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Chapter 1

What is CGmapTool

DNA methylation is crucial for a wide variety of biological processes. With the development of high throughput methylome profiling methods, huge volumes of data are generated and in egent need of computational tools for data analysis. Though several tools have been proposed to fit this need, there is not a mainstream standard for bisulfite sequencing data storage and manipulation. What's more, the performance of available tools needs to be improved.

We proposed **CGmapTool**, a bisulfite sequencing analysis toolset with enhanced features on SNV calling and allele specific methylations and visualizations, in hope to set up a standard for bisulfite sequencing data related manipulation, including better data storage, extraction, visualization and improved performance in SNP calling. We also provide dozens of utilities and a seamless pipeline for bisulfite sequencing data analysis.

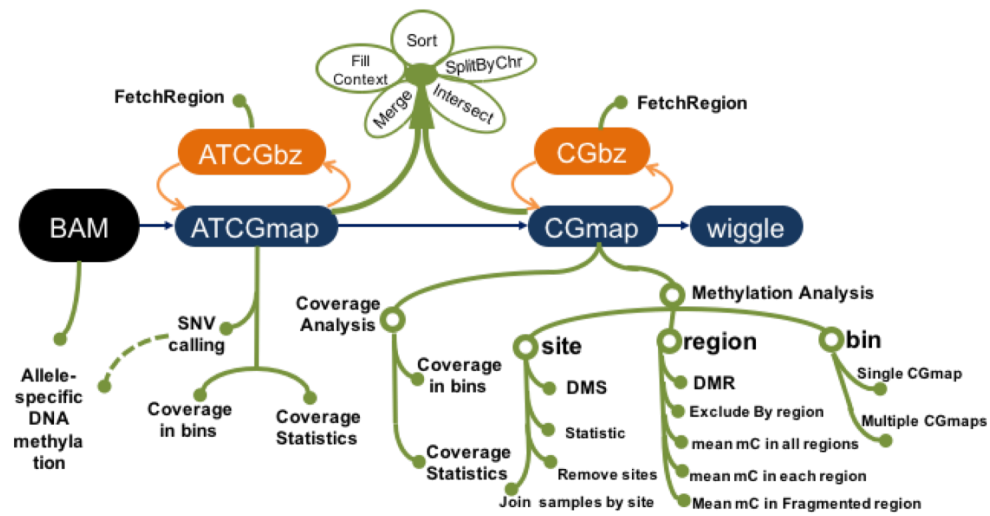


Figure 1.1: Schematic diagram of CGmapTools

Chapter 2

New File Format

To facilitate high throughput data manipulation and reduce storage usage, several file format have been proposed and generally accepted as the standard. Due to these great efforts (e.g. SAM/BAM and VCF), data analysis and tool development become more easier and highly efficient. However, when it comes to bisulfite sequencing data, currently, available tools possess their own tool specific data format. In consequence, integrating results from several tools leads to extra efforts in unifying data format and developing customized tools, which is time consuming and error prone.

As one of the features of CGmapTools, we defined ATCGmap and CGmap file format to simplify downstream DNA methylation analysis and in hope to standardize the storage format of bisulfite sequencing data.

2.1 ATCGmap Format

After alignment of sequencing reads to the reference genome, all the detail information about read coverage and methylation level of a cytosine site are stored in BAM/SAM format files though requiring further interpretation. A well defined file format called **pileup** summarized the information of mapped reads covered on each nucleotide along the reference genome. But the pileup file does not designed for bisulfite sequencing data, which lacks DNA methylation estimation of cytosines.

Here, we defined ATCGmap file format to integrate both mapping and coverage of non-cytosine and cytosine sites with estimated DNA methylation in a single file.

Col	Field	Type	Regexp/Range	Brief description
1	CHR	String	[!-?A-~]{1,118}	Query template NAME
2	NUC	Char	[ATCGN-]	The nucleotide on reference genome
3	POS	Int	[0,232-1]	1-based leftmost mapping position
4	CONT	String	{"-", "\"CG\", \"CHG\", \"CHH\"}	Context
5	DINUC	String	{"-", \"CA\", \"CT\", \"CC\", \"CG\"}	Dinucleotide context
6	WA	Int	[0,214-1]	Counts of reads on Watson strand support Adenine
7	WT	Int	[0,214-1]	Counts of reads on Watson strand support Thymine
8	WC	Int	[0,214-1]	Counts of reads on Watson strand support Cytosine
9	WG	Int	[0,214-1]	Counts of reads on Watson strand support Guanine
10	WN	Int	[0,26-1]	Counts of reads on Watson strand support None
11	CA	Int	[0,214-1]	Counts of reads on Crick strand support Adenine
12	CT	Int	[0,214-1]	Counts of reads on Crick strand support Thymine
13	CC	Int	[0,214-1]	Counts of reads on Crick strand support Cytosine
14	CG	Int	[0,214-1]	Counts of reads on Crick strand support Guanine
15	CN	Int	[0,26-1]	Counts of reads on Crick strand support None
16	METH	Float	[0,1] or "na"	Methylation level or "Not Available"

2.2 CGmap Format

In cases we only want to retain DNA methylation on cytosines to save storage usage, we defined another file format called **CGmap** which provides sequence context and estimated DNA methylation level of any covered cytosines on the reference genome.

