

# Package ‘REMP’

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**Title** Repetitive Element Methylation Prediction

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**Description** Machine learning-based tools to predict DNA methylation of locus-specific repetitive elements (RE) by learning surrounding genetic and epigenetic information. These tools provide genomewide and single-base resolution of DNA methylation prediction on RE that are difficult to measure using array-based or sequencing-based platforms, which enables epigenome-wide association study (EWAS) and differentially methylated region (DMR) analysis on RE.

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REMP-package

*Repetitive Element Methylation Prediction*

---

## Description

Machine learning-based tools to predict DNA methylation of locus-specific repetitive elements (RE) by learning surrounding genetic and epigenetic information. These tools provide genomewide and single-base resolution of DNA methylation prediction on RE that are difficult to measure directly using array-based or sequencing-based platforms, which enables epigenome-wide association study (EWAS) and differentially methylated region (DMR) analysis on RE.

### Overview - standard procedure

- Step 1** Start out generating required dataset for prediction using [initREMP](#). The datasets include RE information, RE-CpG (i.e. CpGs located in RE region) information, and gene annotation, which are maintained in a [REMPParcel](#) object. It is recommended to save these generated data to the working directory so they can be used in the future.
- Step 2** Clean Illumina methylation dataset using [groomMethy](#). This function can help identify and fix abnormal values and automatically impute missing values, which are essential for downstream prediction.
- Step 3** Run [remap](#) to predict genome-wide locus specific RE methylation.
- Step 4** Use the built-in accessors and utilities in [REMPProduct](#) object to get or refine the prediction results.

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### References

Zheng Y, Joyce BT, Liu L, Zhang Z, Kibbe WA, Zhang W, Hou L. Prediction of genome-wide DNA methylation in repetitive elements. *Nucleic Acids Res.* 2017;45(15):8697-711. PubMed PMID: 28911103; PMCID: PMC5587781. <http://dx.doi.org/10.1093/nar/gkx587>.

---

Alu.hg19.demo

*Subset of Alu genomic location dataset (hg19)*

---

### Description

A [GRanges](#) dataset containing 500 Alu sequences that have CpGs profiled by both Illumina 450k and EPIC array. The variables are as follows:

### Usage

Alu.hg19.demo

### Format

A [GRanges](#) object.

**Details**

- seqnames: chromosome number
- ranges: hg19 genomic position
- strand: DNA strand
- swScore: Smith Waterman (SW) alignment score
- repName: Alu name
- repClass: Alu class
- repFamily: Alu family
- Index: internal index (meaningless for external use; not communicable between genome builds)

Alu.hg19.demo has the same format as the data object returned by [fetchRMSK](#).

**Value**

A GRanges object with 500 ranges and 3 metadata columns.

**Source**

RepeatMasker database provided by package AnnotationHub

**See Also**

See [fetchRMSK](#) to obtain the complete Alu/L1 dataset.

---

Alu.hg38.demo

*Subset of Alu genomic location dataset (hg38)*

---

**Description**

A [GRanges](#) dataset containing 500 Alu sequences that have CpGs profiled by both Illumina 450k and EPIC array. The variables are as follows:

**Usage**

```
Alu.hg38.demo
```

**Format**

A [GRanges](#) object.

**Details**

- seqnames: chromosome number
- ranges: hg38 genomic position
- strand: DNA strand
- swScore: Smith Waterman (SW) alignment score
- repName: Alu name
- repClass: Alu class
- repFamily: Alu family
- Index: internal index (meaningless for external use; not communicable between genome builds)

Alu.hg38.demo has the same format as the data object returned by [fetchRMSK](#).

**Value**

A GRanges object with 500 ranges and 3 metadata columns.

**Source**

RepeatMasker database (hg38) is provided by UCSC database downloaded from <http://hgdownload.cse.ucsc.edu/goldenpath>.

**See Also**

See [fetchRMSK](#) to obtain the complete Alu/L1 dataset.

---

fetchRefSeqGene	<i>Get RefSeq gene database</i>
-----------------	---------------------------------

---

**Description**

fetchRefSeqGene is used to obtain refSeq gene database provided by AnnotationHub (hg19) or UCSC web database (hg19/hg38).

**Usage**

```
fetchRefSeqGene(  
  annotation.source = c("AH", "UCSC"),  
  genome = c("hg19", "hg38"),  
  mainOnly = FALSE,  
  verbose = FALSE  
)
```

## Arguments

annotation.source	Character parameter. Specify the source of annotation databases, including the RefSeq Gene annotation database and RepeatMasker annotation database. If "AH", the database will be obtained from the AnnotationHub package. If "UCSC", the database will be downloaded from the UCSC website <a href="http://hgdownload.cse.ucsc.edu/goldenp">http://hgdownload.cse.ucsc.edu/goldenp</a> . The corresponding build ("hg19" or "hg38") will be specified in the parameter genome.
genome	Character parameter. Specify the build of human genome. Can be either "hg19" or "hg38". Note that if annotation.source == "AH", only hg19 database is available.
mainOnly	Logical parameter. See details.
verbose	Logical parameter. Should the function be verbose?

## Details

When mainOnly = FALSE, only the transcript location information will be returned, Otherwise, a [GRangesList](#) object containing gene regions information will be added. Gene regions include: 2000 base pair upstream of the transcript start site (\$tss), 5'UTR (\$fiveUTR), coding sequence (\$cds), exon (\$exon), and 3'UTR (\$threeUTR). The index column is an internal index that is used to facilitate data referral, which is meaningless for external use.

## Value

A single [GRanges](#) (for main refgene data) object or a list incorporating both [GRanges](#) object (for main refgene data) and [GRangesList](#) object (for gene regions data).

## Examples

```
if (!exists("refgene.hg19"))
  refgene.hg19 <- fetchRefSeqGene(annotation.source = "AH",
                                genome = "hg19",
                                verbose = TRUE)

refgene.hg19
```

---

fetchRMSK

*Get RE database from RepeatMasker*

---

## Description

fetchRMSK is used to obtain specified RE database from RepeatMasker Database provided by AnnotationHub.

