- 2 software for DNA methylation
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## **Abstract:**

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DNA methylation (DNAm) is the most commonly studied marker in ecological epigenetics, yet the performance of popular library preparation strategies and bioinformatic tools is seldom assessed and compared in genetically variable natural populations. We profiled DNAm using reduced representation bisulfite sequencing (RRBS) and whole genome bisulfite sequencing (WGBS), including technical and biological replicates from lab-reared and wild-caught threespine stickleback (Gasterosteus aculeatus). We then compared how the most commonly used read mapper and methylation caller (Bismark) performed relative to two alternative pipelines (BWA mem or BWA meth read mappers analyzed with MethyDackel). BWA meth provided 50% higher mapping efficiency than BWA mem and 45% higher efficiency than Bismark. Despite differences in mapping efficiency, BWA meth and Bismark produced highly similar methylation profiles, while BWA mem systematically discarded unmethylated cytosines. Sequencing depth filters had large impacts on CpG sites recovered across multiple individuals, with the largest impact on WGBS data. Notably, the prevalence of CpG sites with intermediate methylation levels is greatly reduced in RRBS data compared to WGBS, which may have important consequences for functional interpretations. We conclude by discussing how library construction and bisulfite alignment wrappers can influence SNP filtering, genomic coverage, and the abundance and reliability of data available for downstream analysis. Our analyses suggest that researchers studying genetically variable populations may prioritize filtering SNPs by constructing RRBS libraries with small insert sizes and paired end reads, which is counter to conventional wisdom.

**Key words:** DNA methylation, RRBS, WGBS, Bismark, BWA meth, Ecological epigenetics

## **Introduction:**

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Epigenetics is an umbrella term for chemical and conformational modifications to DNA and proteins, with DNA methylation (DNAm) being among the most well studied epigenetic marks in ecologically focused studies (Banta & Richards, 2018; Hu & Barrett, 2017; Lea et al., 2017; Rey et al., 2020). As molecular ecologists increasingly explore the role of methylation in natural systems, there is a need for increased attention to how methodological choices impact inferences of between and within individual variation. This is particularly important given that most methylation profiling methods were originally developed and applied in inbred laboratory model organisms. There are multiple forms of DNAm, but the most abundant type in vertebrates is the addition of a methyl group to cytosine at CpG dinucleotides, often found in CpG-dense regions of the genome (CpG islands) (Jaenisch & Bird, 2003). DNAm has a broad range of genomic effects, including silencing transposable elements (Marin et al., 2020), blocking transcription factors and other proteins from binding to DNA, increasing the likelihood of spontaneous deamination of methylated cytosines converting them to thymine (Ord et al., 2023), and slowing down transcriptional machinery (Cholewa-Waclaw et al., 2019). While its role varies, increased methylation is often associated with decreased gene expression (Zemach et al., 2010). Multigenerational common garden experiments show that although the genomic loci that undergo DNAm are largely heritable, they can also be modified by environmental stimuli (Bogan et al., 2023; Heckwolf et al., 2020; Hu et al., 2021). Moreover, environmentally-induced shifts in DNAm can be inherited across generations (Takahashi et al., 2023; Tobi et al., 2018). Recent findings also suggest that methylated regions experience different selection pressures than background genomic DNA (Ord et al., 2023). Despite accumulating evidence that DNAm may

play an important role in evolution, this topic is still an active area for debate (Adrian-Kalchhauser et al., 2020; Charlesworth et al., 2017; Jaenisch & Bird, 2003; Otterdijk & Michels, 2016). As interest in the functional and evolutionary roles of DNAm continues to increase, more attention needs to be directed toward developing a robust understanding of which loci are methylated and at what rates.

DNAm is commonly measured using bisulfite sequencing methods. Treating DNA with bisulfite converts unmethylated cytosines to uracil, which then becomes thymine during PCR amplification. When compared to a reference genome, C/T mismatches are identified as unmethylated cytosines, while any cytosines remaining are inferred to be methylated. Although DNAm at any particular base pair is a binary trait (i.e., present/absent at a CpG site on each read), the same location may not be methylated across cells or tissues. Some CpG sites will be consistently methylated (or unmethylated) while others will have intermediate methylation (Hay et al., 2023). When bisulfite sequencing is used for bulk sequencing of many cells within a particular tissue, the three possible methylation states of a CpG site in a single cell (present/absent/heterozygous) are summed and become a quantitative metric (i.e., proportion of total sites that are methylated). The ability to accurately capture this quantitative variation is highly contingent on the number of times that a given base pair is represented in the sequencing data, often referred to as read depth.

The two most commonly used bisulfite sequencing methods are whole genome bisulfite sequencing (WGBS) and reduced representation bisulfite sequencing (RRBS; Gu et al., 2011). While WGBS offers the potential to profile all CpG sites, it also requires large amounts of sequencing data in order to repeatedly sample individual DNAm sites. This requirement often limits WGBS to experiments involving few biological replicates and relatively low read depth,

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which reduces accuracy of methylation calls and statistical power to detect group-level differences (Seiler Vellame et al., 2021). Additionally, because WGBS does not enrich for methylated sites, much of the sequence data is uninformative for epigenetic studies. For example, 70-80% of mapped reads in human WGBS studies do not contain CpG dinucleotides (Elliott et al., 2015). In contrast, RRBS uses methylation-insensitive restriction enzymes to target sequencing toward CpG islands (e.g., the most commonly used enzyme is MspI, with a cut site of CC/GG). CpG islands are regions of the genome with a high density of CpG sites and believed to be functional hotspots for DNAm (Suzuki & Bird, 2008). Thus, RRBS not only enriches for regions most likely to harbor functionally important methylation differences, but by also profiling less than 10% of the genome for any individual it allows researchers to increase sample sizes and read depth of regions covered (Bock et al., 2010; Laine et al., 2022). The large differences in the types of data recovered from WGBS (wide breadth of the genome sequenced, lower read depth, and smaller sample sizes) and RRBS (specifically targets CpG islands, higher read depth, and larger sample sizes) make each approach better for particular types of research questions. Because evolutionary and ecological studies often assess group-level variation, and therefore require relatively large sample sizes, RRBS is becoming the preferred epigenetic profiling method for these disciplines. However, few RRBS studies have directly compared its results with WGBS, making it difficult to directly assess what biases or limitations are inherent to each method. Apart from the differences in library construction and experimental design, analysis

Apart from the differences in library construction and experimental design, analysis methods also differ for DNAm data. For example, bioinformatic tools apply different approaches to account for unmethylated cytosines no longer matching the reference genome after bisulfite treatment. Bismark (Krueger & Andrews, 2011), the most common tool for mapping bisulfite

