

Journal Club

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Contributions: @JC-STA426-20181022

22th October, 2018

Detection and accurate false discovery rate control of differentially methylated regions from whole genome bisulfite sequencing

Keegan Korthauer, Sutirtha Chakraborty, Yuval Benjamini, Rafael A Irizarry

Biostatistics, kxy007, <https://doi.org/10.1093/biostatistics/kxy007>

Published: 22 February 2018

Table of Contents

dmrseq paper

Epigenetics

Whole Genome Bisulphite Sequencing

Limitations of current methods

dmrseq

Comparison: dmrseq vs other methods

Epigenetics

A layer of **additional information** over genetic information.

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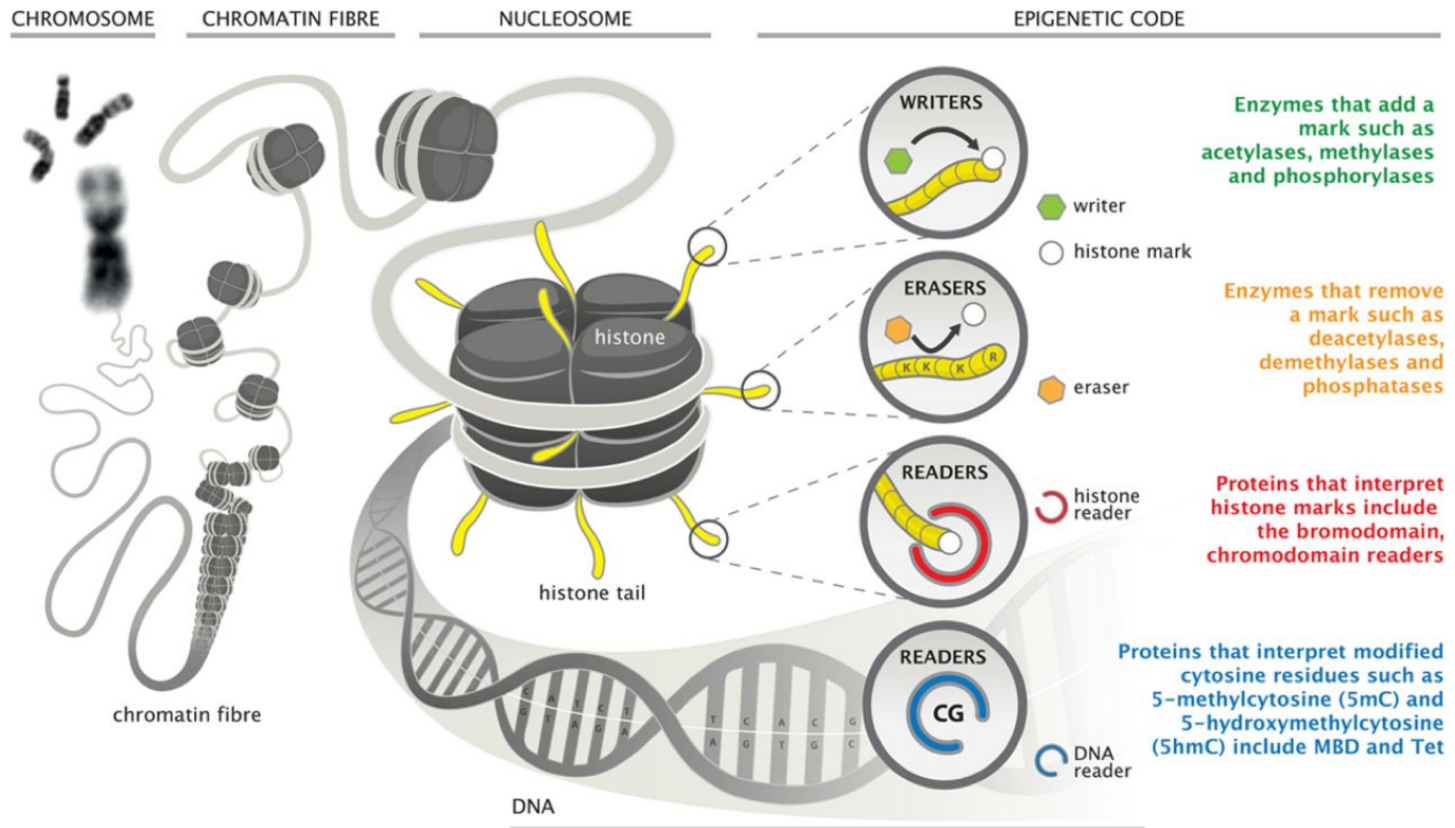
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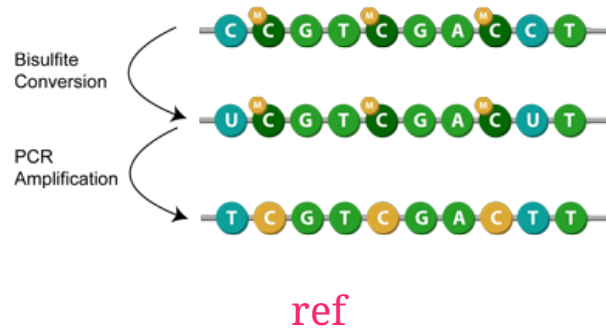
Various Epigenetic Modifications