**Usage**

```
fetchRMSK(
  REtype = c("Alu", "L1", "ERV"),
  annotation.source = c("AH", "UCSC"),
  genome = c("hg19", "hg38"),
  verbose = FALSE
)
```

**Arguments**

REtype	Type of RE. Currently "Alu", "L1", and "ERV" are supported.
annotation.source	Character parameter. Specify the source of annotation databases, including the RefSeq Gene annotation database and RepeatMasker annotation database. If "AH", the database will be obtained from the AnnotationHub package. If "UCSC", the database will be downloaded from the UCSC website <a href="http://hgdownload.cse.ucsc.edu/golden">http://hgdownload.cse.ucsc.edu/golden</a> . The corresponding build ("hg19" or "hg38") will be specified in the parameter genome.
genome	Character parameter. Specify the build of human genome. Can be either "hg19" or "hg38".
verbose	Logical parameter. Should the function be verbose?

**Value**

A [GRanges](#) object containing RE database. 'repName' column indicates the RE name; 'swScore' column indicates the SW score; 'Index' is an internal index for RE to facilitate data referral, which is meaningless for external use.

**Examples**

```
L1 <- fetchRMSK(REtype = "L1",
               annotation.source = "AH",
               genome = "hg19",
               verbose = TRUE)

L1
```

---

findREcpg

*Find RE-CpG genomic location given RE ranges information*


---

**Description**

findREcpg is used to obtain RE-CpG genomic location data.

**Usage**

```
findREcpg(
  RE,
  REtype = c("Alu", "L1", "ERV"),
  genome = c("hg19", "hg38"),
  be = NULL,
  verbose = FALSE
)
```

**Arguments**

RE	A <a href="#">GRanges</a> object of RE genomic location database. This can be obtained by <a href="#">fetchRMSK</a> .
REtype	Type of RE. Currently "Alu", "L1", and "ERV" are supported.
genome	Character parameter. Specify the build of human genome. Can be either "hg19" or "hg38". User should make sure the genome build of RE is consistent with this parameter.
be	A <a href="#">BiocParallel</a> object containing back-end information that is ready for parallel computing. This can be obtained by <a href="#">getBackend</a> . If not specified, non-parallel mode is used.
verbose	logical parameter. Should the function be verbose?

**Details**

CpG site is defined as 5'-C-p-G-3'. It is reasonable to assume that the methylation status across all CpG/CpG dyads are concordant. Maintenance methyltransferase exhibits a preference for hemimethylated CpG/CpG dyads (methylated on one strand only). As a result, methylation status of CpG sites in both forward and reverse strands are usually consistent. Therefore, to accommodate the cytosine loci in both strands, the returned genomic ranges cover the 'CG' sequence with width of 2. The 'strand' information indicates the strand of the RE. Locating CpG sites in RE sequences can be computation intensive. It is recommended to get more than one work in the backend for a faster running speed.

**Value**

A [GRanges](#) object containing identified RE-CpG genomic location data.

**Examples**

```
data(Alu.hg19.demo)
RE.CpG <- findREcpg(RE = Alu.hg19.demo,
  REtype = "Alu",
  genome = "hg19",
  verbose = TRUE)
RE.CpG
```

---

getBackend	<i>Get BiocParallel back-end</i>
------------	----------------------------------

---

### Description

getBackend is used to obtain BiocParallel Back-end to allow parallel computing.

### Usage

```
getBackend(ncore, BPPARAM = NULL, verbose = FALSE)
```

### Arguments

ncore	Number of cores used for parallel computing. By default max number of cores available in the machine will be utilized. If ncore = 1, no parallel computing is allowed.
BPPARAM	An optional BiocParallelParam instance determining the parallel back-end to be used during evaluation. If not specified, default back-end in the machine will be used.
verbose	Logical parameter. Should the function be verbose?

### Value

A [BiocParallel](#) object that can be used for parallel computing.

### Examples

```
# Non-parallel mode
be <- getBackend(ncore = 1, verbose = TRUE)
be

# parallel mode (2 workers)
be <- getBackend(ncore = 2, verbose = TRUE)
be
```

---

getGM12878	<i>Get methylation data of HapMap LCL sample GM12878 profiled by Illumina 450k array or EPIC array</i>
------------	--

---

### Description

getGM12878 is used to obtain public available methylation profiling data of HapMap LCL sample GM12878.

**Usage**

```
getGM12878(arrayType = c("450k", "EPIC"), mapGenome = FALSE)
```

**Arguments**

arrayType	Illumina methylation array type. Currently "450k" and "EPIC" are supported. Default = "450k".
mapGenome	Logical parameter. If TRUE, function will return a <a href="#">GenomicRatioSet</a> object instead of a <code>link{RatioSet}</code> object.

**Details**

Illumina 450k data were sourced and curated from ENCODE <http://hgdownload.cse.ucsc.edu/goldenPath/hg19/encodeDCC/wgEncodeHaibMethyl450/wgEncodeHaibMethyl450Gm12878SitesRep1.bed.gz>. Illumina EPIC data were obtained from data package `minfiDataEPIC`.

**Value**

A [RatioSet](#) or [GenomicRatioSet](#) containing beta value and M value of the methylation data.

**Examples**

```
## Not run:
# Get GM12878 methylation data (450k array)
if (!exists("GM12878_450k")) GM12878_450k <- getGM12878("450k")
GM12878_450k

## End(Not run)

# Get GM12878 methylation data (EPIC array)
if (!exists("GM12878_EPIC")) GM12878_EPIC <- getGM12878("EPIC")
GM12878_EPIC
```

---

GRannot

*Annotate genomic ranges data with gene region information.*


---

**Description**

GRannot is used to annotate a [GRanges](#) dataset with gene region information using refseq gene database

**Usage**

```
GRannot(object.GR, refgene, symbol = FALSE, verbose = FALSE)
```

**Arguments**

object.GR	An <a href="#">GRanges</a> object of a genomic location database.
refgene	A complete refGene annotation database returned by <a href="#">fetchRefSeqGene</a> (with parameter <code>mainOnly = FALSE</code> ).
symbol	Logical parameter. Should the annotation return gene symbol?
verbose	Logical parameter. Should the function be verbose?