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converted sequences (Laine et al., 2022) (Table 1), handles mismatches by producing in silico sense  $(C \to T)$  and antisense  $(G \to A)$  conversions at all CpG sites before aligning reads to a reference genome that is itself converted using the same process. Reads are considered ambiguous and discarded if they align with multiple versions of the *in silico* converted reference genome. In addition to mapping reads, Bismark also streamlines data analysis by extracting methylation information in a file format (i.e., bedGraph) that is compatible with downstream statistical tools. However, this software relies on Bowtie2 (Langmead & Salzberg, 2012) for read mapping, which often produces lower mapping efficiency (percent of reads mapped to the reference) than other commonly used mapping algorithms such as BWA mem (Harrath et al., 2019). Although tuning Bowtie2 mapping parameters can improve mapping rates, this process can pose a substantial hurdle for novice users. Additionally, generating four intermediate in silico conversions for both strands of the reference genome and sample reads leads to longer computational run times and greater memory demands than alternative tools (Farrell et al., 2021; Krueger & Andrews, 2011; Nunn et al., 2021). BWA meth (Pedersen et al., 2014) is an alternative bisulfite sequence alignment script to Bowtie2. This software is faster than Bismark because it only performs in silico conversion of the reference genome prior to read mapping, not the sampled reads, and uses the BWA mem algorithm (Nunn et al., 2021). However, a current limitation of BWA meth is that it stops after mapping reads; the tool MethylDackel (<a href="https://github.com/dpryan79/MethylDackel">https://github.com/dpryan79/MethylDackel</a>) is recommended for extracting methylation information. An advantage of MethylDackel is that it uses overlaps between paired end sequencing data to discriminate between SNPs and unmethylated cytosines. If a site is a bisulfite converted cytosine, the opposite strand should have a G, otherwise it is likely a SNP. MethylDackel allows the user to specify the maximum

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proportion of non-Gs permitted on the opposite strand before a site is considered a SNP and excluded from the dataset. Importantly, this could impact estimates of DNAm variation if polymorphic sites are frequently eliminated from analyses. To prevent erroneous filtering of sites based on sequencing errors, the user can also specify the minimum depth the opposite strand must have in order to be used for filtering. As polymorphism data are often unavailable for natural populations, this feature has the potential to increase the reliability of DNAm research on understudied systems.

Here, we provide a comparison of approaches for generating and analyzing DNAm data that are particularly relevant to molecular ecologists. Specifically, we generated WGBS and RRBS data from five populations of threespine stickleback (Gasterosteus aculeatus). Four individuals were profiled using both WGBS and RRBS to assess how the methods diverge in: 1) which and how many DNAm sites are recovered, 2) depth of coverage, and 3) distributions of percentage DNAm per site. We then applied several bioinformatic pipelines to characterize the performance of bisulfite alignment software and how they impact the mean and distribution of percent DNAm for each individual. While studies have assessed performance of bioinformatic pipelines for human and plant methylation data (Farrell et al., 2021; Nunn et al., 2021; Zhou et al., 2024), to our knowledge this has not yet been done in ecologically-focused vertebrate systems. We predicted that RRBS would provide fewer total DNAm sites, greater per-site read depth, and that most DNAm would be found in CpG islands (Bock et al., 2010). We also predicted that BWA mem-based read mappers would perform better than Bowtie-based mappers in terms of total mapping efficiency, and that MethylDackel would efficiently remove polymorphic sites. After explaining our findings, we discuss how to optimize sequencing resources for ecological epigenetics, particularly for systems where DNA polymorphisms are

## **Methods:**

Literature review of bisulfite library preparation and analysis methods

To assess how frequently Bismark, BWA meth, MethylDackel, and BWA mem are used to analyze methylation data, we used "software name" AND "DNA methylation" to query Web of Science and Google Scholar databases on August 21, 2024. We consolidated the number of search results from each database in Table 1.

### Sample Collection

We sampled wild and lab-reared threespine stickleback from five populations (Table 2). RRBS was applied to all populations, while WGBS was used for common garden fish from two Alaskan populations: Watson and Wik Lakes. Fish were euthanized with an overdose of buffered MS-222 according to IACUC protocol L006460-A01. Field samples were dissected lakeside to avoid introducing transportation stress that may influence DNAm profiles. We dissected and stored liver tissue in RNALater (Sigma-Aldrich). We chose liver as our focal tissue because previous work suggests liver-specific methylation is relatively conserved across vertebrates (Klughammer et al., 2023). Liver also has relatively few cell types (Gumucio & Miller, 1981), which should limit between sample variation due to cell type specific methylation. Two lab fish from each of the Roberts, Gosling, and Sayward populations were exposed to the tapeworm

*Schistocephalus solidus* as part of a separate study, and one of the Sayward individuals was successfully infected. None of these individuals displayed obvious differences in DNAm patterns, and therefore were retained for further analysis (Supplemental Figure 1).

#### Sequencing

We extracted DNA from each liver sample using carboxyl coated magnetic beads (BOMB.bio method: Oberacker et al., 2019). Bisulfite conversion, library preparation, and sequencing were performed by Admera Health Biopharm Services. RRBS samples were bisulfite treated and then converted to Illumina sequencing libraries and using Zymo-Seq RRBS Library Kit. WGBS were similarly bisulfite converted using EZ DNA Methylation-Gold Kit (Zymo Research, California, USA) and library preparation was done with xGen<sup>TM</sup> Methylation-Sequencing DNA Library Preparation Kit (Integrated DNA Technologies, California, USA). Sample quality was assessed using Quibit, agarose gel, and qPCR prior to sequencing. Libraries were sequenced on Illumina Novaseq 6000 using paired end 150bp read chemistry. DNA extractions and sequencing were performed in two batches. Each barcoded RRBS sample was mixed into one of two sequencing pools (Table 2), which produced an average (± SEM) of 10.5 (± 1.3) million and 31.5 (± 15.4) million reads per sample following adapter trimming. All WGBS samples were sequenced in one batch, resulting in 58.5 (± 1.0) million reads per sample after adapter trimming.