Samuel T. Keating et al. Circ Res. 2016;118:1706-1722

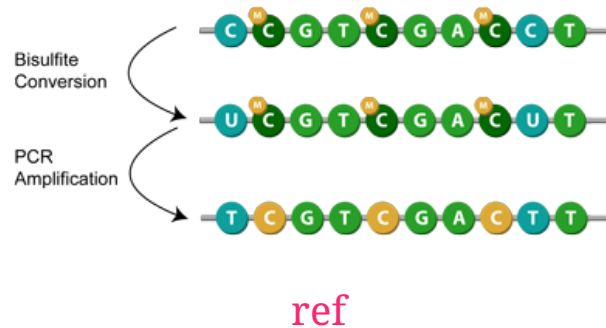
Whole Genome Bisulphite Sequencing (WGBS)

Bisulphite conversion

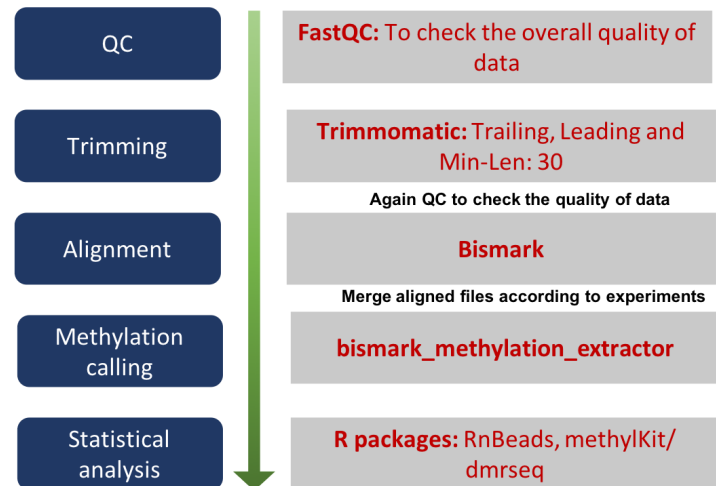


Whole Genome Bisulphite Sequencing (WGBS)

Bisulphite conversion



Basic Data analysis pipeline



Compared to methods developed for RNA-Seq, there are not that many statistical methods available for studying WGBS

Applications of WGBS

- Detection of Differentially Methylated Regions (**DMRs**)
- Detection of Differentially Methylated Loci (**DMLs**)
- Detection of Copy Number Variations (CNVs)
- Detection of Single Nucleotide Polymorphisms (SNPs)
- Detection of Cytosine methylation levels of Transcription Factor Binding Sites (TFBSs)

....cont

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- Multiple testing corrections without considering the spatial correlation can result in a loss of power.
- DMRs more biologically relevant than DMLs.

....cont.

- DML approaches may construct DMRs by chaining together neighboring significant loci, but this type of approach will not yield a proper assessment of the statistical significance of the constructed regions, nor will the FDR be properly controlled [4]. Controlling the FDR at the level of individual loci is not the same as controlling FDR of regions.