**Details**

The annotated gene region information includes: protein coding gene (InNM), noncoding RNA gene (InNR), 2000 base pair upstream of the transcript start site (InTSS), 5'UTR (In5UTR), coding sequence (InCDS), exon (InExon), and 3'UTR (In3UTR). The intergenic and intron regions can then be represented by the combination of these region data. The number shown in these columns represent the row number or 'index' column in the main refgene database obtained by [fetchRefSeqGene](#).

**Value**

A [GRanges](#) or a [GRangesList](#) object containing refSeq Gene database.

**Examples**

```
data(Alu.hg19.demo)
if (!exists("refgene.hg19"))
  refgene.hg19 <- fetchRefSeqGene(annotation.source = "AH",
                                genome = "hg19",
                                verbose = TRUE)
Alu.hg19.demo.refGene <- GRannot(Alu.hg19.demo, refgene.hg19, verbose = TRUE)
Alu.hg19.demo.refGene
```

---

grooMethy

*Groom methylation data to fix potential data issues*

---

**Description**

grooMethy is used to automatically detect and fix data issues including zero beta value, missing value, and infinite value.

**Usage**

```
grooMethy(
  methyDat,
  Seq.GR = NULL,
  impute = TRUE,
  imputebyrow = TRUE,
  mapGenome = FALSE,
  verbose = FALSE
)
```

## Arguments

methyDat	A <a href="#">RatioSet</a> , <a href="#">GenomicRatioSet</a> , <a href="#">DataFrame</a> , <code>data.table</code> , <code>data.frame</code> , or matrix of Illumina BeadChip methylation data (450k or EPIC array) or Illumina methylation percentage estimates by sequencing. If the data are prepared as a <code>data.frame</code> or alike format, for Illumina array data, please make sure there is a column or row names are available to indicate the Illumina probe names (i.e. <code>cg00000029</code> ); for sequencing methylation data, please provide the corresponding CpG location information in <code>Seq.GR</code> .
Seq.GR	A <a href="#">GRanges</a> object containing genomic locations of the CpGs profiled by sequencing platforms. This parameter should not be <code>NULL</code> if the input methylation data <code>methyDat</code> are obtained by sequencing platform. The order of <code>Seq.GR</code> should match the order of <code>methyDat</code> . Note that the genomic location can be in either <code>hg19</code> or <code>hg38</code> build. See details.
impute	If <code>TRUE</code> , K-Nearest Neighbouring imputation will be applied to fill the missing values. Default = <code>TRUE</code> . See Details.
imputebyrow	If <code>TRUE</code> , missing values will be imputed using similar values in row (i.e., across samples); if <code>FALSE</code> , missing values will be imputed using similar values in column (i.e., across CpGs). Default is <code>TRUE</code> .
mapGenome	Logical parameter. If <code>TRUE</code> , function will return a <a href="#">GenomicRatioSet</a> object instead of a <a href="#">RatioSet</a> . This function is not applicable for sequencing data.
verbose	Logical parameter. Should the function be verbose?

## Details

For methylation data in beta value, if zero/one value exists, the logit transformation from beta to M value will produce infinite value. Therefore, zero/one beta value will be replaced with the smallest non-zero beta/largest non-one beta value found in the dataset. `grooMethy` can also handle missing value (i.e. `NA` or `NaN`) using KNN imputation (see [impute.knn](#)). The infinite value will be also treated as missing value for imputation. If the original dataset is in beta value, `grooMethy` will first transform it to M value before imputation is carried out. If the imputed value is out of the original range (which is possible when `imputebyrow = FALSE`), mean value will be used instead. Warning: imputed values for multimodal distributed CpGs (across samples) may not be correct. Please check package `ENmix` to identify the CpGs with multimodal distribution. Please note that `grooMethy` is also embedded in [remap](#) so the user can run [remap](#) directly without explicitly running `grooMethy`. For sequencing methylation data, please specify the genomic location of CpGs in a [GenomicRanges](#) object and specify it in `Seq.GR`. For an example of `Seq.GR`, Please run `minfi::getLocations(IlluminaHumanMethylation450kanno.ilmn12.hg19)` (the row names of the CpGs in `Seq.GR` can be `NULL`). The user should make sure the genome build of `Seq.GR` match the build specified in `genome` parameter of function [initREMP](#) and [remprofile](#) (default is `"hg19"`).

## Value

A [RatioSet](#) or [GenomicRatioSet](#) containing beta value and M value of the methylation data.

## Examples

```
# Get GM12878 methylation data (450k array)
```

```

if (!exists("GM12878_450k")) GM12878_450k <- getGM12878("450k")
GM12878_450k <- grooMethy(GM12878_450k, verbose = TRUE)

# Also works if data input is a matrix
grooMethy(minfi::getBeta(GM12878_450k), verbose = TRUE)

```

---

initREMP

*RE Annotation Database Initialization*


---

### Description

initREMP is used to initialize annotation database for RE methylation prediction. Three RE types in human, Alu element (Alu), LINE-1 (L1), and endogenous retrovirus (ERV) are available.

