## Package 'msgbsR'

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Type Package

**Title** msgbsR: methylation sensitive genotyping by sequencing (MS-GBS)
R functions

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**Depends** R (>= 3.4), GenomicRanges, methods

Imports BSgenome, easyRNASeq, edgeR, GenomicAlignments, GenomicFeatures, GenomeInfoDb, ggbio, ggplot2, IRanges, parallel, plyr, Rsamtools, R.utils, stats, SummarizedExperiment, S4Vectors, utils

Suggests roxygen2, BSgenome.Rnorvegicus.UCSC.rn6

**biocViews** ImmunoOncology, DifferentialMethylation, DataImport, Epigenetics, MethylSeq

**Description** Pipeline for the analysis of a MS-GBS experiment.

License GPL-2

LazyLoad yes

**Collate** 'msgbsR.R' 'rawCounts.R' 'checkCuts.R' 'plotCounts.R' 'diffMeth.R' 'plotCircos.R'

RoxygenNote 5.0.1

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2 checkCuts

## **Contents**

	checkCuts	 																	2
	cuts	 																	3
	diffMeth .	 								 									4
	msgbsR .	 								 									4
	plotCircos	 																	5
	plotCounts																		
	ratdata	 																	6
	ratdata2 .	 																	7
	rawCounts																		8
Index																			9
-														_	_				

Description

checkCuts

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

checkCuts

## Usage

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

## Arguments

cutSites	A GRanges object containing the locations of the cut sites to be checked for sequence match. The names of the correct cut sites will be returned as a GRanges 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

cuts 3

#### **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.

4 msgbsR

#### **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 5

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## **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.colour 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**

6 ratdata

plotCounts

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 7

## **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**

Ranged Summarized Experiment

#### **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

8 rawCounts

rawCounts 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
ratdata2, 7
rawCounts, 8
```