# Package 'minfi'

October 18, 2022

**Version** 1.42.0

Title Analyze Illumina Infinium DNA methylation arrays

**Description** Tools to analyze & visualize Illumina Infinium methylation arrays.

**Depends** methods, BiocGenerics (>= 0.15.3), GenomicRanges, SummarizedExperiment (>= 1.1.6), Biostrings, bumphunter (>= 1.1.9)

Suggests IlluminaHumanMethylation450kmanifest (>= 0.2.0), IlluminaHumanMethylation450kanno.ilmn12.hg19 (>= 0.2.1), minfiData (>= 0.18.0), minfiDataEPIC, FlowSorted.Blood.450k (>= 1.0.1), RUnit, digest, BiocStyle, knitr, rmarkdown, tools

Imports S4Vectors, GenomeInfoDb, Biobase (>= 2.33.2), IRanges, beanplot, RColorBrewer, lattice, nor1mix, siggenes, limma, preprocessCore, illuminaio (>= 0.23.2), DelayedMatrixStats (>= 1.3.4), mclust, genefilter, nlme, reshape, MASS, quadprog, data.table, GEOquery, stats, grDevices, graphics, utils, DelayedArray (>= 0.15.16), HDF5Array, BiocParallel

Collate AllGenerics.R MethylSet-class.R RatioSet-class.R RGChannelSet-class.R RGChannelSetExtended-class.R GenomicMethylSet-class.R GenomicRatioSet-class.R eSet\_methods.R utils.R IlluminaMethylationManifest-class.R IlluminaMethylationAnnotation-class.R minfiQC.R getSex.R dmpFinder.R blocks.R plot.R plotBetasByType.R preprocessRaw.R preprocessIllumina.R detectionP.R preprocessSwan.R preprocessQuantile.R preprocessNoob.R preprocessFunnorm.R qc.R read.450k.R read.meth.R read.meth2.R read.geo.R read.manifest.R bumphunter.R estimateCellCounts.R gaphunter.R compartments.R combineArrays.R DelayedArray\_utils.R

VignetteBuilder knitr

License Artistic-2.0

URL https://github.com/hansenlab/minfi

 ${\bf BugReports} \ {\tt https://github.com/hansenlab/minfi/issues}$ 

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# Description

Tools for analyzing and visualizing Illumina methylation array data. There is special focus on the 450k array; the 27k array is not supported at the moment.

# **Details**

The package contains a (hopefully) useful vignette; this vignette contains a lengthy description of the package content and capabilities.

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	blockFinder	Finds blocks of methylation differences for Illumina methylation arrays
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# Description

Finds blocks (large scale regions) of methylation differences for Illumina methylation arrays

# Usage

# Arguments

object	An object of class GenomicRatioSet.
design	Design matrix with rows representing samples and columns representing covariates. Regression is applied to each row of mat.
coef	An integer denoting the column of the design matrix containing the covariate of interest. The hunt for bumps will be only be done for the estimate of this coefficient.
what	Should blockfinding be performed on M-values or Beta values?
cluster	The clusters of locations that are to be analyzed together. In the case of microarrays, the clusters are many times supplied by the manufacturer. If not available the function clusterMaker can be used to cluster nearby locations.
cutoff	A numeric value. Values of the estimate of the genomic profile above the cutoff or below the negative of the cutoff will be used as candidate regions. It is possible to give two separate values (upper and lower bounds). If one value is given, the lower bound is minus the value.
pickCutoff	Should a cutoff be picked automatically?
pickCutoffQ	The quantile used for picking the cutoff using the permutation distribution.
nullMethod	Method used to generate null candidate regions, must be one of 'bootstrap' or 'permutation' (defaults to 'permutation'). However, if covariates in addition to the outcome of interest are included in the design matrix (ncol(design)>2), the 'permutation' approach is not recommended. See vignette and original paper for more information.
smooth	A logical value. If TRUE the estimated profile will be smoothed with the smoother defined by smoothFunction

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smoothFunction A function to be used for smoothing the estimate of the genomic profile. Two

 $functions\ are\ provided\ by\ the\ package:\ loess By {\tt Cluster}\ and\ run {\tt medBy Cluster}.$ 

B An integer denoting the number of resamples to use when computing null distri-

butions. This defaults to 0. If permutations is supplied that defines the number

of permutations/bootstraps and B is ignored.

permutations is a matrix with columns providing indexes to be used to scramble the data

and create a null distribution. If this matrix is not supplied and B>0 then these

indexes created using the function sample.

verbose Should the function be verbose?

bpSpan Smoothing span. Note that this defaults to a large value because we are search-

ing for large scale changes.

. . . further arguments sent to bumphunterEngine.

#### **Details**

The approximately 170,000 open sea probes on the 450k can be used to detect long-range changes in methylation status. These large scale changes that can range up to several Mb have typically been identified only through whole-genome bisulfite sequencing. blockFinder groups the averaged methylation values in open-sea probe clusters (See cpgCollapse) into large regions in which the bumphunter procedure is applied with a large (250KB+) smoothing window.

Note that estimating the precise boundaries of these blocks are constrained by the resolution of the array.

# Value

**FIXME** 

#### See Also

cpgCollapse, and bumphunter

bumphunter-methods

Methods for function bumphunter in Package minfi

### **Description**

Estimate regions for which a genomic profile deviates from its baseline value. Originally implemented to detect differentially methylated genomic regions between two populations, but can be applied to any CpG-level coefficient of interest.

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#### Usage

#### **Arguments**

object An object of class GenomicRatioSet.

design Design matrix with rows representing samples and columns representing covari-

ates. Regression is applied to each row of mat.

cluster The clusters of locations that are to be analyzed together. In the case of microar-

rays, the clusters are many times supplied by the manufacturer. If not available

the function clusterMaker can be used to cluster nearby locations.

coef An integer denoting the column of the design matrix containing the covariate

of interest. The hunt for bumps will be only be done for the estimate of this

coefficient.

cutoff A numeric value. Values of the estimate of the genomic profile above the cutoff

or below the negative of the cutoff will be used as candidate regions. It is possible to give two separate values (upper and lower bounds). If one value is given,

the lower bound is minus the value.

pickCutoff Should bumphunter attempt to pick a cutoff using the permutation distribution?

pickCutoffQ The quantile used for picking the cutoff using the permutation distribution.

maxGap If cluster is not provided this maximum location gap will be used to define clus-

ter via the clusterMaker function.

nullMethod Method used to generate null candidate regions, must be one of 'boots trap' or

'permutation' (defaults to 'permutation'). However, if covariates in addition to the outcome of interest are included in the design matrix (ncol(design)>2), the 'permutation' approach is not recommended. See vignette and original paper

for more information.

smooth A logical value. If TRUE the estimated profile will be smoothed with the

smoother defined by smoothFunction

smoothFunction A function to be used for smoothing the estimate of the genomic profile. Two

functions are provided by the package: loessByCluster and runmedByCluster.

useWeights A logical value. If TRUE then the standard errors of the point-wise estimates of

the profile function will be used as weights in the loess smoother loessByCluster.

If the runmedByCluster smoother is used this argument is ignored.

B An integer denoting the number of resamples to use when computing null distri-

butions. This defaults to 0. If permutations is supplied that defines the number

of permutations/bootstraps and B is ignored.

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permutations is a matrix with columns providing indexes to be used to scramble the data

and create a null distribution when nullMethod is set to permutations. If the bootstrap approach is used this argument is ignored. If this matrix is not supplied

and B>0 then these indexes are created using the function sample.

verbose logical value. If TRUE, it writes out some messages indicating progress. If FALSE

nothing should be printed.

type Should bumphunting be performed on M-values ("M") or Beta values ("Beta")?

... further arguments to be passed to the smoother functions.

#### **Details**

See help file for bumphunter method in the bumphunter package for for details.

#### Value

An object of class bumps with the following components:

tab The table with candidate regions and annotation for these.

coef The single loci coefficients.

fitted The estimated genomic profile used to determine the regions.

pvaluesMarginal

marginal p-value for each genomic location.

null The null distribution.
algorithm details on the algorithm.

# Author(s)

Rafael A. Irizarry, Martin J. Aryee and Kasper D. Hansen

#### References

AE Jaffe, P Murakami, H Lee, JT Leek, MD Fallin, AP Feinberg, and RA Irizarry. *Bump hunting to identify differentially methylated regions in epigenetic epidemiology studies*. International Journal of Epidemiology (2012) 41(1):200-209. doi:10.1093/ije/dyr238

### See Also

bumphunter

# Examples

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combineArrays

A method for combining different types of methylation arrays into a virtual array.

#### Description

A method for combining different types of methylation arrays into a virtual array. The three generations of Illumina methylation arrays are supported: the 27k, the 450k and the EPIC arrays. Specifically, the 450k array and the EPIC array share many probes in common. This function combines data from the two different array types and outputs a data object of the user-specified type. Essentially, this new object will be like (for example) an EPIC array with many probes missing.

### Usage

```
## S4 method for signature 'RGChannelSet, RGChannelSet'
combineArrays(object1, object2,
                  outType = c("IlluminaHumanMethylation450k",
                               "IlluminaHumanMethylationEPIC"),
                  verbose = TRUE)
## S4 method for signature 'MethylSet, MethylSet'
combineArrays(object1, object2,
                  outType = c("IlluminaHumanMethylation450k",
                               "IlluminaHumanMethylationEPIC",
                               "IlluminaHumanMethylation27k"),
                  verbose = TRUE)
## S4 method for signature 'RatioSet, RatioSet'
combineArrays(object1, object2,
                  outType = c("IlluminaHumanMethylation450k",
                               "IlluminaHumanMethylationEPIC",
                               "IlluminaHumanMethylation27k"),
                  verbose = TRUE)
## S4 method for signature 'GenomicMethylSet, GenomicMethylSet'
combineArrays(object1, object2,
                  outType = c("IlluminaHumanMethylation450k",
                               "IlluminaHumanMethylationEPIC",
                               "IlluminaHumanMethylation27k"),
                  verbose = TRUE)
## S4 method for signature 'GenomicRatioSet, GenomicRatioSet'
combineArrays(object1, object2,
                  outType = c("IlluminaHumanMethylation450k",
                               "IlluminaHumanMethylationEPIC",
                               "IlluminaHumanMethylation27k"),
                  verbose = TRUE)
```

# Arguments

object1 The first object.

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object2 The second object.
----------------------------

outType The array type of the output.
verbose Should the function be verbose?

#### **Details**

FIXME: describe the RCChannelSet combination.

#### Value

The output object has the same class as the two input objects, that is either an RGChannelSet, a MethylSet, a RatioSet, a GemomicMethylSet or a GenomicRatioSet, with the type of the array given by the outType argument.

#### Author(s)

Jean-Philippe Fortin and Kasper D. Hansen.

### **Examples**

```
if(require(minfiData) && require(minfiDataEPIC)) {
  data(RGsetEx.sub)
  data(RGsetEPIC)
  rgSet <- combineArrays(RGsetEPIC, RGsetEx.sub)
  rgSet
}</pre>
```

compartments

Estimates A/B compartments from Illumina methylation arrays

### **Description**

Estimates A/B compartments as revealed by Hi-C by computing the first eigenvector on a binned probe correlation matrix.

# Usage

## **Arguments**

object An object of class (Genomic)MethylSet or (Genomic)RatioSet

resolution An integer specifying the binning resolution what Which subset of probes should be used?

chr The chromosome to be analyzed.

method Method of correlation.

keep Should the correlation matrix be stored or not?

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#### **Details**

This function extracts A/B compartments from Illumina methylation microarrays. Analysis of Hi-C data has shown that the genome can be divided into two compartments (A/B compartments) that are cell-type specific and are associated with open and closed chromatin respectively. The approximately 170,000 open sea probes on the 450k array can be used to estimate these compartments by computing the first eigenvector on a binned correlation matrix. The binning resolution can be specified by resolution, and by default is set to a 100 kb. We do not recommend higher resolutions because of the low-resolution probe design of the 450k array.

