Differential analyses with DSS

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Abstract

This vignette introduces the use of the Bioconductor package DSS (\underline{D} is persion \underline{S} hrinkage for \underline{S} equencing data), which is designed for differential analysis based on high-throughput sequencing data. It performs differential expression analyses for RNA-seq, and differential

methylation analyses for bisulfite sequencing (BS-seq) data. The core of DSS is a procedure based on Bayesian hierarchical model to estimate and shrink gene- or CpG site-specific dispersions, then conduct Wald tests for detecting differential expression/methylation.

1 Introduction

Recent advances in various high-throughput sequencing technologies have revolutionized genomics research. Among them, RNA-seq is designed to measure the the abundance of RNA products, and Bisulfite sequencing (BS-seq) is for measuring DNA methylation. A fundamental question in functional genomics research is whether gene expression or DNA methylation vary under different biological contexts. Thus, identifying differential expression genes (DEGs) or differential methylation loci/regions (DML/DMRs) are key tasks in RNA-seq or BS-seq data analyses.

The differential expression (DE) or differential methylation (DM) analyses are often based on gene- or CpG-specific statistical test. A key limitation in RNA- or BS-seq experiments is that the number of biological replicates is usually limited due to cost constraints. This can lead to unstable estimation of within group variance, and subsequently undesirable results from hypothesis testing. Variance shrinkage methods have been widely applied in DE analyses in microarray data to improve the estimation of gene-specific within group variances. These methods are typically based on a Bayesian hierarchical model, with a prior imposed on the gene-specific variances to provide a basis for information sharing across all genes.

A distinct feature of RNA-seq or BS-seq data is that the measurements are in the form of counts and have to be modeld by discrete distributions. Unlike continuous distributions (such as Gaussian), the variances depend on means in these discrete distributions. This implies that the sample variances do not account for biological variation, and shrinkage cannot be applied on variances directly. In DSS, we assume that the count data are from the Gamma-Poisson (for RNA-seq) or Beta-Binomial (for BS-seq) distribution. We then parameterize the distributions by a mean and a dispersion parameters. The dispersion parameters, which represent the biological variation for replicates within a treatment group, play a central role in the differential analyses.

DSS implements a series of DE/DM detection algorithms based on the dispersion shrinkage method followed by Wald statistical test to test each gene/CpG site for differential expression/methylation. It provides functions for RNA-seq DE analysis for both two group comparision and multi-factor design, BS-seq DM analysis for two group comparision, multifactor design, and data without biological replicate. Simulation and real data results show that the methods provides excellent performance compared to existing methods, especially when the overall dispersion level is high or the number of replicates is small. For more details of the data model, the shrinkage method, and test procedures, please read [4] for differential expression from RNA-seq, [1] for differential methylation for two-group comparison from BS-seq, [2] for differential methylation for data without biological replicate, and [3] for differential methylation for general experimental design.

2 Using DSS for differential expression analysis

2.1 Input data preparation

DSS requires a count table (a matrix of **integers**) for gene expression values (rows are for genes and columns are for samples). This is different from the isoform expression based analysis such as in cufflink/cuffdiff, where the gene expressions are represented as non-integers values. There are a number of ways to obtain the count table from raw sequencing data (fastq file), here we provide some example codes using several Bioconductor packages (the codes require installation of GenomicFeatures, Rsamtools, and GenomicRanges packages).

- 1. Sequence alignment. There are several RNA-seq aligner, for example, tophat or STAR. Assume the alignment result is saved in a BAM file data.bam.
- Choose a gene annotation. GenomicFeatures package provides a convenient way to access different gene annotations. For example, if one wants to use RefSeq annotation for human genome build hg19, one can use following codes:
 - > library(GenomicFeatures)
 - > txdb = makeTranscriptDbFromUCSC(genom="hg19",tablename="refGene")
 - > genes = genes(txdb)
- 3. Obtain count table based on the alignment results and gene annotation. This can be done in several steps. First read in the BAM file using the Rsamtools package:

> bam=scanBam("data.bam")

Next, create GRanges object for the aligned sequence reads.

> IRange.reads=GRanges(seqnames=Rle(bam\$rname), ranges=IRanges(bam\$pos, width=bam\$qwidth))

Finally, use the countOverlaps function in GenomicRanges function to obtain the read counts overlap each gene.

> counts = countOverlaps(genes, IRange.reads)

There are other ways to obtain the counts, for example, using QuasR or easyRNASeq Bioconductor package. Please refer to the package vignettes for more details.

2.2 Single factor experiment

In single factor RNA-seq experiment, DSS also requires a vector representing experimental designs. The length of the design vector must match the number of columns of the count table. Optionally, normalization factors or additional annotation for genes can be supplied.

The basic data container in the package is SeqCountSet class, which is directly inherited from ExpressionSet class defined in Biobase. An object of the class contains all necessary information for a DE analysis: gene expression values, experimental designs, and additional annotations.

