

Correcting Methylation Calls in Low-Mappability Regions

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Abstract

DNA methylation is an important component in vital biological functions such as embryonic development, carcinogenesis, and heritable regulation. Accurate methods to assess genomic methylation status are crucial to its effective use in many scenarios, especially in the detection and diagnosis of disease. Methylation aligners, such as Bismark and bwa-meth, frequently assign MapQ values to reads which are significantly higher than can be supported by the uniqueness of the region they are mapped to. These incorrectly high MapQs result in inappropriate methylation calling in repetitive regions. We observe reads that should map to separate locations (possibly having different methylation states) actually end up mapping to the same locus, causing apparent mixed methylation at such loci. Methylation calling can be improved by using Bismap mappability data to filter out insufficiently unique reads. Simply filtering out Cs in insufficiently unique regions is not adequate as it is prone to overfiltering Cs in small mappability dips. These Cs can in fact be called using reads anchored in a nearby mappable region. We have created a patch for the MethylDackel methylation caller to perform read-based filtering. Read-based filtering resolves some of the apparent mixed methylation to either 0% or 100% methylation. We examined methylation calls with and without read-based filtering in or near the 7830 genes containing ClinVar variants in a methylation sequencing data set from the NA12878 cell line and in tumor samples. Examining low mappability Cs in the NA12878 data set revealed 1405 mixed methylation Cs were corrected to 0% methylation, and 2577 mixed methylation Cs were corrected to 100% methylation.

Introduction

As DNA methylation status can have a significant biological function [19], it is important that there be an accurate way of calling methylation on a genome. Although there are multiple varieties of DNA methylation, a significant type is methylation of cytosine to 5-methylcytosine [2]. Data on DNA cytosine methylation state can be gathered using a methylation sequencing technique (see Figure 1), for example bisulfite sequencing [3]. In bisulfite sequencing,

unmethylated cytosines are deaminated to uracil by the addition of sodium bisulfite. 5-methylcytosines are not affected. Since uracil sequences as thymine and 5-methylcytosine sequences as cytosine, positions of unmethylated Cs in a reference sequence can be identified by C->T transitions[3].

It is also possible to use an enzymatic method employing TET2 to oxidize 5-methylcytosine and an APOBEC enzyme to deaminate unmodified cytosines to uracil. While sodium bisulfite treatment produces other DNA damage, this enzymatic method deaminates with more precision (forthcoming publication).

Whichever method is used to deaminate cytosines, sequence data is typically aligned to a reference genome using a methylation-aware aligner [5], which is specifically designed to handle the C->T transitions in methylation sequencing data during alignment of the reads to a reference. Once aligned, the data can be passed through a methylation caller such as MethylDackel [18] or bismark_methylation_extractor [12], which will use the resulting reads to determine the methylation status of a particular cytosine (see Figure 2). The resulting data shows the methylation status of each cytosine in the genome and can therefore be used to find and study biologically significant DNA methylation sites.

A read must be unambiguously placed if it is to provide information about a specific locus. Reads that equally match more than one area of the reference genome should not be used to assess methylation of any given C. To avoid calling Cs using reads derived from multiple genomic loci, methylation aligners (and read aligners in general) assign a MapQ value to each read alignment (see Figure 2). According to the SAM specification [6], MapQ is defined as: “ $-10 \log_{10} \Pr\{\text{mapping position is wrong}\}$, rounded to the nearest integer”. MapQ indicates how uniquely placed a read alignment is, that is, in how many other places could the read align to the reference. A low MapQ means that the read may align in many places throughout the genome (for instance, a read of centromeric satellite DNA would likely have a very low MapQ). A high MapQ indicates that the read likely aligns where it is placed and nowhere else in the genome.

A methylation caller can use accurate MapQ values to filter out reads with multiple placements in the genome, allowing the resulting methylation calls to accurately reflect their specific loci.

Results

While evaluating the methylation aligner bwa-meth [16], we observed a significant number of reads with unexpectedly high MapQ values in repetitive regions (e.g. centromeres). After observing these high MapQ reads in the centromere and larger repetitive regions, we investigated to see if smaller regions might also be too repetitive to support the high aligner MapQ estimates observed. We identified repetitive regions using data from Bismap [10], a tool that counts the number of occurrences of every single K-mer of a particular length (in this case, k=100) in the genome to create a mappability score for every base in

