

Package ‘compartmap’

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Type Package

Title A/B compartment inference from ATAC-seq and methylation array data

Description Compartmap performs shrunken A/B compartment inference from ATAC-seq and methylation arrays.

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URL <https://github.com/biobenkj/compartmap>

BugReports <https://github.com/biobenkj/compartmap/issues>

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array.data.chr14	<i>Example Illumina 450k methylation array data for compartmap</i>
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Description

This data was generated using the data from the reference via the `sesamize` function from the `SeSAMe` package.

Usage

```
data(meth_array_450k_chr14, package = "compartmap")
```

Author(s)

Benjamin K Johnson <ben.johnson@vai.org>

References

<https://f1000research.com/articles/5-1281/v3>

filtered.data.chr14	<i>Example ATAC-seq data for compartmap</i>
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Description

This data was generated using the data from the reference via `bwa mem` and pre-processing the data using the `ATACseeker` package.

Usage

```
data(bulkATAC_raw_filtered_chr14, package = "compartmap")
```

Author(s)

Benjamin K Johnson <ben.johnson@vai.org>

References

<https://trace.ncbi.nlm.nih.gov/Traces/sra/?study=SRP082417>

fisherZ	<i>Fisher's Z transformation</i>
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Description

fisherZ returns (squeezed) Fisher's Z transformed Pearson's r

Usage

```
fisherZ(cormat)
```

Arguments

cormat Pearson correlation matrix

Details

This function returns (squeezed) Fisher's Z transformed Pearson's r

Value

Fisher Z transformed Pearson correlations

Examples

```
#Generate a random binary (-1, 1) matrix
mat <- matrix(sample(c(1,-1), 10000, replace = TRUE), ncol = 100)

#Correct matrix diag
diag(mat) <- 1

#Transform
mat.transform <- fisherZ(mat)
```

getABSignal	<i>Calculate Pearson correlations of smoothed eigenvectors</i>
-------------	--

Description

This function is used to generate a list x to be passed to getABSignal

Usage

```
getABSignal(x, k = 5, iter = 2, squeeze = FALSE)
```

Arguments

x	A list object from getCorMatrix
k	Value of k for smoothing (default = 2)
iter	Number of iterations for moving average smoothing (default = 2)
squeeze	Whether squeezing was used (implies Fisher's Z transformation)

Value

A list x to pass to getABSignal

Examples

```
library(GenomicRanges)
library(Homo.sapiens)
library(mixOmics)

#Generate random genomic intervals of 1-1000 bp on chr1-22
#Modified from https://www.biostars.org/p/225520/
random_genomic_int <- data.frame(chr = rep("chr14", 100))
random_genomic_int$start <- apply(random_genomic_int, 1, function(x) { round(runif(1, 0, seqlengths(Homo.sapiens$chr14)), 1)})
random_genomic_int$end <- random_genomic_int$start + runif(1, 1, 1000)
random_genomic_int$strand <- "*"

#Generate random counts
counts <- rnbinom(1000, 1.2, 0.4)

#Build random counts for 10 samples
count.mat <- matrix(sample(counts, nrow(random_genomic_int) * 10, replace = FALSE), ncol = 10)
colnames(count.mat) <- paste0("sample_", seq(1:10))

#Bin counts
bin.counts <- getBinMatrix(count.mat, makeGRangesFromDataFrame(random_genomic_int), chr = "chr14", genome = "Homo.sapiens")

#Calculate correlations
bin.cor.counts <- getCorMatrix(bin.counts)

#Get A/B signal
absignal <- getABSignal(bin.cor.counts)
```

getArrayABsignal	<i>Estimate A/B compartments from methylation array data</i>
------------------	--

Description

getArrayABsignal returns estimated A/B compartments from methylation array data.

Usage

```
getArrayABsignal(obj, res = 1e+06, parallel = FALSE, allchrs = FALSE,
  chr = NULL, targets = NULL, ...)
```

Arguments

obj	Input GenomicRatioSet object
res	Compartment resolution (in bp)
parallel	Should the inference be done in parallel?
allchrs	Whether all autosomes should be used for A/B inference
chr	Specific chromosomes to analyze
targets	Specify samples to use as shrinkage targets
...	Additional arguments

Details

This function is modified from the `minfi::compartments` to infer A/B compartments from array data

Value

A $p \times n$ matrix (samples as columns and compartments as rows) of compartments

getATACABsignal	<i>Estimate A/B compartments from ATAC-seq data</i>
-----------------	---

Description

getATACABsignal returns estimated A/B compartments from methylation array data.