Col	Field	Type	Regexp/Range	Brief description
1	CHR	String	[!-?A-~]{1,118}	Query template NAME
2	NUC	Char	[ATCGN-]	The nucleotide on reference genome
3	POS	Int	[0,2 ³² -1]	1-based leftmost mapping position
4	CONT	String	{"-", "CG", "CHG", "CHH"}	Context
5	DINUC	String	{"-", "CA", "CT", "CC", "CG"}	Dinucleotide context
6	METH	Float	[0,1] or "na"	Methylation level or "Not Available"
7	MC	Int	[0,2 ¹² -1]	Counts of reads support methylated Cytosine
8	NC	Int	[0,2 ¹² -1]	Counts of reads support all Cytosine

Chapter 3

File Manipulation

CGmapTool provides multiple utilities to manipulate files in ATCGmap and CGmap format or compressed ATCGbz/CGbz format.

Usage: `cgmaptools <convert|fetch|refill|intersect|merge2|mergelist|sort|split|select|> [options]`

3.1 convert

- **Description:** File format conversion.
- **Usage:** `cgmaptools convert <command> [options]`
- **Commands:**

Commands	From	To
<code>bam2cgmap</code>	BAM	CGmap & ATCGmap
<code>atcgmap2atcgbz</code>	ATCGmap	ATCGbz
<code>atcgbz2atcgmap</code>	ATCGbz	ATCGmap
<code>atcgmap2cgmap</code>	ATCGmap	CGmap
<code>cgmap2cgbz</code>	CGmap	CGbz
<code>cgbz2cgmap</code>	CGbz	CGmap
<code>cgmap2wig</code>	CGmap	WIG

- **Example:**

#1 The commands below will convert bam file to cgmap format.

```
cgmaptools convert bam2cgmap -b <BAM> -g <genome.fa> -o <prefix>
```

#2 This command will convert cgmap to wig format.

```
cgmaptools convert cgmap2wig [-i <CGmap>] [-w <wig>] [-c <INT> -b <float>]
```

Note: please refer to the help message for usage details using `-h` option.

3.2 fetch

- **Description:** Fastly access methylation data in specified region.

- **Usage:** `cgmmaptools fetch <command> [options]`
- **Commands:**
 - atcgbz:** fetch lines from ATCGBz file.
Usage: `cgmmaptools fetch atcgbz -b <ATCGBz> -C <CHR> -L <LeftPos> -R <RightPos>`
 -b, --ATCGBz <arg> input ATCGBz file
 -C, --CHR <arg> specify the chromosome name
 -L, --leftPos <arg> the left position
 -R, --rightPos <arg> the right position
 - cgbz:** fetch lines from CGBz file.
Usage: `cgmmaptools fetch cgbz -b <CGBz> -C <CHR> -L <LeftPos> -R <RightPos>`
 -b, --CGBz <arg> input CGBz file
 -C, --CHR <arg> specify the chromosome name
 -L, --leftPos <arg> the left position
 -R, --rightPos <arg> the right position

3.3 refill

- **Description:** Fill the CG/CHG/CHH and CA/CC/CT/CG context to CGmap or ATCGmap files. Other fields will not be affected.
- **Usage:** `cgmmaptools refill [-i <CGmap>] -g <genome.fa> [-o output]`
 -i STRING Input CGmap file (CGmap or CGmap.gz)
 -g STRING genome file, FASTA format (gzipped if end with '.gz')
 -o STRING Output file name (gzipped if end with '.gz')
 -0, --0-base 0-based genome if specified [Default: 1-based]
- **Example:**

The input CGmap file, which is lacking C context on the 3rd and 4th columns:

```
Chr1    C        3541    -        -        0.0    0        1
```

After `refill` processing, the CGmap file would be as below, added C context information:

```
Chr1    C        3541    CG        CG        0.0    0        1
```