A typical DE analysis contains the following simple steps.

- 1. Create a SeqCountSet object using newSeqCountSet.
- 2. Estimate normalization factor using estNormFactors.
- 3. Estimate and shrink gene-wise dispersion using estDispersion
- 4. Two-group comparison using waldTest.

The usage of DSS is demonstrated in the simple simulation below.

1. First load in the library, and make a SeqCountSet object from some counts for 2000 genes and 6 samples.

```
> library(DSS)
> counts1=matrix(rnbinom(300, mu=10, size=10), ncol=3)
```

- > counts2=matrix(rnbinom(300, mu=50, size=10), ncol=3)
- > X1=cbind(counts1, counts2) ## these are 100 DE genes
- > X2=matrix(rnbinom(11400, mu=10, size=10), ncol=6)
- > X=rbind(X1,X2)

```
> designs=c(0,0,0,1,1,1)
```

```
> seqData=newSeqCountSet(X, designs)
```

```
> seqData
```

SeqCountSet (storageMode: lockedEnvironment)

```
assayData: 2000 features, 6 samples
```

```
element names: exprs
```

```
protocolData: none
```

phenoData

```
sampleNames: 1 2 ... 6 (6 total)
```

```
varLabels: designs
varMetadata: labelDescription
featureData: none
experimentData: use 'experimentData(object)'
Annotation:
```

- 2. Estimate normalization factor.
 - > seqData=estNormFactors(seqData)
- 3. Estimate and shrink gene-wise dispersions

```
> seqData=estDispersion(seqData)
```

4. With the normalization factors and dispersions ready, the two-group comparison can be conducted via a Wald test:

```
> result=waldTest(seqData, 0, 1)
> head(result,5)
                                     lfc
                                           difExpr
   geneIndex
                           muB
                                                        stats
                                                                      pval
                  muA
41
          41 6.000000 61.15152 -2.249696 -55.15152 -5.561982 2.667281e-08
          68 5.000000 54.56313 -2.303732 -49.56313 -5.522555 3.341043e-08
68
15
          15 7.000000 58.89141 -2.069247 -51.89141 -5.368886 7.922442e-08
37
          37 7.333333 61.80556 -2.073662 -54.47222 -5.367189 7.997328e-08
70
          70 8.6666667 63.34343 -1.940860 -54.67677 -5.288659 1.232165e-07
      local.fdr
                         fdr
41 0.0001162828 0.0001162828
68 0.0001174614 0.0001162828
15 0.0001900275 0.0001504715
37 0.0001910996 0.0001504715
70 0.0002433947 0.0001726463
```

A higher level wrapper function DSS.DE is provided for simple RNA-seq DE analysis in a two-group comparison. User only needs to provide a count matrix and a vector of 0's and 1's representing the design, and get DE test results in one line. A simple example is listed below:

- > counts = matrix(rpois(600, 10), ncol=6)
- > designs = c(0, 0, 0, 1, 1, 1)
- > result = DSS.DE(counts, designs)
- > head(result)

geneIndex muA muB lfc difExpr stats pval 61 61 13.000000 6.814815 0.6127880 6.185185 2.292107 0.02189948 57 57 8.000000 13.555556 -0.5029516 -5.555556 -1.966263 0.04926821 92 6.000000 10.555556 -0.5311309 -4.555556 -1.876002 0.06065494 92 36 8.333333 13.481481 -0.4592013 -5.148148 -1.825857 0.06787177 36 80 13.333333 8.259259 0.4569698 5.074074 1.802837 0.07141388 80 70 12.666667 7.777778 0.4641138 4.888889 1.786663 0.07399199 70 local.fdr fdr 61 1.0000000 1.0000000 57 0.5096204 0.9238904 92 0.5753564 0.8743174 36 0.6233741 0.8628584 80 1.0000000 0.8952432 70 1.0000000 0.8743174

2.3 Multifactor experiment

DSS provides functionalities for dispersion shrinkage for multifactor experimental designs. Downstream model fitting (through genearlized linear model) and hypothesis testing can be performed using other packages such as edgeR, with the dispersions estimated from DSS.

Below is an example, based a simple simulation, to illustrate the DE analysis of a crossed design.

- 1. First simulate data for a 2x2 crossed experiments. Note the counts are randomly generated.
 - > library(DSS)
 - > library(edgeR)
 - > counts=matrix(rpois(800, 10), ncol=8)
 - > design=data.frame(gender=c(rep("M",4), rep("F",4)), strain=rep(c("WT", "Mutant"),4))
 - > X=model.matrix(~gender+strain, data=design)
- 2. make SeqCountSet, then estimate size factors and dispersion
 - > seqData=newSeqCountSet(counts, as.data.frame(X))
 - > seqData=estNormFactors(seqData)
 - > seqData=estDispersion(seqData)
- 3. Using edgeR's function to do glm model fitting, but plugging in the estimated dispersion from DSS.

