# Comparison of whole-genome bisulfite sequencing library preparation strategies identifies sources of biases affecting DNA methylation data

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#### **Abstract**

**Background:** Whole-genome bisulfite sequencing (WGBS) is becoming an increasingly accessible technique, used widely for both fundamental and disease-oriented research. Library preparation methods benefit from a variety of available kits, polymerases and bisulfite conversion protocols. Although some steps in the procedure, such as PCR amplification, are known to introduce biases, a systematic evaluation of biases in WGBS strategies is missing.

**Results:** We performed a comparative analysis of several commonly used pre- and post-bisulfite WGBS library preparation protocols for their performance and quality of sequencing outputs. Our results show that bisulfite conversion *per se* is the initial and main trigger of pronounced sequencing biases, and further amplification steps lead to over-representation of these artefacts. The majority of standard library preparation methods lead to a significantly biased genomic sequence representation and a marked overestimation of both absolute and relative methylation levels, with clear implications for DNA methylation studies.

**Conclusions:** We propose that amplification-free and post-bisulfite procedures should become the gold standard for WGBS library preparation. To aid with the quality assessment of existing WGBS datasets, we have integrated a bias diagnostic tool in the Bismark package and offer several approaches for consideration during the preparation and analysis of WGBS datasets.

**Keywords:** WGBS, biases, artefacts, bisulfite conversion, degradation, DNA methylation, NGS, polymerase, GC skew

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## **Background**

Methylation of DNA at the 5<sup>th</sup> position in cytosine (5mC) is a stable epigenetic modification found in many living organisms, from bacteria to higher eukaryotes. It is known to play a role in the regulation of transcriptional activity during embryonic development, in processes such as genomic imprinting, transposon silencing, X-chromosome inactivation and during the differentiation of pluripotent cells.

Since its first use in 1992 [1], bisulfite (BS) sequencing of DNA has become the gold standard for analysis of DNA methylation. BS treatment of DNA leads to the conversion of unmodified cytosines to uracil whilst maintaining 5mC unchanged, which, after PCR and sequencing, can be mapped at single base resolution [2,3]. More recently, BS treatment has been coupled with next generation sequencing (NGS) to yield reduced representation (RRBS) or whole genome (WGBS) data on the global genomic distribution of 5mC [4]. As NGS costs decrease, the WGBS approach becomes increasingly accessible for both fundamental and clinical research. However, the ever-increasing diversity of WGBS library preparation kits, protocols and their variations demands a thorough examination of their outputs and performance, to inform the choice of users from both specialist and non-specialist fields, academia and industry. At present, there is a wealth of publically available WGBS datasets, generated in multiple different ways, and it is commonly assumed that they are equally comparable. We set out to investigate how the different steps of current library preparation protocols affect the final sequence output and, ultimately, the quantitation and interpretation of methylation data.

Biases and artefacts from BS sequencing have been well studied outside the NGS context. These encompass biases associated with cloning and PCR, such as primer selectivity and design, polymerase sequence preferences and errors, and template switch (strand recombination) [3,5,6]. In addition, sources of false positive and false negative signals have been well characterised, i.e. the incomplete cytosine conversion by sodium bisulfite and over-conversion of 5mC, found to be affected by factors like DNA quality, quantity and purification procedures, BS incubation length and temperatures, strand reannealing, polymerase, sequencing errors as well as conversion resistant sequences [2,3,5–8]. Different solutions to these biases and artefacts have been proposed, which improved quantitation of DNA methylation at specific loci by PCR and cloning-based methods [2,3,5,7,9–15]. Only some of these considerations, however, remain relevant for NGS-based approaches (e.g. the improvements of BS conversion conditions) and a systematic investigation of major sources of biases in WGBS protocols has not yet been performed. PCR amplification bias has received significant attention in classical (non-BS) whole-genome sequencing [16–22], however, it has been less studied in BS-based whole-genome sequencing [23] and additional sources of bias, which affect both sequencing coverage and methylation quantitation, have not been investigated.

