# Package 'meffil'

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```
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guess.batch.vars Guess which columns in sample sheet are batch variables

# Description

Guess which columns in sample sheet are batch variables

```
guess.batch.vars(norm.objects)
```

### **Arguments**

norm.objects Output from meffil.normalize.quantiles

#### Value

Array of variable names

```
meffil.add.cell.type.reference

Create a cell type reference object
```

# **Description**

Create a cell type reference object for estimating cell counts with the Infinium HumanMethylation450 BeadChip.

### Usage

```
meffil.add.cell.type.reference(
   name,
   M,
   U,
   cell.types,
   chip = NA,
   featureset = chip,
   number.sites = 50,
   specific.sites = NULL,
   number.quantiles = 500,
   subsets = NULL,
   object = NULL,
   verbose = F
)
```

# Arguments

```
Character string providing the name of the reference.
name
                 Matrix of methylated probe intensities (rows=CpG sites, columns=samples).
Μ
                 Matrix of unmethylatd probe intensities (rows=CpG sites, columns=samples).
U
cell.types
                 Vector of cell type names corresponding to sample basenames.
                 Name returned by meffil.list.chips() (Default: NA).
chip
                 Name returned by meffil.list.featuresets() (Default: chip).
featureset
number.sites Number of probes to characterise cell type methylation (Default: 50). For each
                 cell type, this number of probes with greater methylation than other cell types
                 and the same number with lesser methylation than the other cell types will be
                 included.
specific.sites
```

If not null (default), then number.sites is ignored and the supplied site identifiers are used to differentiate between cell types instead of those maximally different between the cell types within the reference.

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number.quantiles

Length of numeric sequence to specify probe intensity distributions (Default:

500).

object Cell type reference previously created by this function. If not NULL, then

this reference is added and all other function arguments are ignored (Default:

NULL).

verbose If TRUE, then status messages are printed during execution (Default: FALSE).

#### Value

A list specifying a cell type reference object that can be used by meffil.estimate.cell.counts() to estimate cell counts in another dataset. The object is a list containing: - beta The normalized methylation values of sites differentially methylated between cell types. - quantiles The average quantiles of methylated and unmethylated signals of probe sets defined by subsets (see below). e.g. quantiles[[name]] \$M provides the quantiles (number.quantiles quantiles) of the probes specified by subsets[[name]]. - subsets Probes on the microarray partitioned by relationship to CpG islands, either in an island, in a shore or far from an island.

meffil.add.chip

Add a new chip for analysis.

#### **Description**

Add a new chip for analysis.

# Usage

```
meffil.add.chip(name, manifest)
```

### **Arguments**

name Name of the new chip.

 ${\tt manifest} \qquad \quad A \ data \ frame \ obtained \ by \ loading \ the \ Illumina \ manifest \ into \ R.$ 

#### Value

Assuming that manifest contains a satisfactory set of columns, a new feature set and a new chip is made available. Thus, name will be added to the vectors returned by meffil.list.featuresets() and meffil.list.chips().

The manifest must contain the following columns:

- "IlmnID"character
- "Name"character
- "AddressA\_ID"character
- "AddressB ID"character
- "Infinium\_Design\_Type"values "I","II" or ""
- "CHR"values "1"-"22", "X" or "Y"
- "MAPINFO"integer
- "AlleleA\_ProbeSeq"character

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- UCSC\_RefGene\_Namecharacter
- UCSC\_RefGene\_Accessioncharacter
- UCSC\_RefGene\_Groupcharacter
- "UCSC\_CpG\_Islands\_Name"character
- "Relation\_to\_UCSC\_CpG\_Island"character
- "snp.exclude"logical

```
meffil.add.cnv.reference
```

Create a copy number reference object

# Description

Create a copy number reference object for estimating copy number variation with the Infinium HumanMethylation450 BeadChip.

# Usage

```
meffil.add.cnv.reference(
  name,
  M,
  U,
  chip = NA,
  featureset = chip,
  object = NULL,
  verbose = T
)
```

# **Arguments**

name	Character string providing the name of the reference.
M	Matrix of methylated probe intensities (rows=CpG sites, columns=samples).
U	Matrix of unmethylatd probe intensities (rows=CpG sites, columns=samples).
chip	Name returned by meffil.list.chips() (Default: NA).
featureset	Name returned by meffil.list.featuresets() (Default: chip).
object	A previously created copy number reference object created by this function. If not NULL, then this reference is added with the given name and all other function arguments are ignored (Default: NULL).
verbose	If TRUE, then status messages are printed during execution (Default: FALSE).

# Value

A list specifying a copy number reference object that can be used by meffil.calculate.cnv() to estimate copy number variation in another dataset.

meffil.add.copynumber450k.references

Create copy number references from CopyNumber450kData

#### **Description**

Two copy number references are created using data from the Bioconductor CopyNumber450kData R package. Reference "copynumber450k" is created using the "450k" feature set, and reference "copynumber450k-common" is created using the "common" feature set so it can be used with datasets with mixed 450K and EPIC chips.

### Usage

```
meffil.add.copynumber450k.references(verbose = T)
```

meffil.add.featureset

Add a feature set.

# **Description**

Add a feature set.

# Usage

```
meffil.add.featureset(name, features)
```

# **Arguments**

name Name of the new feature set.

features A data frame listing and describing all features.

#### Value

Assuming that features contains a satisfactory set of columns, a new feature set is made available. Thus, name will be added to the vector returned by meffil.list.featuresets().

The features data frame must contain the following columns:

- "name"character
- · "target"character
- "type"values "i", "ii" or "control"
- "chromosome"values "chr1"-"chr22", "chrX" or "chrY"
- "position"integer
- "gene.symbol"character,
- "gene.accession"character,
- "gene.region"character,
- "cpg.island.name"character
- · "relation.to.island"character
- "snp.exclude"logical

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```
meffil.basenames IDAT file basenames
```

# **Description**

List IDAT file basenames in a given directory.

# Usage

```
meffil.basenames(path, recursive = FALSE)
```

# **Arguments**

```
path Directory containing the IDAT files.

recursive If TRUE, search for IDAT files in subdirectories as well (Default: FALSE).
```

#### Value

Character vector of IDAT file basenames (i.e. filenames with "\_Grn.idat" and "\_Red.idat" removed). In other words, each identifies the Cy5 and Cy3 output files corresponding to a single microarray.