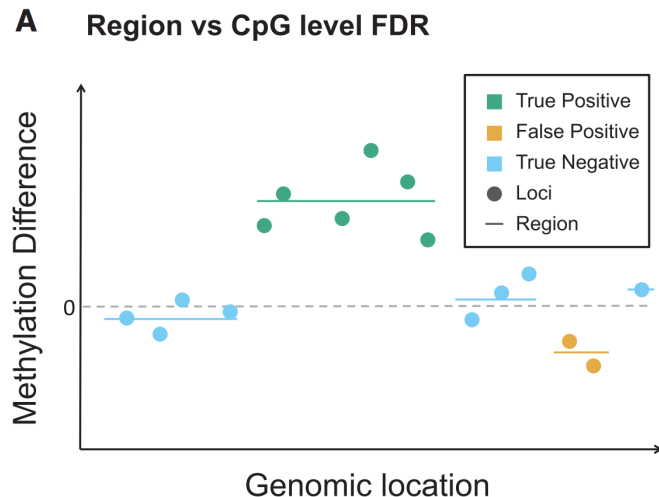
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- Methylation data cannot be modeled neither by Gaussian models (due to low coverage), nor by Binomial models (do not account for biological variability). Beta-Binomial models are computationally difficult.
- Challenges for assessing DMRs:
 - defining region boundaries
 - methods ignore correlation across loci
 - biological variability from sample to sample

FDR at the loci level is not the same as FDR at the region level



Genomic location versus methylation difference estimates at several neighboring loci.

The individual CpGs (points) are shaded by whether they are a true or false positive. Regions are denoted by lines.

The loci FDR is:

$$\text{FDR}_{\text{loci}} = (\text{\#False Positive Loci}) / (\text{Total \# of Significant Loci})$$

The region FDR is:

$$\text{FDR}_{\text{region}} = (\text{\#False Positive Regions}) / (\text{Total \# of Significant Regions})$$

