

# Package ‘exomePeak2’

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**Type** Package

**Title** Bias Awareed Peak Calling and Quantification for MeRIP-Seq

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## Description

exomePeak2 provides bias awareed quantification and peak detection on Methylated RNA immunoprecipitation sequencing data (MeRIP-Seq). MeRIP-Seq is a commonly applied sequencing technology to measure the transcriptome-wide location and abundance of RNA modification sites under a given cellular condition. However, the quantification and peak calling in MeRIP-Seq are sensitive to PCR amplification bias which is prevalent in next generation sequencing (NGS) techniques. In addition, the RNA-Seq based count data exhibits biological variation in small reads count. exomePeak2 collectively address these challenges by introducing a rich set of robust data science models tailored for MeRIP-Seq. With exomePeak2, users can perform peak calling, modification site quantification, and differential analysis with a straightforward one-step function. Alternatively, users could define personalized methods for their own analysis through multi-step functions and diagnostic plots.

**License** GPL (>= 2)

**BugReports** <https://github.com/ZW-xjtlu/exomePeak2/issues>

## Imports

Rsamtools, GenomicAlignments, GenomicRanges, GenomicFeatures, DESeq2, ggplot2, mclust, genefilter, Biostrings, BSgenome

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call_peaks_with_GLM	<i>Statistical Inference with DESeq package based on the provided reads count for exomic bins.</i>
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**Description**

call\_peaks\_with\_GLM conduct inference on every exome bins using negative binomial model, the significant bins will be the merged into peaks.

**Usage**

```
call_peaks_with_GLM(
  SE_bins,
  glm_type = c("Poisson", "NB", "DESeq2"),
  correct_GC_bg = TRUE,
  qtnorm = TRUE,
  txdb,
  count_cutoff = 5,
  p_cutoff = NULL,
  p_adj_cutoff = 0.05,
  log2FC_cutoff = 0,
  consistent_peak = TRUE,
  consistent_log2FC_cutoff = 0,
  consistent_fdr_cutoff = 0.05,
  alpha = 0.05,
```

```

    p0 = 0.8
  )

```

### Arguments

SE_bins	a SummarizedExperiment object. The meta-data column should contain the design information of IP/input + treated/control.
glm_type	a character, which can be one of the "Poisson", "NB", and "DESeq2". This argument specifies the type of generalized linear model used in peak calling; Default to be "Poisson". The DESeq2 method is only recommended for high power experiments with more than 3 biological replicates for both IP and input.
correct_GC_bg	a logical value of whether to estimate the GC content linear effect on background regions; default = TRUE. If correct_GC_bg = TRUE, it could result in a more accurate estimation of the technical effect of GC content for the RNA modifications that are highly biologically related to GC content.
qtnorm	a logical of whether to perform subset quantile normalization after the GC content linear effect correction; default = TRUE. Subset quantile normalization will be applied within the IP and input samples separately to account for the inherent differences between the marginal distributions of IP and input samples.
txdb	the txdb object that is necessary for the calculation of the merge of the peaks.
count_cutoff	an integer value indicating the cutoff of the mean of reads count in a row, inference is only performed on the windows with the row average read count bigger than the cutoff. Default value is 5.
p_cutoff	a numeric value of the p value cutoff used in DESeq inference.
p_adj_cutoff	a numeric value of the adjusted p value cutoff used in DESeq2 inference; if provided, the value of p_cutoff will be ignored; Default = 0.05.
log2FC_cutoff	a non negative numeric value of the log2 fold change (log2 IP/input) cutoff used in the inference of peaks.
consistent_peak	a logical of whether the positive peaks returned should be consistent among replicates; default = TRUE.
consistent_log2FC_cutoff	a numeric for the modification log2 fold changes cutoff in the peak consistency calculation; default = 0.
consistent_fdr_cutoff	a numeric for the BH adjusted C-test p values cutoff in the peak consistency calculation; default = 0.05. Check <a href="#">ctest</a> .
alpha	a numeric for the binomial quantile used in the consistent peak filter; default = 0.05.
p0	a numeric for the binomial proportion parameter used in the consistent peak filter; default = 0.8. For a peak to be consistently methylated, the minimum number of significant enriched replicate pairs is defined as the 1 - alpha quantile of a binomial distribution with $p = p_0$ and $N =$ number of possible pairs between replicates. The consistency defined in this way is equivalent to the rejection of an exact binomial test with null hypothesis of $p < p_0$ and $N =$ replicates number of IP * replicates number of input.

**Details**

call\_peaks\_with\_GLM will perform exome level peak calling using DESeq2 model,

The significant bins will be merged into modification peaks.

The insignificant bins (pass the row means filtering) will also be merged into control peaks.

**Value**

This function will return a list of GRangesList object storing peaks for both modification and control.

---

estimateSeqDepth-methods

*Method estimateSeqDepth*

---

**Description**

estimateSeqDepth estimate sequencing depth size factors for each MeRIP-seq samples used in peak statistics quantification. Under default setting, the sequencing depth are estimated by the robust estimator defined in package DESeq. i.e. the median of the ratios to the geometric means of peak counts.

**Usage**

```
estimateSeqDepth(sep, from = c("Background", "Modification", "All"), ...)
```

```
## S4 method for signature 'SummarizedExomePeak'
estimateSeqDepth(sep, from = c("Background", "Modification", "All"), ...)
```

**Arguments**

sep a [SummarizedExomePeak](#) object.

from a character specify the subset of features for sequencing depth estimation, can be one of c("Background", "Modification", "All").

Background The sequencing depths are estimated from the background control regions. This method could make the IP/input LFC estimates become closer to the true modification proportion.

Modification The sequencing depths are estimated from the modification peaks/sites.

All The sequencing depths are estimated from both the background and the modification regions.

Under the default settings, the sequencing depth size factors are estimated from the background control regions.

... inherited from [estimateSizeFactorsForMatrix](#).

**Details**

The function takes the input of a [SummarizedExomePeak](#) object, and it estimates the sequencing depth size factors by the columns of its [assay](#).

## Value

This function will return a [SummarizedExomePeak](#) object containing newly estimated sequencing depth size factors.

## See Also

[normalizeGC](#)

## Examples

```
### Load the example SummarizedExomPeak object
f1 = system.file("extdata", "sep_ex_mod.rds", package="exomePeak2")

sep <- readRDS(f1)

### Estimate the sequencing depth size factors
sep <- estimateSeqDepth(sep)

sep$sizeFactor
```

---

exomePeak2

*Peak Calling and Peak Statistics Quantification on MeRIP-seq Dataset.*

---

## Description

exomePeak2 conducts peak calling and peak statistics calculation from **BAM** files of a MeRIP-seq experiment. The function integrates the following steps of a standard MeRIP-seq data analysis pipeline.

1. Check and index the BAM files with [scanMeripBAM](#).
2. Call modification peaks on exons with [exomePeakCalling](#).
3. Calculate offset factors of GC content biases with [normalizeGC](#).
4. Calculate (differential) modification statistics with the generalized linear model (GLM) using [glmM](#) or [glmDM](#).
5. Export the peaks/sites statistics with user defined format by [exportResults](#).

See the help pages of the corresponding functions for the complete documentation.

## Usage

```
exomePeak2(
  bam_ip = NULL,
  bam_input = NULL,
  bam_treated_ip = NULL,
  bam_treated_input = NULL,
  txdb = NULL,
  bsgenome = NULL,
  genome = NA,
```

```

gff_dir = NULL,
mod_annot = NULL,
paired_end = FALSE,
library_type = c("unstranded", "1st_strand", "2nd_strand"),
fragment_length = 100,
binding_length = 25,
step_length = binding_length,
peak_width = fragment_length/2,
pc_count_cutoff = 5,
bg_count_cutoff = 50,
p_cutoff = 1e-05,
p_adj_cutoff = NULL,
log2FC_cutoff = 1,
consistent_peak = FALSE,
consistent_log2FC_cutoff = 1,
consistent_fdr_cutoff = 0.05,
alpha = 0.05,
p0 = 0.8,
parallel = FALSE,
background_method = c("all", "Gaussian_mixture", "m6Aseq_prior", "manual"),
manual_background = NULL,
correct_GC_bg = TRUE,
qtnorm = FALSE,
glm_type = c("DESeq2", "Poisson", "NB"),
LFC_shrinkage = c("apeglm", "ashr", "Gaussian", "none"),
export_results = TRUE,
export_format = c("CSV", "BED", "RDS"),
table_style = c("bed", "granges"),
save_plot_GC = TRUE,
save_plot_analysis = FALSE,
save_plot_name = "",
save_dir = "exomePeak2_output",
peak_calling_mode = c("exon", "full_tx", "whole_genome")
)

