An Introduction to the REMP Package

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1 Introduction

REMP predicts DNA methylation of locus-specific repetitive elements (RE) by learning surrounding genetic and epigenetic information. *REMP* provides genomewide single-base resolution of DNA methylation on RE that is 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. *REMP* also provides handy tool to extract methylation data of CpGs that are located within RE sequences.

REMP supports both Illumina methylation BeadChip array platforms (450k and EPIC) and sequencing platforms (e.g. TruSeq Methyl Capture EPIC). Both genome build hg19 and hg38 are supported.

2 Installation

Install *REMP* (release version):

```
> if (!requireNamespace("BiocManager", quietly=TRUE))
```

```
+ install.packages("BiocManager")
```

```
> BiocManager::install("REMP")
```

To install devel version:

> library(devtools)

```
> install_github("YinanZheng/REMP")
```

Load *REMP* into the workspace:

```
> library(REMP)
```

3 REMP: Repetitive Element Methylation Prediction

Currently *REMP* supports Human (hg19/hg38) Alu, LINE-1 (L1), and Long Terminal Repeat (LTR) (including endogenous retroviruses, ERV) repetitive element (RE) methylation prediction using Illumina 450k/EPIC array or sequencing platform.

3.1 Groom methylation data

Appropriate data preprocessing including quality control and normalization of methylation data are recommended before running *REMP*. Many packages are available to carry out these data preprocessing steps, for example, *minfi*, *wateRmelon*, and *methylumi*.

REMP is trying to minimize the requirement of the methylation data format. Users can maintain the methylation data in *RatioSet* or *GenomicRatioSet* object offered by *minfi*, *data.table*, *data.frame*, *DataFrame*, or *matrix*. Users can input either beta value or M-value. There are only two basic requirements of the methylation array data (450k/EPIC):

- 1. Each row should represent CpG probe and each column should represent sample.
- 2. The row names should indicate Illumina probe ID (i.e. cg00000029).

However, there are some other common data issues that may prevent *REMP* from running correctly. For example, if the methylation data are in beta value and contain zero methylation values, logit transformation (to create M-value) will create negative infinite value; or the methylation data contain NA, Inf, or NaN data. To tackle these potential issues, *REMP* includes a handy function grooMethy which can help detect and fix these issues. We highly recommend to take advantage of this function:

```
> # Get GM12878 methylation data (450k array)
> GM12878_450k <- getGM12878('450k')
> GM12878 450k <- grooMethy(GM12878 450k)
> GM12878 450k
class: RatioSet
dim: 482421 1
metadata(0):
assays(2): Beta M
rownames(482421): cq00000029 cq00000108 ... cq27666046
  cq27666123
rowData names(0):
colnames(1): GM12878
colData names(0):
Annotation
  array: IlluminaHumanMethylation450k
  annotation: ilmn12.hg19
Preprocessing
 Method: NA
 minfi version: NA
 Manifest version: NA
```

For zero beta values, grooMethy will replace them with smallest non-zero beta value. For one beta values, grooMethy will replace them with largest non-one beta value. For NA/NaN/Inf values, grooMethy will treat them as missing values and then apply KNN-imputation to complete the dataset. 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.

For sequencing data, the users only need to prepare a methylation data matrix (row = CpGs, column = samples). The corresponding CpG location information (either in hg19 or hg38) should be prepared in a separate *GRanges* object and provide it to the Seq.GR argument in grooMethy. For an example of Seq.GR, please run:

```
> library(IlluminaHumanMethylation450kanno.ilmn12.hg19)
> getLocations(IlluminaHumanMethylation450kanno.ilmn12.hg19)
```

```
GRanges object with 485512 ranges and 0 metadata columns:
                             ranges strand
                  seqnames
                    <Rle> <IRanges> <Rle>
                     chrY 9363356
       cg00050873
                                          *
       cq00212031
                     chrY 21239348
                                          *
       cg00213748
                     chrY 8148233
                                          *
       cg00214611
                    chrY 15815688
                                          *
       cq00455876
                           9385539
                     chrY
                                          *
              . . .
                      . . .
                                 . . .
                                        . . .
    ch.22.909671F chr22 46114168
                                        *
  ch.22.46830341F
                    chr22 48451677
                                          *
   ch.22.1008279F
                    chr22 48731367
                                          *
  ch.22.47579720R
                    chr22 49193714
                                          *
  ch.22.48274842R
                    chr22 49888838
                                          *
  seqinfo: 24 sequences from hg19 genome; no seqlengths
```

