Package 'methyltools'

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methyltools-package

Additional tools for the analysis of methylation data

Description

Additional tools for the analysis of methylation data

Details

Package: methyltools Type: Package Version: 0.1

Date: 2013-03-15 License: GPL (>= 2)

Author(s)

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References

+ add reference related to lumi and minfi

Staaf, Johan and Vallon-Christersson, Johan and Lindgren, David and Juliusson, Gunnar and Rosenquist, Richard and Hoglund, Mattias and Borg, Ake and Ringner, Markus (2008) Normalization of Illumina Infinium whole-genome SNP data improves copy number estimates and allelic intensity ratios, BMC Bioinformatics, 9:409

Louhimo, R. and Hautaniemi, S. (2011) CNAmet: an R package for integration of copy number, expression and methylation data Bioinformatics 27(6):887-888.

auc

Area Under Curves

Description

This function compute the area under curves.

Usage

auc(x,y)

Arguments

x a numeric vector y a numeric vector beta3mix 3

Value

The function returns a numeric value corresponding to the area under the curves.

Examples

```
# y <- rpois(1:10)
# d1 <- density(y)
# auc(d1$x,d1$y)</pre>
```

beta3mix

set of functions related to beta3mix object

Description

set of functions associated with the beta3mix objects, including print, simulate and graphical representation.

Usage

```
beta3mix(x,n=10000,initial=c(0.2,0.75),niter=25,tol=1e-4,...)
beta3mix.light(x,n=10000,initial=c(0.2,0.75),niter=25,tol=1e-4,...)
## S3 method for class 'beta3mix'
print(x,...)
## S3 method for class 'beta3mix'
plot(x,data,nsim=10000,col=c("black","green2"),...)
## S3 method for class 'beta3mix'
predict(object,data,option="postcluster",encoding=NULL,...)
## S3 method for class 'beta3mix'
simulate(object,nsim=1,seed=NULL,...)
```

Arguments

X	a numeric vector or object 'beta3mix'
object	an object 'beta3mix'
data	a numeric vector
n	a numeric value corresponding to the number of values used to fit the model (10000 by default)
nsim	a numeric value corresponding to the number of simulated values
initial	initial condition used to defined a priori classification
niter	a numeric value corresponding to the maximum number of iteration
tol	a numeric value corresponding to the tolerance criteria
cpus	a numeric value providing the number of cpu used (by defualt cpus=4)
option	a character used to defined the type of predicted values (by default, type="postcluster", see details)
encoding	a character used to defined the type of encoding for the predicted values ("123" or "UHM", see details)
seed	a numeric value corresponding to the argument of the function set.seed (by default, seed=NULL).
	further arguments passed to or from other methods

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Details

The function 'beta3mix' uses a modified version of the function 'blc' from R packages (RPMM). The function 'predict' proposes three different types of prediction values: meancluster: classification based on averaged cut-off; postcluster: classification based on posterior probabilities; postproba: posterior probability. The argument call 'encoding' offer the possibilities to replace the default encoding ('c(1,2,3)') by an other (e.g. 'c("U","H","M")').

Value

The function 'beta3mix' returns a list of of classes "beta3mix". The list contains parameters and objects representing mixture model fit, including posterior weights and log-likelihood from the function 'blc' from R package RPMM. This list is completed by element related to the classification of the values in the three categories described below:

idx identification of the values used to fit the model subcluster classification of the values used to fit the model

limit values used to defined 'meancluster' classification

meancluster a vector containing the classification of the values based on averaged cut-off (see

RPMM package)

nclass number of class

postw a matric containing posterior probabilities

postcluster a vector containing the classification of the values based on maximal posterior

probabilities

call the call function

References

Teschendorff AE, Marabita F, Lechner M, Bartlett T, Tegner J, Gomez-Cabrero D, Beck S. A Beta-Mixture Quantile Normalisation method for correcting probe design bias in Illumina Infinium 450k DNA methylation data. Bioinformatics. 2012 Nov 21. Houseman AE and Sc.D. (2012). RPMM: Recursively Partitioned Mixture Model. R package version 1.10. http://CRAN.R-project.org/package=RPMM Koestler DC, Christensen BC, Marsit CJ, Kelsey KT, Houseman EA. (2013) Recursively partitioned mixture model clustering of DNA methylation data using biologically informed correlation structures. Stat Appl Genet Mol Biol. 2013 Mar 5;12(2):225-40.