```

## Arguments

<code>bam_ip</code>	a character vector for the BAM file directories of the (control) IP samples.
<code>bam_input</code>	a character vector for the BAM file directories of the (control) input samples.
<code>bam_treated_ip</code>	a character vector for the BAM file directories of the treated IP samples.
<code>bam_treated_input</code>	a character vector for the BAM file directories of the treated input samples. If the bam files do not contain treatment group, user should only fill the arguments of <code>BAM_ip</code> and <code>BAM_input</code> .
<code>txdb</code>	a <code>TxDb</code> object for the transcript annotation. If the <code>TxDb</code> object is not available, it could be a character string of the UCSC genome name which is acceptable by <code>makeTxDbFromUCSC</code> . For example: "hg19".
<code>bsgenome</code>	a <code>BSgenome</code> object for the genome sequence information. If the <code>BSgenome</code> object is not available, it could be a character string of the UCSC genome name which is acceptable by <code>getBSgenome</code> . For example: "hg19".

genome	<p>a character string of the UCSC genome name which is acceptable by <a href="#">getBSgenome</a> or/and <a href="#">makeTxDbFromUCSC</a>. For example: "hg19".</p> <p>By default, the argument = NA, it should be provided when the BSgenome or/and the TxDb object are not available.</p>
gff_dir	<p>optional, a character which specifies the directory toward a gene annotation GFF/GTF file, it is applied when the TxDb object is not available, default = NULL.</p>
mod_annot	<p>a <a href="#">GRanges</a> object for user provided single based RNA modification annotation. If user provides the single based RNA modification annotation, exomePeak2 will perform reads count and (differential) modification quantification on the provided annotation.</p> <p>The single base annotation will be flanked by <math>\text{length} = \text{floor}(\text{fragment\_length} - \text{binding\_length}/2)</math> to account for the fragment length of the sequencing library.</p>
paired_end	<p>a logical of whether the data comes from the Paired-End Library, TRUE if the data is Paired-End sequencing; default FALSE.</p>
library_type	<p>a character specifying the protocol type of the RNA-seq library, can be one in <code>c("unstranded", "1st_strand", "2nd_strand")</code>; default = "unstranded".</p> <p><b>unstranded</b> The randomly primed RNA-seq library type, i.e. both the strands generated during the first and the second strand sythesis are sequenced; example: Standard Illumina.</p> <p><b>1st_strand</b> The first strand-specific RNA-seq library, only the strand generated during the first strand sythesis is sequenced; examples: dUTP, NSR, NNSR.</p> <p><b>2nd_strand</b> The second strand-specific RNA-seq library, only the strand generated during the second strand sythesis is sequenced; examples: Ligation, Standard SOLiD.</p>
fragment_length	<p>a positive integer number for the expected fragment length in nucleotides; default = 100.</p>
binding_length	<p>a positive integer number for the expected binding length of the anti-modification antibody in IP samples; default = 25.</p>
step_length	<p>a positive integer number for the shift distances of the sliding window; default = binding_length.</p>
peak_width	<p>a numeric value for the minimum width of the merged peaks; default = fragment_length</p>
pc_count_cutoff	<p>a numeric value for the cutoff on average window's reads count in peak calling; default = 5.</p>
bg_count_cutoff	<p>a numeric value for the cutoff on average window's reads count in background identification; default = 50.</p>
p_cutoff	<p>a numeric value for the cutoff on p values in peak calling; default = 1e-05.</p>
p_adj_cutoff	<p>a numeric value for the cutoff on Benjamini Hochberg adjusted p values in peak calling; default = NULL.</p>
log2FC_cutoff	<p>a numeric value for the cutoff on log2 IP over input fold changes in peak calling; default = 1.</p>
consistent_peak	<p>a logical of whether the positive peaks returned should be consistent among all the replicates; default = FALSE.</p>

consistent_log2FC_cutoff	a numeric for the modification log2 fold changes cutoff in the peak consistency calculation; default = 1.
consistent_fdr_cutoff	a numeric for the BH adjusted C-test p values cutoff in the peak consistency calculation; default = 0.05. Check <a href="#">ctest</a> .
alpha	a numeric for the binomial quantile used in the consistent peak filter; default = 0.05.
p0	a numeric for the binomial proportion parameter used in the consistent peak filter; default = 0.8. For a peak to be consistently methylated, the minimum number of significant enriched replicate pairs is defined as the 1 - alpha quantile of a binomial distribution with $p = p_0$ and $N =$ number of possible pairs between replicates. The consistency defined in this way is equivalent to the rejection of an exact binomial test with null hypothesis of $p < p_0$ and $N =$ replicates number of IP * replicates number of input.
parallel	a logical value indicating whether to use parallel computation, it will require > 16GB memory if parallel = TRUE; default = FALSE.
background_method	a character specifies the method of finding background regions for peak detection, i.e. to identify the windows without modification signal. It could be one of "Gaussian_mixture", "m6Aseq_prior", "manual", and "all"; default = "all". In order to accurately account for the technical variations, it is often important to estimate the sequencing depth and GC content linear effects on windows without modification signals. The following methods are supported in ExomePeak2 to differentiate the no modification background windows from the modification containing windows. <b>all</b> Use all windows as the background. This choice assumes no differences in the effects of technical features between the background and the modification regions. <b>Gaussian_mixture</b> The background is identified by Multivariate Gaussian Mixture Model (MGMM) with 2 mixing components on the vectors containing methylation level estimates and GC content, the background regions are predicted by the Bayes Classifier on the learned GMM. <b>m6Aseq_prior</b> The background is identified by the prior knowledge of m6A topology, the windows that are not overlapped with long exons (exon length $\geq 400$ bp) and 5'UTR are treated as the background windows. This type of background should not be used if the MeRIP-seq data is not targetting on m6A methylation. <b>manual</b> The background regions are defined by the user manually at the argument manual_background.
manual_background	a <a href="#">GRanges</a> object for the user provided unmodified background; default = NULL.
correct_GC_bg	a logical value of whether to estimate the GC content linear effect on background regions during modification level quantification; default = TRUE. If = TRUE, it could lead to a more accurate estimation of GC content bias for the RNA modifications that are highly biologically related to GC content.
qtnorm	a logical of whether to perform subset quantile normalization after the GC content linear effect correction; default = FALSE.



If `qtnorm = TRUE`, subset quantile normalization will be applied within the IP and input samples separately to account for the inherent differences between the marginal distributions of IP and input samples.

<code>glm_type</code>	<p>a character specifies the type of Generalized Linear Model (GLM) fitted for the purpose of statistical inference during peak calling, which can be one of the <code>c("DESeq2", "NB", "Poisson")</code>.</p> <p>DESeq2 Fit the GLM defined in the function <code>DESeq</code>, which is the NB GLM with regulated estimation of the overdispersion parameters.</p> <p>NB Fit the ordinary Negative Binomial (NB) GLM.</p> <p>Poisson Fit the Poisson GLM.</p> <p>By default, the DESeq2 GLMs are fitted on the data set with &gt; 1 biological replicates for both the IP and input samples, the Poisson GLM will be fitted otherwise.</p>
<code>LFC_shrinkage</code>	<p>a character for the method of empirical bayes shrinkage on log2FC, could be one of <code>c("apeglm", "ashr", "Gaussian", "none")</code>; Default = "apeglm".</p> <p>see <a href="#">lfcShrink</a> for more details; if "none" is selected, only the MLE will be returned.</p>
<code>export_results</code>	a logical of whether to save the results on disk; default = TRUE.
<code>export_format</code>	a character vector for the format(s) of the result being exported, could be the subset of <code>c("CSV", "BED", "RDS")</code> ; Default = <code>c("CSV", "BED", "RDS")</code> .
<code>table_style</code>	a character for the style of the table being exported, could be one of <code>c("bed", "granges")</code> ; Default = "bed".
<code>save_plot_GC</code>	a logical of whether to generate the plots for GC content bias assessment; default = TRUE.
<code>save_plot_analysis</code>	a logical of whether to generate the plots for genomic analysis on modification sites; default = FALSE.
<code>save_plot_name</code>	a character for the name of the plots being saved; Default = "Plot".
<code>save_dir</code>	a character for the name of the directory being saved; Default = "exomePeak2_output".
<code>peak_calling_mode</code>	<p>a character specifies the scope of peak calling on genome, can be one of <code>c("exon", "full_transcript", "whole_genome")</code>; Default = "exon".</p> <p><code>exon</code> generate sliding windows on exon regions.</p> <p><code>full_transcript</code> generate sliding windows on the full transcripts (include both introns and exons).</p> <p><code>whole_genome</code> generate sliding windows on the whole genome (include introns, exons, and the intergenic regions).</p> <p>P.S. The full transcript mode and the whole genome mode demand big memory size (&gt; 4GB) for large genomes.</p>

## Details

`exomePeak2` call RNA modification peaks and calculate peak statistics from **BAM** files of a MeRIP-seq experiment.

