

Package ‘DMCFB’

April 12, 2022

Type Package

Title Differentially Methylated Cytosines via a Bayesian Functional Approach

Version 1.8.0

Description DMCFB is a pipeline for identifying differentially methylated cytosines using a Bayesian functional regression model in bisulfite sequencing data. By using a functional regression data model, it tries to capture position-specific, group-specific and other covariates-specific methylation patterns as well as spatial correlation patterns and unknown underlying models of methylation data. It is robust and flexible with respect to the true underlying models and inclusion of any covariates, and the missing values are imputed using spatial correlation between positions and samples. A Bayesian approach is adopted for estimation and inference in the proposed method.

Depends R (>= 4.0.0), SummarizedExperiment, methods, S4Vectors, BiocParallel, GenomicRanges, IRanges

Imports utils, stats, speedglm, MASS, data.table, splines, arm, rtracklayer, benchmarkme, tibble, matrixStats, fastDummies, graphics

Suggests testthat, knitr, rmarkdown

VignetteBuilder knitr

biocViews DifferentialMethylation, Sequencing, Coverage, Bayesian, Regression

License GPL-3

Encoding UTF-8

LazyData true

BugReports <https://github.com/shokoohi/DMCFB/issues>

RoxygenNote 7.1.1

git_url <https://git.bioconductor.org/packages/DMCFB>

git_branch RELEASE_3_14

git_last_commit eab8fbe

git_last_commit_date 2021-10-26

Date/Publication 2022-04-12

Author Farhad Shokoohi [aut, cre] (<<https://orcid.org/0000-0002-6224-2609>>)

Maintainer Farhad Shokoohi <shokoohi@icloud.com>

R topics documented:

DMCFB-package	2
BSDMC-class	3
cBSDMC-method	4
combine-method	6
findDMCFB-method	7
methLevels-method	9
methReads-method	10
params	11
plotDMCFB-method	13
readBismark-method	14
totalReads-method	15

Index	17
--------------	-----------

DMCFB-package	<i>Differentially Methylated cytosines using functional Bayesian regression models</i>
---------------	--

Description

DMCFB is a profiling tool for identifying differentially methylated cytosines using Functional Bayesian Model in bisulfite sequencing data.

DMCFB methods

[findDMCFB](#), [plotDMCFB](#), [cBSDMC](#), [readBismark](#).

BSDMC objects

[BSDMC-class](#)

BSDMC-class	<i>BSDMC object</i>
-------------	---------------------

Description

The BSDMC object is an S4 class that represents differentially methylated CpG sites (DMCs) in BS-Seq Data.

Arguments

- | | |
|------------|---|
| methReads | The matrix <code>methReads</code> contains the number of methylated reads spanning a CpG-site. The rows represent the CpG sites in <code>rowRanges</code> and the columns represent the samples in <code>colData</code> . |
| totalReads | The matrix <code>totalReads</code> contains the number of reads spanning a CpG-site. The rows represent the CpG sites in <code>rowRanges</code> and the columns represent the samples in <code>colData</code> . |
| methLevels | The matrix <code>methLevels</code> contains the predicted methylation level spanning a CpG-site using Bayesian functional regression models. The rows represent the CpG sites in <code>rowRanges</code> and the columns represent the samples in <code>colData</code> . |

Value

A [BSDMC-class](#) object

Slots

- `methReads` An integer matrix
- `totalReads` An integer matrix
- `methLevels` A numeric matrix

Author(s)

Farhad Shokoohi <shokoohi@icloud.com>

See Also

[RangedSummarizedExperiment-class](#) [GRanges-class](#)

