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Chapter 1

What is CGmapTools

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.

We proposed **CGmapTools**, 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 performence in SNP calling. We also provide dozens of utilities and a seamless pipeline for bisulfite sequencing data analysis.

```
cgmaptools -h
```

```
Program : cgmaptools (Tools for analysis in CGmap/ATCGmap format)
#
#
    Usage:
              cgmaptools <command> [options]
#
    Commands:
      -- File manipulation
#
#
                     + data format conversion tools
         convert
#
         fetch
                     + fetch a region by random accessing
#
         refill
                       refill the missing columns
#
                       intersect two files
         intersect
#
         merge2
                     + merge two files into one
#
         mergelist
                     + merge a list of files
#
                       sort lines by chromosome and position
         sort
#
         split
                     + split file by chromosomes
#
                     + select lines by region/site
         select
#
      -- SNV analysis
#
         snv
                        snv analysis
#
      -- Methylation analysis
#
         dms
                       differentially methylated site analysis
#
                        differentially methylated region analysis
         dmr
#
                        allele-specific methylation analysis
         asm
                       average methylation level in regions
         mbed
#
         mbin
                     * single sample, mC levels in bins
#
                       multiple samples, mC levels in bins
         mmbin
#
                       methlation levels across fragmented region
         mfg
         mstat
                     * methyaltion statistic
#
                        methylation level to each region
         mtr
```

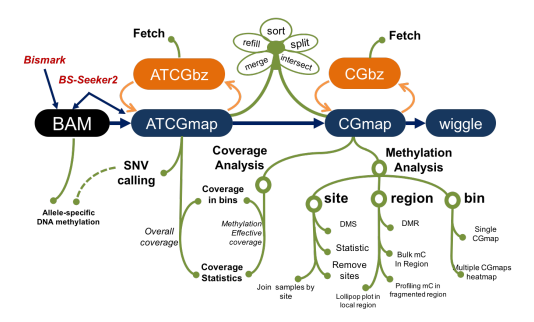


Figure 1.1: Schematic diagram of CGmapTools

```
#
      -- Coverage analysis
#
                    +* overall coverage (for ATCGmap)
         oac
#
         mec
                    +* methylation effective coverage (for CGmap)
      -- Graph related functions
#
                     * show local mC levels as lollipop bars
#
         lollipop
#
         heatmap
                     * global mC distribution for multiple samples
#
                     * show mC profile across fragmented regions
         fragreg
#
                     * show local mapped reads in Tanghulu shape
         tanghulu
#
      -- Other Utils
#
                       get MspI cutting sites for RRBS
         findCCGG
#
         bed2fragreg
                       get fragmented region based on region
#
#
      Commands support figures generation are marked with "*"
#
      Commands contain sub-commands are marked with "+"
#
   Authors:
      GUO, Weilong; guoweilong@126.com; http://guoweilong.github.io
#
#
      ZHU, Ping; pingzhu.work@gmail.com; http://perry-zhu.github.io
```

Chapter 2

File Formats

To facilitate high throughput data manipulation and reduce storage usage, several file format have been proposed and generaly 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 custermized tools, which is time comsuming and error prone.

The widely-used BS-seq alignment software **BS-Seeker2** defines **CGmap** and **ATCGmap** file formats for the representation of DNA methylomes. In CGmapTools, we used **ATCGmap** and **CGmap** as the standard file format interface, so that to simplify the development of downstream DNA methylation analysis tools and to provide standard formats for storing and sharing the DNA methylomes.

In CGmapTools, we designed novel binary formats: CGbz and ATCGbz for less coverage and improvements in random-accessing data in large data in hard-disk.

2.1 ATCGmap Format

Similar with **pileup**, **ATCGmap** format summarizes the information of mapped reads covered on each nucleotide on both strands, specially designed for BS-seq data.

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.

Example

```
chr1
         Τ
             3009410 --
                                0
                                     10
                                         0
                                                            3
                                                                     0
                                                                              na
             3009411 CHH CC
                                     10
                                         0
                                                       0
                                                            4
                                                                     0
                                                                          0
                                                                              0.0
chr1
         C
                                0
         С
             3009412 CHG CC
                                                       0
                                                                          0
chr1
                                0
                                     10
                                         0
                                              0
                                                   0
                                                            9
                                                                 1
                                                                     0
                                                                              0.0
             3009413 CG CG
                                     10
                                         50
chr1
                                                                              0.83
```

• Column Description

2.2 CGmap Format

In cases we only want to retain DNA methylation on cytonsines 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.

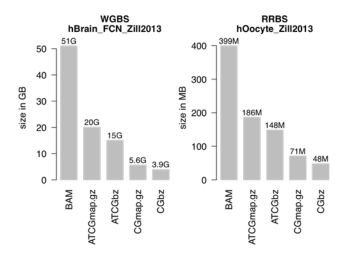


Figure 2.1: Size of multiple file formats

Col	Field	Type	Regexp/Range	Brief description
1	CHR	String	[!-?A-~]{ <mark>1,118</mark> }	Query template NAME
2	NUC	Char	[ATCGN-]	The nucleotide on reference genome
3	POS	Int	[0,2 <mark>32</mark> -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	[0,2 ¹⁴ -1]	Counts of reads on Watson strand support Adenine
7	WT	Int	[0,2 ¹⁴ -1]	Counts of reads on Watson strand support Thymine
8	WC	Int	[0,2 ¹⁴ -1]	Counts of reads on Watson strand support Cytosine
9	WG	Int	[0,2 ¹⁴ -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	[0,2 ¹⁴ -1]	Counts of reads on Crick strand support Adenine
12	СТ	Int	[0,2 ¹⁴ -1]	Counts of reads on Crick strand support Thymine
13	CC	Int	[0,2 ¹⁴ -1]	Counts of reads on Crick strand support Cytosine
14	CG	Int	[0,2 ¹⁴ -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"

Figure 2.2: Description of ATCGmap

2.3. ATCGBZ FORMAT 7

Col	Field	Туре	Regexp/Range	Brief description
1	CHR	String	[!-?A-~]{ <mark>1,118</mark> }	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

Figure 2.3: Description of CGmap

Fie	eld	Description	Туре	Value
N_chr		# chromosome	uint32_t	
		List of ChrInfo		
CHR		Name of chromosome	char [118]	
	count	# of ATCGbzT under this chromosome	uint32_t	
		List of ATCGbzT		
pos		Position on this chromosome	uint32_t	
	info	The mapping information	uint32_t [4]	

Figure 2.4: Data structure of ATCGbz

• Example

chr1	G	3000851	CHH	CC	0.1	1	10
chr1	C	3001624	CHG	CA	0.0	0	9
chr1	C	3001631	CG	CG	1.0	5	5
chr1	G	3001632	CG	CG	0 9	9	10

• Column Description

2.3 ATCGbz Format

ATCGbz format is the binary compressed version for ATCGmap format. ATCGmap format is readable, while quite large for storing, and difficult for fetching information in a specific position. ATCGbz is defined as the sorted binary version, that storing all information of ATCGmap into standard binary form, largely reduced the storage requirement, and also supporting fast retrival of methylation information for any position on genome.

- Data structure
- Related command

Command

```
\operatorname{cgmaptools} fetch \operatorname{atcgbz} -h
```

#

Usage: cgmaptools fetch atcgbz -b <ATCGbz> -C <CHR> -L <LeftPos> -R <RightPos>

	info : uint32_t[4] 128 bit													
	info [0]				info [1] info [2] info			info [3]	nfo [3]					
1	2,3	4	5-18	19-32	1-14	15-28	29-32	1-2	3-16	17-30	31-32	1-12	13-26	27-32
strand	Dinuc	Context	WA	WT	wc	WG	٧W	1	CA	СТ	C	2	CG	CN
0 = + 1 = -	00=CA 01=CC 10=CT 11=CG	=CA =CC 0=CNH 0=CT 0=CNG Count of reads mapped on Watson/Crick strands, supporting A. T. C. G or N												
	111 = ""	not CGG												

Figure 2.5: Data structure of info field of ATCGbz

Field		Description	Туре	Value
N_chr		# chromosome	uint32_t	
		List of ChrInfo		
CHR		Name of chromosome	char [118]	
	count	# of CGbzT under this chromosome	uint32_t	
		List of CGbzT		
	pos	Position on this chromosome	uint32_t	
	info	The mapping information	uint32_t	

Figure 2.6: Data structure of ATCGbz

```
(aka ATCGbzFetchRegion)
#
#
      Description: Convert ATCGbz format to ATCGmap format.
                    Guo, Weilong; guoweilong@126.com
#
      Contact:
      Last update: 2016-12-07
#
      Options:
        -h, --help
                                 output help information
        -b, --ATCGbz <arg> output ATCGbz file
-C, --CHR <arg> specify the chromos
#
#
                                 specify the chromosome name
        -L, --leftPos <arg> the left position
#
        -R, --rightPos <arg> the right position
```

2.4 CGbz Format

CGbz format is the binary compressed version for CGmap format.

