# Package 'methylaction'

March 19, 2015

2 getCounts

```
methylaction-package
```

methylaction: Differentially Methylated Regions (DMRs) from MBD-isolated Genome Sequencing (MiGS/MBD-seq)

## Description

 $methylaction: \ Differentially \ Methylated \ Regions \ (DMRs) \ from \ MBD-isolated \ Genome \ Sequencing \ (MiGS/MBD-seq)$ 

getCounts	Produce a GRanges with counts of overlapping reads for a set of
	ranges

#### **Description**

Will return a GenomicRanges object for non-overlapping windows genome-wide for the genome given as a bsgenome object for the chromosomes given in chrs. The values() of the GRanges will contain a table of counts for each sample at each window.

## Usage

```
getCounts(samp, reads, bsgenome = NULL, ranges = NULL, chrs = NULL,
winsize = 50, ncore = 1)
```

## Arguments

samp	Sample data.frame from readSampleInfo()
bsgenome	
ranges	
winsize	Size of the non-overlapping windows.
fragsize	Average fragment length from the sequencing experiment. Reads will be extended up to this size when computing coverage.

## Value

A GenomicRanges object with values() containing a table of counts for each sample at each window.

getCountsFc 3

getCountsFc	Get counts using featureCounts	
-------------	--------------------------------	--

#### **Description**

desc

#### Usage

```
getCountsFc(samp, bsgenome = NULL, ranges = NULL, chrs = NULL, fragsize,
  winsize = 50, ncore = 1)
```

## **Arguments**

samp Sample data.frame from readSampleInfo()

bsgenome chrs

fragsize Average fragment length from the sequencing experiment. Reads will be ex-

tended up to this size when computing coverage.

#### Value

desc

getReads	Store GenomicRanges of BAM reads in a list

## Description

Will return a list of GenomicRanges objects

#### Usage

```
getReads(samp, bsgenome, chrs, fragsize, ncore)
```

## Arguments

samp Sample data.frame from readSampleInfo()
bsgenome
chrs
fragsize Average fragment length from the sequencing experiment. Reads will be extended up to this size when computing coverage.

#### Value

A list of GenomicRanges objects.

4 maClustering

mased write BED file of DMK region	maBed	Write BED file of DMR region
------------------------------------	-------	------------------------------

#### **Description**

Creates a BED file suitable for uploading as a custom track to the UCSC genome browser.

## Usage

```
maBed(ma, file, dmr.only = F)
```

## **Arguments**

ma Output list from a run of methylaction()

file Name of BED file to create

dmr.only Don't report regions without a significant pattern (show up listed as NS, default:

FALSE)

#### Value

Writes BED file to disk.

maClustering Clustering of samples based on read counts

# Description

Will plot clustering based on some subset of read counts.

## Usage

```
maClustering(samp, ma, mincv, type, pdf)
```

## Arguments

samp Description of samples from readSampleInfo()

ma Output object from methylaction()

mincv Minimum cv (cv=mean/sd) for a window to be included in the clustering.

type One of "mds" or "hca" pdf PDF file to output.

#### Value

Saves plot to disk.

maCompare 5

maCompare	Compare between various runs of methylaction()
-----------	--

## Description

Reports shared and non-shared regions detected by different runs of methylaction(). Useful for comparing between studies/cohorts and paramater tuning.

#### Usage

```
maCompare(malist)
```

#### **Arguments**

malist

List of output objects from methylaction(). Set the names() attribute of this list to unique descriptive names for each run.

#### Value

A list containing both site by site comparisions and a data.frame summary table of total and shared sites for each pairwise comparision between runs.

maHeatmap of the differentially methylated regions run of methylaction()	(DMRs) found by a
--	-------------------

## Description

Will plot a heatmap of the DMRs based on the normalized read counts.

## Usage

```
maHeatmap(samp, ma, pdf)
```

## **Arguments**

samp Description of samples from readSampleInfo()

ma Output object from methylaction()

pdf PDF file to output.

#### Value

Saves plot to disk.

6 maTable

maSummary

Summary stats for a run of methylaction()

# Description

Will return information about number of windows/regions that pass cutoffs at each stage of the analysis. Useful for paramater tuning.

# Usage

```
maSummary (ma)
```

## **Arguments**

ma

Output object from methylaction()

#### Value

A data.frame with the summary statistics.

maTable

Table of differentially methylated regions (DMRs) by pattern

# Description

Will return a table of which patterns were detected and the number of DMRs in each.

#### Usage

```
maTable(samp, ma)
```

## Arguments

samp Description of samples from readSampleInfo()

ma Output object from methylaction()

#### Value

A data frame with the summary statistics.

maTracks 7

	maTracks	Write BED and BIGWIG files for normalized, filter-passed window count values
--	----------	--

## Description

Creates a BED file suitable for uploading as a custom track to the UCSC genome browser.

## Usage

```
maTracks(ma, path = ".", bigwig = FALSE, chrs = NULL, bsgenome = NULL,
    ncore = NULL)
```

#### **Arguments**

ma	Output list from a run of methylaction()
path	Folder to save the files in (defulat: current working directory)
bigwig	Convert to BIGWIG files, requires wigToBigWig in \$PATH (default: FALSE)
chrs	
bsgenome	

#### Value

ncore

Writes BED file to disk.

Detect differentially methylated regions (DMRs) from windowed read counts from MBD-isolated genome sequencing (MiGS/MBD-seq)

# Description

Once the counts have been pre-processed, this function performs all the analysis. Detailed results from intermediate steps are stored in the output list object to analyze method performance and provide input for the summary and plotting functions.

## Usage

```
methylaction(samp, counts, winsize, poifdr, stageone.p, joindist, anodev.p,
   post.p, minsize = 150, nperms = 0, ncore = 1)
```

8 readSampleInfo

#### **Arguments**

samp	Description of samples from readSampleInfo()
counts	Preprocessed count data from getCounts()
winsize	Size of the windows used when counting.
poifdr	False discovery rate to use during initital filtering.
stageone.p	P-value cutoff for stage one testing.
anodev.p	P-value cutoff for the analysis of deviance (ANODEV) in stage two testing (ignored for two group comparisions).
post.p	P-value cutoff for post-tests (or for the single test stage two test in the two group case).
minsize	Minimum size for a reported region.
ncore	Number of cores to use.
bsgenome	b-string genome (bsgenome) object for the genome
fragsize	The average fragment length selected for in the sequencing experiment (used to

extend reads when re-counting for regions in stage two testing).

#### Value

A list containing detailed results from each stage of the analysis.

readSampleInfo	Load a CSV containing required information about each sample
----------------	--

## Description

The CSV file must contain the following columns: "sample" - unique sample IDs, "group" - group IDs, "bam" - path to BAM file containing aligned reads for the sample. Columns with other names will be ignored. Note that in subsequent reporting of pattern strings (where each digit represents a group), digits for each group will be ordered in the order they first appear in this samplesheet.

#### Usage

```
readSampleInfo(file = NULL, colors = NULL)
```

#### **Arguments**

file	Path to the CSV samplesheet to open. Must contain the columns described above.
colors	Vector of colors (one for each group) in same order as groups appear in the sample file. These will be uniform colors used in the plotting functions for these groups. Give colors as hex codes. If none provided they will be auto-selected with RColorBrewer. If there is a column named "color" in the CSV, then this will always be used.

#### Value

A data.frame of the samplesheet that will be valid input to the other function's "samp" arguments.