Examples

+ add example

betaUHM DNA methylation state or Genoptyping based on 3-beta mixture model

Description

Classification of Beta-values in using 3-beta mixture distribution and EM algorithm

Usage

beta UHM (df, n=10000, initial=c(0.2, 0.75), niter=25, tol=1e-4, cpus=4, type="postcluster", moda=c("U", "House of the content of the conte

betaUHM 5

Arguments

df	a sdata.frame containing the beta-value
n	a numeric value corresponding to the number of values used to fit the model (10000 by default)
initial	initial condition used to defined a priori classification
niter	a numeric value corresponding to the maximum number of iteration
tol	a numeric value corresponding to the tolerance criteria
cpus	a numeric value providing the number of cpu used (by defualt cpus=4)
type	a character used to defined the type of output (by default, type="postcluster", see details)
moda	a character used to defined the encoding of genotype(by default, moda= $c("U","H","M")$, see details)
	further arguments passed to or from other methods

Details

For the classification of the values, three different types of prediction values are available: meancluster: classification based on averaged cut-off postcluster: classification based on posterior probabilities postproba: posterior probability

Value

return a numeric

References

Teschendorff AE, Marabita F, Lechner M, Bartlett T, Tegner J, Gomez-Cabrero D, Beck S. A Beta-Mixture Quantile Normalisation method for correcting probe design bias in Illumina Infinium 450k DNA methylation data. Bioinformatics. 2012 Nov 21. Houseman AE and Sc.D. (2012). RPMM: Recursively Partitioned Mixture Model. R package version 1.10. http://CRAN.R-project.org/package=RPMM Koestler DC, Christensen BC, Marsit CJ, Kelsey KT, Houseman EA. (2013) Recursively partitioned mixture model clustering of DNA methylation data using biologically informed correlation structures. Stat Appl Genet Mol Biol. 2013 Mar 5;12(2):225-40.

Examples

```
# data(BetaValues)
## three DNA methylation states
# w <- betaUHM(BetaValues,moda=c("U","H","M"))
## deux DNA methylation states
# w <- betaUHM(BetaValues,moda=c("U","M","M"))
## genotyping with A and B alleles
# w <- betaUHM(BetaValues,moda=c("A/A","A/B","B/B"))
## genotyping with numerical encoding
# w <- betaUHM(BetaValues,moda=c(0,1,2))</pre>
```

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cghGene

Extract the CGH information related to a given gene

Description

The function extracts to object from cghCall class the CGH information for a given gene

Usage

```
cghGene(gene, calls, ...)
```

Arguments

gene a character or a object IRanges
calls an object from cghCall class

... further arguments passed to or from other methods

Details

```
add details (?)
```

Value

The function return an object of the class 'cghCall'.

References

Mark A. van de Wiel, Kyung In Kim, Sjoerd J. Vosse, Wessel N. van Wieringen, Saskia M. Wilting and Bauke Ylstra (2007) CGHcall: calling aberrations for array CGH tumor profiles. Bioinformatics, 23,892-894.

combiUHM

Combination of DNA methylation status of CpG-loci

Description

Combination of DNA methylation status of CpG-loci

Usage

```
combiUHM(x,rare=NULL,mc=1000)
## S3 method for class 'combiUHM'
print(x,...)
## S3 method for class 'combiUHM'
plot(x,cex.label=0.75,main=NULL,add.lines=TRUE,...)
```

entropy.index 7

Arguments

x a sdata.frame
rare a numeric value
mc a numeric value
cex.label a numeric value
main a character value
add.lines a logical value

... further arguments passed to or from other methods

Value

return a combiUHM object

 $\verb"entropy.index"$

Computation of the Entropy/Shannon index

Description

The function computes the Entropy/shannon index.

