BiSeq: A package for analyzing targeted bisulfite sequencing data

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1 Introduction

The *BiSeq* package provides useful classes and functions to handle and analyze targeted bisulfite sequencing (BS) data such as reduced representation bisulfite sequencing (RRBS) data. In particular, it implements an algorithm to detect differentially methylated regions (DMRs), as described in detail in [1]. The package takes already aligned BS data from one or multiple samples. Until now, it was used for the analysis of CpG methylation of human and mouse samples only.

2 Data import and classes

2.1 Sample data

As sample data we use a small part of a recently published data set, see [2]. It comprises RRBS data from 10 samples of CpG sites from genomic regions on p arms of chromosome 1 and 2 covered in at least one sample. Data was obtained from 5 bone marrow probes of patients with acute promyelocytic leukemia (APL) and 5 control samples (APL in remission). RRBS data was preprocessed with the Bismark software version 0.5 [3].

2.2 Import of Bismark's methylation output files

Bismark [3] a bisulfite read mapper and methylation caller. BiSeq allows the import of Bismark output files.

> library(BiSeq)

readBismark imports the CpG context output files created by the methylation extractor of Bismark:

> readBismark(files, colData)

The argument files point to files created by Bismark's methylation_extractor and bismark2bedGraph (see the man page of readBismark for details on how to retrieve the right input files). colData specifies the sample names and additional phenotype information. This method returns a *BSraw* object.

2.3 The BSraw and BSrel classes

The *BiSeq* package contains the classes *BSraw* and *BSrel*, both derived from *SummarizedExperiment*.

2.3.1 The BSraw class

The *BSraw* class is a container for 'raw' RRBS data. It comprises sample information together with CpG positions and numbers of reads spanning the CpG positions as well as the number of methylated reads.

A BSraw object consists of four slots:

- 1. A *SimpleList* of arbitrary content describing the overall experiment, accessible with exptData.
- 2. A *GRanges* of the positions of CpG-sites covered by BS in at least one sample, accessible with rowData.
- 3. A *DataFrame* of samples and the values of variables measured on those samples, accessible with colData.
- 4. An assays slot containing a *SimpleList* of two matrices, one containing the numbers of reads (accessible with totalReads) and the other the numbers of methylated reads (accessible with methReads).

A new *BSraw* object can also be created by hand:

Nevertheless, users will most likely create *BSraw* objects when use **readBis**-**mark** to load data.

We load and show the APL data:

```
> data(rrbs)
> rrbs
class: BSraw
dim: 10502 10
exptData(0):
assays(2): totalReads methReads
rownames(10502): 1456 1457 ... 4970981 4970982
```

```
rowData metadata column names(0):
colnames(10): APL1 APL2 ... APL11624 APL5894
colData names(1): group
```

We show the sample characteristics slot:

> colData(rrbs)

DataFrame with 10 rows and 1 column group <factor> APL1 APL APL2 APL APL APL3 APL7 APL APL8 APL APL10961 control APL11436 control APL11523 control APL11624 control APL5894 control

The first CpG sites on chromosome 1 which were covered:

```
> head(rowData(rrbs))
```

GRanges ob	ject with	6 ranges	and	0 metadata	columns	:	
seq	names	ra	nges	strand			
	<rle></rle>	<iran< td=""><td>ges></td><td><rle></rle></td><td></td><td></td><td></td></iran<>	ges>	<rle></rle>			
1456	chr1 [87	0425, 870	425]	+			
1457	chr1 [87	0443, 870	443]	+			
1458	chr1 [87	0459, 870	459]	+			
1459	chr1 [87	0573, 870	573]	+			
1460	chr1 [87	0584, 870	584]	+			
1461	chr1 [87	0599, 870	599]	+			
seqinfo:	25 seque	nces from	an u	inspecified	genome;	no	se

eqlengths led genome; 1IO: Zo seyi sp ecii q.