#### Value

an object of class GRanges containing the correlation matrix, the compartment eigenvector and the compartment labels (A or B) as metadata.

#### Author(s)

Jean-Philippe Fortin < jfortin@jhsph.edu>, Kasper D. Hansen < kasperdanielhansen@gmail.com>

#### References

JP Fortin and KD Hansen. *Reconstructing A/B compartments as revealed by Hi-C using long-range correlations in epigenetic data*. bioRxiv (2015). doi:10.1101/019000.

# **Examples**

```
if (require(minfiData)) {
   GMset <- mapToGenome(MsetEx)
   ## compartments at 1MB resolution; we recommend 100kb.
   comps <- compartments(GMset, res = 10^6)
}</pre>
```

controlStripPlot

Plot control probe signals.

#### **Description**

Strip plots are produced for each control probe type specified.

### Usage

```
controlStripPlot(rgSet, controls = c("BISULFITE CONVERSION I", "BISULFITE CONVERSION II"), sampNames = <math>NULL, xlim = c(5, 17)
```

### **Arguments**

rgSet An RGChannelSet.

controls A vector of control probe types to plot. sampNames Sample names to be used for labels.

xlim x-axis limits.

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#### **Details**

This function produces the control probe signal plot component of the QC report.

#### Value

No return value. Plots are produced as a side-effect.

#### Author(s)

Martin Aryee <aryee@jhu.edu>.

#### See Also

```
qcReport, mdsPlot, densityPlot, densityBeanPlot
```

### **Examples**

```
if (require(minfiData)) {
  names <- pData(RGsetEx)$Sample_Name
  controlStripPlot(RGsetEx, controls=c("BISULFITE CONVERSION I"), sampNames=names)
}</pre>
```

convertArray

A method for converting a type of methylation arrays into a virtual array of another type.

#### **Description**

A method for converting a type of methylation array into a array of another type. The three generations of Illumina methylation arrays are supported: the 27k, the 450k and the EPIC arrays. Specifically, the 450k array and the EPIC array share many probes in common. For RGChannelSet, this function will convert an EPIC array into a 450k array (or vice-versa) by dropping probes that differ between the two arrays. Because most of the probes on the 27k array have a different chemistry than the 450k and EPIC probes, converting an 27k RGChannelSet into another array is not supported. Each array can be converted into another array at the CpG site level, that is any MethylSet and RatioSet (or GenomicMethylSet and GenomicRatioSet) can be converted to a 27k, 450k or EPIC array. The output array is specified by the outType argument.

# Usage

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```
convertArray(object,
                  outType = c("IlluminaHumanMethylation450k",
                              "IlluminaHumanMethylationEPIC",
                               "IlluminaHumanMethylation27k"),
                  verbose = TRUE)
## S4 method for signature 'RatioSet'
convertArray(object,
                  outType = c("IlluminaHumanMethylation450k",
                               "IlluminaHumanMethylationEPIC",
                              "IlluminaHumanMethylation27k"),
                  verbose = TRUE)
## S4 method for signature 'GenomicMethylSet'
convertArray(object,
                  outType = c("IlluminaHumanMethylation450k",
                               "IlluminaHumanMethylationEPIC",
                               "IlluminaHumanMethylation27k"),
                  verbose = TRUE)
## S4 method for signature 'GenomicRatioSet'
convertArray(object,
                  outType = c("IlluminaHumanMethylation450k",
                               "IlluminaHumanMethylationEPIC",
                              "IlluminaHumanMethylation27k"),
                  verbose = TRUE)
```

### **Arguments**

object The input object.

outType The array type of the output.
verbose Should the function be verbose?

#### **Details**

FIXME: describe the RGChannelSet conversion.

#### Value

The output object has the same class as the input object, that is either an RGChannelSet, a MethylSet, a RatioSet, a GemomicMethylSet or a GenomicRatioSet, with the type of the array given by the outType argument.

#### Author(s)

Jean-Philippe Fortin and Kasper D. Hansen.

### **Examples**

```
if(require(minfiData)) {
  data(RGsetEx.sub)
  rgSet <- convertArray(RGsetEx.sub, outType = "IlluminaHumanMethylationEPIC")</pre>
```

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```
rgSet
}
```

cpgCollapse

Collapse methylation values of adjacent CpGs into a summary value.

### **Description**

This function groups adjacent loci into clusters with a specified maximum gap between CpGs in the cluster, and a specified maximum cluster width. The loci within each cluster are summarized resulting in a single methylation estimate per cluster.

### Usage

### **Arguments**

object An object of class [Genomic]MethylSet or [Genomic]RatioSet.

what Should operation be performed on the M-scale or Beta-scale?

maxGap Maximum gap between CpGs in a cluster

blockMaxGap Maximum block gap

maxClusterWidth

Maximum cluster width

dataSummary Function used to summarize methylation across CpGs in the cluster.

na.rm Should NAs be removed when summarizing? Passed on to the dataSummary

function.

returnBlockInfo

Should the block annotation table be returned in addition to the block table?

islandAnno Which Island annotation should be used. NULL indicates the default. This argu-

ment is only useful if the annotatio object contains more than one island anno-

tation.

verbose Should the function be verbose?

... Passed on to getMethSignal and getCN. Can be used to specify

#### **Details**

This function is used as the first step of block-finding. It groups adjacent loci into clusters with a default maximum gap of 500bp and a maximum cluster width of 1,500bp. The loci within each cluster are then summarized (using the mean by default) resulting in a single methylation estimate per cluster. Cluster estimates from open-sea probes are used in block-finding.

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#### Value

If returnBlockInfo is FALSE: a GenomicRatioSet of collapsed CpG clusters.

If returnBlockInfo is TRUE:

object A GenomicRatioSet of collapsed CpG clusters

blockInfo A cluster annotation data frame

#### Author(s)

Rafael Irizarry

#### See Also

blockFinder

densityBeanPlot

Density bean plots of methylation Beta values.

### **Description**

Density 'bean' plots of methylation Beta values, primarily for QC.

### Usage

```
densityBeanPlot(dat, sampGroups = NULL, sampNames = NULL, main = NULL,
    pal = brewer.pal(8, "Dark2"), numPositions = 10000)
```

### **Arguments**

dat An RGChannelSet, a MethylSet or a matrix. We either use the getBeta func-

tion to get Beta values (for the first two) or we assume the matrix contains Beta

values.

sampGroups Optional sample group labels. See details.

sampNames Optional sample names. See details.

main Plot title.
pal Color palette.

numPositions The density calculation uses numPositions randomly selected CpG positions.

If NULL use all positions.

### **Details**

This function produces the density bean plot component of the QC report. If sampGroups is specified, group-specific colors will be used. For speed reasons the plots are produced using a random subset of CpG positions. The number of positions used is specified by the numPositions option.

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### Value

No return value. Plots are produced as a side-effect.

#### Author(s)

Martin Aryee <aryee@jhu.edu>.

#### References

P Kampstra. *Beanplot: A boxplot alternative for visual comparison of distributions*. Journal of Statistical Software 28, (2008). http://www.jstatsoft.org/v28/c01

#### See Also

```
qcReport, mdsPlot, controlStripPlot, densityPlot
```

### **Examples**

```
if (require(minfiData)) {

names <- pData(RGsetEx)$Sample_Name
groups <- pData(RGsetEx)$Sample_Group
par(mar=c(5,6,4,2))
densityBeanPlot(RGsetEx, sampNames=names, sampGroups=groups)
}</pre>
```

densityPlot

Density plots of methylation Beta values.

### **Description**

Density plots of methylation Beta values, primarily for QC.

#### Usage

```
densityPlot(dat, sampGroups = NULL, main = "", xlab = "Beta",
   pal = brewer.pal(8, "Dark2"), xlim, ylim, add = TRUE, legend = TRUE,
   ...)
```

### Arguments

dat An RGChannelSet, a MethylSet or a matrix. We either use the getBeta func-

tion to get Beta values (for the first two) or we assume the matrix contains Beta

values.

sampGroups Optional sample group labels. See details.

main Plot title.

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xlab	x-axis label.
pal	Color palette.
xlim	x-axis limits.
ylim	y-axis limits.
add	Start a new plot?
legend	Plot legend.
	Additional options to be passed to the plot command.

# **Details**

This function produces the density plot component of the QC report. If sampGroups is specified, group-specific colors will be used.

#### Value

No return value. Plots are produced as a side-effect.

#### Author(s)

Martin Aryee <aryee@jhu.edu>.

#### See Also

```
qcReport, mdsPlot, controlStripPlot, densityBeanPlot
```

# **Examples**

```
if (require(minfiData)) {
  groups <- pData(RGsetEx)$Sample_Group
  densityPlot(RGsetEx, sampGroups=groups)
}</pre>
```

 ${\tt detectionP}$ 

Detection p-values for all probed genomic positions.

# **Description**

This function identifies failed positions defined as both the methylated and unmethylated channel reporting background signal levels.

## Usage

```
detectionP(rgSet, type = "m+u")
```

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### **Arguments**

rgSet An RGChannelSet.

type How to calculate p-values. Only m+u is currently implemented (See details).

#### **Details**

A detection p-value is returned for every genomic position in every sample. Small p-values indicate a good position. Positions with non-significant p-values (typically >0.01) should not be trusted.

The m+u method compares the total DNA signal (Methylated + Unmethylated) for each position to the background signal level. The background is estimated using negative control positions, assuming a normal distribution. Calculations are performed on the original (non-log) scale.

This function is different from the detection routine in Genome Studio.

#### Value

A matrix with detection p-values.

#### Author(s)

Martin Aryee <aryee@jhu.edu>.

### **Examples**

```
if (require(minfiData)) {
  detP <- detectionP(RGsetEx.sub)
  failed <- detP>0.01
  colMeans(failed) # Fraction of failed positions per sample
  sum(rowMeans(failed)>0.5) # How many positions failed in >50% of samples?
}
```

dmpFinder

Find differentially methylated positions

### **Description**

Identify CpGs where methylation is associated with a continuous or categorical phenotype.

# Usage

```
dmpFinder(dat, pheno, type = c("categorical", "continuous"),
    qCutoff = 1, shrinkVar = FALSE)
```

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### **Arguments**

dat A MethylSet or a matrix.

pheno The phenotype to be tested for association with methylation.

type Is the phenotype 'continuous' or 'categorical'?

qCutoff DMPs with an FDR q-value greater than this will not be returned.

shrinkVar Should variance shrinkage be used? See details.

#### **Details**

This function tests each genomic position for association between methylation and a phenotype. Continuous phenotypes are tested with linear regression, while an F-test is used for categorical phenotypes.

Variance shrinkage (shrinkVar=TRUE) is recommended when sample sizes are small (<10). The sample variances are squeezed by computing empirical Bayes posterior means using the **limma** package.

### Value

A table with one row per CpG.

# Author(s)

Martin Aryee <aryee@jhu.edu>.

#### See Also

squeezeVar and the limma package in general.

### **Examples**

```
if (require(minfiData)) {
  grp <- pData(MsetEx)$Sample_Group
  MsetExSmall <- MsetEx[1:1e4,] # To speed up the example
  M <- getM(MsetExSmall, type = "beta", betaThreshold = 0.001)
  dmp <- dmpFinder(M, pheno=grp, type="categorical")
  sum(dmp$qval < 0.05, na.rm=TRUE)
  head(dmp)</pre>
```

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estimateCellCounts

Cell Proportion Estimation

### **Description**

Estimates the relative proportion of pure cell types within a sample. For example, given peripheral blood samples, this function will return the relative proportions of lymphocytes, monocytes, B-cells, and neutrophils.