- Data structure
- Related command

```
cgmaptools fetch cgbz -h
```

```
#
# Usage: cgmaptools fetch cgbz -b <CGbz> -C <CHR> -L <LeftPos> -R <RightPos>
```

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		info : u	int32_t		
1	2,3	4	5-18	19-32	
strand	Dinuc	Context	MC	NC	
0: + 1: -	00=CA 01=CC 0=CNH 10=CT 0=CNG 11=CG		# reads support methylated cytosine	# reads support all cytosine	
	111 = "" n	ot CGG	Cytosine		

Figure 2.7: Data structure of info field of CGbz

```
#
             (aka CGvzFetchRegion)
      Description: Convert CGbz file to CGmap format.
#
      Contact: Guo, Weilong; guoweilong@126.com
      Last update: 2016-12-07
#
#
      Options:
#
       -h, --help
                              output help information
       -b, --CGbz <arg>
#
                              output CGbz file
       -C, --CHR <arg>
                              specify the chromosome name
       -L, --leftPos <arg>
                              the left position
       -R, --rightPos <arg>
                              the right position
```

Chapter 3

File Manipulation

CGmapTools 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 coversion.
- Table of command for converting formats:

Commands	From	То
bam2cgmap	BAM	CGmap & ATCGmap
atcgmap2atcgbz	ATCGmap	ATCGbz
atcgbz2atcgmap	ATCGbz	ATCGmap
atcgmap2cgmap	ATCGmap	CGmap
$\operatorname{\mathbf{cgmap2cgbz}}$	CGamp	CGbz
$\operatorname{cgbz2cgmap}$	CGbz	CGmap
$\operatorname{\mathbf{cgmap2wig}}$	CGmap	WIG
bismark2cgmap	Bismark	CGmap

• Command

cgmaptools convert -h

```
cgmaptools convert <command> [options]
    Usage:
#
    Version:
              0.0.4
#
    Commands:
#
         bam2cgmap
                          BAM
                                  => CGmap & ATCGmap
#
         atcgmap2atcgbz
                          ATCGmap => ATCGbz
#
         atcgbz2atcgmap
                          ATCGbz => ATCGmap
#
         atcgmap2cgmap
                          ATCGmap => CGmap
#
         cgmap2cgbz
                          CGamp
                                  => CGbz
#
         cgbz2cgmap
                          CGbz
                                  => CGmap
#
         cgmap2wig
                          CGmap
                                  => WIG
         bismark2cgmap
                          Bismark => CGmap
```

• Example:

```
    BAM to CGmap

cgmaptools convert bam2cgmap -b WG.bam -g genome.fa --rmOverlap -o WG
  - BAM to CGmap
cgmaptools convert bam2cgmap -b RR.bam -g genome.fa --rm0verlap -o RR
  - ATCGmap to ATCGbz
cgmaptools convert atcgmap2atcgbz -c WG.ATCGmap.gz -b WG.ATCGbz

    ATCGvz to ATCGmap

cgmaptools convert atcgbz2atcgmap -c WG2.ATCGmap.gz -b WG.ATCGbz

    CGmap to CGbz

cgmaptools convert cgmap2cgbz -c RR.CGmap.gz -b RR.CGbz
  - CGbz to CGmap
cgmaptools convert cgbz2cgmap -c RR2.CGmap.gz -b RR.CGbz
  - CGmap to WIG
cgmaptools convert cgmap2wig -i <CGmap> [-w <wig>] [-c <INT> -b <float>]

    bismark output to CGmap

cgmaptools convert bismark2cgmap -i bismark.dat -o output.CGmap
Note: please refer to the help message for usage details using -h option.
```

3.2 fetch

- Description: Fastly acess methylation data in specified region.
- Command

```
cgmaptools fetch -h
```

```
# Usage: cgmaptools fetch <command> [options]
# Version: 0.0.4
# Commands:
# atcgbz fetch lines from ATCGbz
# cgbz fetch lines from CGbz
```

3.2.1 fetch cgbz

cgmaptools fetch cgbz -h

• Command

```
#
# Usage: cgmaptools fetch cgbz -b <CGbz> -C <CHR> -L <LeftPos> -R <RightPos>
# (aka CGvzFetchRegion)
# Description: Convert CGbz file to CGmap format.
# Contact: Guo, Weilong; guoweilong@126.com
# Last update: 2016-12-07
```

3.3. REFILL 13

```
#
#
     Options:
#
#
       -h, --help
                               output help information
#
       -b, --CGbz <arg>
                               output CGbz file
#
       -C, --CHR <arg>
                               specify the chromosome name
#
       -L, --leftPos <arg>
                               the left position
#
        -R, --rightPos <arg>
                               the right position
  • Example:
    cgmaptools fetch cgbz -b RR.CGbz -C chr3 -L 2200 -R 2400
```

3.2.2 fetch atcgbz

Command

```
cgmaptools fetch atcgbz -h
```

```
#
#
      Usage: cgmaptools fetch atcgbz -b <ATCGbz> -C <CHR> -L <LeftPos> -R <RightPos>
#
            (aka ATCGbzFetchRegion)
#
      Description: Convert ATCGbz format to ATCGmap format.
#
                   Guo, Weilong; guoweilong@126.com
#
      Last update: 2016-12-07
#
#
      Options:
#
#
        -h, --help
                               output help information
#
        -b, --ATCGbz <arg>
                               output ATCGbz file
#
        -C, --CHR <arg>
                               specify the chromosome name
#
        -L, --leftPos <arg>
                               the left position
#
        -R, --rightPos <arg>
                               the right position
  • Example:
```

cgmaptools fetch atcgbz -b WG.ATCGbz -C chr2 -L 90 -R 100

3.3 refill

• Command

```
cgmaptools refill -h
```

```
#
   Usage: cgmaptools refill [-i <CGmap>] -g <genome.fa> [-o output]
          (aka CGmapFillContext)
#
   Description: Fill the CG/CHG/CHH and CA/CC/CT/CG context.
#
#
                 Other fields will not be affected.
#
                 Can be applied to ATCGmap file.
                 Guo, Weilong; guoweilong@126.com;
   Contact:
   Last Update: 2016-12-07
#
#
   Index Ex:
#
       Chr1
               С
                       3541
                                               0.0
                                                                1
#
   Output Ex:
#
       Chr1
                       3541
                               CG
                                       CG
                                                       0
               C
                                               0.0
                                                                1
```

```
#
# Options:
# -h, --help show this help message and exit
# -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]
```

• File formats:

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

• Example:

```
zcat RR2.CGmap.gz | gawk -F"\t" -vOFS="\t" '{$4="-"; $5="-"; print;}' | cgmaptools
refill -g genome.fa -o RR3.CGmap.gz
```

3.4 intersect

Command

```
cgmaptools intersect -h
   Usage: cgmaptools intersect [-1 <CGmap_1>] -2 <CGmap_2> [-o <output>]
#
#
         (aka CGmapIntersect)
#
   Description:
#
       Get the intersection of two CGmap files.Contact: Guo, Weilong; guoweilong@126.com
#
   Last Update: 2016-08-18
#
   Output Format:
#
       Chr1 C 3541 CG CG 0.8 4 5 0.4 4 10
#
   When 1st CGmap file is:
#
       Chr1 C 3541 CG CG 0.8 4 5
#
    ,and 2nd CGmap file is:
#
       Chr1 C 3541 CG CG 0.4 4 10
#
#
   Options:
#
     -h, --help
                           show this help message and exit
#
     -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
#
     -C CONTEXT, --context=CONTEXT
#
                           specific context: CG, CH, CHG, CHH, CA, CC, CT, CW
```

• Example

#

cgmaptools intersect -1 WG.CGmap.gz -2 RR.CGmap.gz -C CG -o intersect CG.gz

use all sites if not specified

Output format

- Example

3.5. MERGE2 15

Col	Field	Туре	Regexp/Range	Brief description
1	CHR	String	[!-?A-~]{ <mark>1,118</mark> }	Query template NAME
2	NUC	Char	[ATCGN-]	The nucleotide on reference genome
3	POS	Int	[0,2 <mark>32</mark> -1]	1-based leftmost mapping position
4	CONT	String	{"", "CG", "CHG", "CHH"}	context
5	DINUC	String	{"", "CA", "CT", "CC", "CG"}	Dinucleotide context
6	METH_1	Float	[0,1] or "na"	Methylation level in Sample 1
7	MC_1	Int	[0,2 ¹² -1]	Counts of reads support methylated Cytosine in Sample 1
8	NC_1	Int	[0,2 ¹² -1]	Counts of reads support all Cytosine in Sample 1
9	METH_2	Float	[0,1] or "na"	Methylation level in Sample 2
10	MC_2	Int	[0,2 ¹² -1]	Counts of reads support methylated Cytosine in Sample 2
11	NC_2	Int	[0,2 ¹² -1]	Counts of reads support all Cytosine in Sample 2

Figure 3.1: Output format description for cgmaptools intersect

```
Chr1 C 3541 CG CG 0.8 4 5 0.4 4 10 Chr1 C 3542 CG CG 0.8 3 5 0.2 2 10 Chr1 C 3545 CHG CA 0.0 0 5 0.1 1 10
```

- Column Description

3.5 merge2

Command

```
cgmaptools merge2 -h

# Usage: cgmaptools merge2 <command> [options]
```

Version: 0.0.4

Commands:

atcgmap merge two ATCGmap files into one
cgmap merge two CGmap files into one

3.5.1 merge2 atcgmap

```
cgmaptools merge2 atcgmap -h
```

```
Unknown option: -h
#
   Usage: cgmaptools merge2 atcgmap -1 <ATCGmap> -2 <ATCGmap>
#
           (aka ATCGmapMerge)
                 Guo, Weilong; guoweilong@126.com;
#
   Contact:
#
   Last Update: 2016-12-07
#
   Options:
#
      -1
            Input, 1st ATCGmap file
            Input, 2nd ATCGmap file
#
      -2
#
   Output to STDOUT in ATCGmap format
   Tips: Two input files should have the same order of chromosomes
```

Example

cgmaptools merge2 atcgmap -1 WG.ATCGmap.gz -2 RR.ATCGmap.gz | gzip > merge.ATCGmap.gz

3.5.2 merge2 cgmap

Command

```
cgmaptools merge2 cgmap -h
   Usage: cgmaptools merge2 cgmap -1 <CGmap_1> -2 <CGmap_2> [-o <output>]
#
#
          (aka CGmapMerge)
#
   Description: Merge two CGmap files together.
                 Guo, Weilong; guoweilong@126.com
#
   Contact:
#
   Last Update: 2016-12-07
#
   Note: The two input CGmap files should be sorted in the same order first.
#
#
#
   Options:
#
      -h, --help show this help message and exit
#
     -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').
  • Example
  • Example command:
    cgmaptools merge2 cgmap -1 WG.CGmap.gz -2 RR.CGmap.gz | gzip > merge.CGmap.gz
```

3.6 mergelist

• Command

```
cgmaptools mergelist -h

# Usage: cgmaptools mergelist <command> [options]

# Version: 0.0.4

# Commands:

# tomatrix mC levels matrix from multiple files

# tosingle merge list of input files into one
```

3.6.1 mergelist tomatrix

cgmaptools mergelist tomatrix -h

Command

Index format Ex:

```
# Usage: cgmaptools mergelist tomatrix [-i <index>] -f <IN1,IN2,..> -t <tag1,tag2,..> [-o output]
# (aka CGmapFillIndex)
# Description: Fill methylation levels according to the Index file for CGmap files in list.
# Contact: Guo, Weilong; guoweilong@126.com;
# Last Updated: 2016-12-07
```

