# Package 'beadarraySNP'

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Title Normalization and reporting of Illumina SNP bead arrays

Type Package

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|--|
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# Description

Changes one of the levels of a cn.sum data structure

# Usage

```
alterCN(cn.sum, opa, value, updown)
```

# Arguments

| cn.sum | cn.sum structure to change     |
|--------|--------------------------------|
| opa    | opa panel within the structure |
| value  | the predicted value to change  |

updown the value has a higher (TRUE) or lower (FALSE) cn value

#### **Details**

The state in the cn.sum structure that has a predicted value of value will have it's associated associated inferred copy number increased (updown is TRUE) or decreased (updown is FALSE). The function makes sure that the copynumber values within a OPA panel have the same order as the predicted values.

#### Value

a new cn.sum data structure

## Author(s)

Jan Oosting

#### See Also

interactiveCNselect, createCNSummary, setRealCN

 ${\tt backgroundCorrect.SNP} \ \ \textit{Background correction}$ 

#### **Description**

Perform background correction on Illumina Golden Gate bead arrays

#### Usage

## **Arguments**

object SnpSetIllumina object

method character, method of correction

offset numeric, constant to add after correction

#### **Details**

Code has been ported from the limma package. The matrices Gb and Rb should be available in the arrayData slot of the object.

## Value

This function returns an SnpSetIllumina object with background corrected values in the G and R.

#### Author(s)

Jan Oosting, based on limma package by G. Smyth

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#### See Also

```
SnpSetIllumina-class, backgroundCorrect,
backgroundEstimate, normalizeBetweenAlleles.SNP, normalizeWithinArrays.SNP
```

#### **Examples**

```
## Not run: data.bg<-backgroundCorrect.SNP(data.raw, "subtract")</pre>
```

 $back {\tt groundEstimate}$ 

Estimate background intensities from foreground intensity

#### Description

Background intensity from Illumina Golden Gate bead arrays are estimated based on several data models

#### Usage

```
backgroundEstimate(object,method=c("minimum", "mode","intmin",
   "anglemode"), maxmode=3000, bincount=40, maxangle=0.3, subsample="OPA")
```

## **Arguments**

object SnpSetIllumina object
method chracter, data model to use

maxmode numeric, maximum intensity for mode for method="mode"

bincount numeric, for method="intmin", see details

maxangle numeric in radians, maximum theta for mode for method="anglemode"

subsample factor or column name in featureData slot

# **Details**

The Illumina software does not provide background values in the output. Some models can be used to estimate background from the raw data intensities.

minimum: The allele specific minimum intensity is used.

mode: This model assumes that the first mode of the density of the intensities is determined by the zero-allele in the data, see ref. The signal intensity of the zero-allele should be zero, therefore this is considered the background value.

intmin: This model assumes there is crosstalk between the alleles, and background increases with the intensity of the other allele. The range between 0 and the maximum of the other allele is divided in bincount bins, and the minimum for this allele is determined for probes where the other allele falls in a bin. A linear fit is determined though the minimum values to obtain a gradually increasing value

anglemode: This model finds the density modes closest to 0 and  $\frac{\pi}{2}$  for polar transformed intensities, and uses this to determine background.

BeadstudioQC 5

## Value

This function returns an SnpSetIllumina object. The Rb and Gb matrices in the assayData slot contain estimated background values.

#### Author(s)

Jan Oosting

#### See Also

SnpSetIllumina-class, backgroundCorrect.SNP

BeadstudioQC

Quality control of Beadstudio report files

#### **Description**

When data has been imported using a Beadstudio samplesheet and reportfile, these functions can be used to generate quality measures

## Usage

```
BeadstudioQC(object, QClist = list(), arrayType = "Sentrix96")
pdfBeadstudioQC(QClist, basename = "beadstudio", by = 10)
```

# Arguments

object SnpSetIllumina object.

QClist list, result of previous call to BeadstudioQC

arrayType character, type of array

basename character, prefix for PDF files. This name will be added before the Barcode of

the chip

by integer, number of samples in barplot, see reportSamplePanelQC

## Value

The BeadstudioQC function generates a list of QCIIlumina objects The pdfBeadstudioQC function generates a pdf-file for each QCIIlumina object in the list

# Author(s)

J. Oosting

#### See Also

```
pdfQC,calculateQCarray
```

6 calculateLOH

| calculateLOH | Determine LOH in experiment  |
|--------------|------------------------------|
| Calcalacelon | Determine Boll in experiment |

#### **Description**

Using pairings of normal and tumor samples the LOH pattern is determined

## Usage

```
calculateLOH(object, grouping, NorTum = "NorTum", ...)
calculateLair(object, grouping = NULL, NorTum = "NorTum", min.intensity = NULL,
    use.homozygous.avg = FALSE)
```

## **Arguments**

object SnpSetIllumina object

grouping Factor to show which samples belong together (are of the same individual)

NorTum character vector or factor. Elements containing "N" are considered to be the

normal sample

min.intensity numeric
use.homozygous.avg

logical

... extra arguments for link{heterozygousSNPs}

#### **Details**

The heterozygous SNPs of the normal sample are inspected for changes. SNPs where the genotype of the test sample are homozygous are set to TRUE

#### Value

For calculateLOH a SnpSetIllumina object with 1oh and nor.gt matrices in assayData. 1oh is a logical matrix, and nor.gt is a character matrix containing the genotypes of the corresponding normal sample For calculateLair a SnpSetIllumina object with 1air matrix in assayData. 1air is the lesser allele intensity ratio. If a corresponding normal sample is found, it is taken as reference. Else the genotypes of normal samples are taken as a reference

## Author(s)

Jan Oosting

#### See Also

```
SnpSetIllumina-class
```

calculateQCarray 7

# **Description**

Retrieves QC and identifying information of Illumina Sentrix arrays.

#### Usage

```
calculateQCarray(object, QCobject = NULL, arrayType="Sentrix96")
```

#### **Arguments**

object SnpSetIllumina object. Should contain information of a single Sentrix array

and a single type of OPA panel

QCobject QCIllumina-class object: If set the information in the object is amended with

data from the SnpSetIllumina object

arrayType character, see arrayType

#### **Details**

Sample summary values are mapped to the physical layout of the Sentrix array using the Row and Col columns of the phenoData slot. These will be available when read.SnpSetIllumina is used to create SnpSetIllumina objects.