3.4 intersect

- **Description:** Get the intersection of two CGmap files.
- **Usage:** `cgmmaptools intersect [-1 <CGmap_1>] -2 <CGmap_2> [-o <output>]`
 -1 CGmap File File name, end with .CGmap or .CGmap.gz.
 -2 CGmap File standard input if not specified
 -o OUTFILE To standard output if not specified. Compressed output if end with .gz
- **Example:**
 Suppose you have two CGmap file from two samples, the first one is:

```
Chr1    C    3541    CG    CG    0.8    4    5
```


and the second CGmap file is:

```
Chr1 C 3541 CG CG 0.4 4 10
```

After intersction, the output contains sites covered in both CGmap files. And the last three columns of the output are extracted from the second CGmap file:

```
Chr1 C 3541 CG CG 0.8 4 5 0.4 4 10
```

3.5 merge2

- **Description:** Merge two CGmap or ATCGmap files together.
- **Usage:** `cgmapttools merge2 <command> [options]`
- **Commands:**

atcgmap: merge two ATCGmap files into one.

Usage: `cgmapttools merge2 atcgmap -1 <ATCGmap> -2 <ATCGmap>`

-1 Input, 1st ATCGmap file

-2 Input, 2nd ATCGmap file

Output to STDOUT in ATCGmap format

Tips: Two input files should have the same order of chromosomes

cgmap: merge two CGmap files into one.

Usage: `cgmapttools merge2 cgmap -1 <CGmap_1> -2 <CGmap_2> [-o <output>]`

-1 FILE File name end with .CGmap or .CGmap.gz

-2 FILE If not specified, STDIN will be used.

-o OUTFILE CGmap, output file. Use STDOUT if omitted (gzipped if end with '.gz').

3.6 mergelist

- **Description:** merge a list of files.
- **Usage:** `cgmapttools mergelist <command> [options]`
- **Commands:**

tomatrix: Fill methylation levels according to the Index file for CGmap files in list.

Usage: `cgmapttools mergelist tomatrix [-i <index>] -f <IN1,IN2,...> -t <tag1,tag2,...> [-o output]`

-i FILE TXT file, index file, use STDIN if omitted

-f STRING List of (input) CGmap files (CGmap or CGmap.gz)

-t STRING List of tags, same order with '-f'

-c INT minimum coverage [default: 1]

-C INT maximum coverage [default: 200]

-o STRING Output file name (gzipped if end with '.gz')

tosingle: merge list of input files into one.

Usage: `cgmapttools mergelist tosingle -i f1,f2,...,fn [-o <output>]`

```

-i FILE      List of input files; gzipped file ends with '.gz'
-f FILE      cgmap or atcgmap [Default: cgmap]
-o OUTFILE   To standard output if not specified; gzipped file if end with
              '.gz'

```

3.7 sort

- **Description:** Sort the input files by chromosome and position.
- **Usage:** `cgmaptools sort [-i <input>] [-c 1] [-p 3] [-o output]`

```

-i FILE      File name end with .CGmap or .CGmap.gz.
              If not specified, STDIN will be used.
-c INT, --chr=INT  The column of chromosome [default: 1]
-p INT, --pos=INT  The column of position [default: 2]
-o OUTFILE      To standard output if not specified

```

3.8 split

- **Description:** Split the files by each chromosomes.
- **Usage:** `cgmaptools split -i <input> -p <prefix[.chr.]> -s <[.chr.]suffix>`

```

-i FILE      Input file, CGmap or ATCGmap format, use STDIN when not specified.
              (gzipped if end with 'gz').
-p STRING    The prefix for output file
-s STRING    The suffix for output file (gzipped if end with 'gz').

```

3.9 select

- **Description:** Split the files by each chromosomes.
- **Usage:** `cgmaptools select <command> [options]`
- **Commands:**

region: Lines in input CGmap/ATCGmap be selected/excluded by BED file. Strand is NOT considered. Output to STDOUT in same format with input.

Usage: `cgmaptools select region [-i <CGmap/ATCGmap>] -r <BED> [-R]`

```

-i Input, CGmap/ATCGmap file; use STDIN if not specified
  Please use "gunzip -c <input>.gz " and pipe as input for gzipped file.
  Ex: chr12  G  19898796  ...

```

```

-r Input, Region file, BED file to store regions
  At least 3 columns are required
  Ex: chr12 19898766 19898966 XX XXX XXX

```

```

-R [optional] Reverse selection. Sites in region file will be excluded when specified

```

site: Select lines from input CGmap/ATCGmap in index or reverse.