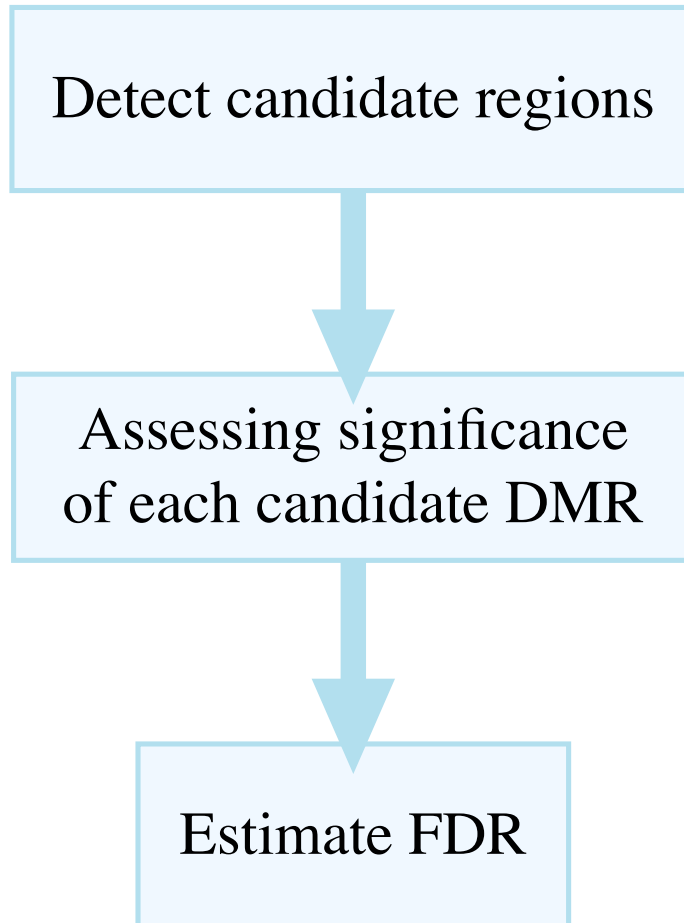
dmrseq



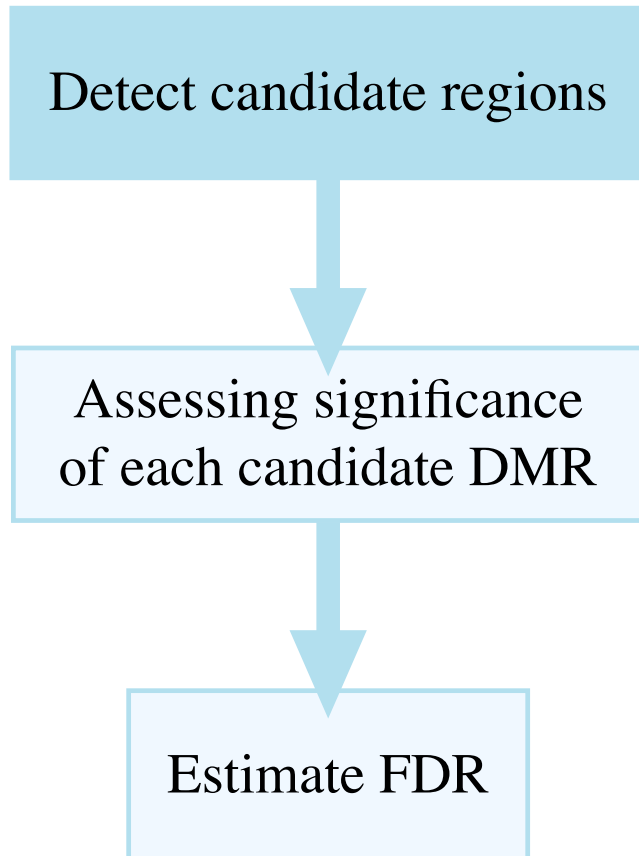
Aim:

1. To **maximize power** while **controlling the FDR** even with sample sizes as small as **two samples per condition**.
1. To develop a procedure to control FDR at the region level and provide an accurate measure of statistical significance for each region.

Identification of DMRs by dmrseq



Detecting candidate regions



- mean methylation proportion for locus i in condition s :

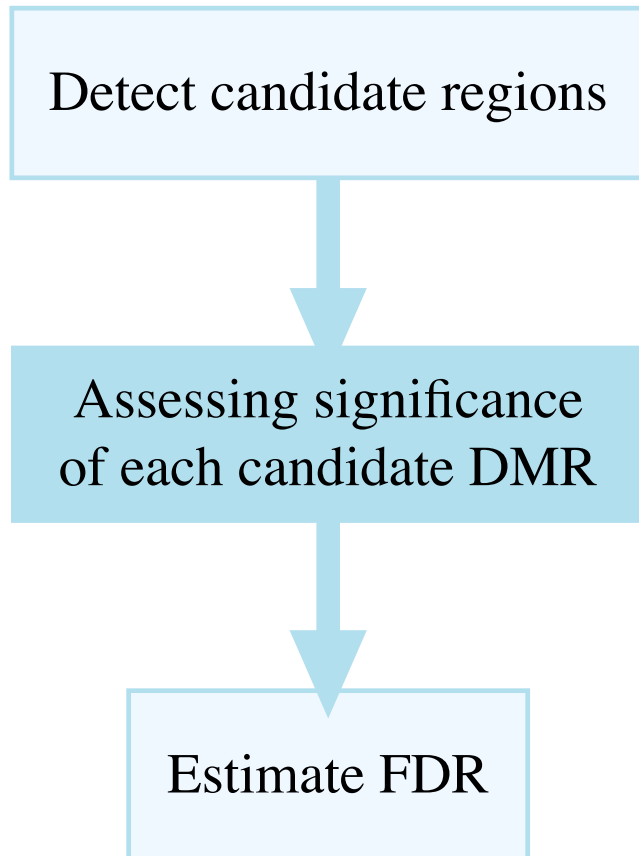
$$\pi_{is} = M_{is}/N_{is}$$

- methylation difference of locus i between biological conditions s and s' :

$$\beta_i = \pi_{is} - \pi_{is'}$$

- β_i 's are smoothed
- candidate regions: segments of genomes with smoothed β_i 's in the same direction and in absolute value greater than a threshold (e.g. 0.1)

Assessing significance of candidate DMRs



- methylation counts modeled by Beta-Binomial distribution
- to stabilize the dependence of variance on mean, data are transformed:

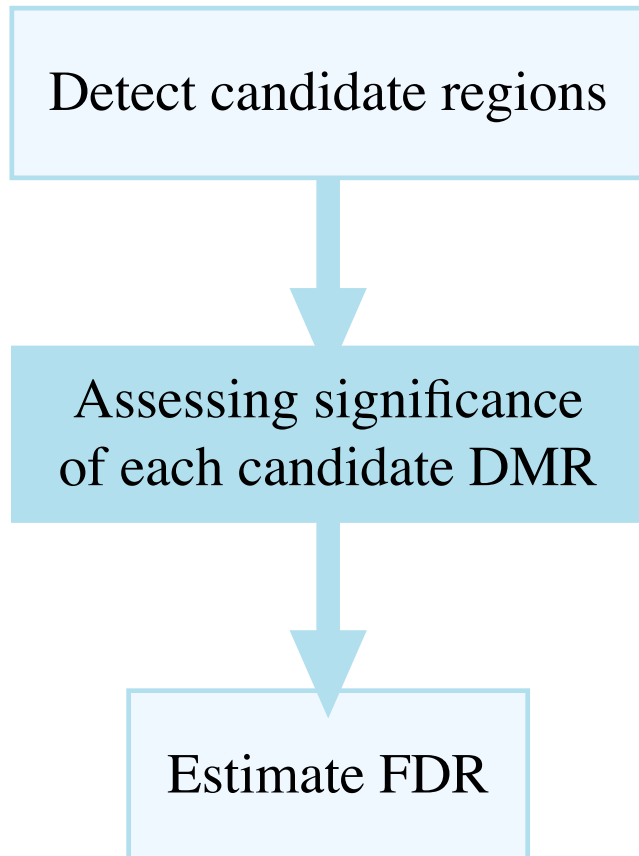
$$Z_{ijr} = \arcsin(2M_{ijr}/N_{ijr} - 1)$$

- mean region methylation level can be then estimated using Generalized Least Squares:

$$Z_r = X\beta_{0r} + \beta_{1r},$$

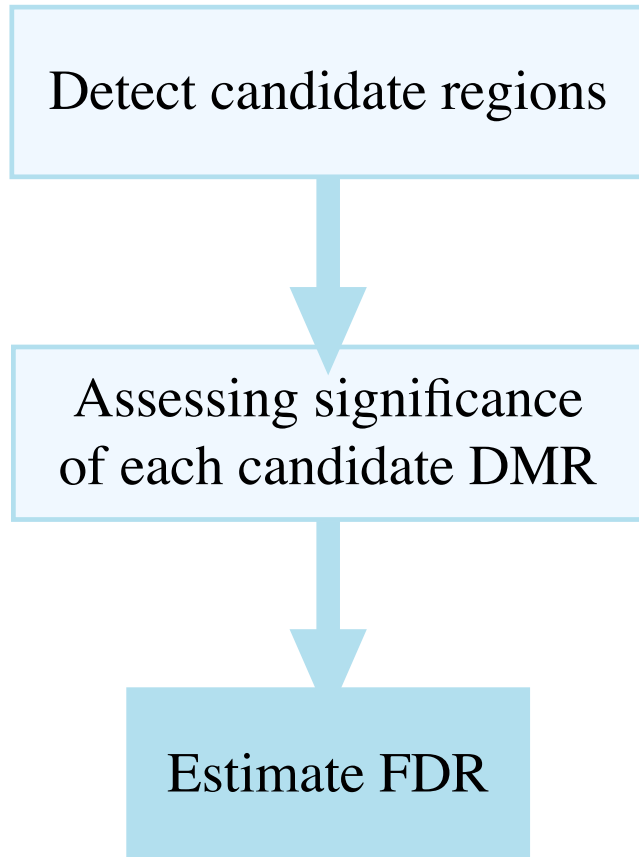
with β_{1r} being the effect of biological group

Assessing significance of candidate DMRs



- Estimation of covariance matrix V_r :
 - variance dependent on coverage
 $\rightarrow \text{Var}(Z_{ijr}) \approx \sigma_r^2 / N_{ijr}$
 - correlation of nearby loci \rightarrow modeled correlation by coefficients dependent of the length of the interval between the two loci

