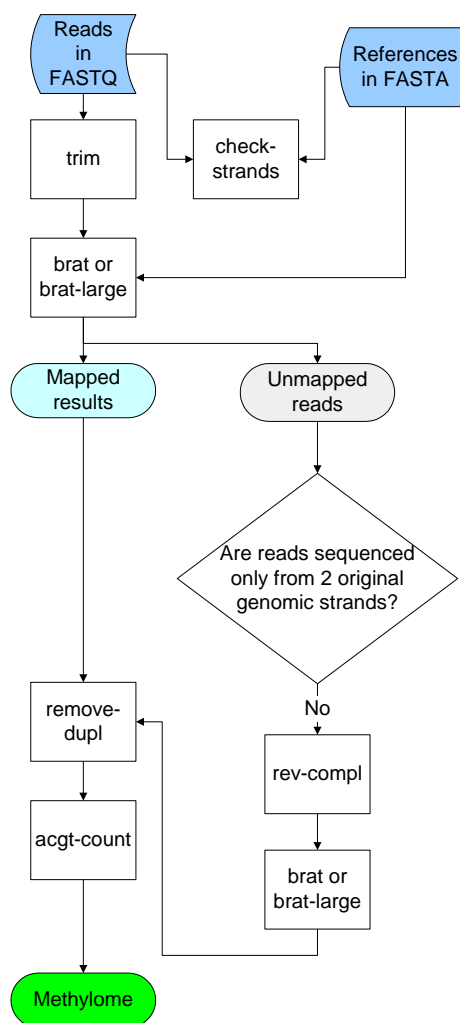


## USING BRAT-1.2.3

This new version has a new tool `convert-to-sam` that converts BRAT format to SAM format. Please use this program as needed after `remove-dupl` in the pipeline below.

### 1 ANALYSIS PIPELINE

Currently BRAT includes the following tools: `brat`, `brat-large`, `brat-large-build`, `acgt-count`, `trim`, `rev-compl`, `check-strands` and `remove-dupl`. The flow of the analysis pipeline is given in the figure below.



First, tool `trim` takes reads in FASTQ format, trims low-quality bases from both ends as well as Ns and outputs reads in raw reads format that are accepted by `brat` and `brat-large`. After `trim`, `brat` or `brat-large` are used to map the reads to the reference genome. If the reads are sequenced from four PCR-product strands, then `rev-compl` is used to take reverse-complements of the unmapped reads followed by mapping of the results with `brat` or `brat-large`. Next, mapped results are used as input for `remove-dupl` that removes copy-duplicates keeping only a randomly chosen one, where copy-duplicates are the reads that are mapped to the same start position in the reference. Finally `acgt-count` processes all the mapped results to produce methylome, a map with methylation status of each cytosine. Tool `check-strands` is used for information on ACGT-distribution in the reference and input reads.

## 2 SYSTEM AND SPACE REQUIREMENTS

BRAT stands for Bisulfite-treated Reads Analysis Tool. There are two versions of BRAT: BRAT and BRAT-large. Both programs run on 64-bit architecture under Linux/Unix operating system. Both versions work with reads of lengths 24 bases and above.

BRAT accepts up to 4.2G references with total size of references up to 4.2G base pairs. BRAT-large can work with up to 4.2G references with the size of the largest reference up to 4.2G base pairs.

BRAT works best with relatively small genomes because it uses significantly more space than BRAT-large. BRAT is faster than BRAT-large that maps reads to the references sequentially: it maps all reads to the first reference, then all reads to the second reference and so on. The space requirement of BRAT-large depends on the size of the largest reference in the set of input references, whereas BRAT's space requirement depends on the total size of all input references (measured in base pairs).

Let  $N$  be the total size of a genome in base pairs,  $T$  be the size of the largest reference in base pairs and  $R$  be the total number of reads (each read is counted, *i.e.* a pair has 2 reads). Then the space required for both programs is bound as follows:

$$\begin{array}{ll}\text{Space}_{\text{BRAT}} = 269 \cdot 10^6 + 2 \cdot 4N + 24R + (3/8)N & \text{Bytes} \\ \text{Space}_{\text{BRAT-large}} = 269 \cdot 10^6 + 4T + 24R + (3/8)T & \text{Bytes} \\ \text{Space}_{\text{BRAT-large}} = 269 \cdot 10^6 + 2 \cdot 4T + 24R + (3/8)T & \text{Bytes (when option } S \text{ is provided)}\end{array}$$

Whenever there is a space limitation, the user can choose to use BRAT-large. Here is an example of space usage with paired-end BS-mapping with non-BS-mismatches (mismatches other than T-to-C and A-to-G): BRAT uses 2.5 GB on 1 million pairs and human chromosome 1, and BRAT-large on 10 million pairs and an entire human genome uses 1.7 GB (or 2.7GB with option  $S$ ).

In this version, BRAT allows for non-BS-mismatches with the restriction on the number of non-BS-mismatches in the first  $X$  bases (for BRAT) and  $X$  bases (for BRAT-large), where  $X$  is a threshold specified with the option  $f$  ( $X$  cannot be smaller than 24 or larger than 64). BRAT guarantees to map all reads that could be mapped to the genome with up to one non-BS-mismatch in the first  $X$  bases of reads. A user can specify any number of non-BS-mismatches with the option  $m$ , and as long as there is only one non-BS-mismatch in the first  $X$  bases of a read, BRAT will map this read. BRAT-large does not allow for non-BS-mismatches in the first  $X$  bases of reads. If a user specifies any number of non-BS-mismatches with the option  $m$ , BRAT-large will map all reads that could be mapped perfectly or with BS-mismatches within the first  $X$  bases, and any number of non-BS-mismatches in the other bases of reads.

The package includes three additional tools: trim, brat-large-build and acgt-count. The latter aligns mapped reads to the genome and counts mapped nucleotides at each base for forward and reverse strands separately. The tool trim accepts FASTQ files with reads/pairs as input, trims the ends of the reads whose base quality scores are lower than the user specified threshold and trims Ns at the ends of the reads. The tool brat-large-build is used with option P with BRAT-large.

## 3 COMMANDS AND INPUT

To unzip run:

```
tar xzvf brat-1.2.*.tar.gz
```

To build:

```
cd brat-1.2.*
make
```

This will create executable programs: brat-large, brat-large-build, acgt-count, trim, rev-compl, check-strands and remove-dupl.

Input format of the reads for BRAT and BRAT-large is raw reads:

- Read <string>: a read after using trim;
- Start <int>: the number of bases trimmed at the beginning of the original read;
- End <int>: the number of bases trimmed at the end of the original read.

To convert reads from FASTQ format to raw reads, one should run trim. If a user does not wish to trim reads' low quality score bases, then he/she should omit the option for the base quality score threshold: the default threshold equals to zero, so all reads will be in the output without change in lengths (except for reads having Ns at the ends). If reads have Ns at the ends, trim trims Ns at the ends and outputs only those reads whose length after trimming is greater or equal to 24 bases.

