methylKit: User Guide

Altuna Akalin Matthias I ala2027@med.cornell.edu mk375@cc

Matthias Kormaksson mk375@cornell.edu

Sheng Li shl2018@med.cornell.edu

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1 Introduction

In this manual, we will show how to use the methylKit package. methylKit is an R package for analysis and annotation of DNA methylation information obtained by high-throughput bisulfite sequencing. The package is designed to deal with sequencing data from RRBS and its variants. But it can potentially handle whole-genome bisulfite sequencing data if proper input format is provided.

1.1 DNA methylation

DNA methylation in vertebrates typically occurs at CpG dinucleotides, however non-CpG Cs are also methylated in certain tissues such as embryonic stem cells. DNA Methylation can act as an epigenetic control mechanism for gene regulation. Methylation can hinder binding of transcription factors and/or methylated bases can be bound by methyl-binding-domain proteins which can recruit chromatin remodeling factors. In both cases, the transcription of the regulated gene will be effected. In addition, aberrant DNA methylation patterns have been associated with many human malignancies and can be used in a predictive manner. In malignant tissues, DNA is either hypo-methylated or hyper-methylated compared to the normal tissue. The location of hyper- and hypo-methylated sites gives a distinct signature to many diseases. Traditionally, hypo-methylation is associated with gene transcription (if it is on a regulatory region such as promoters) and hyper-methylation is associated with gene repression.

1.2 High-throughput bisulfite sequencing

Bisulfite sequencing is a technique that can determine DNA methylation patterns. The major difference from regular sequencing experiments is that, in bisulfite sequencing DNA is treated with bisulfite which converts cytosine residues to uracil, but leaves 5-methylcytosine residues unaffected. By sequencing and aligning those converted DNA fragments it is possible to call methylation status of a base. Usually, the methylation status of a base determined by a highthroughput bisulfite sequencing will not be a binary score, but it will be a percentage. The percentage simply determines how many of the bases that are aligning to a given cytosine location in the genome have actual C bases in the reads. Since bisulfite treatment leaves methylated Cs intact, that percentage will give us percent methylation score on that base. The reasons why we will not get a binary response are 1) the probable sequencing errors in high-throughput sequencing experiments 2) incomplete bisulfite conversion 3) (and a more likely scenario) is heterogeneity of samples and heterogeneity of paired chromosomes from the same sample

2 Basics

2.1 Reading the methylation call files

We start by reading in the methylation call data from bisulfite sequencing with **read** function. Reading in the data this way will return a methylRawList object which stores methylation information per sample for each covered base. The methylation call files are basically text files that contain percent methylation score per base. A typical methylation call file looks like this:

##	chrBase	chr	base	\mathtt{strand}	coverage	freqC	freqT	
## 1	chr21.9764539	chr21	9764539	R	12	25.00	75.00	
## 2	chr21.9764513	chr21	9764513	R	12	0.00	100.00	
## 3	chr21.9820622	chr21	9820622	F	13	0.00	100.00	
## 4	chr21.9837545	chr21	9837545	F	11	0.00	100.00	
## 5	chr21.9849022	chr21	9849022	F	124	72.58	27.42	

Most of the time bisulfite sequencing experiments have test and control samples. The test samples can be from a disease tissue while the control samples can be from a healthy tissue. You can read a set of methylation call files that have test/control conditions giving treatment vector option. For sake of subsequent analysis, file.list, sample.id and treatment option should have the same order. In the following example, first two files are have the sample ids "test1" and "test2" and as determined by treatment vector they belong to the same group. The third and fourth files have sample ids "ctrl1" and "ctrl2" and they belong to the same group as indicated by the treatment vector.

```
library(methylKit)
file.list <- list(system.file("extdata", "test1.myCpG.txt",
    package = "methylKit"), system.file("extdata",
    "test2.myCpG.txt",
    package = "methylKit"), system.file("extdata",
    "control1.myCpG.txt",
    package = "methylKit"), system.file("extdata",
    "control2.myCpG.txt",
    package = "methylKit"))
# read the files to a methylRawList object: myobj
myobj <- read(file.list, sample.id = list("test1",
    "test2", "ctrl1", "ctrl2"), assembly = "hg18", treatment =
c(1,
    1, 0, 0), context = "CpG")</pre>
```

2.2 Reading the methylation calls from sorted Bismark alignments

Alternatively, methylation percentage calls can be calculated from sorted SAM file(s) from Bismark aligner and read-in to the memory. Bismark is a popular aligner for bisulfite sequencing reads [1]. read.bismark function is designed to read-in Bismark SAM files as methylRaw or methylRawList objects which store per base methylation calls. SAM files must be sorted by chromosome and read position columns, using 'sort' command in unix-like machines will accomplish such a sort easily.