Examples

```
nr <- 500
nc <- 16
metht <- matrix(as.integer(runif(nr * nc, 0, nr)), nr)
methc <- matrix(rbinom(n = nr * nc, c(metht), prob = runif(nr * nc)), nr, nc)
meths <- matrix(as.integer(runif(nr * nc, 0, 10)), nr)
methl <- methc / metht
methv <- matrix((runif(nr * nc, 0.1, 0.5)), nr)
```

```
r1 <- GRanges(rep("chr1", nr), IRanges(1:nr, width = 1), strand = "*")
names(r1) <- 1:nr
cd1 <- DataFrame(Group = rep(c("G1", "G2"), each = nc / 2),
  row.names = LETTERS[1:nc])
OBJ2 <- cBSDMC(
  rowRanges = r1, methReads = methc, totalReads = metht,
  methLevels = methl, methStates = meths, methVars = methv, colData = cd1
)
OBJ2
```

cBSDMC-method

cBSDMC method

Description

Creates a [BSDMC-class](#) object

Usage

```
cBSDMC(
  methReads,
  totalReads,
  methLevels,
  rowRanges,
  colData = DataFrame(row.names = colnames(methReads)),
  metadata = list(),
  ...
)

## S4 method for signature 'matrix,matrix,matrix,GRanges'
cBSDMC(
  methReads,
  totalReads,
  methLevels,
  rowRanges,
  colData = DataFrame(row.names = colnames(methReads)),
  metadata = list(),
  ...
)
```

Arguments

- | | |
|------------|---|
| methReads | The matrix <code>methReads</code> contains the number of methylated reads spanning a CpG-site. The rows represent the CpG sites in <code>rowRanges</code> and the columns represent the samples in <code>colData</code> . |
| totalReads | The matrix <code>totalReads</code> contains the number of reads spanning a CpG-site. The rows represent the CpG sites in <code>rowRanges</code> and the columns represent the samples in <code>colData</code> . |

<code>methLevels</code>	The matrix <code>methLevels</code> contains the predicted methylation level spanning a CpG-site using Bayesian functional regression models. The rows represent the CpG sites in <code>rowRanges</code> and the columns represent the samples in <code>colData</code> .
<code>rowRanges</code>	A <code>GRanges</code> or <code>GRangesList</code> object describing the ranges of interest. Names, if present, become the row names of the <code>SummarizedExperiment</code> object. The length of the <code>GRanges</code> or <code>GRangesList</code> must equal the number of rows of the matrices in assays. If <code>rowRanges</code> is missing, a <code>SummarizedExperiment</code> instance is returned.
<code>colData</code>	Object of class 'DataFrame' containing information on variable values of the samples
<code>metadata</code>	A list of storing MCMC samples or DMCs
...	other possible parameters

Details

The rows of a BSDMC object represent ranges (in genomic coordinates) of interest. The ranges of interest are described by a `GRanges` or a `GRangesList` object, accessible using the `rowRanges` function. The `GRanges` and `GRangesList` classes contains sequence (e.g., chromosome) name, genomic coordinates, and strand information. Each range can be annotated with additional data; this data might be used to describe the range or to summarize results (e.g., statistics of differential abundance) relevant to the range. Rows may or may not have row names; they often will not.

Value

A `BSDMC-class`

Author(s)

Farhad Shokoohi <shokoohi@icloud.com>

Examples

```
set.seed(1980)
nr <- 150
nc <- 8
metht <- matrix(as.integer(runif(nr * nc, 0, 100)), nr)
methc <- matrix(rbinom(n = nr * nc, c(metht), prob = runif(nr * nc)), nr, nc)
meths <- matrix(as.integer(runif(nr * nc, 0, 10)), nr)
methl <- methc / metht
methv <- matrix((runif(nr * nc, 0.1, 0.5)), nr)
r1 <- GRanges(rep("chr1", nr), IRanges(1:nr, width = 1), strand = "*")
names(r1) <- 1:nr
cd1 <- DataFrame(
  Group = rep(c("G1", "G2"), each = nc / 2),
  row.names = LETTERS[1:nc]
)
OBJ2 <- cBSDMC(
  rowRanges = r1, methReads = methc, totalReads = metht,
  methLevels = methl, methStates = meths, methVars = methv, colData = cd1
)
```