Here we compare several WGBS library preparation protocols by analysing how their sequence coverage and methylation outputs are affected by: 1) BS-induced DNA degradation, 2) PCR amplification, 3) DNA modifications, and 4) incomplete BS conversion. We find that the BS conversion step is the main trigger of biases, due to a selective and context-specific DNA degradation [24] and incomplete conversion efficiency, while subsequent PCR cycles primarily build on the effect of an already biased sequence composition. We discuss mechanisms to avoid, predict or quantitate biases and artefacts in future or for already available WGBS datasets.

# Results

# Study setup: WGBS library preparation steps and strategies

It is well documented that BS conversion causes DNA fragmentation (also known as degradation) of up to 90% of the DNA input [2,7,8,14,24]. In order to assess biases arising from BS-induced DNA degradation, we first tested five bisulfite conversion protocols directly on synthetic and genomic DNA without sequencing. Building on previous work [2,5,7,14], we chose kits from different manufacturers that vary in two key aspects: 1) DNA denaturation, which can be heat- or alkaline-based, 2) BS treatment temperature, which can be high (65-70°C) or low (50-55°C) and typically associated with

different incubation times (Table 1). Additionally, we also tested a protocol ('Am-BS') that uses high concentration (9 M) of ammonium bisulfite (in contrast with 3-4 M sodium bisulfite used in other protocols) [25].

Table 1. BS conversion protocols and parameters.

Method	Denaturation T°C	Conversion T°C	Incubation Time
'Heat 1'	High heat (99°C)	65°C	90 minutes <sup>a</sup>
'Heat 2'	High heat (99°C)	55°C	10 hours <sup>b</sup>
'Alkaline 1'	Low heat (37°C)	65°C	90 minutes <sup>a</sup>
'Alkaline 2'	Low heat (37°C)	50°C	12-16 hours <sup>c</sup>
'Am-BS'	Low heat (37°C)	70°C	30 minutes <sup>d</sup>

<sup>&</sup>lt;sup>a</sup> Imprint DNA Modification Kit (Sigma-Aldrich), 1-step protocol for 'Heat 1' and 2-step protocol for 'Alkaline 1'

We then coupled the above BS conversion protocols to two strategies for the generation of WGBS libraries: 1) pre-BS, which adds sequencing adaptors by ligation before BS conversion [26,27], and 2) post-BS, which adds adaptors by random priming after BS conversion [28]. In total, we tested seven different combinations of BS conversion and library preparation protocols (Table 2). The pre-BS approach involves two DNA fragmentation steps (DNA sonication before library preparation and subsequent BS-induced degradation), and thus requires larger amounts of DNA input (commonly 0.5-5µg). Post-BS approaches overcome this shortcoming, where BS treatment precedes the adaptor tagging and serves to both convert and fragment the DNA, thus utilising only one fragmentation step. This strategy has led to significant reduction in DNA loss and allowed the successful generation of amplification-free WGBS libraries from as little as 400 oocytes [29,30]. Moreover, adding PCR amplification to the original amplification-free post-BS technique allowed sequencing of even lower cell numbers (~200) and single cells [31,32]. Here we tested the original amplification-free method Post-Bisulfite Adaptor Tagging (PBAT) [28], the PBAT modification with amplification ('ampPBAT') [31,32], and the commercially available EpiGnome (currently TruSeg) post-BS kit [33,34]. To dissect polymerase differences, we have also included a pre-BS approach performed with the low-bias KAPA HiFi Uracil+ [23] to compare with the most commonly used Pfu Turbo Cx polymerase (Table 2).

Table 2. Library preparation parameters of WGBS strategies compared it this study.