```
meffil.calculate.cnv

Calculate CNVs from IDAT files
```

# Description

Based on the algorithm developed in R/CopyNumber450k bioconductor package

# Usage

```
meffil.calculate.cnv(
   samplesheet,
   cnv.reference,
   chip = NA,
   verbose = FALSE,
   ...
)
```

# **Arguments**

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#### **Details**

```
The following default values are being used: - trim = 0.1 - min.width = 5 - nperm = 10000 - alpha = 0.001 - undo.splits = "sdundo" - undo.SD = 2
```

#### Value

Dataframe of segmented results

#### **Description**

Cell count estimates

# Usage

```
meffil.cell.count.estimates(qc.objects)
```

# **Arguments**

```
qc.objects List of objects obtained from meffil.qc() or meffil.create.qc.object().
```

# Description

Cell count estimate quality plot

#### Usage

```
meffil.cell.count.qc.plots(count.objects)
```

# **Arguments**

```
count.objects
```

A list of objects each obtained from meffil.estimate.cell.counts().

#### Value

Two ggplot2 boxplot objects: - betas Contains one box per sample or reference cell type representing the distribution of methylation levels for the CpG sites used to estimate cell counts. - counts Contains one box per reference cell type representing the distribution of cell count estimates across the samples.

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```
meffil.cell.type.specific.methylation

Reduce methylation profiles to most cell-type specific sites
```

# **Description**

Reduce methylation profiles to most cell-type specific sites

### Usage

```
meffil.cell.type.specific.methylation(
  beta,
  cell.types,
  number.sites = 50,
  verbose = F
)
```

# **Arguments**

```
beta Numeric matrix (values = 0..1; rows = CpG sites; columns = samples).

cell.types Name of cell type for each column of beta.

number.sites For each cell type, the number of sites less methylated and the number more methylated than other cell types to include in the reduced methylation profiles.
```

#### Value

Numeric matrix (values = 0.1; rows = CpG sites; columns = cell types) with number.sites CpG sites per cell type more methylated than other cell types and the same number less methylated. Values are the mean CpG site methylation levels of all original samples of the same cell type.#

```
meffil.cnv.matrix Create matrix of CNV values
```

### **Description**

Create matrix of CNV values

# Usage

```
meffil.cnv.matrix(cnv, featureset = "450k")
```

# Arguments

```
cnv     Output from meffil.calculate.cnv().
featureset     Name from meffil.list.featuresets() (Default: "450k").
```

### Value

Matrix of ncpg x nsample

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```
meffil.control.matrix
```

Infinium HumanMethylation450 BeadChip control matrix

# **Description**

Matrix containing control probe intensities from the Infinium HumanMethylation450 BeadChip.

# Usage

```
meffil.control.matrix(
   qc.objects,
   normalize = F,
   fixed.effects = NULL,
   random.effects = NULL)
```

# **Arguments**

```
qc.objects A list of outputs from meffil.create.qc.object().
```

normalize

If TRUE, then control matrix is scaled and specified fixed and random effects removed from the matrix. Otherwise, the raw control matrix is returned. (Default: FALSE).

fixed.effects

Names of columns in samplesheet that should be included as fixed effects along with control matrix principal components (Default: NULL).

random.effects

Names of columns in samplesheet that should be included as random effects (Default: NULL).

# Value

Matrix with one row per object consisting of control probe intensities and summaries.

# Description

Create a quality control object for a given Infinium HumanMethylation450 BeadChip.

#### **Usage**

```
meffil.create.qc.object(
   samplesheet.row,
   number.quantiles = 500,
   dye.intensity = 5000,
   verbose = F,
   detection.threshold = 0.01,
   bead.threshold = 3,
   sex.cutoff = -2,
   chip = NA,
   featureset = chip,
   cell.type.reference = NA
)
```

#### **Arguments**

```
samplesheet.row
                 Row from the data frame containing IDAT file and sample info (see meffil.read.samplesheet
                 or meffil.create.samplesheet).
number.quantiles
                 Number of quantiles to compute for probe subset (Default: 500).
dye.intensity
                 Reference intensity for scaling each color channel (Default: 5000).
verbose
                 If TRUE, then status messages are printed during execution (Default: FALSE).
detection.threshold
                 Default value = 0.01. All probes above this detection threshold detected.
bead.threshold
                 Default value = 3. All probes with less than this number of beads detected.
                 Sex prediction cutoff. Default value = -2.
sex.cutoff
                 Name returned by meffil.list.chips() (Default: NA).
chip
featureset
                 Name returned by meffil.list.featuresets() (Default: chip).
cell.type.reference
                 Character string name of the cell type reference to use for estimating cell counts.
```

# Value

List containing control probe information, probe summaries and quantiles. We call this a "QC object".

Estimates are not generated if set to NA (default). See meffil.list.cell.type.references for a list of available references. New references can be created using meffil.add.cell.type.references.

```
meffil.create.samplesheet

Create sample sheet if an Illumina one isn't available
```

# **Description**

If necessary generates two columns necessary for some functions: Sample\_Name and Sex

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#### Usage

```
meffil.create.samplesheet(
  path,
  basenames = meffil.basenames(path, recursive),
  recursive = FALSE,
  delim = "_"
)
```

# **Arguments**

delim Optional delim character to separate Sample\_Name into multiple columns.

Default: "\_"

#### Value

Sample sheet data frame

# **Description**

Design matrix derived by applying principal components analysis to control probes.

#### Usage

```
meffil.design.matrix(
   qc.objects,
   number.pcs,
   fixed.effects = NULL,
   random.effects = NULL)
```

# Arguments

Names of columns in samplesheet that should be included as random effects (Default: NULL).

### Value

 $Design \ matrix \ with \ one \ column \ for \ each \ of \ the \ first \ \verb|number.pcs| prinicipal \ components.$ 

```
meffil.estimate.cell.counts
```

Estimate cell counts from a reference

# **Description**

Estimate cell type ratios from methylation profiles of purified cell populations (Infinium Human-Methylation450 BeadChip) using the Houseman algorithm (PMID 22568884).

#### **Usage**

```
meffil.estimate.cell.counts(qc.object, cell.type.reference, verbose = T)
```

#### **Arguments**

object

```
Character string name of the cell type reference to use for estimating cell counts.

See meffil.list.cell.type.references() for a list of available references. New references can be created using meffil.add.cell.type.reference().

verbose

If TRUE, then status messages are printed during execution (Default: FALSE).
```

#### Value

A list: - counts Cell count estimates. - beta Normalized methylation levels of sites used to differentiate - reference Name of the cell type reference used. between reference cell types.

Results should be nearly identical to estimateCellCounts().

An object created by meffil.create.qc.object().

```
meffil.estimate.cell.counts.from.betas

Estimate cell counts for a beta matrix from a reference
```

### Description

Estimate cell type ratios from methylation profiles of purified cell populations (Infinium Human-Methylation450 BeadChip).

# Usage

```
meffil.estimate.cell.counts.from.betas(beta, cell.type.reference, verbose = F)
```

# Arguments

```
Matrix of Illumina 450K methylation levels (rows = CpG sites, columns = subjects).

cell.type.reference
Character string name of the cell type reference to use for estimating cell counts.
See meffil.list.cell.type.references() for a list of available references. New references can be created using meffil.add.cell.type.reference().

verbose

If TRUE, then status messages are printed during execution (Default: FALSE).
```

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# Value

A matrix of cell count estimates.

Results should be nearly identical to estimateCellCounts().

meffil.ewas

Epigenome-wide association study

# **Description**

Test association with each CpG site.

# Usage