Usage: `cgmaptools select site -i <index> [-f <CGmap/ATCGmap>] [-r] [-o output]`

-i FILE Name of Index file required (gzipped if end with '.gz').
-r reverse selected, remove site in index if specified
-f STRING Input CGmap/ATCGmap files. Use STDIN if not specified
-o STRING CGmap, Output file name (gzipped if end with '.gz').

Chapter 4

SNP calling

Bisulfite sequencing data contains information of both methylation and genome sequences. In addition to DNA methylation analysis, we can also call variants using bisulfite data. Due to bisulfite conversion and PCR amplification during library preparation, the unmethylated cytosines on the DNA fragments would be converted to thymines. Thus, it's difficult to distinguish thymine produced by bisulfite conversion with the real thymine allele.

In recent years, few tools are adapted to bisulfite data for SNP calling. The main idea is removing vague reads that may contain unmethylated cytosines for a given position. Consequently, the rest reads can be regarded as reads generated from a normal genome DNA without bisulfite treatment and can be used to call variants using regular methods without consideration of bisulfite conversion.

However, removing the vague reads leads to information lost in most cases making variant calling less confident, especially when the sequencing depth is low. To solve this problem, we proposed two independent methods called BinomWC (based on binomial) and BayesWC (based on bayesian), taking vague reads into consideration.

- Usage: cgmaptools snv [-i <ATCGmap>] [-o <output> -v <VCF>]
-i FILE ATCGmap format, STDIN if not specified
-v FILE, --vcf=FILE VCF format file for output
-a, --all_nt Show all sites with enough coverage (-l). Only show SNP sites if not specified.
-o OUTFILE STDOUT if not specified
-m MODE, --mode=MODE Mode for calling SNP [Default: binom]
binom: binomial, separate strands
bayes: bayesian mode
--bayes-e=BAYES_ER (BayesWC mode) Error rate for calling a nucleotide [Default: 0.05]
--bayes-p=BAYES_PV (BayesWC mode) P value as cut-off [Default: 0.001]
--bayes-dynamicP (BayesWC mode) Use dynamic p-value for different coverages instead of specific p-value. (Recommended)
"--bayes-p" will be ignored if "--bayes-dynamicP" is specified.
--binom-e=BINOM_ER (BinomWC mode) Error rate for calling a nucleotide [Default: 0.05]
--binom-p=BINOM_PV (BinomWC mode) P value as cut-off [Default: 0.01]
--binom-cov=BINOM_COV (BinomWC mode) The coverage checkpoint [Default: 10]

Chapter 5

Methylation Analysis

5.1 dms

- **Description:** Get the differentially methylated sites between two samples.
- **Usage:** `cgmactools dms [-i <CGmapInter>] [-m 5 -M 100] [-o output]`

<code>-i FILE</code>	File name for CGmapInter, STDIN if omitted
<code>-m INT, --min=INT</code>	min coverage [default : 0]
<code>-M INT, --max=INT</code>	max coverage [default : 100]
<code>-o OUTFILE</code>	To standard output if omitted. Compressed output if end with .gz
<code>-t STRING, --test-method=STRING</code>	chisq, fisher [default : chisq]

- **Example:**

#1 Using the output of `intersect` as input:

```
Chr1 C 3541 CG CG 0.8 4 5 0.4 4 10
```

The output of `dms` is:

```
chr1 C 4654 CG CG 0.92 1.00 8.40e-01
chr1 C 4658 CHH CC 0.50 0.00 3.68e-04
chr1 G 8376 CG CG 0.62 0.64 9.35e-01
```

5.2 dmr

- **Description:** Get the differentially methylated region by Fisher's exact test.
- **Usage:** `cgmactools dmr [-i <CGmapInter>] [-m 5 -M 100] [-o output]`

<code>-i FILE</code>	File name for CGmapInter, STDIN if omitted
<code>-c INT, --minCov=INT</code>	min coverage [default : 0]
<code>-C INT, --maxCov=INT</code>	max coverage [default : 100]
<code>-s INT, --minStep=INT</code>	min step in bp [default : 100]
<code>-S INT, --maxStep=INT</code>	max step in bp [default : 500]
<code>-n INT, --minNSite=INT</code>	min N sites [default : 2]
<code>-o OUTFILE</code>	To standard output if omitted. Compressed output if end with .gz

- **Example:**

```
#1 Using the output of intersect as input:
```

```
Chr1 C 3541 CG CG 0.8 4 5 0.4 4 10
```

The output of dms is:

```
chr1 1004572 1004574 inf 0.00e+00 0.1100 0.0000
chr1 1009552 1009566 -0.2774 8.08e-01 0.0200 0.0300
chr1 1063405 1063498 0.1435 8.93e-01 0.6333 0.5733
```

