# Package 'msgbsR'

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|---|
| Title msgbsR: methylation sensitive genotyping by sequencing (MS-GBS)<br>R functions  |
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| <b>biocViews</b> ImmunoOncology, DifferentialMethylation, DataImport,<br>Epigenetics, MethylSeq   |
| Description Pipeline for the analysis of a MS-GBS experiment.   |
| License GPL-2   |
| LazyLoad yes  |
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## checkCuts

## Contents

| checkCuts  |   |
|------------|---|
| cuts       | 3 |
| diffMeth   | 4 |
| msgbsR     | 4 |
| plotCircos | 5 |
| plotCounts | 6 |
| ratdata    |   |
| ratdata2   | 7 |
| rawCounts  | 8 |
|            |   |
|            | 9 |

## Index

checkCuts

checkCuts

## Description

Determines the sequence around a cut site using a fasta file or BSgenome

## Usage

checkCuts(cutSites, genome, fasta = FALSE, seq)

## Arguments

| cutSites | A GRanges object containing the locations of the cut sites to be checked for se-<br>quence match. The names of the correct cut sites will be returned as a GRanges<br>object. |
|----------|---|
| genome   | The path to a fasta file or a BSgenome object to check for genomic sequences.   |
| fasta    | TRUE if a fasta file has been supplied. Default = FALSE   |
| seq      | The desired recognition sequence that the enzyme should have cut.   |

## Value

A GRanges object containing the names of the sites that had the correct sequence.

## Author(s)

Benjamin Mayne

2

cuts

#### Examples

cuts

A GRanges object of differentially methylated MspI cut sites on chromosome 20 in Rat from a MS-GBS experiment.

#### Description

The GRanges object was created from a list of differentially methylated cut sites from a MS-GBS experiment between two groups of rats that were fed either a control diet or a high fat diet.

#### Usage

data(cuts)

## Format

A GRanges object of length 10.

#### Details

• Positions of MspI cut sites differentially methylated in the prostate on chromosome 20 in Rats.

The data set contains 10 differentially methylated sites in the prostate between rats fed a control or high fat diet.

#### Value

A GRanges object of length 10.

diffMeth

#### Description

Determines differential methylated sites from a RangedSummarizedExperiment

#### Usage

#### Arguments

| se               | A RangedSummarizedExperiment containing meta data of the samples.  |  |
|------------------|--|--|
| cateogory        | The heading name in the sample data to be tested for differential methylation.   |  |
| condition1       | The reference group within the cateogory.  |  |
| condition2       | The experimental group within the cateogory.   |  |
| block            | The heading name in the sample data if differential methylation is to be tested with a blocking factor. Default is NULL. |  |
| cpmThreshold     | Counts per million threshold of read counts to be filtered out of the analysis.  |  |
| thresholdSamples |  |  |
|                  | Minimum number of samples to contain the counts per million threshold.   |  |

#### Value

A data frame containing which cut sites that are differenitally methylated.

#### Author(s)

Benjamin Mayne

## Examples

msgbsR

msgbsR

## Description

msgbsR

plotCircos

plotCircos

#### Description

Plot a circos representing the cut site locations

#### Usage

```
plotCircos(cutSites, seqlengths, cutSite.colour, seqlengths.colour)
```

## Arguments

| cutSites          | A GRanges object containing the locations of the cut sites to be plotted. |  |
|-------------------|---|--|
| seqlengths        | An integer with the lengths of the chromosomes.                           |  |
| cutSite.co        | Lour The colour of the cut sites.   |  |
| seqlengths.colour |   |  |
|                   | The colour of the chromosomes   |  |

### Value

A circos plot showing the locations of the cut sites.

## Author(s)

Benjamin Mayne

#### Examples

plotCounts

## Description

Plots the total number of reads vs total number of cut sites per sample

## Usage

```
plotCounts(se, cateogory)
```

#### Arguments

| se        | A RangedSummarizedExperiment containing meta data of the samples. |
|-----------|---|
| cateogory | The heading name in the sample data to distinguish groups.        |

#### Value

Produces a plot showing the total number reads vs total number of cut sites per sample.

#### Author(s)

Benjamin Mayne

#### Examples

```
data(ratdata2)
plotCounts(se = ratdata2, cateogory = "Group")
```

| ratdata | Read counts of potential MspI cut sites from a MS-GBS experiment of |
|---------|---|
|         | prostates from rats   |

## Description

A RangedSummarizedExperiment containing read counts generated from a MS-GBS experiment using the restriction enzyme MspI, focusing on chromosome 20 of Rat.

#### Usage

```
data(ratdata)
```

## Format

RangedSummarizedExperiment

#### ratdata2

#### Details

• ratdata A RangedSummarizedExperiment with 16047 potential MspI cut sites on chromosome 20 in Rat and six samples (3 Control and 3 Experimental).

This dataset contains six prostate samples from rats: 3 control and 3 experimental high fat diet.

#### Value

RangedSummarizedExperiment

ratdata2

*Read counts of correct MspI cut sites from a MS-GBS experiment of prostates from rats* 

#### Description

A RangedSummarizedExperiment containing read counts generated from a MS-GBS experiment using the restriction enzyme MspI, focusing on chromosome 20 of Rat. The sites have been checked for the correct recognition site.

#### Usage

data(ratdata2)

## Format

RangedSummarizedExperiment

#### Details

 ratdata2 A RangedSummarizedExperiment containing data for 13983 MspI cut sites on chromosome 20 in Rat and six samples (3 Control and 3 Experimental).

This dataset contains six prostate samples from rats: 3 control and 3 experimental high fat diet. The data can be used for differential methylation analyses.

#### Value

RangedSummarizedExperiment

rawCounts

#### Description

Imports the raw read counts from sorted and indexed bam file(s)

## Usage

```
rawCounts(bamFilepath, threads = 1)
```

## Arguments

| bamFilepath | The path to the location of the bam file(s).                 |
|-------------|--|
| threads     | The total number of usable threads to be used. Default is 1. |

## Value

Produces a RangedSummarizedExperiment. Columns are samples and the rows are cut sites. The cut site IDs are in the format chr:position-position:strand.

### Author(s)

Benjamin Mayne, Sam Buckberry

## Examples

```
my_path <- system.file("extdata", package = "msgbsR")
my_data <- rawCounts(bamFilepath = my_path)</pre>
```

## Index

\* datasets cuts, 3 ratdata, 6 ratdata2, 7 checkCuts, 2 cuts, 3 diffMeth, 4 msgbsR, 4 msgbsR-package (msgbsR), 4 plotCircos, 5 plotCounts, 6 ratdata, 6 ratdata, 6 ratdata, 7

ratdata2,7 rawCounts,8