3.6. MERGELIST

Col	Field	Туре	Regexp/Range	Brief description
1	CHR	String	[!-?A-~]*	Query template NAME
2	POS	Int	[0,2 ³² -1]	Position

Figure 3.2: Format description for INDEX file

```
#
       chr10
               100005504
#
   Output format Ex:
#
       chr
               pos
                               tag2
                                       tag3
                       tag1
#
       Chr1
               111403 0.30
                               nan
                                       0.80
#
       Chr1
               111406 0.66
                               0.40
                                       0.60
#
#
   Options:
#
      -h, --help show this help message and exit
#
      -i FILE
                  TXT file, index file, use STDIN if omitted
#
     -f STRING
                 List of (input) CGmap files (CGmap or CGmap.gz)
#
                 List of tags, same order with '-f'
      -t STRING
      -c INT
#
                  minimum coverage [default: 1]
      -C INT
                  maximum coverage [default: 200]
      -o STRING
                  Output file name (gzipped if end with '.gz')
```

• Example

```
zcat RR*.CGmap.gz WG.CGmap.gz | gawk '$8>=5' | cut -f1,3 | sort -u | cgmaptools sort
-c 1 -p 2 > index
cgmaptools mergelist tomatrix -i index -f RR.CGmap.gz,RR2.CGmap.gz,WG.CGmap.gz -t
```

• Format for Index file

- Example

```
Chr1 940
Chr1 1840
Chr2 9060
```

- Column Description

• Format for output file

- Example

```
chr
                         tag2
        pos
                tag1
                                 tag3
Chr1
        111403 0.05
                         nan
                                 0.02
Chr1
        111500 1.00
                         0.80
                                 0.60
Chr2
        20000
                0.96
                         0.33
                                 0.66
```

RR,RR2,WG -c 5 -C 100 -o matrix.CG.gz

- Column Description

3.6.2 mergelist to single

Command

```
cgmaptools mergelist to single -h
```

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

Col	Field	Туре	Regexp/Range	Brief description
1	CHR	String	[!-?A-~]*	Query template NAME
2	POS	Int	[0,2 ³² -1]	Position
3	METH_1	Float	[0.00, 1.00]	Methylation level in sample 1
n	METH_n	Float	[0.00, 1.00]	Methylation level in sample n

Figure 3.3: Output format description for cgmaptools fill tomatrix

```
(aka MergeListOfCGmap)
   Description: Merge multiple CGmap/ATCGmap files into one.
#
                 Guo, Weilong; guoweilong@126.com
#
   Contact:
#
   Last Update: 2016-12-07
   Note: Large memory is needed.
#
#
#
   Options:
      -h, --help show this help message and exit
#
      -i FILE
                  List of input files; gzipped file ends with '.gz'
                  cgmap or atcgmap [Default: cgmap]
#
      -f FILE
      -o OUTFILE To standard output if not specified; gzipped file if end with
                  '.gz'
```

• Example

3.7 sort

• Command

```
cgmaptools sort -h
```

```
Usage: Sort_chr_pos [-i <input>] [-c 1] [-p 3] [-o output]
   Author: Guo, Weilong; guoweilong@gmail.com; 2014-05-11
   Last Update: 2016-12-07
#
   Description: Sort the input files by chromosome and position.
         The order of chromosomes would be :
#
#
         "chr1 chr2 ... chr11 chr11_random ... chr21 ... chrM chrX chrY"
#
#
   Options:
#
     -h, --help
                         show this help message and exit
#
     -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
```

• Example

```
zcat RR*.CGmap.gz WG.CGmap.gz | gawk '$8>=5' | cut -f1,3 | sort -u | cgmaptools sort
-c 1 -p 2 > index
```

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3.8 split

Command

```
cgmaptools split -h
    Usage: cgmaptools split -i <input> -p <prefix[.chr.]> -s <[.chr.]suffix>
#
          (aka CGmapSplitByChr)
   Description: Split the files by each chromosomes.
#
#
                 Guo, Weilong; guoweilong@126.com
   Contact:
#
   Last Update: 2016-12-07
#
#
   Options:
#
      -h, --help show this help message and exit
#
      -i FILE
                  Input file, CGmap or ATCGmap foramt, 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').

    Example

     cgmaptools split -i WG.CGmap.gz -p WG -s CGmap.gz
```

3.9 select

Command

```
# Usage: cgmaptools select <command> [options]
# Version: 0.0.4
# Commands:
# region select or exclude liens by region lists
# site select or exclude lines by site list
```

3.9.1 select region

• Command

```
cgmaptools select region -h
   Usage: cgmaptools select region [-i <CGmap/ATCGmap>] -r <BED> [-R]
#
          (aka CGmapSelectByRegion)
   Description: Lines in input CGmap/ATCGmap be selected/excluded by BED file.
#
                 Strand is NOT considered.
                 Output to STDOUT in same format with input.
#
#
                 Guo, Weilong; guoweilong@126.com
   Contact:
#
   Last Update: 2016-12-07
#
   Options:
#
      -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
    # -h help
    # Tips: program will do binary search for each site in regions
```

• Example

```
for CHR in 1 2 3 4 5; do (for P in 1 2 3 4 5; do echo | gawk -vC=$CHR -vP=$P -vOFS="\t" '{print "chr"C, P*1000, P*1000+200, "+";}'; done); done > region.bed
zcat WG.CGmap.gz | cgmaptools select region -r region.bed | head
```

3.9.2 select site

```
cgmaptools select site -h
```

```
Usage: cgmaptools select site -i <index> [-f <CGmap/ATCGmap>] [-r] [-o output]
          (aka CGmapSelectBySite)
#
   Description: Select lines from input CGmap/ATCGmap in index or reverse.
                 Guo, Weilong; guoweilong@126.com
   Last Update: 2016-12-07
   Index format example:
#
      chr10
             100504
#
      chr10
             103664
#
#
   Options:
#
     -h, --help show this help message and exit
#
                 Name of Index file required (gzipped if end with '.gz').
     -i FILE
#
                  reverse selected, remove site in index if specified
     -\mathbf{r}
     -f STRING Input CGmap/ATCGmap files. Use STDIN if not specified
     -o STRING
                 CGmap, Output file name (gzipped if end with '.gz').
  • Example
```

```
gawk 'NR%100==50' index > site
cgmaptools select site -f RR.CGmap.gz -i site -o RR_select.CGmap.gz
```

Chapter 4

SNV 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 coversion 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 coversion 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 tried to introduce wild-card in genotype calling. Even for these amiguouse genotypes, we can still learning something.

We proposed two independent methods called BinomWC (based on binomial) and BayesWC (based on bayesian), taking vague reads into consideration.

4.1 BaysWC strategy

BinomWC strategy combines bayesian method and wildcard strategy for predicting the genotype. The likelyhood matrix is designed as following.

Support the	AT	CGm	ap tak	ole
nucleotide on reference genome	Α	Т	С	G
Read count on Watson strand	$A_w^{\#}$	$T_w^{\#}$	$C_w^{\#}$	$G_w^{\#}$
Read count on Crick strand	$A_c^{\#}$	$T_c^{\#}$	$C_c^{\#}$	$G_c^{\#}$

Figure 4.1: ATCGmap table used for SNV calling

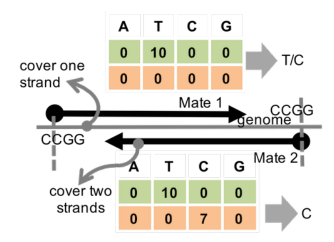


Figure 4.2: Example that alignments refer to vague genotypes

Ambiguous GN symbol	Possible genotypes	Hete- or Homo- zygous	sure to be SNV if reference is
Y	TT / TC / CC	not sure	A, G
R	AA / AG / GG	not sure	T, C
A,Y	AT / AC	heterozygous	A, T, C, G
C,Y	CT / CC	not sure	A, T, G
G,Y	GT / GC	heterozygous	A, T, C, G
T,Y	TT / TC	not sure	A, C, G
A,R	AA / AG	not sure	T, C, G
C,R	CA / CG	heterozygous	A, T, C, G
G,R	GA / GG	not sure	A, T, C
T,R	TA / TG	heterozygous	A, T, C, G

The wildcard characters are defined as: Y=T/C and R=A/G

Figure 4.3: Table for definition of amibiguous genotype

$Pr(I^{\#}=1 \mid g)$	g = A	g = T	g = C	g = G
$A_{w}^{\#} = 1$	p	e	e	e
$T_{\rm w}^{\#}=1$	2e	p + e	p + e	2e
$C_{\rm w}^{\#}=1$	e	e	p	e
$G_{\mathrm{w}}^{\#}=1$	e	e	e	p
$A_c^{\#}=1$	p + e	2e	2e	p + e
$T_{c}^{\#}=1$	e	p	e	e
$C_c^{\#} = 1$	e	e	p	e
$G_c^\#=1$	e	e	e	p

Figure 4.4: The likelyhood matrix for BayesWC strategy

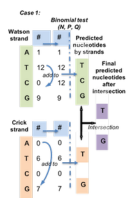


Figure 4.5: Case 1 for BinomWC strategy

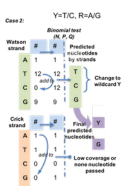


Figure 4.6: Case 2 for BinomWC strategy

4.2 BinomWC strategy

BinomWC (Binomial-WildCard) strategy works as following for 3 cases.