5.3 asm

- **Description:** Allele specific methylation analysis.
- **Usage:** `cgmtools asm [options] -r <ref.fa> -b <input.bam> -l <snv.vcf>`
 - r Samtools indexed reference genome sequence, fasta format. eg. hg19.fa
- use samtools to index reference first: `samtools faidx hg19.fa`
 - b Samtools indexed Bam format file.
- use samtools to index bam file first: `samtools index <input.bam>`
 - l SNPs in vcf file format.
 - s Path to samtools eg. `/home/user/bin/samtools`
- by default, we try to search samtools in your system PATH,
 - o Output results to file. [default: STDOUT]
 - t C context. [default: CG]
- available context: C, CG, CH, CW, CC, CA, CT, CHG, CHH
 - m Specify calling mode. [default: asr]
- alternative: ass
- asr: allele specific methylated region
- ass: allele specific methylated site
 - d Minimum number of read for each allele linked site to call ass. [default: 3]
- ass specific.
 - n Minimum number of C site each allele linked to call asr. [default: 2]
- asr specific.
 - D Minimum read depth for C site to call methylation level when calling asr.
[default: 1]
- asr specific.
 - L Low methylation level threshold. [default: 0.2]
- allele linked region [or site] with low methylation level should be
no greater than this threshold.
 - H High methylation level threshold. [default: 0.8]
- allele linked region[or site] with high methylation level should be
no less than this threshold.
 - q Adjusted p value using Benjamini & Hochberg (1995)
("BH" or its alias "fdr"). [default: 0.05]

5.4 mbed

- **Description:** Calculate average methylation levels in given regions.
- **Usage:** `cgmtools mbed [-i <CGmap>] -b <regin.bed> [-c 5 -C 500 -s]`


```

-i String, CGmap file; use STDIN if not specified
  Ex: chr1    G    3000851 CHH CC  0.1 1    10
-b String, BED file
  Ex: chr1    3000000 3005000 -
-c Int, minimum Coverage [Default: 5]
-C Int, maximum Coverage [Default: 500]
-s Strands would be distinguished when specified

```

- **Example:**

The output format:

5.5 mbin

- **Description:** Generate the methylation level in Bins.

- **Usage:** `cgmactools mbin [-i <CGmap>] [-c 10 --CXY 5 -B 5000000]`

```

-i FILE                File name end with .CGmap or .CGmap.gz.
                        If not specified, STDIN will be used.
-B BIN_SIZE            Define the size of bins [Default: 5000000]
-c COVERAGE           The minimum coverage for site selection [Default: 10]
-C CONTEXT, --context=CONTEXT
                        specific context: CG, CH, CHG, CHH, CA, CC, CT, CW
                        use all sites if not specified
--cXY=COVERAGEXY      Coverage for chrX/Y should be half that of autosome
                        for male [Default: same with -c]
-f FIGTYPE, --figure-type=FIGTYPE
                        png, pdf, eps. Will not generate figure if not
                        specified
-p STRING              Prefix for output figures
-t STRING, --title=STRING
                        title in the output figures

```

- **Example:**

The output format:

```

chr1    1        5000    0.0000
chr1    5001     10000    0.0396
chr2    1        5000    0.0755
chr2    5001     10000    0.0027
chr3    1        5000    na

```

5.6 mmbin

- **Description:** Generate the methylation level in Bins for multiple samples.

- **Usage:** `cgmactools mmbin [-l <1.CGmap[,2.CGmap,...]>] [-c 10 --CXY 5 -B 5000000]`

```

-l FILE                File name list, end with .CGmap or .CGmap.gz. If not
                        specified, STDIN will be used.
-t FILE                List of samples
-B BIN_SIZE            Define the size of bins [Default: 5000000]

```

```
-C CONTEXT, --context=CONTEXT
                        specific context: CG, CH, CHG, CHH, CA, CC, CT, CW
                        use all sites if not specified
-c COVERAGE            The minimum coverage for site selection [Default: 10]
--cXY=COVERAGEXY       Coverage for chrX/Y should be half that of autosome
                        for male [Default: same with -c]
```

- **Example:**

The output format:

```
chr1    1      5000    0.0000
chr1    5001    10000    0.0396
chr2    1      5000    0.0755
chr2    5001    10000    0.0027
chr3    1      5000    na
```

5.7 mfg

- **Description:** Calculated methylation profile across fragmented regions.