```
meffil.ewas(
 beta,
  variable,
  covariates = NULL,
  batch = NULL,
  weights = NULL,
  cell.counts = NULL,
  isva = T,
  sva = T,
  smartsva = F,
  n.sv = NULL,
  isva0 = F,
  isva1 = F,
  winsorize.pct = 0.05,
  robust = TRUE,
  rlm = FALSE,
  outlier.iqr.factor = NA,
  most.variable = min(nrow(beta), 50000),
  featureset = NA,
  random.seed = 20161123,
  lmfit.safer = F,
  verbose = F
)
```

# **Arguments**

beta	Methylation levels matrix, one row per CpG site, one column per sample.
variable	Independent variable vector.
covariates	Covariates data frame to include in regression model, one row per sample, one column per covariate (Default: NULL).
batch	Batch vector to be included as a random effect (Default: NULL).
weights	Non-negative observation weights. Can be a numeric matrix of individual weights of same dimension as $beta$ , or a numeric vector of weights with length $ncol$ (beta), or a numeric vector of weights with length $nrow$ (beta).
cell.counts	Proportion of cell counts for one cell type in cases where the samples are mainly composed of two cell types (e.g. saliva) (Default: NULL).

Apply Independent Surrogate Variable Analysis (ISVA) to the methylation levels and include the resulting variables as covariates in a regression model (Default: TRUE).		
Apply Surrogate Variable Analysis (SVA) to the methylation levels and covariates and include the resulting variables as covariates in a regression model (Default: TRUE).		
Apply the SmartSVA algorithm to the methylation levels and include the resulting variables as covariates in a regression model (Default: FALSE).		
Number of surrogate variables to calculate (Default: NULL).		
t		
Apply all regression models to methylation levels winsorized to the given level. Set to NA to avoid winsorizing (Default: 0.05).		
Test associations with the 'robust' option when limma::eBayes is called (Default: TRUE).		
Test assocaitions with the 'robust' option when limma: lmFit is called (Default: FALSE).		
outlier.igr.factor		
For each CpG site, prior to fitting regression models, set methylation levels less than $Q1$ -outlier.iqr.factor * $IQR$ or more than $Q3$ + outlier.iqr.factor * $IQR$ to NA. Here $IQR$ is the inter-quartile range of the methylation levels at the CpG site, i.e. Q3-Q1. Set to NA to skip this step (Default: NA).		
most.variable		
Apply (Independent) Surrogate Variable Analysis to the given most variable CpG sites (Default: 50000).		
Name from meffil.list.featuresets() (Default: NA).		
Set to TRUE if status updates to be printed (Default: FALSE).		

```
meffil.ewas.bedgraph
```

Save EWAS effect estimates to bedgraph file

# Description

Saves EWAS effect estimates to a bedgraph file for viewing on a genome browser. More file format details can be found here: https://genome.ucsc.edu/goldenPath/help/bedgraph.html

```
meffil.ewas.bedgraph(
  ewas.object,
  filename,
  analysis,
  name,
  description,
  header
)
```

#### **Arguments**

ewas.object	Object returned by meffil.ewas().
filename	Filename for output, typically with a 'bed' extension.
analysis	The particular EWAS analysis from which to obtain summary statistics. This should be one of names (ewas.object $analyses$ ).
name	Text name to be included in the bedgraph header.
description	Text description to be included in the bedgraph header.
header	Bedgraph header. The default header uses the name and description provided.

```
meffil.ewas.covariate.associations
```

Describe associations between EWAS covariates and the variable of interest.

# **Description**

Describe associations between EWAS covariates and the variable of interest.

# Usage

```
meffil.ewas.covariate.associations(ewas.object)
```

# **Arguments**

```
ewas.object Output of meffil.ewas().
```

### Value

A data frame with one or more rows for each covariate.

If both the variable of interest and covariate are continuous or ordinal, then the covariate uses one row showing the name, mean and standard deviation of the covariate following the significance of the association between the covariate and the variable of interest.

If the covariate is categorical, then there is additionally one row for each level showing the mean and standard deviation of the variable of interest for samples at that covariate level.

If the variable of interest is categorical but the covariate is not, then there is one row for each variable level showing the mean and standard deviation of the covariate at the given level.

If both the variable of interest and covariate are categorical, then mean is replaced with the number of samples at each pair of variable/categorical levels and standard deviation with the percentage. P-values indicate the significance of association using Fisher's exact test.

```
meffil.ewas.cpg.plot
```

Scatter plots for a CpG site in an EWAS

# Description

Scatter plots for a CpG site in an EWAS

# Usage

```
meffil.ewas.cpg.plot(ewas.object, cpg, beta, title = cpg)
```

### **Arguments**

```
ewas.object Return object from meffil.ewas().

cpg CpG site to plot.

beta Matrix of methylation levels used to create the ewas.object.

title Title of the plot (Default: cpg).

ggplot object showing the scatterplots of DNA methylation vs the variable of interest in the EWAS. Each plot corresponds to a covariate set. Methylation levels are in fact residuals from fitting a model with DNA methylation and the covariates.
```

# Description

Manhattan plot for EWAS

# Usage

```
meffil.ewas.manhattan.plot(
  ewas.object,
  sig.threshold = 1e-07,
  title = "Manhattan plot"
)
```

# **Arguments**

### Value

```
ggplot showing the Manhattan plot.
```

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```
meffil.ewas.parameters

Specify parameters for QC
```

# **Description**

Specify parameters for QC

# Usage

```
meffil.ewas.parameters(
   sig.threshold = NA,
   max.plots = 10,
   model = "none",
   qq.inflation.method = "median"
)
```

# **Arguments**

sig.threshold

P-value threshold for significance (Default: NA). If NA, then threshold used will be 0.05 divided by the number of tests/probes.

max.plots

Maximum number of plots to generate (Default: 10).

model

Model to use for selecting associations: "none" (no covariates), "all" (all covariates), "isva" (independent surrogate variables), and "sva" (surrogate variables) (Default: "none").

qq.inflation.method

Method for calculating genomic inflation lambda. Valid values are "median", "regression" or "robust" (Default: "median").

#### Value

List of parameter values

```
meffil.ewas.qq.plot QQ \ plot \ for \ EWAS
```

# Description

QQ plot for EWAS

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#### Usage

```
meffil.ewas.qq.plot(
  ewas.object,
  sig.threshold = 1e-07,
  sig.color = "red",
  title = "QQ plot",
  xlab = bquote(-log[10]("expected p-values")),
  ylab = bquote(-log[10]("observed p-values")),
  lambda.method = "median"
)
```

# **Arguments**

### Value

List of ggplot for each analysis in ewas.object.

```
meffil.ewas.report Generate EWAS report.
```

# **Description**

Generate HTML file that summarises the EWAS.

```
meffil.ewas.report(
   ewas.summary,
   output.file = "ewas-report.html",
   author = "Analyst",
   study = "Illumina methylation data",
   ...
)
```

#### **Arguments**

```
meffil.ewas.sample.characteristics
```

Describe EWAS samples using the variable of interest and covariates.

### **Description**

Describe EWAS samples using the variable of interest and covariates.

#### Usage

```
meffil.ewas.sample.characteristics(ewas.object)
```

#### **Arguments**

```
ewas.object Output of meffil.ewas().
```

#### Value

A data frame with one row for each continuous or ordinal variable and one row for each level of each categorical variable. In the first case, each row provides the name, mean value and standard deviation of each variable. In the second case (categorical), each row provides the name of the variable level and the number of cases and percentage of cases at that level.

```
meffil.ewas.summary
```

Summarize EWAS results.