## A command to run trim

This program trims low-quality bases (lower than a threshold given with option *-q*) and Ns from each end of a read: bases are trimmed one at a time from both ends of a read until a base with quality score greater or equal than *q* is encountered (similarly, all consecutive Ns from both ends of a read are trimmed). This tool outputs only those reads whose length is at least 24 after trimming and that have at most *m* internal Ns: the number of allowable internal Ns is set by option *-m*.

To trim single-end reads in the file *reads.fastq* in FASTQ format and output trimmed raw reads into a file with name *prefix\_reads1.txt*, run the command:

```
./trim -s reads.fastq -P prefix -q 20 -L 64 -m 2
```

This will trim bases whose base quality scores are lower than 20 from the ends of reads. The option *L* specifies the smallest value of the range of base quality scores in ASCII representation (please see Commands Options for details). To learn more about Phred scores, please visit <http://www.phrap.com/phred/>. Option *-m* allows each read having at most 2 internal Ns. Option *-P* provides prefix to the output file names (it might contain a path for an output file: *-P /home/directory/prefix*).

If the user does not wish to trim ends with low base quality scores, the *-q* option is not specified. For single-end reads, there is a single output file with trimmed reads.

To trim paired-end reads in the files *reads1.fastq* and *reads2.fastq* in FASTQ format, run the command:

```
./trim -1 reads1.fastq -2 reads2.fastq -P prefix -q 20 -L 64 -m 2
```

Here we assume, that *reads1.fastq* contains sequenced 5` mates, and *reads2.fastq* contains sequenced 3` mates.

The output will be in four files with raw reads: *prefix\_reads1.txt*, *prefix\_reads2.txt*, *prefix\_mates1.txt* and *prefix\_mates2.txt*. To further map paired-end reads, use *prefix\_reads1.txt* and *prefix\_reads2.txt* as input files for paired-end mapping with brat or brat-large. The file *prefix\_mates1.txt* contains reads from the file *reads1.fastq* whose mates have shorter length than 24 bases after trimming. Similarly, the file *prefix\_mates2.txt* contains reads from the file *reads2.fastq* whose mates are shorter than 24 bases. The user can further map these files, *prefix\_mates1.txt* and *prefix\_mates2.txt*, as single-end reads: for BS-mapping of the reads in *prefix\_mates2.txt*, the user must specify **-A** option for mapping to work correctly (the same is true if a user wishes to map the reads in *prefix\_reads2.txt* as single reads).

Additional output files are *prefix\_pair1.fastq*, *prefix\_pair2.fastq*, *prefix\_mates1.fastq* and *prefix\_mates2.fastq*. These files have the same reads as do files *prefix\_reads1.txt*, *prefix\_reads2.txt*, *prefix\_mates1.txt* and *prefix\_mates2.txt* respectively, except the files *prefix\_pair1.fastq*, *prefix\_pair2.fastq*, *prefix\_mates1.fastq* and *prefix\_mates2.fastq* are in FASTQ format. **NOTE:** current version of BRAT and BRAT-large do NOT support FASTQ format. These additional files are for users to track original reads' names and corresponding base quality scores.

## Commands to run brat and brat-large

BRAT and BRAT-large map raw reads (output from trim) to references that have to be in FASTA format: one reference per FASTA file. BRAT accepts a single file that contains names of FASTA files with the references. To map bisulfite single-end reads, run either of the commands:

```
./brat -r references_names.txt -s prefix_reads1.txt -bs -o output_results.txt
```

```
./brat-large -r references_names.txt -s prefix_reads1.txt -bs -o output_results.txt
```

```
./brat -r references_names.txt -s prefix_reads2.txt -bs -o output_results.txt -A
```

```
./brat-large -r references_names.txt -s prefix_reads2.txt -bs -o output_results.txt -A
```

The file *references\_names.txt* contains the names of the FASTA files with the references. The file *output\_results.txt* contains the results of the mapping: only uniquely mapped reads are in this file. The option *-bs* specifies that mapping is for bisulfite sequenced reads. To map normal (not bisulfite-treated) reads, run similar commands but without *-bs* option:

```
./brat -r references_names.txt -s prefix_reads1.txt -o output_results.txt
```

```
./brat-large -r references_names.txt -s prefix_reads1.txt -o output_results.txt
```

To map bisulfite paired-end reads, run either of the following commands:

```
./brat -r references_names.txt -1 prefix_reads1.txt -2 prefix_reads2.txt -pe -bs -o output_results.txt
```

```
./brat-large -r references_names.txt -1 prefix_reads1.txt -2 prefix_reads2.txt -pe -bs -o output_results.txt
```

The option *-pe* specifies paired-end mapping and as with single-end reads, *-bs*, specifies bisulfite mapping. The results of the mapping will be in *output\_results.txt*.

One can choose to pre-build genome index first and then use *brat-large* on this index to speed up mapping. Use option *-P* and tool *brat-large-build*:

```
./brat-large-build -r references_names.txt -P some_directory [options: S, bs, f ]
```

```
./brat-large -P some_directory -1 prefix_reads1.txt -2 prefix_reads2.txt -pe -o output_results.txt [options]
```

To use this option that allows to separate hashing of the genome from mapping of reads, one has to create a directory, *some\_directory* (the name of a directory in the example above), and to run the following commands (1) to build a hashing index and (2) to map reads. If the directory *some\_directory* is the directory inside the directory in which you run the commands above, then providing the name, *some\_directory*, is sufficient. However, if you run these commands in the directory that does not have *some\_directory* in it, then please provide a full path to *some\_directory*.

The tool *brat-large-build* will output hashing index of the references into *some\_directory*. For human genome, space on hard disk for hashing index is 15GB (and 26GB when option *S* is used). The command for tool *brat-large* with option *P* differs only by not specifying option *r*. If option *bs* was passed to *brat-large-build*, then this option must be passed to *brat-large*. In other words, for mapping normal and bisulfite-treated reads, two different hashing indices must be built. The same true when option *S* is used: if a user wishes to use this option and *P* option, then *S* must be used with *brat-large-build*. When *P* is used, during mapping option *f* will have the same value that was set with *brat-large-build*. If a user does not specify this option, the default value, 24, will be used. If a user wishes to speed up mapping time, and to use option *P*, then the user could set option *f* to a higher value with *brat-large-build*.

**General rule for using option *P*:** if a user wishes to change either of the parameters: *f*, *S* or *bs*; the user must apply the same parameters with *brat-large-build*. Default options for *brat-large-build* are *f* = 24, *S* is not set (i.e. preference is given to slower mapping, but also smaller space usage), and normal mapping (i.e. *bs* is not set).