Use successive calls to calculateQCarray to process Sentrix arrays with multiple probe panels.

If data is read using a samplesheet that defines manifest files it is possible to handle data with multiple manifests and/or multiple Sentrix arrays

#### Value

A QCIllumina object, when multiple arrays were combined a list of QCIllumina objects

#### Author(s)

Jan Oosting

#### See Also

```
link{QCIllumina-class}, link{reportSamplePanelQC}, link{plotQC}
```

```
## Not run: QC<-calculateQCarray(data.raw1)
## Not run: QC<-calculateQCarray(data.raw2,QC)</pre>
```

8 compareGenotypes

| 0000000 | Canatunaa  |
|---------|------------|
| combar  | eGenotypes |

Compare genotypes

# Description

Pairwise comparison of genotypes between unaffected and affected tissue from the same subject

## Usage

```
compareGenotypes(genotypeT, genotypeN)
```

## Arguments

genotypeT character or logical vector, genotypes of affected tissue

genotypeN character or logical vector with same length as genotypeT, genotypes of unaf-

fected, normal tissue

#### **Details**

Heterozygous probes have one the following values. TRUE, 'H' or 'AB'. All other values are considered homozygous. The primary purpose of the method is to find probes with loss of heterozygosity (LOH), where the unaffected probe is heterozygous and the affected is called homozygous.

#### Value

A vector with the same length as the arguments where each element can have one of four values

| 'u' | Uninformative: both affected and normal are homozygous |
|-----|--|
| 'i' | Informative: both affected and unaffected heterozygous |
| '1' | Loss: unaffected heterozygous, affected homozygous     |
| 'a' | Artefact: unaffected homozygous, affected heterozygous |

# Author(s)

Jan Oosting

#### See Also

```
heterozygousSNPs
```

```
data(chr17.260)
compareGenotypes(exprs(chr17.260)[,"514TV"],exprs(chr17.260)[,"514NP"])
```

copynumberConversion 9

copynumberConversion Conversion to Copynumber analysis objects

# **Description**

SnpSetIllumina objects are converted to other objects for numerical analysis

## Usage

```
convert2aCGH(object,normalizedTo=2,doLog=TRUE,organism="hsa")
convert2SegList(object,normalizedTo=2,doLog=TRUE,organism="hsa")
```

## Arguments

object SnpSetIllumina object

normalizedTo numeric, 'normal' copynumber datavalue for object doLog logical, perform logarithmic transformation (log2)

organism character, organism used in object. Currently 'hsa' and 'mmu' are recognized.

Used to convert sex chromosomes to their proper numerical representation

#### **Details**

These functions produce objects that can be used by the analysis functions in the aCGH or snapCGH packages. The SnpSetIllumina intensity values are stored in a linear scale. Both types of objects assume a logarithmic scale, so by default the values are transformed to a log2 scale centered around 0.

#### Value

convert2aCGH returns a aCGH object as used in the aCGH package. convert2SegList returns a SegList object as used in the snapCGH package.

#### Author(s)

Jan Oosting

#### See Also

SnpSetIllumina-class, aCGH-class, SegList-class

10 createCNSummary

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## **Description**

Create a summary object of the genomic copy number states in a sample of segmented data

# Usage

```
createCNSummary(object, sample, dnaIndex=1, subsample = "OPA")
```

## Arguments

object SNPSetIllumina object after segmentation segmentate

sample SampleName or index of the sample for which to create the summary

dnaIndex Measured DNA index of the sample

subsample factor or column name in featureData slot

#### **Details**

The segments within a sample are assigned a copy number value. When the inferred slot in assayData is empty, all segments will be set to 2. Otherwise the values are recovered from the inferred slot. Gender is taken into account for the sex chromosomes.

#### Value

list with the following elements

dnaIndex same as parameter dnaIndex

CN. total.nrm Total expected copynumber for a 'normal' specimen ~ 2\*featurecount states data.frame with columns opa, count ,intensity, copynumber

This list can be used as the cn.sum argument for plotGoldenGate40PA, alterCN, getDNAindex and setRealCN

#### Author(s)

Jan Oosting

#### See Also

```
segmentate, alterCN, plotGoldenGate4OPA
```

dist.GT

dist.GT

dist.GT

# Description

Calculate distance matrix based of differences in genotype calls

# Usage

```
dist.GT(object)
```

# Arguments

object

SnpSetIllumina object

# **Details**

Calculates distances between samples as percentage of differences in genotype

## Value

'dist.GT' returns an object of class 'dist'

# Author(s)

Jan Oosting

## See Also

```
dist, hclust
```

```
data(chr17.260)
plot(hclust(dist.GT(chr17.260)))
```

12 GenomicReports

#### **Description**

Create reports for all samples in a dataset.

#### Usage

```
reportChromosomesSmoothCopyNumber(snpdata, grouping, normalizedTo=2,
    smooth.lambda=2, ridge.kappa=0, plotLOH=c("none", "marker", "line", "NorTum"),
    sample.colors = NULL, ideo.bleach=0.25, ...)
reportSamplesSmoothCopyNumber(snpdata, grouping, normalizedTo=2,
    smooth.lambda=2, ridge.kappa=0, plotLOH=c("none", "marker", "line", "NorTum"),
    sample.colors=NULL, ...)
reportGenomeGainLossLOH(snpdata, grouping, plotSampleNames=FALSE, sizeSampleNames=4,
    distance.min, upcolor="red", downcolor="blue", lohcolor="grey", hetcolor="lightgrey",
    lohwidth=1, segment=101, orientation=c("V","H"), ...)
reportChromosomeGainLossLOH(snpdata, grouping, plotSampleNames=FALSE, distance.min,
    upcolor="red", downcolor="blue", lohcolor="grey", hetcolor="lightgrey", proportion=0.2,
    plotLOH=TRUE, segment=101, ...)
reportGenomeIntensityPlot(snpdata, normalizedTo=NULL, subsample=NULL, smoothing=c("mean", "quant"),
    dot.col="black", smooth.col="red", ...)
```

