methylKit: User guide

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1 Introduction

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

2 Basics

2.1 Reading the methylation call files

We start by reading in the methylation call data from bisulfate sequencing with read function. Reading in the data this way will return a methylRawList object which stores methylation information per sample.

2.2 Descriptive statistics on samples

Since we read the 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:

90%

```
Min. 1st Qu.
                  Median
                             Mean 3rd Qu.
                                               Max.
   0.00
           20.00
                   82.79
                             63.17
                                     94.74
                                             100.00
percentiles:
                                                              50%
                  10%
                             20%
                                        30%
                                                   40%
                                                                         60%
       0%
  0.00000
             0.00000
                        0.00000
                                  48.38710
                                             70.00000
                                                        82.78556
                                                                   90.00000
```

99%

96.42857 100.00000 100.00000 100.00000 100.00000 100.00000 100.00000

99.5%

99.9%

70%

93.33333

100%

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.

> library("graphics")

80%

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

95%

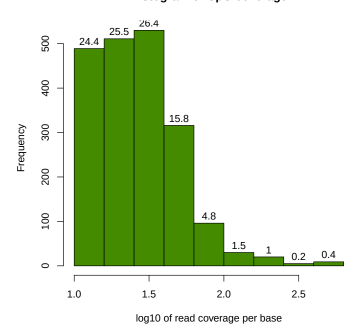
800 38.8 900 23.5 Frequency 400 14.3 6.6 3.9 1.6 0 0 20 40 60 80 100 % methylation per base

Histogram of % 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)

Histogram of CpG coverage



2.3 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.

> filtered.myobj=filterByCoverage(myobj,lo.count=10,lo.perc=NULL,hi.count=NULL,hi.perc=99.

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 base-pair 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). This operation will return a methylBase object which will be our main object for all comparative analysis.

> methidh=unite(myobj,destrand=FALSE)

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

> head(methidh)

| | id | chr | start | end | strand | coverage1 | numCs1 ı | numTs1 |
|-----------------------|------------------------------|----------------|------------|------------------------------|--------------------------|----------------------|----------------------------|-------------------|
| 1 | chr21.10011833 | chr21 | 10011833 | 10011833 | + | 174 | 173 | 1 |
| 2 | chr21.10011841 | chr21 | 10011841 | 10011841 | + | 173 | 164 | 9 |
| 3 | chr21.10011855 | chr21 | 10011855 | 10011855 | + | 175 | 175 | 0 |
| 4 | chr21.10011858 | chr21 | 10011858 | 10011858 | + | 175 | 131 | 44 |
| 5 | chr21.10011861 | chr21 | 10011861 | 10011861 | + | 174 | 147 | 27 |
| 6 | chr21.10011872 | chr21 | 10011872 | 10011872 | + | 167 | 160 | 7 |
| | | | | | | | | |
| | coverage2 numCs | s2 numT | Ss2 covera | age3 numCs | 3 numTs | 3 coverage | 4 numCs4 | l numTs4 |
| 1 | • | s2 num1 18 | Ss2 covera | • | | 3 coverage 6 | | _ |
| 1 2 | 18 | | _ | 40 3 | 34 | • | 4 14 | 1 0 |
| 1 2 3 | 18 1 20 1 | 18 | _ | 40 3 40 1 | 34 18 2 | 6 1 | 1 14 1 8 | 1 0 3 6 |
| 1 2 3 4 | 18 20 21 2 | 18 19 | 0 1 | 40 3 40 3 39 2 | 34 18 2 29 1 | 6 1- 2 1- | 1 14 1 8 1 12 | 1 0 3 6 |
| 1 2 3 4 5 | 18 1 20 2 21 2 21 2 | 18 19 21 | 0 1 | 40 3 40 3 39 2 39 3 | 34 18 2 29 1 31 | 6 14 2 14 0 14 | 4 14 4 8 4 12 3 8 | 1 0 3 6 2 2 |

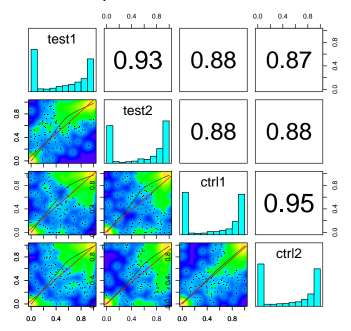
3.2 Sample Correlation

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(methidh,plot=T)

```
test1 test2 ctrl1 ctrl2
test1 1.0000000 0.9252530 0.8767865 0.8737509
test2 0.9252530 1.0000000 0.8791864 0.8801669
ctrl1 0.8767865 0.8791864 1.0000000 0.9465369
ctrl2 0.8737509 0.8801669 0.9465369 1.0000000
```

CpG dinucleotide 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(methidh, dist="correlation", method="ward", plot=TRUE)

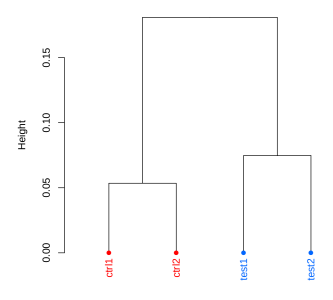
Call:

hclust(d = d, method = HCLUST.METHODS[hclust.method])

Cluster method : ward
Distance : pearson

Number of objects: 4

CpG dinucleotide methylation clustering



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

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(methidh, dist="correlation", method="ward", plot=FALSE)

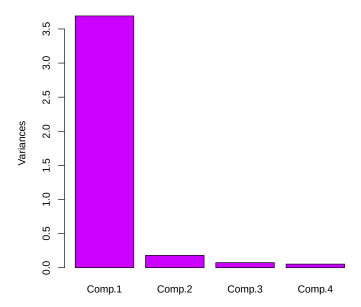
We can also do a PCA analysis on our samples. The following function will plot a scree plot for importance of components.