The transcript annotation (from either the `TxDb` object or the **GFF** file) should be provided to perform analysis on exons.

The `BSSgenome` object is also required to perform the GC content bias adjustment. If the `bssgenome` and the genome arguments are not provided (= NULL), the downstream analysis will proceed without GC content bias corrections.

If the **BAM** files in treated samples are provided at the arguments `bam_treated_ip` and `bam_treated_input`, the statistics of differential modification analysis on peaks/sites will be reported.

Under default setting, `exomePeak2` will save the results of (differential) modification analysis under a folder named by 'exomePeak2\_output'. The results generated include a **BED** file and a **CSV** table that store the locations and statistics of the (differential) modified peaks/sites.

## Value

a `SummarizedExomePeak` object.

## See Also

[exomePeakCalling](#), [glmM](#), [glmDM](#), [normalizeGC](#), [exportResults](#), [plotLfcGC](#)

## Examples

```
#Specify File Directories

GENE_ANNO_GTF = system.file("extdata", "example.gtf", package="exomePeak2")

f1 = system.file("extdata", "IP1.bam", package="exomePeak2")
f2 = system.file("extdata", "IP2.bam", package="exomePeak2")
f3 = system.file("extdata", "IP3.bam", package="exomePeak2")
f4 = system.file("extdata", "IP4.bam", package="exomePeak2")
IP_BAM = c(f1,f2,f3,f4)
f1 = system.file("extdata", "Input1.bam", package="exomePeak2")
f2 = system.file("extdata", "Input2.bam", package="exomePeak2")
f3 = system.file("extdata", "Input3.bam", package="exomePeak2")
INPUT_BAM = c(f1,f2,f3)

# Peak Calling

sep <- exomePeak2(bam_ip = IP_BAM,
                 bam_input = INPUT_BAM,
                 gff_dir = GENE_ANNO_GTF,
                 genome = "hg19",
                 paired_end = FALSE)

sep

# Differential Modification Analysis on Modification Peaks (Comparison of Two Conditions)

f1 = system.file("extdata", "treated_IP1.bam", package="exomePeak2")
TREATED_IP_BAM = c(f1)
f1 = system.file("extdata", "treated_Input1.bam", package="exomePeak2")
TREATED_INPUT_BAM = c(f1)

sep <- exomePeak2(bam_ip = IP_BAM,
                 bam_input = INPUT_BAM,
                 bam_treated_input = TREATED_INPUT_BAM,
                 bam_treated_ip = TREATED_IP_BAM,
                 gff_dir = GENE_ANNO_GTF,
                 genome = "hg19",
```

```

        paired_end = FALSE)
sep

# Modification Level Quantification on Single Based Modification Annotation

f2 = system.file("extdata", "mod_annot.rds", package="exomePeak2")

MOD_ANNO_GRANGE <- readRDS(f2)

sep <- exomePeak2(bam_ip = IP_BAM,
                 bam_input = INPUT_BAM,
                 gff_dir = GENE_ANNO_GTF,
                 genome = "hg19",
                 paired_end = FALSE,
                 mod_annot = MOD_ANNO_GRANGE)

sep

# Differential Modification Analysis on Single Based Modification Annotation

sep <- exomePeak2(bam_ip = IP_BAM,
                 bam_input = INPUT_BAM,
                 bam_treated_input = TREATED_INPUT_BAM,
                 bam_treated_ip = TREATED_IP_BAM,
                 gff_dir = GENE_ANNO_GTF,
                 genome = "hg19",
                 paired_end = FALSE,
                 mod_annot = MOD_ANNO_GRANGE)

sep

```

---

exomePeak2Results-methods

*Method exomePeak2Results*

---

## Description

Method exomePeak2Results

Method exomePeak2Results<-

Accessor to the slot exomePeak2Results in class SummarizedExomePeak.

Accessor to the slot exomePeak2Results in class SummarizedExomePeak.

## Usage

```
exomePeak2Results(x1)
```

```
exomePeak2Results(x2) <- value
```

```
## S4 method for signature 'SummarizedExomePeak'
```

```
exomePeak2Results(x1)
```

```
## S4 replacement method for signature 'SummarizedExomePeak'
```

```
exomePeak2Results(x2) <- value
```

**Arguments**

x1                    A data.frame object.  
x2                    A SummarizedExomePeak object.  
value                 a data.frame object for the DESeq2 Results.

**Value**

A data.frame object for the DESeq2 Results.

**Examples**

```
f1 = system.file("extdata", "sep_ex_mod.rds", package="exomePeak2")  
sep <- readRDS(f1)  
head(exomePeak2Results(sep))
```

---

exomePeakCalling-methods

*Method exomePeakCalling*

---

**Description**

exomePeakCalling call peaks of RNA modification from a MeRIP-seq data set.

**Usage**

```
exomePeakCalling(  
  merip_bams = NULL,  
  txdb = NULL,  
  bsgenome = NULL,  
  genome = NA,  
  mod_annot = NULL,  
  glm_type = c("DESeq2", "NB", "Poisson"),  
  background_method = c("Gaussian_mixture", "m6Aseq_prior", "manual", "all"),  
  manual_background = NULL,  
  correct_GC_bg = TRUE,  
  qtnorm = FALSE,  
  gff_dir = NULL,  
  fragment_length = 100,  
  binding_length = 25,  
  step_length = binding_length,  
  peak_width = fragment_length/2,  
  pc_count_cutoff = 5,  
  bg_count_cutoff = 50,  
  p_cutoff = 1e-05,  
  p_adj_cutoff = NULL,  
  log2FC_cutoff = 0,  
  consistent_peak = FALSE,
```

```

    consistent_log2FC_cutoff = 0,
    consistent_fdr_cutoff = 0.05,
    alpha = 0.05,
    p0 = 0.8,
    parallel = FALSE,
    bp_param = NULL
)

## S4 method for signature 'MeripBamFileList'
exomePeakCalling(
  merip_bams = NULL,
  txdb = NULL,
  bsgenome = NULL,
  genome = NA,
  mod_annot = NULL,
  glm_type = c("DESeq2", "NB", "Poisson"),
  background_method = c("all", "Gaussian_mixture", "m6Aseq_prior", "manual"),
  manual_background = NULL,
  correct_GC_bg = TRUE,
  qtnorm = FALSE,
  gff_dir = NULL,
  fragment_length = 100,
  binding_length = 25,
  step_length = binding_length,
  peak_width = fragment_length/2,
  pc_count_cutoff = 5,
  bg_count_cutoff = 50,
  p_cutoff = 1e-05,
  p_adj_cutoff = NULL,
  log2FC_cutoff = 1,
  consistent_peak = FALSE,
  consistent_log2FC_cutoff = 1,
  consistent_fdr_cutoff = 0.05,
  alpha = 0.05,
  p0 = 0.8,
  parallel = FALSE,
  bp_param = NULL
)

```

### Arguments

<code>merip_bams</code>	a <code>MeripBamFileList</code> object returned by <a href="#">scanMeripBAM</a> .
<code>txdb</code>	a <code>TxDb</code> object for the transcript annotation. If the <code>TxDb</code> object is not available, it could be a character string of the UCSC genome name which is acceptable by <a href="#">makeTxDbFromUCSC</a> . For example: "hg19".
<code>bsgenome</code>	a <code>BSgenome</code> object for the genome sequence information. If the <code>BSgenome</code> object is not available, it could be a character string of the UCSC genome name which is acceptable by <a href="#">getBSgenome</a> . For example: "hg19".
<code>genome</code>	a character string of the UCSC genome name which is acceptable by <a href="#">getBSgenome</a> or/and <a href="#">makeTxDbFromUCSC</a> . For example: "hg19". By default, the argument = NA, it should be provided when the <code>BSgenome</code> or/and the <code>TxDb</code> object are not available.