## Arguments

| snpdata                                 | SnpSetIllumina object.  |
|---|---|
| grouping                                | factor, elements with same value are plotted together. Defaults to groups of 4 in order of the samples in the object. |
| normalizedTo                            | numeric, a horizontal line is drawn at this position.   |
| smooth.lambda                           | smoothing parameter for quantsmooth.  |
| ridge.kappa                             | smoothing parameter for quantsmooth.  |
| plotLOH                                 | indicate regions or probes with LOH, see details.   |
| <pre>sample.colors plotSampleName</pre> | vector of color.  |
|   | logical.  |
| sizeSampleName                          | S   |
|   | numeric, margin size for sample names.  |
| distance.min                            | numerical.  |
| upcolor                                 | color.  |
| downcolor                               | color.  |
| lohcolor                                | color.  |
| hetcolor                                | color.  |

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lohwidth numerical, relative width of the LOH part of the sample segment integer. ["V","H"], vertical or horizontal orientation of plot. orientation proportion numerical, proportion of the plot to use for idiogram annotation subsample character, or factor of length of features smoothing Type of smoothing per chromosome. dot.col color. smooth.col color. ideo.bleach numeric [0,1]

#### **Details**

. . .

The first function creates plots for each group and each chromosome in the dataset. The second function creates full genome plot for each group in the dataset. Beware that a lot of plots can be created, and usually you should prepare for that, by redirecting the plots to pdf or functions that create picture files like jpg, png, bmp.

arguments are forwarded to plot or getChangedRegions.

#### Value

These functions are executed for their side effects

# Author(s)

Jan Oosting

#### See Also

quantsmooth,prepareGenomePlot,pdfChromosomesSmoothCopyNumber,pdfSamplesSmoothCopyNumber

```
data(chr17.260)
chr17nrm <- standardNormalization(chr17.260)
par(mfrow = c(4,2), mar = c(2,4,2,1))
reportChromosomesSmoothCopyNumber(chr17nrm, grouping=pData(chr17.260)$Group,smooth.lambda = 4)</pre>
```

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 ${\tt GetBeadStudioSampleNames}$ 

Extract samplenames from a report file

## **Description**

Extract the samplenames from a report file that was created as a final report from Illumina Beadstudio

# Usage

```
GetBeadStudioSampleNames(reportfile)
```

## **Arguments**

reportfile character, name of report file

#### **Details**

This function will read the report file, and extract the sample names from the Sample ID column

#### Value

character vector

# Author(s)

Jan Oosting

#### See Also

read.SnpSetIllumina

getDNAindex

Calculate the DNA index based on assigned copy number values to probes

## **Description**

Calculate the DNA index based on assigned copy number values to probes

## Usage

```
getDNAindex(cn.sum)
```

heterozygosity 15

#### Arguments

cn.sum list with elements dnaIndex, CN.total.nrm, states, see createCNSummary

#### Value

scalar. DNA index of an unaffected sample is 1

#### Author(s)

Jan Oosting

#### See Also

createCNSummary, plotGoldenGate40PA

heterozygosity

Find regions of homozygous SNPs

## **Description**

Analyze affected material without corresponding unaffected material in order to find regions that contain stretches of homozygous SNPs as an indication of loss of heterozygosity (LOH)

#### Usage

```
heterozygosity(genotype, decay = 0.8, threshold = 0.1)
```

#### **Arguments**

genotype character or logical vector, genotypes of affected tissue

decay numeric in range (0,1)threshold numeric in range (0,1)

# **Details**

The method calculates how long the stretch of homozygous SNPs is for each element delay and threshold can be set to skip individual heterozygous probes in a longer stretch of homozygous probes. The default setting tolerate 1 erroneous heterozygous SNP every 10 homozygous SNPs. Set threshold at 1 to stop discarding hetrozygous SNPs

#### Value

A numeric vector with the same length as genotype is returned. Higher values, of 15 and higher, indicate regions of LOH

#### Author(s)

Jan Oosting

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#### See Also

compareGenotypes, heterozygousSNPs

#### **Examples**

```
data(chr17.260)
plot(heterozygosity(exprs(chr17.260)[,"514TV"]))
```

heterozygousSNPs

Retrieve heterozygous SNPs

#### **Description**

Heterozygous SNPs are determined based on quality score criteria

#### Usage

```
heterozygousSNPs(object, threshold=0.9, useQuality=TRUE, relative=TRUE, percentile=FALSE)
```

#### **Arguments**

object class SnpSetIllumina

threshold numeric (0:1) minimum quality score to be called heterozygous

useQuality logical, use quality score

relative logical, use quality score relative to GTS, see details percentile logical, use percentage of probes above threshold

#### **Details**

This function presumes that the specificity for determining heterozygity is more important than the sensitivity, and will therefore only call probes heterozygous if that can be done with high certainty. The Illumina genotyping software calculates two quality measures: gen train score (GTS) and gen call score (GCS). The GTS is a measure for how well clusters can be recognized in a training set. This value is probe specific, and the same for all samples in an experiment. The GCS is a probe-specific, sample specific value that measures how close a probe in a sample is to the clusters determined in the training step. This value is always lower than the GTS for a probe.

read. SnpSetIllumina will put GCS into the callProbability element of the assaydata slot and the GTS into the featureData slot. The function uses these locations to retrieve the necessary information.

If relative is FALSE then the raw GCS values are compared to the threshold. In this case a threshold of around 0.5 should be used. If relative is TRUE then GCS/GTS is compared to the threshold and threshold should be around 0.9.

With percentile=TRUE the threshold quantile is calculated for each sample, and only probes with higher scores can be called heterozygous. A threshold of around 0.2 seems to work fine usually.

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## Value

This function returns a logical matrix with same dimensions as object.

#### Note

The purpose of the function is to separate heterozygous probes from non-heterozygous probes. In tumor samples the determination of the genotype can be difficult, because of aneuploidy and the fact that a sample is often a mixture of normal and tumor cells.