#### Read Mapping and Methylation Calling

We trimmed adapters and barcodes using the -paired, -rrbs (for RRBS samples only, to remove cytosines filled in at the end of reads as an artifact of library preparation), and -q 20 options in Trim Galore! v. 0.6.10 (https://github.com/FelixKrueger/TrimGalore). We then

aligned reads to the threespine stickleback reference genome (v.5; Nath et al., 2021) using default parameters in three programs: BWA mem v. 0.7.17, BWA meth v. 0.2.7, and Bismark v. 0.24.2. Bismark resulted in a low mapping efficiency 28.10 (± 2.26%). We followed the guidance in the documentation and aligned the data in single-end mode with the flag nondirectional, resulting in a mapping efficiency analogous to other studies. We therefore use the data mapped in single end mode for the rest of the analysis. For BWA mem and BWA meth alignments, samtools v. 1.9 (Danecek et al., 2021) was used to convert SAM outputs to BAM format. Mapping efficiency is included in the Bismark output, and we used Bamtools v.2.5.2 (Barnett et al., 2011) to calculate it for BWA meth and BWA mem runs. To extract methylation from BWA meth and BWA mem results we used MethylDackel v. 0.5.1 (https://github.com/dpryan79/MethylDackel). Specifically, we filtered SNPs, tabulated per base methylation metrics, and created an output file compatible with methylKit in R (-methylKit) (Akalin et al., 2012). As methylation call accuracy scales logistically with read depth and becomes asymptotic at 10x (Seiler Vellame et al., 2021), we used --minDepth 10 to only retain reads that were sequenced at least 10 times. It can be difficult to differentiate between PCR replicates and true biological duplicates in RRBS data due to biased genomic fragmentation (Laine et al., 2022). Therefore, for the RRBS samples only, we followed recommendations in the MethylDackel documentation and applied the --keepDupes option. Three different SNP filtering cut offs were tested (--maxVaraintFrac 0.25, 0.5, and 0.8) with --minOppositeDepth 5. To prioritize capturing potential methylation variation, we used the most lenient SNP cut off (-maxVariantFrac 0.8) for further analyses.

Statistical analyses

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different from 1 while retaining a high  $R^2$ .

We applied a number of statistical tests in R v. 4.3.2 (R Core Team, 2023) to compare read mapping approaches with both RRBS and WGBS data. We used the nonparametric Kruskall-Wallis rank sum test to assess if mapping efficiency differed between Bismark, BWA meth, and BWA mem. Post hoc pairwise comparisons were done using the Dunn test and pvalues were corrected for multiple comparisons using the Holm method in FSA v. 0.9.5 (Ogle et al., 2023). Figures were generated using ggpubr v. 0.6.0 (Kassambara, 2023). For RRBS samples, which were sequenced in two batches, we used Wilcox rank sum exact tests to test whether batch affected mapping efficiency for each of the three read mapping tools. Results were visualized using ggplot2 v. 3.5.1 (Wickham, 2016) and cowplot v. 1.1.3 (Wilke, 2024) (Supplemental Figure 2). The same procedure was used to analyze the number of mapped reads (Supplemental Figure 3) and average read depth for each read mapping approach. To determine if the resulting methylation profiles of individuals were in agreement despite differences in read mapping and SNP calling, we compared both the mean and distribution of percent methylation per individual obtained after using Bismark, BWA meth + MethylDackel, and BWA mem + MethylDackel. We used methylKit v. 1.27.1 (Akalin et al., 2012) to calculate the mean and visualize the distribution of percent methylation for each sample. We then created a linear model in ggpmisc v. 0.6.1 (Aphalo, 2024) to compare the mean percent methylation obtained using each pipeline. If, for every individual, the mean percent methylation obtained from one pipeline reliably matches the mean percent methylation calculated using another pipeline, then the slope and R<sup>2</sup> should be 1. If one method consistently over or underestimates methylation compared to another, then the slope may be significantly

We next assessed differences between RRBS and WGBS. To determine if average read depth per sample varied we used a Wilcox rank sum test. We then used ggvenn v.0.1.10 to tabulate the proportion of CpG sites that were shared (1) between methods within an individual and (2) between individuals using the same method. As we were interested in the breadth of genomic coverage between library preparation methods, we did not filter SNPs prior to this comparison and relaxed the minimum depth to test both 5x and 10x. Finally, the frequency distribution of percent methylation per base was also used to visualize how the overall methylation profile for an individual varied based on the sequencing method.

### **Results:**

Read Mapping: BWA meth outperformed BWA mem and Bismark

BWA meth, BWA mem, and Bismark differed significantly in their mapping efficiency using RRBS data (Kruskal-Wallis chi-squared = 75.95, df = 2, p < 2.2 x  $10^{-16}$ ). With a mean ( $\pm$  SD) mapping efficiency of 99.23% ( $\pm$  0.18), BWA meth outperformed both Bismark (54.38  $\pm$  4.35%) and BWA mem (49.26  $\pm$  4.46%) (Bismark - BWA meth: Z = -5.64, p = 3.41 x  $10^{-8}$ ; BWA mem - BWA meth: Z = -8.57, p = 3.00 x  $10^{-17}$ ). The mapping rate for BWA mem was also significantly lower than Bismark (Bismark - BWA mem: Z = 2.93, p = 3.34 x  $10^{-3}$ ) (Figure 1). Mapping efficiency for Bismark did not differ between sequencing batches (Wilcox rank sum exact test, W = 142, p-value = 0.732), but there were small but significant batch effects for both BWA meth (Wilcox rank sum exact test, W = 261, p = 2.55 x  $10^{-8}$ ) and BWA mem (Wilcox rank sum test, W = 253, p-value = 7.112 x  $10^{-7}$ ) (Supplemental Figure 2). Despite the batch effects, BWA meth produced the highest mapping efficiency (>99%,  $17.8 \pm 13.5$  million mapped reads per RRBS sample), so its output was used for downstream analysis. For the WGBS samples,

BWA meth again performed the best with a mapping efficiency of 99.58 ( $\pm 0.16\%$ ), resulting in a mean of 58.6 (± 1.1) million mapped reads. Although we had too few WGBS libraries to make strong statistical inferences, the average WGBS mapping efficiency of BWA mem and Bismark were again lower than BWA meth: 74.64% ( $\pm 3.43$ ) and 38.7% ( $\pm 1.24$ ), respectively.

BWA mem artificially inflates methylation

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Despite significant differences in mapping efficiency and lack of SNP filtering with Bismark, the mean methylation per putatively methylated site was highly similar between BWA meth and Bismark ( $R^2 = 0.91$ , slope = 0.935; Figure 2a). BWA mem produced inflated methylation rates compared to the other two methods (BWA meth - BWA mem:  $R^2 = 0.67$ , Figure 2b; Bismark - BWA mem:  $R^2 = 0.64$ , Figure 2c). This bias was so extreme that the median percent methylation for every sample mapped with BWA mem was 100%. We also examined distribution of percent methylation per base to further characterize how read mapping wrapper influenced which CpG sites were retained for analysis (Figures 3-5, Supplemental Figures 4-8). BWA mem hardly recovered any CpG sites that were consistently unmethylated (<10% of methylation) while these sites made up 51.6% ( $\pm$  14.8) and 43.7% ( $\pm$  13.9) of the data in BWA meth and Bismark, respectively (Figure 4).

