# Package 'charm'

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|--|
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| Suggests charmData, BSgenome.Hsapiens.UCSC.hg18, corpcor   |
| Description This package implements analysis tools for DNA methylation data generated using Nimblegen microarrays and the McrBC protocol. It finds differentially methylated regions between samples, calculates percentage methylation estimates and includes array quality assessment tools. |
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Description

bgAdjust

Estimate and remove background signal using anti-genomic background probes

Remove background

#### Usage

Index

```
bgAdjust(dat, copy=TRUE)
```

### **Arguments**

dat a TilingFeatureSet

copy Only relevant when using disk-backed objects. If TRUE a copy will be made

leaving the original object (dat) unchanged. The input object will not be pre-

served if copy=FALSE

### **Details**

Background signal removal using a modified version of the RMA convolution model. The background signal level is estimated within GC-strata using anti-genomic background probes.

### Value

 $a\ Tiling Feature Set$ 

clusterMaker 3

#### Author(s)

Martin Aryee <aryee@jhu.edu>

### **Examples**

# See normalizeBetweenSamples

clusterMaker

Redefine array regions given chromosomal coordinates.

### Description

Redefine array regions given chromosomal coordinates.

### Usage

```
clusterMaker(chr,pos,order.it=TRUE,maxGap=300)
```

### **Arguments**

chr Vector of chromosome names

pos Vector of positions within chromosomes.

order.it order probes if they are not already ordered.

maxGap maximum allowable gap between probe start positions for probes to be grouped

into the same region.

### **Details**

Redefine array regions given chromosomal coordinates.

### Author(s)

Martin Aryee <aryee@jhu.edu>, Peter Murakami, Rafael Irizarry

### See Also

dmrFind, plotDMRs, plotRegions

### **Examples**

# See qval

4 cmdsplot

| cmdsplot Classical multi-dimensional scaling plot of charm data. |
|--|
|--|

### Description

Unsupervised clustering of charm data samples by classical multi-dimensional scaling.

### Usage

```
cmdsplot(labcols, expcol, rawData, p, okqc=1:nrow(p), noXorY=TRUE, outfile="./cmds_topN.pdf", top
```

### **Arguments**

| labcols | vector of colors, one for each group being plotted (each unique value of pData(rawData)[,expcol])            |
|---------|--|
| expcol  | name of the column of pData(rawData) the values of which the points are colored differently by.              |
| rawData | the TilingFeatureSet object that p came from.  |
| р       | the matrix of percentage methylation values (scale: 0, 1) from using methp on rawData. One column per sample |

okqc vector of indices identifying which rows of p to use.

noXorY logical value for whether or not (TRUE or FALSE) to ignore probes in chr X

and Y?

outfile file argument to pdf()

topN Use only the

### Author(s)

Martin Aryee <aryee@jhu.edu>, Peter Murakami, Rafael Irizarry

#### See Also

```
regionPlot, dmrFinder, dmrFdr
```

### **Examples**

# See dmrFdr

controlQC 5

### Description

Make boxplots of the non-control probes and the control probes (after spatial and background correction but before any normalization), to confirm that the control probes have a lower distribution of intensities than the non-control probes.

### Usage

```
controlQC(rawData, controlProbes=NULL, controlIndex=NULL, IDcol, expcol, ylimits=c(-6,8), outfile
```

### **Arguments**

| rawData       | a TilingFeatureSet object  |
|---------------|--|
| controlProbes | character string of the label assigned to non-CpG control probes in the annotation file (i.e. the container column of the .ndf file) (getContainer reports the container values for the pm probes). This is used only if controlIndex is not provided. Either this or controlIndex must be provided. |
| controlIndex  | a vector of non-CpG control probe indices (can be obtained using getControlIndex). Either this or controlProbes must be provided.  |
| IDcol         | column name of pData(rawData) with which to label samples in the output plot.  |
| expcol        | name of the column of pData(rawData) by which to order and, if not numeric, color the boxes.   |
| ylimits       | ylim argument to plot()  |
| outfile       | file argument to pdf()   |
| height        | height argument to pdf()   |
| width         | width argument to pdf()  |

#### Value

A data frame with one row per sample and columns

```
non_control Median methylation value among the non-control probescontrol Median methylation value among the control probesdiff non_control-control
```

#### Author(s)

Martin Aryee <aryee@jhu.edu>, Peter Murakami, Rafael Irizarry

### See Also

```
regionPlot, dmrFinder, dmrFdr
```

```
# See dmrFdr
```

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countGC

Count probe GC content

### **Description**

Return the GC content for each probe

### Usage

```
countGC(dat, type = "pm", idx)
```

#### **Arguments**

dat a TilingFeatureSet object

type pm or bg probes

idx An optional vector of probe indices for which to return GC content. If not

specified, values for all pm (or bg) probes will be returned.

### **Details**

This function returns the sum of #G + #C in the pm or bg probes.

### Value

a numeric vector

#### Author(s)

Martin Aryee <aryee@jhu.edu>

### See Also

readCharm

```
if (require(charmData)) {
  phenodataDir <- system.file("extdata", package="charmData")
  pd <- read.delim(file.path(phenodataDir, "phenodata.txt"))
  pd <- subset(pd, sampleID=="441_liver")
  dataDir <- system.file("data", package="charmData")
  setwd(dataDir)
  rawData <- readCharm(files=pd$filename, sampleKey=pd)
  ngc <- countGC(rawData)
  head(ngc)
}</pre>
```

cpgdensity 7

Get CpG density for genomic regions

### **Description**

Calculate the CpG density for a set of windows

#### Usage

```
cpgdensity(subject, chr, pos, windowSize = 500, sequence = "CG")
```

### Arguments

subject BSGenome object (e.g. Hsapiens)

chr character vector
pos numeric vector
windowSize number value
sequence character string

### **Details**

Calculate the CpG density for a set of regions. chr and pos specify the region mid-points and windowSize specifies the size of the window to be centered on these mid-points. i.e. The window will stretch from pos-windowSize/2 to pos+windowSize/2.

#### Value

a numeric vector

### Author(s)

Martin Aryee <aryee@jhu.edu>

```
if (require(BSgenome.Hsapiens.UCSC.hg18)){
    chr <- c("chr1", "chr1", "chr2")
    pos <- c(100000, 100500, 100000)
    cpgd <- cpgdensity(Hsapiens, chr=chr, pos=pos, windowSize = 500)
    cpgd
}</pre>
```

8 dmrFdr

dmrFdr

Calculate FDR q-values for differentially methylated regions (DMRs)

### **Description**

Estimate false discovery rate q-values for a set of differentially methylated regions (found using the dmrFinder function) using a permutation approach. For differentially methylated regions found using the dmrFind function, use the qval function instead.

#### Usage

```
dmrFdr(dmr, compare = 1, numPerms = 1000, seed = NULL, verbose = TRUE)
```

### **Arguments**

dmr a dmr object as returned by dmrFinder

compare The dmr table for which to calculate DMRs. See details.

numPerms Number of permutations

seed Random seed (for reproducibility)

verbose Boolean

#### **Details**

This function estimates false discovery rate q-values for a dmr object returned by dmrFinder. dmrFinder can return a set of DMR tables with one or more pair-wise comparisons between groups. dmrFdr currently only calculated q-values for one of these at a time. The dmr table to use (if the dmr object contains more than one) is specified by the compare option.

#### Value

a list object in the same format as the input, but with extra p-val and q-val columns for the tabs element.