The coverage of the first CpG sites per sample:

> he	ad(to	talRe	ads(r	rbs))				
	APL1	APL2	APL3	APL7	APL8	APL10961	APL11436	APL11523
1456	39	6	10	0	0	48	31	65
1457	39	6	10	0	0	48	31	65
1458	39	6	10	0	0	48	27	65
1459	20	26	49	48	39	27	23	34
1460	20	26	49	48	39	27	23	34
1461	20	26	49	48	39	27	22	34
	APL1:	1624 <i>I</i>	APL589	94				
1456		39		29				
1457		39		29				
1458		39		29				
1459		29	-	15				
1460		28	-	15				
1461		29	-	15				

The number of methylated reads of the first CpG sites per sample:

```
> head(methReads(rrbs))
```

	APL1	APL2	APL3	APL7	APL8	APL10961	APL11436	APL11523
1456	32	6	7	0	0	15	23	16
1457	33	6	7	0	0	18	10	19
1458	33	6	10	0	0	20	10	19
1459	13	20	34	41	32	3	8	8
1460	14	18	35	37	33	2	4	4
1461	14	16	35	40	31	5	5	5
	APL1:	1624 <i>I</i>	APL589	94				
1456		7		7				
1457		2		7				
1458		2		3				
1459		6		4				
1460		3		0				
1461		1		2				

2.3.2 The BSrel class

The *BSrel* is a container for 'relative' methylation levels of BS data. It comprises sample information together with CpG positions and the relative methylation values (between 0 and 1).

- A *BSrel* object consists of four slots:
- 1. A *SimpleList* of arbitrary content describing the overall experiment, accessible with exptData.
- 2. A *GRanges* of the positions of CpG-sites covered by BS in at least one sample, accessible with rowData.
- 3. A *DataFrame* of samples and the values of variables measured on those samples, accessible with colData.
- 4. An assays slot containing a *SimpleList* of a matrix with the relative methylation levels (between 0 and 1), accessible with methLevel.

A new *BSraw* object can be created by:

```
> methLevel <- matrix(c(rep(0.5, 3), rep(1, 3)), ncol = 2)
> BSrel(exptData = exptData,
            rowData = rowData,
            colData = colData,
            methLevel = methLevel)
```

We can convert a *BSraw* object to a *BSrel* object easily:

```
> rrbs.rel <- rawToRel(rrbs)
> rrbs.rel
class: BSrel
dim: 10502 10
exptData(0):
assays(1): methLevel
rownames(10502): 1456 1457 ... 4970981 4970982
rowData metadata column names(0):
colnames(10): APL1 APL2 ... APL11624 APL5894
colData names(1): group
```

The relative methylation values of the first CpG sites:

```
> head(methLevel(rrbs.rel))
```

	APL1	APL2	APL3	APL7	APL8
1456	0.8205128	1.0000000	0.7000000	NaN	NaN
1457	0.8461538	1.0000000	0.7000000	NaN	NaN
1458	0.8461538	1.0000000	1.0000000	NaN	NaN
1459	0.6500000	0.7692308	0.6938776	0.8541667 0	.8205128
1460	0.7000000	0.6923077	0.7142857	0.7708333 0	.8461538
1461	0.7000000	0.6153846	0.7142857	0.8333333 0	.7948718
	APL10961	APL11436	6 APL11523	3 APL11624	APL5894
1456	0.31250000	0.7419355	5 0.2461538	3 0.17948718	0.2413793
1457	0.37500000	0.3225806	6 0.2923077	0.05128205	0.2413793
1458	0.41666667	0.3703704	l 0.2923077	0.05128205	0.1034483
1459	0.11111111	0.3478261	0.2352941	0.20689655	0.2666667
1460	0.07407407	0.1739130	0.1176471	0.10714286	0.0000000
1461	0.18518519	0.2272727	0.1470588	3 0.03448276	0.1333333

2.4 Data handling

All methods for *SummarizedExperiment* objects are applicable for *BSraw* and *BSrel* objects:

> dim(rrbs)

[1] 10502 10

> colnames(rrbs)

[1] "APL1" "APL2" "APL3" "APL7" "APL8" [6] "APL10961" "APL11436" "APL11523" "APL11624" "APL5894"

We can return subsets of samples or CpG sites:

```
> rrbs[,"APL2"]
> ind.chr1 <- which(seqnames(rrbs) == "chr1")
> rrbs[ind.chr1,]
```

We can also subset by overlaps with a *GRanges* object:

```
> findOverlaps(rrbs, region)
> subsetByOverlaps(rrbs, region)
```

