methylSig: A package for whole genome DNA methylation analysis

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1 Introduction

DNA methylation plays critical roles in gene regulation and cellular specification without altering DNA sequences. It is one of the best understood and most intensively studied epigenetic marks in mammalian cells. Treatment of DNA with sodium bisulfite deaminates unmethylated cytosines to uracil while methylated cytosines are resistant to this conversion thus allowing for the discrimination between methylated and unmethylated CpG sites. Sodium bisulfite pre-treatment of DNA coupled with next-generation sequencing has allowed DNA methylation to be studied quantitatively and genome-wide at single cytosine site resolution.

MethylSig is a method for testing for differential methylated cytosines (DMCs) or regions (DMRs) in whole-genome bisulfite sequencing (bis-seq) or reduced representation bisulfite sequencing (RRBS) experiments. MethylSig uses a beta binomial model to test for significant differences between groups of samples. Several options exist for either site-specific or sliding window tests, combining strands, and for variance estimation. It allows annotating the resulting regions to multiple genome features, and visualizing the results for chosen genomic regions.

This document provides a step by step guide for the methylSig package.

2 Installation

MethylSig is available as an R package at http://sartorlab.ccmb.med.umich.edu/software. After downloading the methylSig package, for example, methylSig_0.1.0.tar.gz, the install.packages() function can be used to install methylSig in R. The current version of the methylSig package depends on R (>=2.10).

```
>install.packages("methylSig_0.1.0.tar.gz", repos=NULL, type="source")
* installing *source* package 'methylSig' ...
** R
** data
** inst
** preparing package for lazy loading
** help
*** installing help indices
** building package indices
** testing if installed package can be loaded
* DONE (methylSig)
```

3 Basic

3.1 Loading methylSig package

methylSig package can be loaded using the library() function.

```
>library(methylSig)
Loading required package: boot
Loading required package: parallel
```

This will load methylSig package as well as other dependent packages.

3.2 Methylation score files

CpG methylation score files can be obtained using programs such as bismark. The typical CpG methylation score files are as follows:

```
chrBase
                chr
                        base strand coverage freqC freqT
chr21.43008527 chr21 43008527
                                  F
                                          32 100.00 0.00
chr21.43008531 chr21 43008531
                                  F
                                          32 96.88 3.12
                                          32 90.62 9.38
chr21.43008543 chr21 43008543
                                  F
                                  R
                                          27 100.00 0.00
chr21.43008674 chr21 43008674
chr21.43008710 chr21 43008710
                                  R
                                             94.03 5.97
chr21.43008720 chr21 43008720
                                  R
                                          67
                                             92.54 7.46
```

The CpG methylation score file must contain at least seven columns. Among these, second to seventh column must be, in order, chromosome, base, strand, coverage, percentage of Cs and percentage of Ts. Column names are not important. Strand format is F/R or +/-, where F/+ represents forward and R/- represents reverse strand.