Differences in genomic breadth and depth of coverage between RRBS and WGBS

After applying BWA meth and MethylDackel the average median insert size of RRBS reads was 38.06 ( $\pm$  2.44) compared to 242.25 ( $\pm$  22.47) for WGBS. Despite the small insert size, RRBS resulted in a higher per site depth (mean of  $16.39 \pm 3.46$ ) than WGBS ( $12.73 \pm 0.06$ ). Batch 1 RRBS samples had a significantly lower read depth than batch 2 (14.73  $\pm$  1.07 versus  $19.44 \pm 4.22$ ; Wilcoxon rank sum exact test, W = 24.5, p-value = 0.0001151), but also a slightly higher mapping efficiency (99.34  $\pm$  0.00086% versus 99.03  $\pm$  0.0013%; Wilcoxon rank sum

exact test, W = 261, p-value = 2.553e-08, Supplemental figures). This indicates that the difference in depth of coverage between batches was driven by greater sequencing of batch 2, not by better performance of read mapping software.

We next compared the effects of applying different library preparation methods to technical replicates (i.e. RRBS vs WGBS; Table 3), as well as the similarity of biological replicates from a single population (i.e., RRBS versus RRBS, WGBS versus WGBS; Table 4). Applying a minimum depth filter of 10 reads per site on RRBS and WGBS libraries from the same individual resulted in  $\leq 1.0\%$  shared CpG sites. Lowering the minimum depth to 5 reads caused the proportion of shared CpG sites to increase to 2.25% ( $\pm$  0.42). Notably, at a minimum of 10x depth, >96% of CpG sites were exclusive to WGBS, highlighting the large difference in breadth of coverage between RRBS and WGBS (Table 3).

When assessing CpG sites shared among individuals using RRBS, the percent of overlap depended on fish population. With a minimum of 10x depth, 13.1% and 27.2% of CpG sites were shared between biological replicates from Watson and Wik Lakes, respectively. This population effect was much reduced with WGBS; 18.8% of CpG sites were shared between Watson Lake individuals and 19.7% between Wik individuals. Lowering the minimum depth to 5x increased the proportion of shared CpG sites using RRBS (28.4% for Watson, 40.1% for Wik) and WGBS (57.6% for Watson, 59.0% for Wik) (Table 4).

Apart from comparing overlap in particular sites, the vast majority of the quantitative CpG sites (i.e. the sites that are not always methylated or unmethylated), are not represented in

the RRBS data (Figures 3 - 5). At a minimum depth of 10x, the proportion of CpG sites with methylation rates >10% and < 90% was twice as high in the WGBS samples ( $55.63 \pm 2.49\%$ ) than the RRBS samples ( $28.56 \pm 8.37\%$ ). A higher proportion of intermediately methylated sites were recovered in batch 2 of RRBS sequencing (Supplemental Figures), but none of these samples were also analyzed via WGBS.

## **Discussion:**

The increasing popularity of RRBS for methylation profiling has coincided with a rise in the use of Bismark as the default method for read mapping and methylation calling, particularly in ecologically focused experiments (Table 1). However, little effort has been dedicated to understanding the strengths and limitations of these methodological choices, particularly in genetically variable, wild organisms. By comparing RRBS and Bismark against less commonly used library preparation and analysis alternatives, we identify several options for researchers to maximize their sequencing resources.

Optimizing bisulfite alignment wrappers for ecological epigenetics

As predicted by previous studies (Farrell et al., 2021; Nunn et al., 2021; Zhou et al., 2024), BWA meth outperformed both BWA mem and Bismark in its ability to map the majority of sequenced reads to a reference genome (Figure 1). However, it is noteworthy that this previous work focused on human and plant data. We found no previous studies assessing bisulfite alignment wrappers in wild-caught/non-model systems. Despite differences in mapping efficiency, BWA meth and Bismark provided consistent estimates of the mean percent of CpG sites methylated per individual (Figure 2). In contrast, BWA mem substantially inflated estimates of mean methylation rate (Figures 2 & 4). This difference reflects the optimization of particular bioinformatic tools for bisulfite converted data. Because BWA mem does not account for C/T

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mismatches induced during bisulfite conversion, reads enriched with converted thymines are more likely to differ considerably from the reference sequence and be lost during mapping. Importantly, BWA meth and Bismark only partially address this bias through bioinformatic conversions that improve mapping. Bisulfite conversion damages DNA at unmethylated cytosines during library preparation, which inherently inflates genome-wide methylation levels (Olova et al., 2018). The magnitude of this bias is only recently becoming clear thanks to newer library preparation approaches that forego harsh chemical reactions and instead rely on enzymes to convert unmethylated cytosines (e.g., EM-Seq: Vaisvila et al., 2021). If precise estimates of genomewide methylation are of key concern, then consideration should be given to using these newer, non-bisulfite methods of library preparation. Importantly, EM-seq converts unmethylated cytosines to thymine, making the sequencing data compatible with existing bioinformatic tools for bisulfite converted data. Despite these advances, RRBS remains the most cost effective method and is likely sufficient if the primary goal of a project is to measure or detect large changes in methylation (i.e., those likely to have functional effects or are environmentallyinduced). Regardless of the library preparation method chosen, BWA meth or Bismark will likely provide similar results.

For projects focusing on metrics related to per-site methylation, a key goal is to maximize the number of usable sequencing reads. Bismark produced a mapping efficiency only slightly higher than BWA mem, yet it remains the most commonly used read mapping method (Table 1). Continued reliance on Bismark may partly be attributed to it being one of the first bioinformatic tools designed to handle bisulfite converted data (Table 5). Because Bismark does *in silico* conversions of all reads (sense and antisense, read and reference), its reliability is also agnostic to library directionality. Although Bismark's ability to both map reads and call methylation

levels makes it particularly accessible for researchers new to epigenetics, the tool's strict default parameters may cause difficulties when working with paired end reads, as we observed when we initially had a 28% mapping efficiency. Although rarely discussed in the literature, low mapping rates of paired end reads appears to be a relatively common issue with this tool (e.g.,: Nunn et al., 2021; Zhou et al., 2024). Indeed, the developers provide thorough documentation to help users resolve this mapping problem. The guidance highlights failure to trim adapters prior to read mapping, potential contaminants, and high stringency under default parameters, among other possible sources of error. Under default parameters, both paired end reads must map to one unique version of the *in silico* converted reference genome, otherwise the paired end sequence gets discarded. Following the advice in the documentation, and in-line with Nunn et al. (2021) and Zhou et al. (2024), we relaxed this threshold by mapping our reads as non-directional. This resulted in a mapping rate (54%) that is more in-line with previous DNAm studies in stickleback (Heckwolf et al., 2020; Hu et al., 2021).

