pvalue	from linear regression of means.by.number	

Author(s)

Leslie Cope

affy.scalevalue.exprSet

Examples

```
if (require(affydata)) {
   data(Dilution)
   RNAdeg<-AffyRNAdeg(Dilution)
   plotAffyRNAdeg(RNAdeg)
}</pre>
```

affy.scalevalue.exprSet Scale normalization for expreSets

Description

Normalizes expression values using the method described in the Affymetrix user manual.

Usage

affy.scalevalue.exprSet(eset, sc = 500, analysis="absolute")

Arguments

eset	An ExpressionSet object.
SC	Value at which all arrays will be scaled to.
analysis	Should we do absolute or comparison analysis, although "comparison" is still not implemented.

Details

This is function was implemented from the Affymetrix technical documentation for MAS 5.0. It can be downloaded from the website of the company. Please refer to this document for details.

Value

A normalized ExpressionSet

Author(s)

Laurent

barplot.ProbeSet show a ProbeSet as barplots

Description

displays the probe intensities in a ProbeSet as a barplots

Usage

```
## S3 method for class 'ProbeSet':
barplot(height, xlab = "Probe pair", ylab = "Intensity", main =
NA, col.pm = "red", col.mm = "blue", beside = TRUE, names.arg = "pp",
ask = TRUE, scale, ...)
```

Arguments

height	an object of class ProbeSet
xlab	label for x axis
ylab	label for y axis
main	main label for the figure
col.pm	color for the 'pm' intensities
col.mm	color for the 'mm' intensities
beside	bars beside eachothers or not
names.arg	
ask	ask before ploting the next barplot
scale	put all the barplot to the same scale
	extra parameters to be passed to barplot

Examples

```
if (require(affydata)) {
  data(Dilution)
  gn <- geneNames(Dilution)
  pps <- probeset(Dilution, gn[1])[[1]]
  barplot.ProbeSet(pps)
}</pre>
```

bg.adjust

Description

An internal function to be used by bg.correct.rma.

Usage

```
bg.adjust(pm, n.pts = 2^14, ...)
bg.parameters(pm, n.pts = 2^14)
```

Arguments

pm	a pm matrix
n.pts	number of points to use in call to ${\tt density}.$
	extra arguments to pass to bg.adjust.

Details

Assumes PMs are a convolution of normal and exponentional. So we observe X+Y where X is backround and Y is signal. bg.adjust returns E[Y|X+Y, Y>0] as our backround corrected PM. bg.parameters provides adhoc estimates of the parameters of the normal and exponential distributions.

Value

a matrix

See Also

bg.correct.rma

bg.correct Background Correction

Description

Background corrects probe intensities in an object of class AffyBatch.

Usage

```
bg.correct(object, method, ...)
bg.correct.rma(object, ...)
bg.correct.mas(object, griddim)
bg.correct.none(object, ...)
```

Arguments

object	An object of class AffyBatch.
method	A character that defines what background correction method will be used. Available methods are given by bg.correct.methods.
griddim	grid dimension used for mas background estimate. The array is divided into griddm equal parts. Default is 16.
•••	arguments to pass along to the engine function.

Details

The name of the method to apply must be double-quoted. Methods provided with the package are currently:

- bg.correct.none: returns object unchanged.
- bg.correct.chipwide: noise correction as described in a 'white paper' from Affymetrix.
- bg.correct.rma: the model based correction used by the RMA expression measure.

They are listed in the variable bg.correct.methods. The user must supply the word after "bg.correct", i.e none, subtractmm, rma, etc...

More details are available in the vignette.

R implementations similar in function to the internal implementation used by bg.correct.rma are in bg.adjust.

Value

An AffyBatch for which the intensities have been background adjusted. For some methods (RMA), only PMs are corrected and the MMs remain the same.

Examples

```
if (require(affydata)) {
  data(Dilution)

  ##bgc will be the bg corrected version of Dilution
  bgc <- bg.correct(Dilution, method="rma")

  ##This plot shows the tranformation
  plot(pm(Dilution)[,1],pm(bgc)[,1],log="xy",
  main="PMs before and after background correction")
}</pre>
```

cdfenv.example Example cdfenv

Description

Example cdfenv (environment containing the probe locations).

Usage

```
data(cdfenv.example)
```

cdfFromBioC

Format

An environment cdfenv.example containing the probe locations

Source

Affymetrix CDF file for the array Hu6800

cdfFromBioC

Functions to obtain CDF files

Description

A set of functions to obtain CDF files from various locations.

Usage

```
cdfFromBioC(cdfname, lib = .libPaths()[1], verbose = TRUE)
cdfFromLibPath(cdfname, lib = NULL, verbose=TRUE)
cdfFromEnvironment(cdfname, where, verbose=TRUE)
```

Arguments

cdfname	The CDF desired
lib	Directory to install the CDF package to
where	What environment to search
verbose	Controls extra output

Details

These functions all take a requested CDF environment name and will attempt to locate that environment in the appropriate location (a package's data directory, as a CDF package in the .libPaths(), from a loaded environment or on the Bioconductor website. If the environment can not be found, it will return a list of the methods tried that failed.

Value

The CDF environment or a list detailing the failed locations.

Author(s)

Jeff Gentry

cleancdfname

Description

This function converts Affymetrix's names for CDF files to the names used in the annotation pacakge and in all Bioconductor.

Usage

cleancdfname(cdfname, addcdf = TRUE)

Arguments

cdfname	A character denoting Affymetrix'x CDF file name
addcdf	A logical. If TRUE it adds the string "cdf" at the end of the cleaned CDF
	name. This is used to name the cdfenvs packages.

Details

This function takes a CDF filename obtained from an Affymetrix file (from a CEL file for example) and convert it to a convention of ours: all small caps and only alphanumeric characters. The details of the rule can be seen in the code. We observed exceptions that made us create a set of special cases for mapping CEL to CDF. The object mapCdfName holds information about these cases. It is a data.frame of three elements: the first is the name as found in the CDF file, the second the name in the CEL file and the third the name in bioconductor. mapCdfName can be loaded using data (mapCdfName).

Value

A character

Examples

```
cdf.tags <- c("HG_U95Av2", "HG-133A")
for (i in cdf.tags)
  cat(i, "becomes", cleancdfname(i), "\n")</pre>
```

debug.affy123 Debugging Flag

Description

For developmental use only

expresso

Description

Goes from raw probe intensities to expression values

Usage

```
expresso(
        afbatch,
    # background correction
        bg.correct = TRUE,
        bgcorrect.method = NULL,
        bgcorrect.param = list(),
    # normalize
        normalize = TRUE,
        normalize.method = NULL,
        normalize.param = list(),
    # pm correction
        pmcorrect.method = NULL,
        pmcorrect.param = list(),
    # expression values
        summary.method = NULL,
        summary.param = list(),
        summary.subset = NULL,
    # misc.
        verbose = TRUE,
        widget = FALSE)
```

Arguments

```
afbatch
                 An AffyBatch object
                 a boolean to express whether background correction is wanted or not.
bg.correct
bgcorrect.method
                 the name of the background adjustment method
bgcorrect.param
                 a list of parameters for bgcorrect.method (if needed/wanted)
normalize
                 normalization step wished or not.
normalize.method
                 the normalization method to use
normalize.param
                 a list of parameters to be passed to the normalization method (if wanted).
pmcorrect.method
                 the name of the PM adjustement method
pmcorrect.param
                 a list of parameters for pmcorrect.method (if needed/wanted)
summary.method
                 the method used for the computation of expression values
```

Summary .para	
	a list of parameters to be passed to the summary.method (if wanted).
summary.subs	et
	a list of 'affyids'. If ${\tt NULL},$ a expression summary value is computed for everything on the chip.
verbose	logical value. If TRUE it writes out some messages.
widget	a boolean to specify the use of widgets (the package tkWidget is required).

Details

Some arguments can be left to NULL if the widget=TRUE. In this case, a widget pops up and let the user choose with the mouse. The arguments are: AffyBatch, bgcorrect.method, normalize.method, pmcorrect.method and summary.method.

For the mas 5.0 and 4.0 methods ones need to normalize after obtaining expression. The function affy.scalevalue.exprSet does this.

For the Li and Wong summary method notice you will not get the same results as you would get with dChip. dChip is not open source so it is not easy to reproduce. Notice also that this iterative algorithm will not always converge. If you run the algorithm on thousands of probes expect some non-convergence warnings. These are more likely when few arrays are used. We recommend using this method only if you have 10 or more arrays. Please refer to the fit.li.wong help page for more details.

Value

An object of class ExpressionSet, with an attribute pps.warnings as returned by the method computeExprSet.