OBJ2

combine-method	<i>combine method</i>
----------------	-----------------------

Description

combine two [BSDMC-class](#) or two [BSDMC-class](#)

Usage

```
combine(obj1, obj2)

## S4 method for signature 'BSDMC,BSDMC'
combine(obj1, obj2)
```

Arguments

obj1	A BSDMC-class
obj2	A BSDMC-class

Value

A [BSDMC-class](#) or [BSDMC-class](#)

Author(s)

Farhad Shokoohi <shokoohi@icloud.com>

Examples

```
set.seed(1980)
nr <- 150
nc <- 8
metht <- matrix(as.integer(runif(nr * nc * 2, 0, nr)), nr)
methc <- matrix(
  rbinom(n = nr * nc, c(metht), prob = runif(nr * nc * 2)),
  nr, nc * 2
)
methl <- methc / metht
r1 <- GRanges(rep("chr1", nr), IRanges(1:nr, width = 1), strand = "*")
names(r1) <- 1:nr
cd1 <- DataFrame(Group = rep("G1", each = nc), row.names = LETTERS[1:nc])
OBJ1 <- cBSDMC(
  rowRanges = r1, methReads = methc[, 1:nc], totalReads = metht[, 1:nc],
  methLevels = methl[, 1:nc], colData = cd1
)
cd2 <- DataFrame(
  Group = rep("G2", each = nc),
```

```

    row.names = LETTERS[nc + 1:nc]
  )
OBJ2 <- cBSDMC(
  rowRanges = r1, methReads = methc[, nc + 1:nc], totalReads =
  metht[, nc + 1:nc], methLevels = methl[, nc + 1:nc], colData = cd2
)
OBJ3 <- combine(OBJ1, OBJ2)
OBJ3

```

findDMCFB-method *findDMCFB method*

Description

DMC identification via Bayesian functional regression models

Usage

```

findDMCFB(
  object,
  bwa,
  bwb,
  nBurn,
  nMC,
  nThin,
  alpha,
  sdv,
  nCores,
  pSize,
  sfiles
)

## S4 method for signature 'BSDMC'
findDMCFB(
  object,
  bwa,
  bwb,
  nBurn,
  nMC,
  nThin,
  alpha,
  sdv,
  nCores,
  pSize,
  sfiles
)

```

Arguments

object	A BSDMC-class object
bwa	An integer value specifying the band-width size of B-spline basis matrix for a natural cubic spline for the group-specific effects of the Bayesian functional regression model
bwb	An integer value specifying the band-width size of B-spline basis matrix for a natural cubic spline for the individual-specific effects of the Bayesian functional regression model
nBurn	An integer value specifying the number of burn-in samples
nMC	An integer value specifying the number of MCMC samples after burn-in
nThin	An integer value specifying the thinning number in MCMC
alpha	A numeric value specifying the level of α in credible interval $(1 - \alpha)\%$
sdv	An double value specifying the standard deviation of priors
nCores	An integer value specifying the number of machine cores for parallel computing
pSize	An integer value specifying the number of cytosines in a region to be used in a Bayesian functiona regression model for DMC detection
sfiles	A logical value indicating whether files to be saved or not.

Value

[BSDMC-class](#) object

Author(s)

Farhad Shokoohi <shokoohi@icloud.com>

Examples

```
set.seed(1980)
nr <- 1000
nc <- 4
metht <- matrix(as.integer(runif(nr * nc, 0, 100)), nr)
methc <- matrix(rbinom(n = nr * nc, c(metht), prob = runif(nr * nc)), nr, nc)
methl <- methc / metht
r1 <- GRanges(rep("chr1", nr), IRanges(1:nr, width = 1), strand = "*")
names(r1) <- 1:nr
cd1 <- DataFrame(
  Group = rep(c("G1", "G2"), each = nc / 2),
  row.names = LETTERS[1:nc]
)
OBJ1 <- cBSDMC(
  rowRanges = r1, methReads = methc, totalReads = metht,
  methLevels = methl, colData = cd1
)
OBJ2 <- findDMCFB(OBJ1,
  bwa = 10, bwb = 10, nBurn = 50, nMC = 50, nThin = 1,
  alpha = 0.05, nCores = 2, pSize = 500, sfiles = FALSE
```

```
)  
OBJ2
```

methLevels-method *methLevels method*

Description

Returns `methLevels` stored in [BSDMC-class](#)