Despite requiring additional tools for methylation calling, BWA meth resulted in significantly higher mapping rates than Bismark, leading to more data available for downstream analysis. While a previous study found that BWA meth had higher error rates than alternative tools (Farrell et al., 2021), this analysis was based on both directional and nondirectional libraries. It is plausible that the increased error rate was driven by strand bias rather than being a software limitation. Following BWA meth, we used MethylDackel to tabulate methylation and filter SNPs. Conventional wisdom is that RRBS libraries should be single end, as paired end reads may lead to bias by repeatedly sequencing the same sites (Krueger et al., 2012). However, this bias can be reduced by normalizing the data based on median read depth, and if genetic polymorphisms are unknown, the overlapping nature of paired end reads can be exploited by

software like MethylDackel to call SNPs. Additionally, using MethylDackel for SNP filtering reduces errors associated with SNPs falsely being considered unmethylated cytosines. Similar to most studies of ecological model organisms, we did not have genotype data for either the focal individuals or populations, and were thus unable to verify the accuracy of SNP filtering using MethylDackel.

Recently, there has been an influx of bisulfite alignment wrappers that combine the strengths of BWA meth and Bismark. These software packages use BWA mem for read mapping, are compatible with both directional and nondirectional libraries, and call methylation (Table 5). Although our goal was to assess the most commonly used methods, future efforts should aim to similarly compare the strengths and limitations of new software for ecologically-focused experiments.

RRBS preferentially sequences stable methylation in comparison with WGBS

In addition to assessing how bisulfite alignment wrappers performed with RRBS data, we examined how methylation profiles varied across technical and biological replicates using RRBS and WGBS. Despite receiving relatively low amounts of RRBS data, which drove low read depth, RRBS still reliably resulted in 29% higher sequencing depth than WGBS. However, few CpG sites were commonly sequenced between technical and biological replicates using the two sequencing methods (Tables 3 & 4). In line with previous research (Vaisvila et al., 2021), the number of overlapping CpG sites between replicates greatly increased when the minimum depth threshold was lower. When comparing the methylation patterns recovered from technical replicates, the most striking difference between RRBS and WGBS was the proportion of intermediately methylated sites recovered with both methods (Figures 3-5). Intermediate

methylation comprised 56% of the CpG sites sequenced with WGBS, while it was only 29% of the CpG sites recovered with RRBS.

One potential explanation for the lack of congruence between sites sequenced with RRBS and WGBS is that our sequencing parameters (i.e., 2x150bp reads) were too long for the small inserts (~38bp) produced via RRBS. Therefore, using a longer read length resulted in extraneous sequencing of adapters that were trimmed prior to mapping. However, the greater depth of batch 2 (19x compared to 14x for batch 1) suggests that investing in more sequencing can offset the effects of adapter sequencing. Thus, we suggest selecting shorter read lengths that are still within the constraints of mapping software (70bp) (Harrath et al., 2019). Accuracy of methylation calls requires a minimum of 10x depth (Seiler Vellame et al., 2021). However, due to uneven distribution of sequencing effort across the genome, as exhibited by the low proportion of overlapping loci between our biological and technical replicates, we recommend using a minimum depth threshold of 20x genomic coverage when selecting sequencing parameters.

Perhaps the most striking outcome from using technical replicates to contrast sequencing methods was that the proportion of intermediately methylated sites sequenced with WGBS was almost twice as high as intermediate methylation captured with RRBS (56% versus 29%, respectively). Methylation can degrade if samples are collected multiple hours after mortality (Rhein et al., 2015) or frozen for a long period of time (Lee et al., 2023), but the liver tissue we sequenced was stored in RNALater immediately upon euthanasia and kept at -70°C for less than one year. Therefore, degradation of intermediately methylated sites is unlikely the main driver of this observation. Rather, there are technical and evolutionary explanations that likely interacted to create this pattern. As previously mentioned, batch 2 RRBS samples recovered a much higher proportion of intermediately methylated sites than batch 1 (Supplemental Figures), likely driven

by more sequencing. However, this higher rate  $(37.83 \pm 5.78\%)$  still appears to be substantially lower than the average of WGBS. This pattern suggests that there may be fundamental, biological differences in the methylation patterns surveyed by each method. Future studies could use technical replicates to confirm this reduction of intermediately methylated sites in RRBS data.

In considering how much sequencing to invest in RRBS, there may be evolutionary arguments not to prioritize capturing intermediately methylated sites. A recent study in mouse cell lines found that loci with intermediate methylation probabilistically, but not reliably, transmitted methyl marks during mitosis. Furthermore, their methylation status had little effect on gene expression (Hay et al., 2023). Another study in wild stickleback found that regions of the genome containing stable methylation (<15% and >85% methylation) had stronger signatures of selection than those containing intermediate methylation (Ord et al., 2023). These studies suggest that because CpG islands are often enriched for methylation sites that have functional importance (Jaenisch & Bird, 2003), they are more likely to be subject to selection, and thus may have increased investment by regulatory machinery to maintain methylation state across all cells. Thus, the lower proportion of intermediately methylated cytosines we sequenced with RRBS, which targets CpG islands, may be picking up a biologically interesting pattern. Additionally, given that molecular ecologists are likely interested in identifying functionally important epigenetic effects, RRBS may provide a better option for detecting and measuring these changes. As the field of ecological epigenetics continues to expand, we encourage researchers to continue to interrogate which CpG sites are informative for evolutionary processes and consider how the sequencing and bioinformatic methods they use may influence resolution of methylation profiles.

## **Conclusions:**

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We take a critical approach to analyzing RRBS and WGBS data using several conventional methods. We highlight the strengths and weaknesses of both molecular and bioinformatic methods, all of which should be considered during experimental design. If SNP filtering in the absence of genotype data is a priority, researchers may consider choosing paired end sequencing, contrary to traditional advice regarding RRBS library construction.

Additionally, the most commonly used tool for analyzing bisulfite sequencing data (Bismark) performs significantly worse than alternative methods, such as BWA meth, in terms of percentage of reads mapped. Researchers can increase the availability and reliability of data for downstream analysis by considering alternative analysis pipelines (Table 5). Finally, we highlight how different library preparation methods recover distinct signatures of per-site methylation, and why these patterns warrant additional investigation for understanding how epigenetics influence evolutionary processes.