Usage

```
getATACABsignal(obj, res = 1e+06, parallel = FALSE, allchrs = FALSE,
  chr = NULL, targets = NULL, ...)
```

Arguments

obj	Input GenomicRatioSet object
res	Compartment resolution (in bp)
parallel	Should the inference be done in parallel?
allchrs	Whether all autosomes should be used for A/B inference
chr	Specify a chromosome to analyze
targets	Specify samples as shrinkage targets
...	Additional arguments

Details

This function estimates A/B compartments shrinking towards a global mean of targets or across samples

Value

A $p \times n$ matrix (samples as columns and compartments as rows) of compartments

Examples

```
library(GenomicRanges)
library(SummarizedExperiment)
library(Homo.sapiens)

data(bulkATAC_raw_filtered_chr14, package = "compartmap")
atac_compartments <- getATACABsignal(filtered.data.chr14, chr = "chr14", genome = "hg19")
```

getBinMatrix

Generate bins for A/B compartment estimation

Description

Generate bins across a user defined chromosome for A/B compartment estimation. A/B compartment estimation can be used for non-supported genomes if chr.end is set.

Usage

```
getBinMatrix(x, genloc, chr = "chr1", chr.start = 0, chr.end = NULL,
  res = 1e+05, FUN = sum, genome = "hg19")
```

Arguments

x	A p x n matrix where p (rows) = loci and n (columns) = samples/cells
genloc	GRanges object that contains corresponding genomic locations of the loci
chr	Chromosome to be analyzed
chr.start	Starting position (in bp) to be analyzed
chr.end	End position (in bp) to be analyzed
res	Binning resolution (in bp)
FUN	Function to be used to summarize information within a bin
genome	Genome corresponding to the input data ("hg19" or "mm10")

Details

This function is used to generate a list object to be passed to getCorMatrix

Value

A list object to pass to getCorMatrix

Examples

```
library(GenomicRanges)
library(Homo.sapiens)

#Generate random genomic intervals of 1-1000 bp on chr1-22
#Modified from https://www.biostars.org/p/225520/
random_genomic_int <- data.frame(chr = rep("chr14", 100))
random_genomic_int$start <- apply(random_genomic_int, 1, function(x) { round(runif(1, 0, seqlengths(Homo.sapiens))
```

```

random_genomic_int$end <- random_genomic_int$start + runif(1, 1, 1000)
random_genomic_int$strand <- "*"

#Generate random counts
counts <- rnbinom(1000, 1.2, 0.4)

#Build random counts for 10 samples
count.mat <- matrix(sample(counts, nrow(random_genomic_int) * 10, replace = FALSE), ncol = 10)
colnames(count.mat) <- paste0("sample_", seq(1:10))

#Bin counts
bin.counts <- getBinMatrix(count.mat, makeGRangesFromDataFrame(random_genomic_int), chr = "chr14", genome = "hg19")

```

getCompartments	<i>Estimate A/B compartments</i>
-----------------	----------------------------------

Description

getCompartments returns estimated A/B compartments from ATAC-seq and methylation array data

Usage

```

getCompartments(obj, type = c("atac", "array"), res = 1e+06,
  parallel = FALSE, chrs = "chr1", genome = "hg19", targets = NULL,
  run_examples = FALSE, ...)

```

Arguments

obj	The object with which to perform compartment inference
type	The type of data that obj represents (e.g. atac or array)
res	Resolution of compartments in base pairs (default is 1e6)
parallel	Should the estimates be done in parallel (default is FALSE)
chrs	Chromosomes to operate on (can be individual chromosomes, a list of chromosomes, or all)
genome	Genome to use (default is hg19)
targets	Specify samples to use as shrinkage targets
run_examples	Whether to run ATAC-seq and 450k example analysis
...	Other parameters to pass to internal functions

Details

This is a wrapper function to perform A/B compartment inference. Compartmentalizer implements a Stein estimator to shrink per-sample compartment estimates towards a global mean. The expected input for this function can be generated using packages like SeSAME and ATACseeker.

Value

A p x n matrix (samples as columns and compartments as rows) to pass to embed_compartments

Examples

```

library(GenomicRanges)
library(SummarizedExperiment)
library(Homo.sapiens)

#ATAC-seq data
data(bulkATAC_raw_filtered_chr14, package = "compartmap")
atac_compartments <- getCompartments(filtered.data.chr14, type = "atac", parallel = FALSE, chrs = "chr14")
## Not run:
#450k data
data(meth_array_450k_chr14, package = "compartmap")
array_compartments <- getCompartments(array.data.chr14, type = "array", parallel = FALSE, chrs = "chr14")
## End(Not run)

```

getCorMatrix

Calculate Pearson correlations of a binned matrix

Description

This function is used to generate a list object to be passed to getABSignal

Usage

```
getCorMatrix(binmat, squeeze = FALSE)
```

Arguments

binmat	A binned matrix list object from getBinMatrix
squeeze	Whether to squeeze the matrix for Fisher's Z transformation

Value

A list object to pass to getABSignal

Examples

```

library(GenomicRanges)
library(Homo.sapiens)

#Generate random genomic intervals of 1-1000 bp on chr1-22
#Modified from https://www.biostars.org/p/225520/
random_genomic_int <- data.frame(chr = rep("chr14", 100))
random_genomic_int$start <- apply(random_genomic_int, 1, function(x) { round(runif(1, 0, seqlengths(Homo.sapiens)[x]) * 0.5) })
random_genomic_int$end <- random_genomic_int$start + runif(1, 1, 1000)
random_genomic_int$strand <- "*"

#Generate random counts
counts <- rnbinom(1000, 1.2, 0.4)

#Build random counts for 10 samples
count.mat <- matrix(sample(counts, nrow(random_genomic_int) * 10, replace = FALSE), ncol = 10)