### Usage

```

initREMP(
  arrayType = c("450k", "EPIC", "Sequencing"),
  REtype = c("Alu", "L1", "ERV"),
  annotation.source = c("AH", "UCSC"),
  genome = c("hg19", "hg38"),
  RE = NULL,
  Seq.GR = NULL,
  ncore = NULL,
  BPPARAM = NULL,
  export = FALSE,
  work.dir = tempdir(),
  verbose = FALSE
)

```

### Arguments

arrayType	ILLUMINA methylation array type. Currently "450k", "EPIC", and "Sequencing" are supported. Default = "450k".
REtype	Type of RE. Currently "Alu", "L1", and "ERV" are supported.
annotation.source	Character parameter. Specify the source of annotation databases, including the RefSeq Gene annotation database and RepeatMasker annotation database. If "AH", the database will be obtained from the AnnotationHub package. If "UCSC", the database will be downloaded from the UCSC website <a href="http://hgdownload.cse.ucsc.edu/golden">http://hgdownload.cse.ucsc.edu/golden</a> . The corresponding build ("hg19" or "hg38") can be specified in the parameter genome.
genome	Character parameter. Specify the build of human genome. Can be either "hg19" or "hg38". Note that if annotation.source == "AH", only hg19 database is available.

RE	A <a href="#">GRanges</a> object containing user-specified RE genomic location information. If NULL, the function will retrieve RepeatMasker RE database from <a href="#">AnnotationHub</a> (build hg19) or download the database from UCSC website (build hg19/hg38).
Seq.GR	A <a href="#">GRanges</a> object containing genomic locations of the CpGs profiled by sequencing platforms. This parameter should not be NULL if <code>arrayType == 'Sequencing'</code> . Note that the genomic location can be in either hg19 or hg38 build. See details.
ncore	Number of cores used for parallel computing. By default max number of cores available in the machine will be utilized. If <code>ncore = 1</code> , no parallel computing is allowed.
BPPARAM	An optional <a href="#">BiocParallelParam</a> instance determining the parallel back-end to be used during evaluation. If not specified, default back-end in the machine will be used.
export	Logical. Should the returned <a href="#">REMPParcel</a> object be saved to local machine? See Details.
work.dir	Path to the directory where the generated data will be saved. Valid when <code>export = TRUE</code> . If not specified and <code>export = TRUE</code> , temporary directory <code>tempdir()</code> will be used.
verbose	Logical parameter. Should the function be verbose?

### Details

Currently, we support two major types of RE in the human genome, Alu and L1. The main purpose of `initREMP` is to generate and annotate CpG/RE data using the refSeq Gene (hg19) annotation database (provided by [AnnotationHub](#)). These annotation data are crucial to RE methylation prediction in `remP`. Once generated, the data can be reused in the future (data can be very large). Therefore, we recommend the user to save the output from `initREMP` to the local machine, so that user only need to run this function once as long as there is no change to the RE database. To minimize the size of the resulting data file, the generated annotation data are only for REs that contain RE-CpGs with neighboring profiled CpGs. By default, the neighboring CpGs are confined within 1200 bp flanking window. This window size can be modified using `remP_options`. Note that the refSeq Gene database from UCSC is dynamic (updated periodically) and reflecting the latest knowledge of gene, whereas the database from [AnnotationHub](#) is static and classic. Using different sources will have a slight impact on the prediction results of RE methylation and gene annotation of final results. For sequencing methylation data, please specify the genomic location of CpGs in a `GenomicRanges` object and specify it in `Seq.GR`. For an example of `Seq.GR`, Please run `minfi::getLocations(IlluminaHumanMethylation450kanno.ilmn12.hg19)` (the row names of the CpGs in `Seq.GR` can be NULL). The user should make sure the genome build of `Seq.GR` match the build specified in `genome` parameter (default is "hg19").

### Value

An [REMPParcel](#) object containing data needed for RE methylation prediction.

### See Also

See `remP` for RE methylation prediction.

## Examples

```
if (!exists("remparc1")) {
  data(Alu.hg19.demo)
  remparc1 <- initREMP(arrayType = "450k",
                      REtype = "Alu",
                      annotation.source = "AH",
                      genome = "hg19",
                      RE = Alu.hg19.demo,
                      ncore = 1,
                      verbose = TRUE)
}
```

---

remp

*Repetitive element methylation prediction*


---

## Description

remp is used to predict genomewide methylation levels of locus-specific repetitive elements (RE). Two major RE types in human, Alu element (Alu) and LINE-1 (L1) are available.

## Usage

```
remp(
  methyDat = NULL,
  REtype = c("Alu", "L1", "ERV"),
  Seq.GR = NULL,
  parcel = NULL,
  work.dir = tempdir(),
  win = 1000,
  method = c("rf", "xgbTree", "svmLinear", "svmRadial", "naive"),
  autoTune = TRUE,
  param = NULL,
  seed = NULL,
  ncore = NULL,
  BPPARAM = NULL,
  verbose = FALSE
)
```

## Arguments

**methyDat** A [RatioSet](#), [GenomicRatioSet](#), [DataFrame](#), `data.table`, `data.frame`, or matrix of Illumina BeadChip methylation data (450k or EPIC array) or Illumina methylation percentage estimates by sequencing. See [Details](#). Alternatively, user can also specify a pre-built data template (see [rempTemplate](#)). remp to carry out the prediction. See [rempTemplate](#). With template specified, `methyDat`, `REtype`, `parcel`, and `work.dir` can be skipped.