#### **Usage**

```
estimateCellCounts(rgSet, compositeCellType = "Blood",
                   processMethod = "auto", probeSelect = "auto",
                   cellTypes = c("CD8T","CD4T", "NK","Bcell","Mono","Gran"),
                   referencePlatform = c("IlluminaHumanMethylation450k",
                                         "IlluminaHumanMethylationEPIC",
                                         "IlluminaHumanMethylation27k"),
                   returnAll = FALSE, meanPlot = FALSE, verbose = TRUE, ...)
```

#### **Arguments**

rgSet

The input RGChannelSet for the procedure.

compositeCellType

Which composite cell type is being deconvoluted. Should be one of "Blood", "CordBlood", or "DLPFC". See details.

processMethod

How should the user and reference data be processed together? Default input "auto" will use preprocessQuantile for Blood and DLPFC and preprocessNoob otherwise, in line with the existing literature. Set it to the name of a preprocessing function as a character if you want to override it, like "preprocessFunnorm".

probeSelect

How should probes be selected to distinguish cell types? Options include "both", which selects an equal number (50) of probes (with F-stat p-value < 1E-8) with the greatest magnitude of effect from the hyper- and hypo-methylated sides, and "any", which selects the 100 probes (with F-stat p-value < 1E-8) with the greatest magnitude of difference regardless of direction of effect. Default input "auto" will use "any" for cord blood and "both" otherwise, in line with previous versions of this function and/or our recommendations. Please see the references

for more details.

cellTypes

Which cell types, from the reference object, should be we use for the deconvolution? See details.

referencePlatform

The platform for the reference dataset; if the input rgSet belongs to another platform, it will be converted using convertArray.

returnAll

Should the composition table and the normalized user supplied data be return?

verbose

Should the function be verbose?

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meanPlot Whether to plots the average DNA methylation across the cell-type discrimating

probes within the mixed and sorted samples.

... Passed to preprocessQuantile.

#### **Details**

This is an implementation of the Houseman et al (2012) regression calibration approachalgorithm to the Illumina 450k microarray for deconvoluting heterogeneous tissue sources like blood. For example, this function will take an RGChannelSet from a DNA methylation (DNAm) study of blood, and return the relative proportions of CD4+ and CD8+ T-cells, natural killer cells, monocytes, granulocytes, and b-cells in each sample.

The function currently supports cell composition estimation for blood, cord blood, and the frontal cortex, through compositeCellType values of "Blood", "CordBlood", and "DLPFC", respectively. Packages containing the appropriate reference data should be installed before running the function for the first time ("FlowSorted.Blood.450k", "FlowSorted.DLPFC.450k", "FlowSorted.CordBlood.450k"). Each tissue supports the estimation of different cell types, delimited via the cellTypes argument. For blood, these are "Bcell", "CD4T", "CD8T", "Eos", "Gran", "Mono", "Neu", and "NK" (though the default value for cellTypes is often sufficient). For cord blood, these are "Bcell", "CD4T", "CD8T", "Gran", "Mono", "Neu", and "nRBC". For frontal cortex, these are "NeuN\_neg" and "NeuN\_pos". See documentation of individual reference packages for more details.

The meanPlot should be used to check for large batch effects in the data, reducing the confidence placed in the composition estimates. This plot depicts the average DNA methylation across the cell-type discrimating probes in both the provided and sorted data. The means from the provided heterogeneous samples should be within the range of the sorted samples. If the sample means fall outside the range of the sorted means, the cell type estimates will inflated to the closest cell type. Note that we quantile normalize the sorted data with the provided data to reduce these batch effects.

#### Value

Matrix of composition estimates across all samples and cell types.

If returnAll=TRUE a list of a count matrix (see previous paragraph), a composition table and the normalized user data in form of a GenomicMethylSet.

### Author(s)

Andrew E. Jaffe, Shan V. Andrews, E. Andres Houseman

#### References

EA Houseman, WP Accomando, DC Koestler, BC Christensen, CJ Marsit, HH Nelson, JK Wiencke and KT Kelsey. *DNA methylation arrays as surrogate measures of cell mixture distribution*. BMC bioinformatics (2012) 13:86. doi:10.1186/1471-2105-13-86.

AE Jaffe and RA Irizarry. Accounting for cellular heterogeneity is critical in epigenome-wide association studies. Genome Biology (2014) 15:R31. doi:10.1186/gb-2014-15-2-r31.

KM Bakulski, JI Feinberg, SV Andrews, J Yang, S Brown, S McKenney, F Witter, J Walston, AP Feinberg, and MD Fallin. *DNA methylation of cord blood cell types: Applications for mixed cell birth studies.* Epigenetics (2016) 11:5. doi:10.1080/15592294.2016.1161875.

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#### See Also

preprocessQuantile and convertArray.

### **Examples**

```
## Not run:
if(require(FlowSorted.Blood.450k)) {
  wh.WBC <- which(FlowSorted.Blood.450k$CellType == "WBC")
  wh.PBMC <- which(FlowSorted.Blood.450k$CellType == "PBMC")
  RGset <- FlowSorted.Blood.450k[, c(wh.WBC, wh.PBMC)]
  ## The following line is purely to work around an issue with repeated
  ## sampleNames and Biobase::combine()
  sampleNames(RGset) <- paste(RGset$CellType,
        c(seq(along = wh.WBC), seq(along = wh.PBMC)), sep = "_")
  counts <- estimateCellCounts(RGset, meanPlot = FALSE)
  round(counts, 2)
}
## End(Not run)</pre>
```

fixMethOutliers

Fix methylation outliers

#### **Description**

Methylation outliers (loci with very extreme values of the Meth or Unmeth channel) are identified and fixed (see details).

#### Usage

```
fixMethOutliers(object, K = -3, verbose = FALSE)
```

#### Arguments

object An object of class [Genomic] MethylSet.

K The number of standard deviations away from the median when defining the

outlier cutoff, see details.

verbose Should the function be verbose?

### **Details**

This function fixes outlying methylation calls in the Meth channel and Unmeth channel separately.

Unlike other types of arrays, all loci on a methylation array ought to measure something (apart from loci on the Y chromosome in a female sample). An outlier is a loci with a very low value in one of the two methylation channels. Typically, relatively few loci ought to be outliers.

An outlier is defined in a sample and methylation channel specific way. First the (sample, methylation channel) values are log2(x+0.5) transformed and then the median and mad of these values are computed. An outlier is then defined to be any value less than the median plus K times the mad, and these outlier values are thresholded at the cutoff (on the original scale).

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### Value

An object of the same class as object where outlier values in the methylation channels have been thresholded.

### Author(s)

Rafael A. Irizarry and Kasper D. Hansen

#### See Also

```
minfiQC
```

# **Examples**

```
if(require(minfiData)) {
  MsetEx <- fixMethOutliers(MsetEx)
}</pre>
```

gaphunter

Find gap signals in 450k data

# Description

This function finds probes in the Illumina 450k Array for which calculated beta values cluster into distinct groups separated by a defined threshold. It identifies, for these 'gaps signals' the number of groups, the size of these groups, and the samples in each group.

### Usage

# Arguments

object	An object of class (Genomic)RatioSet, (Genomic)MethylSet, or matrix. If one of the first two, codegetBeta is used to calculate beta values. If a matrix, must be one of beta values.
threshold	The difference in consecutive, ordered beta values that defines the presence of a gap signal. Defaults to 5 percent.
keepOutliers	Should outlier-driven gap signals be kept in the results? Defaults to FALSE
outCutoff	Value used to identify gap signals driven by outliers. Defined as the percentage of the total sample size; the sum of samples in all groups except the largest must exceed this number of samples in order for the probe to still be considered a gap signal. Defaults to 1 percent.
verbose	logical value. If TRUE, it writes some messages indicating progress. If FALSE nothing should be printed.

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#### **Details**

The function can calculate a beta matrix or utilize a user-supplied matrix of beta values.

The function will idenfity probes with a gap in a beta signal greater than or equal to the defined threshold. These probes constitue an additional, dataset-specific subset of probes that merit special consideration due to their tendency to be driven by an underlying SNP or other genetic variant. In this manner, these probes can serve as surrogates for underlying genetic signal locally and/or in a broader (i.e. haplotype) context. Please see our upcoming manuscript for a detailed description of the utility of these probes.

Outlier-driven gap signals are those in which the sum of the smaller group(s) does not exceed a certain percentage of the sample size, defined by the argument outCutoff.

#### Value

A list with three values,

proberesults A data frame listing, for each identified gap signal, the number of groups and

the size of each group.

sampleresults a matrix of dimensions probes (rows) by samples (columns). Individuals are

assigned numbers based onthe groups into which they cluster. Lower number groups indicate lower mean methylation values for the group. For example, individuals coded as '1' will have a lower mean methylation value than those

individuals coded as '2'.

algorithm A list detailing the arguments supplied to the function.

### Author(s)

Shan V. Andrews <sandre17@jhu.edu>.

### References

SV Andrews, C Ladd-Acosta, AP Feinberg, KD Hansen, MD Fallin. '*Gap hunting' to characterize clustered probe signals in Illumina methylation array data*. Epigenetics & Chromatin (2016) 9:56. doi:10.1186/s13072-016-0107-z.

#### **Examples**

```
if(require(minfiData)) {
   gapres <- gaphunter(MsetEx.sub, threshold=0.3, keepOutliers=TRUE)
   #Note: the threshold argument is increased from the default value in this small example
   #dataset with 6 people to avoid the reporting of a large amount of probes as gap signals.
   #In a typical EWAS setting with hundreds of samples, the default arguments should be
   #sufficient.
}</pre>
```

GenomicMethylSet-class

GenomicMethylSet instances

## **Description**

This class holds preprocessed data for Illumina methylation microarrays, mapped to a genomic location.

### Usage

```
## Constructor
GenomicMethylSet(gr = GRanges(), Meth = new("matrix"),
                 Unmeth = new("matrix"), annotation = "",
                 preprocessMethod = "", ...)
## Data extraction / Accessors
## S4 method for signature 'GenomicMethylSet'
getMeth(object)
## S4 method for signature 'GenomicMethylSet'
getUnmeth(object)
## S4 method for signature 'GenomicMethylSet'
getBeta(object, type = "", offset = 0, betaThreshold = 0)
## S4 method for signature 'GenomicMethylSet'
getM(object, type = "", ...)
## S4 method for signature 'GenomicMethylSet'
getCN(object, ...)
## S4 method for signature 'GenomicMethylSet'
pData(object)
## S4 method for signature 'GenomicMethylSet'
sampleNames(object)
## S4 method for signature 'GenomicMethylSet'
featureNames(object)
## S4 method for signature 'GenomicMethylSet'
annotation(object)
## S4 method for signature 'GenomicMethylSet'
preprocessMethod(object)
## S4 method for signature 'GenomicMethylSet'
mapToGenome(object, ...)
```

#### **Arguments**

object A GenomicMethylSet. gr A GRanges object. Meth A matrix of methylation values (between zero and infinity) with each row being

a methylation loci and each column a sample.

Unmeth See the Meth argument.

annotation An annotation character string.

preprocessMethod

A preprocess method character string.

type How are the values calculated? For getBeta setting type="Illumina" sets

offset=100 as per Genome Studio. For getM setting type="" computes M-values as the logarithm of Meth/Unmeth, otherwise it is computed as the logit of

getBeta(object).

offset Offset in the beta ratio, see detail.

betaThreshold Constrains the beta values to be in the inverval between betaThreshold and 1-

betaThreshold.

... For the constructor, additional arguments to be passed to SummarizedExperiment;

of particular interest are colData and metadata. For getM these values gets

passed onto getBeta. For mapToGenome, this is ignored.

#### **Details**

For a detailed discussion of getBeta and getM see the deatils section of MethylSet.

#### Value

An object of class GenomicMethylSet for the constructor.

#### Constructor

Instances are constructed using the GenomicMethylSet function with the arguments outlined above.

#### Accessors

A number of useful accessors are inherited from RangedSummarizedExperiment.

In the following code, object is a GenomicMethylSet.

getMeth(object), getUnmeth(object) Get the Meth or Unmeth matrix.

getBeta(object) Get Beta, see details.

getM(object) get M-values, see details.

getCN(object) get copy number values which are defined as the sum of the methylation and unmethylation channel.

getManifest(object) get the manifest associated with the object.

sampleNames(object), featureNames(object) Get the sampleNames (colnames) or the feature-Names (rownames).

preprocessMethod(object), annotation(object) Get the preprocess method or annotation character.