> fit.edgeR <- glmFit(counts, X, dispersion=dispersion(seqData))</pre>

4. Using edgeR's function to do hypothesis testing on the second parameter of the model (gender).

```
> lrt.edgeR <- glmLRT(glmfit=fit.edgeR, coef=2)
> head(lrt.edgeR$table)
logFC logCPM LR PValue
1 0.67231424 13.78314 3.469009873 0.06252873
2 0.02227149 13.39339 0.002439942 0.96060390
3 -0.25219670 13.45604 0.413084223 0.52040821
4 0.52292872 13.53637 1.815582183 0.17784004
5 0.30727501 13.54557 0.599958678 0.43859379
6 -0.01572223 13.57859 0.001615999 0.96793409
```

3 Using DSS for differential methylation analysis

3.1 Overview

To detect differential methylation, statistical tests are conducted at each CpG site, and then the differential methylation loci (DML) or differential methylation regions (DMR) are called based on user specified threshold. A rigorous statistical tests should account for biological variations among replicates and the sequencing depth. Most existing methods for DM analysis are based on *ad hoc* methods. For example, using Fisher's exact ignores the biological variations, using t-test on estimated methylation levels ignores the sequencing depth. Sometimes arbitrary filtering are implemented: loci with depth lower than an arbitrary threshold are filtered out, which results in information loss

The DM detection procedure implemented in DSS is based on a rigorous Wald test for betabinomial distributions. The test statistics depend on the biological variations (characterized by dispersion parameter) as well as the sequencing depth. An important part of the algorithm is the estimation of dispersion parameter, which is achieved through a shrinkage estimator based on a Bayesian hierarchical model [1]. An advantage of DSS is that the test can be performed even when there is no biological replicates. That's because by smoothing, the neighboring CpG sites can be viewed as "pseudo-replicates", and the dispersion can still be estimated with reasonable precision. DSS also works for general experimental design, based on a beta-binomial regression model with "arcsine" link function. Model fitting is performed on transformed data with generalized least square method, which achieves much improved computational performance compared with methods based on generalized linear model.

DSS depends on bsseq Bioconductor package, which has neat definition of data structures and many useful utility functions. In order to use the DM detection functionalities, bsseq needs to be pre-installed.

3.2 Input data preparation

DSS requires data from each BS-seq experiment to be summarized into following information for each CG position: chromosome number, genomic coordinate, total number of reads, and number of reads showing methylation. For a sample, this information are saved in a simple text file, with each row representing a CpG site. Below shows an example of a small part of such a file:

chr	pos	Ν	Х
chr18	3014904	26	2
chr18	3031032	33	12
chr18	3031044	33	13
chr18	3031065	48	24

One can follow below steps to obtain such data from raw sequence file (fastq file), using bismark (version 0.10.0, commands for newer versions could be different) for BS-seq alignment and count extraction. These steps require installation of bowtie or bowtie2, bismark, and the fasta file for reference genome.

- Prepare Bisulfite reference genome. This can be done using the bismark_genome_preparation function (details in bismark manual). Example command is: bismark_genome_preparation -path_to_bowtie /usr/local/bowtie/ -verbose /path/to/refgenomes/
- 2. BS-seq alignment. Example command is:

bismark -q -n 1 -l 50 -path_to_bowtie /path/bowtie/ BS-refGenome reads.fastq
This step will produce two text files reads.fastq_bismark.sam and reads.fastq_bismark_SE_report.txt.

3. Extract methylation counts using bismark_methylation_extractor function:

bismark_methylation_extractor -s -bedGraph reads.fastq_bismark.sam. This will create multiple txt files to summarize methylation call and cytosine context, a bedGraph file to display methylation percentage, and a coverage file containing counts information. The count file contain following columns:chr, start, end, methylation%, count methylated, count unmethylated. This file can be modified to make the input file for DSS. A typical DML detection contains two simple steps. First one conduct DM test at each CpG site, then DML/DMR are called based on the test result and user specified threshold.

3.3 DML/DMR detection from two-group comparison

Below are the steps to call DML or DMR for BS-seq data in two-group comparison setting.