Assigns `methLevels` to [BSDMC-class](#)

Usage

```
methLevels(object)  
  
methLevels(object) <- value  
  
## S4 method for signature 'BSDMC'  
methLevels(object)  
  
## S4 replacement method for signature 'BSDMC,matrix'  
methLevels(object) <- value
```

Arguments

<code>object</code>	A BSDMC-class object
<code>value</code>	An integer matrix

Value

A matrix
A [BSDMC-class](#) object

Author(s)

Farhad Shokoohi <shokoohi@icloud.com>

Examples

```
nr <- 150  
nc <- 8  
metht <- matrix(as.integer(runif(nr * nc, 0, 100)), nr)  
methc <- matrix(rbinom(n = nr * nc, c(metht), prob = runif(nr * nc)), nr, nc)  
methl <- methc / metht  
r1 <- GRanges(rep("chr1", nr), IRanges(1:nr, width = 1), strand = "*")  
names(r1) <- 1:nr  
cd1 <- DataFrame(  
  Group = rep(c("G1", "G2"), each = nc / 2),
```

```

    row.names = LETTERS[1:nc]
  )
OBJ1 <- cBSDMC(
  rowRanges = r1, methReads = methc, totalReads = metht,
  methLevels = methl, colData = cd1
)
methLevels(OBJ1)
methLevels(OBJ1) <- methl

```

methReads-method *methReads method*

Description

Returns `methReads` stored in [BSDMC-class](#)

Assigns `methReads` to [BSDMC-class](#)

Usage

```

methReads(object)

methReads(object) <- value

## S4 method for signature 'BSDMC'
methReads(object)

## S4 replacement method for signature 'BSDMC,matrix'
methReads(object) <- value

```

Arguments

object	A BSDMC-class object
value	An integer matrix

Value

A matrix
 A [BSDMC-class](#) object

Author(s)

Farhad Shokoohi <shokoohi@icloud.com>

Examples

```

nr <- 150
nc <- 8
metht <- matrix(as.integer(runif(nr * nc, 0, 100)), nr)
methc <- matrix(rbinom(n = nr * nc, c(metht), prob = runif(nr * nc)), nr, nc)
methl <- methc / metht
r1 <- GRanges(rep("chr1", nr), IRanges(1:nr, width = 1), strand = "*")
names(r1) <- 1:nr
cd1 <- DataFrame(
  Group = rep(c("G1", "G2"), each = nc / 2),
  row.names = LETTERS[1:nc]
)
OBJ1 <- cBSDMC(
  rowRanges = r1, methReads = methc, totalReads = metht,
  methLevels = methl, colData = cd1
)
methReads(OBJ1)
methReads(OBJ1) <- methc

```

params

params

Description

parameters name and their descriptions

Arguments

methReads	The matrix <code>methReads</code> contains the number of methylated reads spanning a CpG-site. The rows represent the CpG sites in <code>rowRanges</code> and the columns represent the samples in <code>colData</code> .
totalReads	The matrix <code>totalReads</code> contains the number of reads spanning a CpG-site. The rows represent the CpG sites in <code>rowRanges</code> and the columns represent the samples in <code>colData</code> .
methLevels	The matrix <code>methLevels</code> contains the predicted methylation level spanning a CpG-site using Bayesian functional regression models. The rows represent the CpG sites in <code>rowRanges</code> and the columns represent the samples in <code>colData</code> .
rowRanges	A GRanges or GRangesList object describing the ranges of interest. Names, if present, become the row names of the SummarizedExperiment object. The length of the GRanges or GRangesList must equal the number of rows of the matrices in assays. If <code>rowRanges</code> is missing, a SummarizedExperiment instance is returned.
colData	Object of class 'DataFrame' containing information on variable values of the samples
metadata	A list of storing MCMC samples or DMCs
object	A BSDMC-class object