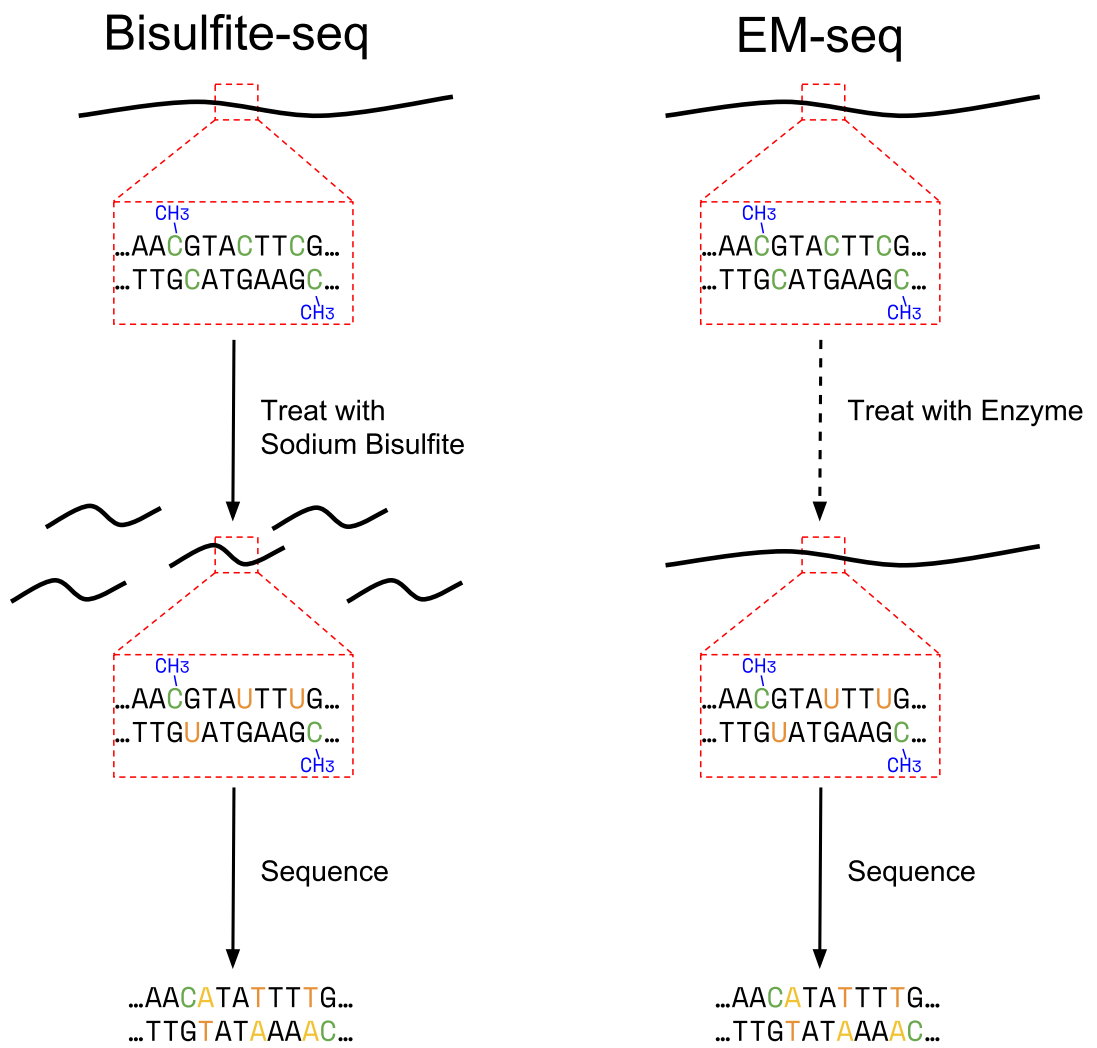


Figure 1: Overview of Methylation Sequencing Methods. <todo: Treat with Enzyme -> Tet2-> APOBEC