4.3 Performance

Performances on simulation data

```
cgmaptools snv -h
    Usage: cgmaptools snv [-i <ATCGmap>] [-o <output> -v <VCF>]
#
          (aka SNVFromATCGmap)
#
    Description: Predict the SNV from ATCGmap file.
#
    Contact:
                 Guo, Weilong; guoweilong@126.com
#
    Last update: 2017-08-24
#
    Output format example:
#
       #chr nuc pos
                         ATCG_watson ATCG_crick predicted_nuc p_value
#
       chr1 G
                  4752
                         17,0,0,69
                                      0,0,0,0
                                                  A,G
                                                                 9.3e-07
                                                                 0.0e+00
#
       chr1 A
                  4770
                         40,0,0,29
                                      0,0,0,0
                                                  A,G
#
       chr1 T
                  8454
                         0,39,0,0
                                      0,0,0,0
                                                  T/C
                                                                 1.00e-01
#
#
```

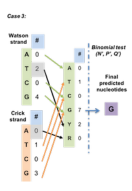


Figure 4.7: Case 3 for BinomWC strategy

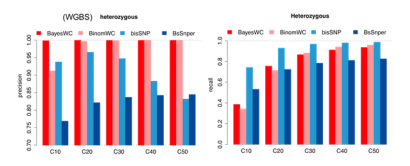


Figure 4.8: Precision-Recall analysis on simulated WGBS data

```
#
   Options:
#
      -h, --help
                            show this help message and exit
                            ATCGmap format, STDIN if not specified
#
      -i FILE
#
      -v FILE, --vcf=FILE
                            VCF format file for output
#
      -a, --all_nt
                            Show all sites with enough coverage (-1). Only show
#
                            SNP sites if not specified.
#
                            STDOUT if not specified
      -o OUTFILE
#
      -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 install of specific p-value. (Recomended)
#
                             "--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]
```

• Example commands :

cgmaptools snv -i WG.ATCGmap.gz -m bayes -v bayes.vcf -o bayes.snv --bayes-dynamicP cgmaptools snv -i WG.ATCGmap.gz -m binom -o binom.snv

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Col	Field	Туре	Regexp/Range	Brief description
1	CHR	String	[!-?A-~]{ <mark>1,118</mark> }	Query template NAME
2	NUC	Char	[ATCGN-]	The nucleotide on reference genome
3	POS	Int	[0,2 <mark>32</mark> -1]	1-based leftmost mapping position
4	W_Count	String	[0-9,]+	Count of reads support A, T, C, G on Watson strand, seprated by ", "
5	C_Count	String	[0-9,]+	Count of reads support A, T, C, G on Crick strand, seprated by ", "
6	PRDNUC	String	[ATCG./]+	Predicted genotype ("," indicate two allele; "/" means "or")
7	PVALUE	Float	[0, 1]	P_value for confidence of this prediction

Figure 4.9: Output format description for cgmaptools snv

• Output format

- Example

#chr	nuc	pos	ATCG_watson	ATCG_crick	<pre>predicted_nuc</pre>	p_value
chr1	G	4752	17, 0, 0, 69	0, 0, 0, 0	A,G	9.3e-07
chr1	Α	4770	40, 0, 0, 29	0, 0, 0, 0	A,G	0.0e+00
chr1	T	8454	0, 39, 0, 0	0, 0, 0, 0	T/C	1.00e-01

⁻ Column Description

Chapter 5

Methylation Analysis

The CGmap Tools supports both differentially methylated site (DMS) analyses and differentially methylated region (DMR) analyses are supported.

As the current available DNA methylome are either low coverage (such as WGBS) or fragmented in covered region (such as RRBS). In *CGmapTools*, we proposed a novel method **dynamic fragmentation strategy** for identifying DMRs between a pair of CGmap files.

5.1 dms

Differentially methylated site analysis, supporting *Chi-square* and *Fisher* tests.

```
cgmaptools dms -h
    Usage: cgmaptools dms [-i <CGmapInter>] [-m 5 -M 100] [-o output]
#
#
          (aka CGmapInterDiffSite)
#
   Description:
#
     Get the differentially methylated sites for two samples.
                 Guo, Weilong; guoweilong@126.com
#
   Last Update: 2017-01-20
#
#
   Input Format, same as the output of CGmapIntersect.py:
#
       Chr1 C 3541 CG CG 0.8 4 5 0.4 4 10
#
   Output Format:
#
                        CG CG 0.92
                                        1.00
                                                8.40e-01
       chr1 C
                4654
#
       chr1 C
               4658
                        CHH CC 0.50
                                        0.00
                                                3.68e-04
#
       chr1 G
              8376
                        CG CG 0.62
                                        0.64
                                                9.35e-01
#
#
   Options:
     -h, --help
                            show this help message and exit
#
     -i FILE
                            File name for CGmapInter, STDIN if omitted
                            min coverage [default : 0]
#
     -m INT, --min=INT
     -M INT, --max=INT
                            max coverage [default : 100]
     -o OUTFILE
                            To standard output if omitted. Compressed output if
#
                            end with .gz
#
     -t STRING, --test-method=STRING
                            chisq, fisher [default : chisq]
```

Col	Field	Туре	Regexp/Range	Brief description
1	CHR	String	[!-?A-~]{ <mark>1,118</mark> }	Query template NAME
2	NUC	Char	[ATCGN-]	The nucleotide on reference genome
3	POS	Int	[0,2 ³² -1]	1-based position
4	CONT	String	{"", "CG", "CHG", "CHH"}	context
5	DINUC	String	{"", "CA", "CT", "CC", "CG"}	Dinucleotide context
6	METH_1	Float	[0,1] or "na"	Methylation level in Sample 1
9	METH_2	Float	[0,1] or "na"	Methylation level in Sample 2
11	PVALUE	Float	[0, 1]	P-value

Figure 5.1: Output format description for cgmaptools dms

• Example

cgmaptools dms -i intersect_CG.gz -m 4 -M 100 -o DMS.gz -t fisher

• Output format

- Example

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

- Column Description

5.2 dmr

Differentially methylated region analysis, using dynamic fragmentation strategy.

• Command

```
cgmaptools dmr -h
```

```
#
   Usage: cgmaptools dmr [-i <CGmapInter>] [-m 5 -M 100] [-o output]
#
          (aka CGmapInterDiffReg)
#
   Description:
#
      Get the differentially methylated sites by Fisher's exact test.
                  Guo, Weilong; guoweilong@126.com;
#
   Author:
   Last Updated: 2017-08-12
#
#
    Input Format, same as the output of CGmapIntersect.py:
       chr1 C 3541 CG CG 0.8 4 5 0.4 4 10
#
#
   Output Format, Ex:
#
      #chr
                start
                        end
                                                    mC_A
                                                            mC_B
                                                                    N_site
                                        pv
#
       chr1 1004572 1004574 inf
                                    0.00e+00
                                                0.1100 0.0000 20
#
       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
#
#
#
   Options:
#
     -h, --help
                            show this help message and exit
#
     -i FILE
                            File name for CGmapInter, STDIN if omitted
      -c INT, --minCov=INT min coverage [default : 4]
#
```

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Col	Field	Туре	Regexp/Range	Brief description
1	CHR	String	[!-?A-~]{1,118}	Query template NAME
2	POS_L	Int	[0,2 <mark>32</mark> -1]	leftmost position for region
3	POS_R	Int	[0,2 ³² -1]	rightmost mapping position for region
4	Т	Float	[0,2 ³² -1) or "inf"	Statistics of T test
5	PV	Float	[0, 1]	P-value of t test
6	METH_1	Float	[0,1] or "na"	Methylation level in Sample 1
7	METH_2	Float	[0,1] or "na"	Methylation level in Sample 2

Figure 5.2: Output format description for cgmaptools dmr

```
-C INT, --maxCov=INT max coverage [default : 500]
#
#
      -s INT, --minStep=INT
#
                            min step in bp [default : 100]
#
      -S INT, --maxStep=INT
#
                            max step in bp [default : 1000]
      -n INT, --minNSite=INT
#
                            min N sites [default : 5]
#
      -o OUTFILE
                            To standard output if omitted. Compressed output if
                             end with .gz
```

• Example

cgmaptools dmr -i intersect_CG.gz -o DMR.gz

• Output format

- Example

```
1004572\ 1004574\ inf
                                0.00e+00
                                            0.1100 0.0000
chr1
        1009552 1009566 -0.2774 8.08e-01
chr1
                                            0.0200 0.0300
chr1
        1063405 1063498 0.1435 8.93e-01
                                            0.6333 0.5733
        1082130 1082133 -0.0822 9.42e-01
chr1
                                            0.5000 0.5550
chr1
        1123931 1123933 inf
                                0.00e+00
                                            0.0600 0.0000
```

- Column Description

• Dynamic Fragment Strategy

5.3 asm

Feeding with the precisely predicted heterozygous SNVs (by $\operatorname{cgmaptools}$ snv), $\operatorname{\mathbf{CGmapTools}}$ can identify Allele-Specific Methylated $(\operatorname{\mathbf{ASM}})$ regions from BAM files.

Following showed an interesting ASM region by analysing a previous cohort (Weilong Guo, et al., Scientific Report, 2016).

• Command

```
cgmaptools asm -h
```

```
# DESCRIPTION
```

```
# Allele specific methylated region/site calling
```

* Fisher exact test for site calling.

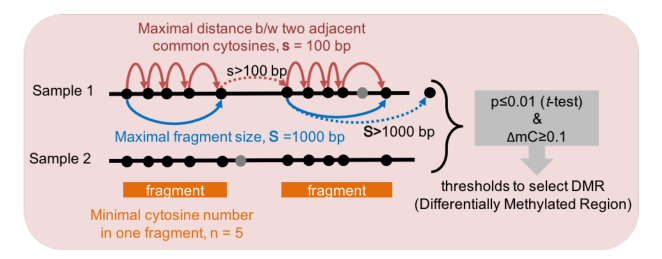


Figure 5.3: Dynamic Fragmentation Strategy

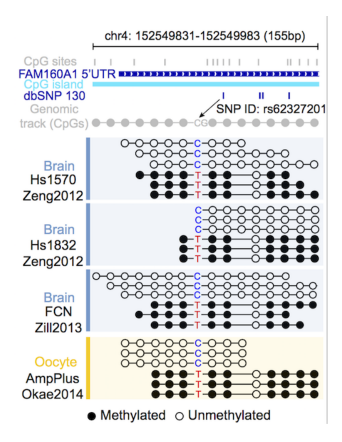


Figure 5.4: Examples showed allele-specific methylated region reported by CGmapTools