See Also

AffyBatch

Examples

expressoWidget A widget for users to pick correction methods

Description

This widget is called by expresso to allow users to select correction methods that will be used to process affy data.

cummary naram

expressoWidget

Usage

```
expressoWidget(BGMethods, normMethods, PMMethods, expMethods, BGDefault,
normDefault, PMDefault, expDefault)
```

Arguments

BGMethods	BGMethods a vector of character strings for the available methods that can be used as a background correction method of affy data
normMethods	normMethods a vector of character strings for the available methods that can be used as a normalization method of affy data
PMMethods	PMMethods a vector of character strings for the available methods that can be used as a PM correction method of affy data
expMethods	expMethodsa vector of character strings for the available methods that can be used as a summary method of affy data
BGDefault	BGDefault a character string for the name of a default background correction method
normDefault	normDefault a character string for the name of a default normalization method
PMDefault	PMDefault a character string for the name of a default PM correction method
expDefault	expDefault a character string for the name of a default summary method

Details

The widget will be invoked when expresso is called with argument "widget" set to TRUE. Default values can be changed using the drop down list boxes. Double clicking on an option from the dropdown list makes an selection. The first element of the list for available methods will be the default method if no default is provided.

Value

The widget returns a list of selected correction methods.

BG	background correction method
NORM	normalization method
PM	PM correction method
EXP	summary method

Author(s)

Jianhua Zhang

References

Documentations of affy package

See Also

expresso

Examples

```
if(interactive()){
  require(widgetTools)
  expressoWidget(c("mas", "none", "rma"), c("constant", "quantiles"),
  c("mas", "pmonly"), c("liwong", "playerout"))
}
```

fit.li.wong Fit Li and Wong Model to a Probe Set

Description

Fits the model described in Li and Wong (2001) to a probe set with I chips and J probes.

Usage

Arguments

data.matrix	
	the PM-MM value for probe pair j in chip i. Another possible use, is to use PM
	instead of PM-MM.
remove.outli	ers
	logical value indicating if the algorithm will remove outliers according to the
	procedure described in Li and Wong (2001).
large.thresh	old
	used to define outliers.
normal.array	.quantile
	quantile to be used when determining what a normal SD is. probes or chips hav-
	ing estimates with SDs bigger than the quantile normal.array.quantile
	of all SDs x large.threshold
normal.resid	.quantile
	any residual bigger than the normal.resid.quantile quantile of all resid-
	uals x large.threshold is considered an outlier
large.variat	ion
	any probe or chip describing more than this much total variation is considered
	an outlier
outlier.fraction	
	this is the maximum fraction of single outliers that can be in the same probe or
	chip.
	1

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fit.li.wong

delta	numerical value used to define the stopping criterion.
maxit	maximum number of iterations when fitting the model.
outer.maxit	maximum number of iterations of defined outliers.
verbose	logical value. If TRUE information is given of the status of the algorithm.

Details

This is Bioconductor's implementation of the Li and Wong algorithm. The Li and Wong PNAS 2001 paper was followed. However, you will not get the same results as you would get with dChip. dChip is not open source so it is not easy to reproduce.

Notice that this iterative algorithm will not always converge. If you run the algorithm on thousands of probes expect some non-convergence warnings. These are more likely when few arrays are used. We recommend using this method only if you have 10 or more arrays.

Please refer to references for more details.

Value

li.wong returns a vector of expression measures (or column effects) followed by their respective standard error estimates. It was designed to work with express which is no longer part of the package.

fit.li.wong returns much more. Namely, a list containing the fitted parameters and relevant information.

theta	fitted thetas.	
phi	fitted phis.	
sigma.eps	estimated standard deviation of the error term.	
sigma.theta	estimated standard error of theta.	
sigma.phi	estimated standard error of phis.	
theta.outlie:	rs	
	logical vector describing which chips (thetas) are considered outliers (TRUE).	
phi.outliers	logical vector describing which probe sets (phis) are considered outliers (TRUE)	
convergencel	logical value. If ${\tt FALSE}$ the algorithm did not converge when fitting the phis and thetas.	
convergence2	logical value. If FALSE the algorithm did not converge in deciding what are outliers.	
iter	number of iterations needed to achieve convergence.	
delta	difference between thetas when iteration stopped.	

Author(s)

Rafael A. Irizarry, Cheng Li, Fred A. Wright, Ben Bolstad

References

Li, C. and Wong, W.H. (2001) Genome Biology 2, 1–11.

Li, C. and Wong, W.H. (2001) Proc. Natl. Acad. Sci USA 98, 31-36.

See Also

li.wong, expresso

Examples

```
x <- sweep(matrix(2^rnorm(600), 30, 20), 1, seq(1, 2, len=30), FUN="+")
fit1 <- fit.li.wong(x)
plot(x[1,])
lines(fit1$theta)</pre>
```

generateExprSet-method

generate a set of expression values

Description

Generate a set of expression values from the probe pair information. The set of expression is returned as an ExpressionSet object.

Usage

```
computeExprSet(x, pmcorrect.method, summary.method, ...)
```

```
generateExprSet.methods()
```

upDate.generateExprSet.methods(x)

Arguments

Х	a AffyBatch holding the probe level informations to generate the expression	
	values, for computeExprSet, and for upDate.generateExprSet.methods it is a	
	character vector	
pmcorrect.method		
	the method used to correct PM values (see section 'details').	
summary.method		
	the method used to generate the expression value (see section 'details').	
	any of the options of the normalization you would like to modify	

Details

An extra argument ids= can be passed. It must be a vector of affids. The expression values will only be computed and returned for these affyids.

The different methods available through this mecanism can be accessed by calling the method generateExprSet.methods with an object of call Cel.container as an argument.

In the Affymetrix design, *MM* probes were included to measure the noise (or background signal). The original algorithm for background correction was to subtract the *MM* signal to the *PM* signal. The methods currently inclluded in the package are "bg.correct.subtractmm", "bg.correct.pmonly" and "bg.correct.adjust".

To alter the available methods for generating ExprSets use upDate.generateExprSet.methods.

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generateExprVal.method.avgdiff

See Also

```
method generateExprSet of the class AffyBatch
expresso
```

Examples

generateExprVal.method.avgdiff

Generate an expression value from the probes informations

Description

Generate an expression from the probes

Usage

```
generateExprVal.method.avgdiff(probes, ...)
generateExprVal.method.medianpolish(probes, ...)
generateExprVal.method.liwong(probes, ...)
generateExprVal.method.mas(probes, ...)
```

Arguments

probes	a matrix of probe intesities with rows representing probes and columns rep-
	resenting samples. Usually pm(probeset) where probeset is a of class
	ProbeSet

```
... extra arguments to pass to the respective function
```

Value

A list containing entries:

exprs	The expression values.
se.exprs	The standard error estimate.

See Also

```
generateExprSet-methods,\code{generateExprSet-methods}, \code{generateExprSet-
methods}, \code{generateExprSet-methods}
```

Examples

```
data(SpikeIn) ##SpikeIn is a ProbeSets
probes <- pm(SpikeIn)
avgdiff <- generateExprVal.method.avgdiff(probes)
medianpolish <- generateExprVal.method.medianpolish(probes)
liwong <- generateExprVal.method.liwong(probes)
playerout <- generateExprVal.method.playerout(probes)
mas <- generateExprVal.method.mas(probes)
concentrations <- as.numeric(sampleNames(SpikeIn))
plot(concentrations,avgdiff$exprs,log="xy",ylim=c(50,10000),pch="a",type="b")
points(concentrations,2^medianpolish$exprs,pch="m",col=2,type="b",lty=2)
points(concentrations,liwong$exprs,pch="l",col=3,type="b",lty=3)
points(concentrations,playerout$exprs,pch="p",col=4,type="b",lty=4)
points(concentrations,mas$exprs,pch="p",col=4,type="b",lty=4)</pre>
```

generateExprVal.method.playerout
 Generate an expression value from the probes informations

Description

Generate an expression from the probes

Usage

generateExprVal.method.playerout(probes, weights=FALSE, optim.method="L-BFGS-E

Arguments

probes	a list of probes slots from PPSet.container
weights	Should the resulting weights be returned ?
optim.method	see parameter 'optim' for the function optim

Details

A non-parametrical method to weight each perfect match probe in the set and to compute a weighted mean of the perfect match values. One will notice this method only makes use of the perfect matches. (see function playerout.costfunction for the cost function).

Value

A vector of expression values.

Author(s)

Laurent <laurent@cbs.dtu.dk> (Thanks to E. Lazaridris for the original playerout code and the discussions about it)

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generateExprVal

References

Emmanuel N. Lazaridis, Dominic Sinibaldi, Gregory Bloom, Shrikant Mane and Richard Jove A simple method to improve probe set estimates from oligonucleotide arrays, Mathematical Biosciences, Volume 176, Issue 1, March 2002, Pages 53-58

generateExprVal Compute a summary expression value from the probes intensities

Description

Compute a summary expression value from the probes intensities

Usage

```
express.summary.stat(x, pmcorrect, summary, ...)
express.summary.stat.methods() # vector of names of methods
upDate.express.summary.stat.methods(x)
```

Arguments

Х	a(ProbeSet
pmcorrect	the method used to correct the PM values before summarizing to an expression value.
summary	the method used to generate the expression value.
•••	other parameters the method might need (see the corresponding methods be-low)

Value

Returns a vector of expression values.