mod_annot	<p>a <a href="#">GRanges</a> or <a href="#">GRangesList</a> object for user provided single based RNA modification annotation, the widths of the ranged object should be all equal to 1.</p> <p>If user provides the single based RNA modification annotation, exomePeak2 will perform reads count and (differential) modification quantification on the provided annotation.</p> <p>The single base annotation will be flanked by <math>\text{length} = \text{floor}(\text{fragment\_length} - \text{binding\_length}/2)</math> to account for the fragment length of the sequencing library.</p>
glm_type	<p>a character specifies the type of Generalized Linear Model (GLM) fitted for the purpose of statistical inference during peak calling, which can be one of the <code>c("DESeq2", "NB", "Poisson")</code>.</p> <p>DESeq2 Fit the GLM defined in function <a href="#">DESeq</a>, which is the NB GLM with regulated estimation of the overdispersion parameters.</p> <p>NB Fit the Negative Binomial (NB) GLM.</p> <p>Poisson Fit the Poisson GLM.</p> <p>By default, the DESeq2 GLMs are fitted on the data set with &gt; 1 biological replicates for both the IP and input samples, the Poisson GLM will be fitted otherwise.</p>
background_method	<p>a character specifies the method for the background finding, i.e. to identify the windows without modification signal. It could be one of the "Gaussian_mixture", "m6Aseq_prior", "manual", and "all"; default = "all".</p> <p>In order to accurately account for the technical variations, it is often necessary to estimate the GC content linear effects on windows without modification signals (background).</p> <p>The following methods are supported in ExomePeak2 to differentiate the no modification background windows from the modification containing windows.</p> <p><b>Gaussian_mixture</b> The background is identified by Multivariate Gaussian Mixture Model (MGMM) with 2 mixing components on the vectors containing methylation level estimates and GC content, the background regions are predicted by the Bayes Classifier on the learned GMM.</p> <p><b>m6Aseq_prior</b> The background is identified by the prior knowledge of m6A topology, the windows that are not overlapped with long exons (exon length <math>\geq 400</math>bp) and 5'UTR are treated as the background windows. This type of background should not be used if the MerIP-seq data is not using anti-m6A antibody.</p> <p><b>manual</b> The background regions are defined by user manually at the argument <code>manual_background</code>.</p> <p><b>all</b> Use all windows as the background. This is equivalent to not differentiating background and signal. It can lead to biases during the sequencing depth and the GC content correction factors estimation.</p>
manual_background	<p>a <a href="#">GRanges</a> object for the user provided unmodified background; default = NULL.</p>
correct_GC_bg	<p>a logical value of whether to estimate the GC content linear effect on background regions; default = TRUE.</p> <p>If = TRUE, it could lead to a more accurate estimation of GC content bias for the RNA modifications that are highly biologically related to GC content.</p>
qtnorm	<p>a logical of whether to perform subset quantile normalization after the GC content linear effect correction; default = FALSE.</p>

	If <code>qtnorm = TRUE</code> , subset quantile normalization will be applied within the IP and input samples separately to account for the inherent differences between the marginal distributions of IP and input samples.
<code>gff_dir</code>	optional, a character which specifies the directory toward a gene annotation GFF/GTF file, it is applied when the <code>TxDb</code> object is not available, default = <code>NULL</code> .
<code>fragment_length</code>	a positive integer number for the expected fragment length in nucleotides; default = <code>100</code> .
<code>binding_length</code>	a positive integer number for the expected binding length of the anti-modification antibody in IP samples; default = <code>25</code> .
<code>step_length</code>	a positive integer number for the shift distances of the sliding window; default = <code>binding_length</code> .
<code>peak_width</code>	a numeric value for the minimum width of the merged peaks; default = <code>fragment_length</code> .
<code>pc_count_cutoff</code>	a numeric value for the cutoff on average window's reads count in peak calling; default = <code>5</code> .
<code>bg_count_cutoff</code>	a numeric value for the cutoff on average window's reads count in background identification; default = <code>50</code> .
<code>p_cutoff</code>	a numeric value for the cutoff on p values in peak calling; default = <code>1e-05</code> .
<code>p_adj_cutoff</code>	a numeric value for the cutoff on Benjamini Hochberg adjusted p values in peak calling; default = <code>NULL</code> .
<code>log2FC_cutoff</code>	a numeric value for the cutoff on log2 IP over input fold changes in peak calling; default = <code>1</code> .
<code>consistent_peak</code>	a logical of whether the positive peaks returned should be consistent among all the replicates; default = <code>FALSE</code> .
<code>consistent_log2FC_cutoff</code>	a numeric for the modification log2 fold changes cutoff in the peak consistency calculation; default = <code>1</code> .
<code>consistent_fdr_cutoff</code>	a numeric for the BH adjusted C-test p values cutoff in the peak consistency calculation; default = <code>0.05</code> . Check <a href="#">ctest</a> .
<code>alpha</code>	a numeric for the binomial quantile used in the consistent peak filter; default = <code>0.05</code> .
<code>p0</code>	a numeric for the binomial proportion parameter used in the consistent peak filter; default = <code>0.8</code> . For a peak to be consistently methylated, the minimum number of significant enriched replicate pairs is defined as the $1 - \alpha$ quantile of a binomial distribution with $p = p0$ and $N =$ number of possible pairs between replicates. The consistency defined in this way is equivalent to the rejection of an exact binomial test with null hypothesis of $p < p0$ and $N =$ replicates number of IP * replicates number of input.
<code>parallel</code>	a logical value of whether to use parallel computation, typically it requires more than 16GB of RAM if <code>parallel = TRUE</code> ; default = <code>FALSE</code> .
<code>bp_param</code>	optional, a <a href="#">BiocParallelParam</a> object that stores the configuration parameters for the parallel execution.

## Details

exomePeakCalling perform peak calling from the MeRIP-seq BAM files on exon regions defined by the user provided transcript annotations. If the [BSgenome](#) object is provided, the peak calling will be conducted with the GC content bias correction.

Under the default setting, for each window, exomePeak2 will fit a GLM of Negative Binomial (NB) with regulated estimation of the overdispersion parameters developed in [DESeq](#). Wald tests with  $H_0$  of IP/input Log2 Fold Change (LFC)  $\leq 0$  are performed on each of the sliding windows. The significantly modified peaks are selected using the cutoff of p value  $< 0.0001$ .

## Value

a [SummarizedExomePeak](#) object.

## See Also

[exomePeak2](#), [glmM](#), [glmDM](#), [normalizeGC](#), [exportResults](#)

## Examples

```
### Define File Directories

GENE_ANNO_GTF = system.file("extdata", "example.gtf", package="exomePeak2")

f1 = system.file("extdata", "IP1.bam", package="exomePeak2")
f2 = system.file("extdata", "IP2.bam", package="exomePeak2")
f3 = system.file("extdata", "IP3.bam", package="exomePeak2")
f4 = system.file("extdata", "IP4.bam", package="exomePeak2")
IP_BAM = c(f1,f2,f3,f4)
f1 = system.file("extdata", "Input1.bam", package="exomePeak2")
f2 = system.file("extdata", "Input2.bam", package="exomePeak2")
f3 = system.file("extdata", "Input3.bam", package="exomePeak2")
INPUT_BAM = c(f1,f2,f3)

f1 = system.file("extdata", "treated_IP1.bam", package="exomePeak2")
TREATED_IP_BAM = c(f1)
f1 = system.file("extdata", "treated_Input1.bam", package="exomePeak2")
TREATED_INPUT_BAM = c(f1)

### Peak Calling

MeRIP_Seq_Alignment <- scanMeripBAM(
  bam_ip = IP_BAM,
  bam_input = INPUT_BAM,
  paired_end = FALSE
)

sep <- exomePeakCalling(
  merip_bams = MeRIP_Seq_Alignment,
  gff_dir = GENE_ANNO_GTF,
  genome = "hg19"
)

sep <- normalizeGC(sep)
```



```
sep <- glmM(sep)

exportResults(sep)

### Differential Modification Analysis (Comparison of Two Conditions)

MeRIP_Seq_Alignment <- scanMeripBAM(
  bam_ip = IP_BAM,
  bam_input = INPUT_BAM,
  bam_treated_ip = TREATED_IP_BAM,
  bam_treated_input = TREATED_INPUT_BAM,
  paired_end = FALSE
)

sep <- exomePeakCalling(
  merip_bams = MeRIP_Seq_Alignment,
  gff_dir = GENE_ANNO_GTF,
  genome = "hg19"
)

sep <- normalizeGC(sep)

sep <- glmDM(sep)

exportResults(sep)
```

---

exonPlot

*plot the distribution for the length of the overlapped exons.*

---

## Description

plot the distribution for the length of the overlapped exons.

## Usage

```
exonPlot(gfeatures, txdb, save_pdf_prefix = NULL, save_dir = ".")
```

## Arguments

<code>gfeatures</code>	a list of GRanges or GRangesList.
<code>txdb</code>	a txdb object.
<code>save_pdf_prefix</code>	provided to save a pdf file.
<code>save_dir</code>	a character indicating the directory to save the plot; default ".".

## Value

a plot for the exons length distribution.