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#### **Utilities**

mapToGenome(object) Since object is already mapped to the genome, this method simply returns object unchanged.

combine: Combines two different GenomicMethylSet, eventually using the cbind method for SummarizedExperiment.

#### Author(s)

Kasper Daniel Hansen <khansen@jhsph.edu>

#### See Also

RangedSummarizedExperiment in the **SummarizedExperiment** package for the basic class structure. Objects of this class are typically created by using the function mapToGenome on a MethylSet.

#### **Examples**

```
showClass("GenomicMethylSet")
```

GenomicRatioSet-class GenomicRatioSet instances

### Description

This class holds preprocessed data for Illumina methylation microarrays, mapped to a genomic location.

#### Usage

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```
## S4 method for signature 'GenomicRatioSet'
featureNames(object)
## S4 method for signature 'GenomicRatioSet'
annotation(object)
## S4 method for signature 'GenomicRatioSet'
preprocessMethod(object)
## S4 method for signature 'GenomicRatioSet'
mapToGenome(object, ...)
```

#### **Arguments**

object A GenomicRatioSet. gr A GRanges object.

Beta A matrix of beta values (optional, see details).

M A matrix of M values (optional, see details).

CN A matrix of copy number values.

An annotation character string.

preprocessMethod

A preprocess method character string.

... For the constructor, additional arguments to be passed to SummarizedExperiment;

of particular interest are colData and metadata. For map To Genome, this is ig-

nored.

### Details

This class holds M or Beta values (or both) together with associated genomic coordinates. It is not possible to get Meth or Unmeth values from this object. The intention is to use this kind of object as an analysis end point.

In case one of M or Beta is missing, the other is computed on the fly. For example, M is computed from Beta as the logit (base 2) of the Beta values.

### Value

An object of class GenomicRatioSet for the constructor.

## Constructor

Instances are constructed using the GenomicRatioSet function with the arguments outlined above.

### Accessors

A number of useful accessors are inherited from RangedSummarizedExperiment.

In the following code, object is a GenomicRatioSet.

```
getBeta(object) Get Beta, see details.
getM(object) get M-values, see details.
```

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```
getCN(object) get copy number, see details.
getManifest(object) get the manifest associated with the object.
sampleNames(object), featureNames(object) Get the sampleNames (colnames) or the feature-
Names (rownames).
preprocessMethod(object), annotation(object) Get the preprocess method or annotation character.
```

#### **Utilities**

```
mapToGenome(object) Since object is already mapped to the genome, this method simply returns object unchanged.
```

combine: Combines two different GenomicRatioSet, eventually using the cbind method for SummarizedExperiment.

#### Author(s)

Kasper Daniel Hansen <khansen@jhsph.edu>

#### See Also

 ${\tt RangedSummarizedExperiment}\ in\ the\ {\bf SummarizedExperiment}\ package\ for\ the\ basic\ class\ structure.$ 

### **Examples**

```
showClass("GenomicRatioSet")
```

getAnnotation

Accessing annotation for Illumina methylation objects

#### **Description**

These functions access provided annotation for various Illumina methylation objects.

# Usage

getAnnotation 29

```
getProbeType(object, withColor = FALSE)
getIslandStatus(object, islandAnno = NULL)
```

#### **Arguments**

object A minfi object.

what Which annotation objects should be returned?

lociNames Restrict the return values to these loci.

orderByLocation

Should the return object be ordered according to genomic location.

dropNonMapping Should loci that do not have a genomic location associated with it (by being

marked as unmapped or multi) be dropped from the return object.

mergeManifest Should the manifest be merged into the return object?

snpAnno The snp annotation you want to use; NULL signifies picking the default.

withColor Should the return object have the type I probe color labelled?

snps The type of SNPs used. maf Minor allelle fraction.

islandAnno Like snpAnno, but for islands.

#### **Details**

getAnnotation returns requested annotation as a DataFrame, with each row corresponding to a methylation loci. If object is of class IlluminaHumanAnnotation no specific ordering of the return object is imposed. If, on the other hand, the class of object imposes some natural order on the return object (ie. if the object is of class [Genomic](Methyl|Ratio)Set), this order is kept in the return object. Note that RGChannelSet does not impose a specific ordering on the methylation loci.

getAnnotationObject returns the annotation object, as opposed to the annotation the object contains. This is useful for printing and examining the contents of the object.

getLocations is a convenience function which returns Locations as a GRanges and which furthermore drops unmapped loci. A user should not need to call this function, instead mapToGenome should be used to get genomic coordinates and granges to return those coordinates.

getSnpInfo is a conevnience function which gets a SNP DataFrame containing information on which probes contains SNPs where. addSnpInfo adds this information to the rowRanges or granges of the object. dropLociWithSnps is a convenience function for removing loci with SNPs based on their MAF.

To see which options are available for what, simply print the annotation object, possibly using getAnnotationObject.

### Value

For getAnnotation, a DataFrame with the requested information.

For getAnnotationObject, a IlluminaMethylationAnnotation object.

For getLocations, a GRanges with the locations.

For getProbeType and getIslandStatus, a character vector with the requested information.

For getSnpInfo, a DataFrame with the requested information. For addSnpInfo, an object of the same class as object but with the SNP information added to the metadata columns of the granges of the object.

For dropLociWithSnps an object of the same kind as the input, possibly with fewer loci.

### Author(s)

Kasper Daniel Hansen<a href="mailto:khansen@jhsph.edu">khansen@jhsph.edu</a>

#### See Also

IlluminaMethylationAnnotation for the basic class, mapToGenome for a better alternative (for users) to getLocations.

### **Examples**

```
if(require(minfiData)) {
  table(getIslandStatus(MsetEx))
  getAnnotation(MsetEx, what = "Manifest")
}
```

getGenomicRatioSetFromGEO

Reading Illumina methylation array data from GEO.

### **Description**

Reading Illumina methylation array data from GEO.

### Usage

```
\label{eq:getGenomicRatioSetFromGEO(GSE = NULL, path = NULL, array = "IlluminaHumanMethylation450k", \\ annotation = .default.450k.annotation, what = c("Beta", "M"), \\ mergeManifest = FALSE, i = 1)
```

### **Arguments**

GSE The GSE ID of the dataset to be downloaded from GEO.

path If data already downloaded, the path with soft files. Either GSE or path are

required.

array Array name.

annotation The feature annotation to be used. This includes the location of features thus

depends on genome build.

what Are Beta or M values being downloaded.

mergeManifest Should the Manifest be merged to the final object.

i If the GEO download results in more than one dataset, it pickes entry i.

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#### **Details**

This function downloads data from GEO using getGEO from the **GEOquery** package. It then returns a GenomicRatioSet object. Note that the rs probes (used for genotyping) are dropped.

#### Value

A GenomicRatioSet object.

### Author(s)

Tim Triche Jr. and Rafael A. Irizarry<rafa@jimmy.harvard.edu>.

#### See Also

If the data is already in memor you can use makeGenomicRatioSetFromMatrix

### **Examples**

```
## Not run:
mset=getGenomicRatioSetFromGEO("GSE42752")
## End(Not run)
```

getMethSignal

Various utilities

### **Description**

Utility functions operating on objects from the minfi package.

### **Usage**

```
getMethSignal(object, what = c("Beta", "M"), ...)
```

# Arguments

object An object from the minfi package supporting either getBeta or getM.

what Which signal is returned.

... Passed to the method described by argument what.

### Value

A matrix.

### Author(s)

Kasper Daniel Hansen <khansen@jhsph.edu>.

getQC

### **Examples**

```
if(require(minfiData)) {
  head(getMethSignal(MsetEx, what = "Beta"))
}
```

getQC

Estimate sample-specific quality control (QC) for methylation data

#### **Description**

Estimate sample-specific quality control (QC) for methylation data.

# Usage

```
getQC(object)
addQC(object, qc)
plotQC(qc, badSampleCutoff = 10.5)
```

### **Arguments**

object An object of class [Genomic]MethylSet.
qc An object as produced by getQC.
badSampleCutoff

The cutoff for identifying a bad sample.

### Value

For getQC, a DataFrame with two columns: mMed and uMed which are the chipwide medians of the Meth and Unmeth channels.

For addQC, essentially object supplied to the function, but with two new columns added to the pheno data slot: uMed and mMed.

# Author(s)

Rafael A. Irizarry and Kasper D. Hansen

### See Also

minfiQC for an all-in-one function.

# **Examples**

```
if(require(minfiData)){
  qc <- getQC(MsetEx)
  MsetEx <- addQC(MsetEx, qc = qc)
  ## plotQC(qc)
}</pre>
```

getSex 33

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Estimating sample sex based on methylation data

### Description

Estimates samples sex based on methylation data.

### Usage

```
getSex(object = NULL, cutoff = -2)
addSex(object, sex = NULL)
plotSex(object, id = NULL)
```

# Arguments

object	An object of class [Genomic]MethylSet.
cutoff	What should the difference in log2 copynumber be between males and females.
sex	An optional character vector of sex (with values M and F).
id	Text used as plotting symbols in the plotSex function. Used for sample identification on the plot.

#### **Details**

Estimation of sex is based on the median values of measurements on the X and Y chromosomes respectively. If yMed - xMed is less than cutoff we predict a female, otherwise male.

#### Value

For getSex, a DataFrame with columns predictedSex (a character with values M and F), xMed and yMed, which are the chip-wide medians of measurements on the two sex chromosomes.

For addSex, an object of the same type as object but with the output of getSex(object) added to the pheno data.

For plotSex, a plot of xMed vs. yMed, which are the chip-wide medians of measurements on the two sex chromosomes, coloured by predictedSex.

### Author(s)

Rafael A. Irizarry, Kasper D. Hansen, Peter F. Hickey

## **Examples**

```
if(require(minfiData)) {
   GMsetEx <- mapToGenome(MsetEx)
   estSex <- getSex(GMsetEx)
   GMsetEx <- addSex(GMsetEx, sex = estSex)
}</pre>
```

```
Illumina Methylation Annotation-class \\ Class~Illumina Methylation Annotation
```

#### **Description**

This is a class for representing annotation associated with an Illumina methylation microarray. Annotation is transient in the sense that it may change over time, wheres the information stored in the IlluminaMethylationManifest class only depends on the array design.

### Usage

### **Arguments**

object An object of class IlluminaMethylationAnnotation.

annotation An annotation character.

defaults A vector of default choices for getAnnotation(what = "everything").

objectNames a character with object names used in the package.

packageName The name of the package this object will be contained in.

### Value

An object of class IlluminaMethylationAnnotation.

### Utilitues

```
In the following code, object is a IlluminaMethylationAnnotation.

getManifest(object) Get the manifest object associated with the array.
```

#### Author(s)

Kasper Daniel Hansen <khansen@jhsph.edu>.

#### See Also

```
Illumina Methylation Manifest\\
```

```
Illumin {\tt aMethylation Manifest-class} \\ Class \ "Illumin {\tt aMethylation Manifest"}
```

# Description

This is a class for representing an Illumina methylation microarray design, ie. the physical location and the probe sequences. This information should be independent of genome build and annotation.

# Usage

```
## Constructor
IlluminaMethylationManifest(TypeI = DataFrame(),
                            TypeII = DataFrame(),
                            TypeControl = DataFrame(),
                            TypeSnpI = DataFrame(),
                            TypeSnpII = DataFrame(),
                            annotation = "")
## Data extraction
## S4 method for signature 'IlluminaMethylationManifest'
getManifest(object)
## S4 method for signature 'character'
getManifest(object)
getProbeInfo(object, type = c("I", "II", "Control",
                              "I-Green", "I-Red", "SnpI", "SnpII"))
getManifestInfo(object, type = c("nLoci", "locusNames"))
getControlAddress(object, controlType = c("NORM_A", "NORM_C",
                                           "NORM_G", "NORM_T"),
                  asList = FALSE)
```

### **Arguments**

object	Either an object of class IlluminaMethylationManifest or class character for getManifest. For getProbeInfo, getManifestInfo and getControlAddress an object of either class RGChannelSet, IlluminaMethylationManifest.
TypeI	A DataFrame of type I probes.
TypeII	A DataFrame of type II probes.
TypeControl	A DataFrame of control probes.
TypeSnpI	A DataFrame of SNP type I probes.
TypeSnpII	A DataFrame of SNP type II probes.
annotation	An annotation character.

type A single character describing what kind of information should be returned.