3.3 Reading methylation score files

MethylSig package provides the methylSigReadData() function to read CpG methylation score files and convert these files into a 'methylSigData' object for further analysis and annotation.

```
> fileList = c(system.file("extdata", "AML_1.txt", package = "methylSig"),
                system.file("extdata", "AML_2.txt", package = "methylSig"),
                system.file("extdata", "AML_3.txt", package = "methylSig"),
system.file("extdata", "AML_4.txt", package = "methylSig"),
system.file("extdata", "NBM_1.txt", package = "methylSig"),
                system.file("extdata", "NBM_2.txt", package = "methylSig"),
                system.file("extdata", "NBM_3.txt", package = "methylSig"),
                system.file("extdata", "NBM_4.txt", package = "methylSig"))
> sample.id = c("AML1", "AML2", "AML3", "AML4", "NBM1", "NBM2", "NBM3", "NBM4")
> treatment = c(1,1,1,1,0,0,0,0)
> #### Read Data ####
> meth <- methylSigReadData(fileList, sample.ids = sample.id, assembly = "hg18",
              treatment = treatment, context = "CpG", destranded=TRUE)
Reading file (1/8) -- /tmp/RtmppYuoUi/Rinst39144d1f5be7/methylSig/extdata/AML_1.txt
Reading file (2/8) -- /tmp/RtmppYuoUi/Rinst39144d1f5be7/methylSig/extdata/AML_2.txt
Reading file (3/8) -- /tmp/RtmppYuoUi/Rinst39144d1f5be7/methylSig/extdata/AML_3.txt
Reading file (4/8) -- /tmp/RtmppYuoUi/Rinst39144d1f5be7/methylSig/extdata/AML_4.txt
Reading file (5/8) -- /tmp/RtmppYuoUi/Rinst39144d1f5be7/methylSig/extdata/NBM_1.txt
Reading file (6/8) -- /tmp/RtmppYuoUi/Rinst39144d1f5be7/methylSig/extdata/NBM_2.txt
Reading file (7/8) -- /tmp/RtmppYuoUi/Rinst39144d1f5be7/methylSig/extdata/NBM_3.txt
Reading file (8/8) -- /tmp/RtmppYuoUi/Rinst39144d1f5be7/methylSig/extdata/NBM_4.txt
(1)(2)(3)(4)(5)(6)(7)(8)
```

It is possible for the user to filter out CpG sites based on the read coverage. CpG sites with very lrage read coverage may be due to PCR bias and hence including CpG sites with very high coverage may distort the statistics of data analysis. The methylSigReadData() function provides 'minCount' and 'maxCount' arguments for defining lower and upper limits for coverage. The default values are 10 and 500 respectively.

Here, the argument quiet controls the progress reports. If quiet=TRUE, no progress reports will be shown.

There are many arguments for the methylSigReadData() funciton. Among these 'fileList', 'sample.ids' and 'treatment' are required. Some options have default values, for example, 'destranded=TRUE', 'num.cores=1', and 'quiet=FALSE'. Other arguments such as 'assembly', 'context' and 'pipeline' are optional and for information purposes only. The data type of treatment is a numeric vector. Each number represents a group. Multiple groups can be stored in one methylSigData object.

The arguemnt 'num.cores' is used for multi-thread reading. See section 5 for more details.

4 Differential methylation analysis

4.1 Usage of the methylSigCalc function

The main function of this package is the differential methylation analysis function methySigCalc(). It calculates differential methylation statistics between two groups of samples. It uses a beta-binomial approach to calculate differential methylation statistics, accounting for coverage and variantion among samples within each group.

Usage:

4.2 Site specific analysis

The default is to do site specific analysis and to use both groups to estimate variances.

```
> myDiffSigboth = methylSigCalc(meth, groups=c(1,0), min.per.group=3)
```

Total number of bases: 3.26k

The differentially methylated cytosines (DMCs) can be defined based on qualues, pvalues and the methylation rate difference between two tested groups.

4.3 Tiled Data analysis

MethylSig package also provides methylSigTile() function to tile data within continuous non-overlapping windows. The default window size is 25bp. After tiling data, the methylSigCalc() function can be used to calculate differential methylation statistics.

```
> ### Tiled analysis
> methTile = methylSigTile(meth,win.size = 25)
> myDiffSigbothTile = methylSigCalc(methTile, groups=c(1,0), min.per.group=3)
Total number of regions: 994
```

4.4 Variance from one group

Using the **dispersion** argument, it is possible to estimate variances from one group rather than from both groups. The following code calculates differential methylation statistics based on estimating variances from group 0 only.

Total number of regions: 994

4.5 Using local information

It is also possible to use information from nearby CpG sites to improve the variance and methylation level estimates. The default winsize.disp and winsize.meth are 200 bps. The winsize.disp argument only takes into effect when local.disp is set to 'TRUE'. Similarly winsize.meth argument only takes into effect when local.meth is set to 'TRUE'.