#### Author(s)

Martin Aryee <aryee@jhu.edu>

#### See Also

```
qval, dmrFinder, dmrPlot, regionPlot
```

```
if (require(charmData) & require(BSgenome.Hsapiens.UCSC.hg18)) {
    phenodataDir <- system.file("extdata", package="charmData")
    pd <- read.delim(file.path(phenodataDir, "phenodata.txt"))
    pd <- subset(pd, tissue %in% c("liver", "colon"))
# Validate format of sample description file
    res <- validatePd(pd)
    dataDir <- system.file("data", package="charmData")
    setwd(dataDir)</pre>
```

dmrFind 9

```
# Read in raw data
rawData <- readCharm(files=pd$filename, sampleKey=pd)</pre>
# Find non-CpG control probes
ctrlIdx <- getControlIndex(rawData, subject=Hsapiens)</pre>
# Estimate methylation
p <- methp(rawData, controlIndex=ctrlIdx)</pre>
# Find differentially methylated regions
grp <- pData(rawData)$tissue</pre>
dmr <- dmrFinder(rawData, p=p, groups=grp,</pre>
                       removeIf=expression(nprobes<4 | abs(diff)<.05 | abs(maxdiff)<.05),</pre>
compare=c("liver", "colon"), cutoff=0.95)
head(dmr$tabs[[1]])
# Estimate false discovery rate for DMRs
dmr <- dmrFdr(dmr, numPerms=3, seed=123)</pre>
head(dmr$tabs[[1]])
                 ##Not run:
                 ## Plot top 10 DMRs:
             #cpg.cur = read.delim("http://rafalab.jhsph.edu/CGI/model-based-cpg-islands-hg18.txt", as.is=T
             #dmrPlot(dmr=dmr, which.table=1, which.plot=1:5, legend.size=1, all.lines=TRUE, all.points=TRUE
             ## plot any given genomic regions using this data, supplying the regions in a data frame that mus
             #mytab = data.frame(chr=as.character(c(dmr$tabs[[1]]$chr[1],"chrY",dmr$tabs[[1]]$chr[-1])), st
             #regionPlot(tab=mytab, dmr=dmr, cpg.islands=cpg.cur, Genome=Hsapiens, outfile="./myregions.pdf
                 ## note that region 2 is not plotted since it is not on the array.
                 ## Example of paired analysis:
             pData(rawData)pair = c(1,2,1,2) ## fake pairing information for this example.
                 dmr2 <- dmrFinder(rawData, p=p, groups=grp,</pre>
                                   compare=c("colon", "liver"),
                            removeIf=expression(nprobes<4 | abs(diff)<.05 | abs(maxdiff)<.05),</pre>
                                   paired=TRUE, pairs=pData(rawData)$pair, cutoff=0.95)
             #dmrPlot(dmr=dmr2, which.table=1, which.plot=c(3), legend.size=1, all.lines=TRUE, all.points=TF
             #regionPlot(tab=mytab, dmr=dmr2, cpg.islands=cpg.cur, Genome=Hsapiens, outfile="myregions.pdf"
}
```

dmrFind

Identify DMR candidates using a regression-based approach and correcting for batch effects.

#### **Description**

Identify DMR candidates using a regression-based approach and correcting for batch effects.

#### Usage

```
dmrFind(p=NULL, logitp=NULL, svs=NULL, mod, mod0, coeff, pns, chr, pos, only.cleanp=FALSE, only.dm
```

#### **Arguments**

matrix of methylation percentage estimates. Either this or logitp must be provided. Can be an ff\_matrix object.

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logitp matrix of logit-transformed methylation percentage estimates. Either this or p must be provided. Can be an ff matrix object. surrogate variables whose effect will be corrected for. This should be svaobj\$sv, SVS where svaobj is the object returned by sva(). Setting svs=0 will result in sva not being used. The mod argument provided to sva() which yielded svs. This should be a design mod matrix with all the adjustment covariates and (in the rightmost column(s)) your covariate of interest. mod0 The mod0 argument provided to sva() which yielded svs. This should be a design matrix with just the adjustment covariates. Thus it should be the same as mod, excluding the rightmost column(s) for the covariate of interest. a character or numeric index for the column of mod that identifies the covariate coeff column of interest. vector of region names for the probes corresponding to rows of p or logitp. pns vector of chromosomal identifiers for the probes corresponding to rows of p or logitp. vector of chromosomal coordinates for the probes corresponding to rows of p or pos logitp. if TRUE, only return the matrix of methylation percentage estimates after reonly.cleanp moving the batch effects (columns of sv) only.dmrs if TRUE, do not return the matrix of methylation percentage estimates that have had the batch effects (columns of sv) removed (called cleanp). rob One of the outputs of dmrFind is cleanp, which is the input p matrix after removing batch effects identified by SVA. By default these are the only effects removed from the p matrix. However, if you set rob=FALSE, then the other adjustment variables in mod and mod0 (all other variables besides the covariate of interest) are also removed. This will affect the methylation levels shown in plots using plotDMRs, plotRegions, and any other function that uses the cleanp output of dmrFind. It does not affect the selection of DMRs, though, except in the application of the filter at the end of the function where DMRs with an average difference between the 2 groups (or the average correlation between methylation and the covariate if the covariate is continuous) of less than the min.value argument are filtered out, since this step uses the cleanp matrix to calculate those averages or that correlation. use.limma Use the linear modeling approach (borrowing strength across probes) of lmFit in the limma package. which method to use for smoothing. "weighted loess", "loess", or "runmed". smoo k argument to runmed() if smoo="runmed". k see DELTA. Only used if smoo="loess" **SPAN** span parameter in loess smoothing will = SPAN/(DELTA \* number of probes in **DELTA** the plotted region). Only used if smoo="loess". use If "sbeta", identify DMRs by segmenting the smoothed effect estimates. If

min.probes The minimum allowable number of probes in a DMR candidate.

quantile.

Q

"swald", identify DMRs by segmenting the smoothed wald statistics.

Identify DMRs as the consecutive groups of probes whose smoothed effect es-

timate (if use="sbeta") or smoothed wald statistics (if use="swald") exceed this

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min.value The minimum allowable average difference in methylation percentage between

the 2 groups if covariate is categorical, or the minimum average correlation be-

tween methylation and the covariate if covariate is continuous.

keepXY if FALSE, exclude DMRs in "chrX" and "chrY".

sortBy column of DMR table to sorty by. verbose print progress messages if TRUE.

vfilter vfilter argument to sva function. The number of most variable probes to use

when building SVs-must be between 100 and m, where m is the total number of probes. vfilter=NULL by default, which means vfilter=m. Setting this to something smaller, like 100000, will typically yield satisfactory SVs and is advisable when the size of the data is very large and memory is limited (as when the p

and/or logitp arguments given to dmrFind are ff\_matrix objects).

nsubsets used if p or logitp are ff\_matrix objects. Rather than doing computations on

the whole logitp matrix, break up its m rows into chunks of m/nsubsets rows. Default is 50, but if even that uses too much memory, set this higher. Results do

not depend on the value chosen.

... Additional arguments passed to sva()

#### **Details**

Identify DMR candidates using a regression-based approach and correcting for batch effects.

#### Value

If only.cleanp=TRUE, only the cleanp matrix (below) is returned. Otherwise, the function returns a list with

dmrs A data frame with all DMR candidates, with columns:

chr chromosome of DMRstart start of DMR (bp)end end of DMR (bp)

value average value of the smoothed effect estimate within the DMR if use="sbeta" (the default), or the average value of the smoothed wald statistic within the DMR if use="swald"

area nprobes x value

**pns** name of the region on the array in which the DMR candidate was identified. **indexStart** index of first probe in DMR. This indexes chr, pos, pns, and cleanp **indexEnd** index of last probe in DMR. This indexes chr, pos, pns, and cleanp **nprobes** number of probes for the DMR, i.e., indexEnd-indexStart+1

**avg** average (across probes) percentage methylation difference within the DMR if covariate is categorical, or average (across probes) correlation between cleanp and covariate if covariate is continuous.

max maximum (across probes) percentage methylation difference within the DMR if covariate is categorical, or maximum (across probes) correlation between cleanp and covariate if covariate is continuous.

area.raw nprobes x avg

pval a vector of p-values for the t-test at each probe (in same order as rows of cleanp)

pns a vector of probe region names corresponding to the rows of cleanp chr a vector of chromosomes corresponding to the rows of cleanp 12 dmrFinder

pos a vector of positions corresponding to the rows of cleanp

args A list containing all the arguments provided to dmrFind. If svs was not provided,

svs here will be the surrogate variables obtained from sva.

