



HPG-Methyl - User Guide - v3.2.4

HPG-Methyl is an ultrafast and highly sensitive Next-Generation Sequencing (NGS) read mapper and methylation context extractor. Compared with other current mapping and methylation extraction tools, HPG-Methyl offers better sensitivity and shorter execution times even for long reads.

Since the files generated by HPG-Methyl are fully compatible with the files generated by other popular tools, it can be used as a drop-in replacement to accelerate existing methylation analysis pipelines.

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Changelog

The following list contains the releases of HPG-Methyl available on the repository. This document is relevant for the 3.2, 3.1 and 3.0 versions, older versions are unsupported but available to download.

v3.2.4 (current)

• Minor fix in BWT index generation.

v3.2.3

• Minor fix in bam generation.

v3.2.2

• Included automatic detection of the number of chromosomes of the reference genome.

v3.2.1

• Included option in order to set the number of chromosomes of the reference genome.

v3.2

• paired-end sequenced reads supported.

v3.1.4

• Fixes a minor bug in the BAM file generation

v3.1.3

• Minor bugfixes and optimizations.

v3.1.2

• Alignment score fixed.

v3.1.1 (current)

• Minor bugfixes and optimizations.

v3.1

- Now methylation context information is stored in the output BAM file, using the same format as Bismark (http://www.bioinformatics.babraham.ac.uk/projects/bismark/).
- Minor bugfixes and optimizations.

v3

 Optimized using bidirectional Burrows-Wheeler Transform and a modified parallel pipeline.

v2

• Improved version with CpG context information switch available as a command line option.

v1

• Original version with CpG context information disabled.

1) Quick Start Guide

1. Downloading and extracting the software

HPG-Methyl runs on a terminal. Get into the destination directory and download the latest compiled version of HPG-Methyl from the public repository:

\$ sftp anonymous@clariano·uv·es:/src/version_3_2-HPG-Methyl2/hpg_methyl_3_2_x86_64·tar·gz ·

When prompted for a password for any download, use anonymous.

Extract the contents of the tarball:

\$ tar xvzf hpg_methyl_3_2_x86_64·tar·gz

2. Process the reference genome

You need a reference genome in FASTA format to compare your sequenced reads with. If you do not have an available genome, an example *homo sapiens* genome can be downloaded from the repository. Create a directory to store your genome and download:

- \$ mkdir genome/
- \$ sftp anonymous@clariano·uv·es:/datasets/Homo_sapiens·GRCh37·68·dna·fa genome/



The example genome dataset size is **3,15 GB**. Make sure that your connection is stable and fast enough to download the file.

HPG-Methyl needs a special index to access the reference genome, which has to be generated before doing any read mapping. This process has to be done **only once per reference genome**.

Create a directory to store the index genome, and let HPG-Methyl generate it. This process will take a while depending on the CPU processing power of your machine. Extensive information on the HPG-Methyl command line options is available at the end of this document:

- \$ mkdir genome/index/
- $$\cdot/hpg-aligner\ build-index\ -g\ \cdot/genome/Homo_sapiens\cdot GRCh37\cdot68\cdot dna\cdot fa\ -i\ genome/index/\ -r\ 10\ --bs-index$



The genome index files take a considerable amount of space. The directory holding the index must have at least 250 GB of available free space. Storing the index in fast media is recommended (like a solid state drive or a fast network drive).

3. Mapping and methylation analysis of the sequenced reads

When the genome index creation has been completed, sequenced read files can be mapped to the reference genome and the methylation context analysis can be performed. The read files must be in the FASTQ format and must contain only single-ended reads. Paired-ended reads are unsupported. If you don't have any available sequenced read data, download an example read from the repository:

\$ sftp anonymous@clariano·uv·es:/datasets/real/SRR309230_1_075nt_15M·fastq genome/



The example sequenced read file size is 3,31 GB. Make sure that your connection is stable and fast enough to download the file.

When the download has been completed, run HPG-Methyl to align the sequence with the reference genome and generate the methylation context analysis. This process will create three output files:

- **10 alignments.bam** Binary SAM file containing the aligned reads and methylation context information for each read. The file can be converted to a text SAM file using *samtools*. The methylation context information is stored in SAM tags, following the *Bismark* tag format.
- **©** CpG_context.txt / CHG_context.txt / CHH_context.txt / MUT_context.txt CSV files containing the context-dependent methylation analysis information.

HPG-Methyl has a multi-threaded pipeline and can fully exploit all the processing cores in your computer platform. Use the following command to run the process with 4 threads:

\$ ·/hpg-aligner bs -i genome/index/ -f genome/SRR309230_1_075nt_15M·fastq -o · --cpu-threads 4 -write-mcontext



Using more threads only improves performance if your system has enough CPU cores or threads to do so. The recommendation is to run as many threads as virtual processors are available in the machine.