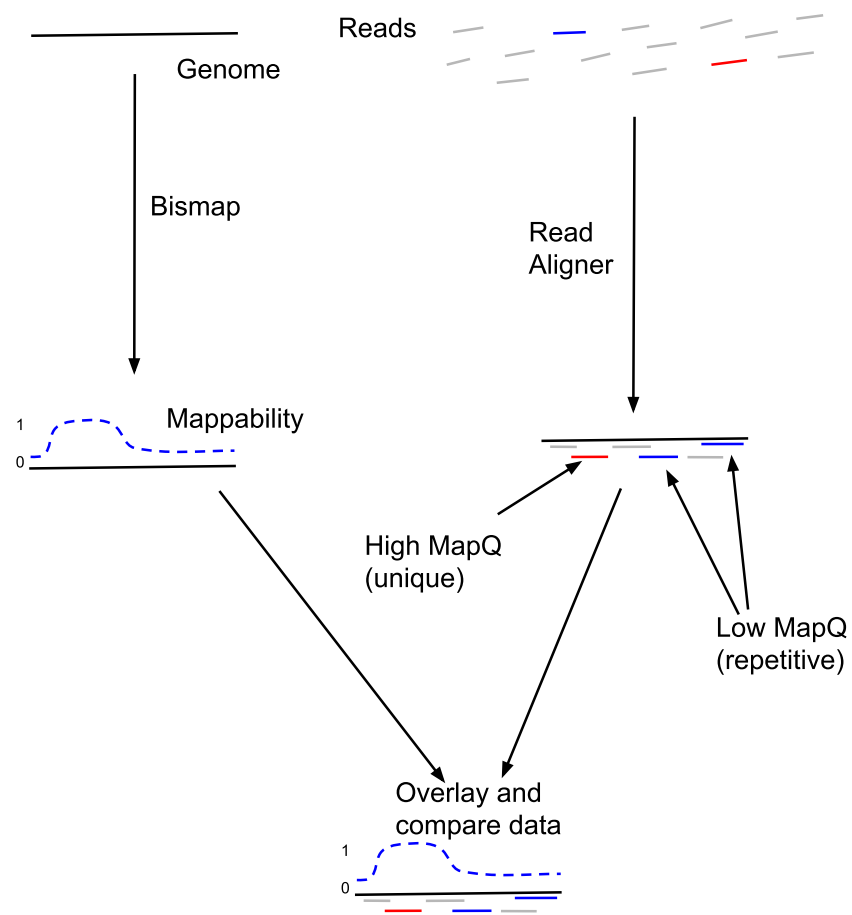


Figure 2: Experimental Overview

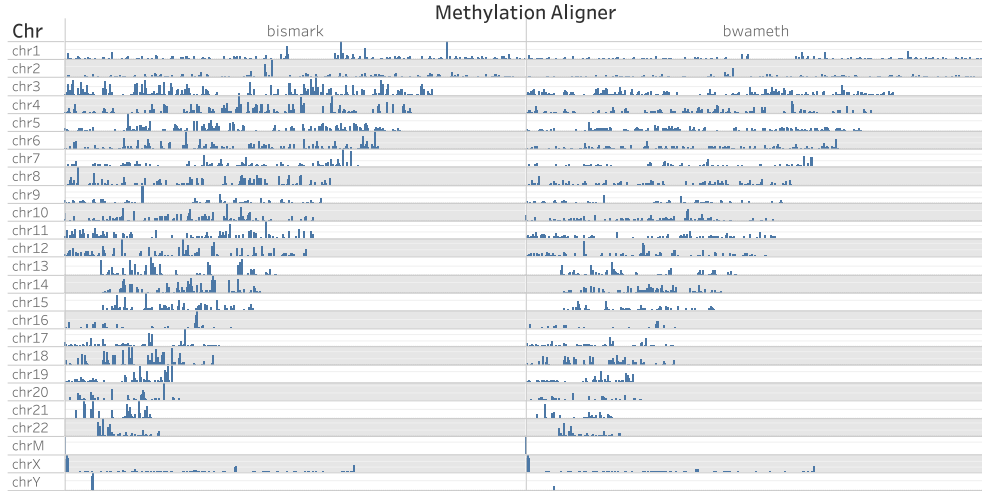


Figure 3: <WRONG CAPTION!!!!> Example of MapQ Issue. On the Cov Difference track, red means that Bismark has higher coverage and blue means bwa-meth has higher coverage.

the Grch38 reference. <bases this on assumption that each locus is either 0 or 100%> Bismap takes the effect of C->T conversion into account and therefore produces data which is applicable in the context of methylation sequencing. Reads entirely contained within a region of low mappability should not have high MapQ values due to their repetitiveness, however we observed many high-MapQ reads in regions with very low or zero Bismap mappability (see Figure 4).

While low MapQ can indicate repetitiveness, even stringent MapQ thresholds cannot reliably select reads for safe methylation calling in regions containing repetitive DNA. We considered excluding methylation calls on Cs found in low-mappability regions (e.g. using bedtools), but rejected this approach because it is prone to both over- and underfiltering. Cs in short, unique regions would be kept (underfiltered) even if the surrounding DNA is repetitive (see Figure 5). However, this does not happen often. Overfiltering of Cs in short repetitive regions is a larger problem though. In this scenario, a C located in a small dip in mappability would be eliminated (overfiltered) despite coverage from read pairs anchored in nearby unique regions (see Figure 7). In practice, underfiltering is rare ($x\%$ of Cs) but overfiltering is much more common ($y\%$ of Cs) (see Figure 6). In cases where reads from multiple low-mappability regions are placed onto one region (creating a coverage spike), filtering by coverage could possibly also remove the problematic region, but in cases where reads are simply mixed among multiple low-mappability regions (not creating a coverage spike), coverage filtering would not be effective at removing the affected regions.