Examples

```
if (require(affydata)) {
   data(Dilution)
   p <- probeset(Dilution, "1001_at")[[1]]
   par(mfcol=c(5,2))
   mymethods <- express.summary.stat.methods()
   nmet <- length(mymethods)
   nc <- ncol(pm(p))
   layout(matrix(c(1:nc, rep(nc+1, nc)), nc, 2), width = c(1, 1))
   barplot(p)
   results <- matrix(0, nc, nmet)
   rownames(results) <- paste("sample", 1:nc)
   colnames(results) <- mymethods
   for (i in 1:nmet) {</pre>
```

justRMA

```
ev <- express.summary.stat(p, summary=mymethods[i], pmcorrect="pmonly")
if (mymethods[[i]] != "medianpolish")
results[, i] <- 2^(ev$exprs)
else
results[, i] <- ev$exprs
}
dotchart(results, labels=paste("sample", 1:nc))
}</pre>
```

hlog

Hybrid Log

Description

Given a constant c this function returns x if x is less than c and sign (x) * (c $* \log (abs (x) / c) + c$) if its not. Notice this is a continuos odd (f(-x)=-f(x)) function with continous first derivative. The main purpose is to perform log transformation when one has negative numbers, for example for PM-MM.

Usage

hlog(x, constant=1)

Arguments

Х	a number.
constant	the constant c (see description).

Details

If constant is less than or equal to $0 \log(x)$ is returned for all x. If constant is infinity x is returned for all x.

Author(s)

Rafael A. Irizarry

justRMA

Read CEL files into an ExpressionSet

Description

Read CEL files and compute an expression measure without using an AffyBatch.

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justRMA

Usage

```
just.rma(..., filenames = character(0),
               phenoData = new("AnnotatedDataFrame"),
               description = NULL,
               notes = "",
               compress = getOption("BioC")$affy$compress.cel,
               rm.mask = FALSE, rm.outliers = FALSE, rm.extra = FALSE,
               verbose=FALSE, background=TRUE, normalize=TRUE,
               bgversion=2, destructive=FALSE, cdfname = NULL)
justRMA(..., filenames=character(0),
             widget=getOption("BioC")$affy$use.widgets,
             compress=getOption("BioC")$affy$compress.cel,
              celfile.path=getwd(),
              sampleNames=NULL,
             phenoData=NULL,
              description=NULL,
              notes="",
              rm.mask=FALSE, rm.outliers=FALSE, rm.extra=FALSE,
             hdf5=FALSE, hdf5FilePath=NULL,verbose=FALSE,
              normalize=TRUE, background=TRUE,
             bgversion=2, destructive=FALSE, cdfname = NULL)
```

Arguments

	file names separated by comma.
filenames	file names in a character vector.
phenoData	a AnnotatedDataFrame object.
description	a MIAME object.
notes	notes.
compress	are the CEL files compressed?
rm.mask	should the spots marked as 'MASKS' set to NA?
rm.outliers	should the spots marked as 'OUTLIERS' set to NA?
rm.extra	if TRUE, then overrides what is in rm.mask and rm.oultiers.
hdf5	use of hdf5? (not available yet)
hdf5FilePath	a filename to use with hdf5 (not available yet).
verbose	verbosity flag.
widget	a logical specifying if widgets should be used.
celfile.path	a character denoting the path ReadAffy should look for cel files.
sampleNames	a character vector of sample names to be used in the AffyBatch.
normalize	logical value. If TRUE, then normalize data using quantile normalization.
background	logical value. If $\ensuremath{\mathtt{TRUE}}$, then background correct using RMA background correction.
bgversion	integer value indicating which RMA background to use 1: use background simi- lar to pure R rma background given in affy version 1.0 - 1.0.2 2: use background similar to pure R rma background given in affy version 1.1 and above

list.celfiles

destructive	logical value. If ${\tt TRUE},$ then works on the PM matrix in place as much as possible, good for large datasets.
cdfname	Used to specify the name of an alternative cdf package. If set to NULL, then the usual cdf package based on Affymetrix' mappings will be used.

Details

justRMA is a wrapper for just.rma that permits the user to read in phenoData, MIAME information, and CEL files using widgets. One can also define files where to read phenoData and MIAME information.

If the function is called with no arguments justRMA(), then all the CEL files in the working directory are read, converted to an expression measure using RMA and put into an ExpressionSet. However, the arguments give the user great flexibility.

phenoData is read using read.AnnotatedDataFrame. If a character is given, it tries to read the file with that name to obtain the AnnotatedDataFrame object as described in read.AnnotatedDataFrame. If left NULL and widget=FALSE (widget=TRUE is not currently supported), then a default object is created. It will be an object of class AnnotatedDataFrame with its pData being a data.frame with column x indexing the CEL files.

description is read using read.MIAME. If a character is given, it tries to read the file with that name to obtain a MIAME instance. If left NULL but widget=TRUE, then widgets are used. If left NULL and widget=FALSE, then an empty instance of MIAME is created..

The arguments rm.masks, rm.outliers, rm.extra are passed along to the function read.celfile.

Value

An ExpressionSet object, containing expression values identical to what one would get from running rma on an AffyBatch.

Author(s)

In the beginning: James MacDonald <jmacdon@med.umich.edu> Supporting routines, maintenance and just.rma: Ben Bolstad
bmb@bmbolstad.com>

See Also

rma,ReadAffy

list.celfiles List the Cel Files in a Directory/Folder

Description

This function produces a vector containing the names of files in the named directory/folder ending in .cel or .CEL.

Usage

list.celfiles(...)

loess.normalize

Arguments

... arguments to pass along to list.files

Value

A character vector of file names.

See Also

list.files

Examples

list.celfiles()

loess.normalize Normalize arrays

Description

This function treats PM and MM as the raw data on each chip. It fits loess curves to MVA plots and tries to normalize the chips with respect to each other by forcing log ratios to be scattered around the same constant.

Usage

Arguments

mat	a matrix with columns containing the values of the chips to normalize.
subset	a subset of the data to fit a loess to.
epsilon	small value used for the stopping criterion.
maxit	maximum number of iterations.
log.it	logical. If TRUE it takes the log2 of mat
verbose	logical. If TRUE displays current pair of chip being worked on.
span	span to be used by loess
family.loess	"gaussian" or "symmetric" as in loess

Details

Experience shows that you only need 1-2 iterations to obtain useful results. This function is not written in an efficient way. In order to make it faster, loess is fit to a sample of the data which we then use to predict the curve for all the data. By setting family.loess="gaussian" the function is faster, but you risk losing information on differentially expressed genes. The function normalize.quantiles is faster.

Value

A matrix with normalized values for chips in columns.

Author(s)

Rafael A. Irizarry

See Also

normalize.quantiles, maffy.normalize, maffy.subset

maffy.normalize Normalize Itensities

Description

Normalizes feauture intensities from AffyBatchs

Usage

```
maffy.normalize(data,subset,verbose=FALSE,span=0.25,family="symmetric",log.it=TR
```

Arguments

data	an matrix of intensities.
subset	a vector of indexes describing which probes to use for normalising.
verbose	logical value.
span	See loess.
family	See loess.
log.it	logical value.

Details

Please refer to references.

Value

The normalized intensities.

Author(s)

Magnus Astrand

References

Astrand, M. (2003) http://www.math.chalmers.se/~magnusaa/maffy/

See Also

maffy.subset

maffy.subset

Examples

```
if (require(affydata)) {
   data(Dilution)
   x <- pm(Dilution)[1:2000,1:3]
   mva.pairs(x)
   x <- maffy.normalize(x,subset=1:nrow(x))
   mva.pairs(x)
}</pre>
```

maffy.subset Select Subset

Description

Select a subset of rows with small rank-range over columns.

Usage

Arguments

data	a matrix
subset.size	desired size of subset
maxit	maximum number of iterations
subset.delta	maximum deviation from subset.size
verbose	logical value.

Details

Please refer to references.

Value

A list with component subset, the indexes for subset.

Author(s)

Magnus Astrand

References

Astrand, M. (2001) http://www.math.chalmers.se/~magnusaa/maffy/

See Also

maffy.normalize

Examples

```
if (require(affydata)) {
   #data(Dilution)
   #x <- log2(pm(Dilution)[,1:3])
   #Index <- maffy.subset(x,subset.size=100)$subset
   #mva.pairs(x[Index,])
}</pre>
```

MAplot

Relative M vs. A plots

Description

Create boxplots of M or M vs A plots. Where M is determined relative to a specified chip or to a pseudo-median reference chip.