The following command reads a sorted SAM file and creates a methylRaw object for CpG methylation. The user has the option to save the methylation call files to a folder given by save.folder option. The saved files can be read-in using the read function when needed.

```
my.methRaw <- read.bismark(location = system.file("extdata",
    "test.fastq_bismark.sorted.min.sam", package = "methylKit"),
    sample.id = "test1", assembly = "hg18", read.context = "CpG",
    save.folder = getwd())
```

It is also possible to read multiple SAM files at the same time, check read.bismark documentation.

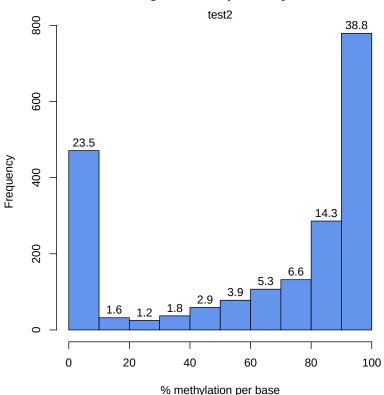
2.3 Descriptive statistics on samples

Since we read the methylation data now, we can check the basic stats about the methylation data such as coverage and percent methylation. We now have a methylRawList object which contains methylation information per sample. The following command prints out percent methylation statistics for second sample: "test2"

```
getMethylationStats(myobj[[2]], plot = F, both.strands = F)
## methylation statistics per base
## summary:
##
      Min. 1st Qu.
                     Median
                                Mean 3rd Qu.
                                                  Max.
                       82.8
##
       0.0
               20.0
                                63.2
                                         94.7
                                                100.0
## percentiles:
##
       0%
              10%
                     20%
                             30%
                                     40%
                                            50%
                                                    60%
                                                           70%
##
     0.00
             0.00
                    0.00
                           48.39
                                  70.00
                                          82.79
                                                  90.00
                                                         93.33
##
      80%
              90%
                     95%
                             99%
                                  99.5%
                                          99.9%
                                                   100%
    96.43 100.00 100.00 100.00 100.00 100.00 100.00
##
##
```

The following command plots the histogram for percent methylation distribution. The figure below is the histogram and numbers on bars denote what percentage of locations are contained in that bin. Typically, percent methylation histogram should have two peaks on both ends. In any given cell, any given base are either methylated or not. Therefore, looking at many cells should yield a similar pattern where we see lots of locations with high methylation and lots of locations with low methylation.

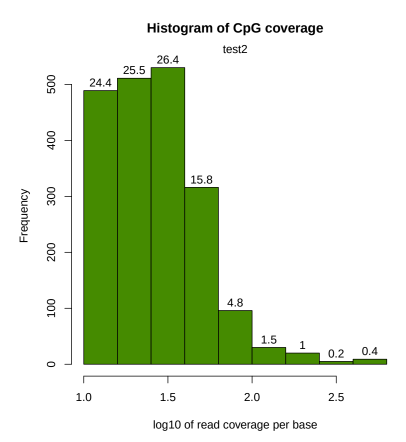
getMethylationStats(myobj[[2]], plot = T, both.strands = F)



Histogram of % CpG methylation

We can also plot the read coverage per base information in a similar way, again numbers on bars denote what percentage of locations are contained in that bin. Experiments that are highly suffering from PCR duplication bias will have a secondary peak towards the right hand side of the histogram.

```
library("graphics")
getCoverageStats(myobj[[2]], plot = T, both.strands = F)
```



2.4 Filtering samples based on read coverage

It might be useful to filter samples based on coverage. Particularly, if our samples are suffering from PCR bias it would be useful to discard bases with very high read coverage. Furthermore, we would also like to discard bases that have low read coverage, a high enough read coverage will increase the power of the statistical tests. The code below filters a methylRawList and discards bases that have coverage below 10X and also discards the bases that have more than 99.9th percentile of coverage in each sample.