# Description

Generates variable and covariate summary tables, QQ plots, Manhattan plots, a list of associations, plots of the strongest associations and plots of selected CpG sites.

```
meffil.ewas.summary(
   ewas.object,
   beta,
   selected.cpg.sites = character(0),
   parameters = meffil.ewas.parameters(),
   verbose = T
)
```

#### **Arguments**

```
ewas.object From meffil.ewas().

beta Matrix of methylation levels used in the analysis.

selected.cpg.sites

Vector of CpG site names to plot (Default: character(0)).

parameters Default = meffil.ewas.parameters(). List of parameter values. See meffil.ewas.parameters()
```

### Value

List

```
meffil.extract.genotypes
```

Extract genotype data from PLINK .raw files for Illumina 450K SNPs

# Description

Extract genotype data from PLINK .raw files for Illumina 450K SNPs

### Usage

```
meffil.extract.genotypes(filenames, verbose = F)
```

# **Arguments**

filenames A vector of filenames of PLINK .raw files from which to extract genotype data.

### Value

Matrix with rows corresponding to SNPs, columns to samples and values equal to 0, 1 or 2 corresponding to genotypes.

# **Examples**

```
R> writeLines(meffil.snp.names("450k"), con="snp-names.txt")
shell> plink --bfile dataset --extract snp-names.txt --recodeA --out genotypes.raw --nowe
R> filenames <- "genotypes.raw"
R> genotypes <- meffil.extract.genotypes(filenames)</pre>
```

meffil.featureset 23

```
meffil.featureset Obtain a list of features in a feature set.
```

# Description

Obtain a list of features in a feature set.

# Usage

```
meffil.featureset(featureset = "450k")
```

# **Arguments**

```
featureset Name returned by meffil.list.featuresets() (Default: "450k").
```

# Value

A data frame with one row for each feature.

# **Examples**

```
x \leftarrow meffil.featureset("450k")
```

```
meffil.get.autosomal.sites
```

Get names of autosomal CpG sites in the feature set.

# Description

Get names of autosomal CpG sites in the feature set.

```
meffil.get.autosomal.sites(featureset = "450k")
```

24 meffil.get.features

meffil.get.beta

Infinium HumanMethylation450 BeadChip methylation levels

# **Description**

Compute beta values (methylation levels) from methylated/unmethylated signals

# Usage

```
meffil.get.beta(M, U, pseudo = 100)
```

# **Arguments**

Methylated signal matrix.

U Unmethylated signal matrix.

pseudo Value to add to the denominator to make the methylation estimate more stable.

#### Value

Matrix of 0..1 methylation level estimates. Equal to methylated/(methylated + unmethylated + pseudo).

```
meffil.get.features
```

Get a list of microarray features from a predefined feature set.

# Description

Get a list of microarray features from a predefined feature set.

# Usage

```
meffil.get.features(featureset = "450k")
```

# **Arguments**

```
featureset A name returned by meffil.list.featuresets() (Default: "450k").
```

# Value

A data frame listing all features in the feature set.

meffil.get.sites 25

```
meffil.get.sites Get names of all CpG sites in the feature set.
```

# **Description**

Get names of all CpG sites in the feature set.

# Usage

```
meffil.get.sites(featureset = "450k")
```

```
meffil.get.typeii.sites
```

Get names of CpG sites corresponding to Infinium Type II probes in the feature set.

# **Description**

Get names of CpG sites corresponding to Infinium Type II probes in the feature set.

# Usage

```
meffil.get.typeii.sites(featureset = "450k")
```

meffil.get.x.sites Get names of chromosome X CpG sites in the feature set.

# Description

Get names of chromosome X CpG sites in the feature set.

# Usage

```
meffil.get.x.sites(featureset = "450k")
```

meffil.get.y.sites *Get names of chromosome Y CpG sites in the feature set*.

### **Description**

Get names of chromosome Y CpG sites in the feature set.

```
meffil.get.y.sites(featureset = "450k")
```

26 meffil.list.cnv.references

# Description

List of available cell type references

# Usage

```
meffil.list.cell.type.references()
```

```
meffil.list.chips List of microarrays formats available.
```

# Description

By default, there is '450k' and 'epic'. Additions can be made using meffil.add.chip().

# Usage

```
meffil.list.chips()
```

```
{\tt meffil.list.cnv.references}
```

List of available copy number references

# Description

List of available copy number references

```
meffil.list.cnv.references()
```

meffil.list.featuresets 27

```
meffil.list.featuresets
```

List of feature sets available.

### **Description**

By default, there is '450k', 'epic' and 'common'. The 'common' feature set contains features in common to both the '450k' and 'epic' feature sets. This feature set can be used to handle datasets with mixed EPIC and HumanMethylation450 microarrays.

# Usage

```
meffil.list.featuresets()
```

#### **Details**

In most cases, a feature corresponds to the two probes from which it's value is derived. Each CpG represented on the chip for example corresponds to a single feature derived from a probe measuring methylated signal and a second probe measuring unmethylated signal.

Each control feature corresponds to a unique control probe.

```
meffil.load.controls
```

Load control probes

# **Description**

Load control probes

# Usage

```
meffil.load.controls(
   samplesheet,
   chip = NA,
   featureset = chip,
   verbose = F,
   ...
)
```

# **Arguments**

```
samplesheet Sample info (see meffil.read.samplesheet or meffil.create.samplesheet).

chip Name returned by meffil.list.chips() (Default: NA).

featureset Name returned by meffil.list.featuresets() (Default: chip).

verbose (Default: FALSE).

Arguments to mclapply.
```

28 meffil.load.raw.data

#### Value

List containing two elements: probes and values. The probes item is a data frame describing the control probes. The values item is a matrix providing the intensities of the control probes for each samples (rows=probes, columns=samples).

# **Description**

Load detection p-value matrix

# Usage

```
meffil.load.detection.pvalues(
    qc.objects,
    max.bytes = 2^30 - 1,
    verbose = F,
    ...
)
```

#### **Arguments**

```
qc.objects A list of outputs from meffil.create.qc.object().
verbose If TRUE, then detailed status messages are printed during execution (Default: FALSE).
... Arguments passed to mclapply().
```

### Value

Matrix of probe detection p-values.

```
meffil.load.raw.data

Load raw beta matrix
```

# Description

Load raw beta matrix

```
meffil.load.raw.data(
   qc.objects,
   pseudo = 100,
   just.beta = T,
   max.bytes = 2^30 - 1,
   verbose = F,
   ...
)
```

meffil.methylation.pcs 29

# **Arguments**

qc.objects	A list of outputs from meffil.create.qc.object().
pseudo	Value to add to the denominator to make the methylation estimate more stable when calculating methylation levels (Default: 100).
just.beta	If TRUE, then return just the methylation levels; otherwise, return the methylated and unmethylated matrices (Default: TRUE).
verbose	If ${\tt TRUE},$ then detailed status messages are printed during execution (Default: ${\tt FALSE}).$
	Arguments passed to mclapply ().

#### Value

If just.beta == TRUE, the matrix of methylation levels between between 0 and 1 equal to methylated signal/(methylated + unmethylated signal + pseudo). Otherwise, a list containing two matrices, the methylated and unmethylated signals.

```
meffil.methylation.pcs
```

Compute principal components of a methylation matrix.