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```
* Students' t-test for region calling.
#
#
   USAGE
#
            cgmaptools asm [options] -r <ref.fa> -b <input.bam> -l <snp.vcf>
#
            (aka ASM)
#
#
            Options:
#
            -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>
#
                  SNPs in vcf file format.
            -1
#
                  Path to samtools eg. /home/user/bin/samtools
            -s
#
                  - by defualt, we try to search samtools in your system PATH,
#
                  Output results to file. [default: STDOUT]
            -0
#
                  C context. [default: CG]
            -t
#
                  - available context: C, CG, CH, CW, CC, CA, CT, CHG, CHH
#
                  Specify calling mode. [default: asr]
            -m
#
                  - alternative: ass
#
                  - asr: allele specific methylated region
#
                  - ass: allele specific methylated site
#
                  Minimum number of read for each allele linked site to call ass. [default: 3]
            -d
#
                  - ass specific.
                  Minimum number of C site each allele linked to call asr. [default: 2]
#
            -n
#
                  - 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
#
                  High methylation level threshold. [default: 0.8]
            -H
#
                  - allele linked region[or site] with high methylation level should be no less than th
#
                  Adjusted p value using Benjamini & Hochberg (1995) ("BH" or its alias "fdr"). [defaul
            -q
#
                  Help message.
            -h
#
#
   AUTHOR
#
            Contact:
                         Zhu, Ping; pingzhu.work@gmail.com
#
            Last update: 2016-12-07

    Example

    gawk '{if(/^#/){print}else{print "chr"$0;}}' bayes.vcf > bayes2.vcf
     cgmaptools asm -r genome.fa -b WG.bam -l bayes2.vcf > WG.asm
  • Output format for ASS (Allele-Specific methylated Site)
       - Example
    Chr
            SNP Pos Ref
                            Allele1
                                        Allele2
                                                      C Pos
                                                              Allele1_linked_C
                                                                                   Allele2_linked_C
                                                                                                       Al:
    Chr1
             8949221 T
                               Τ
                                                          8949252 30,2
                                                                          6,0 0.94
                                                                                       1.00
                                                                                                     1.00
                                             Α
    Chr1
             8965481 A
                                             Т
                                                             8965494 12,3
                                                                                       0.80
                               Α
                                                                               12,4
                                                                                               0.75
       - Column Description
  • Output format for ASR (Allele-Specific methylated Region)
       - Example
```

Allele1_linked_C Allele2_linked_C Allele1

Pos

Allele1

Allele2

Ref

Chr

Col	Field	Туре	Regexp/Range	Brief description
1	CHR	String	[!-?A-~]{ <mark>1,118</mark> }	Query template NAME
2	SNP_Pos	Int	[0,2 ³² -1]	1-based leftmost mapping position of SNP site
3	Ref	Char	[ATCGN-]	The nucleotide on reference genome
4	Allele1	Char	[ATCG]	The nucleotide of allele1
5	Allele2	Char	[ATCG]	The nucleotide of allele2
6	C_Pos	Int	[0,2 <mark>32</mark> -1]	1-based leftmost mapping position of C site
7	Allele1_linked_C	Int	[0, 2 ³² -1]	Comma separated number of reads support methylated and unmethylated C linked by allele1 respectively
8	Allele2_linked_C	Int	[0, 2 ³² -1]	Comma separated number of reads support methylated and unmethylated C linked by allele2 respectively
9	Allele1_linked_C_met	Float	[0,1]	Methylation level of allele1 linked C site
10	Allele2_linked_C_met	Float	[0,1]	Methylation level of allele2 linked C site
11	pvalue	Float	[0,1]	P value of t test
12	fdr	Float	[0,1]	Adjusted p value using Benjamini & Hochberg method
13	ASM	Logical	TRUE/FALSE	TRUE indicates this C site is allele specific methylated. FALSE otherwise.

Figure 5.5: Output format description for cgmaptools asm -m ass

chr1	8943402	Α	Α	T	1-1	0.8-1
chr1	8966879	C	С	G	0.93-0-0 0.81-0-0	0.31

- Column Description

5.4 mbed

The cgmaptools mbed command will calculated one DNA methylation level for all the investiaged regions, which is different from cgmaptools mtr.

For example, this function can be applied when calculating the average DNA methylation levels in regions, such as promoter, gene body, specific Transposon Elements (TEs).

• Command

cgmaptools mbed -h

```
Usage: cgmaptools mbed [-i <CGmap>] -b <regin.bed> [-c 5 -C 500 -s]
          (aka CGmapMethylInBed)
#
   Description: Calculated bulk average methylation levels in given regions.
#
   Contact:
                Guo, Weilong; guoweilong@126.com
   Last Update: 2017-01-20
#
#
   Options:
#
       -i String, CGmap file; use STDIN if not specified
          Please use "gunzip -c <input>.gz " and pipe as input for gzipped file.
                      3000851 CHH CC 0.1 1 10
#
          Ex: chr1 G
#
       -b String, BED file, should have at least 4 columns
          Ex: chr1 3000000 3005000 -
       -c Int, minimum Coverage [Default: 5]
      -C Int, maximum Coverage [Default: 500]
```

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Col	Field	Туре	Regexp/Range	Brief description
1	CHR	String	[!-?A-~]{1,118}	Query template NAME
2	Pos	Int	[0,2 ³² -1]	1-based leftmost mapping position of SNP site
3	Ref	Char	[ATCGN-]	The nucleotide on reference genome
4	Allele1	Char	[ATCG]	The nucleotide of allele1
5	Allele2	Char	[ATCG]	The nucleotide of allele2
6	Allele1_linked_C	Float	[0,1]	'-' separated methylation level of C sites linked by allele1
7	Allele2_linked_C	Float	[0,1]	'-' separated methylation level of C sites linked by allele2
9	Allele1_linked_C_met	Float	[0,1]	Average methylation level of allele1 linked C sites
10	Allele2_linked_C_met	Float	[0,1]	Average methylation level of allele2 linked C sites
11	pvalue	Float	[0,1]	P value of t test
12	fdr	Float	[0,1]	Adjusted p value using Benjamini & Hochberg method
13	ASM	Logical	TRUE/FALSE	TRUE indicates this region is allele specific methylated. FALSE otherwise.

Figure 5.6: Output format description for cgmaptools as $\operatorname{-m}$ asr

```
-s Strands would be distinguished when specified
#
#
       -h help
#
#
    Output to STDOUT:
#
        Title
                      Count
                                mean_mC
#
                      34
                                0.2353
        sense
#
                      54
        antisense
                                0.2778
#
        total
                      88
                                0.2614
#
    Notice:
#
        The overlapping of regions would not be checked.
        A site might be considered multiple times.
#
```

• Example

```
zcat WG.CGmap.gz | cgmaptools mbed -b region.bed
```

• Output format

- Example

chr	sense_0	Count	${\tt sense_mC}$	anti_Cour	nt anti_n	nC all_Count	all_mC
chr1	203	0.081	27 178	0.1148	381	0.09692	
chr2	185	0.070	45 257	0.05586	3 442	0.06197	
chr3	313	0.104	2 250	0.1358	563	0.1182	
chr4	300	0.121	8 271	0.13	571	0.1257	
chr5	282	0.127	2 222	0.1589	504	0.1412	

5.5 mbin

This function will calculated the average methylation levels in equal-length bins, across genome, generating both summary table and distribution graph.

• Command

```
cgmaptools mbin -h
```

```
# Usage: cgmaptools mbin [-i <CGmap>] [-c 10 --CXY 5 -B 5000000]
# (aka CGmapMethInBins)
```

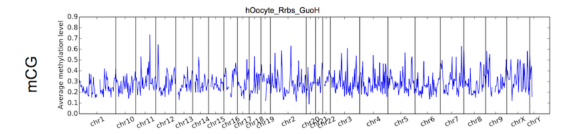


Figure 5.7: Output figure example for cgmaptools mbin

```
#
    Description: Generate the methylation in Bins.
                 Guo, Weilong; guoweilong@126.com
#
#
    Last Update: 2016-10-26
#
    Output Ex:
#
                       5000
                                0.0000
       chr1
               1
#
       chr1
               5001
                       10000
                                0.0396
#
                       5000
                                0.0755
       chr2
               1
#
       chr2
               5001
                       10000
                                0.0027
#
       chr3
                       5000
                                na
#
#
    Options:
#
      -h, --help
                             show this help message and exit
#
      -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
#
      -H FLOAT
                             Height of figure in inch [Default: 4]
#
      -W FLOAT
                             Width of figure in inch [Default: 8]
#
      -p STRING
                             Prefix for output figures
#
      -t STRING, --title=STRING
#
                             title in the output figures
```

• Example

cgmaptools mbin -i WG.CGmap.gz -B 500 -c 4 -f png -t WG -p WG > mbin.WG.data

• File format

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

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5.6 mmbin

This function will calculate the average methylation levels in equal-length bins for **mulitple** samples, generating a summary table.

• Command

```
cgmaptools mmbin -h
```

```
Usage: cgmaptools mmbin [-1 <1.CGmap[,2.CGmap,..]>] [-c 10 --CXY 5 -B 5000000]
#
#
          (aka CGmapsMethInBins)
#
    Description: Generate the methylation in Bins.
#
                 Guo, Weilong; guoweilong@126.com
    Contact:
#
    Last Update: 2016-12-07
#
    Output Ex:
#
       chr1
                        5000
                                0.0000
               1
#
       chr1
               5001
                        10000
                                0.0396
#
                        5000
                                0.0755
       chr2
               1
#
       chr2
               5001
                        10000
                                0.0027
#
       chr3
                        5000
                                na
#
#
    Options:
      -h, --help
#
                             show this help message and exit
#
                             File name list, end with .CGmap or .CGmap.gz. If not
      -1 FILE
#
                             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

```
cgmaptools mmbin -1 WG.CGmap.gz,RR.CGmap.gz,RR2.CGmap.gz,merge.CGmap.gz -c 4 -B 2000 | gawk '{printf("%s:%s-%s", $1, $2, $3); for(i=4;i<=NF;i++){printf("\t%s", $i);} printf("\n");}' > mmbin
```

• Output format

- Example

chr	pos1	pos2	tag1	tag2	tag3
Chr1	111403	113403	0.05	nan	0.02
Chr1	111500	112500	1.00	0.80	0.60
Chr2	20000	20500	0.96	0.33	0.66

- Column Description

5.7 mfg

The cgmaptools mfg supports studys of Methylation in FraGmented regions. The function can be applied to draw mC distribution across gene body, transposon elements, and other user-provided regions. The

Col	Field	Туре	Regexp/Range	Brief description
1	CHR	String	[!-?A-~]*	Query template NAME
2	POS1	Int	[0,2 ³² -1]	Leftmost Position
3	POS2	Int	[0,2 ³² -1]	Rightmost Position
4	METH_1	Float	[0.00, 1.00]	Methylation level in sample 1
n	METH_n	Float	[0.00, 1.00]	Methylation level in sample n

Figure 5.8: Output format description for cgmaptools mmbin

fragmented regions can be generated using cgmaptools bed2fragreg.