To reliably filter out only problematic read pairs (those where both mates

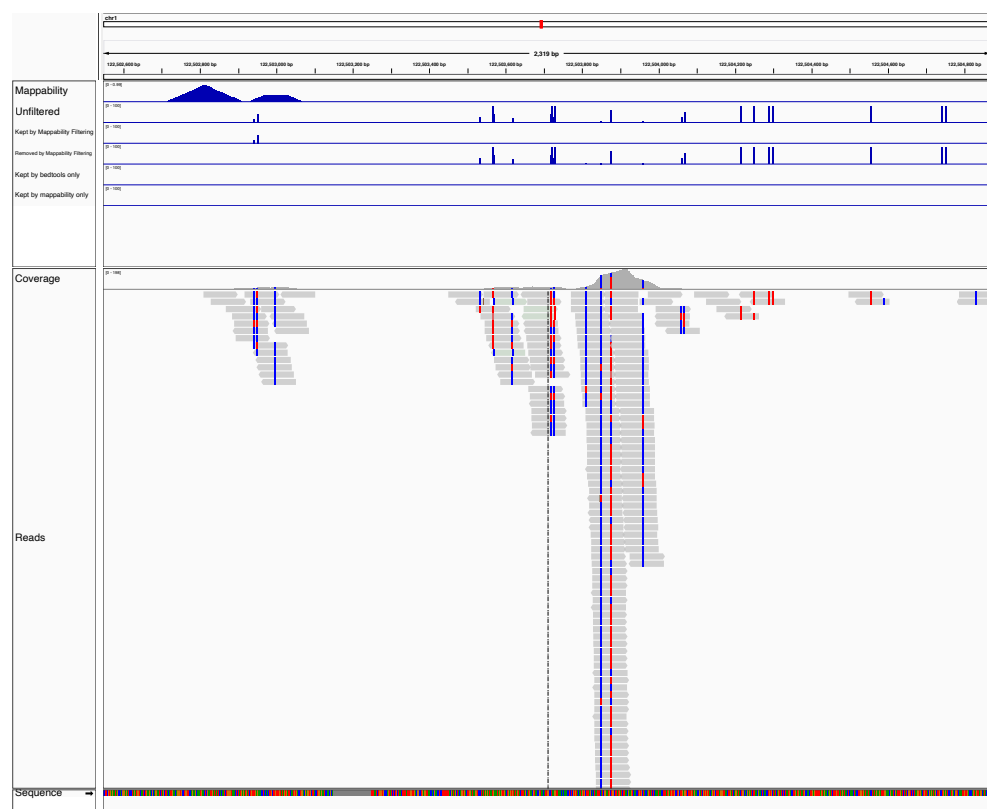


Figure 4: An example of how MapQ filtering does not remove all poorly mapped reads.

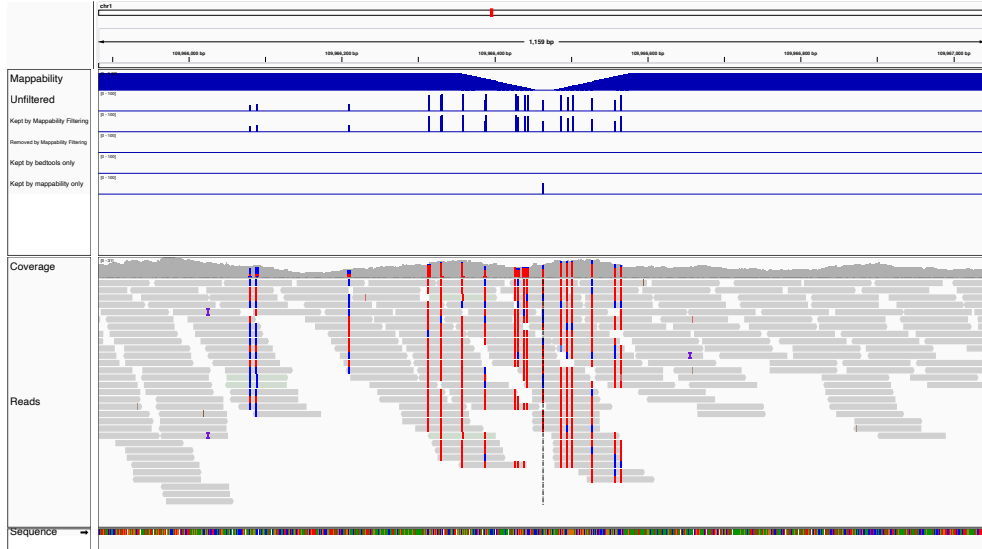


Figure 5: An example of a C incorrectly discarded by coordinate filtering.