3 Comparative analysis

3.1 Merging samples

In order to do further analysis, we will need to get the bases covered in all samples. The following function will merge all samples to one object for basepair locations that are covered in all samples. Setting destrand=TRUE (the default is FALSE) will merge reads on both strands of a CpG dinucleotide. This provides better coverage, but only advised when looking at CpG methylation (for CpH methylation this will cause wrong results in subsequent analyses). In addition, setting destrand=TRUE will only work when operating on base-pair resolution, otherwise setting this option TRUE will have no effect. The unite() function will return a methylBase object which will be our main object for all comparative analysis. The methylBase object contains methylation information for regions/bases that are covered in all samples.

meth <- unite(myobj, destrand = FALSE)</pre>

Let us take a look at the data content of methylBase object:

head(meth)

##			id	chr		start	end	strand	coverage1
##	1	chr21.1	10011833						174
##	2	chr21.2	10011841	chr21	100	011841	10011841	+	173
##	3	chr21.2	10011855	chr21	100	011855	10011855	+	175
##	4	chr21.2	10011858	chr21	100	011858	10011858	+	175
##	5	chr21.2	10011861	chr21	100	011861	10011861	+	174
##	6	chr21.2	10011872	chr21	100	011872	10011872	+	167
##		numCs1	numTs1 c	overag	ge2	numCs2	2 numTs2	coverage	e3 numCs3
##	1	173	1		18	18	8 0	4	10 34
##	2	164	9		20	19) 1	4	10 18
##	3	175	0		21	21	. 0	3	39 29
##	4	131	44		21	20) 1	3	39 31
##	5	147	27		20	15	5 5	3	39 13
##	6	160	7		20	19) 1	3	39 34
##		numTs3	coverage	4 num(Cs4	numTs4	.		
##	1	6	1	4	14	C)		
##	2	22	1	4	8	6	5		
##	3	10	1	4	12	2	2		
##	4	8	1	3	8	5	5		
##	5	26	1	3	9	4			
##	6	5	1	4	8	6	;		

By default, unite function produces bases/regions covered in all samples. That requirement can be relaxed using "min.per.group" option in unite function.

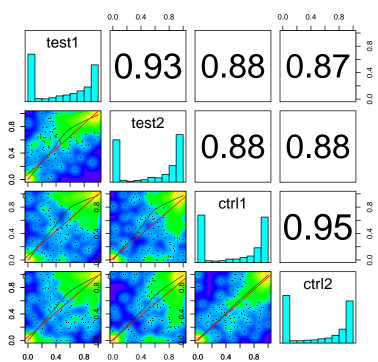
```
# creates a methylBase object. Only CpGs covered at least
# in 1 sample per group will be returned there were two
# groups defined by the treatment vector given during the
# creation of myobj treatment=c(1,1,0,0)
meth.min <- unite(myobj, min.per.group = 1L)</pre>
```

3.2 Sample Correlation

We can check the correlation between samples using getCorrelation. This function will either plot scatter plot and correlation coefficients or just print a correlation matrix

```
getCorrelation(meth, plot = T)
```

```
## test1 test2 ctrl1 ctrl2
## test1 1.0000 0.9253 0.8768 0.8738
## test2 0.9253 1.0000 0.8792 0.8802
## ctrl1 0.8768 0.8792 1.0000 0.9465
## ctrl2 0.8738 0.8802 0.9465 1.0000
```



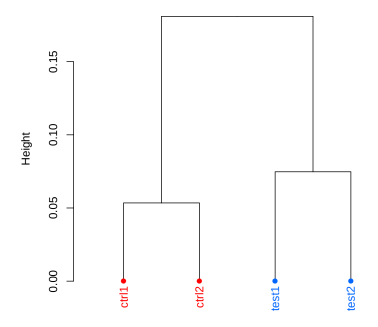
CpG base correlation

3.3 Clustering samples

We can cluster the samples based on the similarity of their methylation profiles. The following function will cluster the samples and draw a dendrogram.

clusterSamples(meth, dist = "correlation", method = "ward",
 plot = TRUE)

CpG methylation clustering



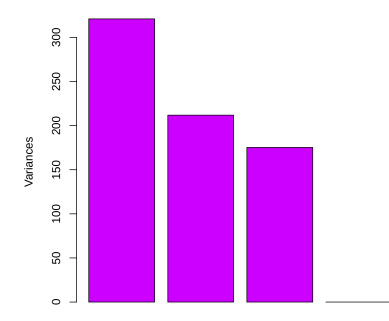
Samples Distance method: "correlation"; Clustering method: "ward"

```
##
## Call:
## hclust(d = d, method = HCLUST.METHODS[hclust.method])
##
## Cluster method : ward
## Distance : pearson
## Number of objects: 4
##
```

Setting the plot=FALSE will return a dendrogram object which can be manipulated by users or fed in to other user functions that can work with dendrograms.

hc <- clusterSamples(meth, dist = "correlation", method = "ward",
 plot = FALSE)</pre>

We can also do a PCA analysis on our samples. The following function will plot a scree plot for importance of components.