• Command

```
cgmaptools mfg -h
```

```
Usage: cgmaptools mfg [-i <CGmap>] -r <region> [-c 5 -C 500]
   Description: Calculated methylation profile across fragmented regions.
#
#
                 Guo, Weilong; guoweilong@126.com
#
   Last Update: 2017-01-20
#
   Options:
#
       -i String, CGmap file; use STDIN if not specified
#
           Please use "gunzip -c <input>.gz " and pipe as input for gzipped file.
#
                    851 CHH CC 0.1 1
           chr1 G
#
       -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
#
           #chr strand U1 R1 R2 D1 End
#
#
           chr1 +
                    600 700 800 900 950
           chr1 -
                            1500
#
                    1600
                                    1400
                                             1300
                                                     1250
#
       -c Int, minimum Coverage [Default: 5]
#
       -C Int, maximum Coverage [Default: 500]
           Sites exceed the coverage range will be discarded
#
#
       -x String, context [use all sites by default]
#
           string can be CG, CH, CHG, CHH, CA, CC, CT, CW
#
       -h help
#
   Output to STDOUT:
#
       Region_ID
                       U1
                               R1
                                        R2
                                                D1
#
       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
#
                       8.0
                               4.0
                                        2.0
       anti_sum_mC
                                                0.0
#
       anti_sum_NO
                       20
                               20
                                        20
#
                       0.43
                               0.27
                                       0.17
                                                0.2
       total_ave_mC
#
                       13.0
                               8.0
                                        5.0
                                                2.0
       total_sum_mC
#
                               30
                                        30
       total_sum_NO
                       30
                                                10
```

• Example:

for CHR in 1 2 3 4 5; do (for P in 1 2 3 4 5; do echo | gawk -vC=\$CHR -vP=\$P -vOFS="\t"

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5.8 mstat

CGmapTools provide resourceful statistic analysis on DNA methylation globally. The cgmaptools mstat command will generated a table summary, together with several graphs:

- mC contributions of different contexts in Pie chart
- Bulk mC levels of different contexts
- Fragmented distribution of mC in different contexts

specified

The methylation contexts are different for plants and animals. - For plants, the contexts for DNA methylations are known as \mathbf{CG} , \mathbf{CHG} and \mathbf{CHH} , where $H = \{A, C, T\}$. - For animals, the situation is different. In 2014, Weilong Guo, et al. showed that it is unnecessary to seperate CHG methylations and CHH methylations in human. In 2016, Weilong Guo, et al. designed \mathbf{MiDD} method and de novo predicted the main separated contexts for non-CG (CH) methylation should be CW ($W = \{A, T\}$) and CC, and mCW is cell-type specific and conserved between human and mice. In $\mathbf{CGmapTools}$, we support both human view (\mathbf{CG} , \mathbf{CW} and \mathbf{CC}) and plant view (\mathbf{CG} , \mathbf{CHG} , and \mathbf{CHH}) for DNA methylation contexts.

Command

#

#

#

#

#

-H FLOAT

-W FLOAT

-p STRING

-t STRING, --title=STRING

```
cgmaptools mstat -h
#
    Usage: cgmaptools mstat [-i <CGmap>]
#
          (aka CGmapStatMeth)
#
    Description: Generate the bulk methylation.
#
                  Guo, Weilong; guoweilong@126.com
#
    Last Update: 2016-12-08
    Output Ex:
#
#
       MethStat
                                         CG
                                                  CHG
                                                            CHH
                                                                    CA
                                                                             CC
                                                                                      CT
                                                                                              CH
                                                                                                       CW
                        context C
#
                                         0.3719
                                                  0.0465
                                                           0.0403
                                                                   0.0891
                                                                            0.0071
                                                                                    0.0241
                                                                                             0.0419
                                                                                                      0.0559
       mean mC
                        global
                                 0.0798
#
                                                                            0.0049
                                         0.0341
                                                  0.0163
                                                           0.0110
                                                                                    0.0076
                                                                                             0.0096
                                                                                                      0.0148
       sd_mCbyChr
                        global
                                 0.0078
                                                                   0.0252
#
       count C
                        global
                                 10000
                                         1147
                                                  2332
                                                           6521
                                                                    3090
                                                                            2539
                                                                                     3224
                                                                                             8853
                                                                                                      6314
#
       contrib mC
                        global
                                 1.0000
                                                  0.1360
                                                           0.3292
                                                                            0.0228
                                                                                    0.0973
                                                                                             0.4652
                                                                                                      0.4424
                                         0.5348
                                                                   0.3452
#
       quant_mC
                        [0]
                                 8266
                                         471
                                                  2012
                                                           5783
                                                                   2422
                                                                            2421
                                                                                     2952
                                                                                             7795
                                                                                                      5374
#
       quant_mC
                                                                                                      426
                   (0.00, 0.20] 705
                                         182
                                                  155
                                                           368
                                                                   272
                                                                            97
                                                                                     154
                                                                                             523
#
       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
#
#
    Options:
#
      -h, --help
                              show this help message and exit
#
      -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
```

Height of figure in inch [Default: 3]

Width of figure in inch [Default: 8]

Prefix for output figures



Figure 5.9: mC contribution example

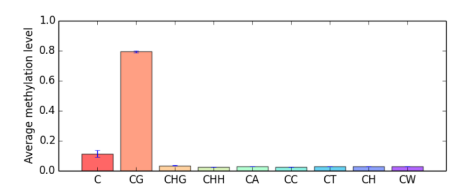


Figure 5.10: Bulk mC example

title in the output figures

• Example

#

cgmaptools mstat -i WG.CGmap.gz -c 4 -f png -p WG -t WG > WG.mstat.data

• File format

The output format:

MethStat	context	C	CG	CHG	CHH	CA	CC	CT	CH	CW
${\tt mean_mC}$	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 (0.0	0,0.20]	705	182	155	368	272	97	154	523	426
${\tt mean_mC_byChr}$	chr1	0.0840	0.4181	0.0340	0.0412	0.0794	0.0065	0.0251	0.0393	0.0513
${\tt mean_mC_byChr}$	chr10	0.0917	0.4106	0.0758	0.0421	0.0968	0.0097	0.0349	0.0502	0.0655

• Output figures

5.9 mtr

The cgmaptools mtr command will calculated the DNA methylation levels for each investigged region.

Command

cgmaptools mtr -h

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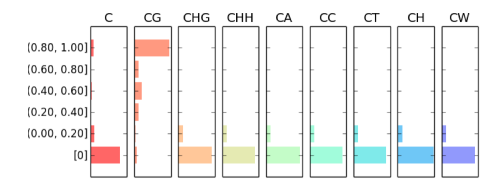


Figure 5.11: mC fragmented distribution example

```
Usage: cgmaptools mtr [-i <CGmap>] -r <region> [-o <output>]
#
          (aka CGmapToRegion)
#
#
   Description: Calculated the methylation levels in regions in two ways.
                 Guo, Weilong; guoweilong@126.com
#
   Contact:
#
   Last Update: 2017-01-20
#
   Format of Region file:
#
      #chr
              start_pos
                         end_pos
#
       chr1
              8275
                         8429
#
   Output file format:
                        end_pos mean(mC) #_C #read(C)/#read(T+C) #read(T+C)
#
       #chr start_pos
#
       chr1
              8275
                         8429
                                  0.34
                                           72
                                                       0.40
                                                                        164
   Note: The two input CGmap files should be sorted by Sort_chr_pos.py first.
#
          This script would not distinguish CG/CHG/CHH contexts.
#
#
#
   Options:
#
      -h, --help show this help message and exit
#
                  File name end with .CGmap or .CGmap.gz. If not specified, STDIN
      -i FILE
#
                  will be used.
#
      -r FILE
                  Filename for region file, support *.gz
      -o OUTFILE To standard output if not specified.
```

• Example

cgmaptools mtr -i WG.CGmap.gz -r region.bed -o WG.mtr.gz

• Input region format

```
#chr start_pos end_pos
chr1 8275 8429
```

Output format

- Example

```
8275
               8429
                        0.34
                                              164
chr1
                                72
                                      0.40
chr1
       8899
               8999
                        0.20
                                40
                                      0.33
                                              198
chr2
       8275
               8429
                        0.50
                                12
                                      0.45
                                              40
```

- Column Description

Col	Field	Type	Regexp/Range	Brief description
1	CHR	String	[!-?A-~]*	Query template NAME
2	POS_L	Int	[0,2 ³² -1]	Leftmost Position
3	POS_R	Int	[0,2 ³² -1]	Rightmost Position
4	MC_s	Float	[0, 2 ³² -1)	Average methylation levels by each site
5	NC_s	Int	[0,2 ³² -1]	Count of Cytosines in this region
6	MC_r	Float	[0, 2 ³² -1)	Average methylation levels recalculated by region
7	NC_r	Int	[0,2 ³² -1]	Sum of effective coverage for all cytosines in this region

Figure 5.12: Output format description for cgmaptools mtr

Chapter 6

Coverage Analysis

Read coverage is an important factor for interpreting DNA methylomes.

It requires different coverage levels for different purpose. For example, SNV calling requires higher coverage than it is required for DMR study. The SNV calling process is depending on all nucleotides (A, T, C and G), whereas DNA methylation levels only depend on T and C read counts aligned to cytosines.

In CGmapTools, we propsed two ways for evaluating the coverages of DNA methylations: OverAll Coverage (OAC) and Methylation-Effective Coverage (MEC).

- OAC is calculated as the average read coverage on all nucleotides on both strands, which are calculated from the ATCGmap file.
- MEC is calculated as the average read coverage only for cytosines, which is calculated from the CGmap file. Generally, the MEC is slightly higher than half of the OAC.