are placed in low mappability regions) we modified the MethylDackel methylation caller to accept a bigWig file of low-mappability regions to exclude from analysis. This alignment filtering approach precisely eliminates only those reads in repetitive regions and does not incur a significant cost in terms of execution speed (10:14 min with the patch 9:55 min without, when run with 20 threads on the Grch38 reference genome). (see computational methods for details)

To focus on the incorrect alignments and methylation calls that have biological and medical significance, we examined methylation calls in GENCODE genes [7] that contain variants listed in the ClinVar [13] database of disease-associated variants. Using our alignment filtering approach we successfully avoided calling 15,539 Cs in these important regions in a 50ng enzymatic sequencing sample which have insufficient mappability to support methylation calling. Briefly, to determine that we avoided calling a region has insufficient mappability to support methylation calling, we intersected read alignments and MapQ data with Bismap low mappability regions, counting Cs in regions where the MapQ is higher than should be possible given the mappability (see Figure 2).

For comparison, we also examined the Bismark methylation aligner in the same manner. We observed that compared to bwa-meth, Bismark produced more high MapQ reads in repetitive regions (see Figure 3). In addition, Bismark and bwa-meth use different systems to define values for MapQ. Bismark (as a result of using Bowtie2 for alignment) reports MapQ based on number of reference mismatches, producing values between 0 and 42 [1, 9]. Bwa-meth (using BWA-MEM for alignments) follows the SAM specification in estimating probabilities in the MapQ field.

todo: filtering effect em-seq vs wgbs (fig) for bwa-meth

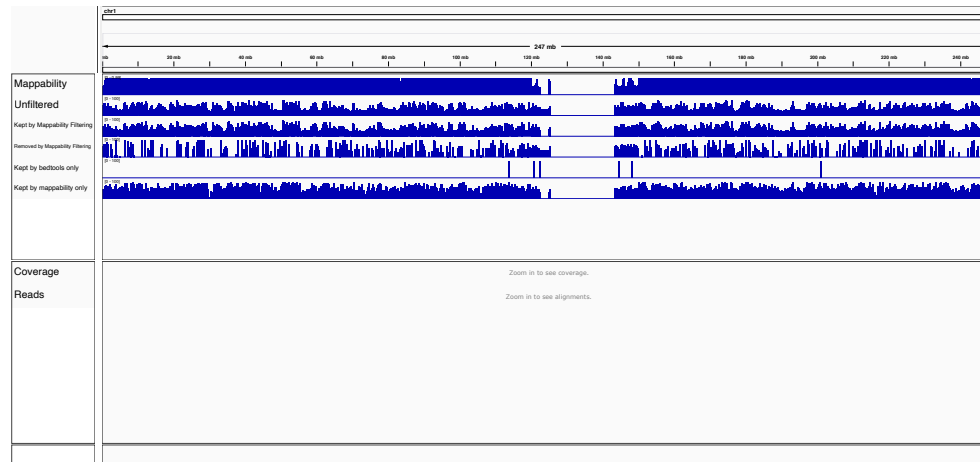


Figure 6: An overview of the occurrences of these scenarios in grch38 chromosome 1.



Figure 7: An example of a C incorrectly kept by coordinate filtering.

todo: filtering effect em-seq-vs wgbs for bismark (this is the data I'm working on)

todo: # cs affected (worst) bismark vs filtered vs bwa-meth vs filtered (best) (fig?) (I can get this from the emseq/wgbs data, it's a simple bedtools operation)

todo: group Cs by magnitude of change in methylation post filtering (fig, heatmap?) (best done by comparing both %methyl and read count, maybe we should discuss how to layout this figure though)

todo: effect on ClinVar

todo: examples of specific effects in ClinVar (I have 2 of these from my talk)

todo: examine GEO dataset (maybe, depends on time/progress)

Materials and Methods

DNA Methylation Sequencing

Materials

Libraries were prepared from 10, 50, and 200 ng of genomic DNA from the NA12878 cell line (Coriel). This input was supplemented with a small amount of fully-methylated Xp-12 DNA <is there a specific source for this, e.g. a company and/or product name?>, lambda phage DNA (NEB #N3011), and a pUC19 plasmid (NEB #N3041) treated with M.SssI CpG Methyltransferase (NEB #M0226).

Whole Genome Bisulfite Libraries

Libraries were prepared using the Ultra II DNA library prep kit <product number?> before being Bisulfite converted according to <protocol>. <anything about the EM-seq samples?>

Sequencing

Libraries were pooled and sequenced with diverse libraries (~10%) on 2 flow cells <is this still correct with the 2 other masses added?> of an Illumina Novaseq 6000 [8] using the S2 chemistry<citation>. We acquired 1.55 billion 99bp paired-end reads for the enzymatic sequencing method and 1.60 billion paired-end reads for the Bisulfite converted libraries.