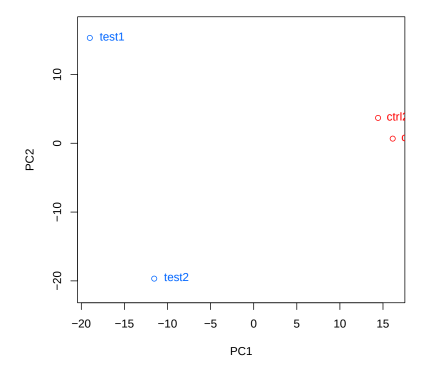


CpG methylation PCA Screeplot

We can also plot PC1 and PC2 axis and a scatter plot of our samples on those axis which will reveal how they cluster.

PCASamples(meth)





3.4 Tiling windows analysis

For some situations, it might be desirable to summarize methylation information over tiling windows rather than doing base-pair resolution analysis. methylKit provides functionality to do such analysis. The function below tiles the genome with windows 1000bp length and 1000bp step-size and summarizes the methylation information on those tiles. In this case, it returns a methylRawList object which can be fed into unite and calculateDiffMeth functions consecutively to get differentially methylated regions.

```
tiles <- tileMethylCounts(myobj, win.size = 1000,</pre>
    step.size = 1000)
head(tiles[[1]])
##
                         id
                              chr
                                    start
                                               end strand
## 1 chr21.9764001.9765000 chr21 9764001 9765000
                                                         *
## 2 chr21.9820001.9821000 chr21 9820001 9821000
                                                         *
##
  3 chr21.9837001.9838000 chr21 9837001 9838000
                                                         *
## 4 chr21.9849001.9850000 chr21 9849001 9850000
                                                         *
```

##	5	chr21.985	53001.9	9854000	chr21	9853001	9854000	*	
##	6	chr21.986	50001.9	9861000	chr21	9860001	9861000	*	
##		coverage	$\verb"numCs"$	numTs					
##	1	24	3	21					
##	2	13	0	13					
##	3	11	0	11					
##	4	124	90	34					
##	5	34	22	12					
##	6	39	38	1					

3.5 Finding differentially methylated bases or regions

calculateDiffMeth() function is the main function to calculate differential methylation. Depending on the sample size per each set it will either use Fisher's exact or logistic regression to calculate P-values. P-values will be adjusted to Q-values using SLIM method [2].

```
myDiff <- calculateDiffMeth(meth)</pre>
```

After q-value calculation, we can select the differentially methylated regions/bases based on q-value and percent methylation difference cutoffs. Following bit selects the bases that have q-value;0.01 and percent methylation difference larger than 25%. If you specify type="hyper" or type="hype" options, you will get hyper-methylated or hypo-methylated regions/bases.

```
# get hyper methylated bases
myDiff25p.hyper <- get.methylDiff(myDiff, difference = 25,
        qvalue = 0.01, type = "hyper")
# get hypo methylated bases
myDiff25p.hypo <- get.methylDiff(myDiff, difference = 25,
        qvalue = 0.01, type = "hypo")
#
#
#
# get all differentially methylated bases
myDiff25p <- get.methylDiff(myDiff, difference = 25,
        qvalue = 0.01)
```

We can also visualize the distribution of hypo/hyper-methylated bases/regions per chromosome using the following function. In this case, the example set includes only one chromosome. The **list** shows percentages of hypo/hyper methylated bases over all the covered bases in a given chromosome.

```
diffMethPerChr(myDiff, plot = FALSE, qvalue.cutoff = 0.01,
    meth.cutoff = 25)
```

```
## $diffMeth.per.chr
##
       chr number.of.hypomethylated
## 1 chr21
                                  59
##
     percentage.of.hypomethylated number.of.hypermethylated
## 1
                             6.127
                                                            75
##
     percentage.of.hypermethylated
## 1
                              7.788
##
## $diffMeth.all
##
     percentage.of.hypermethylated number.of.hypermethylated
## 1
                              7.788
                                                             75
##
     percentage.of.hypomethylated number.of.hypomethylated
## 1
                                                          59
                             6.127
##
```

3.5.1 Finding differentially methylated bases using multiple-cores

The differential methylation calculation speed can be increased substantially by utilizing multiple-cores in a machine if available. Both Fisher's Exact test and logistic regression based test are able to use multiple-core option.