Usage

```
entropy.index(x,standard=FALSE,...)
```

Arguments

X a vector or a data.frame

Standard a logical value indicating if the value is standardized by maximal Entropy (FALSE

by default).

... further arguments passed to or from other methods

Value

The functions returns a numeric or vector containing the entropy values (by row if x is a data.frame).

References

```
add reference fro Shannon index (?) hataway (1992)
```

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Examples

```
## example 1
log(4)
x <- c(0.25,0.25,0.25,0.25)
-sum(x*log(x))
entropy.index(x,standard=FALSE)
entropy.index(x,standard=TRUE)
## example 2
x <- rep(4,4)
entropy.index(x,standard=FALSE)
entropy.index(x,standard=TRUE)</pre>
```

extractCGH

Extraction of gene or genomic regions into an object 'cghCall'

Description

The functions extract of 'cghCall' information for a given gene, chromosomes or genomic regions.

Usage

```
extractCGH.gene(x,gene,...)
extractCGH.chr(x,chr,...)
extractCGH.default(x,chr, start,end,winsize=0.0,info=NULL,mode=NULL,...)
```

Arguments

X	an object 'cghCall' (output of the function ExpandCGHcall from Rpackage CGHcall).
chr	a character corresponding to the chromosome information (e.g. "chr1")
gene	a character value corresponding to gene name (Symbol name)
start	a numeric value corresponding to the start position of the selected genomic region
end	a numeric value corresponding to the end position of the selected genomic region
info	a character value describing the genomic region (NULL by default).
winsize	a numerical value used to increase the windows of the selected genomic region (in pb)
mode	a numeric value to select the selection mode (NULL by default, see details).
	further arguments passed to or from other methods

Details

The estimation is based on mixture models proposed in the R package CGHcall (see functions called 'CGHcall').

The argument called "mode" is used to select the mode of selection: * mode 1 (include): |x---x|

By default, the mode is equal to NULL that corresponds to the use of the four selection modes (c(1,2,3,4)).

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Value

The functions returns a object 'cghCall' with an addition attributes called "loc" (as location). This new attribute is a list describing below:

info	a character value describing the genomic region, for example gene or chromosome name (NULL by default).
chr	a character corresponding to the chromosome information (e.g. "chr1")
start	a numeric value corresponding to the start position of the selected genomic region
end	a numeric value corresponding to the end position of the selected genomic region
winsize	a numerical value used to increase the windows of the selected genomic region (in pb)
mode	a numeric value to select the selection mode (NULL by default, see details).
gene2IR	Convert a gene symbol, geneloc object or cghCall object into IRanges object

Description

The functions convert a gene symbol, geneloc object or cghCall object into IRanges object

Usage

```
gene2IR(gene, calls, chrom = NULL, ...)
cghCall2IR(x, chrom, ...)
geneloc2IR(x, ...)
gene2GR(gene, calls, chrom = NULL, ...)
cghCall2GR(x, chrom, ...)
geneloc2GR(x, ...)
```

Arguments

X	a character value corresponding to a gene symbol or an object 'geneloc'
gene	a character value corresponding to a gene symbol or a object IRanges
calls	an cghCall object
chrom	an integer value given the chromosome location
	further arguments passed to or from other methods

Details

```
+ add details (?)
```

Value

The functions return on IRanges or GRanges object associated with a given gene or an cghCall object.

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References

Mark A. van de Wiel, Kyung In Kim, Sjoerd J. Vosse, Wessel N. van Wieringen, Saskia M. Wilting and Bauke Ylstra (2007) CGHcall: calling aberrations for array CGH tumor profiles. Bioinformatics, 23,892-894.