- **Usage:** `cgmtools mfg [-i <CGmap>] -r <region> [-c 5 -C 500]`

```
-i String, CGmap file; use STDIN if not specified
    chr1    G    851 CHH CC  0.1 1    10
-r String, Region file, at least 4 columns
    Format: chr strand pos_1 pos_2 pos_3 ...
    Regions would be considered as [pos_1, pos_2), [pos_2, pos_3)
    Strand information will be used for distinguish sense/antisense strand
    Ex:
    chr1    +    600 700 800 900 950
    chr1    -    1600 1500 1400 1300 1250
-c Int, minimum Coverage [Default: 5]
-C Int, maximum Coverage [Default: 500]
    Sites exceed the coverage range will be discarded
```

- **Example:**

The output format:

Region_ID	R_1	R_2	R_3	R_4
sense_ave_mC	0.50	0.40	0.30	0.20
sense_sum_mC	5.0	4.0	3.0	2.0
sense_sum_NO	10	10	10	10
anti_ave_mC	0.40	0.20	0.10	NaN
anti_sum_mC	8.0	4.0	2.0	0.0
anti_sum_NO	20	20	20	0
total_ave_mC	0.43	0.27	0.17	0.2
total_sum_mC	13.0	8.0	5.0	2.0
total_sum_NO	30	30	30	10

5.8 mstat

- **Description:** Methylation statistic.

- **Usage:** cgmaptools mstat [-i <CGmap>]

```
-i FILE          File name end with .CGmap or .CGmap.gz. If not
                  specified, STDIN will be used.
-c COVERAGE     The minimum coverage for site selection [Default: 10]
-f FILE, --figure-type=FILE
                  png, pdf, eps. Will not generate figure if not
                  specified
-p STRING        Prefix for output figures
-t STRING, --title=STRING
                  title in the output figures
```

- **Example:**

The output format:

MethStat	context	C	CG	CHG	CHH	CA	CC	CT	CH	CW
mean_mC	global	0.0798	0.3719	0.0465	0.0403	0.0891	0.0071	0.0241	0.0419	0.0559
sd_mCbyChr	global	0.0078	0.0341	0.0163	0.0110	0.0252	0.0049	0.0076	0.0096	0.0148
count_C	global	10000	1147	2332	6521	3090	2539	3224	8853	6314
contrib_mC	global	1.0000	0.5348	0.1360	0.3292	0.3452	0.0228	0.0973	0.4652	0.4424
quant_mC	[0]	8266	471	2012	5783	2422	2421	2952	7795	5374
quant_mC	(0.00 ,0.20]	705	182	155	368	272	97	154	523	426
mean_mC_byChr	chr1	0.0840	0.4181	0.0340	0.0412	0.0794	0.0065	0.0251	0.0393	0.0513
mean_mC_byChr	chr10	0.0917	0.4106	0.0758	0.0421	0.0968	0.0097	0.0349	0.0502	0.0655

5.9 mtr

- **Description:** Calculate the methylation levels in regions in two ways.

- **Usage:** cgmaptools mtr [-i <CGmap>] -r <region> [-o <output>]

```
-i FILE          File name end with .CGmap or .CGmap.gz. If not specified, STDIN
                  will be used.
-r FILE          Filename for region file, support *.gz
-o OUTFILE       To standard output if not specified.
```

- **Example:**

The input file format:

#chr	start_pos	end_pos
chr1	8275	8429

The output format:

#chr	start_pos	end_pos	mean(mC)	#_C	#read(C)/#read(T+C)	#read(T+C)
chr1	8275	8429	0.34	72	0.40	164

Chapter 6

Coverage Analysis

6.1 oac

- **Description:** Overall coverage (for ATCGmap).
- **Usage:** `cgmactools oac <command> [options]`
- **Commands:**

bin: Overall coverage in bins.

Usage: `cgmactools oac bin [-i <ATCGmap>] [-B 5000000]`

`-i FILE` File name end with `.ATCGmap` or `.ATCGmap.gz`. If not specified, STDIN will be used.

`-B BIN_SIZE` Define the size of bins [Default: 5000000]

`-f FILE, --figure-type=FILE` png, pdf, eps. Will not generate figure if not specified

`-p STRING` Prefix for output figures

`-t STRING, --title=STRING` title in the output figures

stat: Get the distribution of overall coverages.

Usage: `cgmactools oac stat [-i <ATCGmap>]`

`-i FILE` File name end with `.ATCGmap` or `.ATCGmap.gz`. If not specified, STDIN will be used.