#### Author(s)

Jan Oosting

#### See Also

```
SnpSetIllumina-class
```

## **Examples**

```
data(chr17.260)
plot(heterozygosity(heterozygousSNPs(chr17.260[,"514TV"])),col="red",pch="x")
points(heterozygosity(exprs(chr17.260)[,"514TV"]))
```

Illumina Genomic data *Illumina example data* 

# Description

These datasets are subsets of an experiment to test the applicability of paraffin embedded material in Illumina SNP arrays

## Usage

```
data(chr17.260)
data(QC.260)
```

## **Format**

chr17.260 is a SnpSetIllumina object with data from chromosome 17 of 24 samples. QC.260 is a QCIllumina object with summary data of 96 samples of a single SAM array

18 interactiveCNselect

interactiveCNselect

Interactive assignment of copynumbers to genomic segments

# Description

This function plots the genomic view of a sample, and allows the assignment of a discrete copy number to each segment

#### Usage

```
interactiveCNselect(object, sample = 1, dnaIndex)
```

# **Arguments**

object class SnpSetIllumina after segmentation

sample Sample identifier within object

dnaIndex numeric, measured DNA index of the sample (1=normal)

#### **Details**

The user can interactively assign discrete, integer copy number values to each segment. This is done by either clicking in the lower part of a panel to decrease the copy number, or in the higher part of a panel to increase the copy number. The order of copy number values is always maintained; a segment with a lower raw value can not get a higher copy number assigned then a segment with a higher raw copy number value.

#### Value

```
list, see createCNSummary
```

# Author(s)

Jan Oosting

#### See Also

```
segmentate, alterCN, plotGoldenGate4OPA createCNSummary
```

normalizeBetweenAlleles.SNP

between Allele normalization

## **Description**

Perform between Allele normalization on Illumina Golden Gate bead arrays

## Usage

```
normalizeBetweenAlleles.SNP(object,method=c("quantile"),subsample="OPA")
```

#### **Arguments**

object class SnpSetIllumina method char, type of normalization

subsample factor with length number of features in object or char, column name in featureData

slot

#### **Details**

This function performs a quantile normalization between the Red and Green channels for each sample. The rationale for this procedure stems from the fact that the allele frequencies within each channel are always very similar, even in the presence of genomic abnormalities.

#### Value

This function returns an SnpSetIllumina object.

# Author(s)

Jan Oosting

#### See Also

 ${\tt SnpSetIllumina-class,} normalize {\tt WithinArrays.SNP,} background {\tt Correct.SNP} \\$ 

```
data(chr17.260)
data.nrm<-normalizeBetweenAlleles.SNP(chr17.260)</pre>
```

normalizeBetweenSubsamples.SNP

Normalization between subsamples

## **Description**

Quantile normalization between subsamples within a single SnpSetIllumina object

#### Usage

```
normalizeBetweenSubsamples.SNP(object, subsample = "OPA")
```

#### **Arguments**

object class SnpSetIllumina

subsample factor with length number of features in object or char, column name in featureData

slot

#### **Details**

Perform quantile normalization of the red and green channel between subsamples. This can be used in situations where multiple different assays that cover the same genomic regions (or whole genome) have been done on the same biological specimen. This function was introduced for version 5 Golden Gate Linkage analysis that consist of 4 assays of ~ 1500 probes. Where previous versions of this assay each targeted a number of chromosomes, in version 5 each assay covers the whole genome.

#### Value

This function returns an SnpSetIllumina object.

#### Author(s)

Jan Oosting

#### See Also

SnpSetIllumina-class, normalize Between Alleles. SNP, normalize Within Arrays. SNP, background Correct. SNP, and the support of the property of the support of the suppor

```
data(chr17.260)
data.nrm<-normalizeBetweenSubsamples.SNP(chr17.260)</pre>
```

normalizeLoci.SNP 21

|--|

# Description

Perform locus normalization on Illumina Golden Gate bead arrays

# Usage

# Arguments

| object      | object class SnpSetIllumina  |
|-------------|--|
| method      | character. If "normals" then all normal samples in the dataset are used as the invariant set. If "paired" then affected samples are normalized to their paired normal samples. "alleles" fits a linear model between the B-allele ratio and the signal intensity and normalizes for that |
| NorTum      | logical or character vector or name of column in pData slot. depicts the normal, unaffected samples in the dataset. In a character vector these should have the value "N"  |
| Gender      | logical or character vector or name of column in pData slot. depicts the female samples in the dataset and is used to normalize the sex chromosomes. In a character vector these should have value "F"   |
| Subject     | factor or name of or column in pData slot. This factor is used to pair the samples when method is "paired"   |
| normalizeTo | normalizeTo numeric. The average copy number of the sample.  |
| trig        | Logical, use geometric distance of intensity. Otherwise use addition of intensities  |

# Details

This function is usually performed in the last step of normalization in order to obtain calculated copy numbers.

## Value

This function returns an SnpSetIllumina object.

# Author(s)

Jan Oosting

#### See Also

 ${\tt SnpSetIllumina,} normalize {\tt WithinArrays.SNP}, normalize {\tt BetweenAlleles.SNP}, normalize {\tt BetweenAlleles.SNP}, normalize {\tt SnpSetIllumina,} norm$ 

## **Examples**

```
data(chr17.260)
data.nrm<-normalizeLoci.SNP(chr17.260)</pre>
```

normalizeWithinArrays.SNP

Within Array normalization

# **Description**

Perform within array normalization on Illumina Golden Gate bead arrays.

## Usage

```
normalizeWithinArrays.SNP(object, callscore=0.5, normprob=0.5, quantilepersample=FALSE, relative=FALSE, fixed=FALSE, useAll=FALSE, subsample="OPA", Q.scores="callProbability")
```

# Arguments

| object            | class SnpSetIllumina.  |  |
|-------------------|--|--|
| callscore         | numeric with range 0:1, threshold for probe inclusion.   |  |
| normprob          | numeric with range 0:1, target quantile for normalization. The default is to divide the sample intensities by the median of the selected subset. |  |
| quantilepersample |  |  |
|                   | logical. If TRUE then the threshold is determined for each sample, else it is experiment wide. This is only relevant when fixed is FALSE.        |  |
| relative          | logical. If TRUE then the ratio of GCS and GTS is used, else only the GCS is used as the quality score.  |  |
| fixed             | logical. If TRUE then callscore is the fixed threshold for the quality score, else the probes above the quantile callscore are used.             |  |
| useAll            | logical. If TRUE then all probes in the dataset are eligible as the invariant set, else only the heterozygous SNPs.                              |  |
| subsample         | factor or column name in featureData slot, the levels of the factor are treated separately.  |  |
| Q.scores          | name of assayData() element, or numeric matrix of appropriate size. Quality scores to select high quality SNPs                                   |  |

## **Details**

The function uses high quality heterozygous SNPs as an invariant set with the assumption that these have the highest probability of coming from unaffected regions of the genome. Most of the arguments are used to determine the quality of the call.