 Load in library. Read in text files and create an object of BSseq class, which is defined in bsseq Bioconductor package. This step requires bsseq Bioconductor package. BSseq class is defined in that package.

```
> library(DSS)
> require(bsseq)
> path <- file.path(system.file(package="DSS"), "extdata")</pre>
> dat1.1 <- read.table(file.path(path, "cond1_1.txt"), header=TRUE)</pre>
> dat1.2 <- read.table(file.path(path, "cond1_2.txt"), header=TRUE)</pre>
> dat2.1 <- read.table(file.path(path, "cond2_1.txt"), header=TRUE)</pre>
> dat2.2 <- read.table(file.path(path, "cond2_2.txt"), header=TRUE)</pre>
> BSobj <- makeBSseqData( list(dat1.1, dat1.2, dat2.1, dat2.2),</pre>
       c("C1","C2", "N1", "N2") )[1:1000,]
+
> BSobj
An object of type 'BSseq' with
  1000 methylation loci
  4 samples
has not been smoothed
All assays are in-memory
```

2. Perform statistical test for DML by calling DMLtest function. This function basically performs following steps: (1) estimate mean methylation levels for all CpG site; (2) estimate dispersions at each CpG sites; (3) conduct Wald test. For the first step, there's an option for smoothing or not. Because the methylation levels show strong spatial correlations, smoothing can help obtain better estimates of mean methylation when the CpG sites are dense in the data (such as from the whole-genome BS-seq). However for data with sparse CpG, such as from RRBS or hydroxyl-methylation, smoothing is not recommended.

To perform DML test without smoothing, do:

> dmlTest <- DMLtest(BSobj, group1=c("C1", "C2"), group2=c("N1", "N2"))
Estimating dispersion for each CpG site, this will take a while ...</pre>

```
> head(dmlTest)
    chr
            pos
                      mu1
                                mu2
                                           diff
                                                   diff.se
                                                                 stat
1 chr18 3014904 0.3817233 0.4624549 -0.08073162 0.24997034 -0.3229648
2 chr18 3031032 0.3380579 0.1417008 0.19635711 0.11086362 1.7711592
3 chr18 3031044 0.3432172 0.3298853 0.01333190 0.12203116 0.1092500
4 chr18 3031065 0.4369377 0.3649218 0.07201587 0.10099395 0.7130711
5 chr18 3031069 0.2933572 0.5387464 -0.24538920 0.13178800 -1.8619996
6 chr18 3031082 0.3526311 0.3905718 -0.03794068 0.07847999 -0.4834440
         phi1
                    phi2
                               pval
                                          fdr
1 0.300542998 0.01706260 0.74672190 0.9985094
2 0.008911745 0.04783892 0.07653423 0.6792127
3 0.010409029 0.01994821 0.91300423 0.9985094
4 0.010320888 0.01603200 0.47580174 0.9985094
5 0.012537553 0.02320887 0.06260315 0.6158797
6 0.007665696 0.01145531 0.62878051 0.9985094
```

To perform statistical test for DML with smoothing, do:

```
> dmlTest.sm <- DMLtest(BSobj, group1=c("C1", "C2"), group2=c("N1", "N2"), smoothing=TRUE)
Smoothing ...
Estimating dispersion for each CpG site, this will take a while ...</pre>
```

User has the option to smooth the methylation levels or not. For WGBS data, smoothing is recommended so that information from nearby CpG sites can be combined to improve the estimation of methylation levels. A simple moving average algorithm is implemented for smoothing. In RRBS since the CpG coverage is sparse, smoothing might not alter the results much. If smoothing is requested, smoothing span is an important parameter which has non-trivial impact on DMR calling. We use 500 bp as default, and think that it performs well in real data tests.

3. With the test results, one can call DML by using callDML function. The results DMLs are sorted by the significance.

```
639 chr18 4431511 0.01327049 0.9430566 -0.9297862 0.09270080 -10.02997
710 chr18 4564237 0.91454619 0.0119300 0.9026162 0.05260037 17.15988
782 chr18 4657576 0.98257334 0.0678355 0.9147378 0.06815000 13.42242
           phi1
                      phi2
                                   pval
                                                 fdr postprob.overThreshold
450 0.052591567 0.02428826 1.029974e-45 2.499403e-43
                                                                           1
451 0.052591567 0.02428826 1.029974e-45 2.499403e-43
                                                                           1
638 0.053172411 0.07746835 1.177826e-23 1.429096e-21
                                                                           1
639 0.053121697 0.07746835 1.125518e-23 1.429096e-21
                                                                           1
710 0.009528898 0.04942849 5.302004e-66 3.859859e-63
                                                                           1
782 0.010424723 0.06755651 4.468885e-41 8.133371e-39
                                                                           1
```