If only.dmrs=FALSE,

cleanp The matrix of percentage methylation estimates, after subtracting batch effects.

If rob=FALSE, the effects of the other adjustment covariates are removed also.

If covariate is continuous,

beta the effect estimate at each probe.

sbeta the smoothed effect estimate at each probe.

#### Author(s)

Martin Aryee <aryee@jhu.edu>, Peter Murakami, Rafael Irizarry

#### See Also

```
plotDMRs, plotRegions, qval
```

#### **Examples**

# See qval

dmrFinder

Find differentially methylated regions (DMRs)

#### **Description**

Find differentially methylated regions (DMRs) from tiling microarray data. If you want to adjust for covariates (including batch effects estimated using SVA) or if your covariate of interest is continuous, use the dmrFind function.

### Usage

#### Arguments

| eset   | a TilingFeatureSet   |
|--------|--|
| groups | a vector of group labels for the samples in eset                               |
| р      | a matrix of percentage methylation values (scale: 0, 1). One column per sample |
| 1      | a matrix of methylation values (scale: -Inf, Inf), typically log-ratios.       |
| chr    | vector of chromosome labels for the probes in eset, p or l                     |

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pos vector of chromosomal coordinates for the probes in eset, p or l

pns vector of region names for the probes in eset, p or l

sdBins not currently implemented

controlIndex vector of indices of non-CpG control probes (which can be obtained using get-

ControlIndex), to be passed on to the methp function if neither the p nor the l arguments are provided, in which case either this or the controlProbes argument

must be provided.

controlProbes names of probe containers corresponding to control probes (e.g., see values re-

turned by getContainer), to be passed on to the methp function if neither the p nor the l arguments are provided, in which case either this or the controlIndex

argument must be provided.

Indexes not currently used

filter smoothing window weights. See details

package annotation package name

ws smoothing window size parameter. See details.

verbose Verbose progress reporting

compare the groups between which to find DMRs.

withinSampleNorm

within-sample normalization method. "loess" or "none"

betweenSampleNorm

between-sample normalization method. "quantile", "sqn" or "none"

cutoff t-statistic cutoff used to identify probes as being in a DMR

sortBy sort column for the DMR table. "area", "ttarea", "avg.diff", or "max.diff".

removeIf expression indicating which DMRs to drop from the DMR tables that get re-

turned. The negation of this is used as the subset argument to the subset function when it is called on the final DMR table before it is returned. If NULL, no DMRs will be subsetted out from the final table before it is returned. DMR table column names to use are listed below. E.g., to drop all DMRs with less than 4

probes, set removeIf=expression(nprobes<4).

paired if TRUE, do comparisons within pairs of samples. FALSE by default.

pairs if paired=TRUE, this must be provided. a vector of pair identifiers for the sam-

ples in eset. values must be the same within pairs and different between pairs.

DD Ignore this argument.

COMPS Ignore this argument.

COMPS. names Ignore this argument.

### **Details**

This function finds differentially methylated regions (DMRs). The sortBy parameter can be used to sort the DMRs by area (# probes x average difference), t-statistic area (# probes x average t-statistic), average difference, or maximum difference.

#### Value

A list with

tabs A list of DMR tables, one per comparison with columns:

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**chr** chromosome of DMR (bp)

start start of DMR (bp)

end end of DMR (bp)

**p1** if paired=FALSE, and p!=NULL or l=NULL, average percentage methylation of all probes between start and end for group 1

**p2** if paired=FALSE, and p!=NULL or l=NULL, average percentage methylation of all probes between start and end for group 2

m1 if paired=FALSE, p=NULL and 1!=NULL, average methylation 1 (logit(percentage methylation) if l=NULL) of all probes between start and end for group 1

**m2** if paired=FALSE, p=NULL and 1!=NULL, average methylation 1 (logit(percentage methylation) if l=NULL) of all probes between start and end for group 2

**regionName** name of the tiling region in which the DMR is found (These names come from the NDF file)

indexStart index of first probe in DMR. This indexes the output of dmrFinder,
 \*not\* the input.

**indexEnd** index of last probe in DMR. This indexes the output of dmrFinder, \*not\* the input.

**nprobes** number of probes for the DMR, i.e., indexEnd-indexStart+1

diff average percentage methylation difference within the DMR (i.e., column p1 - column p2), if p or eset arguments are provided. Otherwise, if only 1 argument is provided, it is the average difference in 1 (i.e, column m1 - column m2). Prior to version 2.0.1, if paired=TRUE, this was the average 1 (logit(percentage) methylation if l=NULL) difference within the DMR regardless of whether p or eset were provided.

maxdiff maximum percentage methylation difference within the DMR, if p or eset arguments are provided. Otherwise, if only 1 argument is provided, it is the maximum difference in 1. Prior to version 2.0.1, if paired=TRUE, this was the maximum 1 (logit(percentage) methylation if l=NULL) difference within the DMR regardless of whether p or eset were provided. Also prior to package version 2.0.1, this column was reported only in absolute value, however, post-version 2.0.1 the sign is retained.

area nprobes x average difference

**ttarea** nprobes x (average probe level t-statistic for between group difference)

A matrix of percentage methylation estimates (NOTE: the probe order may differ from that of the input p matrix since probes are sorted into chromosomal order)

This contains methylation log-ratios if they were passed to the function. Otherwise it contains logit-transformed percentage methylation estimates. (NOTE: the probe order may differ from that of the input l matrix since probes are sorted into chromosomal order)

a vector of chromosomes corresponding to the rows of p and l

a vector of positions corresponding to the rows of p and l

a vector of probe region names corresponding to the rows of p and l

a vector identifying which subset of the input probes (i.e. which elements of the input chr, pos, and pns, and rows of the input p and/or l) were used to search for DMRs. The output objects (chr, pos, pns, p, l, etc) are this subset of probes from the input. Therefore, e.g., while tabs\$indexStart:tabs\$indexEnd indexes the elements or rows of the output objects for each DMR candidate in tabs, index[tabs\$indexStart:tabs\$indexEnd] indexes the elements or rows of the input objects.

g

1

chr

pos

pns

index

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| gm      | if paired=FALSE, group medians of the 1 matrix  |
|---------|---|
| DD      | if paired=TRUE, a list of within-pair differences for each comparison                 |
| sMD     | if paired=TRUE, a matrix of smoothed mean within-pair differences for each comparison |
| groups  | a vector of group labels  |
| args    | the DMR finder parameter vector   |
| comps   | the vector of pairwise group comparisons  |
| package | the array annotation package name   |

### Author(s)

Martin Aryee <aryee@jhu.edu>, Peter Murakami, Rafael Irizarry

#### See Also

dmrFind, readCharm, methp, dmrFdr

### **Examples**

# See dmrFdr

| dmrPlot | Plot differentially methylated regions (DMRs) found using the dm-rFinder function. |
|---------|--|
|         | v  |

### Description

Plot differentially methylated regions (DMRs) from tiling microarray data that were identified using the dmrFinder function. To plot DMRs identified using the dmrFind function, use the plotDMRs function.