The following piece of code will run differential methylation calculation using 2 cores.

myDiff <- calculateDiffMeth(meth, num.cores = 2)</pre>

4 Annotating differentially methylated bases or regions

We can annotate our differentially methylated regions/bases based on gene annotation. In this example, we read the gene annotation from a bed file and annotate our differentially methylated regions with that information. This will tell us what percentage of our differentially methylated regions are on promoters/introns/exons/intergenic region. Similar gene annotation can be fetched using GenomicFeatures package available from Bioconductor.org.

```
gene.obj <- read.transcript.features(system.file("extdata",
                "refseq.hg18.bed.txt", package = "methylKit"))
# annotate differentially methylated Cs with
# promoter/exon/intron using annotation data
annotate.WithGenicParts(myDiff25p, gene.obj)
## summary of target set annotation with genic parts
## 133 rows in target set
```

```
## ------
## -----
   percentage of target features overlapping with annotation :
##
##
                             intron intergenic
     promoter
                   exon
                   15.04
##
        27.82
                              34.59
                                         57.14
##
##
## percentage of target features overlapping with annotation (with promoter>exon>intron pre-
                             intron intergenic
##
     promoter
                    exon
##
        27.82
                    0.00
                              15.04
                                         57.14
##
##
## percentage of annotation boundaries with feature overlap :
## promoter
                exon
                       intron
   0.28604 0.02683 0.17068
##
##
##
## summary of distances to the nearest TSS :
##
     Min. 1st Qu. Median
                              Mean 3rd Qu.
                                              Max.
               828
                                     94600 314000
##
         5
                     45200
                             52000
```

Similarly, we can read the CpG island annotation and annotate our differentially methylated bases/regions with them.

```
# read the shores and flanking regions and name the flanks
# as shores and CpG islands as CpGi
cpg.obj <- read.feature.flank(system.file("extdata",
        "cpgi.hg18.bed.txt", package = "methylKit"),
feature.flank.name = c("CpGi",
        "shores"))
#
diffCpGann <- annotate.WithFeature.Flank(myDiff25p,
        cpg.obj$CpGi, cpg.obj$shores, feature.name = "CpGi",
flank.name = "shores")</pre>
```

4.1 Regional analysis

We can also summarize methylation information over a set of defined regions such as promoters or CpG islands. The function below summarizes the methylation information over a given set of promoter regions and outputs a methylRaw or methylRawList object depending on the input.

```
promoters <- regionCounts(myobj, gene.obj$promoters)
head(promoters[[1]])</pre>
```

##			id	chr	start	end	strand
## 1 c	hr21.178	306094	.17808094.NA	chr21	17806094	17808094	+
## 2 c	hr21.101	19796	.10121796.NA	chr21	10119796	10121796	-
## 3 c	hr21.100	011791	.10013791.NA	chr21	10011791	10013791	-
## 4 c	hr21.101	19808	.10121808.NA	chr21	10119808	10121808	-
## 5 c	hr21.153	357997	.15359997.NA	chr21	15357997	15359997	-
## 6 c	hr21.160	023366	.16025366.NA	chr21	16023366	16025366	+
## c	overage	$\verb"numCs"$	numTs				
## 1	1834	7	1827				
## 2	79	44	35				
## 3	3697	2982	715				
## 4	79	44	35				
## 5	8613	16	8594				
## 6	6296	5	6291				

4.2Convenience functions for annotation objects

After getting the annotation of differentially methylated regions, we can get the distance to TSS and nearest gene name using the getAssociationWithTSS function.

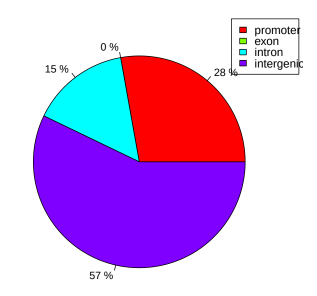
```
diffAnn <- annotate.WithGenicParts(myDiff25p, gene.obj)</pre>
# target.row is the row number in myDiff25p
head(getAssociationWithTSS(diffAnn))
##
        target.row dist.to.feature feature.name feature.strand
## 60
                 1
                                951
                                        NM_199260
                                        NM_199260
## 60.1
                 2
                                931
## 60.2
                 3
                                        NM_199260
                                838
## 60.3
                  4
                                828
                                        NM_199260
## 60.4
                  5
                                802
                                        NM_199260
## 60.5
                  6
                                723
                                        NM_199260
```

It is also desirable to get percentage/number of differentially methylated regions that overlap with intron/exon/promoters

_

```
getTargetAnnotationStats(diffAnn, percentage = TRUE,
    precedence = TRUE)
##
     promoter
                              intron intergenic
                     exon
##
        27.82
                    0.00
                               15.04
                                          57.14
```

We can also plot the percentage of differentially methylated bases overlapping with exon/intron/promoters