See Also

GRanges-class, IRanges-class

Examples

```
##---- Should be DIRECTLY executable !! ----
##-- ==> Define data, use random,
##--or do help(data=index) for the standard data sets.
## The function is currently defined as
function (gene, calls, chrom = NULL, ...)
    if (!inherits(gene, "IRanges") & is.character(gene)) {
        gene1 <- getGeneLocationR(gene)</pre>
        chrom <- gene1$chrom</pre>
        chrom \leftarrow c(1:24, 23, 24)[match(chrom, c(paste("chr", 
             1:24, sep = ""), "chrX", "chrY"))]
        chrom <- as.numeric(chrom)</pre>
        if (length(unique(chrom)) > 1)
            warning("multiple chrosome locations!")
        chrom <- unique(chrom)[1]</pre>
        wgene <- geneloc2IR(gene1)</pre>
    else if (inherits(geneloc2IR, "IRanges") & !is.null(chrom)) {
        wgene <- gene
    }
    else stop("non convenient argument!")
    wcalls <- cghCall2IR(calls, chrom)</pre>
    wintersect <- findOverlaps(wcalls, wgene)</pre>
    res <- wcalls[attributes(wintersect)$queryHits]</pre>
    attr(res, "chrom") <- chrom</pre>
    attr(res, "call") <- match.call()</pre>
    return(res)
```

getGeneLocationR

Get the location of a vector of gene

Description

This function provides the location information of a given gene (character) or a set of gene (vector of character) based on the R packages 'org.Hs.eg.db' and 'TxDb.Hsapiens.UCSC.hg19.knownGene'.

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Usage

```
getGeneLocationR(genelist, group = TRUE, orderby = "genename",...)
getGeneLocationR2(genelist, group = TRUE, orderby = "genename",onlychr=FALSE,...)
getGeneInfoR(genelist, orderby = "genename",exon=TRUE,cds=TRUE,tx=TRUE,ucsckg=TRUE,...)
getVariantInfoR(chrom,start,end=start,labels=NULL,winsize=0,verbose=FALSE,...)
getVariantInfoR.df(df,control=list(),winsize=0,verbose=FALSE,...)
variant.control(label="rownames",chrom="chrom",start="start",end=start)
```

Arguments

genelist	a vector of character
group	a logical value (TRUE by default, see details)
orderby	a character corresponding to the name of the column used to order the output
tx	a logical value (TRUE by default, see details)
exon	a logical value (TRUE by default, see details)
cds	a logical value (TRUE by default, see details)
ucsckg	a logical value (TRUE by default, see details)
df	a data.frame containing label,chromosome, end and start position information
chrom	a character vector containing chromosome information (e.g. chr11)
start	a numeric vector containing start position
end	a numeric vector containing end position
labels	a character vector containing names information (e.g. rsxxxxx for SNP or cgxxxxxx for DNA methylation probes)
labels	a numeric value used to modify the windows size (start-winsize and end-winsize, winsize=0 by default)
verbose	a logical value (FALSE by default)
control	a list of parameters for controlling the selection of label, chromosome, end and start position information. This is passed to 'variant.control'.
onlychr	a logical value for the exclusion of elements with chromosome information containing '_' (by default, onlychr=FALSE).
	further arguments passed to or from other methods

Details

add info on the argument "group"!!!

Value

The function returns an object of the class geneloc corresponding to a data.frame containing gene location information from R packages 'org.Hs.eg.db' and 'TxDb.Hsapiens.UCSC.hg19.knownGene'.

genename the symbol gene
chrom the chromosome location
txStart the physical location of the starting point
txEnd the physical location of the ending point
strand the strand of the gene

The data.frame is ordered by genename by default (see argument orderby). ... add more information for the functions getInfoR() and getVariantInfoR()

12 getTCGAnames

getTCGAnames

TCGA sample name decoding

Description

The function decodes the names of the samples from TCGA project (optimized for GBM).