### Usage

```
\label{lem:main_dmr} $$\dim \operatorname{Plot}(\dim, \ \operatorname{which.table=1:length}(\dim \operatorname{stabs}), \ \operatorname{which.plot=1:30}, \ \operatorname{legend.size=1}, \ \operatorname{all.lines=TRUE}, \ \operatorname{all.lines=TR
```

### Arguments

| dmr         | a list object as returned by dmrFinder.  |
|-------------|--|
| which.table | a vector of indices identifying which tables in the dmr list to plot regions from.                     |
| which.plot  | a vector of indices identifying which regions (rows) from each table to plot.                          |
| legend.size | cex argument for the legend (factor by which to magnify/shrink the legend).                            |
| all.lines   | if TRUE, plot the smooth lines for all groups. If FALSE, only for the 2 groups being compared.         |
| all.points  | if TRUE, plot the points for all groups. If FALSE, only for the 2 groups being compared.               |
| colors.l    | a vector of line colors, one color for each group whose line is to be plotted (in alphabetical order). |

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colors.p a vector of point colors, one color for each group whose points are to be plotted (in alphabetical order). outpath where to save the output pdf file. a table with columns "chr", "start", and "end" for CpG islands to plot in the cpg.islands second panel. Genome the BSgenome object for the organism based upon which your array was designed. plotG if TRUE, a third panel of each DMR plot will show the difference between the median green channel value (after subtracting probe medians and correcting for gc content) between the 2 groups. If true, dat must be provided. dat if plotG=TRUE, this must be provided. It is the TilingFeatureSet object used when estimating the matrix of percent methylation estimates used in dmrFinder when it produced dmr. buffer Number of base pairs to plot on either side of each DMR candidate. set to FALSE if you want to plot the methlation values (the "I" output from plot.p

#### **Details**

This function plots the differentially methylated regions (DMRs). The second panel shows the location of CpG's with ticks on the bottom (islands are colored) and the location of mcrbc recognition sites with ticks on the top.

dmrFinder was run on l instead of p, plot.p=FALSE necessarily.

dmrFinder) instead of the percentage methylation values (the "p" output). If

### Author(s)

Martin Aryee <aryee@jhu.edu>, Peter Murakami, Rafael Irizarry

### See Also

```
plotDMRs, regionPlot, dmrFinder, dmrFdr
```

### **Examples**

# See dmrFdr

| getControlIndex | Get indices of control probes from CpG-free regions |
|-----------------|---|
|                 |   |

### **Description**

Get indices of control probes from CpG-free regions.

### Usage

```
getControlIndex(dat, controlProbes = NULL, noCpGWindow = 1000, subject, onlyGood = FALSE, matrix =
```

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#### **Arguments**

dat TilingFeatureSet

controlProbes vector of names used to denote control probes in the 'container' column of the

Nimblegen annotation (ndf) file. Optional

noCpGWindow Size of the window centered on the probe that must be CpG-free

subject A BSgenome object onlyGood deprecated option matrix deprecated option

#### **Details**

The probes can either be identified as control probes in the microarray annotation package, or alternatively the function will search the genome (given an appropriate BSgenome object) for suitable probes.

#### Value

a vector

#### Author(s)

Martin Aryee <aryee@jhu.edu>

### **Examples**

# See dmrFdr

maxDensity

Find the mode of a density function

#### **Description**

Calculate a density function and find the max point

### Usage

```
maxDensity(x, n.pts = 2^14, minPoints=30)
```

#### Arguments

x a data vector

n.pts Number of points to use in density estimation
minPoints Minimum number of data points to accept

### **Details**

This function finds the maximum of a density function. It is identical to the (unexported) max.density function in affy except that it returns an NA if the number of data points provided is less than minPoints

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#### Value

a numeric value

#### Author(s)

Martin Aryee <aryee@jhu.edu>

#### **Examples**

```
x <- rnorm(1000)
maxDensity(x)</pre>
```

methp

Estimate DNA methylation

### Description

Estimate DNA methylation from McrBC/CHARM microarray data in terms of log-ratios or percentages.

### Usage

```
methp(dat, spatial = TRUE, bgSubtract = TRUE, withinSampleNorm = "loess",
scale = c(0.99, 0.99), betweenSampleNorm = "quantile",
controlProbes = NULL, controlIndex = NULL, excludeIndex = NULL,
commonMethPercentParams = NULL,
verbose = TRUE, returnM = FALSE,
plotDensity = NULL, plotDensityGroups = NULL)
```

### Arguments

dat a TilingFeatureSet object

spatial boolean indicating whether to correct spatial artefacts

bgSubtract boolean indicating whether to estimate and remove background signal before

computing log-ratios

 $within {\tt Sample Norm}$ 

within-sample normalization method. Choices are "loess" and "none". "loess" uses the control-probe loess procedure described in Aryee et al., 2011 (PMID:

20858772).

scale a numeric vector (x,y). The xth percentile of each sample is scaled to represent

y% methylation. The default c(0.99, 0.99) means probes in the 99% percentile

represent 99% methylation.

betweenSampleNorm

between-sample normalization method. Choices are "quantile", "sqn", and "none".

See Details for more fine-grained control.

 $control Probes \quad character \ string \ of \ the \ label(s) \ assigned \ to \ non-CpG \ control \ probes \ in \ the \ array$ 

design file, i.e. in the container column of the .ndf design file for your array. The getContainer function from the oligo package can be used do see the container values for all pm probes. This argument is used only if controlIndex is not

provided. Either this or controlIndex must be provided.

methPercent 19

controlIndex a numeric vector of probe indices indicating which pm probes are the non-CpG

control probes (can be obtained using getControlIndex). Either this or control-

Probes must be provided.

excludeIndex a numeric vector of probe indices indicating which pm probes to ignore when

creating the between-array normalization target distributions. These probes are also removed from all the plots created if plotDensity is specified, however be aware that they ARE still returned in the output of methp (just without having

had any between-array normalization applied to them).

commonMethPercentParams

boolean indicating whether a common set of parameters should be used for all

samples when converting M-values to percentage methylation.

verbose boolean: Verbose output?

returnM boolean. Return M-values without converting to percentage methylation esti-

mates

plotDensity if specified this is the filename of the pdf diagnostic density plots.

plotDensityGroups

numeric vector of group labels used to color lines in the diagnostic density plots

(see plotDensity option)

#### **Details**

This function provides probe-level estimates of percentage DNA methylation from CHARM microarray data.

#### Value

A matrix of probe-level percentage methylation estimates, one column per sample. If the dat argument contains ff rather than matrix objects (as when it is produced by readCharm after loading the ff package), the output will be an ff object.

#### Author(s)

Martin Aryee <aryee@jhu.edu>

### See Also

readCharm

### **Examples**

# See dmrFdr

methPercent Estimate percentage DNA methylation from log-ratios

### **Description**

Estimate percentage DNA methylation from log-ratios

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#### Usage

```
methPercent(m, pmIndex, ngc, commonParams = TRUE)
```

### **Arguments**

m a matrix of M-values (methylation log-ratios). One column per sample.

pmIndex A vector of probe indices to use in the calculation. Usually set to the indices of

the pm probes (excluding background and other non-specific controls) by using

pmIndex=pmindex(dat)

ngc a vector with GC-content of probes. Same length as nrow(m)

commonParams boolean indicating whether a common set of parameters should be used for all

samples when converting M-values to percentage methylation.

#### **Details**

This function estimates percentage DNA methylation from normalized methylation log-ratios (M-values).