For getProbeInfo it represents the following subtypes of probes on the array: Type I, Type II, Controls as well as Type I (methylation measured in the Green channel) and Type II (methylation measured in the Red channel). For getManifestInfo it represents either the number of methylation loci (approx.

number of CpGs) on the array or the locus names.

controlType A character vector of control types.

asList If TRUE the return object is a list with one component for each controlType.

#### Value

An object of class IlluminaMethylationManifest for the constructor.

#### **Details**

The data slot contains the following objects: TypeI, TypeII and TypeControl which are all of class data.frame, describing the array design.

Methylation loci of type I are measured using two different probes, in either the red or the green channel. The columns AddressA, Address describes the physical location of the two probes on the array (with ProbeSeqA, ProbeSeqB giving the probe sequences), and the column Color describes which color channel is used.

Methylation loci of type II are measured using a single probe, but with two different color channels. The methylation signal is always measured in the green channel.

#### **Utilities**

In the following code, object is a IlluminaMethylationManifest.

getManifest(object) Get the manifest object.

getProbeInfo(object) Returns a DataFrame giving the type I, type II or control probes. It is also possible to get the type I probes measured in the Green or Red channel. This function ensures that the return object only contains probes which are part of the input object. In case of a RGChannelSet and type I probes, both addresses needs to be in the object.

getManifestInfo(object) Get some information about the manifest object (the chip design). getControlAddress(object) Get the control addresses for control probes of a certain type. getControlTypes(object) Returns the types and the numbers of control probes of each type.

#### Author(s)

Kasper Daniel Hansen <khansen@jhsph.edu>.

#### See Also

IlluminaMethylationAnnotation for annotation information for the array (information depending on a specific genome build).

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# **Examples**

```
if(require(IlluminaHumanMethylation450kmanifest)) {
    show(IlluminaHumanMethylation450kmanifest)
    head(getProbeInfo(IlluminaHumanMethylation450kmanifest, type = "I"))
    head(IlluminaHumanMethylation450kmanifest@data$TypeI)
    head(IlluminaHumanMethylation450kmanifest@data$TypeII)
    head(IlluminaHumanMethylation450kmanifest@data$TypeControl)
}
```

logit2

logit in base 2.

# **Description**

Utility functions for computing logit and inverse logit in base 2.

# Usage

```
logit2(x)
ilogit2(x)
```

# **Arguments**

Х

A numeric vector.

# Value

A numeric vector.

# Author(s)

Kasper Daniel Hansen <khansen@jhsph.edu>.

```
logit2(c(0.25, 0.5, 0.75))
```

makeGenomicRatioSetFromMatrix

Make a GenomicRatioSet from a matrix

# **Description**

Make a GenomicRatioSet from a matrix.

# Usage

# **Arguments**

mat The matrix that will be converted.

rownames The feature IDs associated with the rows of mat that will be used to match to the

IlluminaHumanMethylation450k feature IDs.

pData A DataFrame or data. frame describing the samples represented by the columns

of mat. If the rownames of the pData don't match the colnames of mat these colnames will be changed. If pData is not supplied, a minimal DataFrame is

created.

array Array name.

annotation The feature annotation to be used. This includes the location of features thus

depends on genome build.

mergeManifest Should the Manifest be merged to the final object.

what Are Beta or M values being downloaded.

### **Details**

Many 450K data is provided as csv files. This function permits you to convert a matrix of values into the class that is used by functions such as bumphunter and blockFinder. The rownames of mat are used to match the 450K array features. Alternatively the rownames can be supplied directly through rownames.

### Value

A GenomicRatioSet object.

### Author(s)

Rafael A. Irizarry<rafa@jimmy.harvard.edu>.

### See Also

getGenomicRatioSetFromGEO is similar but reads data from GEO.

### **Examples**

```
mat <- matrix(10,5,2)
rownames(mat) <- c( "cg13869341", "cg14008030","cg12045430", "cg20826792","cg00381604")
grset <- makeGenomicRatioSetFromMatrix(mat)</pre>
```

mapToGenome-methods

Mapping methylation data to the genome

# Description

Mapping Ilumina methylation array data to the genome using an annotation package. Depending on the genome, not all methylation loci may have a genomic position.

# Usage

```
## S4 method for signature 'MethylSet'
mapToGenome(object, mergeManifest = FALSE)
## S4 method for signature 'MethylSet'
mapToGenome(object, mergeManifest = FALSE)
## S4 method for signature 'RGChannelSet'
mapToGenome(object, ...)
```

### **Arguments**

object Either a MethylSet, a RGChannelSet or a RatioSet.

mergeManifest Should the information in the associated manifest package be merged into the

location GRanges?

... Passed to the method for MethylSet.

### **Details**

FIXME: details on the MethylSet method.

The RGChannelSet method of this function is a convenience function: the RGChannelSet is first transformed into a MethylSet using preprocessRaw. The resulting MethylSet is then mapped directly to the genome.

This function silently drops loci which cannot be mapped to a genomic position, based on the associated annotation package.

### Value

An object of class GenomicMethylSet or GenomicRatioSet.

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### Author(s)

Kasper Daniel Hansen <khansen@jhsph.edu>

### See Also

GenomicMethylSet for the output object and MethylSet for the input object. Also, getLocations obtains the genomic locations for a given object.

### **Examples**

```
if (require(minfiData)) {
    ## MsetEx.sub is a small subset of MsetEx;
    ## only used for computational speed.
    GMsetEx.sub <- mapToGenome(MsetEx.sub)
}</pre>
```

mdsPlot

Multi-dimensional scaling plots giving an overview of similarities and differences between samples.

# **Description**

Multi-dimensional scaling (MDS) plots showing a 2-d projection of distances between samples.

# Usage

```
mdsPlot(dat, numPositions = 1000, sampNames = NULL, sampGroups = NULL, xlim, ylim,
    pch = 1, pal = brewer.pal(8, "Dark2"), legendPos = "bottomleft",
    legendNCol, main = NULL)
```

# **Arguments**

dat An RGChannelSet, a MethylSet or a matrix. We either use the getBeta func-

tion to get Beta values (for the first two) or we assume the matrix contains Beta

values.

numPositions Use the numPositions genomic positions with the most methylation variability

when calculating distance between samples.

sampNames Optional sample names. See details.

sampGroups Optional sample group labels. See details.

xlim x-axis limits. ylim y-axis limits.

pch Point type. See par for details.

pal Color palette.

legendPos The legend position. See legend for details.

legendNCol The number of columns in the legend. See legend for details.

main Plot title.

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### **Details**

Euclidean distance is calculated between samples using the numPositions most variable CpG positions. These distances are then projected into a 2-d plane using classical multidimensional scaling transformation.

### Value

No return value. Plots are produced as a side-effect.

# Author(s)

Martin Aryee <aryee@jhu.edu>.

### References

I Borg, P Groenen. *Modern Multidimensional Scaling: theory and applications (2nd ed.)* New York: Springer-Verlag (2005) pp. 207-212. ISBN 0387948457.

```
http://en.wikipedia.org/wiki/Multidimensional_scaling
```

#### See Also

```
qcReport, controlStripPlot, densityPlot, densityBeanPlot, par, legend
```

### **Examples**

```
if (require(minfiData)) {

names <- pData(MsetEx)$Sample_Name
groups <- pData(MsetEx)$Sample_Group
mdsPlot(MsetEx, sampNames=names, sampGroups=groups)
}</pre>
```

MethylSet-class

MethylSet instances

### **Description**

This class holds preprocessed data for Illumina methylation microarrays.

# Usage

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```
## S4 method for signature 'MethylSet'
getMeth(object)
## S4 method for signature 'MethylSet'
getUnmeth(object)
## S4 method for signature 'MethylSet'
getBeta(object, type = "", offset = 0, betaThreshold = 0)
## S4 method for signature 'MethylSet'
getM(object, type = "", ...)
## S4 method for signature 'MethylSet'
getCN(object, ...)
## S4 method for signature 'MethylSet'
getManifest(object)
## S4 method for signature 'MethylSet'
preprocessMethod(object)
## S4 method for signature 'MethylSet'
annotation(object)
## S4 method for signature 'MethylSet'
pData(object)
## S4 method for signature 'MethylSet'
sampleNames(object)
## S4 method for signature 'MethylSet'
featureNames(object)
## Utilities
dropMethylationLoci(object, dropRS = TRUE, dropCH = TRUE)
```

# Arguments

object A MethylSet.

Meth A matrix of methylation values (between zero and infinity) with each row being

a methylation loci and each column a sample.

Unmeth See the Meth argument.

annotation An annotation string, optional.

preprocessMethod

A character, optional.

type How are the values calculated? For getBeta setting type="Illumina" sets

offset=100 as per Genome Studio. For getM setting type="" computes M-values as the logarithm of Meth/Unmeth, otherwise it is computed as the logit of

getBeta(object).

offset Offset in the beta ratio, see detail.

betaThreshold Constrains the beta values to be in the inverval betwen betaThreshold and 1-

betaThreshold.

dropRS Should SNP probes be dropped?
dropCH Should CH probes be dropped

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For the constructor, additional arguments to be passed to SummarizedExperiment; of particular interest are colData, rowData and metadata. For getM these values gets passed onto getBeta.

### **Details**

This class inherits from eSet. Essentially the class is a representation of a Meth matrix and a Unmeth matrix linked to a pData data frame.

In addition, an annotation and a preprocessMethod slot is present. The annotation slot describes the type of array and also which annotation package to use. The preprocessMethod slot describes the kind of preprocessing that resulted in this dataset.

A MethylSet stores meth and Unmeth. From these it is easy to compute Beta values, defined as

$$\beta = \frac{\text{Meth}}{\text{Meth} + \text{Unmeth} + \text{offset}}$$

The offset is chosen to avoid dividing with small values. Illumina uses a default of 100. M-values (an unfortunate bad name) are defined as

$$M = logit(\beta) = log(Meth/Unmeth)$$

This formula has problems if either Meth or Unmeth is zero. For this reason, we can use betaThreshold to make sure Beta is neither 0 nor 1, before taken the logit. What makes sense for the offset and betaThreshold depends crucially on how the data was preprocessed. Do not expect the default values to be particular good.

### Value

An object of class MethylSet for the constructor.

### Constructor

Instances are constructed using the MethylSet function with the arguments outlined above.

### Accessors

In the following code, object is a MethylSet.

getMeth(object), getUnmeth(object) Get the Meth or the Unmeth matrix

getBeta(object) Get Beta, see details.

getM(object) get M-values, see details.

getCN(object) get copy number values which are defined as the sum of the methylation and unmethylation channel.

getManifest(object) get the manifest associated with the object.

preprocessMethod(object) Get the preprocess method character.

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# Utilities

In the following code, object is a MethylSet.

dropMethylationLoci(**object**) A unifed interface to removing methylation loci. You can drop SNP probes (probes that measure SNPs, not probes containing SNPs) or CH probes (non-CpG methylation).

combine: Combines two different MethylSet, eventually using the combine method for eSet.

# Author(s)

Kasper Daniel Hansen <khansen@jhsph.edu>

### See Also

eSet for the basic class structure. Objects of this class are typically created from an RGChannelSet using preprocessRaw or another preprocessing function.

# **Examples**