Total number of bases: 3.26k

5 Multiple threads computation

MethylSig package provides multi-thread programming to substantially reduce data analysis time. In the functions methylSigDataRead() and methylSigCalc(), multi-core programming will be initiated using 'num.cores' argument. Note that this option depends on R package 'parallel' and hence is not available in the Windows platform. The following example code is using 5 cores.

Total number of bases: 3.26k

6 Annotation

6.1 CpG island Annotation

There are two functions, cpgAnnotation() and cpgAnnotationPlot(), in the methylSig package for CpG island annotation. The CpG island information file can be download from websites such as the UCSC genome browser. The appropriate genome assembly (the same genome assembly of the provided data) should be used.

In a linux server, the user may use the following command to download the annotation file for hg19. Please use appropriate directories for hg18, mm9 or mm10.

```
wget ftp://hgdownload.cse.ucsc.edu/goldenPath/hg19/database/cpgIslandExt.txt.gz
gunzip *.gz
```

Here we use the CpG island annotation file provided in the methylSig package to annotate our example. Note that this is a reduced annotation file and is not appropriate for a full real data analysis.

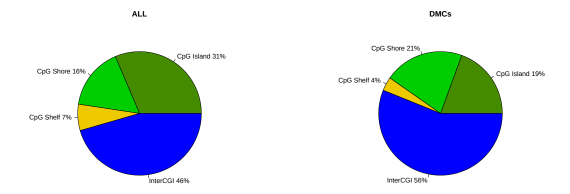


Figure 1: CpG annotation plot

6.2 RefGene Annotation

Again, there are two functions, refGeneAnnotation() and refGeneAnnotationPlot(), in methylSig package for annotation using RefGene models. The refGene information file can be download from websites such UCSC genome browser. The appropriate genome assembly (the same genome assembly of the provided data) should be used.

In a linux server, the user may use the following command to download the annotation file for hg19. Please use appropriate directories for hg18, mm9 or mm10.

```
wget ftp://hgdownload.cse.ucsc.edu/goldenPath/hg19/database/refGene.txt.gz
gunzip *.gz
```

We use refGene annotation file provided in the methylSig package to annotate in our example. Note that this is a reduced annotation file and is not appropriate for the a full real data analysis.

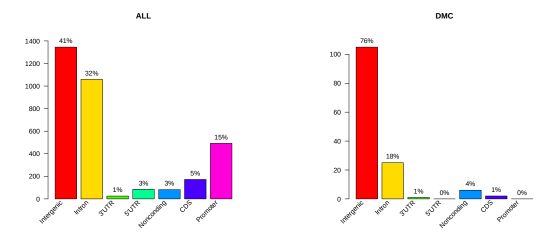


Figure 2: RefGene annotation plot

6.3 Transcription factor enrichment test

The functions getTFBSInfo() and methylSig.tfbsEnrichTest() are provided for reading the TFBS information file and implementing transcription factor enrichment test.

UCSC genome browser provides TFBS conserved track for hg18 and hg19. The following linux server shell command can be used to download these files:

```
wget ftp://hgdownload.cse.ucsc.edu/goldenPath/hg19/database/tfbsConsSites.txt.gz
wget ftp://hgdownload.cse.ucsc.edu/goldenPath/hg19/database/tfbsConsFactors.txt.gz
gunzip *.gz
```

Here, 'tfbsConsSites.txt' is tracking information and can be used directory in function getTFBSInfo(). The explanation of variable names is listed in file 'tfbsConsFactors.txt'.

Another TFBS track is from ENCODE for hg18, hg19 and mm9. However, the methylSig package cannot use this type of track directly. We provide ENCODE TFBS track files that are suitable for methylSig package at http://sartorlab.ccmb.med.umich.edu/software.

To identify which transcription factors (TFs) have significant level of hypermethylation or hypomethylation across their binding sites, which could indicate whether the TF is having a weaker or stronger regulatory effect, respectively, we first tile all reads from regions to which a particular TF is predicted to bind. We then apply our beta-binomial model to the data for each TF to identify TFs with hyper- or hypo-methylated binding sites.