#### Value

a matrix of percentage methylation estimates. Same dimensions as m

#### Author(s)

Martin Aryee <aryee@jhu.edu>

```
if (require(charmData) & require(BSgenome.Hsapiens.UCSC.hg18)) {
    phenodataDir <- system.file("extdata", package="charmData")
    pd <- read.delim(file.path(phenodataDir, "phenodata.txt"))
    pd <- subset(pd, sampleID=="441_liver")
    dataDir <- system.file("data", package="charmData")
    setwd(dataDir)
# Read in raw data
    rawData <- readCharm(files=pd$filename, sampleKey=pd)
# Find non-CpG control probes
    ctrlIdx <- getControlIndex(rawData, subject=Hsapiens)
# Get normalized methylation log-ratios
m <- methp(rawData, controlIndex=ctrlIdx, returnM=TRUE)
# Estimate percentage methylation
    ngc <- countGC(rawData)
p <- methPercent(m, ngc=ngc)
}</pre>
```

```
normalizeBetweenSamples
```

Between-sample normalization

### Description

Between-sample normalization for two-color DNA methylation microarray data.

### Usage

```
normalizeBetweenSamples (dat, copy=TRUE,
m="allQuantiles", untreated="none", enriched="none",
controlProbes=NULL, controlIndex=NULL, excludeIndex=NULL, verbose=FALSE)
```

### Arguments

| dat           | a TilingFeatureSet object   |
|---------------|---|
| сору          | Only relevant when using disk-backed objects. If TRUE a copy will be made leaving the original object (dat) unchanged. The input object will not be preserved if copy=FALSE   |
| m             | normalization method for log-ratios. "allQuantiles" for full quantile normalization, or "none"  |
| untreated     | normalization method for the untreated channel. "complete", "allQuantiles" or "none"  |
| enriched      | normalization method for the untreated channel. "sqn", "allQuantiles" or "none"   |
| controlProbes | character string of the label assigned to non-CpG control probes in the annotation file (i.e. the container column of the .ndf file).   |
| controlIndex  | a vector of non-CpG control probe indices   |
| excludeIndex  | a vector indicating which pm probes to ignore when creating normalization target distributions. Can be a vector of probe indices or a boolean vector of length(pmindex(dat)). |
| verbose       | boolean: Verbose output?  |

### **Details**

This function is used by methp performs between-sample normalization. It is normally not used directly by the user.

### Value

 $a\ Tiling Feature Set$ 

### Author(s)

Martin Aryee <aryee@jhu.edu>

#### See Also

methp

#### **Examples**

```
if (require(charmData) & require(BSgenome.Hsapiens.UCSC.hg18)) {
phenodataDir <- system.file("extdata", package="charmData")</pre>
pd <- read.delim(file.path(phenodataDir, "phenodata.txt"))</pre>
pd <- subset(pd, sampleID=="441_liver")</pre>
dataDir <- system.file("data", package="charmData")</pre>
setwd(dataDir)
rawData <- readCharm(files=pd$filename, sampleKey=pd)</pre>
# Correct spatial artifacts
dat <- spatialAdjust(rawData)</pre>
# Remove background signal
dat <- bgAdjust(dat)</pre>
# Find non-CpG control probes
ctrlIdx <- getControlIndex(rawData, subject=Hsapiens)</pre>
# Within-sample normalization
dat <- normalizeWithinSamples(dat, controlIndex=ctrlIdx)</pre>
# Within-sample normalization
dat <- normalizeBetweenSamples(dat)</pre>
}
```

normalizeWithinSamples

Within-sample normalization for two-color data

#### **Description**

Within-sample (between-channel) normalization for two-color DNA methylation microarray data. This function implements the control probe loess procedure described in Aryee et al., 2011 (PMID: 20858772).

#### Usage

```
normalizeWithinSamples(dat, copy=TRUE,
method = "loess", scale=c(0.99, 0.99),
controlProbes = NULL, controlIndex = NULL, approx=TRUE, breaks=1000, verbose=FALSE)
```

#### **Arguments**

| dat           | a TilingFeatureSet   |
|---------------|--|
| сору          | Only relevant when using disk-backed objects. If TRUE a copy will be made leaving the original object (dat) unchanged. The input object will not be pre-   |
|               | served if copy=FALSE   |
| method        | normalization method. "loess" or "none"  |
| scale         | a numeric vector (x,y). The xth percentile of each sample is scaled to represent y% methylation. The default c(0.99, 0.99) means probes in the 99% percentile represent 99% methylation. Set to NA for no scaling. |
| controlProbes | character string of the label assigned to non-CpG control probes in the annotation file (i.e. the container column of the .ndf file).  |
| controlIndex  | a vector of non-CpG control probe indices  |
| approx        | Bin probes by signal intensity when loess normalizing. Much faster when TRUE   |
| breaks        | Number of bins to use when approx=TRUE   |
| verbose       | boolean: Verbose output?   |

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#### **Details**

This function is used by methp performs within-sample (between-channel) normalization. It is normally not used directly by the user.

#### Value

```
a TilingFeatureSet
```

#### Author(s)

Martin Aryee <aryee@jhu.edu>, Rafael Irizarry

#### **Examples**

# See normalizeBetweenSamples

plotDensity

Log-ratio density plot for all probes and control probes

### **Description**

Make density plots of log-ratios for two-color microarray data. Two plots are produced: one for all probes on the array, and a second for the control probes.

### Usage

### **Arguments**

dat a TilingFeatureSet

rx x-axis range

controlIndex a vector of non-CpG control probe indices

controlProbes vector of names used to denote control probes in the 'container' column of the

Nimblegen annotation (ndf) file.

pdfFile name of output pdf file

main main title

lab vector of sample labels. If not specified the sample names from dat will be used.

excludeIndex a numeric vector of probe indices indicating which pm probes to ignore when

plotting.

#### **Details**

This function makes density plots for a) all probes and b) control probes. It is typically called from within methp when a file name is specified for its plotDensity option. The plots are useful for identifying problematic outlier samples.

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#### Value

No return value. Called for its side-effect of producing a pdf plot.

#### Author(s)

Martin Aryee <aryee@jhu.edu>

### **Examples**

```
if (require(charmData) & require(BSgenome.Hsapiens.UCSC.hg18)) {
phenodataDir <- system.file("extdata", package="charmData")</pre>
pd <- read.delim(file.path(phenodataDir, "phenodata.txt"))</pre>
# Read in raw data
dataDir <- system.file("data", package="charmData")</pre>
rawData <- readCharm(path=dataDir, files=pd$filename,</pre>
 sampleKey=pd)
ctrlIdx <- getControlIndex(rawData, subject=Hsapiens)</pre>
                 ## Not run:
#plotDensity(rawData, controlIndex=ctrlIdx, pdfFile="density.pdf")
```

plotDMRs

Plot differentially methylated regions (DMRs) found using the dmrFind function.

### **Description**

Plot differentially methylated regions (DMRs) from tiling microarray data that were identified using the dmrFind function.

### Usage

```
plotDMRs(dmrs, Genome, cpg.islands, exposure, outfile, which_plot=1:50, which_lines=NULL, which_p
```

### **Arguments**

a list object as returned by dmrFind. dmrs Genome the BSgenome object for the organism based upon which your array was de-

signed.

a table with columns "chr", "start", and "end" for CpG islands to plot in the cpg.islands

second panel.

The covariate of interest. exposure

outfile the name of the file to save (including the full path)

which\_plot numeric vector of indices identifying which DMR candidates from dmrs\$dmrs

to plot.

vector specifying which groups (unique elements of exposure) to plot the lines which\_lines

for. If NULL (the default), plots lines for all groups. Only applies if exposure is

categorical.