```
showClass("MethylSet")
```

minfi-defunct

Defunct functions in package 'minfi'

# Description

These functions are provided now defunct in 'minfi'.

# **Details**

The following functions are now defunct (not working anymore); use the replacement indicated below:

• read.450k: Use read.metharray

• read.450k.sheet: Use read.metharray.sheet

• read.450k.exp: Use read.metharray.exp

# See Also

Defunct.

minfi-deprecated 45

minfi-deprecated Deprecated functions in package 'minfi'
--

# **Description**

These functions are provided for compatibility with older versions of 'minfi' only, and will be defunct at the next release.

### **Details**

No functions are currently deprecated.

The following functions are deprecated and will be made defunct; use the replacement indicated below:

```
read.450k: read.metharray
read.450k.sheet: read.metharray.sheet
read.450k.exp: read.metharray.exp
```

minfiQC

easy one-step QC of methylation object

# **Description**

This function combines a number of functions into a simple to use, one step QC step/

# Usage

```
minfiQC(object, fixOutliers = TRUE, verbose = FALSE)
```

### **Arguments**

object An object of class [Genomic]MethylSet.

fixOutliers Should the function fix outlying observations (using fixMethOutliers) before

running QC?

verbose Should the function be verbose?

### **Details**

A number of functions are run sequentially on the object.

First outlier values are thresholded using fixMethOutliers. Then qc is performed using getQC and then sample specific sex is estimated using getSex.

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### Value

A list with two values,

object The object processed by fixMethOutliers and with a column predictedSex

added to the pheno data.

qc A DataFrame with columns from the output of getQC and getSex

### Author(s)

Kasper D. Hansen

### See Also

```
getSex, getQC, fixMethOutliers
```

# **Examples**

```
if(require(minfiData)) {
  out <- minfiQC(MsetEx)
  ## plotQC(out$qc)
  ## plotSex(out$sex)
}</pre>
```

plotBetasByType

Plot the overall distribution of beta values and the distributions of the Infinium I and II probe types.

### **Description**

Plot the overall density distribution of beta values and the density distributions of the Infinium I and II probe types.

# Usage

# **Arguments**

data A MethylSet or a matrix or a vector. We either use the getBeta function to

get Beta values (in the first case) or we assume the matrix or vector contains

Beta values.

probeTypes If data is a MethylSet this argument is not needed. Otherwise, a data.frame

with a column 'Name' containing probe IDs and a column 'Type' containing

their corresponding assay design type.

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legendPos The x and y co-ordinates to be used to position the legend. They can be specified

by keyword or in any way which is accepted by xy.coords. See legend for

details.

colors Colors to be used for the different beta value density distributions. Must be a

vector of length 3.

main Plot title.

1wd The line width to be used for the different beta value density distributions.

cex.legend The character expansion factor for the legend text.

### **Details**

The density distribution of the beta values for a single sample is plotted. The density distributions of the Infinium I and II probes are then plotted individually, showing how they contribute to the overall distribution. This is useful for visualising how using preprocessSWAN affects the data.

### Value

No return value. Plot is produced as a side-effect.

### Author(s)

Jovana Maksimovic <jovana.maksimovic@mcri.edu.au>.

### See Also

densityPlot, densityBeanPlot, par, legend

# **Examples**

```
## Not run:
if (require(minfiData)) {
   Mset.swan <- preprocessSWAN(RGsetEx, MsetEx)
   par(mfrow=c(1,2))
   plotBetasByType(MsetEx[,1], main="Raw")
   plotBetasByType(Mset.swan[,1], main="SWAN")
}
## End(Not run)</pre>
```

plotCpg

Plot methylation values at an single genomic position

# **Description**

Plot single-position (single CpG) methylation values as a function of a categorical or continuous phenotype

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### Usage

```
plotCpg(dat, cpg, pheno, type = c("categorical", "continuous"),
    measure = c("beta", "M"), ylim = NULL, ylab = NULL, xlab = "",
    fitLine = TRUE, mainPrefix = NULL, mainSuffix = NULL)
```

### **Arguments**

dat An RGChannelSet, a MethylSet or a matrix. We either use the getBeta (or

 $\texttt{getM} \ for \ \texttt{measure="M"}) \ function \ to \ \texttt{get} \ Beta \ values \ (or \ M-values) \ (for \ the \ first$ 

two) or we assume the matrix contains Beta values (or M-values).

cpg A character vector of the CpG position identifiers to be plotted.

pheno A vector of phenotype values.

type Is the phenotype categorical or continuous?

measure Should Beta values or log-ratios (M) be plotted?

ylim y-axis limits. ylab y-axis label. xlab x-axis label.

fitLine Fit a least-squares best fit line when using a continuous phenotype.

mainPrefix Text to prepend to the CpG name in the plot main title.

Text to append to the CpG name in the plot main title.

### **Details**

This function plots methylation values (Betas or log-ratios) at individual CpG loci as a function of a phenotype.

# Value

No return value. Plots are produced as a side-effect.

# Author(s)

Martin Aryee <aryee@jhu.edu>.

```
if (require(minfiData)) {
grp <- pData(MsetEx)$Sample_Group
cpgs <- c("cg00050873", "cg00212031", "cg26684946", "cg00128718")
par(mfrow=c(2,2))
plotCpg(MsetEx, cpg=cpgs, pheno=grp, type="categorical")
}</pre>
```

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preprocessFunnorm	Functional normalization for Illumina 450k arrays

### **Description**

Functional normalization (FunNorm) is a between-array normalization method for the Illumina Infinium HumanMethylation450 platform. It removes unwanted variation by regressing out variability explained by the control probes present on the array.

# Usage

# Arguments

rgSet	An object of class RGChannelSet.
nPCs	Number of principal components from the control probes PCA
sex	An optional numeric vector containing the sex of the samples.
bgCorr	Should the NOOB background correction be done, prior to functional normalization (see preprocessNoob)
dyeCorr	Should dye normalization be done as part of the NOOB background correction (see preprocessNoob)?
keepCN	Should copy number estimates be kept around? Setting to FALSE will decrease the size of the output object significantly.
ratioConvert	Should we run ratioConvert, ie. should the output be a GenomicRatioSet or should it be kept as a GenomicMethylSet; the latter is for experts.
verbose	Should the function be verbose?

### **Details**

This function implements functional normalization preprocessing for Illumina methylation microarrays. Functional normalization extends the idea of quantile normalization by adjusting for known covariates measuring unwanted variation. For the 450k array, the first k principal components of the internal control probes matrix play the role of the covariates adjusting for technical variation. The number k of principal components can be set by the argument nPCs. By default nPCs is set to 2, and have been shown to perform consistently well across different datasets. This parameter should only be modified by expert users. The normalization procedure is applied to the Meth and Unmeth intensities separately, and to type I and type II signals separately. For the probes on the X and Y chromosomes we normalize males and females separately using the gender information provided in the sex argument. For the Y chromosome, standard quantile normalization is used due to the small number of probes, which results in instability for functional normalization. If sex is unspecified (NULL), a guess is made using by the getSex function using copy number information. Note that this algorithm does not rely on any assumption and therefore can be be applicable for cases where global changes are expected such as in cancer-normal comparisons or tissue differences.

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### Value

an object of class GenomicRatioSet, unless ratioConvert=FALSE in which case an object of class GenomicMethylSet.

### Author(s)

Jean-Philippe Fortin < jfortin@jhsph.edu>, Kasper D. Hansen < khansen@jhsph.edu>.

#### References

JP Fortin, A Labbe, M Lemire, BW Zanke, TJ Hudson, EJ Fertig, CMT Greenwood and KD Hansen. Functional normalization of 450k methylation array data improves replication in large cancer studies. (2014) Genome Biology (2014) 15:503. doi:10.1186/s13059-014-0503-2.

### See Also

RGChannelSet as well as IlluminaMethylationManifest for the basic classes involved in these functions. preprocessRaw and preprocessQuantile are other preprocessing functions. Background correction may be done using preprocessNoob.

### **Examples**

```
if (require(minfiData)) {
    ## RGsetEx.sub is a small subset of RGsetEx;
    ## only used for computational speed.
    Mset.sub.funnorm <- preprocessFunnorm(RGsetEx.sub)
}</pre>
```

preprocessIllumina

Perform preprocessing as Genome Studio.

# **Description**

These functions implements preprocessing for Illumina methylation microarrays as used in Genome Studio, the standard software provided by Illumina.

### Usage

```
preprocessIllumina(rgSet, bg.correct = TRUE, normalize = c("controls", "no"),
    reference = 1)
bgcorrect.illumina(rgSet)
normalize.illumina.control(rgSet, reference = 1)
```

# **Arguments**

rgSet An object of class RGChannelSet.

bg.correct logical, should background correction be performed?
normalize logical, should (control) normalization be performed?
reference for control normalization, which array is the reference?

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### **Details**

We have reverse engineered the preprocessing methods from Genome Studio, based on the documentation.

The current implementation of control normalization is equal to what Genome Studio provides (this statement is based on comparing Genome Studio output to the output of this function), with the following caveat: this kind of normalization requires the selection of a reference array. It is unclear how Genome Studio selects the reference array, but we allow for the manual specification of this parameter.

The current implementation of background correction is roughly equal to Genome Studio. Based on examining the output of 24 arrays, we are able to exactly recreate 18 out of the 24. The remaining 6 arrays had a max discrepancy in the Red and/or Green channel of 1-4 (this is on the unlogged intensity scale, so 4 is very small).

A script for doing this comparison may be found in the scripts directory (although it is of limited use without the data files).

### Value

preprocessIllumina returns a MethylSet, while bgcorrect.illumina and normalize.illumina.control both return a RGChannelSet with corrected color channels.

### Author(s)

Kasper Daniel Hansen <khansen@jhsph.edu>.

### See Also

RGChannelSet and MethylSet as well as IlluminaMethylationManifest for the basic classes involved in these functions. preprocessRaw is another basic preprocessing function.

# **Examples**

```
if (require(minfiData)) {

dat <- preprocessIllumina(RGsetEx, bg.correct=FALSE, normalize="controls")
slot(name="preprocessMethod", dat)[1]
}</pre>
```

preprocessNoob

The Noob/ssNoob preprocessing method for Infinium methylation microarrays.

### **Description**

Noob (normal-exponential out-of-band) is a background correction method with dye-bias normalization for Illumina Infinium methylation arrays.

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### Usage

### **Arguments**

rgSet An object of class RGChannelSet.

offset An offset for the normexp background correction.

dyeCorr Should dye correction be done? verbose Should the function be verbose?

dyeMethod How should dye bias correction be done: use a single sample approach (ss-

Noob), or a reference array?

#### Value

An object of class MethylSet.

### Author(s)

Tim Triche, Jr.

### References

TJ Triche, DJ Weisenberger, D Van Den Berg, PW Laird and KD Siegmund *Low-level processing of Illumina Infinium DNA Methylation BeadArrays*. Nucleic Acids Res (2013) 41, e90. doi:10.1093/nar/gkt090.

# See Also

RGChannelSet as well as IlluminaMethylationManifest for the basic classes involved in these functions. preprocessRaw and preprocessQuantile are other preprocessing functions.

preprocessQuantile 53

preprocessQuantile

Stratified quantile normalization for an Illumina methylation array.

### **Description**

Stratified quantile normalization for Illumina amethylation arrays.

This function implements stratified quantile normalization preprocessing for Illumina methylation microarrays. Probes are stratified by region (CpG island, shore, etc.)

### Usage

# **Arguments**

object An object of class RGChannelSet or [Genomic]MethylSet.

fixOutliers Should low outlier Meth and Unmeth signals be fixed?

removeBadSamples

Should bad samples be removed?

badSampleCutoff

Samples with median Meth and Umneth signals below this cutoff will be la-

belled 'bad'.

quantileNormalize

Should quantile normalization be performed?

stratified Should quantile normalization be performed within genomic region strata (e.g.

CpG island, shore, etc.)?

mergeManifest Should the information in the associated manifest package be merged into the

output object?

sex Gender

verbose Should the function be verbose?

# Details

This function implements stratified quantile normalization preprocessing for Illumina methylation microarrays. If removeBadSamples is TRUE we calculate the median Meth and median Unmeth signal for each sample, and remove those samples where their average falls below badSampleCutoff. The normalization procedure is applied to the Meth and Unmeth intensities separately. The distribution of type I and type II signals is forced to be the same by first quantile normalizing the type II probes across samples and then interpolating a reference distribution to which we normalize the type I probes. Since probe types and probe regions are confounded and we know that DNAm distributions vary across regions we stratify the probes by region before applying this interpolation. For the probes on the X and Y chromosomes we normalize males and females separately using

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the gender information provided in the sex argument. If gender is unspecified (NULL), a guess is made using by the getSex function using copy number information. Background correction is not used, but very small intensities close to zero are thresholded using the fixMethOutlier. Note that this algorithm relies on the assumptions necessary for quantile normalization to be applicable and thus is not recommended for cases where global changes are expected such as in cancer-normal comparisons.

Note that this normalization procedure is essentially similar to one previously presented (Touleimat and Tost, 2012), but has been independently re-implemented due to the present lack of a released, supported version.

### Value

a GenomicRatioSet

### Note

A bug in the function was found to affect the Beta values of type I probes, when stratified=TRUE (default). This is fixed in minfi version 1.19.7 and 1.18.4 and greater.

### Author(s)

Rafael A. Irizarry

### References

N Touleimat and J Tost. Complete pipeline for Infinium Human Methylation 450K BeadChip data processing using subset quantile normalization for accurate DNA methylation estimation. Epigenomics (2012) 4:325-341.

# See Also

getSex, minfiQC, fixMethOutliers for functions used as part of preprocessQuantile.