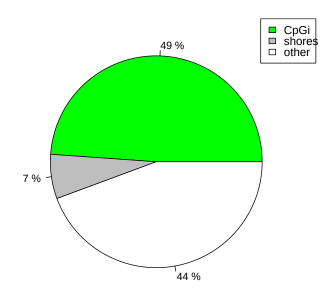
differential methylation annotation

plotTargetAnnotation(diffAnn, precedence = TRUE, main =

"differential methylation annotation")

We can also plot the CpG island annotation the same way. The plot below shows what percentage of differentially methylated bases are on CpG islands, CpG island shores and other regions.

```
plotTargetAnnotation(diffCpGann, col = c("green",
      "gray", "white"), main = "differential methylation
      annotation")
```



differential methylation annotation

It might be also useful to get percentage of intron/exon/promoters that overlap with differentially methylated bases.

```
getFeatsWithTargetsStats(diffAnn, percentage = TRUE)
## promoter exon intron
## 0.28604 0.02683 0.17068
```

5 methylKit convenience functions

5.1 coercion

Most methylKit objects (methylRaw,methylBase and methylDiff) can be coerced to GRanges objects from GenomicRanges package. Coercing methylKit objects to GRanges will give users additional flexiblity when customising their analyses.

class(meth)

[1] "methylBase"
attr(,"package")
[1] "methylKit"

as(meth, "GRanges")

##	GRanges	with 963	ranges	and	13 e	element	Metad	lata cols:
##		seqnames			r	anges	strar	nd
##		<rle></rle>			<ira< th=""><th>inges></th><th><rle< th=""><th>e> </th></rle<></th></ira<>	inges>	<rle< th=""><th>e> </th></rle<>	e>
##	[1]	chr21	[100118	833,	1001	1833]		+
##	[2]	chr21	[100118	841,	1001	.1841]		+
##	[3]	chr21	-	855,	1001	1855]		+
##	[4]	chr21	-			1858]		+
##	[5]	chr21				.1861]		+
##	[6]	chr21				.1872]		+
##	[7]	chr21	[100118			1876]		+
##	[8]	chr21	-			.1878]		+
##	[9]	chr21	[100119	925,	1001	1925]		-
##	•••	•••	_			•••	• •	• • • •
##	[955]	chr21	-	4505,		4505]		+
##	[956]	chr21				4663]		-
##	[957]	chr21				59407]		+
##	[958]	chr21				59541]		-
##	[959]	chr21				59569]		-
##	[960]	chr21	-			59577]		-
##	[961]	chr21	_			59644]		-
##	[962]	chr21				59650]		-
##	[963]	chr21				67634]	- ·	-
##				covei	0		umCs1	numTs1
##	Г 4 Л			<inte< th=""><th>-</th><th></th><th></th><th><numeric></numeric></th></inte<>	-			<numeric></numeric>
##		chr21.100			174		173	1
##	[2]				173		164	9
##	[3]				175		175	0
##	[4]	chr21.100			175		131	44
## ##	[5] [6]	chr21.100			174 167		147 160	27 7
## ##	[7]	chr21.100			160		148	12
## ##	[8]	chr21.100			150		134	12
## ##	[9]	chr21.100			120		65	55
## ##	[9]	01121.100	11920					
##	 [955]	chr21.99			 37		· · · 2	 35
##	[956]	chr21.99			61		19	42
##	[957]	chr21.99			44	-	17	27
##	[958]	chr21.99			26		12	14
nπ	[000]	0111 21.00			20		12	11

##	[959]	chr21.998	59569	25	17	8
##	[960]	chr21.99	59577	25	25	0
##	[961]	chr21.99	59644	21	0	21
##	[962]	chr21.99	59650	21	6	15
##	[963]	chr21.996	67634	10	0	10
##		coverage2	numCs2	numTs2	coverage3	numCs3
##		<integer></integer>	<numeric></numeric>	<numeric></numeric>	<integer></integer>	<numeric></numeric>
##	[1]	18	18	0	40	34
##	[2]	20	19	1	40	18
##	[3]	21	21	0	39	29
##	[4]	21	20	1	39	31
##	[5]	20	15	5	39	13
##	[6]	20	19	1	39	34
##	[7]	21	18	3	38	24
##	[8]	20	19	1	37	20
##	[9]	37	21	16	68	21
##						
##	[955]	147	56	91	86	79
##	[956]	116	71	45	45	35
##	[957]	118	58	60	52	49
##	[958]	76	44	32	39	37
##	[959]	77	69	8	40	40
##	[960]	77	71	6	40	40
##	[961]	97	50	47	59	52
##	[962]	103	57	46	59	51
##	[963]	61	25	36	93	62
##			coverage4	numCs4	numTs4	
##		<numeric></numeric>	<integer></integer>	<numeric></numeric>	<numeric></numeric>	
##	[1]	6	14	14	0	
##	[2]	22	14	8	6	
##	[3]	10	14	12	2	
##	[4]	8	13	8	5	
##	[5]	26	13	9	4	
##	[6]	5	14	8	6	
##	[7]	14	11	9	2	
##	[8]	17	12	12	0	
##	[9]	47	20	6	14	
##						
##	[955]	7	40	25	15	
##	[956]	10	31	25	6	
##	[957]	3	40	27	13	
##	[958]	2	39	32	7	
##	[959]	0	39	35	4	
##	[960]	0	39	36	3	
##	[961]	7	31	14	17	

