

SSCORE

October 25, 2011

OneStepBiweightAlgorithm

One-step Tukey's biweight

Description

Computes one-step Tukey's biweight on a vector. Note that this implementation follows the Affymetrix code, which is different from the Tukey's biweight computed by the affy package.

Usage

```
OneStepBiweightAlgorithm(x, c, epsilon)
```

Arguments

x	vector of data
c	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 numeric value

Note

Based on Affymetrix MAS5 Statistical SDK source code http://www.affymetrix.com/Auth/support/developer/stat_sdk/STAT_SDK_source.zip

Author(s)

Richard Kennedy <rkennedy@vcu.edu>

References

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

SScore

Compute S-Score values

Description

Computes the S-Score values for a pair of Affymetrix GeneChips

Usage

```
SScore(afbatch = stop("No CEL files specified"), classlabel = c(0,1), SF = NULL,
       SDT = NULL, rm.outliers = TRUE, rm.mask = TRUE, rm.extra = NULL,
       digits = NULL, verbose = FALSE, celfile.path = NULL, celfile.names = NULL)
```

Arguments

<code>afbatch</code>	An <code>AffyBatch</code> object
<code>classlabel</code>	A vector identifying the class for each column of the <code>AffyBatch</code> object
<code>SF</code>	a list of Scale Factor (SF) values for each GeneChip
<code>SDT</code>	a list of Standard Difference Threshold (SDT) values for each GeneChip
<code>rm.outliers</code>	should the spots marked as 'OUTLIERS' be excluded from S-Score calculation?
<code>rm.mask</code>	should the spots marked as 'MASKS' be excluded from S-Score calculation?
<code>rm.extra</code>	if TRUE, overrides what is in <code>rm.mask</code> and <code>rm.outliers</code>
<code>digits</code>	number of significant digits for S-Score values
<code>verbose</code>	logical value. If TRUE it provides more detail of the S-Score calculations.
<code>celfile.path</code>	character denoting the path for the *.CEL files corresponding to <code>afbatch</code>
<code>celfile.names</code>	optional character vector containing the names of the *.CEL files

Details

Computes S-Score values as described by Zhang et al. (2002). `SScore` provides a simpler interface for comparing only two classes of GeneChips, while `SScoreBatch` compares multiple pairs of chips.

The `classlabel` consists of a vector with one entry for each column of the `AffyBatch` object. Each entry consists of a 0 or a 1 to identify the class to which the chip for the corresponding column belongs. `SScore` will conduct a two-class test comparing all chips labeled 0 to all chips labeled 1. If `classlabel` is not specified, it defaults to a two-chip comparison, compatible with previous versions of `SScore`.

The SF and SDT factors are required for all calculations. If NULL, these values will be calculated according to the Affymetrix Statistical Algorithms Description Document. `digits` allows the specification of the number of significant digits for the S-Score values; if NULL, the maximum number of significant digits are retained.

Value

An `ExpressionSet` with S-Score values in the `exprs` slot.

Note

Based on C++ code by Li Zhang and Delphi code by Robnet Kerns

Author(s)

Richard Kennedy <rkennedy@vcu.edu>

References

Zhang, L., Wang, L., Ravindranathan, A., Miles, M.F. (2002) A new algorithm for analysis of oligonucleotide arrays: application to expression profiling in mouse brain regions. *Journal of Molecular Biology*, 317(2), pp. 225–35

Kerns, R.T., Zhang, L., Miles, M.F. (2003) Application of the S-score algorithm for analysis of oligonucleotide microarrays. *Methods*, 31(4), pp. 274–81

See Also

[SScoreBatch](#), [computeSFandSDT](#), [computeOutlier](#)

Examples

```
if (length(dir(pattern=".cel$")) != 0) {  
  
  ## Read in the *.CEL files  
  abatch <- ReadAffy()  
  
  ## default calling method  
  SScores <- SScore(abatch)  
  
  ## specifying SF and SDT (gives same results as above)  
  SfSdt <- computeSFandSDT(abatch)  
  SScores <- SScore(abatch, SF=SfSdt$SF, SDT=SfSdt$SDT)  
  
  ## specifying outlier and masked values should be included in calculations  
  SScores <- SScore(abatch, rm.outliers=FALSE, rm.mask=FALSE)  
  
  ## round results to 3 significant digits  
  SScores <- SScore(abatch, digits=3)  
  
  ## show verbose output  
  SScores <- SScore(abatch, verbose=TRUE)  
  
}
```

SScoreBatch

Compute S-Score values

Description

Computes the S-Score values for multiple pairs of Affymetrix GeneChips

Usage

```
SScoreBatch(afbatches = stop("No CEL files specified"), compare = stop("No list of
SF = NULL, SDT = NULL, rm.outliers = TRUE, rm.mask = TRUE, rm.extra
digits = NULL, verbose = FALSE, celfile.path = NULL, celfile.names =
```