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# **Author Contributions:**

E.V.K. and J.N.W. conceived and designed the study and co-wrote the manuscript; E.V.K. wrote the software wrappers, conducted analyses, and produced figures; J.N.W. supervised and funded the work.

### **Tables and Figures:**

Table 1: Search results for different bioinformatic tools associated with bisulfite converted DNA.

The search terms used were "software name" AND "DNA methylation," conducted on August
21, 2024.

Tool	Google Scholar Search Results	Web of Science Search Results	
Bismark	6,030	30	
BWA meth	313	7	
BWA mem	3,570	0	
MethylDackel	347	1	

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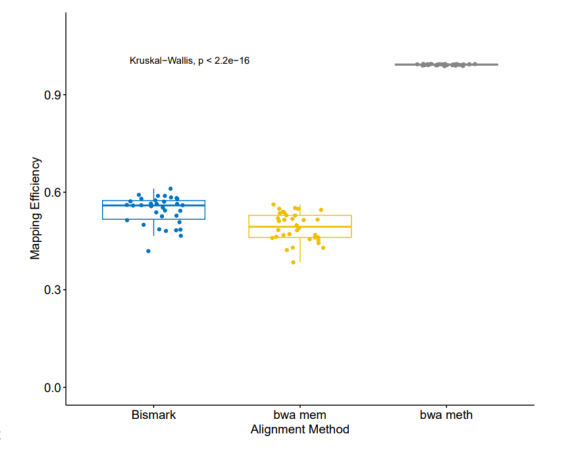
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Table 2: Population, rearing environment, and sample sizes with associated batch of sequencing for RRBS and WGBS. Abbreviations for populations and rearing environments are consistent with figures in the results.

Population	Rearing Environment	Sample Size
		(Sequencing Batch)
Watson (WT, 60.54, -150.47)	Field (FD)	n=4 (1)
	Common Garden (CG)	n=4 (1)
		WGBS (n=2) (1)
Wik (WK, 60.72, -151.25)	Field	n=4 (1)
	Common Garden	n=4 (1)
		WGBS (n=2) (1)
Roberts (ROB, 50.22, -125.54)	Field	n=3 (2)
	Common Garden	n=2 (1), n=2 (2)
Gosling (GOS, 50.05, -125.50)	Field	n=3 (2)
	Common Garden	n=2 (1), n = 2 (2)
Sayward (SAY, 50.39, -125.95)	Common Garden	n=2 (1), n = 2 (2)



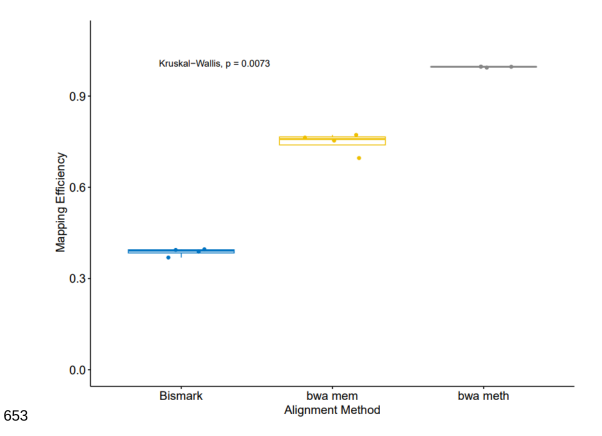
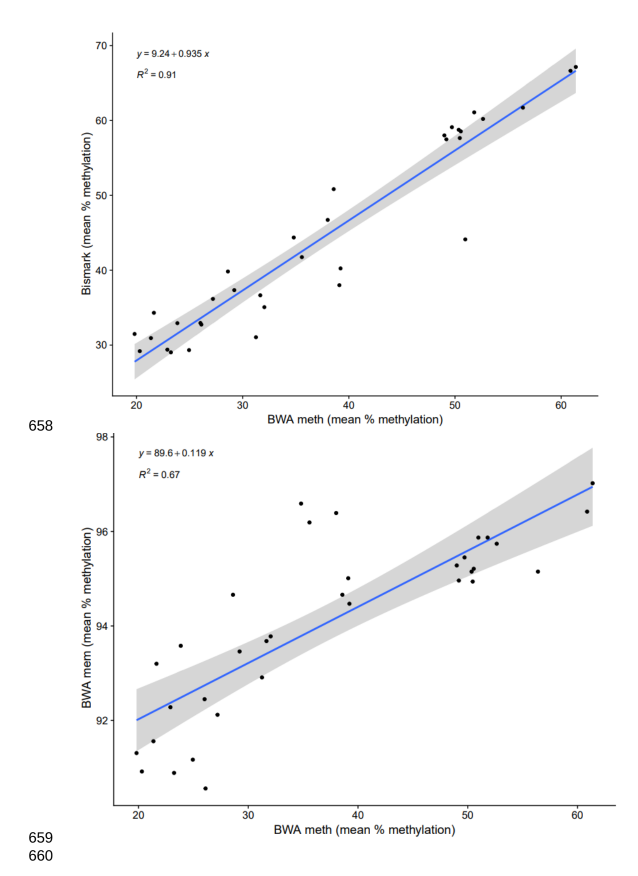


Figure 1: Mapping efficiency of Bismark, BWA mem, and BWA meth on all (A) RRBS and (B) WGBS samples.