---

exportResults-methods *Method exportResults*

---

## Description

Method exportResults

Export the (Differential) Modification Peaks/Sites and their associated Statistics

## Usage

```
exportResults(
  sep,
  format = c("CSV", "BED", "RDS"),
  table_style = c("bed", "granges"),
  save_dir = "exomepeaks_result",
  cut_off_pvalue = NULL,
  cut_off_padj = 0.1,
  cut_off_log2FC = 0,
  min_num_of_positive = 30,
  expected_direction = c("both", "hyper", "hypo"),
  inhibit_filter = FALSE,
  reads_count = TRUE,
  GC_sizeFactors = TRUE
)

## S4 method for signature 'SummarizedExomePeak'
exportResults(
  sep,
  format = c("CSV", "BED", "RDS"),
  table_style = c("bed", "granges"),
  save_dir = "exomePeak2_output",
  cut_off_pvalue = NULL,
  cut_off_padj = 0.1,
  cut_off_log2FC = 0,
  min_num_of_positive = 100,
  expected_direction = c("both", "hyper", "hypo"),
  inhibit_filter = FALSE,
  reads_count = TRUE,
  GC_sizeFactors = TRUE
)
```

## Arguments

sep	a <a href="#">SummarizedExomePeak</a> object.
format	a character for the exported format, could be a vector that contains c("CSV", "BED", "RDS"). CSV export a comma separated values (CSV) table with the genomic location and LFC statistics. BED export a BEDGraph file with the score column = -log2(adjusted p value). RDS export the RDS file of the <a href="#">SummarizedExperiment</a> object.

table_style	a character for the style of the CSV table being exported, could be one in <code>c("bed", "granges")</code> .  bed the genomic locations in the table are represented by BEDgraph style. granges the genomic locations in the table are represented by GRanges style.
save_dir	a character for the name of the directory being saved; Default = "exomePeak2_output".
cut_off_pvalue	a numeric value for the p value cutoff in the exported result; Default = NULL.
cut_off_padj	a numeric value for the adjusted p value cutoff in the exported result; Default = 0.05.
cut_off_log2FC	a numeric value for the log2 fold change cutoff of the exported result, only the sites with <code>abs(LFC)</code> larger than this value are kept; Default = 0.
min_num_of_positive	a numeric value for the minimum number of reported sites. If the number of remaining sites is less than this number after filtering, additional sites will be reported by the increasing order of the p value to meet this number.
expected_direction	a character for the expected differential modification direction, could be one in <code>c("hyper", "hypo", "both")</code> .  hyper only report the peaks/sites with interactive LFC > 0. hypo only report the peaks/sites with interactive LFC < 0. both report the peaks/sites in both directions.  This argument is useful when the treated group involves the perturbation of a known writer or eraser protein; Default "both".
inhibit_filter	a logical of whether to remove all the filters, this option is useful when quantification on single based site annotation; Default = FALSE.
reads_count	a logical of whether to export the reads count for each sample; Default = TRUE.
GC_sizeFactors	a logical of whether to export the GC content correction size factors; Default = TRUE.

**Value**

none, a folder will be saved on the disk.

**Examples**

```
### Load the example SummarizedExomPeak object
f1 = system.file("extdata", "sep_ex_mod.rds", package="exomePeak2")

sep <- readRDS(f1)

### Save the modification peaks/sites statistics on the current directory.
exportResults(sep)
```

---

GCsizeFactors-methods *Method GCsizeFactors*

---

**Description**

Method GCsizeFactors

Method GCsizeFactors<-

Accessor to the slot GCsizeFactors in class SummarizedExomePeak.

Accessor to the slot GCsizeFactors in class SummarizedExomePeak.

**Usage**

```
GCsizeFactors(x1)
```

```
GCsizeFactors(x2) <- value
```

```
## S4 method for signature 'SummarizedExomePeak'  
GCsizeFactors(x1)
```

```
## S4 replacement method for signature 'SummarizedExomePeak'  
GCsizeFactors(x2) <- value
```

**Arguments**

x1                    A SummarizedExomePeak object.

x2                    A SummarizedExomePeak object.

value                A matrix object.

**Value**

a data.frame for the GC content size factors of each sample

**Examples**

```
f1 = system.file("extdata", "sep_ex_mod.rds", package="exomePeak2")  
sep <- readRDS(f1)  
head(GCsizeFactors(sep))
```

glmDM-methods

Method glmDM

**Description**

glmDM perform inference and estimation on RNA differential modification log2FC.

GLMs with interactive design between dummy variables of IP/input and Treatment/control are fitted for each peaks/sites:

$$\log_2(Q) = \text{intercept} + I(\text{Treatment}) + I(\text{IP}) + I(\text{IP}) * I(\text{Treatment})$$

The log2FC and the associated statistics are based on the coefficient estimate of the interactive term:  $I(\text{IP}) * I(\text{Treated})$ .

Under default setting, the returned log2FC are the RR estimates with Couchey priors defined in [apeglm](#).

**Usage**

```
glmDM(
  sep,
  glm_type = c("DESeq2", "NB", "Poisson"),
  LFC_shrinkage = c("apeglm", "ashr"),
  ...
)

## S4 method for signature 'SummarizedExomePeak'
glmDM(
  sep,
  glm_type = c("DESeq2", "NB", "Poisson"),
  LFC_shrinkage = c("apeglm", "ashr", "none"),
  ...
)
```

**Arguments**

sep	a <a href="#">SummarizedExomePeak</a> object.
glm_type	a character specifies the type of Generalized Linear Model (GLM) fitted for the purpose of statistical inference during peak calling, which can be one of the <code>c("DESeq2", "NB", "Poisson")</code> . DESeq2 Fit the GLM defined in the function <a href="#">DESeq</a> , which is the NB GLM with regulated estimation of the overdispersion parameters. NB Fit the Negative Binomial (NB) GLM. Poisson Fit the Poisson GLM. By default, the DESeq2 GLMs are fitted on the data set with > 1 biological replicates for both the IP and input samples, the Poisson GLM will be fitted otherwise.
LFC_shrinkage	a character for the method of empirical bayes shrinkage on log2FC, could be one of <code>c("apeglm", "ashr", "none")</code> ; Default = "apeglm". see <a href="#">lfcShrink</a> for details; if "none" is selected, only the MLE will be returned.
...	Optional arguments passed to <a href="#">DESeq</a>

**Value**

a SummarizedExomPeak object.

**See Also**

[glmM](#)

**Examples**

```
### Load the example SummarizedExomPeak object
f1 = system.file("extdata", "sep_ex_dm.rds", package="exomePeak2")

sep <- readRDS(f1)

### Normalize the GC contents biases
sep <- normalizeGC(sep)

### Calculate GLM Statistics on the Modification Peaks
sep <- glmDM(sep)
```

---

glmM-methods

*Method glmM*

---

**Description**

glmM performs inference and estimation on IP/input log2FC.

GLMs with the design of an indicator of IP samples are fitted for each peaks/sites:

$$\log_2(Q) = \text{intercept} + I(IP)$$

The log2FC and the associated statistics are based on the coefficient estimate of the dummy variable term:  $I(IP)$ .

Under default setting, the returned log2FC are the RR estimates with Couchey priors defined in [apeglm](#).

**Usage**

```
glmM(
  sep,
  glm_type = c("DESeq2", "NB", "Poisson"),
  LFC_shrinkage = c("apeglm", "Gaussian", "ashr"),
  ...
)

## S4 method for signature 'SummarizedExomePeak'
glmM(
  sep,
  glm_type = c("DESeq2", "NB", "Poisson"),
  LFC_shrinkage = c("apeglm", "Gaussian", "ashr"),
  ...
)
```

**Arguments**

sep	a SummarizedExomePeak object.
glm_type	a character specifies the type of Generalized Linear Model (GLM) fitted for the purpose of statistical inference during peak calling, which can be one of the <code>c("DESeq2", "NB", "Poisson")</code> .  DESeq2 Fit the GLM defined in the function <a href="#">DESeq</a> , which is the NB GLM with regulated estimation of the overdispersion parameters. NB Fit the Negative Binomial (NB) GLM. Poisson Fit the Poisson GLM.  By default, the DESeq2 GLMs are fitted on the data set with > 1 biological replicates for both the IP and input samples, the Poisson GLM will be fitted otherwise.
LFC_shrinkage	a character for the method of empirical bayes shrinkage on log2FC, could be one of <code>c("apeglm", "Gaussian", "ashr", "none")</code> ; Default = "apeglm". see <a href="#">lfcShrink</a> for details; if "none" is selected, only the MLE will be returned.
...	Optional arguments passed to <a href="#">DESeq</a>

**Value**

a SummarizedExomPeak object.

**See Also**

[glmDM](#)

**Examples**

```
### Load the example SummarizedExomPeak object
f1 = system.file("extdata", "sep_ex_mod.rds", package="exomePeak2")

sep <- readRDS(f1)

### Select only the control group to avoid warning.
sep <- sep[,!colData(sep)$design_Treatment]

### Normalize the GC contents biases
sep <- normalizeGC(sep)

### Calculate GLM Statistics on the Modification Peaks
sep <- glmM(sep)
```

---

LibraryType-methods    *Method LibraryType*

---

**Description**

Method LibraryType

Accessor to the slot LibraryType in class MeripBamFileList.