Arguments

<code>afbatches</code>	An <code>AffyBatch</code> object
<code>compare</code>	A matrix describing which chips to compare
<code>SF</code>	a list of Scale Factor (SF) values for each GeneChip
<code>SDT</code>	a list of Standard Difference Threshold (SDT) values for each GeneChip
<code>rm.outliers</code>	should the spots marked as 'OUTLIERS' be excluded from S-Score calculation?
<code>rm.mask</code>	should the spots marked as 'MASKS' be excluded from S-Score calculation?
<code>rm.extra</code>	if TRUE, overrides what is in <code>rm.mask</code> and <code>rm.outliers</code>
<code>digits</code>	number of significant digits for S-Score values
<code>verbose</code>	logical value. If TRUE it provides more detail of the S-Score calculations.
<code>celfile.path</code>	character denoting the path for the *.CEL files corresponding to <code>afbatches</code>
<code>celfile.names</code>	optional character vector containing the names of the *.CEL files

Details

Computes S-Score values as described by Zhang et al. (2002). `SScoreBatch` allows comparison of multiple pairs of chips, while `SScore` provides a simpler interface when comparing only two GeneChips.

`compare` specifies how the pairwise comparisons are performed. It is an $N \times 2$ matrix, where N is the number of pairwise comparisons; each row of the matrix contains index in the `AffyBatch` object for the chips to be compared. For example,

```
1    3
4    2
5    9
10   2
5    7
```

would do a comparison of chip 1 to chip 3, a comparison of chip 4 to chip 2, a comparison of chip 5 to chip 9, and so on. The columns in `ExpressionSet` correspond to the rows of `compare`, so that the results of the first comparison are in column 1, the results of the second comparison are in column 2, and so on.

The SF and SDT factors are required for all calculations. If NULL, these values will be calculated according to the Affymetrix Statistical Algorithms Description Document. `probes.digits` allows the specification of the number of significant digits for the S-Score values; if NULL, the maximum number of significant digits are retained.

Value

An `ExpressionSet` with S-Score values in the `exprs` slot.

Note

Based on C++ code by Li Zhang and Delphi code by Robnet Kerns

Author(s)

Richard Kennedy <rkennedy@vcu.edu>

References

Zhang, L., Wang, L., Ravindranathan, A., Miles, M.F. (2002) A new algorithm for analysis of oligonucleotide arrays: application to expression profiling in mouse brain regions. *Journal of Molecular Biology*, 317(2), pp. 225–35

Kerns, R.T., Zhang, L., Miles, M.F. (2003) Application of the S-score algorithm for analysis of oligonucleotide microarrays. *Methods*, 31(4), pp. 274–81

See Also

[SScore](#), [computeSFandSDT](#), [computeOutlier](#)

Examples

```
if (length(dir(pattern=".cel$")) != 0) {  
  
  ## Read in the *.CEL files  
  abatch <- ReadAffy()  
  
  ## default calling method  
  SScores <- SScoreBatch(abatch)  
  
  ## specifying SF and SDT (gives same results as above)  
  SfSdt <- computeSFandSDT(abatch)  
  
  ## specifying outlier and masked values should be included in calculations  
  SScores <- SScoreBatch(abatch, SF=SfSdt$SF, SDT=SfSdt$SDT)  
  SScores <- SScoreBatch(abatch, rm.outliers=FALSE, rm.mask=FALSE)  
  
  ## round results to 3 significant digits  
  SScores <- SScoreBatch(abatch, digits=3)  
  
  ## show verbose output  
  SScores <- SScoreBatch(abatch, verbose=TRUE)  
  
}
```

computeAffxRawQ

Computes RawQ using affxparser routines

Description

Computes the RawQ value of a single Affymetrix GeneChip

Usage

```
computeAffxRawQ(intensity, stdvs, pixels, probe.index, probe.zoneID, bgCells, Nu
```

Arguments

intensity	vector of intensities for the GeneChip
stdvs	matrix of standard deviations for probe intensities from *.CEL file
pixels	matrix of number of pixels for probe intensities from *.CEL file
probe.index	vector of indices for each probe
probe.zoneID	vector of zone ID numbers for each probe
bgCells	number of background cells for the GeneChip
NumberZones	number of zones on the GeneChip

Details

Calculates RawQ using the algorithms described in the Affymetrix Statistical Algorithms Description Document for a single GeneChip. This is an internal function that will generally not be accessed directly.