We can sort BSraw and BSrel objects into ascending order of CpG sites positions on chromosomes:

> sort(rrbs)

BSraw and BSrel objects can be combined and splitted:

```
> combine(rrbs[1:10,1:2], rrbs[1:1000, 3:10])
```

```
> split(rowData(rrbs),
```

f = as.factor(as.character(seqnames(rrbs))))

3 Quality control

Via two very simple methods it is possible to compare the sample's coverages. covStatistics lists the number of CpG sites that were covered per sample together with the median of the coverage of these CpG sites. covBoxplots represent the coverage distributions per sample.

```
> covStatistics(rrbs)
```

\$Covered	_CpG_sites	5			
APL1	APL2	APL3	APL7	APL8	APL10961
5217	4240	4276	3972	3821	5089
APL11436	APL11523	APL11624	APL5894		
5169	6922	6483	7199		
\$Median_0	coverage				
APL1	APL2	APL3	APL7	APL8	APL10961
12	5	12	15	11	10
APL11436	APL11523	APL11624	APL5894		
6	8	4	5		





Figure 1: Sample wise coverage distributions

4 Detection of DMRs within groups of samples

The algorithm to detect differentially methylated regions (DMRs) within two groups of samples (e.g. cancer and control) is described in detail in [1]. To better understand this User's guide it is helpful to know the rough procedure. The DMR detection is a five-steps approach:

- 1. Definition of CpG clusters
- 2. Smooth methylation data within CpG clusters
- 3. Model and test group effect for each CpG site within CpG clusters
- 4. Apply hierarchical testing procedure:
 - (a) Test CpG clusters for differential methylation and control weighted FDR on cluster
 - (b) Trim rejected CpG clusters and control FDR on single CpGs
- 5. Define DMR boundaries

Please see [1] for more details.

4.1 Definition of CpG clusters

In order to smooth the methylation data we first have to detect CpG clusters (regions with a high spatial density of covered CpG sites). Within a *BSraw* object clusterSites searches for agglomerations of CpG sites across all samples. In a first step the data is reduced to CpG sites covered in round(perc.samples*ncol(object)) samples (here: 4 samples), these are called 'frequently covered CpG sites'. In a second step regions are detected where not less than min.sites frequently covered CpG sites are sufficiantly close to each other (max.dist. Note, that the frequently covered CpG sites are sufficiently sequent analysis the methylation data of all CpG sites within these clusters are used.

We perform the analysis on a subset of our data to save time:

rrbs.clust.unlim is a again *BSraw* object but restricted to CpG sites within CpG clusters. Each CpG site is assigned to a cluster:

```
> head(rowData(rrbs.clust.unlim))
```

GRanges object wi	th 6 ranges and 1	metadata column:
seqnames	ranges	<pre>strand cluster.id</pre>
<rle></rle>	<iranges></iranges>	<rle> <character></character></rle>
1513 chr1	[872335, 872335]	* chr1_1
1514 chr1	[872369, 872369]	* chr1_1
401911 chr1	[872370, 872370]	* chr1_1
1515 chr1	[872385, 872385]	* chr1_1
401912 chr1	[872386, 872386]	* chr1_1
1516 chr1	[872412, 872412]	* chr1_1

seqinfo: 25 sequences from an unspecified genome; no seqlengths

The underlying CpG clusters can also be converted to a *GRanges* object with the start and end positions:

> clusterSitesToGR(rrbs.clust.unlim)

GRanges	object	with 6 ra	anges and	d 1 meta	ada	ta column:
Se	eqnames		ranges	strand		cluster.id
	<rle></rle>	<]	[Ranges>	<rle></rle>		<factor></factor>
[1]	chr1	[872335,	872616]	*		chr1_1
[2]	chr1	[875227,	875470]	*		chr1_2
[3]	chr1	[875650,	876028]	*		chr1_3
[4]	chr1	[876807,	877458]	*		chr1_4
[5]	chr1	[877684,	877932]	*		chr1_5
[6]	chr2	[45843,	46937]	*		chr2_1

seqinfo: 25 sequences from an unspecified genome; no seqlengths

4.2 Smooth methylation data

In the smoothing step CpG sites with high coverages get high weights. To reduce bias due to unusually high coverages we limit the coverage, e.g. to the 90% quantile:

```
> ind.cov <- totalReads(rrbs.clust.unlim) > 0
> quant <- quantile(totalReads(rrbs.clust.unlim)[ind.cov], 0.9)
> quant
90%
32
> rrbs.clust.lim <- limitCov(rrbs.clust.unlim, maxCov = quant)</pre>
```