## Commands Options:

**-A** specifies 3`mates (in our examples above, either of *prefix\_reads2.txt* or *prefix\_mates2.txt* files must be used with this option). If the user does not specify this option, and provides either of the files, *prefix\_mates2.txt* or *prefix\_reads2.txt*, as input reads for single-end mapping, the mapping will NOT be correct;

**-u** is used when a user wishes to output unmapped reads. The output will be in *prefix\_reads1.txt.unm* file (assuming *prefix\_reads1.txt* was provided in the command line with option **-s**) for single reads and with paired-end reads in *prefix\_reads1.txt.unm* and *prefix\_reads2.txt.unm* files (if *prefix\_reads1.txt* and *prefix\_reads2.txt* were used with the options **-1** and **-2** in the corresponding command line). Unmapped reads in the output files are in raw reads format: <read, X, Y>, where X and Y are the number of bases trimmed from the beginning and end of original read respectively (there are no ambiguous reads/pairs in the files with unmapped reads);

**-s** <single-end reads file>: to specify the file with input reads for single-end reads mapping;

**-1** <paired-end reads file>: to specify the file with 5` mates for paired-end reads mapping (in our example, *prefix\_reads1.txt*). This option is also used with *acgt-count*;

**-2** <paired-end reads file>: to specify the file with 3` mates for paired-end reads mapping (in our example above, *prefix\_reads2.txt*); This option is also used with *acgt-count*;

**-pe** to specify paired-end reads mapping (default is false, i.e. single-end mapping);

-**bs** to specify bisulfite sequenced reads mapping. Without this option, mapping will be done as for normal, not bisulfite-treated, sequenced reads;

-**i** <positive integer>: to specify minimum insert size for paired-end mapping, the minimum distance allowed between the leftmost ends of the mapped mates on forward strand (default is 100);

-**a** <positive integer>: to specify maximum insert size for paired-end mapping, the maximum distance allowed between the leftmost ends of the mapped mates on forward strand (default is 300);

-**o** <string>: to specify the file with the results of mapping;

-**M** is used when a user wishes to output ambiguous reads. The output will be in *prefix\_reads1.txt.amb* for single reads and in *prefix\_reads1.txt.amb* and *prefix\_reads2.txt.amb* files with paired-end reads: a single read per line. This option does not output the mapping locations for ambiguous reads, but just the reads themselves;

-**m** <integer>: the maximum number of non-BS-mismatches allowed by a user (default is 0).

-**f** <integer>: the number of the first bases of a read, where the restriction on the number of non-BS-mismatches applies: for BRAT, only one non-BS-mismatch is allowed in the first <integer> bases, and for BRAT-large NO non-BS-mismatches are allowed in the first <integer> bases (for BRAT, default is 36, and for BRAT-large, default is 24).

-**S** to specify speed mode for BRAT-large. Space usage with this option doubles, but running time is about three times faster.

-**P** <directory name> To use this option that allows to separate hashing of the genome from mapping of reads.

-**L** <integer>: the smallest value of the range of base quality scores in ASCII representation (default is 33).

The table below gives examples of different quality scores and their range in ASCII representation (from Wikipedia). The option **L** uses the values in the “Smallest Value in ASCII representation” column.

Type	Smallest Score	Largest Score	Smallest Value in ASCII representation	Largest Value in ASCII representation
Phred quality score (Sanger format)	0	93	33	126
Solexa/Illumina, 1.0	-5	62	59	126
Solexa/Illumina, 1.3+	0	62	64	126

-**B**: specifies the second option for output with acgt-count (please read **Output format for acgt-count**).

## A command to run remove-dupl

This program processes the mapping results and removes copy-duplicates: it outputs all reads that are mapped to a unique genomic location and only a randomly chosen one out of copy-duplicates (the reads mapped to the same location).

```
./remove-dupl -r references_names.txt -p pairs_results.txt -1 single_results_mates1.txt -2 single_results_mates2.txt
```

**NOTE: the file *pairs\_results.txt* does not contain the actual results, it contains the names of the files with the results for paired-end mapping, and similarly, *single\_results.txt* file contains the names of the files with the actual results for single-end mapping.**

For example, the content of *pairs\_results.txt* is:

```
output_pairs_lane1.txt
output_pairs_lane2.txt
```

and the content of *single\_results\_mates1.txt* is:

```
output_singles_mates1_lane1.txt
output_singles_mates1_lane2.txt
```

and the content of *single\_results\_mates2.txt* is:

```
output_singles_mates2_lane1.txt
output_singles_mates2_lane2.txt
```

The output of remove-dupl are the files with the same names as before with additional extension “*.nodupl*”:

```
output_pairs_lane1.txt.nodupl
output_pairs_lane2.txt.nodupl
output_singles_mates1_lane1.txt.nodupl
output_singles_mates1_lane2.txt.nodupl
output_singles_mates2_lane1.txt.nodupl
output_singles_mates2_lane2.txt.nodupl
```

For single-end mapping, run the following command:

```
./remove-dupl -r references_names.txt -s single_results.txt
```

The file *single\_results.txt* contains the names of the files with mapping results.

**NOTE: that the output files with extension “.nodupl” are opened with C++ “*app*” option** (opens a file and appends output to the file’s content). This means that once you have run remove-dupl, you will have “.nodupl” files, and if you want for some reason to run remove-dupl on the same files (and possibly some additional files), you need to remove “.nodupl” files for the corresponding files first and only then re-run remove-dupl. For example, as in the example above, you run remove-dupl and obtain “.nodupl” files:

```
output_pairs_lane1.txt.nodupl
output_pairs_lane2.txt.nodupl
output_singles_mates1_lane1.txt.nodupl
output_singles_mates1_lane2.txt.nodupl
output_singles_mates2_lane1.txt.nodupl
output_singles_mates2_lane2.txt.nodupl
```

Then you wish to add *output\_pairs\_lane3.txt* to *pairs\_results.txt*:

```
output_pairs_lane1.txt
output_pairs_lane2.txt
output_pairs_lane3.txt
```

and to re-run remove-dupl on all files.

Make sure you delete existent files with extension “.nodupl”:

```
rm output_pairs_lane1.txt.nodupl
rm output_pairs_lane2.txt.nodupl
rm output_singles_mates1_lane1.txt.nodupl
rm output_singles_mates1_lane2.txt.nodupl
rm output_singles_mates2_lane1.txt.nodupl
rm output_singles_mates2_lane2.txt.nodupl
and only then run remove-dupl.
```

If you don’t remove these files, you will have previous output plus new output (for example: if *output\_pairs\_lane1.txt.nodupl* had 100 lines, then re-running remove-dupl without removing this file will result in 200 lines).