Value

the RawQ value for the given array

Note

Based on Affymetrix MAS5 Statistical SDK source code http://www.affymetrix.com/Auth/support/developer/stat_sdk/STAT_SDK_source.zip, as well as C++ code by Li Zhang and Delphi code by Robnet Kerns

Author(s)

Richard Kennedy <rkennedy@vcu.edu>

References

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

```
computeAffxSFandSDT
```

Compute SF and SDT values using affxparser routines

Description

Computes the scaling factor (SF) and statistical difference threshold (SDT) values of Affymetrix GeneChips, for use in calculating S-Score values

Usage

```
computeAffxSFandSDT(afbatch, stdvs, pixels, TGT = 500, digits = NULL, verbose =
```

Arguments

afbatch	An AffyBatch object
stdvs	matrix of standard deviations for probe intensities from *.CEL file
pixels	matrix of number of pixels for probe intensities from *.CEL file
TGT	the target intensity to which the arrays should be scaled
digits	number of significant digits for SF and SDT values
verbose	logical value. If TRUE it provides more detail of the SF and SDT calculations.
plot.histogram	logical value. if TRUE it plots a histogram of intensities

Details

Calculates SF and SDT factors using the algorithms described in the Affymetrix Statistical Algorithms Description Document. The SF and SDT may be used in the calculation of S-Score values, or may be useful in their own right. One SF and SDT value is calculated for each GeneChip, which are arranged in the same order as the columns in the AffyBatch object.

Value

computeSFandSDT returns a list containing the following components:

SF	SF values, one for each GeneChip
SDT	SDT values, one for each GeneChip

Note

Based on Affymetrix MAS5 Statistical SDK source code http://www.affymetrix.com/Auth/support/developer/stat_sdk/STAT_SDK_source.zip, as well as C++ code by Li Zhang and Delphi code by Robnet Kerns

Author(s)

Richard Kennedy <rkennedy@vcu.edu>

References

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

Examples

```
if (length(dir(pattern=".cel$")) != 0) {
  ## Read in the *.CEL files
  abatch <- ReadAffy()

  ## compute SF and SDT
```

```

SfSdt <- computeSFandSDT(abatch)

## show verbose output
SfSdt <- computeSFandSDT(abatch,verbose=TRUE)

## plot PM and MM histograms for each *.CEL file
SfSdt <- computeSFandSDT(abatch,plot.histogram=TRUE)

}

```

computeOutlier *Compute outlier probes*

Description

Computes outlier and/or mask probes for a set of Affymetrix GeneChips that will be excluded from S-Score calculations

Usage

```
computeOutlier(afbatch, rm.mask = TRUE, rm.outliers = TRUE, rm.extra = TRUE, cel
```

Arguments

afbatch	An AffyBatch object
rm.mask	should the spots marked as 'MASKS' be excluded from S-Score calculation?
rm.outliers	should the spots marked as 'OUTLIERS' be excluded from S-Score calculation?
rm.extra	if TRUE, overrides what is in rm.mask and rm.outliers
celfile.path	character denoting the path for the *.CEL files corresponding to afbatch
celfile.names	optional character vector containing the names of the *.CEL files

Details

Computes the outlier and / or mask probes for an AffyBatch object. These are returned in matrix form, with one probe per row and one chip per column. The value of each location in the matrix will be TRUE if the corresponding probe is an outlier / masked value and FALSE if it is not. The options may be set to exclude only outlier values, only mask values, or both. The probes are arranged in the same row order as the intensity values. Note that this function assumes the *.CEL files are still available in the directory given by celfile.path (or the current directory if celfile.path is not specified). The *.CEL names are given by celfile.names. If celfile.names is not specified, the sample names from the AffyBatch object will be used.

Value

a matrix containing the list of outliers / masked values for the given AffyBatch object.

Note

Based on C++ code by Li Zhang and Delphi code by Robnet Kerns

Author(s)

Richard Kennedy <rkennedy@vcu.edu>

Examples

```
if (length(dir(pattern=".cel$")) != 0) {  
  abatch <- ReadAffy()  
  outlier <- computeOutlier(abatch)  
}
```

computeRawQ	<i>Computes RawQ</i>
-------------	----------------------

Description

Computes the RawQ value of a single Affymetrix GeneChip

Usage

```
computeRawQ(fname, intensity, probe.index, probe.zoneID, bgCells, NumberZones, c
```

Arguments

fname	character string with the filename of the GeneChip
intensity	vector of intensities for the GeneChip
probe.index	vector of indices for each probe
probe.zoneID	vector of zone ID numbers for each probe
bgCells	number of background cells for the GeneChip
NumberZones	number of zones on the GeneChip
celfile.path	character denoting the path for the *.CEL files specified in fname

Details

Calculates RawQ using the algorithms described in the Affymetrix Statistical Algorithms Description Document for a single GeneChip. This is an internal function that will generally not be accessed directly.