# Description

Compute principal components of a methylation matrix.

# Usage

```
meffil.methylation.pcs(
  normalized.beta,
  probe.range = 5000,
  autosomal = T,
  verbose = F
)
```

# **Arguments**

```
normalized.beta
Output from meffil.normalize.samples(), either a matrix or a GDS filename.

probe.range Default = 5000. How many probes to be used in calculating PCs
autosomal Default = TRUE. If true, remove probes on sex chromosomes.

verbose=T Print progress messages?
```

#### Value

the principal components of normalized.beta.

# **Description**

Returns the most variable CpG sites (rows) in the methylation matrix.

# Usage

```
meffil.most.variable.cpgs(beta, n = 1000)
```

### **Arguments**

```
beta Methylation matrix (rows = CpG sites, columns = samples).

n Number of CpG sites to return.
```

#### Value

The n CpG site identifiers (rownames of x) with the greatest variance in x.

```
meffil.normalization.parameters

Specify parameters for testing normalization
```

# **Description**

Specify parameters for testing normalization

# Usage

```
meffil.normalization.parameters(
  norm.objects,
  variables = guess.batch.vars(norm.objects),
  control.pcs = 1:10,
  batch.pcs = 1:10,
  batch.threshold = 1e-50,
  colours = NULL
)
```

# **Arguments**

```
norm.objects Output from meffil.normalize.quantiles

variables Default = guess.batch.vars(norm). Which variables in sample sheet to test

control.pcs Default = 1:10. Number of control PCs to test against batch variables

colours Colours to use for scatterplots.
```

#### Value

List of parameters

```
meffil.normalization.parameters.from.betas

Specify parameters for testing normalization
```

# Description

Specify parameters for testing normalization

# Usage

```
meffil.normalization.parameters.from.betas(
  batch.pcs = 1:10,
  batch.threshold = 1e-50,
  colours = NULL
)
```

# **Arguments**

```
batch.pcs Default = 1:10. Number of PCs to test against batch variables batch.threshold

Default = 1e-50. Which pvalue threshold to show in table colours

Colours to use for scatterplots.
```

#### Value

List of parameters

```
meffil.normalization.report

Generate report on normalization performance
```

# **Description**

Generate HTML file that summarises the normalization.

```
meffil.normalization.report(
  normalization.summary,
  output.file = "normalization-report.md",
  author = "Analyst",
  study = "Illumina methylation data",
  ...
)
```

### **Arguments**

```
meffil.normalization.report.from.betas

Generate report on normalization performance
```

# **Description**

Generate HTML file that summarises the normalization.

# Usage

```
meffil.normalization.report.from.betas(
  normalization.summary,
  output.file = "normalization-report.md",
  author = "Analyst",
  study = "Illumina methylation data",
  ...
)
```

# **Arguments**

```
Output from meffil.normalization.summary.from.betas.

output.file Default = "meffil-normalization-report.html". If the file extension is not .htm, .html, .HTM or .HTML then output will be in markdown format.

author Default = "Analyst". Author name to be specified on report.

study Default = "Illumina methylation data". Study name to be specified on report.

Arguments to be passed to knitr::knit
```

```
meffil.normalization.summary
```

Perform tests to check normalization performance

#### **Description**

Creates scree plot of PCs of control probes, tests for association of control probe PCs with batch variables, tests for association of normalized probes with batch variables, creates PCA plots

#### Usage

```
meffil.normalization.summary(
  norm.objects,
  pcs,
  parameters = meffil.normalization.parameters(norm.objects),
  variables = NULL,
  verbose = TRUE
)
```

### **Arguments**

```
norm.objects Output from meffil.normalize.quantiles

pcs Output from meffil.methylation.pcs() applied to the normalized methylation matrix corresponding to norm.objects.

parameters Default = meffil.post.parameters(norm.objects). Report parameters.

variables Default = NULL. Data frame of variables to compare to principal components (pcs). Must contain length(norm.objects) rows. Columns that are not factors are ignored.

verbose Default = TRUE
```

# Value

List of tables and graphs.

```
meffil.normalization.summary.from.betas

*Perform tests to check normalization performance*
```

# **Description**

Perform tests to check normalization performance

```
meffil.normalization.summary.from.betas(
  pcs,
  parameters = meffil.normalization.parameters.from.betas(),
  samplesheet = samplesheet,
  variables = variables,
  verbose = TRUE
)
```

34 meffil.normalize.dataset

# **Arguments**

pcs	Output from ${\tt meffil.methylation.pcs}$ () applied to the normalized methylation matrix
parameters	$Default = meffil.normalization.parameters.from.betas(). \ Report\ parameters.$
samplesheet	Default = NULL. Data frame of variables to compare to principal components (pcs). Must contain nrow (pcs) == nrow (samplesheet) rows. Columns that are not factors are ignored.
variables	Which variables in sample sheet to test
verbose	Default = TRUE

#### Value

List of tables and graphs.

```
meffil.normalize.dataset

Functional normalization
```

#### **Description**

Apply functional normalization to a set of Infinium HumanMethylation450 BeadChip IDAT files.

```
meffil.normalize.dataset(
  samplesheet,
  number.quantiles = 500,
  detection.threshold = 0.01,
 bead.threshold = 3,
  sex.cutoff = -2,
  chip = NA,
  featureset = chip,
  cell.type.reference = NA,
  qc.parameters = meffil.qc.parameters(),
  qc.file = "meffil-qc-report.md",
  author = "Analyst",
  study = "IlluminaHuman450 data",
  number.pcs = 2,
  fixed.effects = NULL,
  random.effects = NULL,
  pseudo = 100,
  just.beta = T,
  gds.filename = NULL,
  probe.range = 5000,
  autosomal = T,
  norm.parameters = NULL,
  norm.file = "meffil-normalization-report.md",
  verbose = FALSE
)
```

meffil.normalize.dataset 35

#### **Arguments**

```
Output from meffil.read.samplesheet() or meffil.create.samplesheet().
samplesheet
                Arguments to meffil.qc():
cell.type.reference
                Argument to meffil.qc.summary():
qc.parameters
                (parameters)
                Arguments to meffil.qc.report():
qc.file
                (output.file)
study
                Arguments to meffil.normalize.quantiles():
fixed.effects
                Names of columns in samplesheet that should be included as fixed effects along
                with control matrix principal components (Default: NULL).
random.effects
                Names of columns in samplesheet that should be included as random effects
                (Default: NULL).
                Arguments to meffil.normalize.samples():
                Arguments to meffil.methylation.pcs().
pseudo
gds.filename If not NULL (default), then saves the output to a GDS (Genomic Data Structure).
                This is for cases where the output is too large to fit into main memory. The GDS
                option assumes that argument just.beta == TRUE.
probe.range
                (Default: 5000).
                (Default: TRUE).
autosomal
                Arguments to meffil.normalization.summary():
norm.parameters
                (parameters)
norm.file
                (output.file)
                Other:
                If TRUE, then status messages are printed during execution (Default: FALSE).
verbose
                (Default: 1:10).
npcs
```

# Details

Fortin JP, Labbe A, Lemire M, Zanke BW, Hudson TJ, Fertig EJ, Greenwood CM, Hansen KD. Functional normalization of 450k methylation array data improves replication in large cancer studies. Genome Biol. 2014 Dec 3;15(12):503. doi: 10.1186/s13059-014-0503-2. PMID: 25599564

# Value

```
A list: -qc.summary meffil.qc.summary() output. -norm meffil.normalize.quantiles() output. - beta Normalized beta matrix (methylation levels). -norm.summary meffil.normalization.summary( output.
```

```
meffil.normalize.quantiles

Normalize microarray quantiles
```

# **Description**

Normalize microarray quantiles using controls extracted (Infinium HumanMethylation450 Bead-Chip).