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| which_points | vector specifying which groups (unique elements of exposure) to plot the points for. If NULL (the default), plots points for all groups. Only applies if exposure is categorical.  |
|--------------|--|
| SPAN         | see DELTA. Only used if smoo="loess"   |
| DELTA        | span parameter in loess smoothing will = SPAN/(DELTA $*$ number of probes in the plotted region). Only used if smoo="loess".   |
| smoo         | "loess" for loess smoother or "runmed" for running median smoother (runmed with k=3). This does not need to be the same as the smoo argument to dmrFind.   |
| ADD          | Number of base pairs to plot on either side of each DMR candidate (if it is covered on the array).   |
| cols         | vector of colors to use, one for each group (if covariate is categorical)  |
| point.info   | if TRUE, function returns a table identifying which sample is plotted with which number or letter (if pch.groups=NULL, the default).   |
| legend.size  | magnification factor for the legend  |
| pch.groups   | vector whose length is equal to the number of samples. Each unique value will be plotted with a different point type.  |
| panel3       | if panel3="G", the third panel of each DMR plot will show the difference between the median green channel value (after subtracting probe medians and correcting for gc content) between the 2 groups (i.e., the group defined by mod[,coef] in dmrFind minus the reference group). If panel="G", seq argument must be provided. If panel!="G", the 3rd panel will show -log10(dmrs\$pval). G |
| G            | matrix of green channel intensities to use for plotting in the 3rd panel if panel3="G".  |
| seq          | vector of probe sequences corresponding to the rows of G (and dmrs $\c)$ if panel3="G".  |
|              |  |

### **Details**

This function plots the differentially methylated regions (DMRs) that were identified using the dmrFind function.

### Author(s)

Martin Aryee <aryee@jhu.edu>, Peter Murakami, Rafael Irizarry

### See Also

plotRegions, dmrFind, qval

### **Examples**

# See qval

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| plotRegions Plot user-provided regions. |
|---|
| Plot user-provided regions.             |

#### **Description**

Plot user-provided regions.

#### Usage

plotRegions(thetable, cleanp, chr, pos, seq=NULL, Genome, cpg.islands, exposure, exposure.continue

#### **Arguments**

thetable a table with columns for "chr", "start", and "end", identifying the genomic re-

gions to plot (if they are covered on the array).

cleanp the matrix of percent methylation estimates to be used for plotting

chr vector of chromosome labels for the probes in cleanp

pos vector of chromosomal coordinates for the probes in cleanp

seq vector of probe sequences corresponding to the rows of G (and cleanp). Needed

only if panel3="G".

Genome the BSgenome object for the organism based upon which your array was de-

signed.

cpg.islands a table with columns "chr", "start", and "end" for CpG islands to plot in the

second panel.

outfile the name of the file to save (including the full path)

exposure The covariate of interest.

exposure.continuous

set to TRUE if exposure is a continuous variable.

which\_lines vector specifying which groups (unique elements of exposure) to plot the lines

for. If NULL (the default), plots lines for all groups. Only applies if exposure is

categorical.

which\_points vector specifying which groups (unique elements of exposure) to plot the points

for. If NULL (the default), plots points for all groups. Only applies if exposure

is categorical.

ADD Number of base pairs to plot on either side of each DMR candidate (if it is

covered on the array).

cols vector of colors to use, one for each group (if covariate is categorical)

legend.size magnification factor for the legend

smoo "loess" for loess smoother or "runmed" for running median smoother (runmed

with k=3).

SPAN see DELTA. Only used if smoo="loess"

DELTA span parameter in loess smoothing will = SPAN/(DELTA \* number of probes in

the plotted region). Only used if smoo="loess".

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panel3 if panel3="G", the third panel of each DMR plot will show the difference between the median green channel value (after subtracting probe medians and correcting for gc content) between the 2 groups (i.e., the group defined by mod[,coef] in dmrFind minus the reference group). If panel="G", seq argument must be provided. If panel!="G", the 3rd panel will show -log10(dmrs\$pval).

G matrix of green channel intensities to use for plotting in the 3rd panel if panel3="G".

grs if panel3="G", plot difference between the median green channel value (af-

ter subtracting probe medians and correcting for gc content) between these 2

groups.

#### **Details**

This function plots user-provided regions.

#### Author(s)

Martin Aryee <aryee@jhu.edu>, Peter Murakami, Rafael Irizarry

#### See Also

```
plotDMRs, dmrFind, qval
```

#### **Examples**

# See qval

pmQuality

Calculate probe quality scores

### **Description**

pmQuality calculates probe quality for each pm probe by comparing the total DNA signal (green) to the distribution of the background probe signals. 0 means lower than all background probes. 100 means higher than all background probes.

### Usage

```
pmQuality(dat, channel="channel1", verbose=FALSE, idx=NULL)
```

### **Arguments**

dat a TilingFeatureSet object channel which channel to assess

verbose logical value for whether or not to print message to screen

idx vector of probe indices specifying which probes to evaluate. By default all pm

probes are used.

#### Author(s)

Martin Aryee <aryee@jhu.edu>, Peter Murakami, Rafael Irizarry

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#### See Also

```
regionPlot, dmrFinder, dmrFdr
```

#### **Examples**

# See dmrFdr

qcReport

Microarray quality report

#### **Description**

Calculate microarray quality scores and produce an optional pdf report

### Usage

```
qcReport(dat, file = NULL, utRange = c(30, 100), enRange = c(8, 12), numProbes = 5e+05, blockSize)
```

### **Arguments**

dat a TilingFeatureSet file name of output pdf file

utRange color-scale range for the untreated channel plots

enRange color-scale range for the methyl-depleted channel plots

numProbes maximum number of probes to use for plots. If smaller than the number of

probes on the array numProbes are chosen at random, speeding up calculations

for high-density arrays with several million probes.

blockSize The array is divided into a series of blockSize x blockSize rectangular blocks

and the average signal level calculated for each. If blockSize is unspecified a

size is chosen that gives about 1250 probes per block.

#### **Details**

This function calculates microarray quality scores and produces an optional pdf report. Three quality metrics are calculated for each array:

**Average signal strength.** The average percentile rank of untreated channel signal probes among the background (anti-genomic) probes. Since the untreated channel contains total DNA a successful hybridization would have strong signal for all untreated channel genomic probes.

**Untreated channel signal standard deviation.** The array is divided into a series of rectangular blocks and the average signal level calculated for each. Since probes are arranged randomly on the array there should be no large differences between blocks. Arrays with spatial artifacts have a larger standard deviation between blocks.

Methyl-depleted channel signal standard deviation

### Value

a matrix with a row for each sample. The 3 columns contain array signal strength score, untreated channel standard deviation and methyl-depleted channel standard deviation.

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#### Author(s)

Martin Aryee <aryee@jhu.edu>

#### **Examples**

qval

Obtain False Discovery Rate q-values for the DMR candidates returned by dmrFind.

#### **Description**

Obtain False Discovery Rate q-values for the DMR candidates returned by dmrFind.

### Usage

```
qval(p=NULL,\ logitp=NULL,\ dmr,\ numiter=500,\ seed=54256,\ verbose=FALSE,\ mc=1,\ return.permutations=100,\ seed=54256,\ seed=54
```

### Arguments

p The matrix of methylation percentage estimates provided to dmrFind (argument

p) when it returned dmr, if p was provided.

logitp The matrix of logit-transformed methylation percentage estimates provided to

dmrFind (argument logitp) when it returned dmr, if logitp was provided.

dmr The output of dmrFind with the dmr table to obtain q-values for.

numiter The number of times to iterate the resampling procedure.

seed random seed.

verbose verbose argument to dmrFind.

mc Number of CPUs to use for parallel processing.

return.permutations

if TRUE, also return a matrix defining the resampling at each iteration.

method

if "direct" (the default), the resulting table will have an additional column ("qvalue") with the direct estimate of the FDR qvalue as described below. If "pool", the resulting table will have additional columns ("pvalue.pool" and "qvalue.pool") with the p-values and corresponding q-values obtained as the proportion of statistics in the null distribution that are greater than or equal to the observed statistic, where the null distribution is obtained by simply pooling together all the statistic vectors from all iterations. If "fwer", the resulting table will have an additional column ("pvalue.fwer") with the p-value adjusted for control of the family-wise error rate (obtained as the proportion of maximum statistics (across all iterations) that are greater than or equal to the observed statistic). More than one value for method may be specified.