Note that the row names of the CpGs in Seq. GR can be NULL.

3.2 Prepare annotation data

To run *REMP* for RE methylation prediction, users first need to prepare some annotation datasets. The function initREMP is designed to do the job.

Suppose users will predict Alu methylation using Illumina 450k array data:

```
> data(Alu.hg19.demo)
> remparcel <- initREMP(arrayType = "450k",</pre>
                        REtype = "Alu",
+
                        annotation.source = "AH",
+
                        genome = "hg19",
+
+
                        RE = Alu.hg19.demo,
                        ncore = 1)
+
> remparcel
REMParcel object
RE type: Alu
Genome build: hg19
Illumina platform: 450k
Valid (max) Alu-CpG flanking window size: 1200
Number of RE: 500
Number of Alu-CpG: 5138
```

For demonstration, we only use 500 selected Alu sequence dataset which comes along with the package (Alu.hg19.demo). We specify RE = Alu.hg19.demo, so that the annotation dataset will be generated for the 500 selected Alu sequences. Most of the time, specifying RE is not necessary, as the function will fetch the complete RE sequence dataset from package *AnnotationHub* using fetchRMSK. Users can also use this argument RE to provide customized RE dataset.

annotation.source allows the users to switch the source of the annotation databases, including the RefSeq Gene annotation database and RepeatMasker annotation database. If annotation.source = "AH", the database will be obtained from the AnnotationHub package. If annotation.source = "UCSC", the database will be downloaded from the UCSC website http://hgdownload.cse.ucsc.edu/goldenpath. The corresponding build ("hg19" or "hg38") can be specified in the argument genome. Most of the time "hg19" is used for array data. But if "hg38" is specified, the function will liftover the CpG probe location information to "hg38" and obtain annotation databases in "hg38".

If arrayType = "Sequencing", users should provide the genomic location information of the CpGs in a *GRanges* object to Seq.GR. Note that the genome build of Seq.GR provided should match the genome build specified in genome.

All data are stored in the *REMParcel* object:

```
> saveParcel(remparcel)
```

It is recommended to specify a working directory using argument work.dir in initREMP so that the annotation data generated can be re-used. Without specifying working directory, the annotation dataset will be created under the temporal directory tempdir() by default. Users can also turn on the export argument in initREMP to save the data automatically.

3.3 Run prediction

Once the annotation data are ready, users can pass the annotation data parcel to remp for prediction:

+	<pre>parcel = remparcel,</pre>
+	ncore = 1,
+	seed = 777)

If parcel is missing, remp will then try to search the *REMParcel* data file in the directory indicated by work.dir. If work.dir is also missing, remp will try to search the REMParcel data file in the temporal directory tempdir().

By default, remp uses Random Forest (method = 'rf') model (package *ranger* for fast implementation) for prediction. Random Forest model is recommended because it offers more accurate prediction results and it automatically enables Quantile Regression Forest (Nicolai Meinshausen, 2006) for prediction reliability evaluation. remp constructs predictors to carry out the prediction. For Random Forest model, the tuning parameter param = 6 (i.e. mtry in *ranger* or *randomForest*) indicates how many predictors will be randomly selected for building the individual trees. 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. It is recommended to specify a seed for reproducible prediction results.