## Value

This function returns a SnpSetIllumina object.

#### Author(s)

Jan Oosting

#### See Also

 ${\tt SnpSetIllumina, normalizeLoci.SNP, backgroundCorrect.SNP, normalizeBetweenAlleles.SNP}$ 

# Examples

## **Description**

Functions that help create pdf reports

## Usage

```
pdfChromosomesSmoothCopyNumber(object, filename, ...)
pdfSamplesSmoothCopyNumber(object, filename, ...)
pdfChromosomeGainLossLOH(object, filename, ...)
```

# **Arguments**

```
object SnpSetIllumina object
filename filename of output pdf file
... arguments for report functions
```

## **Details**

These functions set up and perfom reporting to pdf files.

#### Value

This function is used for its side effects

## Author(s)

Jan Oosting

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#### See Also

report Chromosomes Smooth Copy Number, report Samples Smooth Copy Number, report Chromosome Gain Loss LOH Copy Number, report Chromosome Gain Loud Chromosome Gain Chromosome Gain Chromosome Gain Chromosome Gain

## **Examples**

```
## Not run: data(chr17.260)
## Not run: data.nrm<-standardNormalization(chr17.260)
## Not run: pdfChromosomesSmoothCopyNumber(data.nrm, "Chr17.pdf", grouping=pData(data.nrm)$Group,smooth.lambda</pre>
```

pdfQC

**QCreport** 

# Description

Create PDF file with experimental quality control plots

#### Usage

```
pdfQC(object, filename = "arrayQC.pdf", by = 10)
```

## **Arguments**

object QCIllumina object, or list of QCIllumina objects

filename character, output pdf filename

by number of samples in barplot, see reportSamplePanelQC

#### **Details**

This function creates a pdf file with QC information. The first page contains 8 plotQC panels showing the spatial distribution of intensities on a SAM plate. The following page(s) contain the output of reportSamplePanelQC

## Value

A PDF file is produced

#### Author(s)

Jan Oosting

#### See Also

```
plotQC, reportSamplePanelQC, QCIllumina-class
```

plotGoldenGate4OPA 25

| plotGoldenGate40PA | Plot Golden Gate genomic view |  |  |
|--------------------|-------------------------------|--|--|
|--------------------|-------------------------------|--|--|

# Description

Plots a full genome view based on 4 subsamples of Illumina Golden Gate data

## Usage

```
plotGoldenGate40PA(object, cn.sum = NULL, sample = 1, plotRaw = FALSE, main = NULL, interact = FALSE,
plotGenomePanels(object, cn.sum = NULL, sample = 1, plotRaw = FALSE, main = NULL, interact = FALSE, al
```

# Arguments

| object   | class SnpSetIllumina  |
|----------|---|
| cn.sum   | list containing genomic states, see createCNSummary   |
| sample   | identifier to select the sample within the object   |
| plotRaw  | logical, plot raw data points   |
| main     | character, Title of plot  |
| interact | logical, plot should be usable for interactive copy number determination interactiveCNselect                                    |
| allLair  | logical, TRUE: plot all LAIR values, FALSE: only plot LAIR values from probes that are heterozygous in the paired normal sample |
| panels   | list, vectors of chromosomes for each panel   |
|          | extra arguments are formwarded to plot  |

#### **Details**

prepare interactive selection

#### Value

list, see createCNSummary

## Author(s)

Jan Oosting

# See Also

```
segmentate, alterCN, interactiveCNselect createCNSummary
```

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plotQC

Spatial plots of array QC information

## **Description**

Plots array wide summary information using the layout of the physical medium

# Usage

```
plotQC(object, type)
```

## **Arguments**

object object that contains QC information. e.g. QCIllumina-class

type character, the type of information to plot, currently the following types are

supported: "intensityMed", "greenMed", "redMed", "validn", "annotation" and "samples"

#### Value

The function is used for its side effects

#### Author(s)

Jan Oosting

#### See Also

```
pdfQC, reportSamplePanelQC
```

# **Examples**

```
data(QC.260)
plotQC(QC.260, "greenMed")
```

PolarTransforms

Polar transformations

## **Description**

Perform polar transforms on Illumina Golden Gate bead arrays

## Usage

```
RG2polar(object,trig=FALSE)
polar2RG(object,trig=FALSE)
```

OCaccessors 27

## **Arguments**

object SnpSetIllumina object

trig Logical, use geometric distance intensity. Otherwise use addition of intensities

#### **Details**

RG2polar transforms the R and G matrices to theta and intensity matrices. Note that the intensity value is the sum of R and G and not the geometric distance to the origin.

polar2RG performs the reverse transformation

## Value

This function returns an SnpSetIllumina object.

#### Author(s)

Jan Oosting

#### See Also

```
SnpSetIllumina-class
```

## **Examples**

```
data(chr17.260)
data.polar<-RG2polar(chr17.260)
plot(assayData(data.polar)$theta,assayData(data.polar)$intensity)</pre>
```

**QCaccessors** 

Accessor methods for QC objects

## **Description**

These generic functions set and retrieve properties of quality control objects like QCIllumina-class

#### Usage

```
arrayType(object)
arrayType(object)<- value
arrayID(object)
arrayID(object)<- value</pre>
```

# Arguments

object Object, possibly derived from class QCIllumina-class.

value character.

QCIllumina-class

#### **Details**

Currently the following types of arrays are supported

"Sentrix96": Sentrix array, 12 columns, 8 rows "Sentrix16": Sentrix array, 2 columns, 8 rows

"Slide12": Slide with 12 samples

#### Value

arrayType and arrayID return a character value

#### Author(s)

Jan Oosting

QCIllumina-class

Class "QCIllumina"

## **Description**

Container of QC information on arrays that contain multiple samples.

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

Objects can be created by calls of the form new("QCIllumina", arrayType, arrayID, intensityMed, greenMed, redMe but are usually created by calculateQCarray.