**Usage**

```
LibraryType(x)

## S4 method for signature 'MeripBamFileList'
LibraryType(x)
```

**Arguments**

x                    a MeripBamFileList object.

**Value**

a value for the library type of MeRIP-seq experiment.

**Examples**

```
GENE_ANNO_GTF = system.file("extdata", "example.gtf", package="exomePeak2")

f1 = system.file("extdata", "IP1.bam", package="exomePeak2")
f2 = system.file("extdata", "IP2.bam", package="exomePeak2")
f3 = system.file("extdata", "IP3.bam", package="exomePeak2")
f4 = system.file("extdata", "IP4.bam", package="exomePeak2")
IP_BAM = c(f1,f2,f3,f4)
f1 = system.file("extdata", "Input1.bam", package="exomePeak2")
f2 = system.file("extdata", "Input2.bam", package="exomePeak2")
f3 = system.file("extdata", "Input3.bam", package="exomePeak2")
INPUT_BAM = c(f1,f2,f3)

f1 = system.file("extdata", "treated_IP1.bam", package="exomePeak2")
TREATED_IP_BAM = c(f1)
f1 = system.file("extdata", "treated_Input1.bam", package="exomePeak2")
TREATED_INPUT_BAM = c(f1)

MeRIP_Seq_Alignment <- scanMeripBAM(
  bam_ip = IP_BAM,
  bam_input = INPUT_BAM,
  paired_end = FALSE
)

LibraryType(MeRIP_Seq_Alignment)
```

---

MeripBamFileList-class

*Maintain and use BAM files in MeRIP-Seq experiment*

---

**Description**

An S4 object defined in exomePeak2 that summarizes the BAM files used in a MeRIP-Seq experiment.

MeripBamFileList() provide a convenient format to store and manage the BAM file directories for MeRIP-Seq data set.

This class contains [BamFileList](#) from the package Rsamtools.



**Details****Constructors:**

MeripBamFileList can be constructed by scanMeripBAM()

**Accessors:**

MeripBamFileList object share all the accessors with the [BamFileList](#) class, please check it for more information.

The frequently used accessors include:

metadata(): Return a list storing the design of MeRIP-Seq experiment.

Parameter(): Access to the BAM FLAG parameters used for BAM file filtering.

asMate(): Return a logical value, TRUE if the BAM file is paired end.

It has one additional accessor LibraryType()

LibraryType() retrieves the strand specificity information of the RNA-Seq library.

**Examples**

```
### Define BAM File Directories

f1 = system.file("extdata", "IP1.bam", package="exomePeak2")
f2 = system.file("extdata", "IP2.bam", package="exomePeak2")
f3 = system.file("extdata", "IP3.bam", package="exomePeak2")
f4 = system.file("extdata", "IP4.bam", package="exomePeak2")
IP_BAM = c(f1,f2,f3,f4)
f1 = system.file("extdata", "Input1.bam", package="exomePeak2")
f2 = system.file("extdata", "Input2.bam", package="exomePeak2")
f3 = system.file("extdata", "Input3.bam", package="exomePeak2")
INPUT_BAM = c(f1,f2,f3)

f1 = system.file("extdata", "treated_IP1.bam", package="exomePeak2")
TREATED_IP_BAM = c(f1)
f1 = system.file("extdata", "treated_Input1.bam", package="exomePeak2")
TREATED_INPUT_BAM = c(f1)

### For MeRIP-Seq Experiment Without the Treatment Group

MeRIP_Seq_Alignment <- scanMeripBAM(
  bam_ip = IP_BAM,
  bam_input = INPUT_BAM,
  paired_end = FALSE
)

### For MeRIP-Seq Experiment With the Treatment Group

MeRIP_Seq_Alignment <- scanMeripBAM(
  bam_ip = IP_BAM,
  bam_input = INPUT_BAM,
  bam_treated_ip = TREATED_IP_BAM,
  bam_treated_input = TREATED_INPUT_BAM,
  paired_end = FALSE
)

LibraryType(MeRIP_Seq_Alignment)
```

Parameter(MeRIP\_Seq\_Alignment)

---

normalizeGC-methods     *Method normalizeGC*

---

## Description

normalizeGC estimates the feature specific size factors in order to reduce the technical variation during modification peak statistics quantification.

## Usage

```
normalizeGC(
  sep,
  bsgenome = "hg19",
  txdb = "hg19",
  gff_dir = NULL,
  fragment_length = 100,
  binding_length = 25,
  feature = c("Background", "Modification", "All"),
  qtnorm = FALSE,
  effective_GC = FALSE
)

## S4 method for signature 'SummarizedExomePeak'
normalizeGC(
  sep,
  bsgenome = NULL,
  txdb = NULL,
  gff_dir = NULL,
  fragment_length = 100,
  binding_length = 25,
  feature = c("Background", "Modification", "All"),
  qtnorm = FALSE,
  effective_GC = FALSE
)
```

## Arguments

sep	a <a href="#">SummarizedExomePeak</a> object returned by <a href="#">exomePeak2</a> or <a href="#">exomePeakCalling</a> .
bsgenome	a <a href="#">BSgenome</a> object for the genome reference, If the BSgenome object is not available, it could be a character string of the UCSC genome name which is acceptable by <a href="#">getBSgenome</a> , example: "hg19".
txdb	a <a href="#">TxDb</a> object for the transcript annotation, If the TxDb object is not available, it could be a character string of the UCSC genome name which is acceptable by <a href="#">makeTxDbFromUCSC</a> , example: "hg19".
gff_dir	optional, a character which specifies the directory toward a gene annotation GFF/GTF file, it is applied when the TxDb object is not available, default = NULL.

fragment_length	a positive integer number for the expected fragment length in nucleotides; default = 100.
binding_length	a positive integer number for the expected binding length of the anti-modification antibody in IP samples; default = 25.
feature	a character specifies the region used in the GC content linear effect estimation, can be one in c("Background", "All", "Modification"); default is "Background". Background The GC content linear effects will be estimated on the background regions. By default, the background is defined as the exon regions not overlapping with peaks / modification sites flanked by the fragment length. You could select alternative background finding methods with background at <a href="#">exomePeakCalling</a> . Modification The GC content linear effects will be estimated on the regions of modification peaks/sites. All The GC content linear effects will be estimated on all regions, i.e. both the region of modification and the background control regions.
qtnorm	a logical of whether to perform subset quantile normalization after the GC content linear effect correction; default = FALSE. If qtnorm = TRUE, subset quantile normalization will be applied within the IP and input samples separately to account for the inherent differences between the marginal distributions of IP and input samples.
effective_GC	a logical of whether to calculate the effective GC content weighted by the fragment alignment probabilities; default = FALSE.

### Details

PCR amplification bias related to GC content is a major source of technical variation in RNA-seq. The GC content biases are usually correlated within the same laboratory environment, and this will result in the batch effect between different studies.

The GC content normalization can result in an improvement of peak accuracy for most published m6A-seq data, and it is particularly recommended if you want to compare the quantifications on methylation levels between different laboratory conditions.

### Value

a SummarizedExomePeak object with the updated slot GCsizeFactors.

### See Also

[estimateSeqDepth](#)

### Examples

```
### Load the example SummarizedExomPeak object
f1 = system.file("extdata", "sep_ex_mod.rds", package="exomePeak2")

sep <- readRDS(f1)

### Normalize the GC content biases
sep <- normalizeGC(sep)
```

---

Parameter-methods	<i>Method Parameter</i>
-------------------	-------------------------

---

**Description**

Method Parameter

Accessor to the slot Parameter in class MeripBamFileList.

**Usage**

```
Parameter(x)

## S4 method for signature 'MeripBamFileList'
Parameter(x)
```

**Arguments**

x                    a MeripBamFileList object.

**Value**

a list for the additional parameters of the MeRIP-seq experiment.

**Examples**

```
GENE_ANNO_GTF = system.file("extdata", "example.gtf", package="exomePeak2")

f1 = system.file("extdata", "IP1.bam", package="exomePeak2")
f2 = system.file("extdata", "IP2.bam", package="exomePeak2")
f3 = system.file("extdata", "IP3.bam", package="exomePeak2")
f4 = system.file("extdata", "IP4.bam", package="exomePeak2")
IP_BAM = c(f1,f2,f3,f4)
f1 = system.file("extdata", "Input1.bam", package="exomePeak2")
f2 = system.file("extdata", "Input2.bam", package="exomePeak2")
f3 = system.file("extdata", "Input3.bam", package="exomePeak2")
INPUT_BAM = c(f1,f2,f3)

f1 = system.file("extdata", "treated_IP1.bam", package="exomePeak2")
TREATED_IP_BAM = c(f1)
f1 = system.file("extdata", "treated_Input1.bam", package="exomePeak2")
TREATED_INPUT_BAM = c(f1)

MeRIP_Seq_Alignment <- scanMeripBAM(
  bam_ip = IP_BAM,
  bam_input = INPUT_BAM,
  paired_end = FALSE
)

Parameter(MeRIP_Seq_Alignment)
```

---

plotExonLength-methods

*Method plotExonLength*

---

## Description

This function plot the distribution of the exon length for peaks containing exons.