Computational Methods

In order to run the analysis, the following data and tools were used:

The GRCh38.p11 <re-do with .p13? .p11 is out of date> analysis set (hereafter referred to as “grch38”) supplemented with phage T4, phage lambda <should lambda be written out as a word or as λ?>, phage Xp12, and pUC19 contigs was used throughout [4]. A VCF file of disease-associated variant sites was downloaded from ClinVar (see supplemental data for file date) and a GFF

file of the GENCODE v31 <re-do with v38? v31 is out of date> gene annotations were used. The mappability data was the 100bp multi-read bigWig file downloaded from Bismap (see supplemental materials for link). A detailed diagram of the analysis pipeline is found in Figures 8 and 9. Tools used include bedtools [17], samtools [14], GNU awk, MethylDackel, bigWigToBedGraph [11], BEDOPS [15], GNU sort, GNU head, bwa-meth, and Bismark. GNU sort and head are required since some of the functionality needed (specifically, the ability to parallelize sorting and the ability to use a negative value with the -n option for head to count lines from the end of the file) is not present in BSD sort and head.

The sequencing reads were aligned using the bwa-meth aligner using default options. Methylation calling was performed on the resulting BAMs using MethylDackel v0.5.2. MethylDackel uses a default value of 10 as a minimum MapQ which should include mostly single locus reads, however inaccurate MapQs lead to reads being incorrectly included in methylation calling. In order to eliminate reads in low-mappability regions, a patch was created for MethylDackel which allows it to take as an input a bigWig file which is then used to filter out read pairs (this patch currently only supports paired-end reads) where neither mate intersects a high-mappability region. The patch allows for user configuration of the low mappability threshold and the number of bases which must be equal to or above that threshold in order for the read pair to be kept (the defaults are a low mappability threshold of 0.01 and to require 15 bases that are greater than or equal to that threshold in a single read). The filtering algorithm has been optimized both by loading the mappability data into memory before calling, and through the use of a custom run-length compressed binary file format that we here term BBM (Binary BisMap). This format can store the mappability data for the hg38/grch38 human genome in 143 MB, compared to 1.11 GB when stored as a bigWig, giving a compression ratio for this dataset of 7.78:1.

The ClinVar VCF and GENCODE GFF were combined using bedtools and BEDOPS into a single BED file listing all GENCODE annotation regions that overlap one or more ClinVar variants (these regions will be referred to as “ClinVar regions”), which will be called the “ClinVar regions BED”. The Bismap bedGraph (which, as is standard for bedGraph files, did not contain zeroes) and an FAI index of the grch38 reference genome FASTA file were processed with bedtools, awk, and GNU sort to obtain a bedGraph containing all mappability data, including zeroes. This file will be referred to as the “Bismap complete bedGraph”. It was then combined with the ClinVar regions BED using bedtools map to create a file containing the minimum and mean mappability for each gene with a ClinVar variant.

The Bismap complete bedGraph was then filtered using awk to produce a file containing only low mappability regions (mappability < 0.01). The file with minimum and mean mappability for every ClinVar region was filtered likewise on minimum mappability. The two resulting files (one of low-mappability regions, one of ClinVar regions with low minimum mappability) were combined to produce a file of all low mappability regions that are in ClinVar regions. <what