REtype	Type of RE. Currently "Alu", "L1", and "ERV" are supported. If NULL, the type of RE will be extracted from parcel.
Seq.GR	A <a href="#">GRanges</a> object containing genomic locations of the CpGs profiled by sequencing platforms. This parameter should not be NULL if the input methylation data methyDat are obtained by sequencing. Note that the genomic location can be in either hg19 or hg38 build. See details in <a href="#">initREMP</a> .
parcel	An <a href="#">REMPParcel</a> object containing necessary data to carry out the prediction. If NULL, REtype must specify a type of RE so that the function can search the .rds data file in work.dir exported by <a href="#">initREMP</a> (with export = TRUE) or <a href="#">saveParcel</a> .
work.dir	Path to the directory where the annotation data generated by <a href="#">initREMP</a> are saved. Valid when the argument parcel is missing. If not specified, temporary directory tempdir() will be used. If specified, the directory path has to be the same as the one specified in <a href="#">initREMP</a> or in <a href="#">saveParcel</a> .
win	An integer specifying window size to confine the upstream and downstream flanking region centered on the predicted CpG in RE for prediction. Default = 1000. See Details.
method	Name of model/approach for prediction. Currently "rf" (Random Forest), "xgbTree" (Extreme Gradient Boosting), "svmLinear" (SVM with linear kernel), "svmRadial" (SVM with radial kernel), and "naive" (carrying over methylation values of the closest CpG site) are available. Default = "rf" (Random Forest). See Details.
autoTune	Logical parameter. If TRUE, a 3-time repeated 5-fold cross validation will be performed to determine the best model parameter. If FALSE, the param option (see below) must be specified. Default = TRUE. Auto-tune will be disabled using Random Forest. See Details.
param	A list specifying fixed model tuning parameter(s) (not applicable for Random Forest, see Details). For Extreme Gradient Boosting, param list must contain '\$nrounds', '\$max_depth', '\$eta', '\$gamma', '\$colsample_bytree', '\$min_child_weight', and '\$subsample'. See xgbTree in package caret. For SVM, param list must contain '\$C' (cost) for linear kernel or '\$sigma' and '\$C' for radial basis function kernel. This parameter is valid only when autoTune = FALSE.
seed	Random seed for Random Forest model for reproducible prediction results. Default is NULL, which generates a seed.
ncore	Number of cores used for parallel computing. By default, max number of cores available in the machine will be utilized. If ncore = 1, no parallel computing is allowed.
BPPARAM	An optional <a href="#">BiocParallelParam</a> instance determining the parallel back-end to be used during evaluation. If not specified, default back-end in the machine will be used.
verbose	Logical parameter. Should the function be verbose?

## Details

Before running `remp`, user should make sure the methylation data have gone through proper quality control, background correction, and normalization procedures. Both beta value and M value are

allowed. Rows represents probes and columns represents samples. For array data, please make sure to have row names that specify the Illumina probe ID (i.e. cg00000029). For sequencing data, please provide the genomic location of CpGs in a [GRanges](#) object and specify it using `Seq.GR` parameter. `win = 1000` is based on previous findings showing that neighboring CpGs are more likely to be co-modified within 1000 bp. User can specify narrower window size for slight improvement of prediction accuracy at the cost of less predicted RE. Window size greater than 1000 is not recommended as the machine learning models would not be able to learn much useful information for prediction but introduce noise. Random Forest model (`method = "rf"`) is recommended as it offers more accurate prediction and it also enables prediction reliability functionality. Prediction reliability is estimated by conditional standard deviation using Quantile Regression Forest. Please note that if parallel computing is allowed, parallel Random Forest (powered by package [ranger](#)) will be used automatically. The performance of Random Forest model is often relatively insensitive to the choice of `mtry`. Therefore, auto-tune will be turned off using Random Forest and `mtry` will be set to one third of the total number of predictors. For SVM, if `autoTune = TRUE`, preset tuning parameter search grid can be access and modified using [remp\\_options](#).

### Value

A [REMPProduct](#) object containing predicted RE methylation results.

### See Also

See [initREMP](#) to prepare necessary annotation database before running `remp`.

### Examples

```
# Obtain example Illumina example data (450k)
if (!exists("GM12878_450k"))
  GM12878_450k <- getGM12878("450k")

# Make sure you have run 'initREMP' first. See ?initREMP.
if (!exists("remparcel")) {
  data(Alu.hg19.demo)
  remparcel <- initREMP(arrayType = "450k",
                       REtype = "Alu",
                       annotation.source = "AH",
                       genome = "hg19",
                       RE = Alu.hg19.demo,
                       ncore = 1,
                       verbose = TRUE)
}

# With data template pre-built. See ?rempTemplate.
if (!exists("template"))
  template <- rempTemplate(GM12878_450k,
                          parcel = remparcel,
                          win = 1000,
                          verbose = TRUE)

# Run remp with pre-built template:
remp.res <- remp(template, ncore = 1)
```

```
# Or run remp without pre-built template (identical results):
## Not run:
  remp.res <- remp(GM12878_450k,
                  REtype = "Alu",
                  parcel = remparcel,
                  ncore = 1,
                  verbose = TRUE)

## End(Not run)

remp.res
details(remp.res)
rempB(remp.res) # Methylation data (beta value)

# Extract CpG location information.
# This accessor is inherit from class 'RangedSummarizedExperiment')
rowRanges(remp.res)

# RE annotation information
rempAnnot(remp.res)

# Add gene annotation
remp.res <- decodeAnnot(remp.res, type = "symbol")
rempAnnot(remp.res)

# (Recommended) Trim off less reliable prediction
remp.res <- rempTrim(remp.res)

# Obtain RE-level methylation (aggregate by mean)
remp.res <- rempAggregate(remp.res)
rempB(remp.res) # Methylation data (beta value)

# Extract RE location information
rowRanges(remp.res)

# Density plot across predicted RE
rempplot(remp.res)
```

---

REMPParcel-class

*REMPParcel instances*

---

## Description

REMPParcel is a container class to organize required datasets for RE methylation prediction generated from [initREMP](#) and used in [remp](#).

**Usage**

```
REMPParcel(  
  REtype = "Unknown",  
  genome = "Unknown",  
  platform = "Unknown",  
  RefGene = GRanges(),  
  RE = GRanges(),  
  RECPG = GRanges(),  
  ILMN = GRanges()  
)  
  
getParcelInfo(object)  
  
getRefGene(object)  
  
getRE(object)  
  
getRECPG(object)  
  
getILMN(object, ...)  
  
saveParcel(object, ...)  
  
## S4 method for signature 'REMPParcel'  
saveParcel(object, work.dir = tempdir(), verbose = FALSE, ...)  
  
## S4 method for signature 'REMPParcel'  
getParcelInfo(object)  
  
## S4 method for signature 'REMPParcel'  
getRefGene(object)  
  
## S4 method for signature 'REMPParcel'  
getRE(object)  
  
## S4 method for signature 'REMPParcel'  
getRECPG(object)  
  
## S4 method for signature 'REMPParcel'  
getILMN(object, REonly = FALSE)
```

**Arguments**

REtype	Type of RE ("Alu", "L1", or "ERV").
genome	Specify the build of human genome. Can be either "hg19" or "hg38".
platform	Illumina methylation profiling platform ("450k" or "EPIC").
RefGene	refSeq gene annotation data, which can be obtained by <a href="#">fetchRefSeqGene</a> .