## Usage

```
plotExonLength(  
  sep,  
  txdb = NULL,  
  save_pdf_prefix = NULL,  
  include_control_regions = TRUE,  
  save_dir = "."  
)
```

```
## S4 method for signature 'SummarizedExomePeak'  
plotExonLength(  
  sep,  
  txdb = NULL,  
  save_pdf_prefix = NULL,  
  include_control_regions = TRUE,  
  save_dir = "."  
)
```

## Arguments

sep	a <a href="#">SummarizedExomePeak</a> object.
txdb	a <a href="#">TxDb</a> object containing the transcript annotation.
save_pdf_prefix	a character if provided, a pdf file with the given name will be saved under the current working directory.
include_control_regions	a logical for whether to include the control regions or not; Default = TRUE.
save_dir	a character for the directory to save the plot; Default = ".".

## Details

If the [SummarizedExomePeaks](#) object contains LFC statistics, the significantly modified peaks with IP to input  $\log_2FC > 0$  and GLM Wald test  $padj < .05$  will be plotted .

If the [SummarizedExomePeaks](#) object contains interactive LFC statistics, both the hyper modification and hypo modification peaks with GLM Wald test p values  $< .05$  will be plotted.

## Value

a ggplot object

## Examples

```
### Make TxDb object from the gff file
library(GenomicFeatures)
GENE_ANNO_GTF = system.file("extdata", "example.gtf", package="exomePeak2")

txdb <- makeTxDbFromGFF(GENE_ANNO_GTF)

### Load the example SummarizedExonPeak object
f1 = system.file("extdata", "sep_ex_mod.rds", package="exomePeak2")

sep <- readRDS(f1)

### Visualize the linear relationships between GC content and normalized reads count under different regions
plotExonLength(sep,txdb)
```

---

plotLfcGC-methods      *Method plotLfcGC*

---

## Description

plotLfcGC plot the scatter plot between GC content and the (differential) modification LFCs.

## Usage

```
plotLfcGC(
  sep,
  bsgenome = NULL,
  txdb = NULL,
  save_pdf_prefix = NULL,
  fragment_length = 100,
  binding_length = 25,
  effective_GC = FALSE,
  save_dir = "."
)

## S4 method for signature 'SummarizedExomePeak'
plotLfcGC(
  sep,
  bsgenome = NULL,
  txdb = NULL,
  save_pdf_prefix = NULL,
  fragment_length = 100,
  binding_length = 25,
  effective_GC = FALSE,
  save_dir = "."
)
```

## Arguments

sep	a <a href="#">SummarizedExomePeak</a> object.
bsgenome	a <a href="#">BSgenome</a> object for the genome sequence, it could be the name of the reference genome recognized by <a href="#">getBSgenome</a> .
txdb	a <a href="#">TxDb</a> object for the transcript annotation, it could be the name of the reference genome recognized by <a href="#">makeTxDbFromUCSC</a> .
save_pdf_prefix	a character, if provided, a pdf file with the given name will be saved under the current directory; Default = NULL.
fragment_length	a numeric value for the expected fragment length in the RNA-seq library; Default = 100.
binding_length	a numeric value for the expected antibody binding length in IP samples; Default = 25.
effective_GC	a logical value of whether to calculate the weighted GC content by the probability of reads alignment; default = FALSE.
save_dir	a character for the directory to save the plot; default ".".

## Details

By default, this function will generate a scatter plot between GC content and the log2FC value. The significant modification sites will be labeled in different colours.

## Value

a ggplot object.

## Examples

```
### Load the example SummarizedExomPeak object
f1 = system.file("extdata", "sep_ex_mod.rds", package="exomePeak2")

sep <- readRDS(f1)

### Visualize the relationship between GC content and the (differential) LFC
plotLfcGC(sep)
```

---

plotReadsGC-methods     *Method plotReadsGC*

---

## Description

plotReadsGC visualizes the local regression curves between the normalized reads abundance and the local GC content.

**Usage**

```

plotReadsGC(
  sep,
  bsgenome = NULL,
  txdb = NULL,
  save_pdf_prefix = NULL,
  fragment_length = 100,
  binding_length = 25,
  effective_GC = FALSE,
  pool_replicates = FALSE,
  save_dir = "."
)

## S4 method for signature 'SummarizedExomePeak'
plotReadsGC(
  sep,
  bsgenome = NULL,
  txdb = NULL,
  save_pdf_prefix = NULL,
  fragment_length = 100,
  binding_length = 25,
  effective_GC = FALSE,
  pool_replicates = FALSE,
  save_dir = "."
)

```

**Arguments**

<code>sep</code>	a <a href="#">SummarizedExomePeak</a> object.
<code>bsgenome</code>	a <a href="#">BSgenome</a> object for the genome sequence, it could be the name of the reference genome recognized by <a href="#">getBSgenome</a> .
<code>txdb</code>	a <a href="#">TxDb</a> object for the transcript annotation, it could be the name of the reference genome recognized by <a href="#">makeTxDbFromUCSC</a> .
<code>save_pdf_prefix</code>	a character, if provided, a pdf file with the given name will be saved under the current directory; Default = NULL.
<code>fragment_length</code>	a numeric value for the expected fragment length in the RNA-seq library; Default = 100.
<code>binding_length</code>	a numeric value for the expected antibody binding length in IP samples; Default = 25.
<code>effective_GC</code>	a logical value of whether to calculate the weighted GC content by the probability of reads alignment; default = FALSE.
<code>pool_replicates</code>	a logical value of whether to pool the replicates in the local regression fit; default = FALSE.
<code>save_dir</code>	a character for the directory to save the plot; default ".".



## Details

The read abundances of both the control and the modification site regions are plotted, the read counts are normalized using the following method:

$$\text{normalizedfeatureabundance} = ((\text{readcount}/\text{sizefactor})/\text{regionlength}) * 500$$

By default, it will use the sequencing depth size factor defined in the [SummarizedExomePeak](#) object, if the sequencing depth size factor is not found, new size factors will be estimated with the default method in [estimateSeqDepth](#).

## Value

a ggplot object

## Examples

```
### Load the example SummarizedExomPeak object
f1 = system.file("extdata", "sep_ex_mod.rds", package="exomePeak2")

sep <- readRDS(f1)

### Visualize the linear relationships between GC content and normalized reads count
plotReadsGC(sep)
```

---

plotSizeFactors-methods

*Method plotSizeFactors*

---

## Description

Method plotSizeFactors

plot the size factors using different strategies.

## Usage

```
plotSizeFactors(sep)
```

```
## S4 method for signature 'SummarizedExomePeak'
plotSizeFactors(sep)
```

## Arguments

sep                    a [SummarizedExomePeak](#) object.

## Value

A ggplot.

**Examples**

```
### Load the example SummarizedExomPeak object
f1 = system.file("extdata", "sep_ex_mod.rds", package="exomePeak2")

sep <- readRDS(f1)

### Visualize the size factors estimated using different regions
plotSizeFactors(sep)
```

---

quiet	<i>Silencing unwanted function output.</i>
-------	--

---

**Description**

Silencing unwanted function output.

**Usage**

```
quiet(x)
```

**Arguments**

x                    any R expression.

**Value**

none.

---

Results-methods	<i>Method Results</i>
-----------------	-----------------------

---

**Description**

Method Results

Report the (Differential) Modification Peaks/Sites and their associated LFC Statistics

**Usage**

```
Results(
  sep,
  cut_off_pvalue = NULL,
  cut_off_padj = 0.05,
  cut_off_log2FC = 0,
  min_num_of_positive = 30,
  expected_direction = c("both", "hyper", "hypo"),
  inhibit_filter = FALSE,
  table_style = c("bed", "granges")
)
```

```
## S4 method for signature 'SummarizedExomePeak'
Results(
  sep,
  cut_off_pvalue = NULL,
  cut_off_padj = 0.1,
  cut_off_log2FC = 0,
  min_num_of_positive = 100,
  expected_direction = c("both", "hyper", "hypo"),
  inhibit_filter = FALSE,
  table_style = c("bed", "granges")
)
```

### Arguments

**sep** a [SummarizedExomePeak](#) object.

**cut\_off\_pvalue** a numeric value for the p value cutoff in the exported result; Default = NULL.

**cut\_off\_padj** a numeric value for the adjusted p value cutoff in the exported result; Default = 0.05.

**cut\_off\_log2FC** a numeric value for the log2 fold change (LFC) cutoff of the exported result, only the sites with abs(LFC) larger than this value are kept; Default = 0.