2) Building

1. Downloading the source and file structure

First, download the source tarball from the SFTP repository. This manual corresponds to the latest version available, HPG-Methyl v3.2, located on the /src/version_3_2-HPG-Methyl2 directory. The complete connection details to download the tarball are:

Address	clariano·uv·es
Username	anonymous
Password	anonymous
Protocol	SFTP
Complete path	sftp://anonymous:anonymous@clariano·uv·es/src/version_3_2-HPG- Methyl2/hpg-methyl2_v3·2·tar·gz

Extract the source tarball into the desired directory as usual:

\$ tar xvzf hpg-methyl2_v3_2git a·tar·gz

The extracted tarball has the following directory structure:

/	Root directory containing the SCons build script, README and COPYING	
	notices.	
/src	Source code of HPG-Methyl	
/lib	Source code of OpenCB bioformats and common libraries. The version of libraries included with HPG-Methyl is modified and might not be compatible.	
	the version available at GitHub.	
/bin	Directory where the compiled binary will be generated.	

2. Compilation and installation

Building HPG-Methyl requires having a working installation of GCC 4.4 or greater, SCons and the following software packages, which can be installed using *apt-get* on Ubuntu or Debian based distributions or *yum* on Red Hat based distributions:

Ubuntu / Debian

Fedora / CentOS / Red Hat

zlib1g-dev libcurl4-gnutls-dev libxml2-dev libncurses5-dev libgs10-dev check zlib-devel
libcurl-devel
libxml2-devel
ncurses-devel
gsl-devel
check-devel

When all packages are installed, get into the directory that was created and build the software using the command:

\$ scons

When the process finishes, the compiled binary will be available at the */bin* folder, and it can be run from there. If a system-wide installation of the tool is desired, copy the binary to the applications directory:

\$ cd bin/

\$ cp ·/hpg-methyl /user/bin/hpg-methyl

3. Extending and debugging



This section is only relevant if you plan to extend or debug HPG-Methyl. Otherwise, you can skip it.

In order to make changes to HPG-Methyl, modify the code on the /src or /lib directories and build the toolchain. To compile without optimizations and with debug symbols, use the command:

\$ scons debug=1

The source is bundled with Eclipse and Visual Studio Code projects solutions to ease the development process. Under Eclipse, import the tarball directory into the workspace to work with the source code. Under Visual Studio Code, use the Open Folder command and open the tarball directory.

The Eclipse solution requires the SConsolidator plugin to integrate with SCons. The Visual Studio Code doesn't have any additional dependencies. Check the included VS Code tasks for building, debugging and running information.

3) Running

The syntax for running HPG-Methyl is:

\$ hpg-methyl <mode> <options>

The available modes are **bs**, used to map bisulphite-treated sequences and **build-index**, used to create the Burrows-Wheeler index of the reference genome.

The bisulphite mapping mode aligns the sequenced reads using the Burrows-Wheeler Transform, the Smith-Waterman algorithm and several post-process and pre-process filters. The details of the pipeline are beyond the scope of this manual, and they are available on an article. Contact the developers for more information.

The pipeline has been designed to be easily tunable to any particular application, and most algorithm parameters can be adjusted through command line options. For most applications, the default values should suffice since all algorithms have been throughly tested and optimized to obtain the best performance in a variety of situations.

Following is the description of the different algorithms on HPG-Methyl and their tunable parameters. The annex available at the end of this document shows the complete list of optional parameters and their description.

1. Smith-Waterman Algorithm parameters

The Smith-Waterman algorithm is used to perform the mapping on reads that were not fully mapped on the Burrows-Wheeler stage, allowing gaps and mismatches with the reference genome.

The weight matrix for the Smith-Waterman algorithm can be tuned through the sw-match, sw-mismatch, sw-gap-open, sw-gap-extend and sw-min-score optional parameters. The default SWA weight table has been optimized by hand to have the best performance on most cases.

Minimum score	0,8
Match score	5,0
Mismatch score	-4,0
Gap open	10,0
Gap extend	0,5

Default SWA weight table.

2. Filtering and seeding parameters

Between the different mapping stages and at the end of the pipeline, there are several filtering and seeding stages which are used to accelerate the mapping process.