We then smooth the methylation values of CpG sites within the clusters with the default bandwidth h = 80 base pairs. It is possible - and recommended - to parallelize this step by setting mc.cores, to 6 cores for instance, if there are 6 available.

```
> predictedMeth <- predictMeth(object = rrbs.clust.lim)</pre>
```

predictedMeth is a *BSrel* object with smoothed relative methylation levels for each CpG site within CpG clusters:

```
> predictedMeth
```

```
class: BSrel
dim: 344 10
exptData(0):
assays(1): methLevel
rownames(344): 1 2 ... 343 344
rowData metadata column names(1): cluster.id
colnames(10): APL1 APL2 ... APL11624 APL5894
colData names(1): group
```

The effect of the smoothing step can be shown with the plotMeth function:

> covBoxplots(rrbs.clust.lim, col = "cornflowerblue", las = 2)



Figure 2: Sample wise coverage distributions after coverage limitation

```
> plotMeth(object.raw = rrbs[,6],
        object.rel = predictedMeth[,6],
        region = region,
        lwd.lines = 2,
        col.points = "blue",
        cex = 1.5)
```



APL10961

Figure 3: Raw data together with smoothed methylation levels

4.3 Model and test group effect

We observe a differential methylation between cancer and control for some CpG sites:



Figure 4: Smoothed methylation levels in APL and control samples

To detect the CpG sites where the DNA methylation differs between APL and control samples we model the methylation within a beta regression with the group as explanatory variable and use the Wald test to test if there is a group effect:

> ## OR:

> data(betaResults)

betaResults is a *data.frame* containing model and test information for each CpG site:

> head(betaResults)

ch	r pos	p.v	val meth.g	roup1 meth.group2
chr1.1 chr	1 872335	0.00113176	652 0.95	25098 0.8635983
chr1.2 chr	1 872369	0.00076780	0.94	14368 0.8444060
chr1.3 chr	1 872370	0.00083474	451 0.94	14314 0.8448725
chr1.4 chr	1 872385	0.00103374	477 0.94	12217 0.8521862
chr1.5 chr	1 872386	0.0010975	571 0.94	10544 0.8526863
chr1.6 chr	1 872412	0.00351148	339 0.93	78250 0.8718024
me	th.diff	estimate	std.error	pseudo.R.sqrt
chr1.1 0.08	3891149	-0.5730618	0.1760266	0.6304051
chr1.2 0.09	9703074	-0.5542171	0.1647422	0.6291904
chr1.3 0.09	9655890	-0.5522164	0.1652843	0.6246474
chr1.4 0.08	3903549	-0.5192564	0.1582531	0.6108452
chr1.5 0.08	3836810	-0.5156622	0.1579728	0.6090794
chr1.6 0.00	6602261	-0.4018161	0.1376551	0.5851744
clus	ster.id			
chr1.1	chr1_1			
chr1.2	chr1_1			
chr1.3	chr1_1			
chr1.4	chr1_1			
chr1.5	chr1_1			
chr1.6	chr1_1			

By setting type = "BR" the maximum likelihood with bias reduction is called. This is especially useful, when the sample size is small, see [4]. The

mean of the response (methylation) is linked to a linear predictor described by $\tilde{x} + x^2$ using a link function while the precision parameter is assumed to be constant. Sometimes the variance of DNA methylation is dependent on the group factor, e.g. the methylation variance in cancer samples is often higher than in normal samples. These additional regressors can be linked to the precision parameter within the formula of type $\tilde{x} + x^2 + y^2 + y^2$ where the regressors in the two parts can be overlapping, see the documentation in the *betareg* package.