Estimating FDR



- permutation to get an approximate null distribution
- p-value computed by comparing the observed test statistics to the null distribution

Data used for comparison

- six samples of normal human dendritic cells, partitioned into groups of two or three → no DMRs, **negative control**

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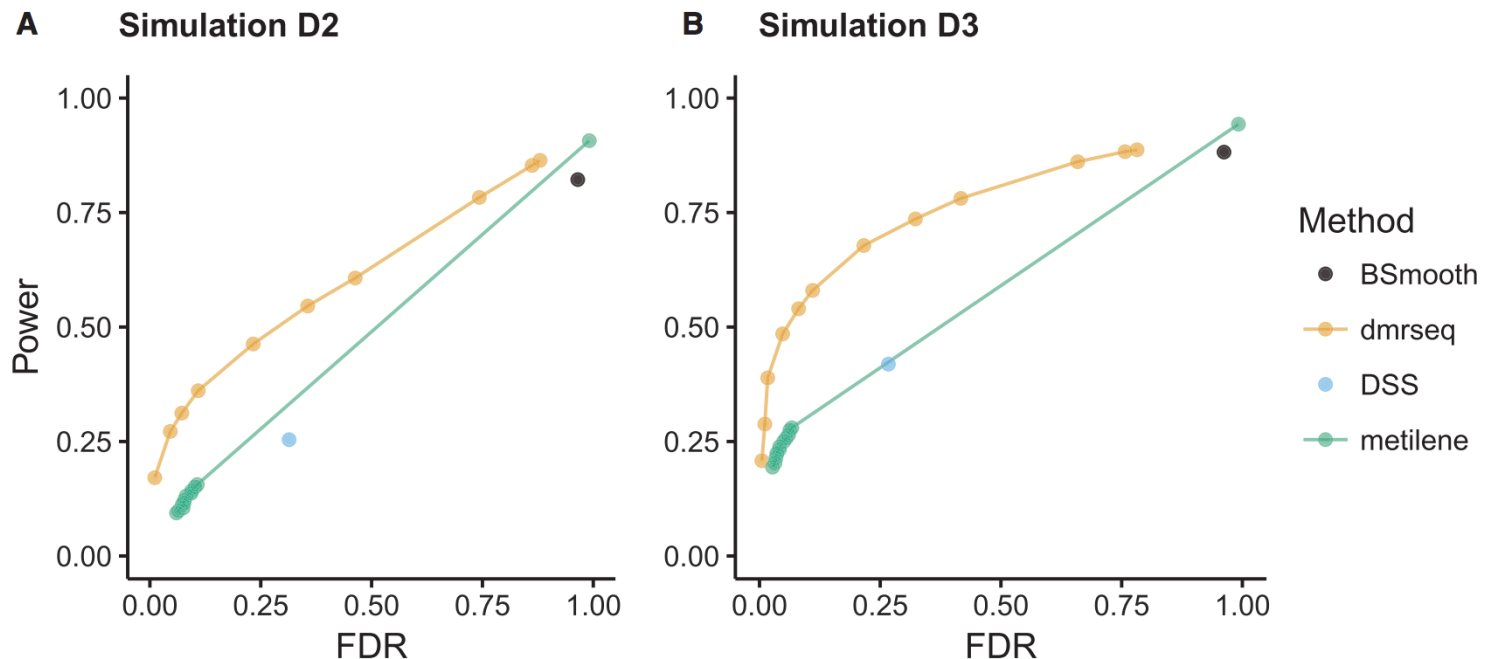
- six samples of normal human dendritic cells, partitioned into groups of two or three → no DMRs, **negative control**
- **simulated data:** 3000 simulated DMRs added to each data set
 - DMR is created by choosing a cluster of CpGs and simulating the number of methylated reads from a binomial distribution
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- **real data:** methylation data from several human tissues and from two mouse strains