## A command to run convert-to-sam

This program converts BRAT format to SAM format.

```
./convert-to-sam -P prefix -p pairs_results.txt -1 single_results_mates1.txt
-2 single_results_mates2.txt -s single_results.txt
```

**NOTE: the file *pairs\_results.txt* does not contain the actual results, it contains the names of the files with the results for paired-end mapping, and similarly, *single\_results\*.txt* files contain the names of the files with the actual results for single-end mapping (see examples in “A command to run remove-dupl”).**

**NOTE: Use options -1 and -2 only when mapping paired-end reads as singles (for 5’ mates and 3’ mates respectively), and option s if the original sequenced reads were single reads.**

The output will be in two files with names *prefix\_forw.sam* and *prefix\_rev.sam* if option **P** is provided, or *mapped\_to\_forw.sam* and *mapped\_to\_rev.sam* if not. The reason behind separating the mapped reads into two separate files is the following. To distinguish between methylated and unmethylated cytosines in aligned reads, one has to look at C mapped to C and T mapped to C respectively if reads come from forward strand, and one has to look at G mapped to G and A mapped to G respectively if reads come from reverse strand. If one has a mixture of reads from forward and reverse strands, it would be impossible to track methylated and unmethylated cytosines when visualizing aligned reads.

Paired-end mapped mates mapped as pairs are both either in forward output file (*prefix\_forw.sam*) if 5’ mate (the first mate) is mapped to positive strand, or both in reverse output file (*prefix\_rev.sam*). Reads from files with option **1** are in

forward output file if they are mapped to positive strand, and in reverse output file if they are mapped to negative strand. Reads from files with option 2 are in forward output file if they are mapped to negative strand, and in reverse output file if they are mapped to positive strand. Single reads from files with option s are in forward output file if they are mapped to positive strand, and in reverse output file if to negative strand.

**NOTE: that the output files with extension “.sam” are opened with C++ “app” option.** Please see details how to be careful with re-running convert-to-sam in “A command to run remove-dupl” (since it is similar situation as with remove-dupl).

## A command to run acgt-count

To count mapped As, Cs, Gs and Ts at each base of forward and reverse strands of the references, use acgt-count.

```
./acgt-count -r references_names.txt -P prefix -p pairs_results.txt -s single_results.txt
```

the file *references\_names.txt* contains the names of FASTA files with the references, which are needed to calculate the sizes of the references. The output will be in two files per a reference: *prefix\_forw\_aReference\_name* and *prefix\_rev\_aReference\_name*. The option **-p** is to specify the results of paired-end mapping (if any), and **-s** is to specify the results of single-end mapping (if any). **NOTE: the file *pairs\_results.txt* does not contain the actual results, it contains the names of the files with the results for paired-end mapping, and similarly, *single\_results.txt* file contains the names of the files with the actual results for single-end mapping.** At least one of these options must be provided. The files whose names are listed in the files *pairs\_results.txt* and *single\_results.txt* must be in BRAT’s output format.

To make this point clear, assume, a user ran brat on paired-end reads and had the output file with the results in *output\_pairs\_results.txt*; to run acgt-count, the user must store the name of this file in *pairs\_results.txt* file and run acgt\_count using *pairs\_results.txt* (i.e. *pairs\_results.txt* will have in this case one line, namely, *output\_pairs\_results.txt*). The command for this example is:

```
./acgt-count -r references_names.txt -P prefix -p pairs_results.txt
```

**Please note that if a user has paired-end reads (files with mates 1 and mates 2) and wishes to map the mates as single-end reads, then the user must provide names of the files with results for mates 1 and mates 2 separately using options -1 and -2. This will ensure unbiased ACGT-counting when reads are sequenced from two original genomic strands:**

```
./acgt-count -r references_names.txt -P prefix -p pairs_results.txt -1 single_results_mates1.txt -2 single_results_mates2.txt
```

To produce a more concise output, use option -B (choose an appropriate command from the commands below):

```
./acgt-count -r references_names.txt -P prefix -p pairs_results.txt -s single_results.txt -B
```

```
./acgt-count -r references_names.txt -P prefix -p pairs_results.txt -1 single_results_mates1.txt -2 single_results_mates2.txt -B
```

This program takes care of overlapping mates: if two mates of a pair overlap, then ACGT-count is done only for one mate in the overlapped region. This program also takes care of producing un-biased ACGT-count from mates 2 (3` mates). Please see Details on ACGT-count section for details.

## A command to run rev-compl

Please read subsection *How to use BRAT with reads sequenced from four strands*.

## A command to run check-strands

Please read section *Details on ACGT-count*.

## 4 OUTPUT FORMAT

## Output format for **brat** and **brat-large** for single-end mapping

- Read id < integer >: a consecutive number of a read in the reads input file that starts with 0;
- Read 1 < string >: the read given as in the input file (*prefix\_reads1.txt*);
- Reference name < string >: a name of a reference to which the read is mapped (the first word following ">" in a FASTA file);
- Strand "+" if the read is mapped to forward strand, and "-" if the read is mapped to reverse strand;
- Position < integer >: position within the reference starting with 0, where the read is mapped (the leftmost position on forward strand).
- The number of non-BS-mismatches <int>
- Original position <integer>: position within the reference starting with 0, where the *original* read is mapped (the leftmost position on forward strand), where original read is the read before its ends have been trimmed. For example, if the number of trimmed bases at the beginning of a read is 2, and the read is mapped to positive strand at the position 10, then original position is  $10 - 2 = 8$ . If the number of trimmed bases at the end of a read was 3 and the reverse-complement of the read is mapped to the position 10 on positive strand, then original position =  $\text{position} - 3 = 10 - 3 = 7$ . The original positions are used to identify copy-duplicates.

## Output format for **brat** and **brat-large** for paired-end mapping

- Read id < integer >: a consecutive number of a read in the reads input file that starts with 0;
- Read 1 < string >: the first mate of a pair given as in the input file (*prefix\_reads1.txt*);
- Read 2 < string >: the second mate of a pair given as in the input file (*prefix\_reads2.txt*);
- Reference name < string >: a name of the reference to which the pair is mapped (the first word following ">" in a FASTA file);
- Strand "+" if 5' mate (from *prefix\_reads1.txt*) is mapped to forward strand (consecutively, 3' mate, from *prefix\_reads2.txt*, is mapped to reverse strand), and "-" if the 5' mate is mapped to reverse strand (and 3' mate to forward strand);
- Position 1 < integer >: position within the reference starting with 0, where 5' mate is mapped (the leftmost position on forward strand);
- Position 2 < integer >: position within the reference starting with 0, where 3' mate is mapped (the leftmost position on forward strand).
- The number of non-BS-mismatches < integer >: the number of mismatches in the alignment for 5' mate
- The number of non-BS-mismatches < integer >: the number of mismatches in the alignment for 3' mate
- Original position 1 <integer>: original position for 5' mate (see definition of original position above)
- Original position 2 <integer>: original position for 3' mate (see definition of original position above)

## Output format for **acgt-count**

Starting with version *brat-1.1.17*, there are two choices for output format.

The first choice: The number of output files will be double the number of input references: two for each reference listed in *references\_names.txt* file (one file for forward strand and the other for reverse strand). In each file, there are  $M$  lines, where  $M$  is the size of a corresponding reference in base pairs. Each line corresponds to a base of a strand and contains counts for As, Cs, Gs and Ts at that base for all mapped reads (*i.e.* there are four integers per line: from left to right for As, Cs, Gs and Ts).