```
if (require(minfiData)) {
    # NOTE: RGsetEx.sub is a small subset of RGsetEx; only used for computational
    # speed
    GMset.sub.quantile <- preprocessQuantile(RGsetEx.sub)
}
## Not run:
if(require(minfiData)) {
    GMset <- preprocessQuantile(RGsetEx)
}
## End(Not run)</pre>
```

preprocessRaw 55

preprocessRaw

Creation of a MethylSet without normalization

# Description

Converts the Red/Green channel for an Illumina methylation array into methylation signal, without using any normalization.

### Usage

```
preprocessRaw(rgSet)
```

# Arguments

rgSet

An object of class RGChannelSet.

# **Details**

This function takes the Red and the Green channel of an Illumina methylation array, together with its associated manifest object and converts it into a MethylSet containing the methylated and unmethylated signal.

### Value

An object of class MethylSet

# Author(s)

Kasper Daniel Hansen<a href="https://khansen@jhsph.edu">khansen@jhsph.edu</a>>.

# See Also

RGChannelSet and MethylSet as well as IlluminaMethylationManifest.

```
if (require(minfiData)) {
dat <- preprocessRaw(RGsetEx)
slot(name="preprocessMethod", dat)[1]
}</pre>
```

56 preprocessSWAN

preprocessSWAN	Subset-quantile Within Array Normalisation for Illumina Infinium HumanMethylation450 BeadChips

### **Description**

Subset-quantile Within Array Normalisation (SWAN) is a within array normalisation method for the Illumina Infinium HumanMethylation450 platform. It allows Infinium I and II type probes on a single array to be normalized together.

### Usage

```
preprocessSWAN(rgSet, mSet = NULL, verbose = FALSE)
```

### **Arguments**

rgSet An object of class RGChannelSet.

mSet An optional object of class MethylSet. If set to NULL preprocessSwan uses

preprocessRaw on the rgSet argument. In case mSet is supplied, make sure it

is the result of preprocessing the rgSet argument.

verbose Should the function be verbose?

# Details

The SWAN method has two parts. First, an average quantile distribution is created using a subset of probes defined to be biologically similar based on the number of CpGs underlying the probe body. This is achieved by randomly selecting N Infinium I and II probes that have 1, 2 and 3 underlying CpGs, where N is the minimum number of probes in the 6 sets of Infinium I and II probes with 1, 2 or 3 probe body CpGs. If no probes have previously been filtered out e.g. sex chromosome probes, etc. N=11,303. This results in a pool of 3N Infinium I and 3N Infinium II probes. The subset for each probe type is then sorted by increasing intensity. The value of each of the 3N pairs of observations is subsequently assigned to be the mean intensity of the two probe types for that row or "quantile". This is the standard quantile procedure. The intensities of the remaining probes are then separately adjusted for each probe type using linear interpolation between the subset probes.

### Value

an object of class MethylSet

### Note

SWAN uses a random subset of probes to do the between array normalization. In order to achive reproducible results, the seed needs to be set using set.seed.

# Author(s)

Jovana Maksimovic<jovana.maksimovic@mcri.edu.au>

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### References

J Maksimovic, L Gordon and A Oshlack (2012). SWAN: Subset quantile Within-Array Normalization for Illumina Infinium HumanMethylation450 BeadChips. Genome Biology 13, R44.

### See Also

RGChannelSet and MethylSet as well as IlluminaMethylationManifest.

### **Examples**

```
if (require(minfiData)) {
    ## RGsetEx.sub is a small subset of RGsetEx;
    ## only used for computational speed.
    MsetEx.sub.swan <- preprocessSWAN(RGsetEx.sub)
}
## Not run:
if (require(minfiData)) {
    dat <- preprocessRaw(RGsetEx)
    preprocessMethod(dat)
    datSwan <- preprocessSWAN(RGsetEx, mSet = dat)
    datIlmn <- preprocessIllumina(RGsetEx)
    preprocessMethod(datIlmn)
    datIlmnSwan <- preprocessSWAN(RGsetEx, mSet = datIlmn)
}
## End(Not run)</pre>
```

qcReport

QC report for Illumina Infinium Human Methylation 450k arrays

### **Description**

Produces a PDF QC report for Illumina Infinium Human Methylation 450k arrays, useful for identifying failed samples.

# Usage

```
qcReport(rgSet, sampNames = NULL, sampGroups = NULL, pdf = "qcReport.pdf",
    maxSamplesPerPage = 24, controls = c("BISULFITE CONVERSION I",
    "BISULFITE CONVERSION II", "EXTENSION", "HYBRIDIZATION",
    "NON-POLYMORPHIC", "SPECIFICITY I", "SPECIFICITY II", "TARGET REMOVAL"))
```

### **Arguments**

```
rgSet An object of class RGChannelSet.
sampNames Sample names to be used for labels.
sampGroups Sample groups to be used for labels.
```

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pdf Path and name of the PDF output file. maxSamplesPerPage

Maximum number of samples to plot per page in those sections that plot each sample separately.

controls The control probe types to include in the report.

### **Details**

This function produces a QC report as a PDF file. It is a useful first step after reading in a new dataset to get an overview of quality and to flag potentially problematic samples.

### Value

No return value. A PDF is produced as a side-effect.

### Author(s)

Martin Aryee <aryee@jhu.edu>.

### See Also

```
mdsPlot, controlStripPlot, densityPlot, densityBeanPlot
```

### **Examples**

```
if (require(minfiData)) {
names <- pData(RGsetEx)$Sample_Name
groups <- pData(RGsetEx)$Sample_Group

## Not run:
qcReport(RGsetEx, sampNames=names, sampGroups=groups, pdf="qcReport.pdf")

## End(Not run)
}</pre>
```

ratioConvert-methods Converting methylation signals to ratios (Beta or M-values)

# **Description**

Converting methylation data from methylation and unmethylation channels, to ratios (Beta and M-values).

RatioSet-class 59

# Usage

```
## S4 method for signature 'MethylSet'
ratioConvert(object, what = c("beta", "M", "both"), keepCN = TRUE, ...)
## S4 method for signature 'GenomicMethylSet'
ratioConvert(object, what = c("beta", "M", "both"), keepCN = TRUE, ...)
```

### **Arguments**

object Either a MethylSet, or a GenomicRatioSet.

what Which ratios should be computed and stored?

keepCN A logical, should copy number values be computed and stored in the object?

Passed to getBeta, getM methods.

### Value

An object of class RatioSet or GenomicRatioSet.

### Author(s)

Kasper Daniel Hansen <khansen@jhsph.edu>

### See Also

RatioSet or codeGenomicRatioSet for the output object and MethylSet or codeGenomicMethylSet for the input object.

# **Examples**

```
if (require(minfiData)) {
    ## MsetEx.sub is a small subset of MsetEx;
    ## only used for computational speed.
    RsetEx.sub <- ratioConvert(MsetEx.sub, keepCN = TRUE)
}</pre>
```

RatioSet-class

RatioSet instances

# **Description**

This class holds preprocessed data for Illumina methylation microarrays.

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# Usage

```
## Constructor
RatioSet(Beta = NULL, M = NULL, CN = NULL,
        annotation = "", preprocessMethod = "", ...)
## Data extraction / Accessors
## S4 method for signature 'RatioSet'
getBeta(object)
## S4 method for signature 'RatioSet'
getM(object)
## S4 method for signature 'RatioSet'
getCN(object)
## S4 method for signature 'RatioSet'
preprocessMethod(object)
## S4 method for signature 'RatioSet'
annotation(object)
## S4 method for signature 'RatioSet'
pData(object)
## S4 method for signature 'RatioSet'
sampleNames(object)
## S4 method for signature 'RatioSet'
featureNames(object)
```

# Arguments

object	A RatioSet.	
Beta	A matrix of beta values (between zero and one) with each row being a methylation loci and each column a sample.	
М	A matrix of log-ratios (between minus infinity and infinity) with each row being a methylation loci and each column a sample.	
CN	An optional matrix of copy number estimates with each row being a methylation loci and each column a sample.	
annotation	An annotation string, optional.	
preprocessMethod		
	A character, optional.	
	For the constructor, additional arguments to be passed to SummarizedExperiment; of particular interest are colData, rowData and metadata. For getM these values gets passed onto getBeta.	

### **Details**

This class inherits from eSet. Essentially the class is a representation of a Beta matrix and/or a M matrix and optionally a CN (copy number) matrix linked to a pData data frame.

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In addition, an annotation and a preprocessMethod slot is present. The annotation slot describes the type of array and also which annotation package to use. The preprocessMethod slot describes the kind of preprocessing that resulted in this dataset.

For a RatioSet, M-values are defined as logit2 of the Beta-values if the M-values are not present in the object. Similarly, if only M-values are present in the object, Beta-values are ilogit2 of the M-values.

### Value

An object of class RatioSet for the constructor.

#### Constructor

Instances are constructed using the RatioSet function with the arguments outlined above.

### Accessors

```
In the following code, object is a RatioSet.
```

```
getBeta(object), getM(object), CN(object) Get the Beta, M or CN matrix. getManifest(object) get the manifest associated with the object. preprocessMethod(object) Get the preprocess method character.
```

### **Utilities**

In the following code, object is a RatioSet.

combine: Combines two different RatioSet, eventually using the combine method for eSet.

### Author(s)

Kasper Daniel Hansen <khansen@jhsph.edu>

### See Also

eSet for the basic class structure. Objects of this class are typically created from an MethylSet using ratioConvert.