#### **Slots**

```
arrayType: character, Type of array. See arrayType
arrayID: character, Array ID
intensityMed: numeric matrix, Median of intensity of samples
greenMed: numeric matrix, Median of green values
redMed: numeric matrix, Median of red values
callrate: numeric matrix, callrate of genotyping
hetPerc: numeric matrix, Percentage of heterozygotes
ptpdiff: numeric matrix, point-to-point difference, local estimate of variability
validn: numeric matrix, Number of valid probe values in samples
annotation: character matrix, Annotation of samples
samples: character matrix, Sample IDs
```

read.SnpSetIllumina 29

#### Methods

```
arrayID signature(object = "QCIllumina"): Returns type of array
arrayID<- signature(object = "QCIllumina"): Sets type of array. Currently only "Sentrix"
    is supported
arrayType signature(object = "QCIllumina"): Returns ID of array
arrayType<- signature(object = "QCIllumina"): Sets ID/Barcode of array
initialize signature(.0bject = "QCIllumina")
plotQC signature(object = "QCIllumina") character: plots spatial overview of QC information, type is one of c("intensityMed", "greenMed", "redMed", "validn", "annotation", "samples")</pre>
```

#### Author(s)

Jan Oosting

#### See Also

calculateQCarray

## **Description**

A SnpSetIllumina object is created from the textfiles created by the Illumina GenCall or BeadStudio software.

## Usage

```
read.SnpSetIllumina(samplesheet, manifestpath=NULL, reportpath=NULL,
  rawdatapath=NULL, reportfile=NULL, briefOPAinfo=TRUE, readTIF=FALSE,
  nochecks=FALSE, sepreport="\t", essentialOnly=FALSE, ...)
```

#### **Arguments**

| samplesheet  | a data.frame or filename, contains the sample sheet  |
|--------------|--|
| manifestpath | a character string for the path containing the manifests / OPA definition files, defaults to path of samplesheet |
| reportpath   | a character string for the path containing the report files, defaults to path of samplesheet                     |
| rawdatapath  | a character string for the path containing the intensity data files, defaults to path of samplesheet             |
| reportfile   | a character string for the name of BeadStudio reportfile   |

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brief0PAinfo logical, if TRUE then only the SNP name, Illumi code, chromosome and basepair

position are put into the featureData slot of the result, else all information

from the OPA file is put into the featureData slot

readTIF logical, uses beadarray package and raw TIF files to read data

nochecks logical, limited validity checks on beadstudio report files. See details

sepreport character, field separator character for beadstudio report files

essentialOnly logical, if TRUE then only the essential columns from a reportfile are included

into the result. See details

... arguments are forwarded to readIllumina and can be used to perform bead-

level normalization

#### **Details**

The text files from Illumina software are imported to a SnpSetIllumina object. Both result files from GenCall and BeadStudio can be used. In both cases the sample sheets from the experiments are used to select the proper data from the report or data files. The following columns from the sample sheet file are used for this purpose: 'Sample\_Name', 'Sentrix\_Position', and 'Pool\_ID'. The values in columns 'Sample\_Plate', 'Pool\_ID', and 'Sentrix\_ID' should be the same for all samples in the file, as this is the case for processed experiments. The contents of the sample sheet are put into the phenoData slot.

Zero values in the raw data signals are set to NA

Ideally the OPA manifest file containing SNP annotation should be available, these files are provided by Illumina. Columns 'IllCode', 'CHR', and 'MapInfo' are put into the featureData slot.

#### GenCall Data

In order to process experiments that were genotyped using the GenCall software, the arrays should be scanned with the setting <SaveTextFiles>true</SaveTextFiles> in the Illumina configuration file Settings.XML. 3 Types of files need to be present in the same folder: The sample sheet, .csv files containing signal intensity data, and the report file that contains the genotype information. For each sample in the sample sheet there should be a .csv file with the following file mask: [sam\_id]\_R00[yy]\_C00[xx].csv, where sam\_id is the Illumina ID for the SAM, and xx and yy are the column and row number respectively. From the report files the file with mask [Pool\_ID]\_LocusByDNA[\_ExpName].csv is used. 'Pool\_ID' is the OPA panel used, and '\_ExpName' is optional.

#### BeadStudio Data

To process experiments that were processed with BeadStudio, only two files are needed. The sample sheet and the Final Report file. The sample sheet must contain the same columns as for GenCall, the report file should contain the following columns: 'SNP Name', 'Sample ID', 'GC Score', 'Allelel - AB', 'Allelel - AB', 'GT Score', 'X Raw', and 'Y Raw'. 'SNP Name' and 'Sample ID' are used to form rows and columns in the experimental data, 'GC Score' is put in the callProbability matrix, 'Allelel - AB' and 'Allelel - AB' are combined into the call matrix, 'GT Score' is added to the featureData slot, 'X Raw' is put in the R matrix and 'Y Raw' in the G matrix. Other columns in the report file are added as matrices in the assayData slot, or columns in the featureData slot if values are identical for all samples in the reportfile. When nochecks is TRUE then only the 'SNP Name' and 'Sample ID' columns are required. The resulting object is now of class MultiSet

Sample sheets

To help generate a sample sheet for BeadStudio data a Sample\_Map.txt file can be converted to a sample sheet with the Sample\_Map2Samplesheet function. For Beadstudio reportfiles it is also possible to set samplesheet=NULL. In this case the phenoData slot will be fabricated from the sample names in the reportfile.

Manifest/OPA/annotation files

For BeadStudio reportfiles it is not necessary to have a Manifest file if the columns 'Chr' and 'Position' are available in the report file. Currently this is the only way to import data from Infinium arrays, because Illumina does not supply Manifest files for these arrays.

#### Value

This function returns an SnpSetIllumina object, or a MultiSet object when nochecks is TRUE.

#### Author(s)

Jan Oosting

#### See Also

```
SnpSetIllumina-class, Sample_Map2Samplesheet, readIllumina
```

#### **Examples**

```
# read a SnpSetIllumina object using example textfiles in data directory
datadir <- system.file("testdata", package="beadarraySNP")
SNPdata <- read.SnpSetIllumina(paste(datadir,"4samples_opa4.csv",sep="/"),datadir)</pre>
```

removeLowQualityProbes

Quality control of SnpSetIllumina objects

# **Description**

Remove probes form a SnpSetIllumina object that show a low quality throughout the experiment

#### Usage

```
removeLowQualityProbes(object, cutoff = 0.25)
```

#### **Arguments**

object SnpSetIllumina object

cutoff numeric

#### **Details**

Probes that have a median value below cutoff \* median value for the whole experiment are deleted from the object.