For the reverse strand, the counts of As, Cs, Gs and Ts are given for the reads that are mapped to the reverse strand, but the counts are obtained by aligning the reverse-complements of these reads with the forward strand.

Following is an example to illustrate this point.

Let a read ACCGTT be mapped to a reverse strand at position  $i$ , then the corresponding forward strand starting at position  $i$  is AACGGT, and the counts for the reverse strand at positions  $i \dots i+5$  from this read are incremented for the following nucleotides:  $i(A)$ ,  $i+1(A)$ ,  $i+2(C)$ ,  $i+3(G)$ ,  $i+4(G)$  and  $i+5(T)$ .

The second choice: If a user provides option *-B*:

```
./acgt-count -r references_names.txt -P prefix -p pairs_results.txt -s single_results.txt -B
```



then the output is in two files: one file for positive strand and another for negative strand (output files will contain words “forw” and “rev” to distinguish between strands). Each line in the output corresponds to a base in the genome that is either a cytosine on positive strand or cytosine on negative strand (given in separate files). Output format:

*chrom, start, end, total, methylation\_level, strand*

where *chrom* is the reference name, *start* and *end* are positions in the genome (Note: base count in a reference starts with 0), *total* takes one of the values: CHH:X, CHG:X or CpG:X, where X is the sum of counts of Cs and Ts mapped to this base, methylation level is calculated as the number of Cs over the *total* (methylation level =  $\text{count\_C}/(\text{count\_C} + \text{count\_T})$ ). CHH, CHG and CpG describe the sequence content following C: if two consecutive bases that follow C are not G, then *total* = CHH:X; if the first consecutive base following C is non-G and the second consecutive is G, then *total* = CHG:X; and finally, if G follows C (we have CG di-nucleotide), then *total* = CpG:X.

## Output format for trim.

The tool trim accepts FASTQ files with reads/pairs as input, trims the ends of the reads whose base quality scores are lower than the user specified threshold or whose ends are Ns. The output for single reads is a single file with reads whose lengths might be different and whose lengths are greater than or equal to 24 bases. The output for pairs is four files: two for paired-end mapping, and two for single-end mapping. Trimming of paired-end reads produces two files with single reads: if one mate is shorter than the minimum length allowed, and the other’s length is correct, then the mate with the correct length will be output into a corresponding file with single reads. Two files for single reads are necessary because BS-mapping for 5` and 3` mates is different. Each file contains a single line (raw reads format) with the following fields:

- Read <string>: a read after using trim;
- Start <int>: the number of bases trimmed at the beginning of the original read;
- End <int>: the number of bases trimmed at the end of the original read.

## 5 DETAILS ON ACGT-COUNT

Let us denote a mapping of a base in a read to a base in the reference as  $C \rightarrow C$ ,  $T \rightarrow C$ ,  $A \rightarrow G$ ,  $G \rightarrow G$ . Initially, we thought that if a read maps to a positive strand, then the mappings  $C \rightarrow C$  and  $T \rightarrow C$  contribute to the count of methylation level of cytosines of the positive strand. Similarly, if the reverse-complement of a read maps to the positive strand (equivalently, a read maps to the negative strand), then the mappings  $A \rightarrow G$  and  $G \rightarrow G$  contribute to the count of methylation level of cytosines of the negative strand. Let us illustrate this idea in Figures 1-3 below.

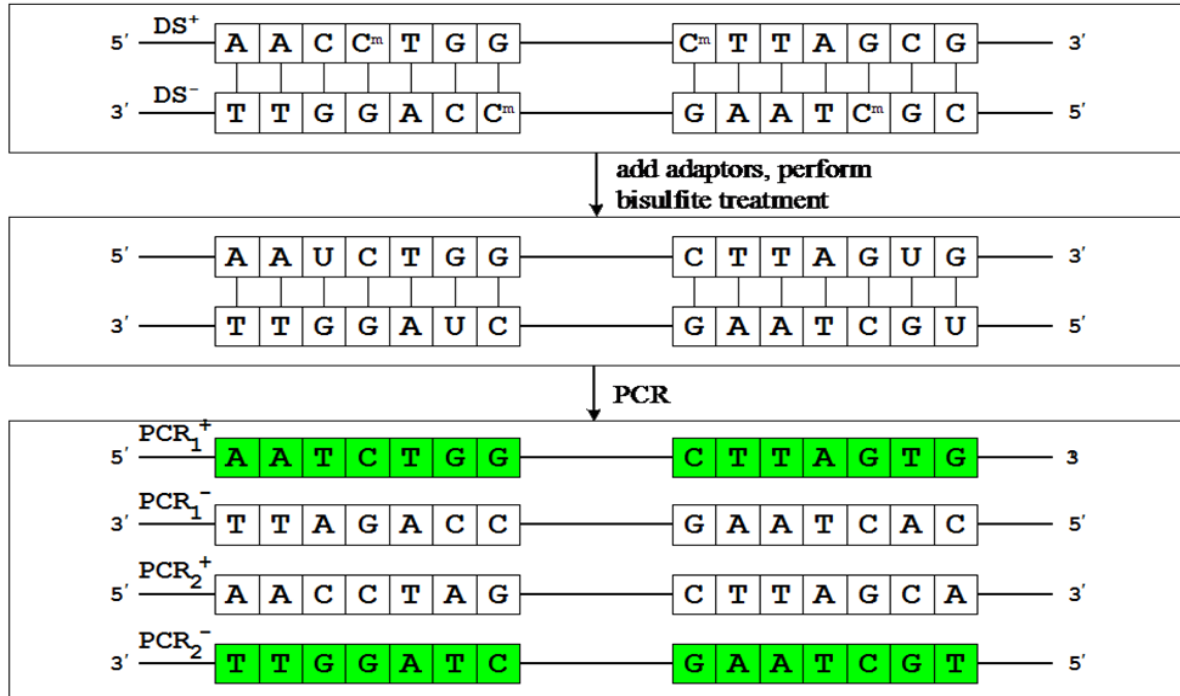


Figure 1. After the special adaptors with methylated cytosines are ligated to DNA fragments, sodium bisulfite treatment (BS-treatment) is applied to DNA fragments, after which unmethylated cytosines are converted to uracils and later to thymines during PCR for library amplification. Note, that after BS-treatment, there are four distinct PCR-product strands: PCR1<sup>+</sup> and PCR2<sup>-</sup> (correspond to original genomic strands) and PCR1<sup>-</sup> and PCR2<sup>+</sup> (the reverse-complements of PCR1<sup>+</sup> and PCR2<sup>-</sup> respectively). Note, that PCR1<sup>+</sup> and PCR2<sup>-</sup> are T-rich and C-depleted (since unmethylated cytosines converted to thymines), and PCR1<sup>-</sup> and PCR2<sup>+</sup> are A-rich and G-depleted (as the reverse-complements of PCR1<sup>+</sup> and PCR2<sup>-</sup> respectively).