In *CGmapTools*, we provides function for basic statistics of coverages (cgmaptools oac stat and cgmaptools mec stat) and visualization of coverages in bins across genome (cgmaptools oac bin and cgmaptools mec stat).

6.1 oac

Command

```
cgmaptools oac -h
```

```
# Usage: cgmaptools oac <command> [options]
# Version: 0.0.4
# Commands:
# bin * overall coverage in bins
# stat * overall coverage statistics globally
```

6.1.1 oac bin

```
cgmaptools oac bin -h
```

```
# Usage: cgmaptools oac bin [-i <ATCGmap>] [-B 5000000]
# (aka ATCGmapCovInBins)
# Description: Generate the overall coverage in Bins.
```

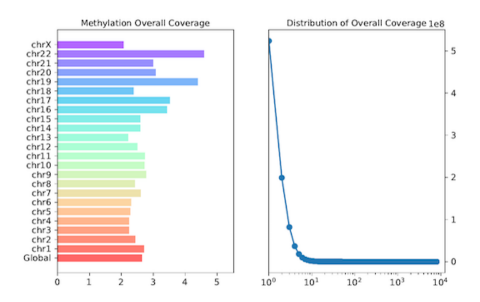


Figure 6.1: MEC example

```
#
    Contact:
                 Guo, Weilong; guoweilong@126.com;
    Last Update: 2016-12-07
#
#
    Output Ex:
#
                        5000
                                29.0000
       chr1
               1
#
       chr1
               5001
                        10000
                                30.0396
#
                        5000
                                35.0755
       chr2
               1
#
       chr2
               5001
                        10000
                                40.0027
#
       chr3
                        5000
                                na
#
#
    Options:
#
      -h, --help
                             show this help message and exit
#
      -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
#
                             Height of figure in inch [Default: 4]
      -H FLOAT
#
      -W FLOAT
                             Width of figure in inch [Default: 8]
#
      -p STRING
                             Prefix for output figures
#
      -t STRING, --title=STRING
                             title in the output figures
```

• Example

cgmaptools oac bin -i WG.ATCGmap.gz -B 1000 -f png -p WG -t WG > WG.oac_bin.data

• Output figure

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6.1.2 oac stat

• Command

```
cgmaptools oac stat -h
```

```
Usage: cgmaptools oac stat [-i <ATCGmap>]
#
           (aka ATCGmapStatCov)
#
   Description: Get the distribution of overall coverages.
                Guo, Weilong; guoweilong@126.com;
#
   Last Update: 2016-12-16
#
   Output Ex:
#
       OverAllCov
                       global 47.0395
#
       OverAllCov
                       chr1
                               45.3157
#
       OverAllCov
                     chr10 47.7380
#
       CovAndCount
                               1567
                     1
                               655
#
       CovAndCount
                       2
#
       CovAndCount
                      3
                               380
#
#
   Options:
#
      -h, --help
                            show this help message and exit
#
     -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
     -H FLOAT
                            Scale ratio for the Height of figure [Default: 4]
#
     -W FLOAT
                            Width of figure in inch [Default: 8]
     -p STRING
                            Prefix for output figures
```

• Example

cgmaptools oac stat -i WG.ATCGmap.gz -p WG -f png > WG.oac_stat.data

• output format:

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
${\tt CovAndCount}$	1	1567
CovAndCount	2	655
CovAndCount	3	380

6.2 mec

```
# Usage: cgmaptools mec <command> [options]
# Version: 0.0.4
# Commands:
# bin * methylation effective coverage in bins
# stat * methylation effective coverage statistics globally
```

6.2.1 mec bin

• Command

```
cgmaptools mec bin -h
    Usage: cgmaptools mec bin [-i <CGmap>] [-B 5000000]
#
#
          (aka CGmapCovInBins)
#
    Description: Generate the methylation-effective coverage in Bins.
                 Guo, Weilong; guoweilong@126.com;
    Contact:
#
    Last Update: 2016-12-07
#
    Output Ex:
#
       chr1
               1
                       5000
                                29.0000
#
       chr1
               5001
                       10000
                                30.0396
#
       chr2
                       5000
                                35.0755
#
               5001
                       10000
                                40.0027
       chr2
#
       chr3
                       5000
#
#
    Options:
#
      -h, --help
                             show this help message and exit
#
      -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
                             Height of figure in inch [Default: 4]
#
      -H FLOAT
#
      -W FLOAT
                             Width of figure in inch [Default: 8]
      -p STRING
                             Prefix for output figures
#
#
      -t STRING, --title=STRING
#
                             title in the output figures
#
      -C CONTEXT, --context=CONTEXT
#
                             specific context: CG, CH, CHG, CHH, CA, CC, CT, CW
```

Example

#

#

cgmaptools mec bin -i WG.CGmap.gz -B 1000 -f png -p WG -t WG > WG.mec_bin.data

use all sites if not specified

6.2.2 mec stat

• Command

(aka CGmapStatCov)

```
cgmaptools mec stat -h

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

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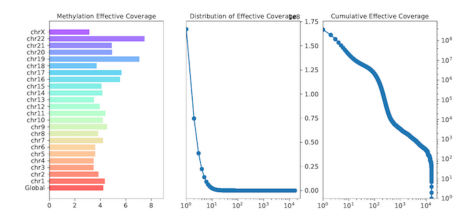


Figure 6.2: MEC example

```
#
    Description: Get the distribution of methylation-effective coverages.
                 Guo, Weilong; guoweilong@126.com
#
    Contact:
#
    Last Update: 2016-12-16
#
    Output Ex:
#
       MethEffectCove global
                                47.0395
#
       MethEffectCove chr1
                                45.3157
#
       MethEffectCove chr10
                                47.7380
#
       CovAndCount
                                1567
                       1
       {\tt CovAndCount}
#
                        2
                                655
#
       CovAndCount
                                380
                       3
#
#
    Options:
#
      -h, --help
                             show this help message and exit
#
      -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
#
      -H FLOAT
                             Scale factor for the Height of figure [Default: 4]
#
      -W FLOAT
                             Width of figure in inch [Default: 11]
#
      -p STRING
                             Prefix for output figures
#
      -C CONTEXT, --context=CONTEXT
                             specific context: CG, CH, CHG, CHH, CA, CC, CT, CW
#
                             use all sites if not specified
#
```

• Example

cgmaptools mec stat -i WG.CGmap.gz -p WG -f png > WG.mec_stat.data

• Output figure

Chapter 7

Graphics

7.1 lollipop

The consideration that we design novel **Lollipop** plot, is to be able to distinguish **un-methylated sites** and **un-detected sites**. Each covered cytosine would have a large round head; color and height of the bar represent DNA methylation level.

• Command

```
cgmaptools lollipop -h
```

```
Usage: cgmaptools lollipop [options] file
#
          (aka mCLollipop)
#
    Description: Plot local mC level for multiple samples
                 Guo, Weilong; guoweilong@126.com
#
    Last Update: 2017-09-16
#
    Example:
#
        mCLollipop [-i input] -o gene.png
#
    -Input Format (-i)
#
        Can be output by "cgmaptools mergelist tomatrix". Use STDIN if omitted.
#
        The 1st line (header line) is required.
#
        Example:
#
           chr
                            tag1
                                    tag2
                                            tag3
                   pos
#
                           0.30
                                            0.80
           Chr1
                   111403
                                    nan
                   111406 0.66
#
           Chr1
                                    0.40
                                            0.60
#
    -Site File (-s)
#
        >= 3 columns, the 1st line (header line) is required, using R color name or "NaN".
#
        To show specific sites (such as DMS, SNV) at the bottom as triangles.
#
        Example:
#
            chr
                             A_vs_B
                                     B_vs_C
                                             A_vs_C
                  pos
#
            chr1
                  13116801
                             NaN
                                     NaN
                                              darkgreen
#
            chr1
                  13116899
                             NaN
                                              NaN
#
    -Region File (-b)
#
        the first 4 columns are required.
#
        To show specific region (such as DMR, Repeats) at the bottom as blocks.
#
        Example:
#
            chr1
                  213941196 213942363
                                         hyper-DMR
#
                             213943530
                                         hypo-DMR
            chr1
                  213942363
#
                                         region-description
            chr
                  left
                              right
```

• Figure examples

• refFlat format

• Example

```
#
    -annotation file (-a), refFlat Format:
#
        To show the structure of genes/transcripts. One-line in annotation, one-track in figure.
#
        Example:
#
                                          1000
                                                    2000
                                                                1100
                                                                                  3
                                                                                        1100,1500,1700, 1
            {\tt GeneA}
                    TransA chr2 +
                                                                        1950
#
            GeneID TrandID ChrID Strand TransLeft TransRight CDSLeft CDSRight nExon ExonLefts
#
#
#
    Options:
#
        -i INFILE, --infile=INFILE
#
            input file
#
#
        -a ANNOTATION, --annotation=ANNOTATION
#
            [opt] annotation file name, refFlat format
#
#
        -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
#
#
        -s SITE, --site=SITE
#
            [opt] file of site to be marked
#
#
        -b BED, --bed=BED
#
            [opt] BED file for region to be markered
#
#
        -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.
#
#
        -h, --help
#
            Show this help message and exit
  • Example
    cgmaptools lollipop -i matrix.CG.gz -a anno.refFlat -f pdf
```

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TRIM59

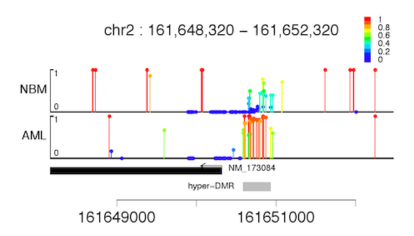


Figure 7.1: Lollipop example-1

VCAN

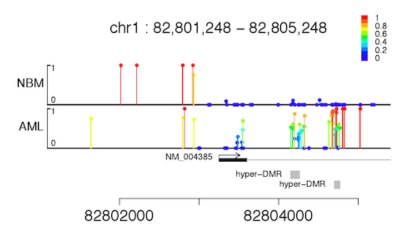


Figure 7.2: Lollipop example-2

GeneA TransA chr2 + 1000 2000 1100 1950 3 1100,1500,1700, 1200,1580,1950,

```
• Description
```

```
Col 1: Gene ID
Col 2: Transcript ID
Col 3: chromatine ID
Col 4: strand, "+" or "-"
Col 5: The left-most position of transcript
Col 6: The right-most position of transcript
Col 7: The left-most position of CDS
Col 8: The right-most position of CDS
Col 9: Number of exons
Col 10: List of left-most position of exons, seperated by ","
Col 11: List of right-most position of exons, seperated by ","
```

• Convert GTF format to refFlat format

The following is an example for Z. mays.

