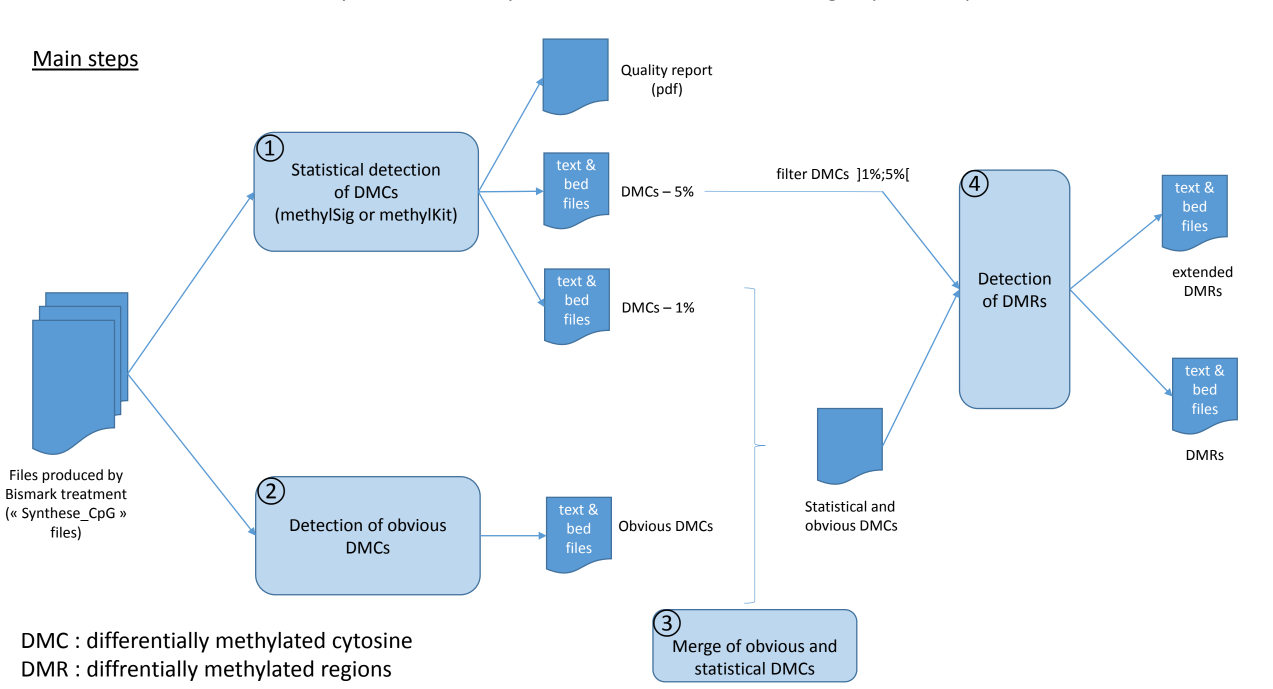
# Analysis of DNA methylation differences between two groups of samples



## Figure comments

- Detection of CpGs significantly differentially methylated between two conditions
   This analysis requires to have replicates within group of each condition.
   User can choose between analysis conducted by either methylSig or methylKit packages.
   The analysis produces a quality report file and two {text;bed} files containing the significant DMCs for two p-value/q-value thresholds (default: <1% and <5%). Bed files can be imported in genome browers (e.g. IGV)</p>
- Detection of CpGs obviously differentially methylated between two conditions. This analysis does not require replicates within the conditions compared. If parameter methdiff\_threshold2 is set to 0.95, all samples in condition C1 must have a % of methylation >=95%, while samples in condition C2 must have a % of methylation <=5% (idem with inversion of C1 and C2 conditions).</p>
- 3 : DMCs statistically significants (statistical value: pValue or qValue<1% ) detected in step 1 are merged with obvious DMCs detected in step 2
- ④: Adjacents DMCs collected in step ③ are grouped in DMRs (DMRs output file).

  DMCs detected in step ① with a p-value/q-value ∈ ]1%;5%[ and adjacent to DMRs are used to extend these DMRs (« extended DMRs » output file)

## Requirements

## Before to launch analysis one need to:

- have a set of files produced by bismark (synthese\_CpG.txt), one for each replicate from each condition
- prepare a configuration file describing the parameters used for the analysis (e.g : analysis\_config.txt)

## To launch analysis, type following command:

```
RRBS_HOME/Scripts/Differential_analysis/get_methylation_differences.sh <relative or absolute path to config file>
```

## Structure of analysis config file:

```
#Analysis parameters:
#output dir
                        ./out
                                                                  Part dedicated to analysis parametrization
#title
                       Male vs Female
                                                                  (parameters and values are separated by one or several <TAB> character)
#parameter n
                        value
Sample File
                   Condition
        M1/extract/synthese CpG.txt
                                                Male
                                                                  Definition of the two groups to compare and localization of analysis input files
        M2/extract/synthese CpG.txt
                                                Male
M2
                                                                  (fileds in this table are separated by a <TAB> character)
        F1/extract/synthese CpG.txt
                                                Female
F1
        F2/extract/synthese CpG.txt
F2
                                                Female
```

'output\_dir' and 'title' parameters are the two global parameters needed for the analysis ('title' is mandatory). All other parameters are step specific and will be presented hereafter.