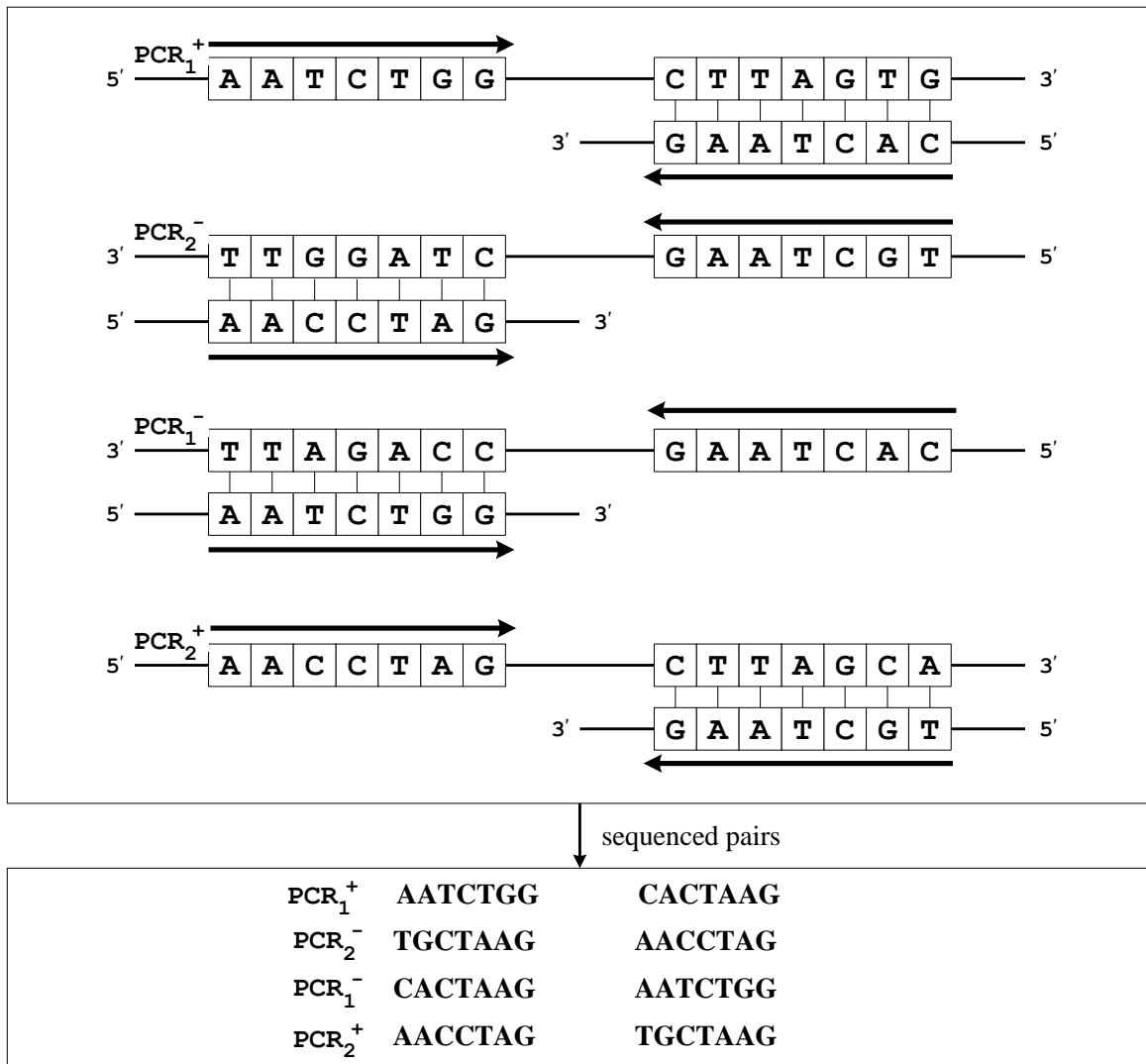


Figure 2. Here we show the sequenced pairs resulted from sequencing all four PCR-product strands, each of which serves as a template during sequencing. If PCR1+ is attached to the flow cell, then in single-end sequencing, 5'-end of this strand is sequenced; in paired-end sequencing, 5'-end of PCR1+ is sequenced, then PCR1+ is copied into its reverse-complement PCR1- and 5'-end of PCR1- is sequenced as the second mate of a pair. Similarly, sequencing produces reads for the rest of strands. If procedure has some technical details that ensure that only original genomic strands serve as templates for sequencing, then even though we have four PCR-product strands, we'll have reads sequenced only from PCR1+ and PCR2- with mates ordering shown above (in paired-end sequencing, the left column of reads are mates 1 will have reads IDs end with "1" and will be in one file, the right column of reads are mates 2 will have reads IDs end with "2" and will be in the other file of Illumina's sequenced reads).