# Usage

```
meffil.normalize.quantiles(
   qc.objects,
   number.pcs = 2,
   fixed.effects = NULL,
   random.effects = NULL,
   verbose = F
)
```

#### Arguments

```
qc.objects A list of outputs from meffil.create.qc.object().

number.pcs Number of control matrix principal components to adjust for (Default: 2).

fixed.effects

Names of columns in samplesheet that should be included as fixed effects along with control matrix principal components (Default: NULL).

random.effects

Names of columns in samplesheet that should be included as random effects (Default: NULL).

verbose

If TRUE, then status messages are printed during execution (Default: FALSE).
```

#### Value

Same list as input with additional elements added for each sample including normalized quantiles needed for normalizing each sample.

```
meffil.normalize.sample

Normalize Infinium HumanMethylation450 BeadChip
```

# **Description**

Normalize sample methylation data using normalized quantiles.

```
meffil.normalize.sample(norm.object, verbose = F)
```

#### **Arguments**

```
norm.object An element of meffil.normalize.quantiles().

verbose If TRUE, then status messages are printed during execution (Default: FALSE).
```

#### Value

List containing normalized methylated and unmethylated signals.

```
meffil.normalize.samples

Normalize Infinium HumanMethylation450 BeadChips
```

# **Description**

Normalize a set of samples using their normalized quality control objects.

# Usage

```
meffil.normalize.samples(
  norm.objects,
  pseudo = 100,
  just.beta = T,
  cpglist.remove = NULL,
  max.bytes = 2^30 - 1,
  gds.filename = NULL,
  verbose = F,
  ...
)
```

# Arguments

```
norm.objects The list or sublist of meffil.normalize.quantiles().
                 Value to add to the denominator to make the methylation estimate more stable
pseudo
                 when calculating methylation levels (Default: 100).
                 If TRUE, then return just the normalized methylation levels; otherwise, return
just.beta
                 the normalized methylated and unmethylated matrices (Default: TRUE).
cpglist.remove
                 Optional list of CpGs to exclude from final output
gds.filename If not NULL (default), then saves the output to a GDS (Genomic Data Structure).
                 This is for cases where the output is too large to fit into main memory. The GDS
                 option assumes that argument just.beta == TRUE.
                 If TRUE, then detailed status messages are printed during execution (Default:
verbose
                 FALSE).
                 Arguments passed to mclapply ().
. . .
```

#### Value

If <code>just.beta</code> == TRUE, the normalized matrix of methylation levels between between 0 and 1 equal to methylated signal/(methylated + unmethylated signal + pseudo). Otherwise, a list containing two matrices, the normalized methylated and unmethylated signals. If <code>gds.filename</code> is not <code>NULL</code>, then the output is saved to the GDS file rather than retained in memory and returned to the caller. The library 'gdsfmt' must be installed.

meffil.pcs

Calculate control probe PCs

### **Description**

Calculate control probe PCs

# Usage

```
meffil.pcs(qc.objects, fixed.effects = NULL, random.effects = NULL)
```

# **Arguments**

```
{\tt qc.objects} A list of outputs from meffil.create.qc.object(). fixed.effects
```

Names of columns in samplesheet that should be included as fixed effects along with control matrix principal components (Default: NULL).

random.effects

Names of columns in samplesheet that should be included as random effects (Default: NULL).

# Value

PCA of control probes

```
meffil.plot.beadnum.cpgs
```

Manhattan plot of number of beads by probe - percentage of probes with beads < 3 for each sample

# Description

Manhattan plot of number of beads by probe - percentage of probes with beads < 3 for each sample

# Usage

```
meffil.plot.beadnum.cpgs(qc.objects, threshold = 0.05)
```

# **Arguments**

```
qc.objects From meffil.qc
```

threshold Cut off value for proportion of samples with poor detection p values. Default 0.05.

### Value

Data frame of results plus plot

```
meffil.plot.beadnum.samples

Plot number of beads per sample
```

### **Description**

Plot number of beads per sample

### Usage

```
meffil.plot.beadnum.samples(qc.objects, threshold = 0.05, colour.code = NULL)
```

# **Arguments**

```
qc.objects From meffil.qc

threshold Cut off value for proportion of CpGs with low bead numbers. Default 0.05

colour.code Array of length n samples to colour code points. Defaults to NULL
```

#### Value

Data frame of results plus plot

```
meffil.plot.cell.counts

Cell count estimate quality plot
```

# **Description**

Cell count estimate quality plot

# Usage

```
meffil.plot.cell.counts(qc.objects)
```

### **Arguments**

```
qc.objects Output from meffil.qc().
reference Object describing methylation profiles of purified cell populations obtained from
    meffil.add.cell.type.reference().
```

### Value

Two ggplot2 boxplot objects: - betas Contains one box per sample or reference cell type representing the distribution of methylation levels for the CpG sites used to estimate cell counts. - counts Contains one box per reference cell type representing the distribution of cell count estimates across the samples.

```
meffil.plot.control.batch
```

Test for association of control matrix probes with known batch variables

# **Description**

Performs association of each of n PCs calculated from the control matrix against each of m measured batch variables

# Usage

```
meffil.plot.control.batch(
  norm.objects,
  npcs = 1:10,
  variables = guess.batch.vars(norm.objects),
  additional = NULL,
  batch.threshold = 1e-50,
  cols = NULL,
  verbose = TRUE
)
```

# **Arguments**

```
norm.objects From meffil.normalize.quantiles

pcs Which PCs to plot. Default first 10

variables. Default = guess.batch.vars(norm.objects). Array spacifying column names in samplesheet to test for association with control matrix PCs.

additional. Default = NULL. Data frame containing variables to test for association with control matrix PCs. Must have nrow (additional) == length(norm.objects).

verbose=T Print progress messages?
```

#### Value

Data frame of results plus plot

```
meffil.plot.control.scree

Plot scree plot of control matrix
```

# **Description**

Plot scree plot of control matrix

```
meffil.plot.control.scree(norm.objects)
```

### **Arguments**

```
{\tt norm.objects} {\tt From} {\tt meffil.normalize.quantiles}
```

### Value

Data frame of results plus plot

```
meffil.plot.controlmeans
```

Plot the means of control probes for each sample and for each control probe type

# **Description**

Plot the means of control probes for each sample and for each control probe type

# Usage

```
meffil.plot.controlmeans(
   qc.objects,
   control.categories = NULL,
   colour.code = NULL,
   outlier.sd = 5
)
```