4.4 Test CpG clusters for differential methylation

The aim is to detect CpG clusters containing at least one differentially methylated location. To do so the P values p from the Wald tests are transformed to Z scores: $z = \Phi^{-1}(1-p)$, which are normally distributed under Null hypothesis (no group effect). As cluster test statistic a standardized Z score average is used. To estimate the standard deviation of the Z scores we have to estimate the correlation and hence the variogram of methylation between two CpG sites within a cluster. The estimation of the standard deviation requires that the distribution of the Z scores follows a standard normal distribution. However, if methylation in both groups differs for many CpG sites the density distribution of P values shows a peak near 0. To ensure that the P values are roughly uniformly distributed to get a variance of the Z scores that is Gaussian with variance 1 we recommend to estimate the variogram (and hence the correlation of Z scores) under the null hypothesis. To do so we model the beta regression again for resampled data:

```
> ## Both resampled groups should have the same number of
```

```
> ## cancer and control samples:
```

```
> predictedMethNull <- predictedMeth[,c(1:4, 6:9)]</pre>
```

```
> colData(predictedMethNull)$group.null <- rep(c(1,2), 4)</pre>
```

```
> ## To shorten the run time, please set mc.cores, if possible!
```

```
> betaResultsNull <- betaRegression(formula = ~group.null,</pre>
```

```
link = "probit",
object = predictedMethNull,
```

type="BR")

> ## OR:
> data(betaResultsNull)

We estimate the variogram for the Z scores obtained for the resampled data:

```
> vario <- makeVariogram(betaResultsNull)
> ## OR:
> data(vario)
```

Based on the variogram plot we evaluate the sill (usually near 1) of the variogram and smooth the curve:



Figure 5: Estimated variogram together with the smoothed curve

The vario.sm object is a list of two:

```
> names(vario.sm)
```

```
[1] "variogram" "pValsList"
```

> head(vario.sm\$variogram)

hv.sm110.000000000220.000000000330.000000000440.0004532616550.0019789519660.0038612942

> head(vario.sm\$pValsList[[1]])

	chr	pos	p.val n	meth.group1	meth.group2
chr1.1	chr1	872335	0.8567258	0.9267336	0.9191601
chr1.2	chr1	872369	0.7910594	0.9180561	0.9056264
chr1.3	chr1	872370	0.7855283	0.9187319	0.9060236
chr1.4	chr1	872385	0.7462536	0.9225722	0.9080750
chr1.5	chr1	872386	0.7462667	0.9225955	0.9081423
chr1.6	chr1	872412	0.8575813	0.9165966	0.9093737
	met	th.diff	estimate	e std.error	pseudo.R.sqrt
chr1.1	0.00	7573505	-0.05244358	8 0.2904759	0.006696118
chr1.2	0.012	2429670	-0.07781982	2 0.2937315	0.013223677
chr1.3	0.012	2708263	-0.07993265	5 0.2937384	0.013863179
chr1.4	0.014	4497213	-0.09359433	1 0.2892434	0.020416968
chr1.5	0.014	4453228	-0.09334700	0.2884945	0.020484807
chr1.6	0.00	7222875	-0.04562887	7 0.2542651	0.006457349
	clust	ter.id	z.score p	pos.new	
chr1.1	(chr1_1 -	-1.0657239	1	
chr1.2	(chr1_1 -	-0.8101026	35	
chr1.3	(chr1_1 -	-0.7910010	36	
chr1.4	(chr1_1 -	-0.6627467	51	
chr1.5	(chr1_1 -	-0.6627876	52	
chr1.6	(chr1_1 -	-1.0695155	78	

We replace the **pValsList** object (which consists of the test results of the resampled data) by the test results of interest (for group effect):

```
> ## auxiliary object to get the pValsList for the test
```

```
> ## results of interest:
```

```
> vario.aux <- makeVariogram(betaResults, make.variogram=FALSE)</pre>
```

```
> vario.sm$pValsList <- vario.aux$pValsList</pre>
```

```
> head(vario.sm$pValsList[[1]])
```