By default, the test is based on the null hypothesis that the difference in methylation levels is 0. Alternatively, users can specify a threshold for difference. For example, to detect loci with difference greater than 0.1, do:

```
dmls2 <- callDML(dmlTest, delta=0.1, p.threshold=0.001)</pre>
>
    head(dmls2)
>
      chr
                                              diff
                                                      diff.se
              pos
                         mu1
                                    mu2
                                                                    stat
450 chr18 3976129 0.01027497 0.9390339 -0.9287590 0.06544340 -14.19179
451 chr18 3976138 0.01027497 0.9390339 -0.9287590 0.06544340 -14.19179
638 chr18 4431501 0.01331553 0.9430566 -0.9297411 0.09273779 -10.02548
639 chr18 4431511 0.01327049 0.9430566 -0.9297862 0.09270080 -10.02997
710 chr18 4564237 0.91454619 0.0119300 0.9026162 0.05260037 17.15988
782 chr18 4657576 0.98257334 0.0678355 0.9147378 0.06815000 13.42242
           phi1
                                                  fdr postprob.overThreshold
                      phi2
                                    pval
450 0.052591567 0.02428826 1.029974e-45 2.499403e-43
                                                                            1
451 0.052591567 0.02428826 1.029974e-45 2.499403e-43
                                                                            1
638 0.053172411 0.07746835 1.177826e-23 1.429096e-21
                                                                            1
639 0.053121697 0.07746835 1.125518e-23 1.429096e-21
                                                                            1
710 0.009528898 0.04942849 5.302004e-66 3.859859e-63
                                                                            1
782 0.010424723 0.06755651 4.468885e-41 8.133371e-39
                                                                            1
```

When delta is specified, the function will compute the posterior probability that the difference of the means is greater than delta. So technically speaking, the threshold for p-value here actually refers to the threshold for 1-posterior probability, or the local FDR. Here we use the same parameter name for the sake of the consistence of function syntax.

4. DMR detection is also Based on the DML test results, by calling callDMR function. Regions with many statistically significant CpG sites are identified as DMRs. Some restrictions are provided by users, including the minimum length, minimum number of CpG sites, percentage of CpG site being significant in the region, etc. There are some *post hoc* procedures to merge nearby DMRs into longer ones.

```
> dmrs <- callDMR(dmlTest, p.threshold=0.01)
> head(dmrs)
```

chr start end length nCG meanMethyl meanMethy2 diff.Methy areaStat 27 chr18 4657576 4657639 64 4 0.506453 0.318348 0.188105 14.34236

Here the DMRs are sorted by "areaStat", which is defined in bsseq as the sum of the test statistics of all CpG sites within the DMR.

Similarly, users can specify a threshold for difference. For example, to detect regions with difference greater than 0.1, do:

```
dmrs2 <- callDMR(dmlTest, delta=0.1, p.threshold=0.05)</pre>
>
    head(dmrs2)
>
     chr
           start
                      end length nCG meanMethy1 meanMethy2 diff.Methy areaStat
31 chr18 4657576 4657639
                              64
                                    4
                                       0.5064530 0.3183480
                                                               0.188105 14.34236
19 chr18 4222533 4222608
                              76
                                    4
                                      0.7880276 0.3614195
                                                               0.426608 12.91667
```

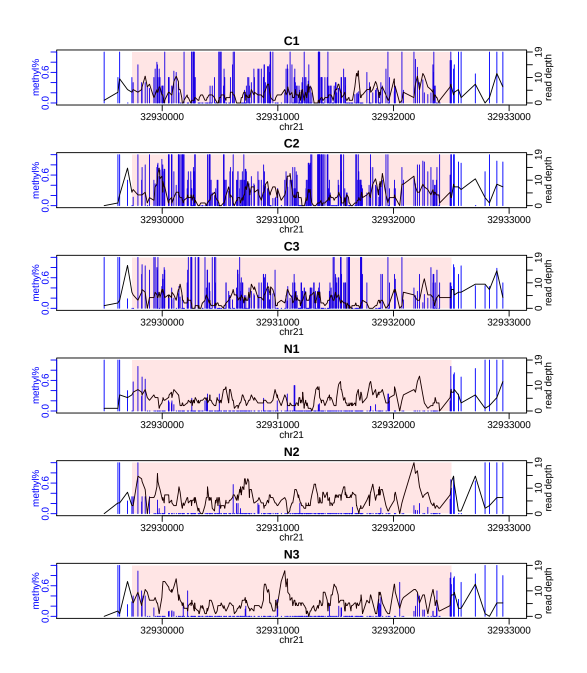
Note that the distribution of test statistics (and p-values) depends on the differences in methylation levels and biological variations, as well as technical factors such as coverage depth. It is very difficulty to select a natural and rigorous threshold for defining DMRs. We recommend users try different thresholds in order to obtain satisfactory results.

5. The DMRs can be visualized using showOneDMR function, This function provides more information than the plotRegion function in bsseq. It plots the methylation percentages as well as the coverage depths at each CpG sites, instead of just the smoothed curve. So the coverage depth information will be available in the figure.

To use the function, do

```
> showOneDMR(dmrs[1,], BSobj)
```

The result figure looks like the following. Note that the figure below is not generated from the above example. The example data are from RRBS experiment so the DMRs are much shorter.



3.4 DML/DMR detection from general experimental design

In DSS, BS-seq data from a general experimental design (such as crossed experiment, or experiment with covariates) is modeled through a generalized linear model framework. We use "arcsine" link function instead of the typical logit link for it better deals with data at boundaries (methylation levels close to 0 or 1). Linear model fitting is done through ordinary least square on transformed methylation levels. Variance/covariance matrices for the estimates are derived with consideration of count data distribution and transformation.