```



```
colnames(count.mat) <- paste0("sample_", seq(1:10))

#Bin counts
bin.counts <- getBinMatrix(count.mat, makeGRangesFromDataFrame(random_genomic_int), chr = "chr14", genome = "H")

#Calculate correlations
bin.cor.counts <- getCorMatrix(bin.counts)
```

ifisherZ

Fisher's Z transformation

Description

fisherZ returns the inverse (squeezed) Fisher's Z transformed Pearson's r. This will fail if a matrix is used as input instead of a vector.

Usage

```
ifisherZ(cormat)
```

Arguments

cormat vector of Fisher's Z transformed Pearson correlations or an eigenvector

Details

This function returns the inverse (squeezed) Fisher's Z transformed Pearson's r

Value

Back transformed Fisher's Z

Examples

```
#Generate a random binary (-1, 1) matrix
mat <- matrix(sample(c(1,-1), 10000, replace = TRUE), ncol = 100)

#Correct matrix diag
diag(mat) <- 1

#Transform
mat.transform <- fisherZ(mat)

#Back transform
mat.transform.inverse <- apply(mat.transform, 1, ifisherZ)
```

plotAB

*Plots A/B compartment estimates on a per chromosome basis***Description**

Plot A/B compartments bins

Usage

```
plotAB(x, main = "", ylim = c(-1, 1), unitarize = FALSE,
       reverse = FALSE, top.col = "deeppink4", bot.col = "grey50")
```

Arguments

x	The matrix object returned from getCompartments
main	Title for the plot
ylim	Y-axis limits (default is -1 to 1)
unitarize	Should the data be unitarized? (not explicitly necessary for arrays)
reverse	Reverse the sign of the PC values?
top.col	Top (pos. PC values) chromatin color to be plotted
bot.col	Bottom (neg. PC values) chromatin color to be plotted

Value

invisibly, the compartment estimates from the plot

Examples

```
library(GenomicRanges)
library(Homo.sapiens)

#Generate random genomic intervals of 1-1000 bp on chr1-22
#Modified from https://www.biostars.org/p/225520/
random_genomic_int <- data.frame(chr = rep("chr14", 100))
random_genomic_int$start <- apply(random_genomic_int, 1, function(x) { round(runif(1, 0, seqlengths(Homo.sapiens)[x]), 1) })
random_genomic_int$end <- random_genomic_int$start + runif(1, 1, 1000)
random_genomic_int$strand <- "*"

#Generate random counts
counts <- rnbinom(1000, 1.2, 0.4)

#Build random counts for 10 samples
count.mat <- matrix(sample(counts, nrow(random_genomic_int) * 10, replace = FALSE), ncol = 10)
colnames(count.mat) <- paste0("sample_", seq(1:10))

#Bin counts
bin.counts <- getBinMatrix(count.mat, makeGRangesFromDataFrame(random_genomic_int), chr = "chr14", genome = "Homo.sapiens")

#Calculate correlations
bin.cor.counts <- getCorMatrix(bin.counts)
```

```
#Get A/B signal
absignal <- getABSignal(bin.cor.counts)

#Plot the A/B signal
par(mar=c(1,1,1,1))
par(mfrow=c(1,1))
plotAB(absignal$pc, ylim = c(-0.2, 0.2), unitarize = TRUE)

## Not run:
#If plotting individual A/B signals using output from getCompartments
#Note: this function currently only supports plotting individual chromosomes from single samples
bin.chr1.ab <- getCompartments(data, "array", chrs = "chr1", genome = "hg19")

#For 7 samples
#Adjust ylim as necessary
par(mar=c(1,1,1,1))
par(mfrow=c(7,1))
plotAB(bin.chr1.ab[,1], ylim = c(-0.2, 0.2), unitarize = TRUE)
plotAB(bin.chr1.ab[,2], ylim = c(-0.2, 0.2), unitarize = TRUE, top.col = "goldenrod")
plotAB(bin.chr1.ab[,3], ylim = c(-0.2, 0.2), unitarize = TRUE, top.col = "darkblue")
plotAB(bin.chr1.ab[,4], ylim = c(-0.2, 0.2), unitarize = TRUE, top.col = "red")
plotAB(bin.chr1.ab[,5], ylim = c(-0.2, 0.2), unitarize = TRUE, top.col = "black")
plotAB(bin.chr1.ab[,6], ylim = c(-0.2, 0.2), unitarize = TRUE, top.col = "cyan")
plotAB(bin.chr1.ab[,7], ylim = c(-0.2, 0.2), unitarize = TRUE, top.col = "seagreen")

## End(Not run)
```

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