#### Value

SnpSetIllumina object

#### Author(s)

Jan Oosting

 ${\tt removeLowQualitySamples}$ 

Quality control of SnpSetIllumina objects

# **Description**

Remove samples from a SnpSetIllumina object that show a low quality

#### Usage

```
removeLowQualitySamples(object, min.intensity = 1500, min.gt = 100, subsample = "OPA")
```

# Arguments

object SnpSetIllumina-class object

min.intensity numeric. Samples that show a median intensity below this value in either Red

or Green channel are removed

min.gt numeric. Samples that have less than this amount of valid genotypes are re-

noved

subsample factor or column name in featureData slot of object

## Value

This function returns an SnpSetIllumina object.

#### Author(s)

Jan Oosting

```
data(chr17.260)
chr17.260<-removeLowQualitySamples(chr17.260,min.gt=10)</pre>
```

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renameOPA

Change the linkage panel in a dataset

# Description

Change the linkage panel in a dataset

## Usage

```
renameOPA(snpdata, newOPA)
```

# **Arguments**

snpdata SnpSetIllumina object

newOPA character, new linkage panel

## **Details**

In order to combine different versions of the linkage panels, this function makes it possible to map the equivalent SNPs in both datasets.

# Value

SnpSetIllumina object

# Author(s)

Jan Oosting

 ${\tt reportGenotypeSegmentation}$ 

plot genomic view

# Description

Create a figure that can be used for interactive work

## Usage

reportGenotypeSegmentation(object, plotRaw = TRUE, subsample = NULL, panels = 0, minProbes = 10, maxY

#### **Arguments**

object class SnpSetIllumina after segmentation

plotRaw logical subsample factor

panels number of panels on a page

minProbes minimum number of probes for a chromosome within a panel

maxY maximum value on vertical scale within panels

... arguments are forwrded to plot

#### Value

this function is used for its side effects

#### Author(s)

Jan Oosting

reportSamplePanelQC-methods

reportSamplePanelQC

# Description

Show raw intensity values for green and red channel for all samples in an experiment

# Usage

```
reportSamplePanelQC(object, by=10, legend=TRUE, ...)
```

## Arguments

object QCIllumina object

by numeric, number of samples in each plot

legend logical, create a final plot with a common legend for the barplots

... arguments are forwarded to barplot

```
data(QC.260)
par(mfrow=c(2,2))
reportSamplePanelQC(QC.260,by=8)
```

Sample\_Map2Samplesheet

Convert Beadstudio Sample Map file to samplesheet

#### **Description**

Create a samplesheet that can be used to import Illumina beadstudio data

#### Usage

```
Sample_Map2Samplesheet(samplemapfile, saveas = "")
```

#### **Arguments**

```
samplemapfile character, name of the SampleMap file
saveas character, optional, name of samplesheet file that can be used directly by read.SnpSetIllumina
```

#### **Details**

During the creation of a final reportfile from Beadstudio there is an option to create Map files. The Sample\_Map.txt file can be used to create an initial samplesheet for use in the read.SnpSetIllumina function

#### Value

A data. frame with the samplesheet

#### Author(s)

J. Oosting

#### See Also

read.SnpSetIllumina

segmentate

Segmentation for SnpSetIllumina objects

## **Description**

Use snapCGH package to perform segmentation

## Usage

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#### **Arguments**

object class SnpSetIllumina method char, type of segmentation

normalizedTo numeric

doLog logical, perform transformation before segmentation, see convert2seglist

doMerge logical, perform merging of close states

useLair logical, Also segmentate on lair

subsample factor

alpha numeric, probability threshold to distinguish segments

#### Value

 $SnpSetIllumina\ object\ with\ elements\ observed,\ \ states\ \ and\ \ predicted\ set\ in\ the\ AssayData\ slot$ 

# Author(s)

Jan Oosting

setRealCN

Integrate state information into SNP object

#### **Description**

Set calculated values of copy numbers in inferred element of AssayData slot

# Usage

```
setRealCN(object, sample, cn.sum, subsample="OPA")
```

# Arguments

object class SnpSetIllumina sample sample identifier

cn.sum list, see createCNSummary

subsample "OPA"

#### Value

SnpSetIllumina object with inferred element of AssayData slot set

#### Author(s)

Jan Oosting

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#### See Also

```
segmentate, alterCN, plotGoldenGate40PA createCNSummary
```

smoothed.intensity

Smooth intensity data

# Description

Create a table of smoothe intensity values

## Usage

```
smoothed.intensity(snpdata, smooth.lambda = 4, tau = 0.5)
```

#### **Arguments**

snpdata SnpSetIllumina object smooth.lambda smoothing parameter tau quantile to smooth

#### Value

Numerical matrix with same dimensions as data

# Author(s)

Jan Oosting

#### See Also

SnpSetIllumina-class

SnpSetIllumina

Class to Contain Objects Describing High-Throughput SNP Assays.

#### **Description**

Container for high-throughput assays and experimental metadata. SnpSetIllumina class is derived from eSet, and requires matrices R, G, call, callProbability as assay data members.

It supports featureData. Several visualization methods use columns CHR and MapInfo. The CHR column is used to handle sex chromosomes in a specific way. The OPA column is the default way to specify subsamples.

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#### **Extends**

Directly extends class eSet.

#### **Creating Objects**

```
new('SnpSetIllumina', phenoData = [AnnotatedDataFrame], experimentData = [MIAME], annotation =
```

SnpSetIllumina instances are usually created through new("SnpSetIllumina", ...). Arguments to new include call (a matrix of gentoypic calls, with features (SNPs) corresponding to rows and samples to columns), callProbability, G, R, phenoData, experimentData, and annotation. phenoData, experimentData, and annotation can be missing, in which case they are assigned default values.