3.4.1 Hypothesis testing in general experimental design

In a general design, the data are modeled through a multiple regression thus there are multiple regression coefficients. In contrast, there is only one parameter in two-group comparison which is the difference between two groups. Under this type of design, hypothesis testing can be performed for one, multiple, or any linear combination of the parameters.

DSS provides flexible functionalities for hypothesis testing. User can test one parameter in the model through a Wald test, or any linear combination of the parameters through an F-test.

The DMLtest.multiFactor function provide interfaces for testing one parameter (through coef parameter), one term in the model (through term parameter), or linear combinations of the parameters (through Contrast parameter). We illustrate the usage of these parameters through a simple example below. Assume we have an experiment from three strains (A, B, C) and two sexes (M and F), each has 2 biological replicates (so there are 12 datasets in total).

```
> Strain = rep(c("A", "B", "C"), 4)
```

- > Sex = rep(c("M", "F"), each=6)
- > design = data.frame(Strain,Sex)
- > design

Strain Sex

1	Α	Μ
2	В	М
3	С	М
4	Α	М
5	В	М
6	С	М
7	Α	F
8	В	F

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9	С	F
10	А	F
11	В	F
12	С	F

To test the additive effect of Strain and Sex, a design formula is Strain+Sex, and the corresponding design matrix for the linear model is:

```
> X = model.matrix(~Strain+ Sex, design)
> X
   (Intercept) StrainB StrainC SexM
1
              1
                       0
                               0
                                    1
2
              1
                      1
                               0
                                    1
3
              1
                       0
                               1
                                    1
4
              1
                       0
                               0
                                    1
5
              1
                      1
                               0
                                    1
6
              1
                       0
                               1
                                    1
7
              1
                                    0
                       0
                               0
8
              1
                               0
                                    0
                      1
9
              1
                       0
                               1
                                    0
10
              1
                       0
                               0
                                    0
11
              1
                      1
                               0
                                    0
12
              1
                       0
                               1
                                    0
attr(,"assign")
[1] 0 1 1 2
attr(,"contrasts")
attr(,"contrasts")$Strain
[1] "contr.treatment"
attr(,"contrasts")$Sex
[1] "contr.treatment"
```

Under this design, we can do different tests using the DMLtest.multiFactor function:

If we want to test the sex effect, we can either specify coef=4, coef="SexM", or term="Sex". Notice that when using character for coef, the character must match the column name of the design matrix, cannot do coef="Sex". It is also important to note that using term="Sex" only tests a single paramter in the model because sex only has two levels.

- If we want to test the effect of Strain B versus Strain A (this is also testing a single parameter), we do coef=2 or coef="StrainB".
- If we want to test the whole Strain effect, it becomes a compound test because Strain has three levels. We do term="Strain", which tests StrainB and StrainC simultaneously. We can also make a Contrast matrix L as following. It's clear that testing $L^T \beta = 0$ is equivalent to testing StrainB=0 and StrainC=0.

```
> L = cbind(c(0,1,0,0),c(0,0,1,0))
> L
       [,1] [,2]
[1,] 0 0
[2,] 1 0
[3,] 0 1
[4,] 0 0
```

 One can perform more general test, for example, to test StrainB=StrainC, or that strains B and C has no difference (but they could be different from Strain A). In this case, we need to make following contrast matrix:

```
> matrix(c(0,1,-1,0), ncol=1)
    [,1]
[1,] 0
[2,] 1
[3,] -1
[4,] 0
```

3.4.2 Example analysis for data from general experimental design

 Load in data distributed with DSS. This is a small portion of a set of RRBS experiments. There are 5000 CpG sites and 16 samples. The experiment is a 2 design (2 cases and 2 cell types). There are 4 replicates in each case:cell combination.

```
> data(RRBS)
> RRBS
An object of type 'BSseq' with
  5000 methylation loci
  16 samples
has not been smoothed
All assays are in-memory
```