Usage

```
MAplot(object,...)
Mbox(object,...)
ma.plot(A, M, subset = sample(1:length(M), min(c(10000, length(M)))),
show.statistics=TRUE, span=2/3, family.loess="gaussian", cex = 2,plot.method=c("
```

Arguments

Additional parameters for the routineAA vector to plot along the horizonal axisMA vector to plot along vertical axissubsetA set of indices to use when drawing the loess curveshow.statistics
MA vector to plot along vertical axissubsetA set of indices to use when drawing the loess curve
subset A set of indices to use when drawing the loess curve
-
show.statistics
If true some summary statistics of the M values are drawn
span span to be used for loess fit.
family.loess "guassian" or "symmetric" as in loess.
cex Size of text when writing summary statistics on plot
plot.method a string specifying how the plot is to be drawn. "normal" plots points, "smoothScatter" uses the smoothScatter function. Specifying "add" means that the MAplot should be added to the current plot
add.loess add a loess line to the plot
lwd width of loess line
lty line type for loess line
loess.col color for loess line

See Also

mva.pairs

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mas5calls

Examples

```
if (require(affydata)) {
   data(Dilution)
   MAplot(Dilution)
   Mbox(Dilution)
}
```

mas5calls

MAS 5.0 Absolute Detection

Description

Performs the Wilcoxon signed rank-based gene expression presence/absence detection algorithm first implemented in the Affymetrix Microarray Suite version 5.

Usage

Arguments

object	An object of class AffyBatch or ProbeSet	
ids	probeset IDs for which you want to compute calls	
mat	an n-by-2 matrix of paired values (pairs in rows), PMs first col	
verbose	logical. It TRUE status of processing is reported	
tau	a small positive constant	
alpha1	a significance threshold in (0,alph2)	
alpha2	a significance threshold in (alpha1,0.5)	
exact.pvals	a boolean controlling whether exact p-values are computed (irrelevant if n<50 and there are no ties). Otherwise the normal approximation is used	
ignore.saturated		
	if true do the saturation correction described in the paper, with a saturation level of 46000	
cont.correct	a boolean controlling whether continuity correction is used in the p-value normal approximation	
	any of the above arguments that applies	

Details

This function performs the hypothesis test:

H0: median(Ri) = tau, corresponding to absence of transcript H1: median(Ri) > tau, corresponding to presence of transcript

where Ri = (PMi - MMi) / (PMi + MMi) for each i a probe-pair in the probe-set represented by data.

Currently exact.pvals=TRUE is not supported, and cont.correct=TRUE works but does not give great results (so both should be left as FALSE). The defaults for tau, alpha1 and alpha2 correspond to those in MAS5.0.

The p-value that is returned estimates the usual quantity:

Pr(observing a more "present looking" probe-set than data | data is absent)

So that small p-values imply presence while large ones imply absence of transcript. The detection call is computed by thresholding the p-value as in:

call "P" if p-value < alpha1 call "M" if alpha1 <= p-value < alpha2 call "A" if alpha2 <= p-value

This implementation has been validated against the original MAS5.0 implementation with the following results (for exact.pvals and cont.correct set to F):

Average Relative Change from MAS5.0 p-values:38% Proportion of calls different to MAS5.0 calls:1.0%

where "average/proportion" means over all probe-sets and arrays, where the data came from 11 bacterial control probe-sets spiked-in over a range of concentrations (from 0 to 150 pico-mols) over 26 arrays. These are the spike-in data from the GeneLogic Concentration Series Spikein Dataset.

Clearly the p-values computed here differ from those computed by MAS5.0 – this will be improved in subsequent releases of the affy package. However the p-value discrepancies are small enough to result in the call being very closely aligned with those of MAS5.0 (99 percent were identical on the validation set) – so this implementation will still be of use.

The function mas5.detect is no longer the engine function for the others. C code is no available that computes the wilcox test faster. The function is kept so that people can look at the R code (instead of C)

Value

mas5.detect returns a list containing the following components:

pval	a real p-value in [0,1] equal to the probability of observing probe-level intensities that are more present looking than data assuming the data represents an absent transcript; that is a transcript is more likely to be present for p-values closer 0.
call	either "P", "M" or "A" representing a call of present, marginal or absent; com- puted by simply thresholding pval using alpha1 and

The mas5calls method for AffyBatch returns an ExpressionSet with calls accessible with exprs(obj) and p-values available with assayData(obj)[["se.exprs"]]. The codemas5calls for ProbeSet returns a list with vectors of calls and pvalues.

Author(s)

Crispin Miller, Benjamin I. P. Rubinstein, Rafael A. Irizarry

mas5

References

Liu, W. M. and Mei, R. and Di, X. and Ryder, T. B. and Hubbell, E. and Dee, S. and Webster, T. A. and Harrington, C. A. and Ho, M. H. and Baid, J. and Smeekens, S. P. (2002) Analysis of high density expression microarrays with signed-rank call algorithms, Bioinformatics, 18(12), pp. 1593–1599.

Liu, W. and Mei, R. and Bartell, D. M. and Di, X. and Webster, T. A. and Ryder, T. (2001) Rankbased algorithms for analysis of microarrays, Proceedings of SPIE, Microarrays: Optical Technologies and Informatics, 4266.

Affymetrix (2002) Statistical Algorithms Description Document, Affymetrix Inc., Santa Clara, CA, whitepaper. http://www.affymetrix.com/support/technical/whitepapers/sadd_whitepaper.pdf, http://www.affymetrix.com/support/technical/whitepapers/sadd_whitepaper.pdf

Examples

```
if (require(affydata)) {
   data(Dilution)
   PACalls <- mas5calls(Dilution)
}</pre>
```

mas5

MAS 5.0 expression measure

Description

This function converts an instance of AffyBatch into an instance of ExpressionSet using our implementation of Affymetrix's MAS 5.0 expression measure.

Usage

mas5(object, normalize = TRUE, sc = 500, analysis = "absolute", ...)

Arguments

object	an instance of AffyBatch
normalize	logical. If $\ensuremath{\texttt{TRUE}}$ scale normalization is used after we obtain an instance of $\ensuremath{\texttt{ExpressionSet}}$
SC	Value at which all arrays will be scaled to.
analysis	should we do absolute or comparison analysis, although "comparison" is still not implemented.
	other arguments to be passed to expresso.

Details

This function is a wrapper for expresso and affy.scalevalue.exprSet.

Value

ExpressionSet

The methods used by this function were implemented based upon available documentation. In particular a useful reference is Statistical Algorithms Description Document by Affymetrix. Our implementation is based on what is written in the documentation and as you might appreciate there are places where the documentation is less than clear. This function does not give exactly the same results. All source code of our implementation is available. You are free to read it and suggest fixes.

For more information visit this URL: http://stat-www.berkeley.edu/users/bolstad/

See Also

expresso,affy.scalevalue.exprSet

Examples

```
if (require(affydata)) {
   data(Dilution)
   eset <- mas5(Dilution)
}</pre>
```

merge.AffyBatch merge two AffyBatch objects

Description

merge two AffyBatch objects into one.

Usage

Arguments

	additional arguments
notes	a character
description	a characterORmiame, eventualy NULL
annotation	a character
У	an AffyBatch
Х	an AffyBatch

Details

To be done.

Value

A object if class AffyBatch.

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multiloess

See Also

AffyBatch-class

multiloess

Local Polynomial Regression Fitting

Description

A modified version of of loess. Perform loess for every column of Y, but with the robust weights calculated using all columns

Usage

```
multiloess(formula, data=NULL, weights, subset, na.action, model = FALSE,
    span = 0.75, enp.target, degree = 2,
    normalize = TRUE,
    family = c("gaussian", "symmetric"),
    method = c("loess", "model.frame"),
    control = loess.control(...), ...)
```

Arguments

See loess.

Details

Please refer to loess.

Value

See loess.

Author(s)

Magnus Astrand

References

Astrand, M. (2001) http://www.math.chalmers.se/~magnusaa/maffy/

See Also

loess

mva.pairs

Description

A matrix of M vs. A plots is produced. Plots are made on the upper triangle and the IQR of the Ms are displayed in the lower triangle

Usage

Arguments

Х	A matrix containing the chip data in the columns.
labels	the names of the variables.
log.it	logical. If TRUE uses log scale.
span	span to be used for loess fit.
family.loess	"gaussian" or "symmetric" as in loess.
digits	number of digits to use in the display of IQR.
line.col	color of the loess line.
main	an overall title for the plot.
cex	size for text
•••	graphical parameters can be given as arguments to mva.plot

See Also

pairs

Examples

```
x <- matrix(rnorm(4000),1000,4)
x[,1] <- x[,1]^2
dimnames(x) <- list(NULL,c("chip 1","chip 2","chip 3","chip 4"))
mva.pairs(x,log=FALSE,main="example")</pre>
```

normalize.constant Scale probe intensities

Description

Scale array intensities in a AffyBatch.