Besides random forest, remp provides other machine learning engines for users to explore, including Extreme Gradient Boosting, SVM with linear kernel, and SVM with radial kernel).

remp will return a *REMPset* object, which inherits Bioconductor's *RangedSummarizedExperiment* class:

```
> remp.res
```

```
class: REMProduct
dim: 4952 1
metadata(8): REannotation RECpG ... GeneStats Seed
assays(3): rempB rempM rempQC
rownames: NULL
rowData names(1): RE.Index
colnames(1): GM12878
colData names(1): mtry
> # Display more detailed information
> details(remp.res)
RE type: Alu
Genome build: hg19
Methylation profiling platform: 450k
Flanking window size: 1000
Prediction model: Random Forest
QC model: Quantile Regression Forest
Seed: 777
Covered 4952 CpG sites in 500 Alu
Number of Alu-CpGs by chromosome:
chr1 chr2 chr3 chr4 chr5 chr6 chr7 chr8
 429 195 257 197 142 382 202 153
 chr9 chr10 chr11 chr12 chr13 chr14 chr15 chr16
             192
                  355
   93
       172
                           52
                                122
                                      201
                                            280
chr17 chr18 chr19 chr20 chr21 chr22
  361
        65
            803
                  107
                           56
                                136
Training information:
  500 profiled Alu are used for model training.
```

490 Alu-CpGs that have at least 2 neighboring profiled CpGs are used for model training.

Coverage information: The data cover 500 Alu (4952 Alu-CpG). Gene coverage by Alu (out of total # of RefSeq genes): 530 (2.13%) total genes; 460 (2.4%) protein-coding genes; 103 (1.43%) non-coding RNA genes. Distribution of methylation value (beta value): Median 3rd Qu. Min. lst Ou. Mean Max. 0.01178818 0.43739915 0.64585305 0.57035834 0.75523415 0.90391551 Distribution of reliability score (lower score = higher reliability): Min. 1st Qu. Median Mean 3rd Qu. Max. 0.6405955 1.3005384 1.5831950 1.6681776 2.0136318 5.9084103 Prediction results can be obtained by accessors: > # Predicted RE-CpG methylation value (Beta value) > rempB(remp.res) DataFrame with 4952 rows and 1 column GM12878 <numeric> 1 0.658459 2 0.655791 3 0.658361 4 0.665226 5 0.664966 4948 0.809959 4949 0.810494 4950 0.811938 4951 0.811706 4952 0.825812 > # Predicted RE-CpG methylation value (M value) > rempM(remp.res) DataFrame with 4952 rows and 1 column GM12878 <numeric> 0.947032 1 2 0.929951 3 0.946409 4 0.990658 5 0.988975 4948 2.09154 4949 2.09656 4950 2.11016 4951 2.10797

4952 2.24516

> # Genomic location information of the predicted RE-CpG

> # Function inherit from class 'RangedSummarizedExperiment'

> rowRanges(remp.res)

GRanges object wi	th 4952 ranges and	1 metadata column:
seqnames	ranges	strand RE.Index
<rle></rle>	<iranges></iranges>	<rle> <rle></rle></rle>
[1] chr1	1149956-1149957	+ Alu_0000214
[2] chr1	1149966-1149967	+ Alu_0000214
[3] chr1	1149999-1150000	+ Alu_0000214
[4] chr1	1150049-1150050	+ Alu_0000214
[5] chr1	1150057-1150058	+ Alu_0000214
[4948] chr22	50914839-50914840	- Alu_1118131
[4949] chr22	50914853-50914854	- Alu_1118131
[4950] chr22	50914860-50914861	- Alu_1118131
[4951] chr22	50914869-50914870	- Alu_1118131
[4952] chr22	50914907-50914908	- Alu_1118131

seqinfo: 24 sequences from an unspecified genome; no seqlengths

> # Standard error-scaled permutation importance of predictors
> rempImp(remp.res)