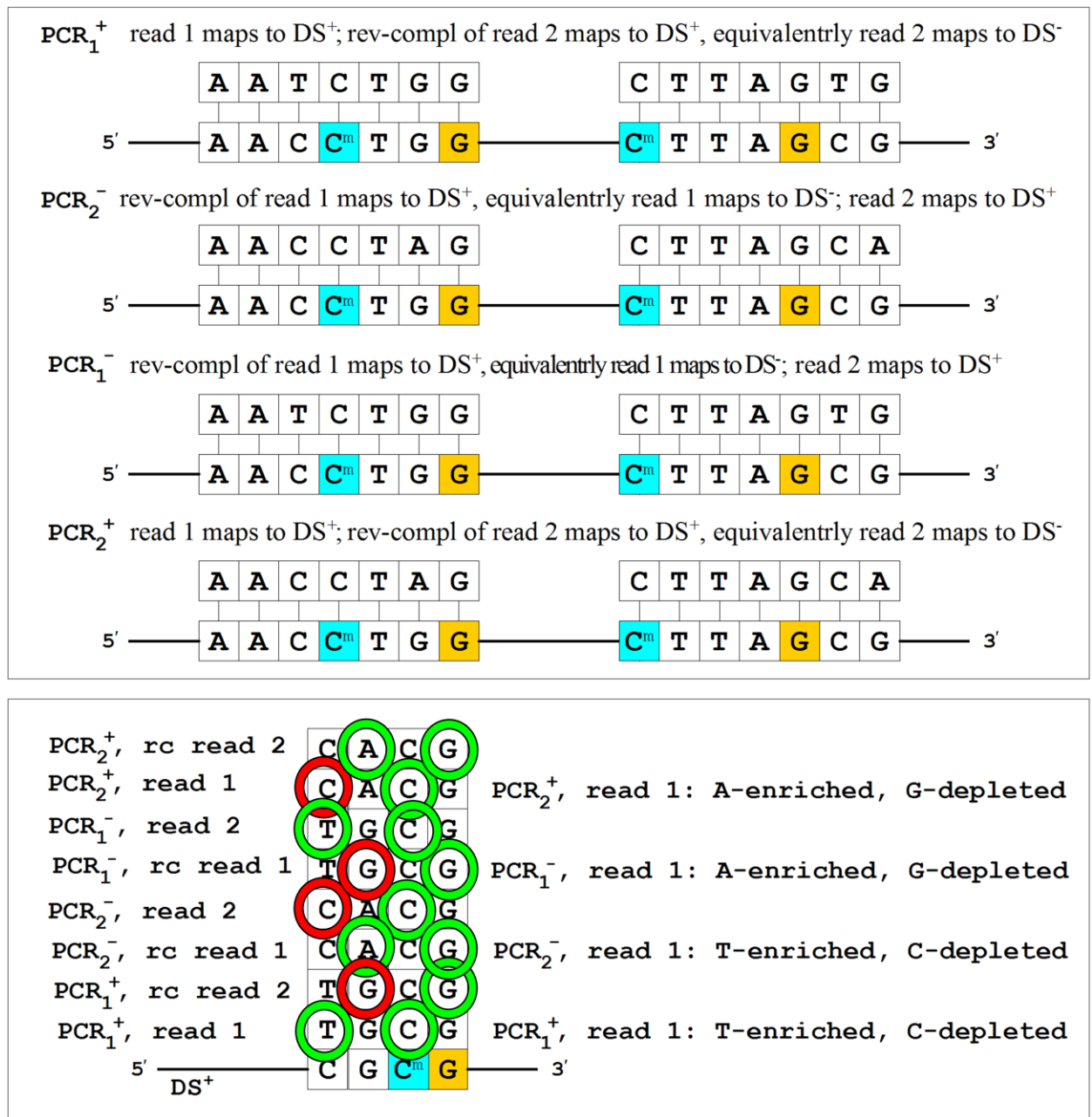
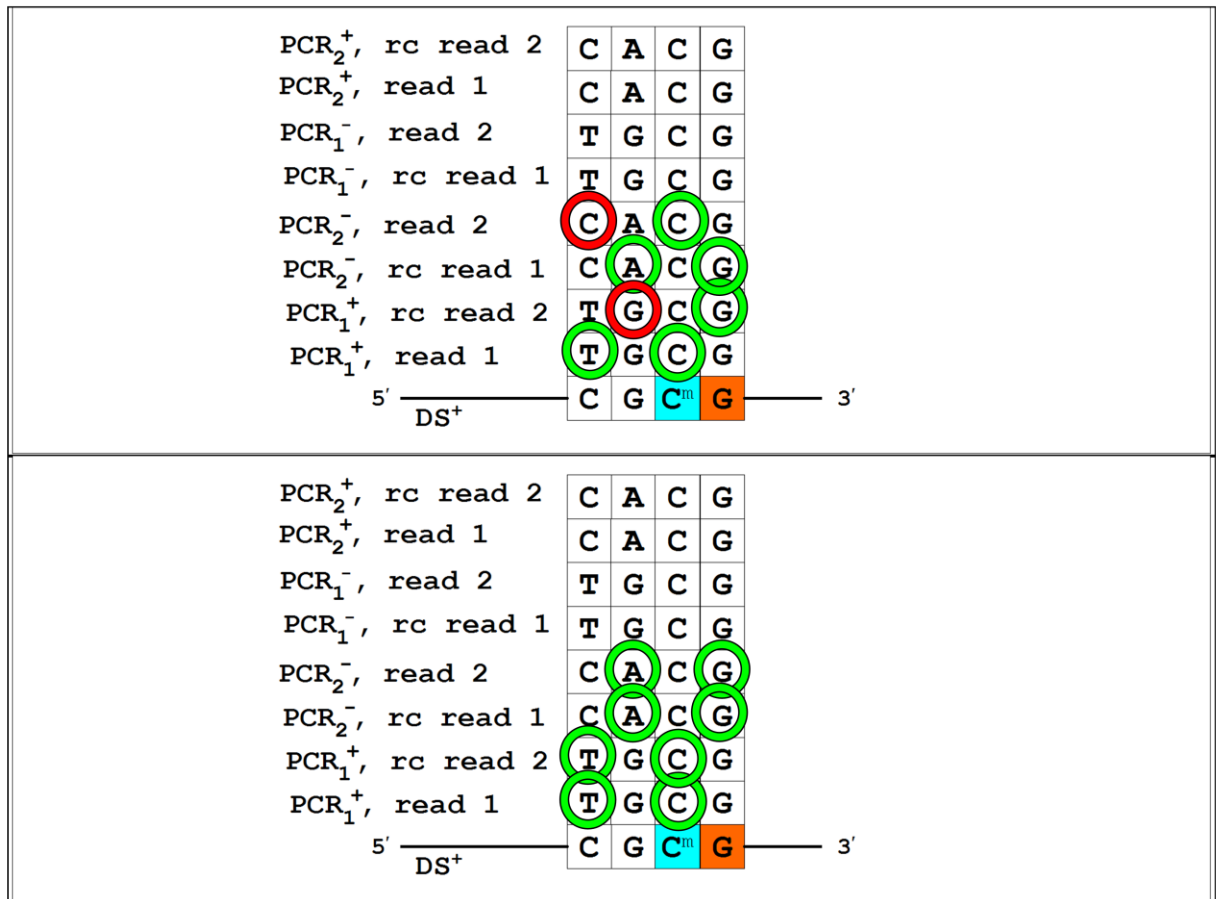


Figure 3. Here, we show mapping of the pairs sequenced from all four strands. The bottom strand shows positive strand of the reference with methylated cytosine shown in blue and methylated cytosine on negative strand shown as G in orange. This figure demonstrates that if we use simplified approach of counting methylated level of cytosines from positive and negative strands, then we introduce a bias in counting methylation level of unmethylated cytosines (either cytosines that are partially methylated across cells of a sample or completely unmethylated). For example, read 2 (mate 2) from PCR1<sup>+</sup> maps to negative strand (its reverse-complement maps to positive strand), and therefore, initially we contributed ACGT-counts from this read toward negative strand. This was incorrect: in this Figure, bias is introduced by counting G from this read (in red circle) toward unmethylated G (i.e. unmethylated cytosine on negative strand). The count of another G from this read (in green circle) toward methylated G did not introduce a bias (since methylated G has only Gs mapped to it), but the coverage of methylated Gs is also not what it would have been were we using the correct counting. Similar bias is introduced when counting contribution of ACGT from mates 2 from PCR2<sup>+</sup>.

From Figures 1-3, we can observe that mate 2 of PCR1<sup>+</sup> strand is reverse-complement 3'-end of PCR1<sup>+</sup>; thus, mate 2 must reflect methylation on positive strand rather than on negative strand. For example, methylated C on PCR1<sup>+</sup> will be G on PCR1<sup>-</sup>, and unmethylated C (which is T) on PCR1<sup>+</sup> will be A on PCR1<sup>-</sup>. Thus, T→C and C→C, where T and C respectively belong to PCR1<sup>-</sup>, must contribute to the count of methylation level of cytosines on positive strand. Similarly,

These changes have been made to **acgt-count** and now ACGT-counting from second mates of pairs sequenced from PCR1+ and PCR2- is done without bias, which is demonstrated in Figure 4.