chr	pos	p.v	val meth.g	roup1 meth.group2
chr1.1 chr1	872335 (0.00113176	352 0.95 ⁴	25098 0.8635983
chr1.2 chr1	872369 (0.00076780	0.94	14368 0.8444060
chr1.3 chr1	872370 (0.00083474	151 0.94	14314 0.8448725
chr1.4 chr1	872385 (0.00103374	177 0.94	12217 0.8521862
chr1.5 chr1	872386 (0.00109755	571 0.94	10544 0.8526863
chr1.6 chr1	872412 (0.00351148	339 0.93 [.]	78250 0.8718024
meth	.diff	estimate	std.error	pseudo.R.sqrt
chr1.1 0.088	91149 -0	0.5730618	0.1760266	0.6304051
chr1.2 0.097	03074 -0	0.5542171	0.1647422	0.6291904
chr1.3 0.096	55890 -0	0.5522164	0.1652843	0.6246474
chr1.4 0.089	03549 -0	0.5192564	0.1582531	0.6108452
chr1.5 0.088	36810 -0	0.5156622	0.1579728	0.6090794
chr1.6 0.066	02261 -0	0.4018161	0.1376551	0.5851744
clust	er.id z	z.score po	os.new	
chr1.1 c	hr1_1 3.	.053282	1	
chr1.2 c	hr1_1 3.	.167869	35	
chr1.3 c	hr1_1 3.	. 143485	36	
chr1.4 c	hr1_1 3.	.080361	51	
chr1.5 c	hr1_1 3.	.062480	52	
chr1.6 c	hr1_1 2.	.695753	78	

vario.sm now contains the smoothed variogram under the Null hypothesis together with the P values (and Z scores) from the Wald test, that the group has no effect on methylation. The correlation of the Z scores between two locations in a cluster can now be estimated:

> locCor <- estLocCor(vario.sm)</pre>

We test each CpG cluster for the presence of at least one differentially methylated location at q what can be interpreted as the size-weighted FDR on clusters:

```
> clusters.rej <- testClusters(locCor,</pre>
                                FDR.cluster = 0.1)
3 CpG clusters rejected.
> clusters.rej$clusters.reject
GRanges object with 3 ranges and 2 metadata columns:
      segnames
                          ranges strand |
                                                 value
         <Rle>
                                  <Rle> | <character>
                       <IRanges>
  [1]
          chr1 [872335, 872616]
                                       * |
                                                chr1_1
  [2]
          chr1 [875227, 875470]
                                       * |
                                                chr1_2
  [3]
          chr2 [ 45843, 46937]
                                       * |
                                                chr2_1
           p.value
         <numeric>
  [1] 6.867163e-03
  [2] 4.295184e-07
  [3] 1.881082e-09
  _____
```

seqinfo: 2 sequences from an unspecified genome; no seqlengths

4.5 Trim significant CpG clusters

We then trim the rejected CpG clusters that is to remove the not differentially methylated CpG sites at q_1 what can be interpreted as the location-wise FDR:

```
chr
                                  p.val meth.group1
                       pos
                                          0.9525098
chr1_1.chr1.1 chr1 872335 0.0011317652
chr1_1.chr1.2 chr1 872369 0.0007678027
                                          0.9414368
chr1_1.chr1.3 chr1 872370 0.0008347451
                                          0.9414314
chr1_1.chr1.4 chr1 872385 0.0010337477
                                          0.9412217
chr1_1.chr1.5 chr1 872386 0.0010975571
                                          0.9410544
chr1_2.chr1.21 chr1 875227 0.0003916052
                                          0.7175829
               meth.group2 meth.diff
                                        estimate std.error
                 0.8635983 0.08891149 -0.5730618 0.1760266
chr1_1.chr1.1
```

```
chr1_1.chr1.2
                 0.8444060 0.09703074 -0.5542171 0.1647422
                 0.8448725 0.09655890 -0.5522164 0.1652843
chr1_1.chr1.3
chr1_1.chr1.4
                 0.8521862 0.08903549 -0.5192564 0.1582531
                 0.8526863 0.08836810 -0.5156622 0.1579728
chr1_1.chr1.5
                 0.3648251 0.35275781 -0.9212669 0.2598282
chr1_2.chr1.21
               pseudo.R.sqrt cluster.id z.score pos.new
chr1_1.chr1.1
                   0.6304051
                                  chr1_1 3.053282
                                                         1
chr1_1.chr1.2
                                  chr1_1 3.167869
                   0.6291904
                                                        35
chr1_1.chr1.3
                                  chr1_1 3.143485
                   0.6246474
                                                        36
chr1_1.chr1.4
                   0.6108452
                                  chr1_1 3.080361
                                                        51
                                  chr1_1 3.062480
                                                        52
chr1_1.chr1.5
                   0.6090794
chr1_2.chr1.21
                   0.6818046
                                  chr1_2 3.358661
                                                         1
                      p.li
chr1_1.chr1.1
               0.019767638
chr1_1.chr1.2
               0.019953150
chr1_1.chr1.3
               0.021679300
chr1_1.chr1.4
               0.028817819
chr1_1.chr1.5
               0.030594942
chr1_2.chr1.21 0.004852623
```