Method	Strategy	BS conversion	PCR cycles	Polymerase
'Alkaline'	Pre-bisulfite	Alkaline	15-18	Pfu Turbo Cx
'Heat'	Pre-bisulfite	Heat	10-16	Pfu Turbo Cx
'KAPA'	Pre-bisulfite	Heat & Alkaline	11-15	KAPA Uracil+
'Am-BS'	Pre-bisulfite	Am-BS	9	JumpStart
'PBAT'	Post-bisulfite	Heat	None*	-
'ampPBAT'	Post-bisulfite	Alkaline†	9-12*	KAPA HiFi
'EpiGnome'	Post-bisulfite	Heat	9-10*	FailSafe

<sup>\*</sup> Includes one, three or five steps of strand synthesis and pre-PCR enrichment (see Additional File 1). † modified from manufacturer's protocol (see Additional File 1).

<sup>&</sup>lt;sup>b</sup> EpiTect Bisulfite kit (Qiagen), FFPE protocol and doubled incubation time (see Materials and Methods)

<sup>&</sup>lt;sup>c</sup> EZ DNA Methylation Kit (Zymo Research)

<sup>&</sup>lt;sup>d</sup> protocol conditions as described in Hayatsu et al. [25]

In order to increase relevance, comparability and robustness of our analyses, but also to avoid analysing batch variability instead of method variability, we have combined datasets generated by our lab with a wealth of publically available datasets generated by different labs (~200 datasets in total, see Additional file 1) [28–32,34–42]. This ensures that the trends described herein are observed across data generated by a wide scientific community and not inherent to a single lab's results.

# Effect of BS-induced DNA degradation

DNA degradation is a well-known concomitant effect of BS conversion, which has made challenging its usability for low cell numbers, but has never been reported as a factor creating sequence biases. BS-induced fragmentation was initially attributed to loss of purines [1,7], but was later shown to result from random base loss at unmethylated cytidines, which causes backbone breakage upon exposure to heat and alkali [24]. Such cytosine-specific effect could lead to two possible biases: 1) depletion of cytosine-rich DNA from the total sequence pool, resulting in a skewed representation of genomic sequences, and 2) depletion of unmethylated fragments, leading to an overestimation of the absolute 5mC values. To test these possibilities, we BS treated synthetic DNA fragments of low (15%, 'C-poor') or high (30%, 'C-rich') cytosine content (see sequences in Additional file 2: Table S1). Strikingly, the recovery of the C-poor fragment was 2-fold higher than that of the C-rich fragment when using the 'Heat' BS treatment (Fig. 1a). The milder 'Alkaline' denaturation showed higher recovery and reduced bias across cytosine contents (1.3-fold difference), whereas the 'Am-BS' protocol showed no significant difference between C-contents, despite its relatively low recovery (Fig. 1a). Reducing BS incubation temperature from 65°C to 50-55°C at the expense of longer incubation (Table 1) did not produce a difference in yields (Additional file 2: Fig. S1a). Results from 'Heat' 1 and 2 or 'Alkaline' 1 and 2 pairs have therefore been pooled as 'Heat' and 'Alkaline' in subsequent analyses, unless otherwise stated. These results suggest that BS conversion conditions could have an impact in genomic coverage in WGBS.

To test whether DNA degradation leads to uneven sequence coverage in WGBS data, we sought genomic regions where the relative strand coverage could be affected by the depletion of cytosines. Both the major (pericentric) satellite repeat and mitochondrial DNA (mtDNA) display substantial differences in cytosine content between their lower and upper strands (Additional file 2: Table S2). To exclude interference of PCR bias, we only analysed amplification-free PBAT datasets, which employ heat-based DNA denaturation (Table 2 and Additional File 1). Both mouse major satellites and mtDNA showed significantly higher coverage of their C-poor (12-14% cytosine) in comparison to their C-rich (23-24% cytosine) strand (Fig. 1b). We also examined the telomere repeat, comprised of 50% cytosines on one strand ([CCCTAA]<sub>n</sub>, 'C-strand') and none on the other ([TTAGGG]<sub>n</sub>, 'G-strand'). BS sequencing reads from tandem telomere repeats showed up to 1000-fold higher coverage of the G-strand compared to the C-strand (Fig. 1c and Additional file 2: Fig. S1b). Notably, whilst BS sequencing cannot distinguish BS-converted CCCTAA repeats from genomic TTTTAA repeats, the abundance of the latter cannot explain the observed bias (not shown). These results confirm that the WGBS output of unmethylated C-rich sequences is affected by BS-induced degradation.