Usage

```
getTCGAnames(x, rownames = TRUE, ...)
```

Arguments

```
    a character
    a logical values (by default TRUE). If rownames is equal to TRUE, the rownames of output is the sample names.
    further arguments passed to or from other methods
```

Details

+ description de l'orgasnisation du code (?) need to check the type of tissue available in the TCGA databases.

Value

a data frame containing the decoded information related to the name of sample from TCGA project.

References

reference for TCGA project

Examples

```
##---- Should be DIRECTLY executable !! ----
##-- ==> Define data, use random,
##--or do help(data=index) for the standard data sets.
## The function is currently defined as
function (x, rownames = TRUE, ...)
{
    if (is.factor(x))
        x <- as.character(x)</pre>
    if (!is.character(x))
        stop("non convenient argument!")
    fdecode <- function(z) {</pre>
        w \leftarrow list(Name = substring(z, 1, 12), Project = substring(z, 1, 12))
            1, 4), CollectionCenter = substring(z, 6, 7), Patient = substring(z,
            9, 12), SampleType = substring(z, 14, 15), SampleSequence = substring(z,
            16, 16), PortionSequence = substring(z, 18, 19),
            PortionAnalyte = substring(z, 20, 20), PlateID = substring(z,
                 22, 25), DataGeneratingCenter = substring(z,
        w$type <- c("solid tumor", "normal blood", "normal tissue",
```

logoffset 13

logoffset

Log link function with an offset

Description

Computes the log transformation with an offset

Usage

```
logoffset(x, ..., offset = 0, method = "only0")
```

Arguments

```
    x a numeric
    ... further arguments passed to or from other methods
    offset a numeric corresponding to the offset (equal to 0 by default)
    method a character
```

Details

+ description of the argument 'method'

Value

return a numeric

Examples

```
##---- Should be DIRECTLY executable !! ----
##-- ==> Define data, use random,
##--or do help(data=index) for the standard data sets.

## The function is currently defined as
function (x, ..., offset = 0, method = "only0")
{
    switch(method, only0 = ifelse(x == 0, 0 + offset, log(x, ...)), all = log(x + offset, ...), stop("non convient method!"))
}
```

14 maxCNAcalls

maxCNAcalls	Estimation of the CNA for a given region based on Mixture Model at the sample scale.

Description

set of functions used to estimate the CNA of a given region, chromosome or gene for each individual samples (samples scale estimation). The estimation is based on mixture models proposed in the R package CGHcall.

Usage

```
maxCNAcalls(X,chrom, start,end,winsize=0,mode=NULL,nclass=5,...)
maxCNAcalls.chr(X,chrom,nclass=5,...)
maxCNAcalls.gene(X,gene,nclass=5,...)
## S3 method for class 'CNAcalls'
print(x,nclass=5,...)
```

Arguments

X	an object 'cghCall' (output of the function ExpandCGHcall from Rpackage CGHcall).
X	an object 'CNAcalls'
chrom	a character corresponding to the chromosome information (e.g. "chr1")
gene	a character value corresponding to gene name (Symbol name)
start	a numeric value corresponding to the start position of the selected genomic region
end	a numeric value corresponding to the end position of the selected genomic region
winsize	a numerical value used to increase the windows of the selected genomic region (in pb)
mode	a numeric value to select the selection mode (NULL by default, see details).
nclass	a numeric value to select the number of CNA state (5 by default, see the Function CGHcall from R package CGHcall).
	further arguments passed to or from other methods

Details

The estimation is based on mixture models proposed in the R package CGHcall (see functions called 'CGHcall').

```
* mode 2 (incomplete overlap): x——|——x |
```

By default, the mode is equal to NULL that corresponds to the use of the four selection modes (c(1,2,3,4)).

^{*} mode 3 (incomplete overlap): | x——|——x

^{*} mode 4 (total overlap): x——l—x

meth2norm 15

Value

The functions returns a list of of classes "CNAcalls":

a character vector containing the CNA status for each sample

a numeric value corresponding to the number of samples

info a character value describing the selected genomic regions

winsize a numerical value used to increase the windows of the selected genomic region (in pb)

location a list containing information about the location of the selected region (chr, start and end positions)

summary a numerical vector containing a summarize of the CNA information

call the call function

References

Mark A. van de Wiel, Kyung In Kim, Sjoerd J. Vosse, Wessel N. van Wieringen, Saskia M. Wilting and Bauke Ylstra. CGHcall: calling aberrations for array CGH tumor profiles. Bioinformatics, 23,892-894.