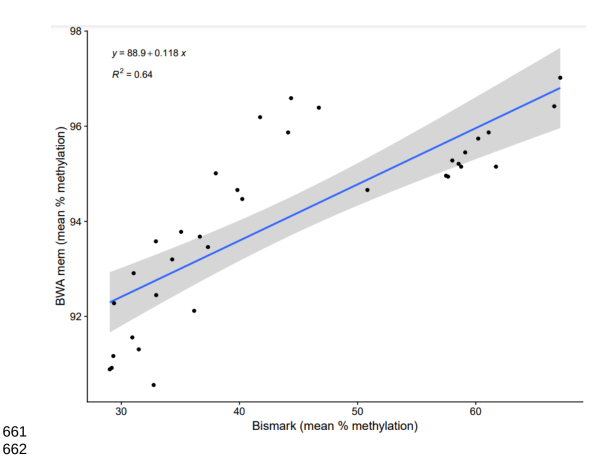
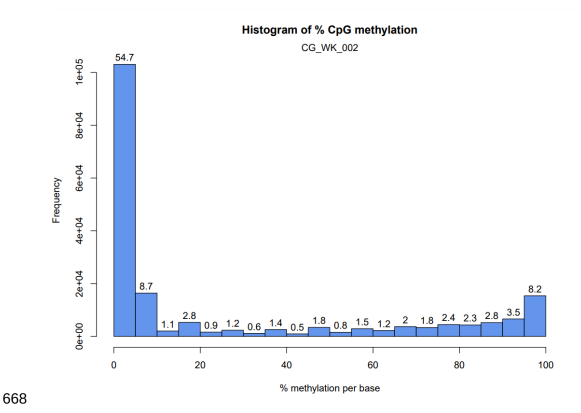
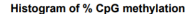
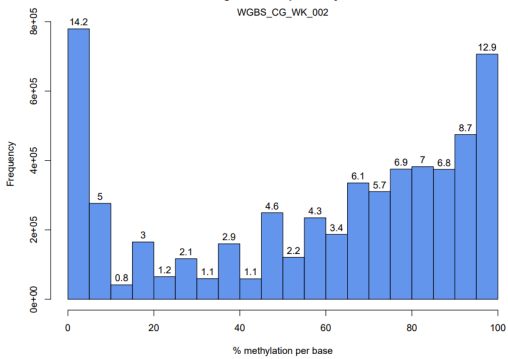
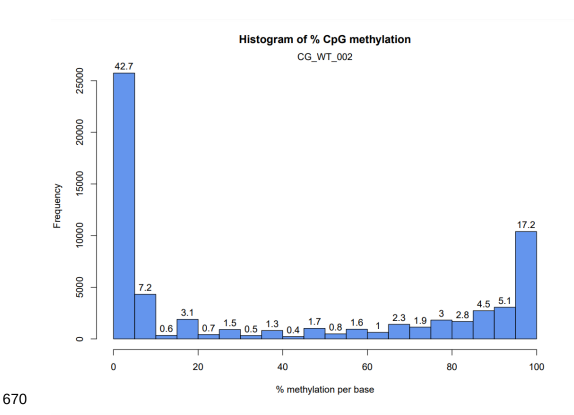


Figure 2: Mean percent methylation per base for each individual sequenced with RRBS using (A) BWA meth and Bismark, (B) BWA meth and BWA mem, and (C) Bismark and BWA mem. A slope and  $R^2$  that is close to one indicates agreement in mean percent methylation between methods.









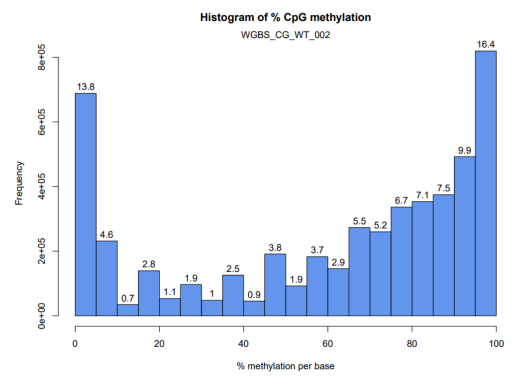
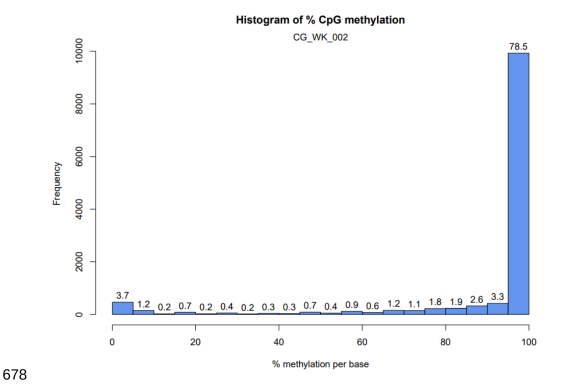
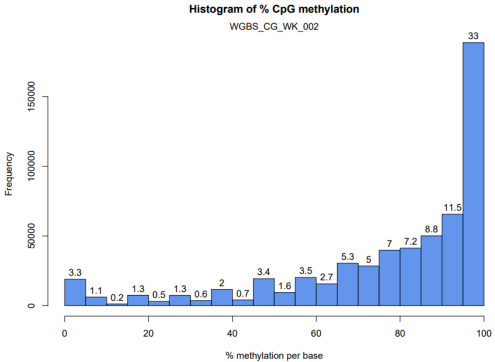
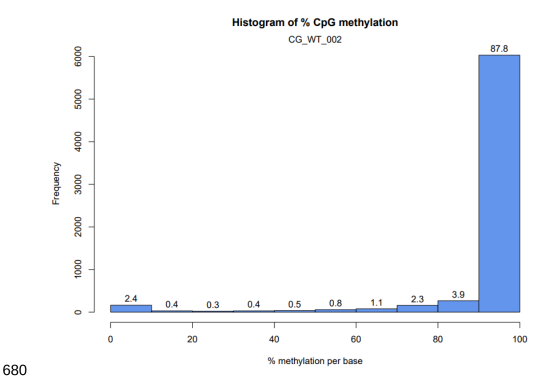


Figure 3: Frequency histograms of percent methylation per CpG site for technical replicates from Wik Lake (A: RRBS, B: WGBS) and Watson Lake (C: RRBS, D: WGBS). The numbers on

top of each bin are the percentage of CpG sites that fall within that bin. Obtained from BWA meth aligned reads with SNP filtering parameters in MethylDackel set to MaxVariantFrac 0.8.