clusters.trimmed is a *data.frame* object containing all differentially methylated CpG sites. The p.li column contains the P values estimated in the cluster trimming step, see [1].

4.6 Definition of DMR boundaries

We can now define the boundaries of DMRs as rejected CpG sites within which rejected CpG sites solely are located. Within the DMRs the distance between neighbored rejected CpG sites should not exceed max.dist base pairs (usually the same as for max.dist in clusterSites), otherwise, the DMR is splitted. DMRs are also splitted if the methylation difference switches from positive to negative, or vice versa, if diff.dir = TRUE. That way we ensure that within a DMR all CpG sites are hypermethylated, and hypomethylated respectively.

```
> DMRs <- findDMRs(clusters.trimmed,</pre>
                    max.dist = 100,
                    diff.dir = TRUE)
```

> DMRs

GRanges object with 4 ranges and 4 metadata columns: seqnames ranges strand | <Rle> <Rle> | <IRanges> [1] chr1 [872335, 872386] * | [2] chr1 [875227, 875470] * | [3] chr2 [46126, 46718] * | [4] chr2 [46915, 46937] * | median.p median.meth.group1 <numeric> <numeric> [1] 0.0010337476937105 0.94143135870377 [2] 6.67719258580069e-06 0.502443544206678 [3] 3.9604046820675e-05 0.438629367812053 [4] 0.0148459621327497 0.13636374336524 median.meth.group2 median.meth.diff <numeric> <numeric> [1] 0.852186207117906 0.0890354930388882 [2] 0.182082007625853 0.319474180010203 [3] 0.0776818250851119 0.355363178661131 [4] 0.0369197526986237 0.099443990666616 _____

seqinfo: 2 sequences from an unspecified genome; no seqlengths

5 Detection of DMRs between two samples

If there are two samples only to be compared we can use the compareTwoSamples function which determines the differences per CpG site and aggregates the sites surpassing the minimum difference minDiff:

Some of the DMRs detected within these two samples overlap with the groupwise DMRs:

```
> sum(overlapsAny(DMRs.2,DMRs))
```

[1] 1

6 Further data processing

The plotMethMap function is helpful to evaluate DMRs graphically. Via zlim = c(0,1) that is passed to the heatmap function we ensure that green stands for a relative methalytion of 0 and red stands for a relative methylation of 1:

```
> rowCols <- c("magenta", "blue")[as.numeric(colData(predictedMeth)$group)]
> plotMethMap(predictedMeth,
            region = DMRs[3],
            groups = colData(predictedMeth)$group,
            intervals = FALSE,
            zlim = c(0,1),
            RowSideColors = rowCols,
            labCol = "", margins = c(0, 6))
```





To represent the smoothed methylation curves we can use the plotSmooth-

Meth function:



Figure 7: The smoothed methylation curves for all samples within a detected DMR

We can annotate the detected DMRs by means of a *GRanges* object, e.g. a list of promoter regions. In case of an overlapping of both *GRanges* objects the DMR is marked as **TRUE**, or with the respective identifier in the promoter list:

```
> data(promoters)
```

```
> head(promoters)
```