# Arguments

# Value

Data frame of results plus plot

# Description

Manhattan plot of detection pval per probe - percentage with pvalue < 0.01

```
meffil.plot.detectionp.cpgs(qc.objects, threshold = 0.05)
```

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# **Arguments**

```
qc.objects From meffil.qc threshold Cut off value for proportion of samples with poor detection p values. Default 0.05.
```

#### Value

Data frame of results plus plot

```
meffil.plot.detectionp.samples

Plot detection p values from idat files
```

# **Description**

Plot detection p values from idat files

# Usage

```
meffil.plot.detectionp.samples(
   qc.objects,
   threshold = 0.05,
   colour.code = NULL
)
```

# **Arguments**

```
qc.objects From meffil.qc

threshold Cut off value for proportion of CpGs with poor detection p values. Default 0.05

colour.code Array of length n samples to colour code points. Defaults to NULL
```

# Value

Data frame of results plus plot

```
meffil.plot.genotypes

Plot SNP beta and sample genotype concordances
```

# **Description**

Plot SNP beta and sample genotype concordances

```
meffil.plot.genotypes(
   qc.objects,
   genotypes = NULL,
   sample.threshold = 0.9,
   snp.threshold = 0.99
)
```

#### **Arguments**

```
qc.objects Output from meffil.qc().

genotypes Optional output from meffil.extract.genotypes(). Sample genotypes
are matched to sample qc.objects using colnames (genotypes) and names (qc.objects).

sample.threshold
Concordance threshold below which the Illumina 450K and genetic profiles for
a sample are deemed a mismatch (Default: 0.9).

snp.threshold
Concordance threshold below which the Illumina 450K and genetic profiles for
a SNP are deemed a mismatch (Default: 0.99).
```

#### Value

A list consisting of: -graphs A list of ggplot2 objects. The first snp.betas plots the beta distributions of each SNP probe in the microarray. The second and third plots are added only if the genotypes parameter is not NULL. The second plot shows the distribution of SNP concordances, and the third plot shows the distribution of sample concordances. -tabs Contains two data frames if the genotypes parameter is not NULL. The first samples lists the concordances of each sample, the second snps lists the concordances of each SNP.

```
meffil.plot.meth.unmeth
```

Plot average methylated vs unmethylated levels for each individuals

# **Description**

plot raw control probes and fit linear regression, remove samples that have sd(y - yhat) > mean\*3

### Usage

```
meffil.plot.meth.unmeth(qc.objects, outlier.sd = 3, colour.code = NULL)
```

### **Arguments**

```
qc.objects From meffil.qc
outlier.sd Cut off for declaring outliers. Default = 3
colour.code Array of length n samples to colour code points. Defaults to NULL
```

### Value

Data frame of results plus plot

```
meffil.plot.pc.fit Number of control matrix principal components
```

# **Description**

Fits probe intensities to principal components of the microarray control matrix and calculates the resulting mean squared residuals for different numbers of principal components.

# Usage

```
meffil.plot.pc.fit(
   qc.objects,
   fixed.effects = NULL,
   random.effects = NULL,
   n.cross = 10,
   name = "autosomal.ii"
)
```

# **Arguments**

```
qc.objects A list of outputs from meffil.create.qc.object().

fixed.effects

Names of columns in samplesheet that should be included as fixed effects along with control matrix principal components (Default: NULL).

random.effects

Names of columns in samplesheet that should be included as random effects (Default: NULL).
```

number.pcs Number of principal components to include in the design matrix (Default: all).

# Value

A list containing a data frame with the mean squared residuals for different numbers of principal components and a plot of these residuals.

```
meffil.plot.probe.batch
```

Test normalized betas for association with known batch variables

# **Description**

Performs association of each of n PCs calculated from most variable CpG sites (after normalization) against each of m measured batch variables

#### Usage

```
meffil.plot.probe.batch(
  norm.objects,
  pcs,
  variables = guess.batch.vars(norm.objects),
  additional = NULL,
  batch.threshold = 1e-50,
  cols = NULL,
  verbose = T
)
```

### **Arguments**

```
norm.objects Output from meffil.normalize.quantiles().

pcs Output from meffil.methylation.pcs() applied to the normalized methylation matrix corresponding to norm.objects.

variables Default = guess.batch.vars(norm). Which variables in sample sheet to test

additional. Default = NULL. Data frame containing variables to test for association with control matrix PCs. Must have nrow (additional) == length (norm.objects).

verbose=T Print progress messages?
```

### Value

List of table of results and graph

```
meffil.plot.probe.batch.from.betas

Test normalized betas for association with known batch variables
```

# Description

Performs association of each of n PCs calculated from most variable CpG sites (after normalization) against each of m measured batch variables

```
meffil.plot.probe.batch.from.betas(
   samplesheet,
   variables,
   pcs,
   batch.threshold = batch.threshold,
   cols = NULL,
   verbose = T
)
```

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#### **Arguments**

variables Which variables in sample sheet to test

pcs Output from meffil.methylation.pcs() applied to the normalized methy-

lation matrix

samplesheet. Data frame containing variables to test for association with PCs. Must have

nrow(pcs) == nrow(samplesheet).

verbose=T Print progress messages?

### Value

List of table of results and graph

### **Description**

Plot predicted sex

### Usage

```
meffil.plot.sex(qc.objects, outlier.sd = 3)
```

### **Arguments**

```
qc.objects From meffil.qc
```

#### Value

Data frame of results plus plot

```
meffil.probe.info Obtain a list of probes for a given feature set (chip).
```

# **Description**

Obtain a list of probes for a given feature set (chip).

### Usage

```
meffil.probe.info(chip = "450k", featureset = chip)
```

# Arguments

```
chip Name returned by meffil.list.chips() (Default: "450k").

featureset Name returned by meffil.list.featuresets() (Default: chip).
```

### Value

A data frame with one row per probe. The full set of probes for a chip is returned if chip == featureset; otherwise, the probes are restricted to those corresponding to features in the feature set.

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meffil.qc

Perform QC on HumanMethylation450 idat files

# Description

Read in control matrices for each sample. Perform background correction and R/G dye bias correction. Predict sex

# Usage

```
meffil.qc(
   samplesheet,
   number.quantiles = 500,
   dye.intensity = 5000,
   detection.threshold = 0.01,
   bead.threshold = 3,
   sex.cutoff = -2,
   chip = NA,
   featureset = chip,
   cell.type.reference = NA,
   max.bytes = 2^30 - 1,
   verbose = F,
   ...
)
```

# **Arguments**

```
samplesheet
                 Data frame containing IDAT file and sample info (see meffil.read.samplesheet
                 pr meffil.create.samplesheet).
number.quantiles
                 Number of quantiles to compute for probe subset (Default: 500).
dye.intensity
                 Reference intensity for scaling each color channel (Default: 5000).
detection.threshold
                 Default value = 0.01. All probes above this detection threshold detected.
bead.threshold
                 Default value = 3. All probes with less than this number of beads detected.
                 Sex prediction cutoff. Default value = -2.
sex.cutoff
                 Name returned by meffil.list.chips() (Default: NA).
chip
featureset
                 Name returned by meffil.list.featuresets() (Default: chip).
cell.type.reference
                 Character string name of the cell type reference to use for estimating cell counts.
                 Estimates are not generated if set to NA (default). See meffil.list.cell.type.references
                 for a list of available references. New references can be created using meffil.add.cell.type.
verbose
                 If TRUE, then status messages are printed during execution (Default: FALSE).
```

# Value

List containing control probe information, probe summaries and quantiles.