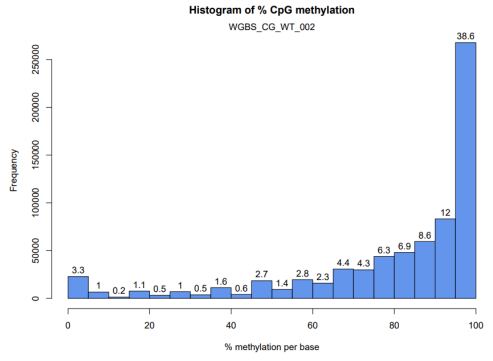
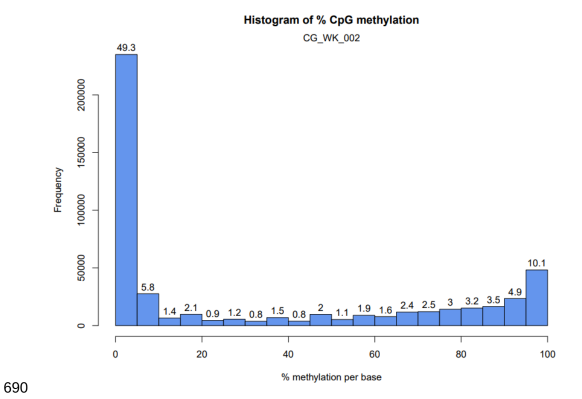
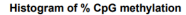
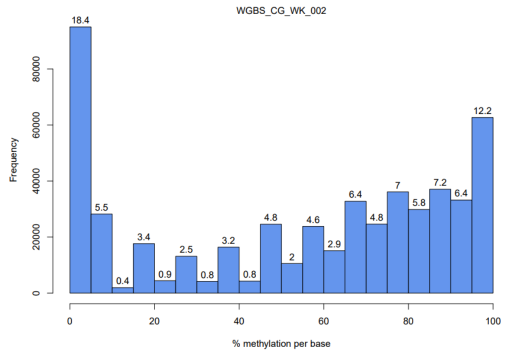
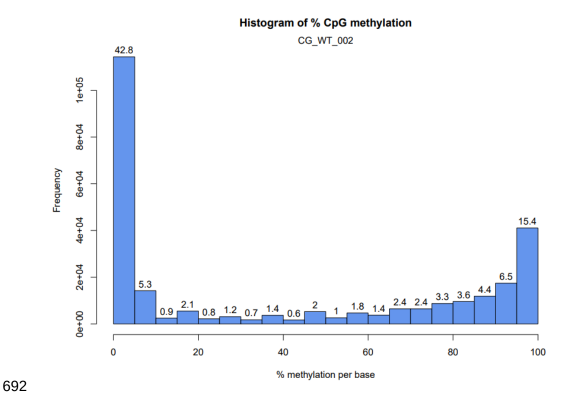


Figure 4: Frequency histograms of percent methylation per CpG site for technical replicates from Wik Lake (**A: RRBS, B: WGBS**) and Watson Lake (**C: RRBS, D: WGBS**).. The numbers on top of each bin are the percentage of CpG sites that fall within that bin. Obtained from BWA mem aligned reads and SNP filtering parameters in MethylDackel set to MaxVariantFrac .8.









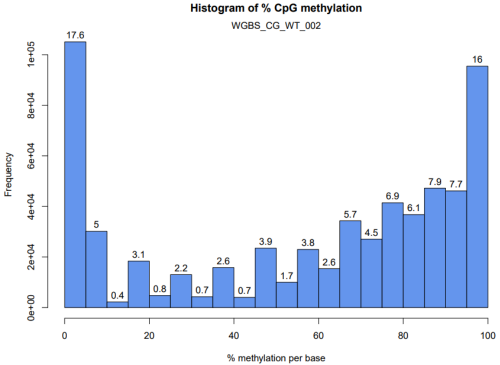


Figure 5: Histograms of percent methylation per CpG site for technical replicates from Wik Lake (A: RRBS, B: WGBS) and Watson Lake (C: RRBS, D: WGBS). The numbers on top of each

bin are the percentage of CpG sites that fall within that bin. Obtained from Bismark aligned reads and no SNP filtering.

Table 3: CpG sites that were in common or unique to WGBS and RRBS for technical replicates with a minimum of 10x depth and 5x depth.

Sample ID	Minimum	Unique to WGBS	Unique to RRBS	Overlapping Sites	
	Depth	(Number of CpG	(Number of CpG	(Number of CpG	
		sites)	sites)	sites)	
CG_WK_002	10x	96.6% (5,417,478)	2.4% (134,797)	1.0% (55,001)	
	5x	96.0% (18,501,995)	1.1% (219,188)	2.8% (546,037)	
CG_WK_003	10x	97.1% (5,160,689)	2.2% (116,016)	0.7% (36,879)	
	5x	96.9% (18,194,034)	1.0% (188,716)	2.1% (402,195)	
CG_WT_002	10x	98.8% (4,966,649)	0.9% (46,124)	0.3% (14,544)	
	5x	97.1% (17,915,424)	1.1% (194,016)	1.8% (337,685)	
CG_WT_004	10x	97.1% (4,410,125)	2.3% (102,397)	0.7% (31,069)	
	5x	96.5% (17,312,003)	1.2% (209,426)	2.3% (418,643)	

Table 4: CpG sites that were in common between two individuals within the using RRBS and WGBS with a minimum of 10x and 5x depth.

Sample Comparison	Sequencing Method	Minimum Depth	Unique to First Sample	Unique to Second	Overlapping Sites
-			(Number of CpG sites)	Sample (Number of CpG sites)	(Number of CpG sites)
CG_WT_002 & CG_WT_004	RRBS	10x	22.2% (38,133)	64.6% (110,931)	13.1% (22,535)
		5x	30.4% (274,939)	41.1% ( <i>371,307</i> )	28.4% (256,762)
	WGBS	10x	44.0% (3,493,419)	37.2% (2,953,420)	18.8% (1,487,774)
		5x	22.3% (5,099,302)	20.0% (4,576,839)	57.6% (13,153,807)
CG_WK_002 & CG_WK_003	RRBS	10x	43.2% (116,469)	29.5% (79,556)	27.2% (73,329)
		5x	39.0% ( <i>377</i> , <i>396</i> )	21.0% (203,082)	40.1% (387,829)
	WGBS	10x	41.7% (3,715,901)	38.6% (3,440,990)	19.7% (1,756,578)

	5x	21.5%	19.6%	59.0%
		(5,085,066)	(4,633,263)	(13,962,966)

Table 5: A summary of the limitations and strengths of alignment wrappers available for bisulfite converted DNA. Qualitative descriptions of performance (Low, Moderate, High) are relative to the performance of other mapping tools for bisulfite converted libraries and collated from the results of multiple studies.

Tool	Algorithm	Mapping Efficiency	Library Type	SNP calling with bisulfite sequencing data	Sources
Bismark (Krueger & Andrews, 2011)	Bowtie2, HiSAT2	Low	Directional & Nondirectional	No	Present study, (Farrell et al., 2021; Guo et al., 2013; Nunn et al., 2021; Zhou et al., 2024)
BWA meth (Pedersen et al., 2014) + MethylDackel (https://github. com/dpryan79/ MethylDackel)	BWA mem	High	Directional	Yes (can also use BISCUIT in place of MethylDackel)	Present study, (Farrell et al., 2021; Nunn et al., 2021; Zhou et al., 2024)
BISCUIT (Zhou et al., 2024)	BWA mem	High	Directional & Nondirectional	Yes	(Farrell et al., 2021; Zhou et al., 2024)
<b>BiSulfite Bolt</b> (Farrell et al., 2021)	BWA mem	High	Directional & Nondirectional	Yes	(Farrell et al., 2021; Zhou et al., 2024)