Usage

```
normalize.AffyBatch.constant(abatch, refindex=1, FUN=mean, na.rm=TRUE)
normalize.constant(x, refconstant, FUN=mean, na.rm=TRUE)
```

Arguments

abatch	an instance of the AffyBatch-class.
Х	a vector of intensities on a chip (to normalize to the reference).
refindex	the index of the array used as a reference.
refconstant	the constant used as a reference
FUN	A function generating a value from the intensities on an array. Typically mean or median.
na.rm	Paramater passed to the function FUN.

Value

An AffyBatch with an attribute "constant" holding the value of the factor used for scaling.

Author(s)

L. Gautier <laurent@cbs.dtu.dk>

See Also

AffyBatch

normalize.contrasts

Normalize intensities using the contrasts method

Description

Scale chip objects in an AffyBatch-class.

Usage

Arguments

abatch	an AffyBatch-class
span	parameter to be passed to the function loess.
choose.subset	
subset.size	
verbose	verbosity flag
family	parameter to be passed to the function loess.
type	A string specifying how the normalization should be applied.

Value

An object of the same class as the one passed.

See Also

maffy.normalize

```
normalize.invariantset
```

Invariante Set normalization

Description

Normalize arrays in an AffyBatch using an invariant set.

Usage

```
normalize.invariantset(data, ref, prd.td=c(0.003,0.007))
```

Arguments

abatch	an AffyBatch
data	a vector of intensities on a chip (to normalize to the reference).
ref	a vector of reference intensities.
prd.td	cutoff parameter (details in the bibliographic reference)
baseline.type	
	Specify how to determine the baseline array
type	A string specifying how the normalization should be applied. See details for more.
verbose	A flag to have a dumps throughout the normalization

normalize.loess

Details

The set of invariant intensities between data and ref is found through an iterative process (based on the respective ranks the intensities). This set of intensities is used to generate a normalization curve by smoothing.

The type argument should be one of "separate", "pmonly", "mmonly", "together" which indicates whether to normalize only one probe type (PM,MM) or both together or separately.

Value

Respectively a AffyBatch of normalized objects, or a vector of normalized intensities, with an attribute "invariant.set" holding the indexes of the 'invariant' intensities.

Author(s)

L. Gautier <laurent@cbs.dtu.dk> (Thanks to Cheng Li for the discussions about the algorithm.)

References

Cheng Li and Wing Hung Wong, Model-based analysis of oligonucleotides arrays: model validation, design issues and standard error application. Genome Biology 2001, 2(8):research0032.1-0032.11

See Also

normalize to normalize AffyBatch objects.

normalize.loess Scale microarray data

Description

Normalizes arrays using loess.

Usage

Arguments

abatchan AffyBatch object.subseta subset of the data to fit a loess to.epsilona tolerance value (supposed to be a small value - used as a stopping criterium)maxitmaximum number of iterations.log.itlogical. If TRUE it takes the log2 of mat	mat	a matrix with columns containing the values of the chips to normalize.
epsilona tolerance value (supposed to be a small value - used as a stopping criterium)maxitmaximum number of iterations.	abatch	an AffyBatch object.
maxit maximum number of iterations.	subset	a subset of the data to fit a loess to.
	epsilon	a tolerance value (supposed to be a small value - used as a stopping criterium)
log.it logical. If TRUE it takes the log2 of mat	maxit	maximum number of iterations.
	log.it	logical. If TRUE it takes the log2 of mat

verbose	logical. If TRUE displays current pair of chip being worked on.
span	parameter to be passed the function loess
family.loess	parameter to be passed the function loess. "gaussian" or "symmetric" are acceptable values for this parameter.
type	A string specifying how the normalization should be applied. See details for more.
	any of the options of normalize.loess you would like to modify (described above).

Details

The type arguement should be one of "separate", "pmonly", "mmonly", "together" which indicates whether to normalize only one probe type (PM,MM) or both together or separately.

See Also

normalize

Examples

```
if (require(affydata)) {
  #data(Dilution)
  #x <- pm(Dilution[,1:3])
  #mva.pairs(x)
  #x <- normalize.loess(x,subset=1:nrow(x))
  #mva.pairs(x)
}</pre>
```

normalize-methods Normalize Affymetrix Probe Level Data - methods

Description

Method for normalizing Affymetrix Probe Level Data

Usage

```
normalize.methods(object)
bgcorrect.methods()
upDate.bgcorrect.methods(x)
pmcorrect.methods()
upDate.pmcorrect.methods(x)
```

Arguments

object	An AffyBatch.
х	A character vector that will replace the existing one.

normalize.qspline

Details

If object is an AffyBatch then normalize (object) returns an AffyBatch with the intensities normalized using the methodology specified by getOption ("BioC") \$affy\$normalize.method. The affy package default is quantiles.

Other methodologies can be used by specifying them with the method argument. For example to use the invariant set methodology described by Li and Wong (2001) one would type: normalize (object, method="invariantset").

Further arguments passed by ..., apart from method, are passed along to the function responsible for the methodology defined by the method argument.

A character vector of *nicknames* for the methodologies available is returned by normalize.methods (object)), where object is an AffyBatch, or simply by typing normalize.AffyBatch.methods. If the nickname of a method is called "loess", the help page for that specific methodology can be accessed by typing ?normalize.loess.

For more on the normalization methodologies currently implemented please refer to the vignette 'Custom Processing Methods'.

To add your own normalization procedures please refer to the customMethods vignette.

The functions: bgcorrect.methods, pmcorrect.methods, provide access to internal vectors listing the corresponding capabilities.

See Also

AffyBatch-class, normalize.

Examples

```
if (require(affydata)) {
   data(Dilution)
   normalize.methods(Dilution)
   generateExprSet.methods()
   bgcorrect.methods()
}
```

normalize.qspline Normalize arrays

Description

normalizes arrays in an AffyBatch each other or to a set of target intensities

Usage

```
normalize.AffyBatch.qspline(abatch,type=c("together", "pmonly", "mmonly",
    "separate"), ...)
normalize.qspline(x, target = NULL, samples = NULL,
    fit.iters = 5, min.offset = 5,
    spline.method = "natural", smooth = TRUE,
    spar = 0, p.min = 0, p.max = 1.0,
    incl.ends = TRUE, converge = FALSE,
    verbose = TRUE, na.rm = FALSE)
```

Arguments

Х	a data.matrix of intensities
abatch	an AffyBatch
target	numerical vector of intensity values to normalize to. (could be the name for one of the celfiles in 'abatch')
samples	numerical, the number of quantiles to be used for spline. if $(0,1]$, then it is a sampling rate
fit.iters	number of spline interpolations to average
min.offset	minimum span between quantiles (rank difference) for the different fit iterations
spline.methc	od
	specifies the type of spline to be used. Possible values are "fmm", "natural", and "periodic".
smooth	logical, if 'TRUE', smoothing splines are used on the quantiles
spar	smoothing parameter for 'splinefun', typically in (0,1].
p.min	minimum percentile for the first quantile
p.max	maximum percentile for the last quantile
incl.ends	include the minimum and maximum values from the normalized and target arrays in the fit
converge	(currently unimplemented)
verbose	logical, if 'TRUE' then normalization progress is reported
na.rm	logical, if 'TRUE' then handle NA values (by ignoring them)
type	A string specifying how the normalization should be applied. See details for more.
•••	Optional parameters to be passed through

Details

This normalization method uses the quantiles from each array and the target to fit a system of cubic splines to normalize the data. The target should be the mean (geometric) or median of each probe but could also be the name of a particular chip in the abatch object.

Parameters setting can be of much importance when using this method. The parameter fit.iter is used as a starting point to find a more appropriate value. Unfortunately the algorithm used do not converge in some cases. If this happens, the fit.iter value is used and a warning is thrown. Use of different settings for the parameter samples was reported to give good results. More specifically, for about 200 data points use samples = 0.33, for about 2000 data points use samples = 0.02 (thanks to Paul Boutros).

The type argument should be one of "separate", "pmonly", "mmonly", "together" which indicates whether to normalize only one probe type (PM,MM) or both together or separately.

Value

a normalized AffyBatch.

Author(s)

Laurent and Workman C.

References

Christopher Workman, Lars Juhl Jensen, Hanne Jarmer, Randy Berka, Laurent Gautier, Henrik Bjorn Nielsen, Hans-Henrik Saxild, Claus Nielsen, Soren Brunak, and Steen Knudsen. A new non-linear normal-ization method for reducing variability in dna microarray experiments. Genome Biology, accepted, 2002

normalize.quantiles

Quantile Normalization

Description

Using a normalization based upon quantiles, this function normalizes a matrix of probe level intensities.

