

Package ‘methFuse’

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Title Functional Segmentation of the Methylome

Version 1.1.0

Description

Implements FUSE (Functional Segmentation of DNA methylation data), a data-driven method for identifying spatially coherent DNA methylation segments from whole-genome bisulfite sequencing (WGBS) count data. The method performs hierarchical clustering of CpG sites based on methylated and unmethylated read counts across multiple samples and determines the optimal number of segments using an information criterion (AIC or BIC). Resulting segments represent regions with homogeneous methylation profiles across the input cohort while allowing sample-specific methylation levels. The package provides functions for clustering, model selection, tree cutting, segment-level summarization, and visualization. Input can be supplied as count matrices or extracted directly from 'BSseq' and 'methrix' objects.

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Imports stats, methods

Suggests bsseq, methrix, beachmat, GenomicRanges,
SummarizedExperiment, DelayedArray, testthat (>= 3.0.0), knitr,
rmarkdown, graphics

VignetteBuilder knitr

Config/testthat/edition 3

URL <https://holmsusa.github.io/methFuse/>

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Contents

| | |
|--------------------|---|
| fuse.cluster | 2 |
| fuse.cut.tree | 3 |
| fuse.segment | 4 |
| fuse.summary | 5 |
| number.of.clusters | 6 |
| plot.fuse_summary | 7 |

| | |
|--------------|----------|
| Index | 9 |
|--------------|----------|

| | |
|--------------|--|
| fuse.cluster | <i>Perform Hierarchical Clustering on Methylation Data</i> |
|--------------|--|

Description

Produces a hierarchical clustering tree based on the input matrices of counts.

Usage

```
fuse.cluster(x, ...)
```

Default S3 method:
 fuse.cluster(x, K1, chr = NULL, pos = NULL, ...)

S3 method for class 'BSseq'
 fuse.cluster(x, ...)

S3 method for class 'methrix'
 fuse.cluster(x, ...)

Arguments

| | |
|-----|--|
| x | Input object. One of: matrix Unmethylated count matrix (K0). BSseq A BSseq object. methrix A methrix object. |
| ... | Additional arguments if K0 is a matrix. |
| K1 | Methylated count matrix (if x is matrix). |
| chr | Chromosome labels (if x is matrix). |
| pos | Genomic positions (if x is matrix). |

Value

A clustering tree of class hclust.

Examples

```
# Example: Clustering generated data
set.seed(1234)
K0 <- matrix(
  rep(c(sample(0:20, 200, replace = TRUE), sample(20:40, 200, replace = TRUE)), 2),
  nrow = 100, byrow = TRUE
)
K1 <- matrix(
  rep(c(sample(20:40, 200, replace = TRUE), sample(0:20, 200, replace = TRUE)), 2),
  nrow = 100, byrow = TRUE
)
tree <- fuse.cluster(K0, K1)
tree
```

fuse.cut.tree

Cut Hierarchical Clustering Tree into Clusters

Description

Divides the clustering tree into a specified number of clusters.

Usage

```
fuse.cut.tree(tree, k)
```

Arguments

| | |
|------|----------------------------------|
| tree | Clustering tree of class hclust. |
| k | Number of clusters |

Value

A vector indicating which cluster each element in the original data frame belonged to

Examples

```
# Example: Cutting small tree in 2 segments
set.seed(1234)
K0 <- matrix(
  rep(c(sample(0:20, 200, replace = TRUE), sample(20:40, 200, replace = TRUE)), 2),
  nrow = 100, byrow = TRUE
)
K1 <- matrix(
  rep(c(sample(20:40, 200, replace = TRUE), sample(0:20, 200, replace = TRUE)), 2),
  nrow = 100, byrow = TRUE
)
tree <- fuse.cluster(K0, K1)
```

```
segments <- fuse.cut.tree(tree, 2)
segments
```

| | |
|--------------|--|
| fuse.segment | <i>Full FUSE segmentation pipeline</i> |
|--------------|--|

Description

Performs the full FUSE segmentation workflow: hierarchical clustering, model selection, tree cutting, and genomic segment summarization.

Usage

```
fuse.segment(x, ...)
```

Default S3 method:

```
fuse.segment(x, K1, chr, pos, method = c("BIC", "AIC"), ...)
```

S3 method for class 'BSseq'

```
fuse.segment(x, method = c("BIC", "AIC"), ...)
```

S3 method for class 'methrix'

```
fuse.segment(x, method = c("BIC", "AIC"), ...)
```

Arguments

| | |
|--------|--|
| x | Input object. One of: matrix Unmethylated count matrix (K0). BSseq A BSseq object. methrix A methrix object. |
| ... | Additional arguments (matrix input only) |
| K1 | Methylated count matrix (if x is matrix). |
| chr | Chromosome labels (if x is matrix). |
| pos | Genomic positions (if x is matrix). |
| method | Information criterion for model selection: "BIC" (default) or "AIC". For internal use, x corresponds to the unmethylated count matrix (K0). |

Details

fuse.segment() is an S3 generic with methods for:

matrix Raw count matrices (K0, K1) with genomic annotation.