```
showClass("RatioSet")
```

62 read.metharray

ı	read.metharray	Parsing IDAT files from Illumina methylation arrays.

### **Description**

Parsing IDAT files from Illumina methylation arrays.

# Usage

read.metharray(basenames, extended = FALSE, verbose = FALSE, force = FALSE)

# **Arguments**

basenames The basenames or filenames of the IDAT files. By basenames we mean the file-

name without the ending \_Grn.idat or \_Red.idat (such that each sample occur once). By filenames we mean filenames including \_Grn.idat or \_Red.idat

(but only one of the colors)

extended Should a RGChannelSet or a RGChannelSetExtended be returned.

verbose Should the function be verbose?

force Should reading different size IDAT files be forced? See Details.

# **Details**

The type of methylation array is guess by looking at the number of probes in the IDAT files.

We have seen IDAT files from the same array, but with different number of probes in the wild. Specifically this is the case for early access EPIC arrays which have fewer probes than final release EPIC arrays. It is possible to combine IDAT files from the same inferred array, but with different number of probes, into the same RGChannelSet by setting force=TRUE. The output object will have the same number of probes as the smallest array being parsed; effectively removing probes which could have been analyzed.

### Value

An object of class RGChannelSet or RGChannelSetExtended.

### Author(s)

Kasper Daniel Hansen<a href="khansen@jhsph.edu">khansen@jhsph.edu</a>>.

### See Also

read.metharray.exp for a convenience function for reading an experiment, read.metharray.sheet for reading a sample sheet and RGChannelSet for the output class.

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# **Examples**

```
if(require(minfiData)) {
baseDir <- system.file("extdata", package = "minfiData")
RGset1 <- read.metharray(file.path(baseDir, "5723646052", "5723646052_R02C02"))
}</pre>
```

read.metharray.exp

Reads an entire metharray experiment using a sample sheet

### **Description**

Reads an entire methylation array experiment using a sample sheet or (optionally) a target like data.frame.

# Usage

```
read.metharray.exp(base = NULL, targets = NULL, extended = FALSE,
    recursive = FALSE, verbose = FALSE, force = FALSE)
```

### **Arguments**

base The base directory.

targets A targets data. frame, see details

extended Should the output of the function be a "RGChannelSetExtended" (default is

"RGChannelSet").

recursive Should the search be recursive (see details)

verbose Should the function be verbose?

force Should reading different size IDAT files be forced? See the documentation for

read.metharray

### **Details**

If the targets argument is NULL, the function finds all two-color IDAT files in the directory given by base. If recursive is TRUE, the function searches base and all subdirectories. A two-color IDAT files are pair of files with names ending in \_Red.idat or \_Grn.idat.

If the targets argument is not NULL it is assumed it has a columned named Basename, and this is assumed to be pointing to the base name of a two color IDAT file, ie. a name that can be made into a real IDAT file by appending either \_Red.idat or \_Grn.idat.

The type of methylation array is guess by looking at the number of probes in the IDAT files.

### Value

An object of class "RGChannelSet" or "RGChannelSetExtended".

64 read.metharray.sheet

### Author(s)

Kasper Daniel Hansen <khansen@jhsph.edu>.

#### See Also

read.metharray for the workhorse function, read.metharray.sheet for reading a sample sheet and RGChannelSet for the output class.

### **Examples**

```
if(require(minfiData)) {
baseDir <- system.file("extdata", package = "minfiData")
RGset <- read.metharray.exp(file.path(baseDir, "5723646052"))
}</pre>
```

read.metharray.sheet Reading an Illumina methylation sample sheet

**Description** 

Reading an Illumina methylation sample sheet, containing pheno-data information for the samples in an experiment.

### Usage

```
read.metharray.sheet(base, pattern = "csv$", ignore.case = TRUE,
    recursive = TRUE, verbose = TRUE)
```

### **Arguments**

base The base directory from which the search is started.

pattern What pattern is used to identify a sample sheet file, see list.files

ignore.case Should the file search be case sensitive?

recursive Should the file search be recursive, see list.files?

verbose Should the function be verbose?

### **Details**

This function search the directory base (possibly including subdirectories depending on the argument recursive for "sample sheet" files (see below). These files are identified solely on the base of their filename given by the arguments pattern and ignore.case (note the use of a dollarsign to mean end of file name).

In case multiple sheet files are found, they are all read and the return object will contain the concatenation of the files.

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A sample sheet file is essentially a CSV (comma-separated) file containing one line per sample, with a number of columns describing pheno-data or other important information about the sample. The file may contain a header, in which case it is assumed that all lines up to and including a line starting with \[Data\] should be dropped. This is modelled after a sample sheet file Illumina provides. It is also very similar to the targets file made used by the popular limma package (see the extensive package vignette).

An attempt at guessing the file path to the IDAT files represented in the sheet is made. This should be doublechecked and might need to manually changed.

The type of methylation array is guess by looking at the number of probes in the IDAT files.

#### Value

A data.frame containing the columns of all the sample sheets. As described in details, a column named Sentrix\_Position is renamed to Array and Sentrix\_ID is renamed to Slide. In addition the data.frame will contain a column named Basename.

### Author(s)

Kasper Daniel Hansen<a href="https://khansen@jhsph.edu">khansen@jhsph.edu</a>>.

### See Also

read.metharray.exp and read.metharray for functions reading IDAT files. list.files for help on the arguments recursive and ignore.case.

### **Examples**

```
if(require(minfiData)) {
baseDir <- system.file("extdata", package = "minfiData")
sheet <- read.metharray.sheet(baseDir)
}</pre>
```

readGEORawFile

Read in Unmethylated and Methylated signals from a GEO raw file.

# **Description**

Read in Unmethylated and Methylated signals from a GEO raw file.

# Usage

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# **Arguments**

filename The name of the file to be read from.

sep The field separator character. Values on each line of the file are separated by this

character.

Uname A string that uniquely identifies the columns containing the unmethylated sig-

nals.

Mname A string that uniquely identifies the columns containing the methylated signals.

row.names The column containing the feature (CpG) IDs.

pData A DataFrame or data. frame describing the samples represented by the columns

of mat. If the rownames of the pData don't match the colnames of mat these colnames will be changed. If pData is not supplied, a minimal DataFrame is

created.

array Array name.

annotation The feature annotation to be used. This includes the location of features thus

depends on genome build.

mergeManifest Should the Manifest be merged to the final object.

showProgress TRUE displays progress on the console. It is produced in fread's C code.

... Additional arguments passed to data.table::fread().

### **Details**

450K experiments uploaded to GEO typically include a raw data file as part of the supplementary materials. Unfortunately there does not appear to be a standard format. This function provides enough flexibility to read these files. Note that you will likely need to change the sep, Uname, and Mname arguments and make sure the first column includes the feature (CpG) IDs. You can use the readLines function to decipher how to set these arguments.

Note that the function uses the fread function in the **data.table** package to read the data. To install **data.table** type install.packages("data.table"). We use this package because the files too large for read.table.

### Value

A GenomicMethylSet object.

### Author(s)

Rafael A. Irizarry<rafa@jimmy.harvard.edu>.

### See Also

getGenomicRatioSetFromGEO

readTCGA 67

### **Examples**

readTCGA

Read in tab deliminited file in the TCGA format

# **Description**

Read in tab deliminited file in the TCGA format

# Usage

### **Arguments**

filename	The name of the file to be read from.
sep	The field separator character. Values on each line of the file are separated by this character.
keyName	The column name of the field containing the feature IDs.
Betaname	The character string contained all column names of the beta value fields.
pData	A DataFrame or data. frame describing the samples represented by the columns of mat. If the rownames of the pData don't match the colnames of mat these colnames will be changed. If pData is not supplied, a minimal DataFrame is created.
array	Array name.
annotation	The feature annotation to be used. This includes the location of features thus depends on genome build.
mergeManifest	Should the Manifest be merged to the final object.
showProgress	TRUE displays progress on the console. It is produced in fread's C code.

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### **Details**

This function is a wrapper for makeGenomicRatioSetFromMatrix. It assumes a very specific format, used by TCGA, and then uses the fread function in the **data.table** package to read the data. To install **data.table** type install.packages("data.table"). We use this package because the files too large for read.table.

Currently, an example of a file that this function reads is here: http://gdac.broadinstitute.org/runs/stddata\_\_2014\_10\_17/data/UCEC/20141017/gdac.broadinstitute.org\_UCEC.Merge\_methylation\_humanmethylation450\_\_jhu\_usc\_edu\_\_Level\_3\_\_within\_bioassay\_data\_set\_function\_\_data.Level\_3.2014101700.0.0.tar.gz. Note it is a 8.1 GB archive.

### Value

A GenomicRatioSet object.

### Author(s)

Rafael A. Irizarry<rafa@jimmy.harvard.edu>.

### See Also

makeGenomicRatioSetFromMatrix

# **Examples**

```
## Not run:
    filename <- "example.txt" ##file must be in the specicif TCGA format
    readTCGA(filename)
## End(Not run)</pre>
```

RGChannelSet-class

Class "RGChannelSet"

# **Description**

These classes represents raw (unprocessed) data from a two color micro array; specifically an Illumina methylation array.

# Usage

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```
NBeads = new("matrix"), annotation = "", ...)
## Accessors
## S4 method for signature 'RGChannelSet'
annotation(object)
## S4 method for signature 'RGChannelSet'
pData(object)
## S4 method for signature 'RGChannelSet'
sampleNames(object)
## S4 method for signature 'RGChannelSet'
featureNames(object)
## S4 method for signature 'RGChannelSet'
getBeta(object, ...)
getGreen(object)
getRed(object)
getNBeads(object)
## S4 method for signature 'RGChannelSet'
getManifest(object)
## Convenience functions
get00B(object)
getSnpBeta(object)
```

### **Arguments**

object	An RGChannelSet (or RGChannelSetExtended).
Green	A matrix of Green channel values (between zero and infinity) with each row being a methylation loci and each column a sample.
Red	See the Green argument, but for the Green channel.
GreenSD	See the Green argument, but for standard deviations of the Green channel summaries.
RedSD	See the Green, but for standard deviations of the Red channel summaries.
NBeads	See the Green argument, but contains the number of beads used to summarize the Green and Red channels.
annotation	An annotation string, optional.
•••	For the constructor(s), additional arguments to be passed to SummarizedExperiment; of particular interest are colData, rowData and metadata. For getBeta these values gets passed onto getBeta.

### Value

An object of class RGChannelSet or RGChannelSetExtended for the constructors.

# Constructors

Instances are constructed using the RGChannelSet or RGChannelSetExtended functions with the arguments outlined above.

70 RGChannelSet-class

as(object, "RGChannelSet") coerces a RGChannelSetExtended object into a RGChannelSet.

#### Accessors

getGreen: Gets the Green channel as a matrix.

getRed: Gets the Red channel as a matrix.

getNBeads: Gets the number of beads as a matrix, this requires an RGChannelSetExtended.

getManifest: Gets the manifest object itself associated with the array type

### **Convenience functions**

get00B: Retrives the so-called "out-of-band" (OOB) probes. These are the measurements of Type I probes in the "wrong" color channel. Return value is a list with two matrices, named Red and Grn.

getSnpBeta: Retrives the measurements of the 65 SNP probes located on the array. These SNP probes are intended to be used for sample tracking and sample mixups. The return value is a matrix of beta values. Each SNP probe ought to have values clustered around 3 distinct values corresponding to homo-, and hetero-zygotes.

combine: Combines two different RGChannelSet, eventually using the combine method for eSet.

# **Tips**

The class inherits a number of useful methods from SummarizedExperiment. In earliers versions of minfi, this class inherited from eSet, and we have kept of number of methods related to this, for example pData.

The best way to access phenotype data and sample names are colData and colnames.

Amongst the useful methods are

dim, nrow, ncol The dimension (number of probes by number of samples) of the experiment.

colData, colnames, pData, sampleNames Phenotype information and sample names.

rownames, featureNames This is the addresses (probe identifiers) of the array.

### Author(s)

Kasper Daniel Hansen < khansen@jhsph.edu>

### See Also

See SummarizedExperiment for the basic class that is used as a building block for "RGChannelSet(Extended)". See IlluminaMethylationManifest for a class representing the design of the array.

```
showClass("RGChannelSet")
```

subsetByLoci 71

subsetByLoci Subset an RGChannelset by CpG loci.
--

# **Description**

Subset an RGChannelSet by CpG loci.

# Usage

# Arguments

rgSet An object of class RGChannelSet (or RGChannelSetExtended).
includeLoci A character vector of CpG identifiers which should be kept.

A character vector of CpG identifiers which should be excluded.
keepControls Should control probes be kept?

keepSnps Should SNP probes be kept?

### **Details**

This task is non-trivial because an RGChannelSet is indexed by probe position on the array, not by loci name.

# Value

An object of class RGChannelSet, which some probes removed.

```
if(require(minfiData)) {
  loci <- c("cg00050873", "cg00212031", "cg00213748", "cg00214611")
  subsetByLoci(RGsetEx.sub, includeLoci = loci)
  subsetByLoci(RGsetEx.sub, excludeLoci = loci)
}</pre>
```

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