[962] 32 8 21 11 ## [963] 31 56 29 27 ## ___ ## seqlengths: ## chr21 ## NA class(myDiff) ## [1] "methylDiff" ## attr(,"package") ## [1] "methylKit" as(myDiff, "GRanges") ## GRanges with 963 ranges and 3 elementMetadata cols: ## seqnames ranges strand ## <Rle> <IRanges> <Rle> chr21 [10011833, 10011833] ## [1] + ## [2] chr21 [10011841, 10011841] + ## [3] chr21 [10011855, 10011855] + ## chr21 [10011858, 10011858] [4] + chr21 [10011861, 10011861] ## [5] + ## chr21 [10011872, 10011872] [6] + ## chr21 [10011876, 10011876] [7] + chr21 [10011878, 10011878] ## [8] + ## [9] chr21 [10011925, 10011925] ## [9944505, 9944505] ## [955] chr21 + [9944663, 9944663] ## [956] chr21 ## [957] chr21 [9959407, 9959407] + [9959541, 9959541] ## [958] chr21 ## [959] chr21 [9959569, 9959569] [9959577, 9959577] ## [960] chr21 _ ## [9959644, 9959644] [961] chr21 ## [962] chr21 [9959650, 9959650] ## [963] [9967634, 9967634] chr21 ## id qvalue meth.diff ## <factor> <numeric> <numeric> ## [1] chr21.10011833 8.543e-04 10.590 ## [2] chr21.10011841 6.050e-13 46.671 ## [3] chr21.10011855 4.579e-09 22.642 ## [4] chr21.10011858 5.922e-01 2.041 ## [5] chr21.10011861 8.163e-08 41.197 ## [6] chr21.10011872 1.238e-03 16.477 [7] chr21.10011876 1.933e-04 ## 24.366

##	[8]	chr21.10011878	3.489e-04	24.694				
##	[9]	chr21.10011925	8.543e-04	24.095				
##	• • •							
##	[955]	chr21.9944505	0.000e+00	-51.018				
##	[956]	chr21.9944663	7.678e-05	-28.100				
##	[957]	chr21.9959407	4.839e-08	-36.312				
##	[958]	chr21.9959541	3.145e-06	-33.560				
##	[959]	chr21.9959569	3.702e-02	-10.623				
##	[960]	chr21.9959577	4.923e-01	-2.085				
##	[961]	chr21.9959644	3.291e-05	-30.960				
##	[962]	chr21.9959650	6.575e-05	-28.314				
##	[963]	chr21.9967634	1.028e-03	-25.863				
##								
##	seqlengths:							
##	chr21	1						
##	NA	Α						

5.2 select

We can also select rows from methylRaw, methylBase and methylDiff objects with "select" function. An appropriate methylKit object will be returned as a result of "select" function.

se	lect	t(meth, 1	:10) #	‡ seled	ct first	10 rows o	of a methy	lBase object
##			id	chr	start	ene	d strand c	overage1
##	1	chr21.10	011833	chr21	10011833	1001183	3 +	174
##	2	chr21.10	011841	chr21	10011841	1001184	1 +	173
##	3	chr21.10	011855	chr21	10011855	1001185	5 +	175
##	4	chr21.10	011858	chr21	10011858	1001185	8 +	175
##	5	chr21.10	011861	chr21	10011861	1001186	1 +	174
##	6	chr21.10	011872	chr21	10011872	1001187	2 +	167
##	7	chr21.10	011876	chr21	10011876	1001187	6 +	160
##	8	chr21.10	011878	chr21	10011878	1001187	3 +	150
##	9	chr21.10	011925	chr21	10011925	1001192	5 –	120
##	10	chr21.10	011938	chr21	10011938	1001193	8 –	134
##		numCs1 n	umTs1 d	coverag	ge2 numCs	2 numTs2	coverage3	numCs3
##	1	173	1		-	8 0	40	
##	2	164	9		20 1	9 1	40	18
##	3	175	0		21 2	1 0	39	29
##	4	131	44		21 2	0 1	39	31
##	5	147	27		20 1	5 5	39	13
##	6	160	7		20 1	9 1	39	34
##	7	148	12		21 1	8 3	38	24
##	8	134	16		20 1	9 1	37	20