Examples

```
#+ add examples
```

meth2norm	Normalization and preparation of the Illumina infinium DNA methyla- tion data
	tion adia

Description

The function proposes the normalization of the Illumina infinium DNA Methylation data and computation of total intensity (Methylation + Unmethylation). Before the computation of the total intensity, the Unmethylated and Methylated information can be normalized in the same way and time (quantile normalization by default).

Usage

```
meth2norm(x,...)
## Default S3 method:
meth2norm(x,y,method="normalize.quantiles",scaling=FALSE,
chemistry=NULL,rev=FALSE,verbose=TRUE,add.UM=FALSE,...)
## S3 method for class 'MethySet'
meth2norm(x,method="normalize.quantiles",scaling=FALSE,
chemistry=NULL,rev=FALSE,verbose=TRUE,add.UM=FALSE,...)
```

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Arguments

X	an object 'MethySet' from preprocess functions of package 'minfi' or a matrix or data.frame containing probes for unmethylation
У	a matrix or data.frame containing probes for methylation (if x is a matrix or data.frame containing probes for unmethylation).
method	a character corresponding to normalization method (by default 'normalize.quantiles' from package preprocessCore) $$
scaling	a logical value indicating if the scaling factor procedure is used to correct the chemistry effect (by default 'FALSE')
	further arguments passed to or from other methods

Details

The unmethylated and methylated tables are normalized in the same tables after concatenation of them (by default, we propose the use of quantile normalization). The total intensity is obtained by sum of the normalized unmethylated table and normalized methylated table.

Value

The function returns a list of three normalized matrix

T	an object 'matrix' containing the total intensity
U	an object 'matrix' containing the normalized unmethylation intensity
М	an object 'matrix' containing the normalized unmethylation intensity
colnames	a vector of character containing the sample names
rownames	a vector of character containing the probe names
call	generally 'match.call()'

References

Staaf, Johan and Vallon-Christersson, Johan and Lindgren, David and Juliusson, Gunnar and Rosenquist, Richard and Hoglund, Mattias and Borg, Ake and Ringner, Markus (2008) Normalization of Illumina Infinium whole-genome SNP data improves copy number estimates and allelic intensity ratios, BMC Bioinformatics, 9:409 Louhimo, R. and Hautaniemi, S. (2011) CNAmet: an R package for integration of copy number, expression and methylation data Bioinformatics 27(6):887-888.

Examples

```
##---- Should be DIRECTLY executable !! ----
##-- ==> Define data, use random,
##--or do help(data=index) for the standard data sets.
## The function is currently defined as
function (...)
{
    UseMethod("meth2norm")
}
```

mixCNAcalls 17

mixCNAcalls	Estimation of the CNA for a given region based on Mixture Model at the sample scale.

Description

set of functions used to estimate the CNA of a given region, chromosome or gene for each individual samples (samples scale estimation). The estimation is based on mixture models proposed in the R package CGHcall.

Usage

```
mixCNAcalls(X,chrom, start,end,winsize=0,mode=NULL,nclass=5,...)
mixCNAcalls.chr(X,chrom,nclass=5,...)
mixCNAcalls.gene(X,gene,nclass=5,...)
## S3 method for class 'mixCNAcalls'
print(x,nclass=5,...)
```

Arguments

X	an object 'cghCall' (output of the function ExpandCGHcall from Rpackage CGHcall).
x	an object 'CNAcalls'
chrom	a character corresponding to the chromosome information (e.g. "chr1")
gene	a character value corresponding to gene name (Symbol name)
start	a numeric value corresponding to the start position of the selected genomic region
end	a numeric value corresponding to the end position of the selected genomic region
winsize	a numerical value used to increase the windows of the selected genomic region (in pb)
nclass	a numeric value to select the number of CNA state (5 by default, see the Function CGHcall from R package CGHcall).
	further arguments passed to or from other methods