DataFrame with 18 rows and 1 column GM12878 <numeric> RE.swScore 7.48388 RE.Length 3.42704 RE.CpG.density 6.81685 RE.InTSS 2.11638 RE.In5UTR 1.78648 Methy.mean.mov1 20.84179 Methy.mean.mov2 14.77652 Methy.mean.mov3 9.14708 Methy.mean.mov4 10.58181 Methy.std 7.29195

> # Retrive seed number used for the reesults
> metadata(remp.res)\$Seed

[1] 777

Trim off less reliable predicted results:

> # Any predicted CpG values with quality score less than > # threshold (default = 1.7) will be replaced with NA. > # CpGs contain more than missingRate * 100% (default = 20%) > # missing rate across samples will be discarded. > remp.res <- rempTrim(remp.res, threshold = 1.7, missingRate = 0.2) > details(remp.res) RE type: Alu Genome build: hg19 Methylation profiling platform: 450k Flanking window size: 1000 Prediction model: Random Forest - trimmed (1.7) QC model: Quantile Regression Forest Seed: 777 Covered 2848 CpG sites in 397 Alu Number of Alu-CpGs by chromosome: chr1 chr2 chr3 chr4 chr5 chr6 chr7 chr8 295 145 155 138 88 134 102 93 chr9 chr10 chr11 chr12 chr13 chr14 chr15 chr16 117 223 31 63 107 168 39 89 chr17 chr18 chr19 chr20 chr21 chr22 199 40 459 73 13 77 Coverage information: The data cover 397 Alu (2848 Alu-CpG). Gene coverage by Alu (out of total # of RefSeq genes): 415 (1.67%) total genes; 356 (1.86%) protein-coding genes; 85 (1.18%) non-coding RNA genes. Distribution of methylation value (beta value): Min. 1st Qu. Median Mean 3rd Qu. Max. 0.03459213 0.62096126 0.72580862 0.66668785 0.79453724 0.90391551 Distribution of reliability score (lower score = higher reliability): 1st Qu. Median Mean 3rd Ou. Min. Max. 0.6405955 1.1777745 1.3414789 1.3299838 1.4916722 1.6996203

(Optional) Aggregate the predicted methylation of CpGs in RE by averaging them to obtain the RE-specific methylation level:

> remp.res <- rempAggregate(remp.res, NCpG = 2)</pre> > details(remp.res) RE type: Alu (aggregated by mean: min # of CpGs: 2) Genome build: hg19 Methylation profiling platform: 450k Flanking window size: 1000 Prediction model: Random Forest - trimmed (1.7) QC model: Quantile Regression Forest Seed: 777 Covered 339 Alu (aggregated by mean: min # of CpGs: 2) Number of Alu (aggregated by mean: min # of CpGs: 2) by chromosome: chr1 chr2 chr3 chr4 chr5 chr6 chr7 chr8 32 17 18 14 9 21 14 10 chr9 chr10 chr11 chr12 chr13 chr14 chr15 chr16 7 13 14 25 5 6 13 19

```
chr17 chr18 chr19 chr20 chr21 chr22
             24
                                         4
                                                              55
                                                                                           8
                                                                                                                    2
                                                                                                                                              9
Coverage information:
        The data cover 339 Alu (aggregated by mean: min # of CpGs: 2)
        Gene coverage by Alu (aggregated by mean: min # of CpGs: 2) (out of total # of RefSeq generation of the coverage by Alu (aggregated by mean: min # of CpGs: 2) (out of total # of RefSeq generation of the coverage by Alu (aggregated by mean: min # of CpGs: 2) (out of total # of RefSeq generation of the coverage by Alu (aggregated by mean: min # of CpGs: 2) (out of total # of RefSeq generation of the coverage by Alu (aggregated by mean: min # of CpGs: 2) (out of total # of RefSeq generation of the coverage by Alu (aggregated by mean: min # of CpGs: 2) (out of total # of RefSeq generation of the coverage by Alu (aggregated by mean: min # of CpGs: 2) (out of total # of RefSeq generation of the coverage by Alu (aggregated by mean: min # of CpGs: 2) (out of total # of RefSeq generation of the coverage by Alu (aggregated by mean: min # of CpGs: 2) (out of total # of RefSeq generation of the coverage by Alu (aggregated by mean: min # of CpGs: 2) (out of total # of RefSeq generation of the coverage by Alu (aggregated by mean: min # of CpGs: 2) (out of total # of RefSeq generation of the coverage by Alu (aggregated by mean: min # of CpGs: 2) (out of total # of RefSeq generation of the coverage by Alu (aggregated by mean: min # of CpGs: 2) (out of total # of RefSeq generation of the coverage by Alu (aggregated by mean: min # of CpGs: 2) (out of total # of RefSeq generation of the coverage by Alu (aggregated by mean: min # of CpGs: 2) (out of total # of RefSeq generation of RefSeq generation of total # of RefSeq generation of RefSeq 
                352 (1.41%) total genes;
                298 (1.56%) protein-coding genes;
                74 (1.03%) non-coding RNA genes.
Distribution of methylation value (beta value):
                        Min.
                                                         1st Qu.
                                                                                                          Median
                                                                                                                                                                 Mean
                                                                                                                                                                                                    3rd Qu.
                                                                                                                                                                                                                                                              Max.
0.04786608 0.59569784 0.70904833 0.64428356 0.78028163 0.85384885
Distribution of reliability score (lower score = higher reliability):
                                                lst Qu.
                                                                                              Median
                                                                                                                                                Mean
                                                                                                                                                                               3rd Qu.
                    Min.
                                                                                                                                                                                                                                    Max.
0.8250507 1.2487521 1.3836146 1.3658707 1.4913583 1.6902399
```

Aggregating CpGs in the same RE for RE-level methylation data is beneficial because 1) it greatly reduces the data dimension for downstream analysis and 2) it may produce more robust RE methylation estimation. Note that by default, RE with 2 or more predicted CpG sites will be aggregated. Therefore, the downside of doing this is the reduced coverage of RE. The assumption of doing this is the CpG methylation level within each RE are similar.

To add genomic regions annotation of the predicted REs:

```
> # By default gene symbol annotation will be added
> remp.res <- decodeAnnot(remp.res)
> rempAnnot(remp.res)
```

GRanges object with 339 ranges and 12 metadata columns:

		5				-	
seqnames		range	es strand		swScore	repName	
<rle></rle>		<iranges< td=""><td>s> <rle></rle></td><td> <</td><td><integer></integer></td><td><character></character></td><td></td></iranges<>	s> <rle></rle>	<	<integer></integer>	<character></character>	
chr1	114994	7-115024	12 +	1	2216	AluSx1	
chr1	188594	1-188623	30 +	1	1962	AluJb	
chr1	14032197	-1403250)0 +		2768	AluSg	
chr1	22378622	-2237892	26 +		2142	AluSx1	
chrl	39408080	-3940838	32 +		2493	AluSp	
		• •	• •••	•			
chr22	44964234	-4496453	39 +		2284	AluSq2	
chr22	50311037	-5031133	33 +	1	2251	AluSx1	
chr22	50493205	-5049351	-5 +	1	2229	AluSx	
chr22	42078691	-4207895	52 -		1881	AluSx1	
chr22	50914608	-5091491	.7 –		2526	AluY	
repCla	ass rep	Family	Inde	∋x	InNM.symb	ol InNR.symb	ool
<characte< td=""><td>er> <char< td=""><td>acter></td><td><rle< td=""><td>∋></td><td><characte< td=""><td>r> <characte< td=""><td>er></td></characte<></td></characte<></td></rle<></td></char<></td></characte<>	er> <char< td=""><td>acter></td><td><rle< td=""><td>∋></td><td><characte< td=""><td>r> <characte< td=""><td>er></td></characte<></td></characte<></td></rle<></td></char<>	acter>	<rle< td=""><td>∋></td><td><characte< td=""><td>r> <characte< td=""><td>er></td></characte<></td></characte<></td></rle<>	∋>	<characte< td=""><td>r> <characte< td=""><td>er></td></characte<></td></characte<>	r> <characte< td=""><td>er></td></characte<>	er>
SI	INE	Alu A	Alu_000021	L4	TNFRS	F4 <n< td=""><td>VA></td></n<>	VA>
SI	INE	Alu A	Alu_000063	34	CFAP	74 <n< td=""><td>VA></td></n<>	VA>
SI	INE	Alu A	Alu_000456	65	PRD	M2 <n< td=""><td>VA></td></n<>	VA>
SI	INE	Alu A	Alu_000753	35	CDC	42 <1	VA>
SI	INE	Alu A	Alu_001417	70	RHBD	L2 <n< td=""><td>VA></td></n<>	VA>
SI	INE	Alu A	Alu_110453	34	<n.< td=""><td>A> LINC002</td><td>207</td></n.<>	A> LINC002	207
SI	INE	Alu A	lu_110570) 3 (CRELD2 ALG	12 CREI	LD2
S	INE	Alu A	lu_110573	37	<n.< td=""><td>A> <n< td=""><td>VA></td></n<></td></n.<>	A> <n< td=""><td>VA></td></n<>	VA>
SI	INE					13 <1	VA>
	<pre><rle> chr1 chr1 chr1 chr1 chr1 chr1 chr22 chr2 chr</rle></pre>	chr1 114994 chr1 188594 chr1 14032197 chr1 22378622 chr1 39408080 chr22 44964234 chr22 50311037 chr22 50493205 chr22 42078691 chr22 50914608 repClass rep	<pre><rle> <iranges 1149947-115024="" 14032197-1403250="" 1885941-188623="" 22378622-2237892="" 39408080-3940838="" 42078691-4207895="" 44964234-4496453="" 50311037-5031133="" 50493205-5049351="" 50914608-5091491="" <character="" chr1="" chr22="" repclass="" repfamily=""> SINE Alu # SINE</iranges></rle></pre>	<pre><rle> <iranges> <rle> chr1 1149947-1150242 + chr1 1885941-1886230 + chr1 14032197-14032500 + chr1 22378622-22378926 + chr1 39408080-39408382 + chr22 44964234-44964539 + chr22 50311037-50311333 + chr22 50493205-50493515 + chr22 42078691-42078952 - chr22 50914608-50914917 - repClass repFamily Inde <character> <character> <rle alu="" alu_000062="" alu_000456="" alu_00066="" alu_001417="" alu_110455="" alu_110576="" alu_110576<="" pre="" sine=""></rle></character></character></rle></iranges></rle></pre>	<pre><rle> <iranges> <rle> - chr1 1149947-1150242 + chr1 1885941-1886230 + chr1 14032197-14032500 + chr1 22378622-22378926 + chr1 39408080-39408382 + chr1 39408080-39408382 + chr22 44964234-44964539 + chr22 50311037-50311333 + chr22 50493205-50493515 + chr22 50914608-50914917 - repClass repFamily Index <character> <character> <rle> SINE Alu Alu_0000214 SINE Alu Alu_0004565 SINE Alu Alu_0014170 SINE Alu Alu_1104534 SINE Alu Alu_1105703 SINE Alu Alu_1105703</rle></character></character></rle></iranges></rle></pre>	<rle> <iranges> <rle> <integer> chr1 1149947-1150242 + 2216 chr1 1885941-1886230 + 1962 chr1 14032197-14032500 + 2768 chr1 22378622-22378926 + 2142 chr1 239408080-39408382 + 2493 chr22 44964234-44964539 + 2284 chr22 50311037-50311333 + 2251 chr22 50493205-50493515 + 2229 chr22 50914608-50914917 - 2526 repClass repFamily Index InNM.symb <character> <character> <rle> <characte< td=""> SINE Alu Alu_00000214 TNFRS SINE Alu Alu_0000634 CFAP SINE Alu Alu_00007535 CDC SINE Alu Alu_00014170 RHBD SINE Alu Alu_1104534 <n.< <="" td=""><td><rle> <iranges> <rle> <integer> <character> chr1 1149947-1150242 + 2216 AluSx1 chr1 1885941-1886230 + 1962 AluJb chr1 14032197-14032500 + 2768 AluSg chr1 22378622-22378926 + 2142 AluSx1 chr1 39408080-39408382 + 2493 AluSp chr22 44964234-44964539 + 2284 AluSq2 chr22 50311037-50311333 + 2251 AluSx1 chr22 50493205-50493515 + 2229 AluSx1 chr22 50914608-50914917 - 2526 AluY repClass repFamily Index InNM.symbol InNR.symbol chr22 50914608-50914917 - 2526 AluY repClass repFamily Index InNM.symbol InNR.symbol sINE Alu Alu_0000214 TNFRSF4 <n< td=""> SINE A</n<></character></integer></rle></iranges></rle></td></n.<></characte<></rle></character></character></integer></rle></iranges></rle>	<rle> <iranges> <rle> <integer> <character> chr1 1149947-1150242 + 2216 AluSx1 chr1 1885941-1886230 + 1962 AluJb chr1 14032197-14032500 + 2768 AluSg chr1 22378622-22378926 + 2142 AluSx1 chr1 39408080-39408382 + 2493 AluSp chr22 44964234-44964539 + 2284 AluSq2 chr22 50311037-50311333 + 2251 AluSx1 chr22 50493205-50493515 + 2229 AluSx1 chr22 50914608-50914917 - 2526 AluY repClass repFamily Index InNM.symbol InNR.symbol chr22 50914608-50914917 - 2526 AluY repClass repFamily Index InNM.symbol InNR.symbol sINE Alu Alu_0000214 TNFRSF4 <n< td=""> SINE A</n<></character></integer></rle></iranges></rle>