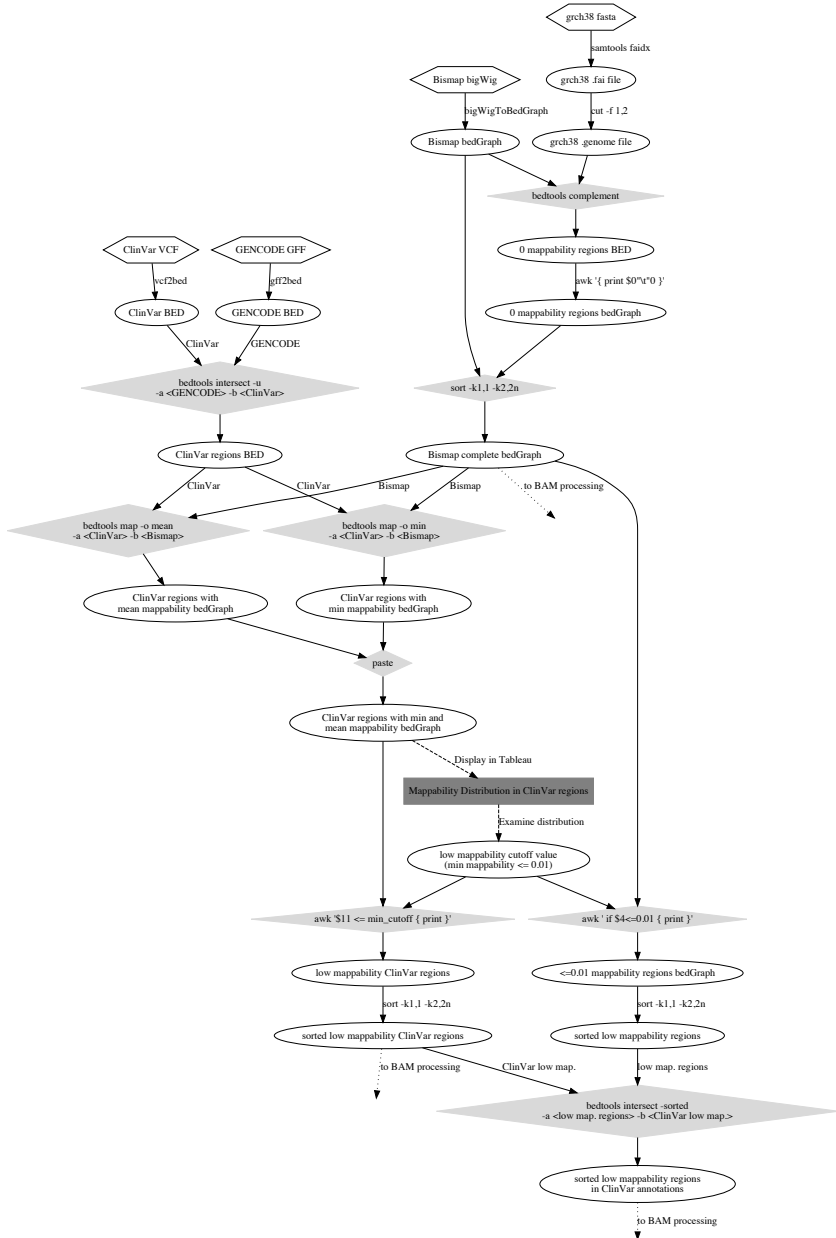


Figure 8: Detailed description of preprocesing steps in data analysis. <todo: add legend with figure features, boxes, ovals, edge types, etc. parts A and B with >