BSseq Bioconductor BSseq objects.

methrix Bioconductor methrix objects (supports DelayedMatrix).

Value

An object of class `fuse_summary`, containing:

summary Data frame with one row per genomic segment.

betas_per_segment Matrix of per-sample methylation estimates.

raw_beta Per-CpG methylation estimates.

raw_pos Genomic positions of CpGs.

Automatic data extraction

For BSseq objects:

- Methylated counts are obtained via `getCoverage(x, "M")`
- Unmethylated counts via `getCoverage(x, "Cov") - M`
- Chromosome and position from `rowRanges(x)`

For `methrix` objects:

- Methylated counts via `get_matrix(x, "M")`
- Total coverage via `get_matrix(x, "C")`
- Unmethylated counts computed as `C - M`
- Genomic coordinates extracted from locus metadata

`fuse.summary`

Summarize FUSE Segmentation Results

Description

Summarizes FUSE segmentation results into one row per segment, including genomic coordinates, CpG count, segment length, average methylation (beta), and stability flag based on likelihood testing. Also returns per-sample methylation estimates for each segment. Result can be visualized using `plot(result)`.

Usage

```
fuse.summary(K0, K1, chr, pos, segments)
```

Arguments

| | |
|-----------------------|--|
| <code>K0</code> | Integer or numeric matrix of unmethylated counts. |
| <code>K1</code> | Integer or numeric matrix of methylated counts. |
| <code>chr</code> | Character vector giving the chromosome for each site. |
| <code>pos</code> | Numeric vector giving genomic coordinates for each site. |
| <code>segments</code> | Integer vector giving segment IDs for each site in <code>K0</code> and <code>K1</code> . |

Value

A list with four elements:

summary A data frame with one row per segment and the following columns:

- Segment: Segment ID
- Chr: Chromosome
- Start: Start genomic coordinate
- End: End genomic coordinate
- CpGs: Number of CpGs in the segment
- Length: Genomic length (End - Start + 1)
- Beta: Average methylation across samples and CpGs
- Coherent: Logical indicator (TRUE if segment is coherently methylated, else FALSE)

betas_per_segment Matrix of per-sample methylation estimates for each segment (rows = segments, columns = samples).

raw_beta Average beta per CpG site, used for plotting.

raw_pos Genomic position for every given CpG site, used for plotting.

Examples

```
set.seed(1234)
K0 <- matrix(
  rep(c(sample(0:20, 200, replace = TRUE), sample(20:40, 200, replace = TRUE)), 2),
  nrow = 100, byrow = TRUE
)
K1 <- matrix(
  rep(c(sample(20:40, 200, replace = TRUE), sample(0:20, 200, replace = TRUE)), 2),
  nrow = 100, byrow = TRUE
)
tree <- fuse.cluster(K0, K1)
segments <- fuse.cut.tree(tree, 4)
res <- fuse.summary(K0, K1, rep("chr1", nrow(K0)), 1:nrow(K0), segments)
head(res$summary)
head(res$betas_per_segment)
```

number.of.clusters *Find Optimal Number of Clusters*

Description

Determines the optimal number of clusters to cut a hierarchical clustering tree, based on the selected information criterion (e.g., BIC or AIC).

Usage

```
number.of.clusters(tree, n, method = c("BIC", "AIC"))
```

Arguments

| | |
|--------|--|
| tree | Clustering tree of class hclust. |
| n | Number of samples in the original data. |
| method | Information criterion method. One of "BIC" or "AIC". |

Value

An integer representing the optimal number of clusters.

Examples

```
# Example: Determine number of clusters in dummy data set
set.seed(1234)
K0 <- matrix(
  rep(c(sample(0:20, 200, replace = TRUE), sample(20:40, 200, replace = TRUE)), 2),
  nrow = 100, byrow = TRUE
)
K1 <- matrix(
  rep(c(sample(20:40, 200, replace = TRUE), sample(0:20, 200, replace = TRUE)), 2),
  nrow = 100, byrow = TRUE
)
tree <- fuse.cluster(K0, K1)
k <- number.of.clusters(tree, ncol(K0), 'BIC')
k
```

plot.fuse_summary *Plot method for FUSE segmentation results*

Description

Plotting method for fuse_summary.

Usage

```
## S3 method for class 'fuse_summary'
plot(x, ..., segments_to_plot = 1:50)
```

Arguments

| | |
|------------------|-----------------------------------|
| x | A fuse_summary object |
| ... | Additional arguments |
| segments_to_plot | Integer vector of segment indices |

Details

Raw CpG-level methylation values are shown as grey points. Segment-level methylation is shown as horizontal bars (red = hypermethylated, blue = hypomethylated).

Value

No return value, called for side effects.

Index

`fuse.cluster`, [2](#)

`fuse.cut.tree`, [3](#)

`fuse.segment`, [4](#)

`fuse.summary`, [5](#)

`number.of.clusters`, [6](#)

`plot.fuse_summary`, [7](#)