Usage

```
normalize.AffyBatch.quantiles(abatch, type=c("separate", "pmonly", "mmonly", "too
```

Arguments

abatch	An AffyBatch
type	A string specifying how the normalization should be applied. See details for
	more.

Details

This method is based upon the concept of a quantile-quantile plot extended to n dimensions. No special allowances are made for outliers. If you make use of quantile normalization either through rma or expresso please cite Bolstad et al, Bioinformatics (2003).

The type argument should be one of "separate", "pmonly", "mmonly", "together" which indicates whether to normalize only one probe type (PM,MM) or both together or separately.

Value

A normalized AffyBatch.

Author(s)

Ben Bolstad, (bmbolstad.com)

References

Bolstad, B (2001) Probe Level Quantile Normalization of High Density Oligonucleotide Array Data. Unpublished manuscript http://bmbolstad.com/stuff/qnorm.pdf

Bolstad, B. M., Irizarry R. A., Astrand, M, and Speed, T. P. (2003) *A Comparison of Normalization Methods for High Density Oligonucleotide Array Data Based on Bias and Variance*. Bioinformatics 19(2), pp 185-193. http://bmbolstad.com/misc/normalize/normalize.html

See Also

normalize

normalize.quantiles.robust

Robust Quantile Normalization

Description

Using a normalization based upon quantiles, this function normalizes a matrix of probe level intensities. Allows weighting of chips

Usage

```
normalize.AffyBatch.quantiles.robust(abatch,
type=c("separate","pmonly","mmonly","together"),
weights=NULL,remove.extreme=c("variance","mean","both","none"),n.remove=1,use.me
```

Arguments

abatch	An AffyBatch
type	A string specifying how the normalization should be applied. See details for more.
weights	A vector of weights, one for each chip
remove.extre	me
	If weights is null, then this will be used for determining which chips to remove from the calculation of the normalization distribution, See details for more info
n.remove	number of chips to remove
use.median	if TRUE use the median to compute normalization chip, otherwise uses a weighted mean
use.log2	work on log2 scale. This means we will be using the geometric mean rather than ordinary mean

Details

This method is based upon the concept of a quantile-quantile plot extended to n dimensions. Note that the matrix is of intensities not log intensities. The function performs better with raw intensities.

Choosing **variance** will remove chips with variances much higher or lower than the other chips, **mean** removes chips with the mean most different from all the other means, **both** removes first extreme variance and then an extreme mean. The option **none** does not remove any chips, but will assign equal weights to all chips.

The type argument should be one of "separate", "pmonly", "mmonly", "together" which indicates whether to normalize only one probe type (PM,MM) or both together or separately.

Value

a matrix of normalized intensites

Note

This function is still experimental.

normalize

Author(s)

Ben Bolstad, (bmb@bmbolstad.com)

See Also

normalize, normalize.quantiles

normalize

Normalize - generic

Description

A generic function which normalizes microarray data. Normalization is intended to remove from the intensity measures any systematic trends which arise from the microarray technology rather than from differences between the probes or between the target RNA samples hybridized to the arrays.

Usage

```
normalize(object, ...)
```

Arguments

object	a data object containing microarray data
	any other arguments

See Also

Type showMethods ("normalize") at the R prompt to see what methods are available. Help on individual methods is generally available as normalize.<class> where <class> is the class of the data object. For example, for the main class in the affy package use ?normalize.AffyBatch.

Other Bioconductor packages include some related generic functions: normalizeWithinArrays, and normalizeBetweenArrays, in the LIMMA package, and maNorm in the marrayNorm package.

pairs.AffyBatch plot intensities using 'pairs'

Description

Plot intensities using the function 'pairs'

Usage

Arguments

х	an AffyBatch object
panel	a function to produce a plot (see pairs)
	extra parameters for the 'panel' function
transfo	a function to transform the intensity values before generating the plot. 'log' and 'log2' are popular choices.
main	title for the plot
oma	see 'oma' in par.
font.main	see pairs
cex.main	see pairs
cex.labels	see pairs
lower.panel	a function to produce the plots in the lower triangle (see pairs).
upper.panel	a function to produce the plots in the upper triangle (see pairs).
diag.panel	a function to produce the plots in the diagonal (see pairs).
font.labels	see pairs
rowlattop	see pairs
gap	see pairs

Details

Plots with several chips can represent zillions of points. They require a lot of memory and can be very slow to be displayed. You may want to try to split of the plots, or to plot them in a device like 'png' or 'jpeg'.

plotDensity Plot Densities

Description

Plots the non-parametric density estimates using values contained in the columns of a matrix.

Usage

plotLocation

Arguments

mat	A matrix containing the values to make densities in the columns.
Х	A object of clase AffyBatch
log	logical value. If ${\tt TRUE}$ the log of the intensities in the ${\tt AffyBatch}$ are plotted.
which	should a histogram of the PMs, MMs, or both be made?
col	The colors to use fot the different arrays
ylab	a title for the y axis.
xlab	a title for the x axis.
type	type for the plot.
na.rm	handling of NA values.
	graphical parameters can be given as arguments to plot

Details

The list returned can be convenient for plotting large input matrices with different colors/line types schemes (the computation of the densities can take some time).

To match other functions in base R, this function should probably be called matdensity, as it is sharing similarities with matplot and matlines.

Value

It returns invisibly a list of two matrices 'x' and 'y'.

Author(s)

Ben Bolstad and Laurent Gautier

Examples

```
if (require(affydata)) {
   data(Dilution)
   plotDensity(exprs(Dilution), log="x")
}
```

plotLocation Plot a location on a cel image

Description

Plots a location on a previously plotted cel image. This can be used to locate the physical location of probes on the array.

Usage

```
plotLocation(x, col="green", pch=22, ...)
```

Arguments

х	a 'location'. It can be obtained by the method of AffyBatch indexProbes, or made elsewhere (basically a location is nrows and two columns array. The first column corresponds to the x positions and the second columns corresponds to the y positions of n elements to locate)
col	colors for the plot
pch	plotting type (see function plot)
	Other parameters passed to the function points

Author(s)

Laurent

See Also

AffyBatch

Examples

```
if (require(affydata)) {
   data(Dilution)

   ## image of the celfile
   image(Dilution[, 1])

   ## genenames, arbitrarily pick the 101th
   n <- geneNames(Dilution)[101]

   ## get the location for the gene n
   l <- indexProbes(Dilution, "both", n)[[1]]

   ## convert the index to X/Y coordinates
   xy <- indices2xy(l, abatch=Dilution)

   ## plot
   plotLocation(xy)
}</pre>
```

plot.ProbeSet plot a probe set

Description

Plot intensities by probe set.

Usage

```
## S3 method for class 'ProbeSet':
plot(x, which=c("pm", "mm"), xlab = "probes", type = "l", ylim = NULL, ...)
```

pmcorrect

Arguments

Х	a ProbeSet
which	get the PM or the MM
xlab	label on x-axis
type	plot type
ylim	range of the y-axis
	optional arguments to be passed to matplot

Value

This function is only used for its (graphical) side-effect.

See Also

ProbeSet

Examples

```
data(SpikeIn)
plot(SpikeIn)
```

pmcorrect

PM Correction

Description

Corrects the PM intensities in a ProbeSet for nonspecific binding.

Usage

```
pmcorrect.pmonly(object)
```

```
pmcorrect.subtractmm(object)
```

pmcorrect.mas(object, contrast.tau=0.03, scale.tau=10, delta=2^(-20))

Arguments

object	An object of class ProbeSet.
contrast.tau	a number denoting the contrast tau parameter in the MAS 5.0 pm correction algorithm.
scale.tau	a number denoting the scale tau parameter in the MAS 5.0 pm correction algorithm.
delta	a number denoting the detla parameter in the MAS 5.0 pm correction algorithm.

Details

These are the pm correction methods perfromed by Affymetrix MAS 4.0 (subtractmm) and MAS 5.0 (mas). See the Affymetrix Manual for details. pmonly does what you think: does not change the PM values.

Value

A ProbeSet for which the pm slot contains the corrected PM values.

References

Affymetrix MAS 4.0 and 5.0 manual

Examples

```
if (require(affydata)) {
  data(Dilution)
  gn <- geneNames(Dilution)
  pps <- probeset(Dilution, gn[1])[[1]]
  pps.pmonly <- pmcorrect.pmonly(pps)
  pps.subtractmm <- pmcorrect.subtractmm(pps)
  pps.mas5 <- pmcorrect.mas(pps)
}</pre>
```

ppsetApply Apply a function over the ProbeSets in an AffyBatch

Description

Apply a function over the ProbeSets in an AffyBatch

Usage

```
ppsetApply(abatch, FUN, genenames = NULL, ...)
```

ppset.ttest(ppset, covariate, pmcorrect.fun = pmcorrect.pmonly, ...)