value	An integer matrix
name	A character list
obj1	A BSDMC-class
obj2	A BSDMC-class
files	A character list
file	A character
nCores	An integer value specifying the number of machine cores for parallel computing
mc.cores	An integer greater than 0
pSize	An integer value specifying the number of cytosines in a region to be used in a Bayesian functional regression model for DMC detection
bwa	An integer value specifying the band-width size of B-spline basis matrix for a natural cubic spline for the group-specific effects of the Bayesian functional regression model
bwb	An integer value specifying the band-width size of B-spline basis matrix for a natural cubic spline for the individual-specific effects of the Bayesian functional regression model
nBurn	An integer value specifying the number of burn-in samples
nThin	An integer value specifying the thinning number in MCMC
nMC	An integer value specifying the number of MCMC samples after burn-in
sdv	An double value specifying the standard deviation of priors
alpha	A numeric value specifying the level of α in credible interval $(1 - \alpha)\%$
col	A character vector indicating which colors to alternate.
sfiles	A logical value indicating whether files to be saved or not.
region	An integer vector of length two specifying which subset of the object to be plotted
nSplit	A integer value specifying the number of subsets must be done for plotting the results of DMC identification
parList	A list specifying plots parameters, see par
...	other possible parameters

Author(s)

Farhad Shokoohi <shokoohi@icloud.com>

`plotDMCFB-method` *plotDMCFB method*

Description

Plotting the results of DMC identification stored in a [BSDMC-class](#) object

Usage

```
plotDMCFB(object, region, nSplit, parList)

## S4 method for signature 'BSDMC'
plotDMCFB(object, region, nSplit, parList)
```

Arguments

object	A BSDMC-class object
region	An integer vector of length two specifying which subset of the object to be plotted
nSplit	A integer value specifying the number of subsets must be done for plotting the results of DMC identification
parList	A list specifying plots parameters, see par

Value

Plot

Author(s)

Farhad Shokoohi <shokoohi@icloud.com>

Examples

```
set.seed(1980)
nr <- 1000
nc <- 4
metht <- matrix(as.integer(runif(nr * nc, 0, 100)), nr)
methc <- matrix(rbinom(n = nr * nc, c(metht), prob = runif(nr * nc)), nr, nc)
methl <- methc / metht
r1 <- GRanges(rep("chr1", nr), IRanges(1:nr, width = 1), strand = "*")
names(r1) <- 1:nr
cd1 <- DataFrame(
  Group = rep(c("G1", "G2"), each = nc / 2),
  row.names = LETTERS[1:nc]
)
OBJ1 <- cBSDMC(
  rowRanges = r1, methReads = methc, totalReads = metht,
  methLevels = methl, colData = cd1
```

```

)
OBJ2 <- findDMCFB(OBJ1,
  bwa = 10, bwb = 10, nBurn = 50, nMC = 50, nThin = 1,
  alpha = 0.05, nCores = 2, pSize = 500, sfiles = FALSE
)
plotDMCFB(OBJ2)

```

readBismark-method *readBismark method*

Description

reads BS-Seq data

Usage

```

readBismark(files, colData, mc.cores)

## S4 method for signature 'character,DataFrame,numeric'
readBismark(files, colData, mc.cores)

## S4 method for signature 'character,data.frame,numeric'
readBismark(files, colData, mc.cores)

## S4 method for signature 'character,character,numeric'
readBismark(files, colData, mc.cores)

```

Arguments

files	A character list
colData	Object of class 'DataFrame' containing information on variable values of the samples
mc.cores	An integer greater than 0

Value

A [BSDMC-class](#) object

Author(s)