##		65	55		37	21		16		68	21
## ##	10	127	7	1	36	34 numTa/		2	-	74	64
## ##	1	11un 153 6	coverage	4 11uiii) 4	14		±)				
##		22		4	8	e					
##	3	10		4	12		2				
##	4	8	1	3	8	Ę	5				
##	5	26	1	3	9	4	1				
##	6	5	1	4	8	6	3				
##	7	14		1	9	2	2				
##	8	17	1	2	12	()				
##		47		:0	6	14					
##	10	10	2	:0	17	3	3				
sel	<pre>select(myDiff, 20:30) # select rows 10 of a methylDiff object</pre>										
##			id	chr		start		end	strand	р	value
##	20	chr21.2	10012079	chr21	100	012079	100	12079	+	1.32	5e-07
##	21	chr21.2	10012089	chr21	100	012089	100	12089	+	6.79	7e-02
##	22	chr21.2	10012095	chr21	100	012095	100	12095	+	9.12	5e-02
			10012101								2e-16
			10012696								3e-03
			10012699								3e-09
			10012876								1e-01
			10012881								0e+00
			10012883								7e-01
			10012887 10012891								5e-02 1e-01
##	50		lue meth.		100	512051	100	12091		0.09	16 01
	20	1.050e-		.617							
		1.048e-		.564							
		1.324e-		.726							
##	23	4.221e-	-14 39	.808							
##	24	6.033e-	-03 9	.685							
##	25	1.955e-	-08 44	.703							
##	26	4.224e-	-01 3	.888							
##	27	5.922e-	-01 0	.000							
		4.252e-		.750							
		3.316e-		.808							
##	30	5.922e-	-01 0	.686							

5.3 reorganize

methylBase and methylRawList can be reorganized by reorganize function. The function can subset the objects based on provided sample ids, it also cre-

ates a new treatment vector determining which samples belong to which group. Order of sample ids should match the treatment vector order.

```
# creates a new methylRawList object
myobj2 <- reorganize(myobj, sample.ids = c("test1",
    "ctrl2"), treatment = c(1, 0))
# creates a new methylBase object
meth2 <- reorganize(meth, sample.ids = c("test1",
    "ctrl2"), treatment = c(1, 0))</pre>
```

5.4 percMethylation

Percent methylation values can be extracted from methylBase object by using percMethylation function.

```
# creates a matrix containing percent methylation values
perc.meth <- percMethylation(meth)</pre>
```

6 Acknowledgements

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7 R session info

```
sessionInfo()
```

```
## R version 2.15.0 (2012-03-30)
## Platform: x86_64-apple-darwin9.8.0/x86_64 (64-bit)
##
## locale:
## [1] en_US.UTF-8/en_US.UTF-8/en_US.UTF-8/C/en_US.UTF-8/en_US.UTF-8
##
## attached base packages:
## [1] stats
                 graphics grDevices utils
                                                datasets
## [6] methods
                 base
##
## other attached packages:
## [1] data.table_1.8.0 methylKit_0.5
                                         knitr_0.4
```

```
##
## loaded via a namespace (and not attached):
##
    [1] BiocGenerics_0.2.0
                             codetools_0.2-8
    [3] digest_0.5.2
                             evaluate_0.4.2
##
    [5] formatR_0.4
                             GenomicRanges_1.8.3
##
##
    [7] highlight_0.3.1
                             IRanges_1.14.2
    [9] KernSmooth_2.23-7
                             parallel_2.15.0
##
## [11] parser_0.0-14
                             plyr_1.7.1
                             stats4_2.15.0
## [13] Rcpp_0.9.10
                             tools_2.15.0
## [15] stringr_0.6
```

References

- Felix Krueger and Simon R Andrews. Bismark: a flexible aligner and methylation caller for Bisulfite-Seq applications. *Bioinformatics (Oxford, England)*, 27(11):1571–2, June 2011.
- [2] Hong-Qiang Wang, Lindsey K Tuominen, and Chung-Jui Tsai. SLIM: a sliding linear model for estimating the proportion of true null hypotheses in datasets with dependence structures. *Bioinformatics (Oxford, England)*, 27(2):225–31, January 2011.