RE	Annotated RE genomic range data, which can be obtained by <a href="#">fetchRMSK</a> and annotated by <a href="#">GRannot</a> .
RECPG	Genomic range data of annotated CpG site identified in RE DNA sequence, which can be obtained by <a href="#">findRECPG</a> and annotated by <a href="#">GRannot</a> .
ILMN	Illumina CpG probe genomic range data.
object	A REMPParcel object.
...	For <code>saveParcel</code> : other parameters to be passed to the <code>saveRDS</code> method. See <a href="#">saveRDS</a> .
<code>work.dir</code>	For <code>saveParcel</code> : path to the directory where the generated data will be saved. If not specified, temporary directory <code>tempdir()</code> will be used.
<code>verbose</code>	For <code>saveParcel</code> : logical parameter. Should the function be verbose?
<code>REonly</code>	For <code>getILMN</code> : see <code>Accessors</code> .

**Value**

An object of class `REMPParcel` for the constructor.

**Accessors**

- `getParcelInfo(object)` Return data type, RE type, and flanking window size information of the parcel.
- `getRefGene(object)` Return RefSeq gene annotation data.
- `getRE(object)` Return RE genomic location data for prediction (annotated by refSeq gene database).
- `getRECPG(object)` Return RE-CpG genomic location data for prediction.
- `getILMN(object, REonly = FALSE)` Return Illumina CpG probe genomic location data for prediction (annotated by refSeq gene database). If `REonly = TRUE`, only probes within RE region are returned.

**Utilities**

- `saveParcel(object, work.dir = tempdir(), verbose = FALSE, ...)` Save the object to local machine.

**Examples**

```
showClass("REMPParcel")
```

---

REMPProduct-class	<i>REMPProduct instances</i>
-------------------	------------------------------

---

### Description

Class `REMPProduct` is to maintain RE methylation prediction results. `REMPProduct` inherits `Bioconductor's RangedSummarizedExperiment` class.

### Usage

```
REMPProduct(  
  REtype = "Unknown",  
  genome = "Unknown",  
  platform = "Unknown",  
  win = "Unknown",  
  predictModel = "Unknown",  
  QCModel = "Unknown",  
  rempM = NULL,  
  rempB = NULL,  
  rempQC = NULL,  
  cpgranges = GRanges(),  
  sampleInfo = DataFrame(),  
  REannotation = GRanges(),  
  RECpG = GRanges(),  
  regionCode = DataFrame(),  
  refGene = GRanges(),  
  varImp = DataFrame(),  
  REStats = DataFrame(),  
  GeneStats = DataFrame(),  
  Seed = NULL  
)  
  
rempM(object)  
  
rempB(object)  
  
rempQC(object)  
  
rempAnnot(object)  
  
rempImp(object)  
  
rempStats(object)  
  
rempplot(object, ...)  
  
details(object)
```

```

decodeAnnot(object, ...)

rempTrim(object, ...)

rempAggregate(object, ...)

rempCombine(object1, object2)

## S4 method for signature 'REMPProduct'
rempM(object)

## S4 method for signature 'REMPProduct'
rempB(object)

## S4 method for signature 'REMPProduct'
rempQC(object)

## S4 method for signature 'REMPProduct'
rempImp(object)

## S4 method for signature 'REMPProduct'
rempAnnot(object)

## S4 method for signature 'REMPProduct'
rempStats(object)

## S4 method for signature 'REMPProduct'
rempIplot(object, type = c("individual", "overall"), ...)

## S4 method for signature 'REMPProduct'
details(object)

## S4 method for signature 'REMPProduct'
decodeAnnot(object, type = c("symbol", "entrez"), ncore = 1, BPPARAM = NULL)

## S4 method for signature 'REMPProduct'
rempTrim(object, threshold = 1.7, missingRate = 0.2)

## S4 method for signature 'REMPProduct'
rempAggregate(object, NCpG = 2, ncore = 1, BPPARAM = NULL)

## S4 method for signature 'REMPProduct,REMPProduct'
rempCombine(object1, object2)

```

### Arguments

REtype            Type of RE ("Alu", "L1", or "ERV").

genome	Specify the build of human genome. Can be either "hg19" or "hg38".
platform	Illumina methylation profiling platform ("450k" or "EPIC").
win	Flanking window size of the predicting RE-CpG.
predictModel	Name of the model used for prediction.
QCModel	Name of the model used for prediction quality evaluation.
rempM	Predicted methylation level in M value.
rempB	Predicted methylation level in beta value (optional).
rempQC	Prediction quality scores, which is available only when Random Forest model is used in <a href="#">remp</a> .
cpgRanges	Genomic ranges of the predicting RE-CpG.
sampleInfo	Sample information.
REannotation	Annotation data for the predicting RE.
RECpG	Annotation data for the RE-CpG profiled by Illumina platform.
regionCode	Internal index code defined in refGene for gene region indicators.
refGene	refSeq gene annotation data, which can be obtained by <a href="#">fetchRefSeqGene</a> .
varImp	Importance of the predictors.
REStats	RE coverage statistics, which is internally generated in <a href="#">remp</a> .
GeneStats	Gene coverage statistics, which is internally generated in <a href="#">remp</a> .
Seed	Random seed for Random Forest model for reproducible prediction results.
object	A REMProduct object.
...	For plot: <a href="#">graphical parameters</a> to be passed to the plot method.
object1	A REMProduct object.
object2	A REMProduct object.
type	For plot and decodeAnnot: see Utilities.
ncore	For decodeAnnot and rempAggregate: number of cores used for parallel computing. By default no parallel computing is allowed (ncore = 1).
BPPARAM	For decodeAnnot and rempAggregate: an optional <a href="#">BiocParallelParam</a> instance determining the parallel back-end to be used during evaluation. If not specified, default back-end in the machine will be used.
threshold	For rempTrim: see Utilities.
missingRate	For rempTrim: see Utilities.
NCpG	For rempAggregate: see Utilities.