Details

The estimation is based on mixture models proposed in the R package CGHcall (see functions called 'CGHcall'). explain the definition of the limits

Value

The functions returns a list of of classes "mixCNAcalls":

```
cna a character vector containing the CNA status for each sample
n a numeric value corresponding to the number of samples
info a charcter value describing the selected genomic regions
```

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winsize a numerical value used to increase the windows of the selected genomic region

(in pb)

location a list containing information about the location of the selected region (chr, start

and end positions)

summary a numerical vector containing a summarize of the CNA information

call the call function

References

Mark A. van de Wiel, Kyung In Kim, Sjoerd J. Vosse, Wessel N. van Wieringen, Saskia M. Wilting and Bauke Ylstra. CGHcall: calling aberrations for array CGH tumor profiles. Bioinformatics, 23,892-894.

Examples

```
#+ add examples
```

pathway.test

Pathway test by hyperfeometric test

Description

This function provide a pathway test by hyperfeometric test

Usage

```
pathway.test(DEgene,pathway,p.method="bonferroni",...)
pathway.mctest(DEgene,pathway,p.method="bonferroni",cpus=2,...)
pathway2df(gene,pathway,...)
```

Arguments

DEgene a vector of binary values. the value is 0 for unselected genes and 1 for selected

genes. the names of the vector can be EntrezID.

gene a vector of character values corresponding to the gene names (the vector can be

EntrezID).

pathway list of genes (same type/id used to name of DEgene or gene elements) from PID,

GO, Kegg, etc ...

p.method see the argument 'method' related to the R functions p.adjust

cpus an integer corresponding to the number of cpus used for computation (see pack-

age snowfall)

... further arguments passed to or from other methods

Details

+ details ?! fisher.test(matrix(c(a,c,b,d),2,2),alternative="greater") phyper(a-1,a+b,c+d,a+c,lower.tail=FALSE)

prep.annot450k

Value

The function returns a data.frame containing the information related to pathway test.

	n	total number of gene
	in1	number of selected gene in pathway
	in0	number of unselected gene in pathway
	out1	number of unselected gene not in pathway
	out0	number of unselected gene not in pathway
	oddsratio	estimate odds ratio
	p.raw	p.value from hypergeometric test
	p.adj	adjusted p.value (see argument 'p.method')
	+	
### Example with GO with R packages gage # per type MF, CC or BP gomf <- go.hs\$go.sets[go.hs\$go.subs\$MF] degene1 <- sample(c(0,1),length(unique(unlist(gomf)[1:10000])),replace=TRUE) names(degene1) <- unique(unlist(gomf)[1:10000]) table(degene1)		
	#Go analysis require(gage) go.hs <- go.gsets(species="human") names(go.hs) gogs <- go.hs\$go.sets	
	ted_go <- pathway.test(degene1,gomf[1:8]) ted_go	
	### Example with PID and NCI from R package CePa require(CePa) data(PID.db) names(PID.db) ted_nci <- pathway.test(degene1,PID.db\$NCI\$pathList[1:8]) ted_nci	

Description

Preparation of annotation file

Usage

```
prep.annot450k(x, snp.rm = TRUE, sex.rm = TRUE,in27k=FALSE,chemistry=NULL)
```

Arguments

Х	a data.frame containing the annotation of Infinium HM-450K platform	
snp.rm	a logical value. If TRUE, probes located on snp locations are excluded (by default TRUE).	
sex.rm	a logical value. If TRUE, probes located on sex chromosomes are excluded (by default TRUE).	
in27k	a logical value. If TRUE, probes common to HM-27K and HM-450K platforms (by default FALSE).	
chemistry	a character value. Selection of for chemistry type I ("I"), for chemistry type II ("II") or the both chemistry type (by default NULL).	