#### Slots

Inherited from Biobase: eSet:

assayData: Contains matrices with equal dimensions, and with column number equal to nrow(phenoData). assayData must contain a matrix call with rows representing features (e.g., SNPs) and columns representing samples, a matrix callProbability describing the certainty of the call, and matrices R and G to describe allele specific intensities. The contents of these matrices are not enforced by the class. The assayData matrices Gb, Rb, intensity, theta are optional, but are either results or input for several methods of the class. Additional matrices of identical size may also be included in assayData. Class:AssayData.

```
phenoData: See eSet.
experimentData: See eSet.
annotation: See eSet.
```

featureData: annotation for SNPs, usually will contain a CHR and a MapInfo column for genomic localization.

#### Methods

Class-specific methods:

exprs(SnpSetIllumina), exprs(SnpSetIllumina, matrix)<- Access and set elements named
 call in the AssayData slot.</pre>

combine(SnpSetIllumina, SnpSetIllumina): performs union-like combination in both dimensions of SnpSetIllumina objects.

fData(SnpSetIllumina), fData(SnpSetIllumina, data.frame)<- Access and set the pData in the featureData slot.

calculateGSR(SnpSetIllumina) calculate ratio of Gentrain score and Gencall score. Creates GSR matrix in assayData. Should be performed before combining datasets.

calculateSmooth(object,smoothType) calculate smoothed data, creates smoothed matrix in assayData. smoothType can only be "quantsmooth" at the moment

sortGenomic(SnpSetIllumina) order the data by chromosome and position on the chromosome.

Derived from eSet:

SnpSetIllumina 39

```
sampleNames(SnpSetIllumina) and sampleNames(SnpSetIllumina)<-: See eSet.</pre>
featureNames(SnpSetIllumina), featureNames(SnpSetIllumina, value)<-: See eSet.</pre>
dims(SnpSetIllumina): See eSet.
phenoData(SnpSetIllumina), phenoData(SnpSetIllumina, value)<-: See eSet.</pre>
varLabels(SnpSetIllumina), varLabels(SnpSetIllumina, value)<-: See eSet.</pre>
varMetadata(SnpSetIllumina), varMetadata(SnpSetIllumina, value)<-: See eSet.</pre>
pData(SnpSetIllumina), pData(SnpSetIllumina, value) <-: See eSet.
varMetadata(SnpSetIllumina), varMetadata(SnpSetIllumina, value) See eSet.
experimentData(SnpSetIllumina),experimentData(SnpSetIllumina,value)<-: See eSet.
pubMedIds(SnpSetIllumina), pubMedIds(SnpSetIllumina, value) See eSet.
abstract(SnpSetIllumina): See eSet.
annotation(SnpSetIllumina), annotation(SnpSetIllumina, value) <- See eSet.
storageMode(eSet), storageMode(eSet, character)<-: See eSet.
featureData(SnpSetIllumina), featureData(SnpSetIllumina, AnnotatedDataFrame) <- See
object[(index): Conducts subsetting of matrices and phenoData and featureData components.
Standard generic methods:
initialize(SnpSetIllumina): Object instantiation, used by new; not to be called directly by the
validObject(SnpSetIllumina): Validity-checking method, ensuring that call, callProbability,
    G, and R are members of assayData. checkValidity(SnpSetIllumina) imposes this valid-
    ity check, and the validity checks of Biobase: class.eSet.
show(SnpSetIllumina) See eSet.
dim(SnpSetIllumina), ncol See eSet.
SnpSetIllumina[(index): See eSet.
SnpSetIllumina$, SnpSetIllumina$<- See eSet.</pre>
```

#### Author(s)

J. Oosting, based on Biobase eSet class

#### See Also

eSet

```
SnpSetSegments-class Class "SnpSetSegments"
```

#### **Description**

The SnpSetSegments class is a direct descendant of the SnpSetIllumina class, with an extra slot to define the genomic segments in each sample.

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

Objects can be created by calls of the form new("SnpSetSegments", assayData, phenoData, experimentData, annota

#### Slots

```
cn.segments: Object of class "list"
assayData: Object of class "AssayData" see "SnpSetIllumina"
phenoData: Object of class "AnnotatedDataFrame" see "SnpSetIllumina"
featureData: Object of class "AnnotatedDataFrame" see "SnpSetIllumina"
experimentData: Object of class "MIAME" see "SnpSetIllumina"
annotation: Object of class "character" see "SnpSetIllumina"
protocolData: Object of class "AnnotatedDataFrame" see "SnpSetIllumina"
.__classVersion__: Object of class "Versiones" "VersionedBiobase"
```

#### **Extends**

```
Class "SnpSetIllumina", directly. Class "eSet", by class "SnpSetIllumina", distance 2. Class "VersionedBiobase", by class "SnpSetIllumina", distance 3. Class "Versioned", by class "SnpSetIllumina", distance 4.
```

#### Methods

```
cn.segments signature(object = "SnpSetSegments"): ...
cn.segments<- signature(object = "SnpSetSegments", value = "list"): ...
initialize signature(.Object = "SnpSetSegments"): ...</pre>
```

#### Note

This class is under development, and not usable in the current form

#### Author(s)

Jan Oosting

#### References

Corver et.al. Can Res dec 2008

standardNormalization 41

#### See Also

```
segmentate
```

## **Examples**

```
showClass("SnpSetSegments")
```

standardNormalization Default complete normalization

## **Description**

Performs all steps in normalization at best settings as determined in ref.

# Usage

```
standardNormalization(snpdata)
```

## **Arguments**

snpdata

SnpSetIllumina object with raw data

#### **Details**

```
The function performs in the following steps snpdata<-normalizeBetweenAlleles.SNP(snpdata)
snpdata<-normalizeWithinArrays.SNP(snpdata,callscore = 0.8, relative = TRUE, fixed = FALSE, quan snpdata<-normalizeLoci.SNP(snpdata,normalizeTo = 2)
```

## Value

A SnpSetIllumina object with the G, R and intensity elements in assayData normalized to obtain values close to 2 on a linear scale for unaffected material.

# Author(s)

Jan Oosting

#### See Also

normalizeBetweenAlleles.SNP,normalizeWithinArrays.SNP,normalizeLoci.SNP

```
data(chr17.260)
data.nrm<-standardNormalization(chr17.260)</pre>
```

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