`-f FILE, --figure-type=FILE` png, pdf, eps. Will not generate figure if not specified

`-p STRING` Prefix for output figures

- **Example:**

The output format of `bin`:

chr1	1	5000	29.0000
chr1	5001	10000	30.0396
chr2	1	5000	35.0755
chr2	5001	10000	40.0027
chr3	1	5000	na

The output format of `stat`:

```
OverAllCov    global  47.0395
OverAllCov    chr1    45.3157
OverAllCov    chr10   47.7380
CovAndCount   1        1567
CovAndCount   2        655
CovAndCount   3        380
```

6.2 mec

- **Description:** Methylation effective coverage (for CGmap).
- **Usage:** `cgmmaptools mec <command> [options]`
- **Commands:**

bin: Generate the methylation-effective coverage in Bins.

Usage: `cgmmaptools mec bin [-i <CGmap>] [-B 5000000]`

```
-i FILE                File name end with .CGmap or .CGmap.gz. If not
                        specified, STDIN will be used.
-B BIN_SIZE            Define the size of bins [Default: 5000000]
-f FILE, --figure-type=FILE
                        png, pdf, eps. Will not generate figure if not
                        specified
-p STRING              Prefix for output figures
-t STRING, --title=STRING
                        title in the output figures
```

stat: Get the distribution of methylation-effective coverages.

Usage: `cgmmaptools mec stat [-i <CGmap>]`

```
-i FILE                File name end with .CGmap or .CGmap.gz. If not
                        specified, STDIN will be used.
-f FILE, --figure-type=FILE
                        png, pdf, eps. Will not generate figure if not
                        specified
-p STRING              Prefix for output figures
```

- **Example:**

The output format of `bin`:

```
chr1    1        5000    29.0000
chr1    5001     10000    30.0396
chr2    1        5000    35.0755
chr2    5001     10000    40.0027
chr3    1        5000     na
```

The output format of `stat`:

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OverAllCov    global  47.0395
OverAllCov    chr1    45.3157
OverAllCov    chr10   47.7380
CovAndCount   1        1567
CovAndCount   2        655
```

CovAndCount	3	380
-------------	---	-----

Chapter 7

Graphics

7.1 lollipop

- **Description:** Plot local mC level for multiple samples.

- **Usage:** `cgmtools lollipop [options] file`

```
-i INFILE, --infile=INFILE
    input file
-a ANNOTATION, --annotation=ANNOTATION
    [opt] sample name
-o OUTFILE, --outfile=OUTFILE
    [opt] output file
-f FORMAT, --format=FORMAT
    [opt] the format for output figure: pdf (default), png, eps
-l LEFT, --left=LEFT
    [opt] Left-most position
-r RIGHT, --right=RIGHT
    [opt] Right-most position
-c CHR, --chr=CHR
    [opt] chromosome name
-t TITLE, --title=TITLE
    [opt] text shown on title
-w WIDTH, --width=WIDTH
    [opt] width (in inch). Default: 8.
--height=HEIGHT
    [opt] height (in inch). Default: 8.
-s SITE, --site=SITE
    [opt] file of site to be marked
-b BED, --bed=BED
    [opt] BED file for region to be marked
```

- **Example:**

The input file format:

>= 3 columns, 1st line is the header, using R color name or “NaN”. Can be output by CGmapFillIndex.py. Use STDIN if omitted.

chr	pos	E_vs_EMT	EMT_vs_M	E_vs_M
chr1	13116801	NaN	NaN	darkgreen

```
chr1    13116899    NaN      red      NaN
```

The bed file format:

the first 4 columns are required.

```
chr1 213941196 213942363 REGION-1
chr1 213942363 213943530 REGION-2
```

7.2 heatmap

- **Description:** Plot methylation dynamics of target region for multiple samples heatmap.

- **Usage:** `cgmmaptools heatmap [options]`

```
-i INFILE, --infile=INFILE
    input file
-o OUTFILE, --outfile=OUTFILE
    [opt] output file name. [default: mCBinHeatmap.SysDate.pdf]
-c, --cluster
    [opt] cluster samples by methylation in regions. [default: FALSE]
-l COLORLOW, --colorLow=COLORLOW
    [opt] color used for the lowest methylation value. [default: cyan3]
-m COLORMID, --colorMid=COLORMID
    [opt] color used for the middle methylation value. [default: null]
-b COLORHIGH, --colorHigh=COLORHIGH
    [opt] color used for the highest methylation value. [default: coral2]
-n COLORNUMBER, --colorNumber=COLORNUMBER
    [opt] desired number of color elements in the panel. [default: 10]
-W WIDTH, --width=WIDTH
    [opt] width of figure (inch). [default: 7]
-H HEIGHT, --height=HEIGHT
    [opt] height of figure (inch). [default: 7]
-f FORMAT, --format=FORMAT
    [opt] format of output figure. Alternative: png. [default: pdf]
-R RESOLUTION, --resolution=RESOLUTION
    [opt] Resolution in ppi. Only available for png format. [default: 300]
```

- **Example:**

The input file format:

The 1st line is the header. Each column contains methylation measurements of a sample.