Arguments

abatch	An object inheriting from AffyBatch.	
ppset	An object of class ProbeSet.	
covariate	the name a covariate in the slot phenoData.	
pmcorrect.fun		
	a function to corrrect PM intensities	
FUN	A function working on a ProbeSet	
genenames	A list of Affymetrix probesets ids to work with. All probe set ids used when $\ensuremath{\texttt{NULL}}$.	
	Optional parameters to the function FUN	

Value

Returns a list of objects, or values, as returned by the function FUN for each ProbeSet it processes.

probeMatch-methods

Author(s)

Laurent Gautier <laurent@cbs.dtu.dk>

See Also

ProbeSet-class

Examples

```
ppset.ttest <- function(ppset, covariate, pmcorrect.fun = pmcorrect.pmonly, ...) {
    probes <- do.call("pmcorrect.fun", list(ppset))
    my.ttest <- function(x) {
        y <- split(x, get(covariate))
        t.test(y[[1]], y[[2]])$p.value
    }
    r <- apply(probes, 1, my.ttest)
    return(r)
}
##this takes a long time - and rowttests is a good alternative
## eg: rt = rowttests(exprs(Dilution), Dilution$liver)
## Not run:
    data(Dilution)
    all.ttest <- ppsetApply(Dilution, ppset.ttest, covariate="liver")
## End(Not run)</pre>
```

probeMatch-methods *Methods for accessing perfect matches and mismatches*

Description

Methods for perfect matches and mismatches probes

Methods

object = AffyBatch All the *perfect match* (pm) or *mismatch* (mm) probes on the arrays the object represents are returned.

object = ProbeSet The pm or mm of the object are returned

probeNames-methods Methods for accessing the Probe Names

Description

Methods for accessing Probe Names

Methods

object = Cdf An accesor function for the name slot.

object = probeNames Returns the probe names associated with the rownames of the intensity matrices one gets with the pm and mm methods.

ProbeSet-class Class ProbeSet

Description

A simple class that contains the PM and MM data for a probe set from one or more samples

Objects from the Class

Objects can be created by applying the method probeset to instances of AffyBatch.

Slots

- id: Object of class "character" containing the probeset ID
- **pm:** Object of class "matrix" containing the PM intensities. Columns represent samples and rows the different probes.
- mm: Object of class "matrix" containing the MM intensities

Methods

- colnames signature(x = "ProbeSet"): the column names of the pm matrices which are the sample names
- express.summary.stat signature(x = "ProbeSet", pmcorrect = "character", summary = "character"): applies a summary statistic to the probe set.
- sampleNames signature(object = "ProbeSet"): the column names of the pm matrices
 which are the sample names

Note

More details are cotained in the vignette

See Also

probeset, AffyBatch-class

Examples

```
if (require(affydata)) {
   data(Dilution)
   ps <- probeset(Dilution, geneNames(Dilution)[1:2])
   names(ps)
   print(ps[[1]])
}</pre>
```

ProgressBarText-class

Class "ProgressBarText"

Description

A class to handle progress bars in text mode

Objects from the Class

Objects can be created by calls of the form new ("ProgressBarText", steps).

Slots

steps: Object of class "integer". The total number of steps the progress bar should represent

barsteps: Object of class "integer". The size of the progress bar.

internals: Object of class "environment". For internal use.

Methods

close signature(con = "ProgressBarText"): Terminate the progress bar (i.e. print
what needs to be printed). Note that closing the instance will ensure the progress bar is plotted
to its end.

initialize signature(.Object = "ProgressBarText"): initialize a instance.

- **open** signature(con = "ProgressBarText"): Open a progress bar (i.e. print things). In the case open is called on a progress bar that was 'progress', the progress bar is resumed (this might be useful when one wishes to insert text output while there is a progress bar running).
- updateMe signature(object = "ProgressBarText"): Update the progress bar (see examples).

Author(s)

Laurent

Examples

```
f <- function(x, header = TRUE) {
   pbt <- new("ProgressBarText", length(x), barsteps = as.integer(20))
   open(pbt, header = header)
   for (i in x) {
     Sys.sleep(i)
     updateMe(pbt)
   }
   close(pbt)
}
## if too fast on your machine, change the number
x <- runif(15)</pre>
```

```
f(x)
f(x, header = FALSE)
## 'cost' of the progress bar:
q <- function(x) {
  z <- 1
  for (i in 1:x) {
    z <- z + 1
  }
}
h <- function(x) {
  pbt <- new("ProgressBarText", as.integer(x), barsteps = as.integer(20))</pre>
  open(pbt)
  for (i in 1:x) {
   updateMe(pbt)
  }
  close(pbt)
}
system.time(q(10000))
system.time(h(10000))
```

read.affybatch Read CEL files into an AffyBatch

Description

Read CEL files into an Affybatch.

Usage

```
read.affybatch(..., filenames = character(0),
               phenoData = new("AnnotatedDataFrame"),
               description = NULL,
               notes = "",
               compress = getOption("BioC")$affy$compress.cel,
               rm.mask = FALSE, rm.outliers = FALSE, rm.extra = FALSE,
               verbose = FALSE, sd=FALSE, cdfname = NULL)
ReadAffy(..., filenames=character(0),
              widget=getOption("BioC")$affy$use.widgets,
              compress=getOption("BioC")$affy$compress.cel,
              celfile.path=NULL,
              sampleNames=NULL,
              phenoData=NULL,
              description=NULL,
              notes="",
              rm.mask=FALSE, rm.outliers=FALSE, rm.extra=FALSE,
              verbose=FALSE, sd=FALSE, cdfname = NULL)
```

read.affybatch

Arguments

	file names separated by comma.
filenames	file names in a character vector.
phenoData	an AnnotatedDataFrame object, a character of length one, or a data.frame.
description	a MIAME object.
notes	notes.
compress	are the CEL files compressed?
rm.mask	should the spots marked as 'MASKS' set to NA?
rm.outliers	should the spots marked as 'OUTLIERS' set to NA?
rm.extra	if TRUE, then overrides what is in rm.mask and rm.oultiers.
verbose	verbosity flag.
widget	a logical specifying if widgets should be used.
celfile.path	a character denoting the path ReadAffy should look for cel files.
sampleNames	a character vector of sample names to be used in the AffyBatch.
sd	should the standard deviation values in the CEL file be read in? Since these are typically not used default is not to read them in. This also save lots of memory.
cdfname	used to specify the name of an alternative cdf package. If set to NULL, then the usual cdf package based on Affymetrix' mappings will be used.

Details

ReadAffy is a wrapper for read.affybatch that permits the user to read in phenoData, MI-AME information, and CEL files using widgets. One can also define files where to read phenoData and MIAME information.

If the function is called with no arguments ReadAffy() all the CEL files in the working directory are read and put into an AffyBatch. However, the arguments give the user great flexibility.

If phenoData is a character vector of length 1, the function read.AnnotatedDataFrame is called to read a file of that name and produce the AnnotationDataFrame object with the sample metadata. If phenoData is a data.frame, it is converted to an AnnotatedDataFrame. If it is NULL and widget=FALSE (widget=TRUE is not currently supported), then a default object of class AnnotatedDataFrame is created, whose pData is a data.frame with rownames being the names of the CEL files, and with one column sample with an integer index.

AllButCelsForReadAffy is an internal function that gets called by ReadAffy. It gets all the information except the cel intensities.

description is read using read.MIAME. If a character is given, then it tries to read the file with that name to obtain a MIAME instance. If left NULL but widget=TRUE, then widgets are used. If left NULL and widget=FALSE, then an empty instance of MIAME is created.

Value

An AffyBatch object.

Author(s)

Ben Bolstad (bmb@bmbolstad.com) (read.affybatch), Laurent Gautier, and Rafael A. Irizarry (ReadAffy)

See Also

AffyBatch

Examples

```
if(require(affydata)){
    celpath <- system.file("celfiles", package="affydata")
    fns <- list.celfiles(path=celpath,full.names=TRUE)
    cat("Reading files:\n",paste(fns,collapse="\n"),"\n")
    ##read a binary celfile
    abatch <- ReadAffy(filenames=fns[1])
    ##read a text celfile
    abatch <- ReadAffy(filenames=fns[2])
    ##read all files in that dir
    abatch <- ReadAffy(celfile.path=celpath)
}</pre>
```

read.probematrix Read CEL file data into PM or MM matrices

Description

Read CEL data into matrices.