Farhad Shokoohi <shokoohi@icloud.com>

Examples

```
fn <- list.files(system.file("extdata", package = "DMCFB"))
fn.f <- list.files(system.file("extdata", package = "DMCFB"),
  full.names = TRUE
)
OBJ <- readBismark(fn.f, fn, mc.cores=1)
cdOBJ <- DataFrame(Cell = factor(c("BC", "TC", "Mono"),
  labels = c("BC", "TC", "Mono"))
), row.names = c("BCU1568", "BCU173", "BCU551"))
colData(OBJ) <- cdOBJ
OBJ
```

totalReads-method *totalReads method*

Description

Returns totalReads stored in [BSDMC-class](#)
Assigns totalReads to [BSDMC-class](#)

Usage

```
totalReads(object)

totalReads(object) <- value

## S4 method for signature 'BSDMC'
totalReads(object)

## S4 replacement method for signature 'BSDMC,matrix'
totalReads(object) <- value
```

Arguments

object	A BSDMC-class object
value	An integer matrix

Value

A matrix
A [BSDMC-class](#) object

Author(s)

Farhad Shokoohi <shokoohi@icloud.com>

Examples

```
nr <- 150
nc <- 8
metht <- matrix(as.integer(runif(nr * nc, 0, 100)), nr)
methc <- matrix(rbinom(n = nr * nc, c(metht), prob = runif(nr * nc)), nr, nc)
methl <- methc / metht
r1 <- GRanges(rep("chr1", nr), IRanges(1:nr, width = 1), strand = "*")
names(r1) <- 1:nr
cd1 <- DataFrame(
  Group = rep(c("G1", "G2"), each = nc / 2),
  row.names = LETTERS[1:nc]
)
OBJ1 <- cBSDMC(
  rowRanges = r1, methReads = methc, totalReads = metht,
  methLevels = methl, colData = cd1
)
totalReads(OBJ1)
totalReads(OBJ1) <- metht
```

Index

* **object**
 BSDMC-class, 3

 BSDMC (BSDMC-class), 3
 BSDMC-class, 3

 cBSDMC, 2
 cBSDMC (cBSDMC-method), 4
 cBSDMC,matrix,matrix,matrix,GRanges-method
 (cBSDMC-method), 4
 cBSDMC-method, 4
 combine (combine-method), 6
 combine,BSDMC,BSDMC-method
 (combine-method), 6
 combine-method, 6

 DMCFB (DMCFB-package), 2
 DMCFB-package, 2

 findDMCFB, 2
 findDMCFB (findDMCFB-method), 7
 findDMCFB,BSDMC-method
 (findDMCFB-method), 7
 findDMCFB-method, 7

 GRanges, 5, 11
 GRangesList, 5, 11

 methLevels (methLevels-method), 9
 methLevels,BSDMC-method
 (methLevels-method), 9
 methLevels-method, 9
 methLevels<- (methLevels-method), 9
 methLevels<-,BSDMC,matrix-method
 (methLevels-method), 9
 methReads (methReads-method), 10
 methReads,BSDMC-method
 (methReads-method), 10
 methReads-method, 10
 methReads<- (methReads-method), 10

 methReads<-,BSDMC,matrix-method
 (methReads-method), 10

 par, 12, 13
 params, 11
 plotDMCFB, 2
 plotDMCFB (plotDMCFB-method), 13
 plotDMCFB,BSDMC-method
 (plotDMCFB-method), 13
 plotDMCFB-method, 13

 readBismark, 2
 readBismark (readBismark-method), 14
 readBismark,character,character,numeric-method
 (readBismark-method), 14
 readBismark,character,data.frame,numeric-method
 (readBismark-method), 14
 readBismark,character,DataFrame,numeric-method
 (readBismark-method), 14
 readBismark-method, 14

 SummarizedExperiment, 5, 11

 totalReads (totalReads-method), 15
 totalReads,BSDMC-method
 (totalReads-method), 15
 totalReads-method, 15
 totalReads<- (totalReads-method), 15
 totalReads<-,BSDMC,matrix-method
 (totalReads-method), 15