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fwer.num

positive integer vector, needed if method argument includes "fwer". For each number, a separate output column will be returned with the FWER(n)-controlling adjusted p-values, where FWER(n) is defined as the probability of making n or more false discoveries (type I errors).

#### **Details**

The q-value for region i is estimated by a resampling procedure. It is estimated as x/N, where N is the number of dmrs in the DMR table returned by dmrFind whose statistic (defined by the sortBy argument to dmrFind) is >= the statistic for region i, and x is the average (across all iterations' DMR tables) number of DMRs with statistic >= the statistic for region i. Monotonicity is then enforced and q-values are capped at 1. For example, to estimate the q-value for the 5th dmr in your dmr table that you ranked by area, N is 5. If it has area=100, then x is the average number of dmrs with area>=100, averaging over all the tables produced by random-resampling iterations. To the extent that the proportion of the genome covered by the array that is truly non-differentially methylated is less than 1, x will slightly overestimate the true number of false DMR candidates expected to be returned from the data, and hence the q-value will be slightly conservative.

The resampling procedure used to obtain null data for each iteration is as follows. 1. Add the surrogate variables identified by SVA to the null model defined by the mod0 argument to dmrFind, and obtain fitted methylation values M, where rows correspond to probes and columns correspond to arrays. 2. Add the surrogate variables identified by SVA to the full model defined by the mod argument to dmrFind, and obtain residuals R, where rows correspond to probes and columns correspond to arrays. Then, for iteration k, obtain a column-resampled version of R, Q (resampling with replacement), and obtain the methylation values to be used in iteration k as M+Q.

### Value

the dmr table in your dmr object (dmr\$dmrs), with an additional column ("qvalue") for the Fdr q-value (if method includes "direct"), additional columns for the pooled estimates of the p-values and q-values (if method includes "pool"), and an additional column for the FWER-adjusted p-values (if method includes "fwer") (see method argument). There is also a column called qvalue0 which is just the qvalue column before enforcing monotonicity.

### Author(s)

Martin Aryee, Peter Murakami <pmurakam@jhsph.edu>, Rafael Irizarry

### See Also

```
dmrFind, plotDMRs, plotRegions
```

```
## See vignette.
if (require(charmData) & require(BSgenome.Hsapiens.UCSC.hg18)) {
    dataDir <- system.file("data", package="charmData")
    phenodataDir <- system.file("extdata", package="charmData")
    pd <- read.delim(file.path(phenodataDir, "phenodata.txt"))
    res <- validatePd(pd)

## Read in raw data:
rawData <- readCharm(files=pd$filename, path=dataDir, sampleKey=pd, sampleNames=pd$sampleID)
    ## Check quality of arrays:
#qual <- qcReport(rawData, file="qcReport.pdf")</pre>
```

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```
## Assess individual probe qualities:
 pmq = pmQuality(rawData)
 rmpmq = rowMeans(pmq)
 okqc = which(rmpmq>75)
## Identify control probes as the probes at positions surrounded by a CpG-free 600bp window:
 ctrlIdx <- getControlIndex(rawData, subject=Hsapiens, noCpGWindow=600)</pre>
## Check that these control probes do indeed have lower intensities than the non-control probes (after
#controlQC(rawData=rawData, controlIndex=ctrlIdx, IDcol="sampleID", expcol="tissue", ylimits=c(-6,8),
 chr = pmChr(rawData)
 pns = probeNames(rawData)
 pos = pmPosition(rawData)
 seq = pmSequence(rawData)
 pd = pData(rawData)
 ## Estimate percent methylation:
 p <- methp(rawData, controlIndex=ctrlIdx, plotDensityGroups=pd$tissue)</pre>
 ## unsupervised clustering of samples:
#cmdsplot(labcols=c("red","black","blue"), expcol="tissue", rawData=rawData, p=p, okqc=okqc, noXorY=T
## Do not look for DMRs among control probes or probes with average probe quality score less than or equ
 Index=setdiff(which(rmpmq>75),ctrlIdx)
 Index = Index[order(chr[Index], pos[Index])]
 p = p[Index,]
 seq = seq[Index]
 chr = chr[Index]
 pos = pos[Index]
 pns = pns[Index]
 pns=clusterMaker(chr,pos)
## Identify DMR candidates between colon and liver (in this example not adjusting for any other covaria
 mod0 = matrix(1,nrow=nrow(pd),ncol=1)
 mod = model.matrix(~1 +factor(pd$tissue,levels=c("liver","colon","spleen")))
 thedmrs = dmrFind(p=p, mod=mod, mod0=mod0, coeff=2, pns=pns, chr=chr, pos=pos)
## Obtain FDR q-values for each DMR candidate. In practice, numiter should be set much higher.
 withq = qval(p=p, dmr=thedmrs, numiter=2, verbose=FALSE, mc=1)
 #### Plotting not run:
## Plot DMR candidates 1,2, and 4, for example. First have to load a table of CpG islands.
#cpg.cur = read.delim("http://rafalab.jhsph.edu/CGI/model-based-cpg-islands-hg18.txt", as.is=TRUE)
#plotDMRs(dmrs=thedmrs, Genome=Hsapiens, cpg.islands=cpg.cur, exposure=pd$tissue, outfile="./colon-li
## Plot DMR candidates, and in the 3rd panel plot the difference in average green channel:
 dat0 = spatialAdjust(rawData, copy=FALSE)
 dat0 = bgAdjust(dat0, copy=FALSE)
 G = pm(dat0)[,,1] #from oligo
 G = G[Index,]
#plotDMRs(dmrs=thedmrs, Genome=Hsapiens, cpg.islands=cpg.cur, exposure=pd$tissue, outfile="./colon-li
 ## Example if covariate of interest is continuous:
 pd$x = c(1,2,3,4,5,6)
 mod0 = matrix(1,nrow=nrow(pd),ncol=1)
```

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```
mod = model.matrix(~1 + pd$x)
    coeff = 2
  thedmrs2 = dmrFind(p=p, mod=mod, mod0=mod0, coeff=coeff, pns=pns, chr=chr, pos=pos)
  ## If covariate of interest is continuous, you can still plot it like it is categorical by categorizing
  groups = as.numeric(cut(mod[,coeff],c(-Inf,2,4,Inf))) #You can change these cutpoints.
    pd$groups = c("low","medium","high")[groups]
  #plotDMRs(dmrs=thedmrs2, Genome=Hsapiens, cpg.islands=cpg.cur, exposure=pd$groups, outfile="./test.pc
  ## Otherwise, if covariate of interest is continuous, plot will show correlation with covariate:
  #plotDMRs(dmrs=thedmrs2, Genome=Hsapiens, cpg.islands=cpg.cur, exposure=pd$x, outfile="./x.pdf", whic
    ## Plot arbitrary regions:
    mytable = thedmrs$dmrs[,c("chr","start","end")]
    mytable[2,] = c("chr1",1,1000) #not on array
    mytable$start = as.numeric(mytable$start)
    mytable$end = as.numeric(mytable$end)
  #plotRegions(thetable=mytable[1:5,], cleanp=thedmrs$cleanp, chr=chr, pos=pos, Genome=Hsapiens, cpg.is
}
```

readCharm

Read in McrBC/CHARM DNA methylation microarray data

#### **Description**

Read in DNA methylation microarray data from the McrBC/CHARM platform

### Usage

```
readCharm(files, path = ".", ut = "_532.xys", md = "_635.xys",
sampleKey, sampleNames = NULL, pkgname, type = NULL, ...)
```

### **Arguments**

files a vector of xys filenames path the path to the xys files

ut the file ending that designates untreated channel files

md the file ending that designates methyl-depleted channel files

sampleKey a data frame with sample description information. One line per xys file.

sampleNames a vector of names to use for the samples. One line per xys file.

pkgname the annotation package name

type deprecated option

... additional options passed on to read.xysfiles2

#### **Details**

This function is a convenience wrapper to read.xysfiles2 to simplify reading in DNA methylation data from the Nimblegen McrBC/CHARM microarray platform. It makes guesses about the extensions used for the methyl-depleted (md) and untreated channels (ut).