We received a couple of e-mails concerned that the sequenced reads must be products of sequencing of all four strands. The best way to be sure how many strands have been sequenced is talking to Illumina tech support, but there is also a quick and dirty method to answer this question. This approach will work best with genomes whose ACGT-distribution is close to uniform, or with genomes that CG-rich. We cannot be positive that the way we will explain shortly can be successfully applied to AT-rich genomes. We added a new tool, `check-strands`, that calculates ACGT-distribution in the genome, and in mates 1 and mates 2 of paired-end reads. For the case, when pairs are sequenced only from two original genomic strands, you should notice significant increase of Ts in mates 1 (and decrease of Cs) compared to Ts and Cs in the genome, and significant increase of As in mates 2 (and consequently decrease of Gs) compared to As and Gs in the genome. If four strands are sequenced, then the file with mates 1 contains a mixture of reads sequenced from original genomic strands as well as their reverse-complements. Thus, the increase of both As and Ts (and decrease of counts of Cs and Gs) should be observed in both files: with mates 1 and mates2. This conclusion is derived from the fact that after BS-treatment original

genomic strands are T-rich and C-depleted, and consequently their reverse-complements PCR1- and PCR2+ are A-rich and G-depleted.

Below are the commands to run check-strands:

Single-end reads:

```
./check-strands -r references_names.txt -s reads1.fastq -o output_results.txt
```

Paired-end reads:

```
./check-strands -r references_names.txt -1 reads1.fastq -2 reads2.fastq -o output_results.txt
```

Here, *reference\_names.txt* as before: it is the file with names of files in FASTA with references; *reads1.fastq* and *reads2.fastq* are FASTQ files from Illumina with sequenced reads (for paired-end reads, with mates 1 and mates 2). The output is in *output\_results.txt*; it is self-explanatory with proportions of A, C, G and T (in that order) in references, in reads from *reads1.fastq* and in reads from *reads2.fastq*.

## How to use BRAT with reads sequenced from four strands.

For single-end reads, if reads are sequenced from four PCR-product strands, namely PCR1+, PCR2-, PCR1- and PCR2+, then run BRAT as explained before with option *-u* that will output unmapped reads into a separate file, say *unmapped.txt*. Then use our new tool rev-compl with *unmapped.txt* as follows:

```
./rev-compl -s unmapped.txt -o rev_compl_of_unmapped.txt
```

The output file *rev\_compl\_of\_unmapped.txt* will contain reverse-complement of reads from *unmapped.txt* file. Then map reads from *rev\_compl\_of\_unmapped.txt* as usual (see how to map single-end reads with BRAT).

For paired-end reads, map reads as usual, but use *-u* option to output unmapped reads (use appropriate command of these two):

```
./brat -r references_names.txt -1 prefix_reads1.txt -2 prefix_reads2.txt -pe -bs -o output_results.txt -u  
./brat-large -r references_names.txt -1 prefix_reads1.txt -2 prefix_reads2.txt -pe -bs -o output_results.txt -u
```

The unmapped reads from *prefix\_reads1.txt* will be in *prefix\_reads1.txt.unm* and unmapped reads from *prefix\_reads2.txt* will be in *prefix\_reads2.txt.unm*

Then take reverse-complements of unmapped reads (please provide output files with option *-o*):

```
./rev-compl -s prefix_reads1.txt.unm -o rc_prefix_reads1.txt.unm  
./rev-compl -s prefix_reads2.txt.unm -o rc_prefix_reads2.txt.unm
```

Then run BRAT again on reverse-complements of unmapped reads and **use option A** (use either of the commands below):

```
./brat -r references_names.txt -1 rc_unm_prefix_reads1.txt -2 rc_unm_prefix_reads2.txt -o output_results2.txt -A  
./brat-large -r references_names.txt -1 rc_unm_prefix_reads1.txt -2 rc_unm_prefix_reads2.txt -o output_results2.txt -A
```

The rest of the tools are used as before.

Why does this work? BRAT allows only T-C BS-mismatches if mate 1 maps to positive strand, and only A-G BS-mismatches if reverse-complement of mate 1 maps to positive strand. Thus, when we first time map paired-end reads, only pairs sequenced from the original strands are mapped; after we take reverse-complement, then mapping is done for the pairs sequenced from PCR1- and PCR2+. However, if a read does not have BS-mismatches or the number of inappropriate BS-mismatches (A-G in case when mate 1 maps to positive strand and T-C when reverse-complement of mate 1 maps to positive strand) is less than the total of allowed non-BS-mismatches, then some of the pairs from PCR1- and PCR2+ can be mapped together with the paired-end reads sequenced from the original strands inducing ACGT-count bias showed in Figure 5.

BRAT does not have any counting bias for the case when only original strands are sequenced, however in case of four strands being sequenced, there might be a bias that users must account for by some other means (statistics, etc.) This is shown in Figure 5.

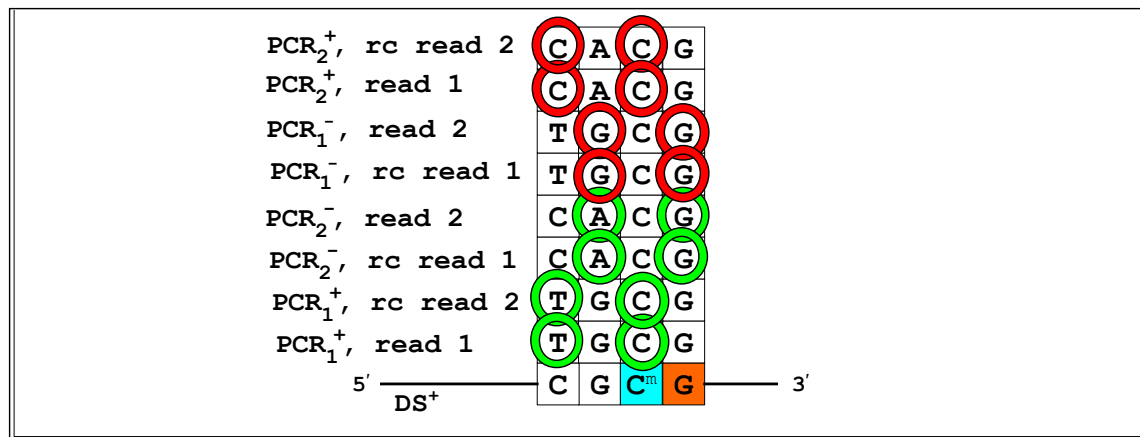


Figure 5. There is no acgt-count bias for the case when two original strands are sequenced, but there might be a bias from mapped paired-end reads sequenced from PCR1- and PCR2+ strands for unmethylated cytosines (or for partially methylated cytosines) if the mates have no BS-mismatches or the number of inappropriate (see above) BS-mismatches is less than or equal to the number of allowed non-BS mismatches.