The seeding and CAL stages default parameters are shown below:

Minimum CAL size	30
Maximum distance between seeds	100
Maximum seed size	23
Minimum seed size	16
CAL flank size	5
Number of seeds per read	10
CAL umbral length factor	4
Read minimum discard length	100
Read maximum inner gap	-1

Seeding and CAL stages default parameters

When the mapping (and optionally methylation analysis) have been completed, there is a last filtering stage where the alignments being written to the BAM file can be filtered (only for single-end mode). The default parameters are shown below:

Report all hits	No
Report only best hits	Yes
Report only n first best hits	No
Report only <i>n</i> first hits	No

Reporting stage default parameters

3. Methylation context analysis parameters

If the *write-mcontext* command line option is enabled, the last pipeline stage will obtain the methylation status of the current read and will store it on the tags of the output BAM file. After all reads have been mapped, the context-dependent methylation information is stored into text files.

When generating the Burrows-Wheeler index, use the *--bs-index* option to create an index compatible with the methylation analysis stage.

4. Paired-end mode

HPG-Methyl is able to process paired-end sequenced reads. In order to do that, the `--paired-mode` command line option must be set to 1 (by default is 0, single-end mode).

The paired-end sequences must be separated in two differents fastq files, but only one bam file with the mapped alignments will be created.

If paired-end mode is set, a mandatory command line `-j` / `--fq2` / `--fastq2` is required, in order to set the path to the second FASTQ sequence file.

Following is an example of use:

hpg-methyl bs -i <index-directory> --paired-mode 1 -f <fastq-file-path-1> -j <fastq-file-path-2> -o <output-directory> --cpu-threads <thread-count>

Annex: Complete list of command line options

-f /fq /fastq <path></path>	Path to the sequenced read file in FASTQ format.
-i /bwt-index <path></path>	Path to the BWT index directory. It must have been
-l /log-level <level></level>	previously generated using the <i>build-index</i> option. Console debug message level. <level> must be one of the following numeric values, from more verbose to less verbose.</level>
	 0: Show all debug messages (used to troubleshoot the application). 1: Show information messages, warnings and errors. This is the default log level. 2: Report only warnings or worse. 3: Report only errors or worse. 4: Report only fatal errors. 5: Don't log any messages.
report-all	Report all alignments.
-o /outdir	Path to the output directory.
cpu-threads	Number of CPU threads to use.
-r /index-ratio	Index compression ratio for the Burrows-Wheeler Transform. Used when creating the genome index.
min-cal-size	Minimum CAL size.
max-distance-seeds	Maximum distance between seeds.
read-batch-size	Read batch size in bytes.
write-batch-size	Write batch size in bytes.
seed-size	Number of nucleotides in a seed.
min-seed-size	Minimum number of nucleotides in a seed.
cal-flank-size	Flank length for CALs.
sw-match	Match value for Smith-Waterman algorithm.
sw-mismatch	Mismatch value for Smith-Waterman algorithm.
sw-gap-open	Gap open penalty for Smith-Waterman algorithm.
sw-gap-extend	Gap extend penalty for Smith-Waterman algorithm.
sw-min-score	Minimum score for valid mappings.
-t /time	Profile timing of all the application stages and log the elapsed time when the execution finishes through the console.
-s /stats	Generate and save execution statistics.
-h /help	Print the application help.

File prefix name.

--prefix

-g /ref-genome -j /fq2 /fastq2 <path></path>	Path to the reference genome. Path to the second sequenced read file in FASTQ format (paired-end mode).
paired-mode <n></n>	 Enable paired-end mode. 0 = single-end mode (default). 1 = paired-end mode. Minimum distance between pairs in order to consider that
paired-min-distante <n>paired-max-distante <n></n></n>	reads map in proper pair (paired-end mode). Maximum distance between pairs in order to consider that reads map in proper pair (paired-end mode).
report-n-best <n></n>	Report the <n> best alignments. Report <n> hits.</n></n>
report-n-hits <n></n>	Number of seeds per read.
num-seeds <n></n>	•
min-num-seeds <n></n>	Minimum number of seeds to create a CAL (if -1, the maximum will be taken).
filter-read-mappings <n></n>	Reads that map in more than <n> locations are discarded.</n>
filter-seed-mappings <n></n>	Seeds that map in more than <n> locations are discarded.</n>
report-best	Report all alignments with best score.
bs-index	Indicate the use of bisulphite generation of the index. Used when creating the genome index.
write-mcontext	Save the methylation context report files (CHG, CHH, CpG, MUT) and embed the alignment methylation information in the BAM file.
umbral-cal-length-factor <f></f>	CAL is accepted as a candidate if it has been mapped more than CAL size / <f></f>
min-read-discard <n></n>	Discard small anchors for reads larger or equal than <n>.</n>
max-inner-gap <n></n>	Recursive Burrows-Wheeler stops when inner gap is lower than <n>.</n>