### Value

An object of class REMProduct for the constructor.

**Accessors**

- remM(object) Return M value of the prediction.
- remB(object) Return beta value of the prediction.
- remQC(object) Return prediction quality scores.
- remImp(object) Return relative importance of predictors.
- remStats(object) Return RE and gene coverage statistics.
- remAnnot(object) Return annotation data for the predicted RE.

**Utilities**

- remplot(object, type = c("individual", "overall"), ...) Make a density plot of predicted methylation (beta values) in the REMProduct object. If type = "individual", density curves will be plotted for each of the samples; If type = "overall", one density curve of the mean methylation level across the samples will be plotted. Default type = "individual".
- details(object) Display detailed descriptive statistics of the prediction results.
- decodeAnnot(object, type = c("symbol", "entrez"), ncore = NULL, BPPARAM = NULL) Decode the RE annotation data by Gene Symbol (when type = "Symbol") or Entrez Gene (when type = "Entrez"). Default type = "Symbol". Annotation data are provided by [org.Hs.eg.db](http://org.hs.eg.db).
- remTrim(object, threshold = 1.7, missingRate = 0.2) Any predicted CpG values with quality score < threshold (default = 1.7, specified by threshold = 1.7) will be replaced with NA. CpGs contain more than missingRate \* 100 rate across samples will be discarded. Relevant summary statistics will be re-evaluated.
- remAggregate(object, NCpG = 2, ncore = NULL, BPPARAM = NULL) Aggregate the predicted RE-CpG methylation by RE using mean. To ensure the reliability of the aggregation, by default only RE with at least 2 predicted CpG sites (specified by NCpG = 2) will be aggregated.
- remCombine(object1, object2) Combine two REMProduct objects by column.

**Examples**

```
showClass("REMPProduct")
```

---

```
remprofile
```

```
Extract DNA methylation data profiled in RE
```

---

**Description**

remprofile is used to extract profiled methylation of CpG sites in RE.

**Usage**

```
remprofile(
  methyDat,
  REtype = c("Alu", "L1", "ERV"),
  annotation.source = c("AH", "UCSC"),
  genome = c("hg19", "hg38"),
  Seq.GR = NULL,
  RE = NULL,
  impute = FALSE,
  imputebyrow = TRUE,
  verbose = FALSE
)
```

**Arguments**

methyDat	A <a href="#">RatioSet</a> , <a href="#">GenomicRatioSet</a> , <a href="#">DataFrame</a> , <code>data.table</code> , <code>data.frame</code> , or matrix of Illumina BeadChip methylation data (450k or EPIC array) or Illumina methylation percentage estimates by sequencing.
REtype	Type of RE. Currently "Alu", "L1", and "ERV" are supported.
annotation.source	Character parameter. Specify the source of annotation databases, including the RefSeq Gene annotation database and RepeatMasker annotation database. If "AH", the database will be obtained from the AnnotationHub package. If "UCSC", the database will be downloaded from the UCSC website <a href="http://hgdownload.cse.ucsc.edu/goldenp">http://hgdownload.cse.ucsc.edu/goldenp</a> . The corresponding build ("hg19" or "hg38") will be specified in the parameter genome.
genome	Character parameter. Specify the build of human genome. Can be either "hg19" or "hg38". For 450k/EPIC array, "hg19" is used more often while specifying "hg38" will lift over the Illumina CpG probe location to build "hg38". For sequencing data, please make sure the specified genome build is consistent with the actual genome build of Seq.GR.
Seq.GR	A <a href="#">GRanges</a> object containing genomic locations of the CpGs profiled by sequencing platforms. This parameter should not be NULL if the input methylation data methyDat are obtained by sequencing. Note that the genomic location can be in either hg19 or hg38 build. The user should make sure the parameter genome is correctly specified.
RE	A <a href="#">GRanges</a> object containing user-specified RE genomic location information. If NULL, the function will retrieve RepeatMasker RE database from <a href="#">AnnotationHub</a> (build hg19) or download the database from UCSC website (build hg19/hg38).
impute	Parameter used by <a href="#">groomMethy</a> . If TRUE, K-Nearest Neighbouring imputation will be applied to fill the missing values. Default = FALSE.
imputebyrow	Parameter used by <a href="#">groomMethy</a> . If TRUE, missing values will be imputed using similar values in row (i.e., across samples); if FALSE, missing values will be imputed using similar values in column (i.e., across CpGs). Default is TRUE.
verbose	Logical parameter. Should the function be verbose?

**Value**

A `REMPProduct` object containing profiled RE methylation results.

**Examples**

```
data(Alu.hg19.demo)
if (!exists("GM12878_450k")) GM12878_450k <- getGM12878("450k")
remprofile.res <- remprofile(GM12878_450k,
                             RType = "Alu",
                             annotation.source = "AH",
                             genome = "hg19",
                             RE = Alu.hg19.demo,
                             verbose = TRUE)

details(remprofile.res)
remprB(remprofile.res) # Methylation data (beta value)

remprofile.res <- rempAggregate(remprofile.res)
details(remprofile.res)
remprB(remprofile.res) # Methylation data (beta value)
```

---

rempTemplate

*Prepare data template for REMP*


---

**Description**

`rempTemplate` is used to build a set of data templates for prediction. The data templates include RE-CpGs and their methylation data for model training, neighboring CpGs of RE-CpGs and their methylation data for model prediction, and other necessary information about the prediction. This function is useful when one needs to experiment different tuning parameters so that these pre-built data templates can be re-used and substantially improve efficiency.

