

# USING BRAT-1.1.5

## 1 SYSTEM AND SPACE REQUIREMENTS

This version differs from the previous only by space usage. This version uses significantly less space than the previous version. In addition, the number of references is practically unlimited.

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,  $S$  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:

$$\text{Space}_{\text{BRAT}} = 269 \cdot 10^6 + 2 \cdot 4N + 24R + (3/8)N \text{ Bytes}$$

$$\text{Space}_{\text{BRAT-large}} = 269 \cdot 10^6 + 4S + 24R + (3/8)N \text{ Bytes}$$

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 2.3 GB on 1 million pairs and human chromosome 1, and BRAT-large on 1 million pairs and an entire human genome uses 1.5 GB.

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 two additional tools: trim 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.

## 2 COMMANDS AND INPUT

To uncompress run:

```
tar zxvf brat-1.1.*.tar.gz
```

To build:

```
cd brat-1.1.*
make
```

This will create four executable programs: brat, brat-large, acgt-count and trim.

**Input format of the reads for BRAT and BRAT-large is raw reads, *i.e.* reads given one per line.** 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

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
```

This will trim bases whose base quality scores are lower than 20 Phred from the ends of reads: bases are trimmed one at a time from both ends of a read until a base with quality score greater or equal than 20 is encountered. To learn more about Phred scores, please visit <http://www.phrap.com/phred/>

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
```

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).

## Commands to run brat and brat-large

BRAT and BRAT-large map raw reads 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*.

## 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 *unm\_prefix\_reads1.txt* file (assuming *prefix\_reads1.txt* was provided in the command line with option `-s`) for single reads and with paired-end reads in *unm\_prefix\_reads1.txt* and *unm\_prefix\_reads2.txt* 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: one per line, 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*);

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

**-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 *amb\_prefix\_reads1.txt* for single reads and in *amb\_prefix\_reads1.txt* and *amb\_prefix\_reads2.txt* files with paired-end reads: raw reads one 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 on 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).

## 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 has 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

### 3 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>

#### 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

#### Output format for acgt-count

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  $i+5(T)$ .

#### 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 read per line (raw reads format).