To achieve this purpose, we provide fuction methylSigTileTFBS() to tile all data corresponding to the same TF.

```
> methTileTFs = methylSigTileTFBS(meth, tfbsInfo)
> myDiffTFs = methylSigCalc(methTileTFs, groups=c(1,0))
```

Total number of TFs: 126

7 Data visualization

MethylSig offers a unique two-tiered visualization of the methylation data depending on the zoom level. When the chromosome range is large (>1 million bp), the visualization function does not show individual sample data.

For narrow regions where at most $500~\rm CpG$ sites have data reads, users can visualize sample-specific coverage levels and % methylation at each site, together with group averages, significance levels and a number of genomic annotations.

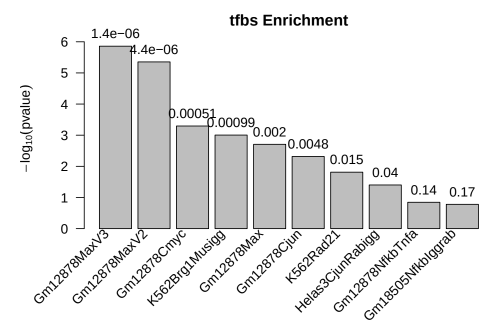


Figure 3: TFBS enrichment pvalue plot

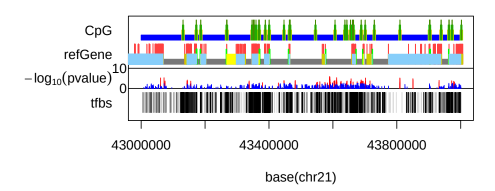


Figure 4: Data visualization in large range

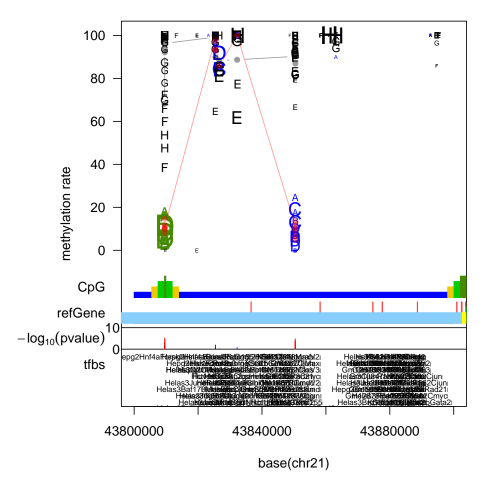


Figure 5: Data visualization within narrow range

8 Other

8.1 'methylSigData' object

8.1.1 S4 data structure

the methylSig package uses S4 object. The contents of 'methylSigData' can be shown using the 'show()' function in R or just type the object itself.

> meth

methylSigData object with 7,571 rows

	chr	start	end	strand	coverage1	${\tt numCs1}$	${\tt numTs1}$		coverage8	${\tt numCs8}$	numTs8		
1	chr21	43000564	43000564	_	NA	NA	NA		NA	NA	NA		
2	chr21	43000820	43000820	_	NA	NA	NA		NA	NA	NA		
3	chr21	43008527	43008527	+	32	32	0		124	122	2		
4	chr21	43008531	43008531	+	32	31	1		125	125	0		
5	chr21	43008543	43008543	+	32	29	3		125	117	8		
6	chr21	43008665	43008665	_	NA	NA	NA		NA	NA	NA		
7	chr21	43008673	43008673	-	27	27	0		29	25	4		
8	chr21	43008709	43008709	_	67	63	4		31	26	5		
9	chr21	43008719	43008719	_	67	62	5		33	33	0		
10	chr21	43014041	43014041	+	NA	NA	NA		207	202	5		

sample.ids: AML1 AML2 AML3 AML4 NBM1 NBM2 NBM3 NBM4

treatment: 1 1 1 1 0 0 0 0

destranded: TRUE
resolution: base

options: maxCount=500 & minCount=10 & assembly=hg18 & context=CpG

'NA' here means no data at this base location on the related sample.