**Usage**

```
rempTemplate(
  methyDat = NULL,
  Seq.GR = NULL,
  parcel = NULL,
  win = 1000,
  verbose = FALSE
)
```

**Arguments**

`methyDat` A `RatioSet`, `GenomicRatioSet`, `DataFrame`, `data.table`, `data.frame`, or matrix of Illumina BeadChip methylation data (450k or EPIC array) or Illumina methylation percentage estimates by sequencing.

Seq.GR	A <a href="#">GRanges</a> object containing genomic locations of the CpGs profiled by sequencing platforms. This parameter should not be NULL if the input methylation data <code>methyDat</code> are obtained by sequencing. Note that the genomic location can be in either hg19 or hg38 build, but must be consistent with the build as <code>parcel</code> . See details in <a href="#">initREMP</a> .
parcel	An <a href="#">REMPParcel</a> object containing necessary data to carry out the prediction. If NULL, the function will search the <code>.rds</code> data file in <code>work.dir</code> exported by <a href="#">initREMP</a> (with <code>export = TRUE</code> ) or <a href="#">saveParcel</a> .
win	An integer specifying window size to confine the upstream and downstream flanking region centered on the predicted CpG in RE for prediction. Default = 1000.
verbose	Logical parameter. Should the function be verbose?

### Value

A template object containing a [GRanges](#) object of neighboring CpGs of RE-CpGs to be predicted (`$NBCpG_GR`) and their methylation dataset matrix (`$NBCpG_methyDat`); a [GRanges](#) object of RE-CpGs for model training (`$REcPcG_GR`) and their methylation dataset matrix (`$REcPcG_methyDat`); [GRanges](#) objects of RefSeq Gene database (`$refgene`) and RE (`$RE`) that are extracted from the `parcel` input; a string of RE type (`$REtype`) and a string of methylation platform (`$arrayType`). Note: the subset operator `[]` is supported.

### Examples

```
if (!exists("GM12878_450k"))
  GM12878_450k <- getGM12878("450k")
if (!exists("remparcel")) {
  data(Alu.hg19.demo)
  remparcel <- initREMP(arrayType = "450k",
                       REtype = "Alu",
                       annotation.source = "AH",
                       genome = "hg19",
                       RE = Alu.hg19.demo,
                       ncore = 1,
                       verbose = TRUE)
}

template <- rempTemplate(GM12878_450k,
                        parcel = remparcel,
                        win = 1000,
                        verbose = TRUE)

template

## To make a subset
template[1]
```

---

remp_options	<i>Set or get options for REMP package</i>
--------------	--

---

**Description**

Tools to manage global setting options for REMP package.

**Usage**

```
remp_options(...)
```

```
remp_reset()
```

**Arguments**

... Option names to retrieve option values or [key]=[value] pairs to set options.

**Value**

NULL

**Supported options**

The following options are supported

- .default.AluFamily.grep Regular expression for 'grep' to extract Alu family to be included in the prediction.
- .default.L1Family.grep Regular expression for 'grep' to extract L1 family to be included in the prediction.
- .default.ERVFamily.grep Regular expression for 'grep' to extract ERV family to be included in the prediction.
- .default.chr List of human chromosome.
- .default.GM12878.450k.URL URL to download GM12878 450k methylation profiling data.
- .default.RMSK.hg19.URL URL to download RepeatMasker database in hg19 genome.
- .default.RMSK.hg38.URL URL to download RepeatMasker database in hg38 genome.
- .default.refGene.hg19.URL URL to download refSeq gene database in hg19 genome.
- .default.refGene.hg38.URL URL to download refSeq gene database in hg38 genome.
- .default.AH.repeatmasker.hg19 AnnotationHub data ID linked to RepeatMasker annotation database (Mar 2020, build hg19).
- .default.AH.repeatmasker.hg38 AnnotationHub data ID linked to RepeatMasker annotation database (Sep 2021, build hg38).
- .default.AH.refgene.hg19 AnnotationHub data ID linked to refSeq gene database (build hg19)
- .default.AH.hg38ToHg19.over.chain AnnotationHub hg38 to hg19 liftover chain data ID.
- .default.AH.hg19ToHg38.over.chain AnnotationHub hg19 to hg38 liftover chain data ID.

- .default.TSS.upstream Define the upstream range of transcription start site region.
- .default.TSS.downstream Define the downstream range of transcription start site region.
- .default.max.flankWindow Define the max size of the flanking window surrounding the predicted RE-CpG.
- .default.27k.total.probes Total number of probes designed in Illumina 27k array.
- .default.450k.total.probes Total number of probes designed in Illumina 450k array.
- .default.epic.total.probes Total number of probes designed in Illumina EPIC array.
- .default.450k.annotation A character string associated with the Illumina 450k array annotation dataset.
- .default.epic.annotation A character string associated with the Illumina EPIC array annotation dataset.
- .default.genomicRegionColNames Define the names of the genomic regions for prediction.
- .default.predictors Define the names of predictors for RE methylation prediction.
- .default.svmLinear.tune Define the default C (Cost) parameter for Support Vector Machine (SVM) using linear kernel.
- .default.svmRadial.tune Define the default parameters (C and sigma) for SVM using Radial basis function kernel.
- .default.xgbTree.tune Define the default parameters (nrounds, eta, max\_depth, gamma, colsample\_bytree, min\_child\_weight, and subsample) for Extreme Gradient Boosting.

## Examples

```
# Display all default settings
remp_options()

# Display a specified setting
remp_options(".default.max.flankWindow")

# Change default maximum flanking window size to 2000
remp_options(.default.max.flankWindow = 2000)

# Reset all options
remp_reset()
```

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