Usage

```
read.probematrix(..., filenames = character(0),
    phenoData = new("AnnotatedDataFrame"),
    description = NULL,
    notes = "",
    compress = getOption("BioC")$affy$compress.cel,
    rm.mask = FALSE, rm.outliers = FALSE, rm.extra = FALSE,
    verbose = FALSE, which="pm", cdfname = NULL)
```

Arguments

	file names separated by comma.
filenames	file names in a character vector.
phenoData	a AnnotatedDataFrame object
description	a MIAME object
notes	notes
compress	are the CEL files compressed ?
rm.mask	should the spots marked as 'MASKS' set to NA ?
rm.outliers	should the spots marked as 'OUTLIERS' set to NA
rm.extra	if TRUE, overrides what is in rm.mask and rm.oultiers
verbose	verbosity flag
which	should be either "pm", "mm" or "both"
cdfname	Used to specify the name of an alternative cdf package. If set to NULL, the usual cdf package based on Affymetrix' mappings will be used.

rma

Value

A list of one or two matrices. Each matrix is either PM or MM data. No AffyBatch is created.

Author(s)

Ben Bolstad (bmb@bmbolstad.com)

See Also

AffyBatch, read.affybatch

rma

Robust Multi-Array Average expression measure

Description

This function converts an AffyBatch into an ExpressionSet using the robust multi-array average (RMA) expression measure.

Usage

rma(object,subset=NULL, verbose=TRUE, destructive = TRUE,normalize=TRUE,backgrou

Arguments

object	an AffyBatch
subset	a character vector with the the names of the probesets to be used in expression calculation.
verbose	logical value. If TRUE it writes out some messages indicating progress. If FALSE nothing should be printed.
destructive	logical value. If TRUE works on the PM matrix in place as much as possible, good for large datasets.
normalize	logical value. If TRUE normalize data using quantile normalization
background	logical value. If TRUE background correct using RMA background correction
bgversion	integer value indicating which RMA background to use 1: use background simi- lar to pure R rma background given in affy version 1.0 - 1.0.2 2: use background similar to pure R rma background given in affy version 1.1 and above
	further arguments to be passed (not currently implemented - stub for future use)

Details

This function computes the RMA (Robust Multichip Average) expression measure described in Irizarry et al Biostatistics (2003).

Note that this expression measure is given to you in log base 2 scale. This differs from most of the other expression measure methods.

Please note that the default background adjustment method was changed during the lead up to the bioconductor 1.2 release. This means that this function and expresso should give results that directly agree.

Value

An ExpressionSet

Author(s)

Ben Bolstad (bmb@bmbolstad.com)

References

Rafael. A. Irizarry, Benjamin M. Bolstad, Francois Collin, Leslie M. Cope, Bridget Hobbs and Terence P. Speed (2003), Summaries of Affymetrix GeneChip probe level data Nucleic Acids Research 31(4):e15

Bolstad, B.M., Irizarry R. A., Astrand M., and Speed, T.P. (2003), A Comparison of Normalization Methods for High Density Oligonucleotide Array Data Based on Bias and Variance. Bioinformatics 19(2):185-193

Irizarry, RA, Hobbs, B, Collin, F, Beazer-Barclay, YD, Antonellis, KJ, Scherf, U, Speed, TP (2003) Exploration, Normalization, and Summaries of High Density Oligonucleotide Array Probe Level Data. Biostatistics .Vol. 4, Number 2: 249-264

See Also

expresso

Examples

```
if (require(affydata)) {
   data(Dilution)
   eset <- rma(Dilution)
}</pre>
```

.setAffyOptions ~~function to set options ~~

Description

~~ Set the options for the package

Usage

```
.setAffyOptions(affy.opt = NA)
```

Arguments

affy.opt A list structure of options. If NA, the default options are set.

Details

See the vignettes to know more. This function could disappear in favor of a more general one the package Biobase

simplemultiLoess

Value

The function is used for its side effect. Nothing is returned.

Author(s)

Laurent

Examples

```
affy.opt <- getOption("BioC")$affy</pre>
```

```
.setAffyOptions(affy.opt)
```

simplemultiLoess Internal function for multiloess

Description

A modified version of of simpleLoess. Perform loess for every column of Y, but with the robust weights calculated using all columns

Usage

```
simplemultiLoess(y, x, weights, span = 0.75, degree = 2,
    normalize = TRUE,
    statistics = "approximate", surface = "interpolate",
    cell = 0.2, iterations = 1, trace.hat = "exact")
```

Arguments

See loess.

Details

Please refer to loess.

Value

See loess.

Author(s)

Magnus Astrand

References

Astrand, M. (2001) http://www.math.chalmers.se/~magnusaa/maffy/

See Also

loess

SpikeIn

Description

This ProbeSet represents part of SpikeIn experiment data set.

Usage

```
data(SpikeIn)
```

Format

SpikeIn is ProbeSet containing the PM and MM intensities for a gene spiked in at different concentrations (given in the vector colnames (pm(SpikeIn))) in 12 different arrays.

Source

This comes from an experiments where 11 different cRNA fragments have been added to the hybridization mixture of the GeneChip arrays at different pM concentrations. The 11 control cRNAs were BioB-5, BioB-M, BioB-3, BioC-5, BioC-3, BioDn-5 (all *E. coli*), CreX-5, CreX-3 (phage P1), and DapX-5, DapX-M, DapX-3 (*B. subtilis*) The cRNA were chosen to match the target sequence for each of the Affymetrix control probe sets. For example, for DapX (a *B. subtilis* gene), the 5', middle and 3' target sequences (identified by DapX-5, DapX-M, DapX-3) were each synthesized separately and spiked-in at a specific concentration. Thus, for example, DapX-3 target sequence may be added to the total hybridization solution of 200 micro-liters to give a final concentration of 0.5 pM.

For this example we have the PM and MM for BioB-5 obtained from the arrays where it was spiked in at 0.0, 0.5, 0.75, 1, 1.5, 2, 3, 5, 12.5, 25, 50, and 150 pM.

For more information see Irizarry, R.A., et al. (2001) http://biosun01.biostat.jhsph.edu/~ririzarr/papers/index.html

summary

Probe Set Summarizing Functions

Description

These were used with the function express which is no longer part of the package. Some are still used by the generateExprVal functions. But you should avoid using them directly.

See Also

expresso

tukey.biweight One-step Tukey's biweight

Description

One-step Tukey's biweight on a matrix

Usage

tukey.biweight(x, c = 5, epsilon = 1e-04)

Arguments

Х	a matrix
С	tuning constant (see details)
epsilon	fuzz value to avoid division by zero (see details)

Details

The details can be found in the given reference.

Value

a vector of values (one value per column in the input matrix).

References

Statistical Algorithms Description Document, 2002, Affymetrix.

See Also

pmcorrect.mas and generateExprVal.method.mas

whatcdf

Find which CDF corresponds

Description

Find which kind of CDF corresponds to a CEL file.

Usage

```
whatcdf(filename, compress = getOption("BioC")$affy$compress.cel)
```

Arguments

filename	a '.CEL' file name
compress	boolean (file compressed or not

Details

Information concerning the corresponding CDF file seems to be found in CEL files. This allows us to try to link CDF information automatically.

Value

a character with the name of the CDF

See Also

```
getInfoInAffyFile, read.celfile
```

xy2indices Functions to convert indices to x/y (and reverse)

Description

Functions to convert indices to x/y (and reverse)

Usage

xy2indices(x, y, nr = NULL, cel = NULL, abatch = NULL, cdf = NULL, xy.offset = N indices2xy(i, nr = NULL, cel = NULL, abatch = NULL, cdf = NULL, xy.offset = NULL

Arguments

Х	X position for the probes
У	Y position for the probes
i	indices in the AffyBatch for the probes
nr	total number of Xs on the chip
cel	a corresponding object of class Cel
abatch	a corresponding object of class AffyBatch
cdf	character - the name of the corresponding cdf package
xy.offset	an eventual offset for the XY coordinates. See Details

Details

The probes intensities for given probe set ids are extracted from an AffyBatch object using the indices stored in the corresponding cdfenv.

The parameter xy.offset is there for compatibility. For historical reasons, the xy-coordinates for the features on Affymetrix chips were decided to start at 1 (one) rather than 0 (zero). One can set the offset to 1 or to 0. Unless the you _really_ know what you are doing, it is advisable to let it at the default value NULL. This way the package-wide option xy.offset is always used.

Value

A vector of indices or a two-columns matrix of Xs and Ys.

xy2indices

Warning

Even if one really knows what is going on, playing with the parameter xy.offset could be risky. Changing the package-wide option xy.offset appears much more sane.

Author(s)

L.

See Also

indexProbes

Examples

```
if (require(affydata)) {
   data(Dilution)
   pm.i <- indexProbes(Dilution, which="pm", genenames="AFFX-BioC-5_at")[[1]]
   mm.i <- indexProbes(Dilution, which="mm", genenames="AFFX-BioC-5_at")[[1]]
   pm.i.xy <- indices2xy(pm.i, abatch = Dilution)
   mm.i.xy <- indices2xy(mm.i, abatch = Dilution)
   image(Dilution[1], transfo=log2)
   ## plot the pm in red
   plotLocation(pm.i.xy, col="red")
   plotLocation(mm.i.xy, col="blue")
}</pre>
```

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