GRanges object with 6 ranges and 1 metadata column: seqnames ranges strand | acc_no <Rle> <IRanges> <Rle> | <character> chr1 [66998824, 67000324] [1] * NM_032291 [2] chr1 [8383389, 8384889] * | NM_001080397 [3] chr1 [16766166, 16767666] | NM_001145277 * chr1 [16766166, 16767666] [4] * | NM_001145278 [5] chr1 [16766166, 16767666] * | NM_018090 [6] chr1 [50489126, 50490626] * | NM_032785 seqinfo: 24 sequences from an unspecified genome; no seqlengths > DMRs.anno <- annotateGRanges(object = DMRs,</pre> regions = promoters, name = 'Promoter', regionInfo = 'acc_no') > DMRs.anno GRanges object with 4 ranges and 5 metadata columns: seqnames ranges strand | median.p <Rle> <IRanges> <Rle> | <numeric> [1] chr1 [872335, 872386] * | 1.033748e-03 [2] chr1 [875227, 875470] * | 6.677193e-06 chr2 [46126, [3] 46718] * | 3.960405e-05 [4] chr2 [46915, 46937] * | 1.484596e-02 median.meth.group1 median.meth.group2 <numeric> <numeric> [1] 0.9414314 0.85218621 [2] 0.5024435 0.18208201 [3] 0.4386294 0.07768183 [4] 0.1363637 0.03691975

	median.meth.diff	Promoter	
	<numeric></numeric>	<character></character>	
[1]	0.08903549	<na></na>	
[2]	0.31947418	<na></na>	
[3]	0.35536318	NM_001077710	
[4]	0.09944399	NM_001077710	
		_	

seqinfo: 2 sequences from an unspecified genome; no seqlengths

plotBindingSites plots the average methylation around given genomic regions, e.g. protein binding sites. Here, we compare the methyation in and around promoter regions between APL and controls:



Figure 8: Methylation around 1,000 promoters; Position 0 refers to the centers of the promoters

The raw and relative methylation data can also be viewed in the Integra-

tive Genomics Viewer (IGV; freely available for download from www.broadinstitute.org/igv) [5]. To do so we first write the methylation information of each sample within the *BSraw* or *BSrel* object to a bed file:

We can load the bed files of the raw data in the IGV. The integers beneath the CpG marks represent the numbers of sequencing reads covering the CpG sites:

	NoME DATA TYPE	hr2 p25.1 p24.1 p22.3 p21 p16.1 p13.3 p11.2 q11.2 q13 q14.3 q22.1 q23.3 q24.3 q31.2 q32.3 q33.3 q35 q37.1 4,682 bp bp 44,000 bp 45,000 bp 45,000 bp 46,000 bp 47,000 bp	- 4 -
RefSeq Genes	28		
APL_1		I IIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIII	
APL_2		I I	
APL_3		I I	
APL_7		1 1 1 1 2 5 3 7 7 12 10 1 2 14 17 3 1 1	
APL_8		I I	
control_10961		I IIIIIIII IIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIII	
control_11436		I III IIII IIIIIIII IIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIII	
control_11523		II III IIIIII IIIIIII IIIIIIII IIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIIII	
control_11624		I II III III.00011000000000000000000000000000000	
control_5894			

Figure 9: IGV snapshot of the raw data in and around a detected DMR

We can also load the smoothed methylation levels:

	Nome Data TrPE	chr2 p25.1 p24.1 p22.3 p21 p16.1 p13.3 p11.2 q13 q14.3 q22.1 q23.3 q24.3 q31.2 q32.3 q33.3 q35 q37.1
RefSeq Genes		
APL_1		
APL_2		
APL_3		
APL_7		
APL_8		
control_10961		
control_11436		
control_11523		
control_11624		
control_5894		

Figure 10: IGV snapshot of the smoothed data in and around a detected $\rm DMR$

References

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- [2] Till Schoofs, Christian Rohde, Katja Hebestreit, Hans-Ulrich Klein, Stefanie Göllner, Isabell Schulze, Mads Lerdrup, Nikolaj Dietrich, Shuchi Agrawal-Singh, Anika Witten, Monika Stoll, Eva Lengfelder, Wolf-Karsten Hofmann, Peter Schlenke, Thomas Büchner, Klaus Hansen, Wolfgang E Berdel, Frank Rosenbauer, Martin Dugas, and Carsten Müller-Tidow. Dna methylation changes are a late event in acute promyelocytic leukemia and coincide with loss of transcription factor binding. *Blood*, Nov 2012. URL: http://dx.doi.org/10.1182/ blood-2012-08-448860, doi:10.1182/blood-2012-08-448860.
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