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### Value

A TilingFeatureSet object. If the ff package is loaded before using this function, the output will contain ff rather than matrix objects.

### Author(s)

Martin Aryee <aryee@jhu.edu>

### References

www.biostat.jhsph.edu/~maryee/charm

### See Also

methp, dmrFinder

### **Examples**

# See normalizeBetweenSamples

| regionMatch | Given two data frames with columns "chr", "start", and "end", identify the nearest region in one to the other. |
|-------------|--|
|             |  |

### Description

Given two data frames with columns "chr", "start", and "end", identify the nearest region in one (object2) to the other (object1).

### Usage

```
regionMatch(object1, object2, verbose=TRUE)
```

### Arguments

| object1 | Data frame with columns "chr", "start", and "end". For each of the rows of this data frame, find the nearest regions in object2. |
|---------|--|
| object2 | Data frame with columns "chr", "start", and "end".   |
| verbose | print progress.  |

### **Details**

Given two data frames with columns "chr", "start", and "end", identify the nearest region in one (object2) to the other (object1).

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#### Value

A data frame with one row corresponding for each row of object1, and columns:

dist bp separating the region in object1 and the nearest region in object2

matchIndex row index of object2 for the region that is closest to the region in object1

**type** "inside" if nearest region in object2 is wholly contained inside the region in object 1, "cover" if nearest region in object2 covers the whole region in object1, "disjoint" if there is no overlap between the region in object1 and the nearest region in object1, and "overlap" if the region in object1 and the nearest region in object2 overlap but one does not wholly cover the other.

amountOverlap amount of overlap between the region in object1 and the nearest region in object2

**insideDist** for type="inside" regions, the smaller (fewest bp) of 1., the end position of region in object1 - the end position of the nearest region in object2, and 2., the start position of region in object1 - the start position of the nearest region in object2.

size1 number of bp in region in object1

size2 number of bp in nearest region in object2

#### Author(s)

Martin Aryee <aryee@jhu.edu>, Peter Murakami, Rafael Irizarry

#### See Also

```
dmrFind, plotDMRs, plotRegions
```

### **Examples**

# See qval

| regionPlot | Plot user-supplied genomic regions using data returned by the dm- |
|------------|---|
|            | rFinder function.   |

### **Description**

Plot any given genomic regions from tiling microarray data using data returned by the dmrFinder function.

### Usage

```
regionPlot(tab, dmr, cpg.islands, Genome, outfile="./regions.pdf", which.plot=1:10, plot.these, c
```

### **Arguments**

tab a data frame with columns chr, start, and end identifying the regions to be plotted

from the data.

dmr a list object as returned by dmrFinder, providing the data to be plotted.

cpg.islands a table with columns "chr", "start", and "end" for CpG islands to plot in the

second panel.

regionPlot 35

| Genome      | the BSgenome object for the organism based upon which your array was designed.   |
|-------------|--|
| outfile     | a character string giving the name of the pdf file that will be saved. Include the full path if file is not to be saved in the current working directory.  |
| which.plot  | a vector of indices identifying which regions (rows) from tab to plot.   |
| plot.these  | if dmr results are from unpaired analysis, specify the groups to plot. Default is colnames(dmr\$gm). If dmr results are from paired analysis, specify the comparisons to plot. Default is colnames(dmr\$sMD).                |
| cl          | a vector of line and point colors, one for each element of plot.these in alphabetical order by group name.   |
| legend.size | cex argument for the legend (factor by which to magnify/shrink the legend).  |
| buffer      | An integer to control how many basepairs to show on either side of the plotted regions.  |
| plot.p      | set to FALSE if you want to plot the methlation values (the "l" output from dmrFinder) instead of the percentage methylation values (the "p" output). If dmrFinder was run on l instead of p, plot.p=FALSE necessarily.      |
| plotG       | if TRUE, a third panel of each DMR plot will show the difference between the median green channel value (after subtracting probe medians and correcting for gc content) between the 2 groups. If true, dat must be provided. |
| dat         | if plotG=TRUE, this must be provided. It is the TilingFeatureSet object used when estimating the matrix of percent methylation estimates used in dmrFinder when it produced dmr.   |
| grs         | if plotG=TRUE, this must be provided. It is the names of the two groups whose difference in G should be plotted in the 3rd panel.  |

### Details

This function enables plotting of any regions, not just DMRs, using the results of dmrFinder. The second panel shows the location of CpG's with ticks on the bottom (islands are colored) and the location of mcrbc recognition sites with ticks on the top.

### Author(s)

Martin Aryee <aryee@jhu.edu>, Peter Murakami, Rafael Irizarry

### See Also

plotRegions, dmrPlot, dmrFinder, dmrFdr

### **Examples**

# See dmrFdr

36 spatialAdjust

|--|

### Description

Remove spatial artifacts from microarray data stored in TilingFeatureSet objects

### Usage

```
spatialAdjust(dat, copy=TRUE, blockSize, theta = 1)
```

### **Arguments**

| dat       | TilingFeatureSet  |
|-----------|---|
| copy      | Only relevant when using disk-backed objects. If TRUE a copy will be made leaving the original object (dat) unchanged. The input object will not be preserved if copy=FALSE   |
| blockSize | The array is divided into a series of blockSize x blockSize rectangular blocks and the average signal level calculated for each. If blockSize is unspecified a size is chosen that gives about 1250 probes per block. |
| theta     | smoothing parameter   |

### **Details**

The array is divided into a set of blockSize x blockSize squares. A kernel smoother is then used to even out spatial artifacts.

### Value

a TilingFeatureSet

### Author(s)

Martin Aryee <aryee@jhu.edu>

### **Examples**

# See normalizeBetweenSamples

validatePd 37

| validatePd | Validate a sample description file for two-color microarray data |  |
|------------|--|--|
|            |  |  |

### **Description**

Checks a sample description file describing two-color arrays for proper formatting and if requested guesses column numbers for file names, sample labels and group labels.

#### Usage

```
validatePd(pd, fileNameColumn, sampleNameColumn, groupColumn,
ut = "_532.xys", md = "_635.xys")
```

### **Arguments**

pd A data frame containing the sample description table

fileNameColumn Number or name of column containing file names (optional)

sampleNameColumn Number or name of column containing sample names (optional)

groupColumn Number or name of column containing group labels (optional)

ut the file ending that designates untreated channel files

md the file ending that designates methyl-depleted channel files

#### **Details**

This function checks the formatting of a sample description file to make sure it has suitable columns for file names, sample names and (optionally) group labels. The sample description file should have one line per channel, i.e. two lines per sample corresponding to the red and green channel data files. Values in the sample name column are used to pair the two channels together. If fileNameColumn, sampleNameColumn and/or groupColumn are unspecified a guess will be made.

### Value

If the input data frame is valid: a list containing the fileNameColumn, sampleNameColumn and groupColumn. If the input data frame is invalid: FALSE

### Author(s)

Martin Aryee <aryee@jhu.edu>

### See Also

readCharm

```
# See dmrFdr
```

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