8.1.2 Subsetting

Data can be subset using matrix style operations. Row represents base location and each column is a sample. Below is an example to obtain data for samples 1 to 4:

> meth1_4 = meth[,1:4]

This example returns the first 100 methylation reads in the data:

> methSub1_100 = meth[1:100,]

Two arguments can be used together. This example returns the first 100 methylation reads for samples 1 and 2.

- > methSubData = meth[1:100,1:2]
- > methSubData

methylSigData object with 60 rows

end strand coverage1 numCs1 numTs1 coverage2 numCs2 numTs2 chr start 1 chr21 43008527 43008527 32 32 0 124 122 2 chr21 43008531 43008531 32 125 125 0 31 1 32 3 chr21 43008543 43008543 29 3 125 117 8 chr21 43008673 43008673 27 27 0 29 25 4 5 chr21 43008709 43008709 67 63 4 31 26 5 6 chr21 43008719 43008719 67 62 5 33 33 Λ NA207 7 chr21 43014041 43014041 NANA202 5 8 chr21 43014044 43014044 NA207 12 NANA195 9 chr21 43014076 43014076 NANA 202 14 NA188

```
10 chr21 43016439 43016439 + 55 51 4 93 90 3
```

sample.ids: AML1 AML2

treatment: 1 1
destranded: TRUE
resolution: base

options: maxCount=500 & minCount=10 & assembly=hg18 & context=CpG

8.1.3 Getting values

If the second argument is a string that matches one of the column names in the methylSigData object, it gives the values of that column. Valid column names are "chr", "start", "end", "strand", "coverage1",..., "numCs1",..., and "numTs1"...

```
> coverage1 = meth[,"coverage1"]
> startTop200 = meth[1:200,"start"]
```

8.2 'methylSigDiff' object

8.2.1 S4 data structure

The contents of 'methylSigDiff' are

> myDiffSigboth

methylSigDiff object with 3,260 rows

```
qvalue meth.diff logLikRatio
       chr
                 start
                                 end strand
                                                      pvalue
   chr21 43008527 43008527 + 0.3219880 0.6664426 -2.4167475 1.27484057
2 chr21 43008531 43008531
                                              + 0.8718018 1.0000000 0.4158129 0.02957797
3 chr21 43008543 43008543
                                             + 0.5583045 0.8746145 -2.8071020 0.40674783
4 chr21 43014041 43014041
                                            + 0.3670413 0.7056865 2.3585424 0.98274395

      5
      chr21
      43014044
      43014044
      + 0.9166589
      1.0000000
      -0.6133413
      0.01210852

      6
      chr21
      43014076
      43014076
      + 0.1040236
      0.4228389
      12.1681339
      4.39674527

      7
      chr21
      43016439
      43016439
      + 0.2318347
      0.5885776
      -2.1713344
      1.76894807

      8
      chr21
      43016517
      43016517
      - 0.4054191
      0.7342590
      -2.0930416
      0.86326517

9 chr21 43018213 43018213
                                              + 0.8344303 1.0000000 -1.4186090 0.04765519
                                              - 0.3425235 0.6784244 2.0009532 1.09884498
10 chr21 43018274 43018274
              theta df
                                                muO
                                  m111
1 3.689198e+01 4 96.49842 98.91517
2 1.172001e+01 4 98.45532 98.03950
3 3.131946e+01 4 90.97520 93.78231
4 2.829499e+01 5 97.82788 95.46934
5 1.269081e+01 5 90.45700 91.07034
   1.904543e+01 4 93.23638 81.06825
7 9.151486e+01 6 96.96983 99.14116
8 1.000000e+06 4 93.10351 95.19656
```

9 7.094155e+00 6 91.07918 92.49778 10 7.115526e+01 5 98.16913 96.16818

sample.ids: AML1 AML2 AML3 AML4 NBM1 NBM2 NBM3 NBM4

treatment: 1 1 1 1 0 0 0 0

destranded: TRUE resolution: base

options: dispersion=both & local.disp=FALSE & local.meth=FALSE& min.per.group=c(3,3)& Total: 3260