Details

this function is optimized to works on annotation files from Illumina compagny:

 $- Human Methylation 450_15017482_v.1.1_For Excel.csv-human methylation 450_15017482_v1-2.csv-human methylation 450_15017482_$

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Value

a data.frame containing the annotation of Infinium HM-450K platform filtered for SNP and Sex information $\frac{1}{2}$

Examples

REout

Remove Bracket, sapce or zero values contained in a character object

Description

The functions remove some specific symbols (such as bracket, space or zero) in a character object.

Usage

```
REbracketout(x, replaceby = "", ...)
REspaceout(x, replaceby = "", ...)
REzeroout(x, replaceby = "", ...)
```

Arguments

```
    x a vector of character
    replaceby a character value replacing bracket, space or zero value contained in x.
    ... further arguments passed to or from other methods
```

Value

return a vector of character (string) where bracket, space or zero symbols was removed.

See Also

gsub

searchGene 21

searchGene search "gene" containing in a character vector	
-----------------------------------------------------------	--

Description

This function offers the possibility to search gene (character pattern) containing in a character vector separted by a given separator (semi-colon by default).

Usage

```
searchGene(x,gene,sep=";",type="logical")
```

Arguments

X	a character vector
gene	the gene (character pattern) of interest
sep	separator between the gene (by default, sep=";")
type a character indicating the type of values returned by the function. By de the function return a logical vector (type="logical"). The type "numeric" "value" return position and value respectively.	

Value

The function returns a vector of logical (type="logical"), numeric (type="numeric") or character (type="value") depending of the selected type.

Examples

```
genelist <-c("MGMT","MGMT;MGMT","MGMT;EGFR","HOXA1;EGFR;HOXA2","EGFR;MGMT",NA)
searchGene(genelist,"EGFR",sep=";",type="logical")
searchGene(genelist,"EGFR",sep=";",type="numeric")
searchGene(genelist,"EGFR",sep=";",type="value")

seg2tab

Convert Segmentation data of an object 'cghSeg' or 'cghCall' in a</pre>
```

matrix (or list of matrix)

Description

The function converts segmentation data ("chrom", "start", "end", "num.mark", "seg.mean") of an object 'cghSeg' or 'cghCall' in a matrix (or list of matrix)

Usage

```
seg2tab(x,...)
```

Arguments

```
x an object 'cghSeg' or 'cghCall'... further arguments passed to or from other methods
```

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Details

```
+ details ?!
```

Value

The function returns an matrix of a list of matrix (if there are several sample) containing the segmentation information obtained by the the function DNAcopy from the R package DNAcopy.

chrom the chromosome within the sample start the starting map location of the segment end the ending map location of the segment num.mark the number of markers in the segment

seg.mean the segment mean

References

Mark A. van de Wiel, Kyung In Kim, Sjoerd J. Vosse, Wessel N. van Wieringen, Saskia M. Wilting and Bauke Ylstra (2007) CGHcall: calling aberrations for array CGH tumor profiles. Bioinformatics, 23,892-894.

Examples

```
##--- Should be DIRECTLY executable !! ----
##-- ==> Define data, use random,
##--or do help(data=index) for the standard data sets.
## The function is currently defined as
```

UHM

Three-Mixture models for DNA methylation genotyping

Description

These functions fitting models for DNA methylation genotyping based on M-values (gamma or normal version) or Beta-values (beta version).

Usage

```
normalUHM(df,cpus=3,verbose=FALSE,...)
gammaUHM(df,lim=c(0.95,0.95),cpus=3,...)
```

Arguments

df	a data.frame containing	beta or M-values

cpus an integer corresponding to the number of cpus used for computation (see pack-

age snowfall)

1im a numeric vector containing the limit used to defined the DNA methylation geno-

type

verbose logical, if TRUE information are printed during each step of the algorithm

... further arguments passed to or from other methods

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Details

+ details ?!

Value

add retunr

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