> PCASamples(methidh, screeplot=TRUE)

Importance of components:

| | Comp.1 | Comp.2 | Comp.3 | Comp.4 |
|------------------------|-----------|------------|-----------|------------|
| Standard deviation | 1.9211651 | 0.42545636 | 0.2735654 | 0.23081050 |
| Proportion of Variance | 0.9227188 | 0.04525328 | 0.0187095 | 0.01331837 |
| Cumulative Proportion | 0.9227188 | 0.96797213 | 0.9866816 | 1.00000000 |

CpG dinucleotide methylation PCA Screeplot

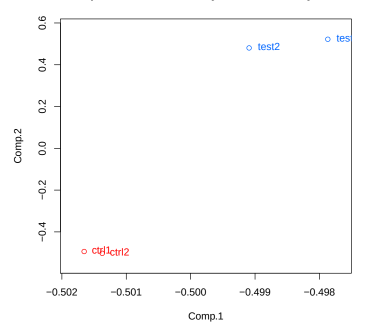


We can also plot PC1 and PC2 axis and a scatter plot of our samples on those axis which will reveal how they cluster.

> PCASamples(methidh)

Importance of components:





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 to get differentially methylated regions

```
> tiles=tileMethylCounts(myobj,win.size=1000,step.size=1000)
> head(tiles[[1]])
```

| | id | chr | start | end | strand | coverage | numCs | numTs |
|---|-----------------------|-------|---------|---------|--------|----------|-------|-------|
| 1 | chr21.9764001.9765000 | chr21 | 9764001 | 9765000 | * | 24 | 3 | 21 |
| 2 | chr21.9820001.9821000 | chr21 | 9820001 | 9821000 | * | 13 | 0 | 13 |
| 3 | chr21.9837001.9838000 | chr21 | 9837001 | 9838000 | * | 11 | 0 | 11 |
| 4 | chr21.9849001.9850000 | chr21 | 9849001 | 9850000 | * | 124 | 90 | 34 |
| 5 | chr21.9853001.9854000 | chr21 | 9853001 | 9854000 | * | 34 | 22 | 12 |
| 6 | chr21.9860001.9861000 | chr21 | 9860001 | 9861000 | * | 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 method¹.

> myDiff=calculateDiffMeth(methidh)

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%.

```
> 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.

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.

percentage of target features overlapping with annotation:
promoter exon intron intergenic
27.81955 15.03759 34.58647 57.14286

```
percentage of target features overlapping with annotation (with promoter>exon>intron prece promoter exon intron intergenic 27.81955 0.00000 15.03759 57.14286

percentage of annotation boundaries with feature overlap: promoter exon intron 0.018129079 0.001589593 0.010038738
```

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

Mean 3rd Qu.

94640

52030

> # read the shores and flanking regions and name the flanks as shores and CpG islands as > cpg.obj=read.feature.flank(system.file("extdata", "cpgi.hg18.bed.txt", package = "methyl")

Max. 313500

> diffCpGann=annotate.WithFeature.Flank(myDiff25p,cpg.obj\$CpGi,cpg.obj\$shores,feature.name

4.1 Regional analysis

Min. 1st Qu.

828

5

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]])

summary of distances to the nearest TSS:

45160

Median

- id chr start end strand coverage numCs 1 chr21.17806094.17808094.NA chr21 17806094 17808094 1834 2 chr21.10119796.10121796.NA chr21 10119796 10121796 79 44 3 chr21.10011791.10013791.NA chr21 10011791 10013791 3697 2982 4 chr21.10119808.10121808.NA chr21 10119808 10121808 79 44 5 chr21.15357997.15359997.NA chr21 15357997 15359997 8613 16 6 chr21.16023366.16025366.NA chr21 16023366 16025366 6296 5 numTs
- 1 1827
- 2 35
- 3 715
- 4 35
- 5 8594
- 6 6291

4.2 Convenience 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.

- > # read the annotation with unique.prom=FALSE does not collapse identical
- > # promoter boundaries and preserves gene names associated with TSS
- > gene.obj=read.transcript.features(system.file("extdata", "refseq.hg18.bed.txt",
- package = "methylKit"),unique.prom=FALSE)
- > diffAnn=annotate.WithGenicParts(myDiff25p,gene.obj)
- > # 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 | - |
|------|---|-----|-----------|---|
| 60.1 | 2 | 931 | NM_199260 | - |
| 60.2 | 3 | 838 | NM_199260 | - |
| 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

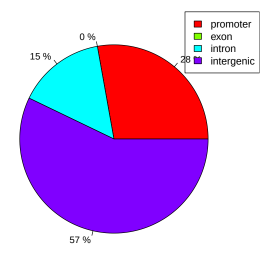
 $\verb|>| getTargetAnnotationStats(diffAnn,percentage=FALSE,precedence=TRUE)| \\$

| promoter | exon | intron | intergenic |
|----------|------|--------|------------|
| 37 | 0 | 20 | 76 |

We can also plot the percentage of differentially methylated bases over lapping with exon/intron/promoters $\,$

- > plotTargetAnnotation(diffAnn,precedence=TRUE,
- + main="differential methylation annotation")

differential methylation annotation



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

> getTargetAnnotationStats(diffAnn,percentage=FALSE,precedence=TRUE)

| promoter | exon | intron | intergenic |
|----------|------|--------|------------|
| 37 | 0 | 20 | 76 |

References

[1] 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.