```
Region Sample1 Sample2 ...
Region1 0.1      0.1      ...
Region2 0.1      0.1      ...
```

7.3 fragreg

- **Description:** Plot methylation dynamics of target and flanking region for multiple samples.

- **Usage:** `cgmmaptools fragreg [options]`

```
-i INFILE, --infile=INFILE
    input file
```

```

-r RATIO, --ratio=RATIO
    [opt] range ratio between target region and flanking region in plot. [default: 5]
-o OUTFILE, --outfile=OUTFILE
    [opt] output file name. [default: FragRegView.SysDate.pdf]
-W WIDTH, --width=WIDTH
    [opt] width of figure (inch). [default: 7]
-H HEIGHT, --height=HEIGHT
    [opt] height of figure (inch). [default: 7]
-f FORMAT, --format=FORMAT
    [opt] format of output figure. Alternative: png. [default: pdf]
-R RESOLUTION, --resolution=RESOLUTION
    [opt] Resolution in ppi. Only available for png format. [default: 300]

```

- **Example:**

The input file format:

The 1st line is the header. Each row contains methylation measurements of a sample.

Sample	Up1	Up2	...	Region1	Region2	...	Down1	Down2	...
Sample1	0.1	0.1	...	0.2	0.2	...	0.3	0.3	...
Sample2	0.1	0.1	...	0.2	0.2	...	0.3	0.3	...

7.4 tanghulu

- **Description:** Show local mapped reads in Tanghulu shape.
- **Usage:** `cgmmaptools tanghulu [options] -r <ref> -b <bam> -l chr1:133-144`

```

-r      Samtools indexed reference genome sequeunce, fasta format. eg. hg19.fa
        - use samtools to index reference: samtools faidx <hg19.fa>
-b      Samtools indexed Bam file to view.
        - use samtools to index bam file: samtools index <input.bam>
-l      Region in which to display DNA methylation.
        - or specify a single position (eg. heterozygous SNP site),
          we will show allele specific methylation.
-s      Path to samtools eg. /home/user/bin/samtools
        - by default, we try to search samtools in your system PATH.
-o      Output results to file [default: CirclePlot.Ctype.region.Date.pdf].
-t      C context. [default: CG]
        - available context: C, CG, CH, CW, CC, CA, CT, CHG, CHH
-d      Ouput device. [default: pdf]
        - alternative: png
-c      Seperate reads by chain. [default: OFF]
        - specify this option to turn ON.
-v      Show vague allele linked reads. [ default: OFF]
-g      Genotype of heterozygous SNP site.
        - This option provides two alleles of htSNP site. eg. AT
        - The genotype information can be used to reduce vague alleles.
        - This option is specific to display methylation in allele specific mode.
-D      Minimum number of reads (depth) covered in this region or allele linked.
        [default: 0|OFF]
-C      Minimum number of C (specified type) covered in this region or allele
        linked. [default: 0|OFF]
-W      Width of graphics reigon in inches. [default: 4]

```

- H Height of graphics reigon in inches. [default: 4]
- R Resolution in ppi. [default: 300]
 - only available for png device.

Chapter 8

Other Utilities

8.1 findCCGG

- **Description:** Get MspI cutting sites for RRBS.
- **Usage:** `cgmtools findCCGG -i <genome.fa> [-o <output>]`
`-i INFILE, --infile=INFILE`
`-i FILE` Genome sequence file in Fasta format
`-o FILE` Name of the output file (standard output if not specified).Format: chr cCgg_pos ccGg_pos (0-base)
- **Example:**

The output file format:

```
chr1    4025    5652
chr1    8274    8431
```

8.2 bed2fragreg

- **Description:** Generate fragmented regions from BED file.
- **Usage:** `cgmtools bed2fragreg [-i <BED>] [-n <N>] [-F <50,50,...> -T <50,...>] [-o output]`
`-i FILE` BED format, STDIN if omitted
`-F INT_list` List of region lengths in upstream of 5' end, Ex: 10,50. List is from 5'end->3'end
`-T INT_list` List of region lengths in downstream of 3' end, Ex: 40,20. List is from 5'end->3'end
`-n INT` Number of bins to be equally split [Default:1]
- **Example:**

The input file format:

```
chr1    1000    2000    +
chr2    9000    8000    -
```

The output file format:

chr1	+	940	950	1000	1200	1400	1600	1800	1850
chr2	-	9060	9050	9000	8800	8600	8400	8200	8150