case cell 1 HC rN 2 HC rN 3 HC rN 4 SLE aN 5 SLE rN 6 SLE aN 7 SLE rN 8 SLE aN 9 SLE rN 10 SLE aN 11 SLE rN 12 HC aN 13 HC aN 14 HC aN 15 HC aN	> d	esign	
1 HC rN 2 HC rN 3 HC rN 4 SLE aN 5 SLE rN 6 SLE aN 7 SLE rN 8 SLE aN 9 SLE rN 10 SLE aN 11 SLE rN 12 HC aN 13 HC aN 14 HC aN		case (cell
2HCrN3HCrN4SLEaN5SLErN6SLEaN7SLErN8SLEaN9SLErN10SLEaN11SLErN12HCaN13HCaN14HCaN			
4SLEaN5SLErN6SLEaN7SLErN8SLEaN9SLErN10SLEaN11SLErN12HCaN13HCaN14HCaN	2	НС	
5 SLE rN 6 SLE aN 7 SLE rN 8 SLE aN 9 SLE rN 10 SLE aN 11 SLE rN 12 HC aN 13 HC aN 14 HC aN	3	HC	rN
6SLEaN7SLErN8SLEaN9SLErN10SLEaN11SLErN12HCaN13HCaN14HCaN	4	SLE	aN
7 SLE rN 8 SLE aN 9 SLE rN 10 SLE aN 11 SLE rN 12 HC aN 13 HC aN 14 HC aN	5	SLE	rN
8 SLE aN 9 SLE rN 10 SLE aN 11 SLE rN 12 HC aN 13 HC aN 14 HC aN 15 HC aN	6	SLE	aN
9 SLE rN 10 SLE aN 11 SLE rN 12 HC aN 13 HC aN 14 HC aN 15 HC aN	7	SLE	rN
10 SLE aN 11 SLE rN 12 HC aN 13 HC aN 14 HC aN 15 HC aN	8	SLE	aN
11 SLE rN 12 HC aN 13 HC aN 14 HC aN 15 HC aN	9	SLE	rN
12 HC aN 13 HC aN 14 HC aN 15 HC aN	10	SLE	aN
13 HC aN 14 HC aN 15 HC aN	11	SLE	rN
14 HC aN 15 HC aN	12	HC	aN
15 HC aN	13	HC	aN
	14	HC	aN
16 HC rN	15	HC	aN
	16	HC	rN

 Fit a linear model using DMLfit.multiFactor function, include case, cell, and case:cell interaction.

> DMLfit = DMLfit.multiFactor(RRBS, design=design, formula=~case+cell+case:cell)
Fitting DML model for CpG site:

3. Use DMLtest.multiFactor function to test the cell effect. It is important to note that the coef parameter is the index of the coefficient to be tested for being 0. Because the model (as specified by formula in DMLfit.multiFactor) include intercept, the cell effect is the 3rd column in the design matrix, so we use coef=3 here.

```
> DMLtest.cell = DMLtest.multiFactor(DMLfit, coef=3)
```

Alternatively, one can specify the name of the parameter to be tested. In this case, the input coef is a character, and it must match one of the column names in the design matrix. The column names of the design matrix can be viewed by

```
> colnames(DMLfit$X)
[1] "(Intercept)" "caseSLE" "cellrN" "caseSLE:cellrN"
```

The following line also works. Specifying coef="cellrN" is the same as specifying coef=3.

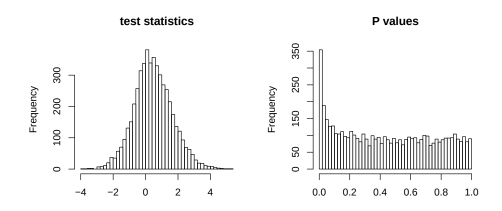
> DMLtest.cell = DMLtest.multiFactor(DMLfit, coef="cellrN")

Result from this step is a data frame with chromosome number, CpG site position, test statistics, p-values (from normal distribution), and FDR. Rows are sorted by chromo-some/position of the CpG sites. To obtain top ranked CpG sites, one can sort the data frame using following codes:

Below is a figure showing the distributions of test statistics and p-values from this example dataset

```
> par(mfrow=c(1,2))
```

- > hist(DMLtest.cell\$stat, 50, main="test statistics", xlab="")
- > hist(DMLtest.cell\$pvals, 50, main="P values", xlab="")



4. DMRs for multifactor design can be called using callDMR function:

```
> callDMR(DMLtest.cell, p.threshold=0.05)
     chr
           start
                    end length nCG
                                      areaStat
33 chr1 2793724 2793907
                            184
                                 5 12.619968
413 chr1 3309867 3310133
                            267
                                 7 -12.093850
250 chr1 3094266 3094486
                                  4 11.691413
                           221
262 chr1 3110129 3110394
                                 5 11.682579
                           266
180 chr1 2999977 3000206
                                 4 11.508302
                           230
121 chr1 2919111 2919273
                           163
                                 4 9.421873
298 chr1 3146978 3147236
                                      8.003469
                           259
                                 5
248 chr1 3090627 3091585
                           959
                                 5 -7.963547
346 chr1 3200758 3201006
                           249
                                 4 -4.451691
213 chr1 3042371 3042459
                            89
                                 5
                                     4.115296
169 chr1 2995532 2996558
                                  4 -2.988665
                           1027
```

Note that for results from for multifactor design, delta is NOT supported. This is because in multifactor design, the estimated coefficients in the regression are based on a GLM framework (loosely speaking), thus they don't have clear meaning of methylation level differences. So when the input DMLresult is from DMLtest.multiFactor, delta cannot be specified.