dmrseq is more powerful

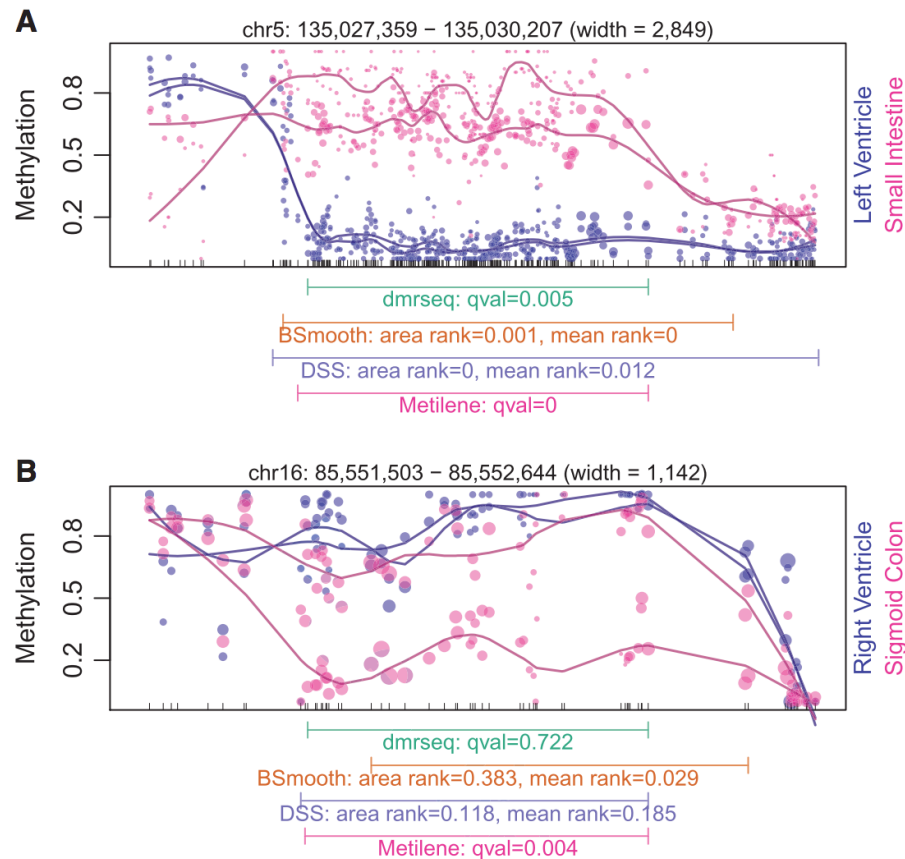
power vs. FDR on simulated data



Power is calculated as the proportion of simulated DMRs overlapped by at least one identified DMR. FDR is calculated as the proportion of DMRs identified that do not overlap with any of the simulated DMRs.

dmrseq performs better

Examples of real biological data and their ranking by different DMRs identification methods.



dmrseq availability

- GitHub: [kdkorthauer/dmrseq](#)
- Bioconductor: [dmrseq](#)

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- Consideration of all possible shortcomings of the available methods for DMRs detection.
- Evaluation of the software using experimental data and Monte Carlo simulations.
- Implementation in **R**.
- Comparison of the new method with the "famous" methods.
- Comparison is not done with **RnBeads**. Probably because RnBeads performs differential methylation analysis (only) at single nucleotide level.
- Comparison is not done with **methyKit**, although methylKit performs differential methylation analysis at regional level.

References

- [1] Smith Z. D. and Meissner A. (2013). DNA methylation: roles in mammalian development. *Nature Reviews Genetics* 14, 204–220.
- [2] Leek J. T., Scharpf R. B., Bravo H. C., Simcha D., Langmead B., Johnson W. E., Geman D., Baggerly K. and Irizarry R. A. (2010). Tackling the widespread and critical impact of batch effects in high-throughput data. *Nature Reviews Genetics* 11, 733–739.
- [3] Aryee M. J., Jaffe A. E., Corrada-Bravo H., Ladd-Acosta C., Feinberg A. P., Hansen K. D. and Irizarry R. A. (2014). Minfi: a flexible and comprehensive Bioconductor package for the analysis of infinium DNA methylation microarrays. *Bioinformatics* 30, 1363–1369.
- [4] Robinson M. D., Kahraman A., Law C. W., Lindsay H., Nowicka M., Weber L. M. and Zhou X. (2014). Statistical methods for detecting differentially methylated loci and regions. *Frontiers in Genetics* 5, 324.

Thank you!

Slides are prepared using **xaringan** R package