[339]	SINE	Alu Alı	Alu Alu_1118131		<na></na>
	InTSS.symbol	In5UTR.symbol	InCDS.symbol	InExon.symbol	
	<character></character>	<character></character>	<character></character>	<character></character>	
[1]	TNFRSF4	<na></na>	<na></na>	<na></na>	
[2]	<na></na>	<na></na>	<na></na>	CFAP74	
[3]	<na></na>	PRDM2	<na></na>	<na></na>	
[4]	CDC42	<na></na>	<na></na>	<na></na>	
[5]	RHBDL2	<na></na>	<na></na>	<na></na>	
• • •					
[335]	LINC00207	<na></na>	<na></na>	<na></na>	
[336]	CRELD2	ALG12	<na></na>	<na></na>	
[337]	<na></na>	<na></na>	<na></na>	<na></na>	
[338]	<na></na>	SNU13	SNU13	<na></na>	
[339]	SBF1	<na></na>	<na></na>	<na></na>	
	In3UTR.symbol	L			
	<character></character>	>			
[1]	<na></na>	>			
[2]	CFAP74	1			
[3]	<na></na>	>			
[4]	<na></na>	>			
[5]	<na></na>	>			
• • •	• • •				
[335]	<na></na>	>			
[336]	<na></na>	>			
[337]	<na></na>	>			
[338]	<na></na>	>			
[339]	<na></na>	>			
		C 1 1 0			

seqinfo: 24 sequences from hg19 genome

Seven genomic region indicators will be added to the annotation data in the input REMProduct object:

- InNM: in protein-coding genes (overlap with refSeq gene's "NM" transcripts + 2000 bp upstream of the transcription start site (TSS))
- InNR: in noncoding RNA genes (overlap with refSeq gene's "NR" transcripts + 2000 bp upstream of the TSS)
- InTSS: in flanking region of 2000 bp upstream of the TSS. Default upstream limit is 2000 bp, which can be modified globally using remp_options
- In5UTR: in 5'untranslated regions (UTRs)
- InCDS: in coding DNA sequence regions
- InExon: in exon regions
- In3UTR: in 3'UTRs

Note that intron region and intergenic region information can be derived from the above genomic region indicators: if "InNM" and/or "InNR" is not missing but "InTSS", "In5UTR", "InExon", and "In3UTR" are missing, then the RE is strictly located within intron region; if all indicators are missing, then the RE is strictly located in intergenic region.

3.4 Plot prediction

Make a density plot of the predicted methylation (beta values):

> remplot(remp.res, main = "Alu methylation (GM12878)", col = "blue")

4 Extract RE-CpG methylation profiled by Illumina BeadChip array

REMP offers a handy tool to extract methylation data of CpGs that are located in RE. Similar as remp, users can choose the source of annotation database (AH: AnnotationHub or UCSC: UCSC website) and genome build (hg19 or hg38).

```
> # Use Alu.hg19.demo for demonstration
> remp.res <- remprofile(GM12878_450k,</pre>
                         REtype = "Alu",
+
                         annotation.source = "AH",
+
+
                         genome = "hg19",
                         RE = Alu.hg19.demo)
+
> details(remp.res)
RE type: Alu
Genome build: hg19
Methylation profiling platform: 450k
Flanking window size: N/A
Prediction model: Profiled
OC model: N/A
Covered 602 CpG sites in 500 Alu
Number of Alu-CpGs by chromosome:
chr1 chr2 chr3 chr4 chr5 chr6 chr7 chr8
  58
      22
          33
               23
                    17
                           59
                                26
                                     18
 chr9 chr10 chr11 chr12 chr13 chr14 chr15 chr16
   12
      27 20
                    42
                            8
                                11
                                       20
                                             30
chr17 chr18 chr19 chr20 chr21 chr22
   39
          9
               97
                     13
                           5
                                 13
Coverage information:
  The data cover 500 Alu (602 Alu-CpG).
  Gene coverage by Alu (out of total # of RefSeq genes):
    530 (2.13%) total genes;
    460 (2.4%) protein-coding genes;
    103 (1.43%) non-coding RNA genes.
Distribution of methylation value (beta value):
    Min. 1st Qu.
                  Median
                              Mean 3rd Qu.
                                                 Max.
0.001000 0.344500 0.649500 0.567397 0.814000 0.959000
> # All accessors and utilites for REMProduct are applicable
> remp.res <- rempAggregate(remp.res)</pre>
> details(remp.res)
RE type: Alu (aggregated by mean: min # of CpGs: 2)
Genome build: hq19
Methylation profiling platform: 450k
Flanking window size: N/A
```

Prediction model: Profiled QC model: N/A Covered 86 Alu (aggregated by mean: min # of CpGs: 2) Number of Alu (aggregated by mean: min # of CpGs: 2) by chromosome: chr1 chr2 chr3 chr4 chr5 chr6 chr7 chr8 13 2 6 2 1 14 3 2 chr10 chr11 chr12 chr14 chr15 chr16 chr17 chr18 2 3 9 1 2 4 4 3 chr19 chr20 chr21 chr22 9 3 1 2 Coverage information: The data cover 86 Alu (aggregated by mean: min # of CpGs: 2) Gene coverage by Alu (aggregated by mean: min # of CpGs: 2) (out of total # of RefSeq gen 98 (0.39%) total genes; 83 (0.43%) protein-coding genes; 24 (0.33%) non-coding RNA genes. Distribution of methylation value (beta value): Min. 1st Qu. Median Mean 3rd Qu. Max.

0.0408311 0.1983081 0.6649476 0.5265950 0.7829270 0.9288642