5. More flexible way to specify a hypothesis test. Following 4 tests should produce the same results, since 'case' only has two levels. However the p-values from F-tests (using term or Contrast) are slightly different, due to normal approximation in Wald test.

```
> ## fit a model with additive effect only
> DMLfit = DMLfit.multiFactor(RRBS, design, ~case+cell)
Fitting DML model for CpG site:
> ## test case effect
> test1 = DMLtest.multiFactor(DMLfit, coef=2)
> test2 = DMLtest.multiFactor(DMLfit, coef="caseSLE")
> test3 = DMLtest.multiFactor(DMLfit, term="case")
> Contrast = matrix(c(0,1,0), ncol=1)
> test4 = DMLtest.multiFactor(DMLfit, Contrast=Contrast)
> cor(cbind(test1$pval, test2$pval, test3$pval, test4$pval))
     [,1] [,2] [,3] [,4]
        1
             1
                  1
                       1
[1,]
[2,]
        1
                       1
             1
                  1
[3,]
        1
             1
                  1
                       1
                  1
                       1
[4,]
        1
             1
```

The model fitting and hypothesis test procedures are computationally very efficient. For a typical RRBS dataset with 4 million CpG sites, it usually takes less than half hour. In comparison, other similar software such as RADMeth or BiSeq takes at least 10 times longer.

4 Session Info

```
> sessionInfo()
```

```
R version 3.5.0 (2018-04-23)
Platform: x86_64-pc-linux-gnu (64-bit)
Running under: Ubuntu 16.04.4 LTS
```

Matrix products: default BLAS: /home/biocbuild/bbs-3.7-bioc/R/lib/libRblas.so LAPACK: /home/biocbuild/bbs-3.7-bioc/R/lib/libRlapack.so

locale:

	[1]	LC_CTYPE=en_US.UTF-8	LC_NUMERIC=C
	[3]	LC_TIME=en_US.UTF-8	LC_COLLATE=C
	[5]	LC_MONETARY=en_US.UTF-8	LC_MESSAGES=en_US.UTF-8
	[7]	LC_PAPER=en_US.UTF-8	LC_NAME=C
	[9]	LC_ADDRESS=C	LC_TELEPHONE=C
г	111		LC TRENTTETCATTON C

[11] LC_MEASUREMENT=en_US.UTF-8 LC_IDENTIFICATION=C

attached base packages:

```
[1] splines stats4 parallel stats graphics grDevices utils[8] datasets methods base
```

other attached packages:

[1]	edgeR_3.22.0	limma_3.36.0
[3]	DSS_2.28.0	bsseq_1.16.0
[5]	$SummarizedExperiment_1.10.0$	DelayedArray_0.6.0
[7]	BiocParallel_1.14.0	<pre>matrixStats_0.53.1</pre>
[9]	GenomicRanges_1.32.0	GenomeInfoDb_1.16.0
[11]	IRanges_2.14.0	S4Vectors_0.18.0
[13]	Biobase_2.40.0	BiocGenerics_0.26.0

loaded via a namespace (and not attached):

[1] Rcpp_0.12.16	compiler_3.5.0	plyr_1.8.4
[4] XVector_0.20.0	R.methodsS3_1.7.1	R.utils_2.6.0
[7] bitops_1.0-6	tools_3.5.0	<pre>DelayedMatrixStats_1.2.0</pre>
[10] zlibbioc_1.26.0	digest_0.6.15	rhdf5_2.24.0
[13] evaluate_0.10.1	lattice_0.20-35	Matrix_1.2-14

[16] yaml_2.1.18	GenomeInfoDbData_1.1.0	stringr_1.3.0
[19] knitr_1.20	gtools_3.5.0	locfit_1.5-9.1
[22] rprojroot_1.3-2	grid_3.5.0	data.table_1.10.4-3
[25] HDF5Array_1.8.0	rmarkdown_1.9	Rhdf5lib_1.2.0
[28] magrittr_1.5	backports_1.1.2	scales_0.5.0
<pre>[31] htmltools_0.3.6</pre>	permute_0.9-4	BiocStyle_2.8.0
[34] colorspace_1.3-2	stringi_1.1.7	RCurl_1.95-4.10
[37] munsell_0.4.3	R.oo_1.22.0	

References

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- [2] HAO WU, TIANLEI XU, HAO FENG, LI CHEN, BEN LI, BING YAO, ZHAOHUI QIN, PENG JIN AND KAREN N. CONNEELY. (2015). Detection of differentially methylated regions from whole-genome bisulfite sequencing data without replicates. *Nucleic Acids Research*. doi: 10.1093/nar/gkv715.
- [3] YONGSEOK PARK, HAO WU. (2016). Differential methylation analysis for BS-seq data under general experimental design. *Bioinformatics*. doi:10.1093/bioinformatics/btw026.
- [4] HAO WU, CHI WANG AND ZHIJING WU. (2013). A new shrinkage estimator for dispersion improves differential expression detection in RNA-seq data. *Biostatistics.* 14(2), 232–243.