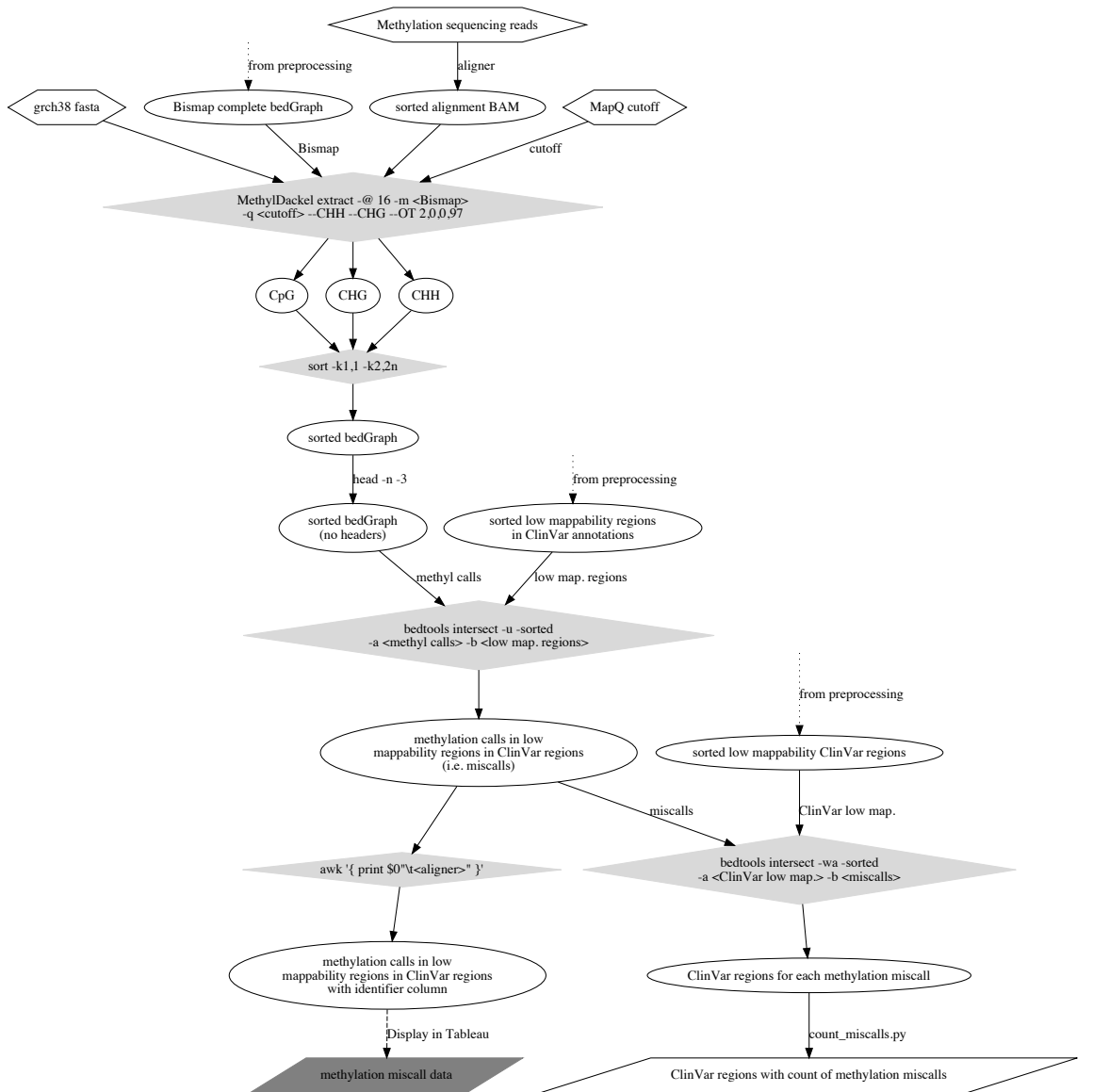


Figure 9: Detailed description of BAM processing steps in data analysis. <todo: add legend with figure features, boxes, ovals, edge types, etc. parts A and B with >

did I do for partial overlaps here??>

Alignments of 2x99-bp paired-end enzymatic sequencing and whole genome bisulfite reads were processed using MethylDackel (with and without the custom patch) using a minimum MapQ cutoff of 10 and the default settings mentioned above and combined with the file of all low mappability regions in ClinVar regions to produce a list of all miscalls (this will be referred to as the “miscalls file”). A miscall, as used here, is defined as a methylation call that was filtered out using the new MethylDackel mappability filtering, but was not filtered out by basic MapQ filtering.

The miscalls file was intersected with the file of ClinVar regions with low minimum mappability to produce a file of ClinVar regions with miscalls. The -wa option for bedtools intersect was used here, which writes a copy of the ClinVar region to the output file for each miscall in the region, in order that this file would contain multiple copies of each region, one per miscall in the region. These duplicates were then used to count miscalls by feeding the data to a custom Python script (invoked in and found alongside with the Nextflow script in the supplemental materials) which counted and combined the duplicates, producing a list of all ClinVar regions with miscalls and how many miscalls are in each region.

Since this analysis was run for three different input masses and two sequencing protocols, the miscalls files were also processed through awk to add a field specifying which input mass, sequencing protocol, and MethylDackel filtering setting were used. The same field was added to the lists of all ClinVar regions with miscalls and counts described previously. All files were examined and compared in Tableau[®].

To compare the behavior of Bismark with bwa-meth, this analysis was re-run with the Bismark aligner <with what alignment options>. The name of the aligner was added to the field specifying the input mass, protocol, and MethylDackel filtering settings present in the miscalls files and the lists of all ClinVar regions with miscalls and counts.

<still getting data>

Discussion

Reads placed with falsely high confidence have cascading detrimental effects on methylation calling, differential methylation assessment, and assessment of phenotypes associated with methylation status. Because of the more accurate MapQ values, decreased run time, and more flexibility to separate methylation calling from alignment, we recommend the use of bwa-meth for alignment and MethylDackel with MapQ > 20 for methylation calling <where is the test of MapQ 10 vs 20? or was >10 intended?>. To further improve accuracy of methylation assessment, reads with both mates in low mappability regions should be excluded.

<waiting on results>

Supplemental Materials

The Nextflow script used to analyze this data can be found at <to be added>. In order to run, it requires vcf2bed, gff2bed, bigWigToBedGraph, cut, MethylDackel (0.5.0 or newer, we performed this analysis with version 0.5.2), GNU sort, GNU head, bedtools, BEDOPS, paste, awk, and Python 3.

The pull request for the patch adding mappability support to MethylDackel can be found at <https://github.com/dpryan79/MethylDackel/pull/80>. It was merged into MethylDackel in version 0.5.0.

The Bismap file used for this analysis was downloaded from <https://www.ncbi.nlm.nih.gov/bioproject/52390>.

The ClinVar VCF was the July 22, 2019 version of ClinVar’s variants VCF, with a file name of clinvar_20190722.vcf.gz <should we update this to a newer version?>

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