```
"gtfToGenePred" is a command tool downloaded from UCSC utility.
```

```
gtfToGenePred -genePredExt -geneNameAsName2 -allErrors AGPv4.gtf AGPv4.GenePred
paste <(cut -f13 AGPv4.GenePred) <(cut -f1-10 AGPv4.GenePred) > AGPv4.refFlat
paste <(cut -f13 AGPv4.GenePred) <(cut -f1-10 AGPv4.GenePred) | sed -i s/transcript://g | cut -f9 | gaw
cut -f1-10 AGPv4.GenePred > AGPv4.refFlat.tmp
gawk -F"\t" -v0FS="\t" 'ARGIND==1{GeneID[$1]=$2;} ARGIND==2{printf GeneID[$1]"\t"$0}' trans_gene_ID AGP
rm ${GN}.refFlat.txt AGPv4.GenePred
```

7.2 heatmap

```
cgmaptools heatmap -h

# Usage: cgmaptools heatmap [options]
```

```
#
          (aka mCBinHeatmap)
#
    Description: Plot methylation dynamics of target region for multiple samples [heatmap]
                 Zhu, Ping; pingzhu.work@gmail.com
#
   Last update: 2017-09-16
#
#
    Example:
#
      mCBinHeatmap.R -i input -m white -o chr1.xxx-xxx.pdf
#
      -Input File Format:
#
      1st line is the header.
      Each column contains methylation measurements of a sample.
#
#
      Example:
#
      Region Sample1 Sample2 ...
#
      Region1 0.1
                       0.1
#
      Region2 0.1
                       0.1
#
#
#
    Options:
#
        -i INFILE, --infile=INFILE
#
            input file
#
#
        -o OUTFILE, --outfile=OUTFILE
```

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```
[opt] output file name. [default: mCBinHeatmap.SysDate.pdf]
#
        -c, --cluster
#
#
            [opt] cluster samples by methylation in regions. [default: FALSE]
#
#
        -1 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]
#
#
        -h, --help
            Show this help message and exit
  • Example:
     cgmaptools mmbin -1 1.CGmap,2.CGmap,3.CGmap > mmbin.tab cgmaptools heatmap -i mmbin.tab
     -c -o cluster.pdf -f pdf
```

• Figure examples

7.3 fragreg

• Command

```
cgmaptools fragreg -h
#
   Usage: cgmaptools fragreg [options]
#
          (aka mCFragRegView)
#
   Description: Plot methylation dynamics of target and flanking region for multiple samples
#
   Contact:
                 Zhu, Ping; pingzhu.work@gmail.com
   Last update: 2017-09-16
#
   Example:
      FragRegView.R -i input -r 5 -o genebody.pdf
#
   -Input File Format:
      1st line is the header.
   Each row contains methylation measurements of a sample.
```

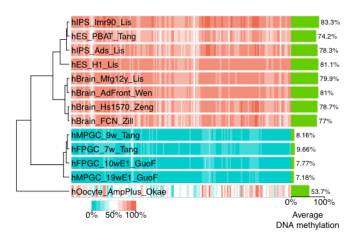


Figure 7.3: heatmap example-1

```
#
   Example:
#
      Sample Up1 Up2 ...
                             Region1 Region2 ...
                                                    Down1
                                       0.2
#
      Sample1 0.1 0.1
                             0.2
                                                    0.3
                                                           0.3
                        . . .
      Sample2 0.1 0.1 ...
                             0.2
                                       0.2
#
                                                    0.3
                                                           0.3
#
#
#
   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]
#
#
        -h, --help
            Show this help message and exit
```

• Example

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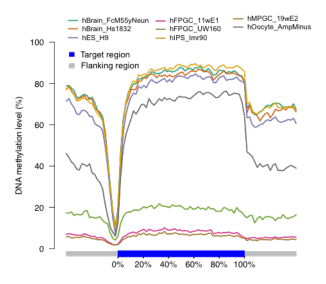


Figure 7.4: DNA methylation distribution across gene body

• Output format

- Example

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

- Column Description

- * Tab-delimited file with header line
- * Content in 1st col is fixed
- * Column number is dynamic
- $\ast\,$ Invalid number is annotated as "NaN"

• Output figure

7.4 tanghulu

The **Tanghulu** plot is designed as show the methylation state on each cytosine by reads. (See what does "*Tanghulu*" strand for? Wikipedia)

• Command

```
cgmaptools tanghulu -h
  DESCRIPTION
           Circle plot representing DNA methylation of each C [defualt CpG] site
#
#
           on each mapped reads.
#
#
  USAGE
#
           cgmaptools tanghulu [options] -r <ref> -b <bam> -l chr1:133-144
#
           or: cgmaptools tanghulu [options] -r <ref> -b <bam> -l chr1:133
           (aka mCTanghulu)
#
#
#
           Options:
#
                 Samtools indexed reference genome sequence, fasta format. eg. hg19.fa
                 - use samtools to index reference: samtools faidx <hg19.fa>
#
                 Samtools indexed Bam file to view.
           -h
                 - use samtools to index bam file: samtools index <input.bam>
#
           -1
                 Region in which to display DNA methylation.
#
                 - or specify a single position (eg. heterozygous SNP site), we will show allele specif
#
                 Path to samtools eg. /home/user/bin/samtools
           -s
#
                 - by defualt, we try to search samtools in your system PATH.
                 Output results to file [default: CirclePlot.Ctype.region.Date.pdf].
#
#
                 C context. [default: CG]
           -t.
#
                 - available context: C, CG, CH, CW, CC, CA, CT, CHG, CHH
#
                 Ouput device. [default: pdf]
           -d
#
                 - alternative: png
#
                 Seperate reads by chain. [default: OFF]
           -c
                 - specify this option to turn ON.
#
                 Show vague allele linked reads. [ default: OFF]
           -v
#
                 Genotype of heterozygous SNP site.
           -g
#
                 - 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.
                 Minimum number of reads (depth) covered in this region or allele linked. [default: 0|0
#
           -D
#
           -C
                 Minimum number of C (specified type) covered in this region or allele linked. [default
                 Width of graphics reigon in inches. [default: 4]
#
           -W
                 Height of graphics reigon in inches. [default: 4]
#
           -H
#
           -R
                 Resolution in ppi. [default: 300]
#
                 - only available for png device.
#
           -h
                 Help message.
#
#
  AUTHOR
#
                        Zhu, Ping; pingzhu.work@gmail.com
#
           Last update: 2016-12-07
```

Example

cgmaptools tanghulu -r genome.fa -b WG.bam -l chr1:2000-2400 -t CG

· Output figure

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chr1:3017150-3017200

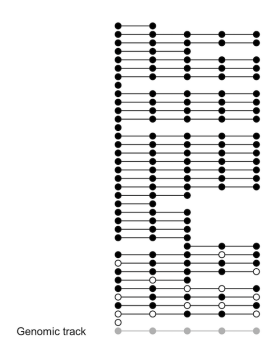


Figure 7.5: Tanghulu plot example

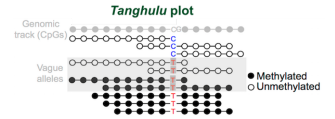


Figure 7.6: Tanghulu plot show vague-reads

We also designed **Tanghulu** plot for visualizing reads that are support methylated, un-methylated, and vague reads for Allele-Specific Methylation (ASM) region.

Chapter 8

Other Ultilities

8.1 findCCGG

• Command

```
cgmaptools findCCGG -h
   Usage: cgmaptools findCCGG -i <genome.fa> [-o <output>]
          (aka FiindCCGG)
#
   Description: Get the positions of all the C'CGG---CCG'G fragments.
                 Guo, Weilong; guoweilong@126.com
   Last Update: 2017-01-20
#
   Output Ex:
#
        chr1
                4025
                        5652
#
        chr1
                8274
                        8431
#
#
   Options:
      -h, --help show this help message and exit
      -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

    cgmaptools findCCGG -i genome.fa -o genome.ccgg
```

8.2 bed2fragreg

```
cgmaptools bed2fragreg -h

# Usage: cgmaptools bed2fragreg [-i <BED>] [-n <N>] [-F <50,50,...> -T <50,...>] [-o output]

# (aka FragRegFromBED)

# Description: Generate fragmented regions from BED file.

# Contact: Guo, Weilong; guoweilong@126.com

# Last Update: 2017-01-20

# Split input region into N bins, get fragments from 5' end and 3' end.

# Input Ex:
```

Col	Field	Type	Regexp/Range	Brief description
1	CHR	String	[!-?A-~]*	Query template NAME
2	STRAND	Char	[+-]	Strand
3	POS_1	Int	[0,2 ³² -1]	The 1st position from 5' end
4	POS_2	Int	[0,2 ³² -1]	The 2 nd position from 5' end
			•••	
n+2	POS_n	Int	[0,2 ³² -1]	The nth position from 5' end

Figure 8.1: Output format description for cgmaptools bed2fragreg

```
#
       chr1
              1000
                      2000
                              +
#
       chr2
              9000
                      8000
#
    Output Ex:
#
                  940 950 1000 1200 1400 1600 1800 1850
       chr1
#
       chr2
                  9060 9050 9000 8800 8600 8400 8200 8150
#
#
#
    Options:
      -h, --help
                   show this help message and exit
#
      -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]
      -o OUTFILE
                   To standard output if omitted. Compressed output if end with
#
```

• Example

• Output format

- Example

```
chr1 + 940 950 1000 1200 1400 1600 1800 2000 2060 2080
chr2 - 9060 9050 9000 8800 8600 8400 8200 8000 7960 7940
```

- Column Description

[POS_1, POS_2), [POS_2, POS_3), ... [POS_(n-1), POS_n) will be used as input for $\mathbf{cgmaptools}$ \mathbf{mfg}