8.2.2 Subsetting

This object can also subset by row to obtain results from part of CpG sites or regions. However, the qualues will not be readjusted.

```
> myDiff100 = myDiffSigboth[1:100,]
```

8.2.3 Getting values

Similar to the 'methSigData' object, if the second argument is a string that is the same as one of the column names, it will return the results for that column. The valid variable names are "chr", "start", "end", "strand", "pvalue", "qvalue", "meth.diff", "logLikRatio", "theta" "df", "mu1", and "mu0". Here, for group methylation mean estimates "mu1" and "mu0", 1 and 0 come from the 'groups' argument in the methylSigCalc() function. So if one has run the methylSigCalc() function with 'groups=c(4,0)', then "mu4" and "mu0" will appear in the results.

```
> qvalues = myDiffSigboth[,"qvalue"]
```

8.2.4 How to subtract DMCs or DMRs

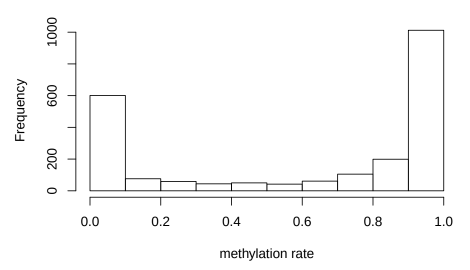
This 'methylSigDiff' object is very flexible to use by combining functions of subsetting and getting values. For example, the following code can obtain differentially methylated cytosines or regions defined as qualue < 0.05 and difference of methylation rate > 25%.

8.3 Summarizing data

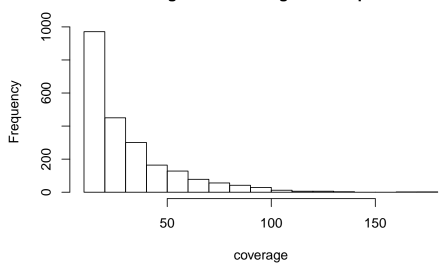
You can easily use other R functions to summarize or draw plots.

```
> methRaw = methylSigReadData(fileList, sample.ids = sample.id, assembly = "hg18",
               treatment = treatment, context = "CpG", minCount = 0,
               maxCount=Inf, destranded=F, quiet=T)
> summary(methRaw[,"numCs1"]/methRaw[,"coverage1"])
   Min. 1st Qu.
                 Median
                           Mean 3rd Qu.
                                                    NA's
                                            Max.
  0.000
          0.069
                  0.861
                          0.612
                                   1.000
                                           1.000
                                                    3606
> summary(methRaw[,"coverage1"])
  Min. 1st Qu. Median
                           Mean 3rd Qu.
                                            Max.
                                                    NA's
  10.00
          14.00
                  23.00
                          30.68
                                  39.00
                                         177.00
                                                    3606
> hist(methRaw[,"numCs1"]/methRaw[,"coverage1"],
                   main="Histogram of methylation rate for sample 1",
                   xlab="methylation rate")
```

Histogram of methylation rate for sample 1



Histogram of coverage for sample 1



8.4 Generating heatmap

Here we provide an example to generate a correlation heatmap.

Here is another example to generate a correlation heatmap based on differentially methylated cytosines.

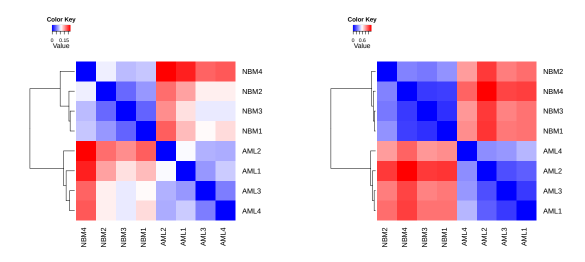


Figure 6: Heatmap based on methylation rate at all CpG sites (left) and at differentially methylated CpG sites (right)

9 References

Park, Y., Figueroa, M. E., Rozek, L. S., and sartor M.A., "methylSig: a whole genome DNA methylation analysis pipeline". Submitted.