Value

the RawQ value for the given array

Note

Based on Affymetrix MAS5 Statistical SDK source code http://www.affymetrix.com/Auth/support/developer/stat_sdk/STAT_SDK_source.zip, as well as C++ code by Li Zhang and Delphi code by Robnet Kerns

Author(s)

Richard Kennedy <rkennedy@vcu.edu>

References

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

computeSFandSDT *Compute SF and SDT values*

Description

Computes the scaling factor (SF) and statistical difference threshold (SDT) values of Affymetrix GeneChips, for use in calculating S-Score values

Usage

```
computeSFandSDT(afbatch, TGT = 500, digits = NULL, verbose = FALSE, plot.histogram
```

Arguments

afbatch	An AffyBatch object
TGT	the target intensity to which the arrays should be scaled
digits	number of significant digits for SF and SDT values
verbose	logical value. If TRUE it provides more detail of the SF and SDT calculations.
plot.histogram	logical value. if TRUE it plots a histogram of intensities
celfile.path	character denoting the path for the *.CEL files corresponding to afbatch

Details

Calculates SF and SDT factors using the algorithms described in the Affymetrix Statistical Algorithms Description Document. The SF and SDT may be used in the calculation of S-Score values, or may be useful in their own right. One SF and SDT value is calculated for each GeneChip, which are arranged in the same order as the columns in the AffyBatch object.

Value

computeSFandSDT returns a list containing the following components:

SF	SF values, one for each GeneChip
SDT	SDT values, one for each GeneChip

Note

Based on Affymetrix MAS5 Statistical SDK source code http://www.affymetrix.com/Auth/support/developer/stat_sdk/STAT_SDK_source.zip, as well as C++ code by Li Zhang and Delphi code by Robnet Kerns

Author(s)

Richard Kennedy <rkennedy@vcu.edu>

References

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

Examples

```
if (length(dir(pattern=".cel$")) != 0) {  
  
  ## Read in the *.CEL files  
  abatch <- ReadAffy()  
  
  ## compute SF and SDT  
  SfSdt <- computeSFandSDT(abatch)  
  
  ## show verbose output  
  SfSdt <- computeSFandSDT(abatch,verbose=TRUE)  
  
  ## plot PM and MM histograms for each *.CEL file  
  SfSdt <- computeSFandSDT(abatch,plot.histogram=TRUE)  
  
}
```

computeZoneIInfo *Compute zone background and noise*

Description

Computes the background and noise for a given zone of a single Affymetrix GeneChip

Usage

```
computeZoneIInfo(ZoneInfo, NumberBGCells)
```

Arguments

ZoneInfo vector of intensities in a given zone
NumberBGCells number of background cells for the GeneChip

Details

Calculates background and noise for a zone using the algorithms described in the Affymetrix Statistical Algorithms Description Document. This is an internal function that will generally not be accessed directly.

Value

computeZoneIInfo returns a list containing the following components:

background background value for the given zone
noise noise value for the given zone

Note

Based on Affymetrix MAS5 Statistical SDK source code http://www.affymetrix.com/Auth/support/developer/stat_sdk/STAT_SDK_source.zip, as well as C++ code by Li Zhang and Delphi code by Robnet Kerns

Author(s)

Richard Kennedy <rkennedy@vcu.edu>

References

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

computeZonenoise *Computes zone noise*

Description

Computes the noise (average standard error) of the probe intensities for a single Affymetrix GeneChip

Usage

```
computeZonenoise(index, intensity, stdv, npixels, bgCells)
```

Arguments

index	vector of indices for probes in the given zone
intensity	vector of intensities for the GeneChip
stdv	vector of standard deviations for the GeneChip
npixels	vector containing number of pixels for each probe of the GeneChip
bgCells	number of background cells on the GeneChip

Details

Calculates the noise (average standard error) of the probes in a given zone, using the algorithms described in the Affymetrix Statistical Algorithms Description Document, for a single GeneChip. This is an internal function that will generally not be accessed directly.

Value

the noise of the probes for the given array

Note

Based on Affymetrix MAS5 Statistical SDK source code http://www.affymetrix.com/Auth/support/developer/stat_sdk/STAT_SDK_source.zip, as well as C++ code by Li Zhang and Delphi code by Robnet Kerns

Author(s)

Richard Kennedy <rkennedy@vcu.edu>

References

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

trimMean

Compute trimmed mean for a vector

Description

Computes the trimmed mean for a vector. Note that this implementation follows the Affymetrix code, which gives different results than the standard R function mean().

Usage

```
trimMean(vec, p1, p2)
```

Arguments

vec	vector of values
p1	lower percentage for trimming
p2	upper percentage for trimming

Details

The details can be found in the given reference.

Value

A numeric value

Note

Based on Affymetrix MAS5 Statistical SDK source code http://www.affymetrix.com/Auth/support/developer/stat_sdk/STAT_SDK_source.zip

Author(s)

Richard Kennedy <rkennedy@vcu.edu>

References

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

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