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```
meffil.qc.parameters
```

Specify parameters for QC

# **Description**

Specify parameters for QC

### Usage

```
meffil.qc.parameters(
  colour.code = NULL,
  control.categories = NULL,
  sex.outlier.sd = 3,
  meth.unmeth.outlier.sd = 3,
  control.means.outlier.sd = 5,
  detectionp.samples.threshold = 0.2,
  beadnum.samples.threshold = 0.2,
  detectionp.cpgs.threshold = 0.2,
  beadnum.cpgs.threshold = 0.2,
  snp.concordance.threshold = 0.9,
  sample.genotype.concordance.threshold = 0.9
```

# **Arguments**

Sets the standard deviation multiple at which sex outliers are identified. Default value = 3.

meth.unmeth.outlier.sd

Sets the standard deviation multiple at which methylated/unmethylated signal outliers are identified. Default value = 3.

control.means.outlier.sd

Sets the standard deviation multiple at which control probe signals are identified as outliers. Default value = 5 <what param does>

detectionp.samples.threshold

Detection p-value threshold. Probes with values above this are considered undetected. Default value = 0.05

beadnum.samples.threshold

A sample is excluded if the given proportion of probes has low bead number. Default value = 0.05

detectionp.cpgs.threshold

A sample is excluded if the given proporition of probes are undetected. Default value = 0.05

beadnum.cpgs.threshold

A probe is excluded if the given proportion of samples have low bead number. Default value = 0.05

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```
snp.concordance.threshold
```

Minimum required concordance between supplied genotypes and genotypes estimated from a SNP probe. Default value = 0.99

```
sample.genotype.concordance.threshold
```

Minimum required concordance between supplied genotypes and genotypes estimated from SNP probes for a given individual. Default value = 0.9 <what param does>

### Value

List of parameter values

```
meffil.qc.report Generate QC report
```

# **Description**

Generate HTML file that summarises the QC.

# Usage

```
meffil.qc.report(
   qc.summary,
   output.file = "qc-report.html",
   author = "Analyst",
   study = "Illumina methylation data",
   ...
)
```

#### **Arguments**

```
qc.summary Output from meffil.qc.summary.
output.file Default = "meffil-qc-report.html". If the file extension is not .htm, .html, .HTM
or .HTML then output will be in markdown format.
author Default = "Analyst". Author name to be specified on report.
study Default = "Illumina methylation data". Study name to be specified on report.
... Arguments to be passed to knitr::knit
```

```
meffil.qc.summary Perform QC analysis on idat files
```

# **Description**

Performs a number of QC analyses including checking for sex differences, methylated vs unmethylated levels, deviation from control probe means, detection p-values and bead numbers per sample and probe.

#### Usage

```
meffil.qc.summary(
   qc.objects,
   genotypes = NULL,
   parameters = meffil.qc.parameters(),
   verbose = TRUE
)
```

### **Arguments**

```
qc.objects From meffil.qc
genotypes Optional output from meffil.extract.genotypes(). Sample genotypes
are matched to sample qc.objects using colnames (genotypes) and names (qc.objects).
parameters Default = meffil.qc.parameters(). List of parameter values. See meffil.qc.parameters
```

#### **Details**

Also returns list of sample IDs and CPGs that are low quality.

#### Value

List

```
meffil.read.samplesheet

Function to read Illumina "Sample Sheet" adapted from read.450k.sheet in R/minfi
```

# Description

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

#### Usage

```
meffil.read.samplesheet(
  base,
  pattern = "csv$",
  ignore.case = TRUE,
  recursive = TRUE,
  verbose = TRUE
```

# **Arguments**

```
The base directory from which the search is started.

pattern = "csv$" What pattern is used to identify a sample sheet file, see list.files
ignore.case = TRUE Should the file search be case sensitive?

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

verbose = TRUE Should the function be verbose?

Dutput from meffil.basenames
```

#### **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.

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.

#### 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.

```
meffil.remove.samples
```

Remove samples from QC objects

# Description

Remove samples from QC objects

### Usage

```
meffil.remove.samples(qc.objects, sample.ids)
```

# **Arguments**

```
qc.objects Output from meffil.qc
sample.ids Array of sample.name IDs to be removed
```

### Value

qc.objects with samples removed

```
meffil.snp.betas Matrix of SNP 'beta' values
```

# Description

Matrix of SNP 'beta' values

# Usage

```
meffil.snp.betas(qc.objects)
```

# Arguments

```
qc.objects List of objects obtained from meffil.qc() or meffil.create.qc.object().
```

```
meffil.snp.concordance
```

Concordance between genotypes and SNP betas

# **Description**

genotypes <- meffil.extract.genotypes(raw.filenames) snp.betas <- meffil.snp.betas(qc.objects) meffil.snp.concordance(snp.betas, genotypes[rownames(snp.betas),colnames(snp.betas)])

# Usage

```
meffil.snp.concordance(
   snp.betas,
   genotypes,
   snp.threshold = 0.99,
   sample.threshold = 0.9
```

### Value

Returns a list of two vectors: - one providing concordances between genotypes and SNP betas for matched samples, - a second providing concordances between genotypes and SNP betas for matched SNPs. as well as the genotype matrix derived from 'snp.betas'.

meffil.snp.names 53

meffil.snp.names Obtain the list of identifiers for the SNPs on the microarray.

# **Description**

Obtain the list of identifiers for the SNPs on the microarray.

# Usage

```
meffil.snp.names(featureset = "450k")
```

### **Arguments**

```
featureset Name from meffil.list.featuresets() (Default: "450k").
```

```
meffil.summarize.relationship
```

Describe the relationship between two variables.

# **Description**

Describe the relationship between two variables.

# Usage

```
meffil.summarize.relationship(vars)
```

# Arguments

vars

A data frame with at least two columns. The first two columns will be compared.

#### Value

A list whose elements depends on the types of the two variables. In each case, the list contains the following elements:

**var1** Name of the first variable, i.e. colnames(vars)[1].

var2 Name of the second variable.

- **r** Spearman's correlation between the variables. This may be meaningless if one variable is an unordered factor.
- **r.p** P-value corresponding to the correlation between the variables.

output The contents of the list formatted to be printed as markdown text.

**plot** A plot (ggplot2) visualizing the relationship.

If both variables are factors, then the list will include a frequency table (freq) and a corresponding matrix of p-values (p.values) obtained using Fisher's test to test for enrichment in each cell of the frequency table.

If one variable is a factor and the other numeric, then list will include the F-statistic (F.stat) and p-value (p.value) from one-way analysis of variance. There will also be a data frame (cases) with each row providing statistics from a t-test comparing the numeric variable within and without each level of the factor variable.

If both variables are numeric, then the list will include the F-statistic (F . stat) and p-value (p . value) from the linear model fitting the variables.

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