Several examples of config files and their corresponding analysis results are available in Differential\_analysis/analysis\_examples directory.

## Explanations on configuration parameters

# 1 Detection of DMCs

<pre>#MethylSig/Kit parameters: #</pre>	
#stat method	methylKit
#min_coverage1	10
#max_coverage1	500
#min_per_group	2
#stat_value	pvalue
#stat_threshold1	0.01
#methdiff_threshold1	0.25

#### stat method:

Either 'methylSig' or 'methylKit' (no default value)

### min coverage1, max coverage1 and min per group :

Minimal and maximal coverage for a CpG to be taken into account for the analysis.

A minimal number of samples (min per group) in each group must satisfy this coverage range.

(default values: 10 for min\_coverage1, no limit for max\_coverage1, smallest group size for min\_per\_group).

#### stat value:

Either 'pValue' or 'qValue' depending whether one wants to use the raw or the adjusted p-value for significant result selection. (default value: qValue)

### stat threshold1:

Threshold used for significant result selection (used in conjunction with stat\_value parameter) (default value : 0.01)

### methdiff\_threshold1:

Minimum value accepted for the absolute difference between average methylation in the two groups (default value: 0.25)

## Explanations on configuration parameters

# 2 Detection of obvious DMCs

#Obvious DMCs parameters:	
#	1.0
#min_coverage2	10
#max_coverage2	500
#methdiff_threshold2	0.95

## min\_coverage2, max\_coverage2 :

Minimal and maximal coverage for a CpG to be taken into account for the analysis. All samples must satisfy these criteria. (default value: 10 for min, 500 for max).

## methdiff\_threshold2:

Minimum value accepted for the methylation in all samples of one condition, while all samples in the other condition will have a maximal methylation of 1-methdiff\_threshold2 (default value : 1)

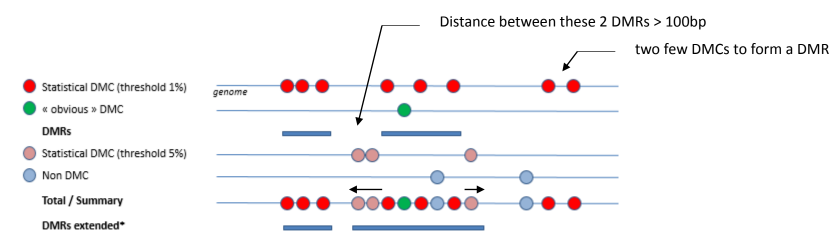
# Explanations on configuration parameters

# (3) Merge of significant and obvious DMCs

No specific parameter is required for this step.

# 4 Detection of DMRs

#DMCs -> DMRs parameters:	
#nb_min_DMCs_in_DMRs	3
#max_distance_between_DMCs	100
#stat_threshold2	0.05



\*DMRs are extended to neighbouring DMCs which are distant from less than 100 bp and for which p-value/q-value< 0.05

#### max distance between DMCs:

Maximum distance accepted between two DMCs to decide that they are neighbours (default value: 100)

### nb min DMCs in DMRs :

Minimal number of neighbours DMCs needed to get a DMRs (default value : 3).

### stat threshold2:

Used to select DMCs detected in step 1. These DMCs will be used to try to extend DMRs.

# Possible analyses

# 1°) More than one sample per condition

Please refer to:

DMCs config methylKit.txt

or:

DMCs config methylSig.txt

# 2°) Only one sample per condition

Please refer to:

DMCs config nostat.txt

# 3°) Generation of DMRs from a list of DMCs

Please refer to:

DMCs\_config\_DMRonly.txt

(these configuration files are available in RRBS\_HOME/Scripts/Differential\_analysis/analysis\_examples directory)

## Comparing analyses results

The script RRBS\_HOME/Scripts/Differential\_analysis/venn\_DMCs\_sets.py allows you to compare a list of DMCs.

Command used to launch the comparison:

python venn DMCs sets.py file DMC set1.txt file DMC set2.txt file DMC set3.txt > output file

You can compare 2 sets or 3 sets of results.

Input file must have following format:

Chromosome Start <any other columns> Difference in methylation Methylation state

Second to last column (Difference in methylation) should be filled with a value corresponding to difference in methylation between the two conditions compared.

### Example:

Chromosome	Start	<any columns="" other=""></any>	Difference in methylation	Methylation state
1	123456	••••	-22.345	hypometh
1	345678		50.3461	hypermeth

Output file must format (example for a comparison of two files (A and B):

Chromosome	Start	End	Only in A	Only in B	Common A B	Α	В	
1	123456	123457	*			-22.345		
1	234567	234568		*			44.8277	
1	345678	345679			*	50.3461	50.1234	

difference in methylation found in A and B