**min\_num\_of\_positive** a numeric value for the minimum number of reported sites. If the number of remaining sites is less than this number after the filter, additional sites will be reported by the increasing order of the p value to meet this number.

**expected\_direction** a character for the expected direction of the differential modification, could be one in c("hyper", "hypo", "both").  
**hyper** only report the peaks/sites with interactive LFC > 0.  
**hypo** only report the peaks/sites with interactive LFC < 0.  
**both** report the peaks/sites in both directions.  
 This argument is useful when the treated group involves the perturbation of a known writer or eraser protein; Default "both".

**inhibit\_filter** a logical for whether to remove all the filters, this option is useful when quantification on single based site annotation; Default = FALSE.

**table\_style** a character for the style of the table being returned, could be one in c("bed", "granges").  
**bed** The genomic locations in the table are represented by BEDgraph style.  
**granges** The genomic locations in the table are represented by GRanges style.

### Value

a data.frame containing the genomic locations of modification peaks/sites, gene ids, and their statistics.

### Examples

```
### Load the example SummarizedExomPeak object
f1 = system.file("extdata", "sep_ex_mod.rds", package="exomePeak2")
```

```

sep <- readRDS(f1)

### Check the modification peaks/sites statistics.
head(Results(sep))

```

---

scanMeripBAM

*Organize the BAM Files Information of a MeRIP-seq Data Set.*


---

## Description

scanMeripBAM check and organize the BAM files in MeRIP-seq data before peak calling using [exomePeakCalling](#). The library types of the RNA-seq and the filters such as SAM FLAG score are specified in this function.

## Usage

```

scanMeripBAM(
  bam_ip = NULL,
  bam_input = NULL,
  bam_treated_ip = NULL,
  bam_treated_input = NULL,
  paired_end = FALSE,
  library_type = c("unstranded", "1st_strand", "2nd_strand"),
  index_bam = TRUE,
  bam_files = NULL,
  design_ip = NULL,
  design_treatment = NULL,
  mapq = 30L,
  isSecondaryAlignment = FALSE,
  isNotPassingQualityControls = FALSE,
  isDuplicate = FALSE,
  isPaired = NA,
  isProperPair = NA,
  hasUnmappedMate = NA,
  ...
)

```

## Arguments

bam_ip	a character vector for the BAM file directories of the (control) IP samples.
bam_input	a character vector for the BAM file directories of the (control) input samples.
bam_treated_ip	a character vector for the BAM file directories of the treated IP samples.
bam_treated_input	a character vector for the BAM file directories of the treated input samples. If the bam files do not contain treatment group, user should only fill the arguments of BAM_ip and BAM_input.
paired_end	a logical of whether the data comes from the Paired-End Library, TRUE if the data is Paired-End sequencing; default = FALSE.

library_type	<p>a character specifying the protocol type of the RNA-seq library, can be one in <code>c("unstranded", "1st_strand", "2nd_strand")</code>; default = "unstranded".</p> <p><b>unstranded</b> The randomly primed RNA-seq library type, i.e. both the strands generated during the first and the second strand sythesis are sequenced; example: Standard Illumina.</p> <p><b>1st_strand</b> The first strand-specific RNA-seq library, only the strand generated during the first strand sythesis is sequenced; examples: dUTP, NSR, NNSR.</p> <p><b>2nd_strand</b> The second strand-specific RNA-seq library, only the strand generated during the second strand sythesis is sequenced; examples: Ligation, Standard SOLiD.</p>
index_bam	<p>a logical value indicating whether to sort and index BAM files automatically if the bam indexes are not found; default = TRUE.</p> <p>The BAM index files will be named by adding ".bai" after the names of the corresponding BAM files.</p>
bam_files	optional, a character vector for all the BAM file directories, if it is provided, the first 4 arguments above will be ignored.
design_ip	optional, a logical vector indicating the information of IP/input, TRUE represents IP samples.
design_treatment	optional, a logical vector indicating the design of treatment/control, TRUE represents treated samples.
mapq	a non-negative integer specifying the minimum reads mapping quality. BAM records with mapping qualities less than mapq are discarded; default = 30L.
isPaired, isProperPair, hasUnmappedMate, isSecondaryAlignment, isNotPassingQualityControls, isDupl.	arguments specifying the filters on SAM FLAG scores, inherited from <a href="#">ScanBamParam</a> .

## Details

scanMeripBAM takes the input of the BAM file directories for the MeRIP-seq datasets. It first checks the completeness of the BAM files and the BAM indexes. Then, the design information of IP/input and treated/control are returned as a `MeripBamFileList` object. If the BAM file indexes are missing, the BAM files will be automatically indexed with the package `Rsamtools`.

## Value

a `MeripBamFileList` object.

## See Also

[exomePeakCalling](#)

## Examples

```
### Define BAM File Directories

f1 = system.file("extdata", "IP1.bam", package="exomePeak2")
f2 = system.file("extdata", "IP2.bam", package="exomePeak2")
f3 = system.file("extdata", "IP3.bam", package="exomePeak2")
f4 = system.file("extdata", "IP4.bam", package="exomePeak2")
IP_BAM = c(f1,f2,f3,f4)
f1 = system.file("extdata", "Input1.bam", package="exomePeak2")
```

```

f2 = system.file("extdata", "Input2.bam", package="exomePeak2")
f3 = system.file("extdata", "Input3.bam", package="exomePeak2")
INPUT_BAM = c(f1,f2,f3)

f1 = system.file("extdata", "treated_IP1.bam", package="exomePeak2")
TREATED_IP_BAM = c(f1)
f1 = system.file("extdata", "treated_Input1.bam", package="exomePeak2")
TREATED_INPUT_BAM = c(f1)

### For MeRIP-Seq Experiment Without the Treatment Group

MeRIP_Seq_Alignment <- scanMeripBAM(
  bam_ip = IP_BAM,
  bam_input = INPUT_BAM,
  paired_end = FALSE
)

### For MeRIP-Seq Experiment With the Treatment Group

MeRIP_Seq_Alignment <- scanMeripBAM(
  bam_ip = IP_BAM,
  bam_input = INPUT_BAM,
  bam_treated_ip = TREATED_IP_BAM,
  bam_treated_input = TREATED_INPUT_BAM,
  paired_end = FALSE
)

```

---

SummarizedExomePeak-class

*SummarizedExomePeak*

---

## Description

An S4 object defined in `exomePeak2` that summarizes the information of modification peaks/sites, reads counts, size factors, GC contents, and the LFC related statistics.

This class contains [SummarizedExperiment](#).

## Usage

```
SummarizedExomePeak(...)
```

## Arguments

... arguments passed to `new()`.

## Details

### Constructors:

The `SummarizedExomePeak` object can be constructed by 3 functions.

1. [SummarizedExomePeak](#)
2. [exomePeakCalling](#)

### 3. [exomePeak2](#)

#### Accessors:

The SummarizedExomePeak object share all the accessors with the [SummarizedExperiment](#) class.

It has 2 additional accessors:

1. [GCsizeFactors](#)
2. [exomePeak2Results](#)

#### Value

SummarizedExomePeak object

#### Examples

```
# Generate the SummarizedExomePeak object by peak calling

GENE_ANNO_GTF = system.file("extdata", "example.gtf", package="exomePeak2")

f1 = system.file("extdata", "IP1.bam", package="exomePeak2")
f2 = system.file("extdata", "IP2.bam", package="exomePeak2")
f3 = system.file("extdata", "IP3.bam", package="exomePeak2")
f4 = system.file("extdata", "IP4.bam", package="exomePeak2")
IP_BAM = c(f1,f2,f3,f4)
f1 = system.file("extdata", "Input1.bam", package="exomePeak2")
f2 = system.file("extdata", "Input2.bam", package="exomePeak2")
f3 = system.file("extdata", "Input3.bam", package="exomePeak2")
INPUT_BAM = c(f1,f2,f3)

sep <- exomePeak2(bam_ip = IP_BAM,
                 bam_input = INPUT_BAM,
                 gff_dir = GENE_ANNO_GTF,
                 genome = "hg19",
                 paired_end = FALSE)

#Access to the slots in the SummarizedExomePeak object

## Access to reads count
assays(sep)

## Access to the sequencing depth size factors and experimental design
colData(sep)

## Access to the GC content and feature length information
elementMetadata(sep)

## Access to the genomic locations of the modification peaks/sites and the background control regions
rowRanges(sep)

## Access to the feature specific size factors
GCsizeFactors(sep)

## Access to the statistics on (differential) modification LFC
exomePeak2Results(sep)
```

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