To test the effect of cytosine modifications on DNA fragmentation, we generated 5mC- and 5hmC-modified C-poor and C-rich fragments. Both modifications yielded a ~4-fold increase in recovery for the C-rich sequence and more than two-fold for the C-poor sequence with the 'Heat' BS-conversion protocol (Fig. 1d). A weaker protective effect on the C-rich fragment was observed for the 'Alkaline' conversion protocol, whereas the 'Am-BS' protocol showed 3-4 fold increase in recovery for both fragments (Additional file 2: Fig. S2a). This indicates that cytosine modifications have a protective effect against BS-induced DNA degradation, especially in C-rich sequences. Finally, analysis of BS converted DNA from mouse embryonic stem cells (mESC) by liquid chromatography coupled to mass spectrometry (LC/MS) revealed that both highly degrading 'Heat' and 'Am-BS' protocols cause a direct 5-10% increase in the global estimate of DNA methylation, whilst no such effect was observed for the milder 'Alkaline' procedure (Fig. 1e). Differences in DNA clean-up procedures affected overall

yields (Additional file 2: Fig. S2b), but were not responsible for the observed differences in the estimation of methylation by LC/MS (Additional file 2: Fig. S2c).

In summary, BS-induced DNA degradation leads to depletion of genomic regions enriched for unmethylated cytosines, which creates a biased sequence representation and directly affects the final estimation of 5mC levels. DNA degradation is strong in harsher BS conversion protocols that utilise high denaturation temperatures ('Heat') or high BS molarity ('Am-BS').

# Effect of PCR amplification bias

PCR amplification is a notorious source of bias in massively parallel sequencing, known to affect primarily sequences with highly skewed base composition on the extreme ends of GC content [18]. This has led to technical difficulties, which have partially been resolved by new amplification-free approaches [22], PCR buffer additives [16,18–20], better temperature control over PCR steps and cycle ramp rates [16,21], and extensive screens for low bias polymerases [16–20,23]. The mammalian BS converted genome has ~80% AT content and ~20% G content, which makes it a real challenge for polymerases. Although the commonly used Pfu Turbo Cx polymerase is not the worst performer among its counterparts [17], the KAPA HiFi family of polymerases used together with the PCR additive TMAC (tetramethylammonium chloride) have shown the best tolerance for AT-rich regions, albeit at the expense of higher error rates [18–20,23]. A more recent study suggested that the bias resulting from PCR in NGS stems primarily from factors such as stochasticity of amplification of low-copy sequences and polymerase errors [21]. These factors seem especially relevant for BS converted DNA, given the high degradation of input material and the higher reported rate of sequencing errors by polymerases in high AT content DNA.

To evaluate the effect of PCR biases on sequence representation, we quantitated the dinucleotide coverage in all datasets from our panel of WGBS methods (Table 2 and Additional file 1) against the expected genomic value (Additional file 3) and compared to a non-converted control. All methods showed a highly significant dinucleotide coverage bias relative to control (Fig. 2a and Additional file 2: Fig. S3a). There was a clear enrichment for G-containing dinucleotides and depletion of AT-rich sequences in all methods, with the exception of 'KAPA', which showed a balanced G content, an unexpected depletion of C content and enrichment of AT content (Fig. 2a and Additional file 2: Fig. S3a). The effects of KAPA HiFi Uracil+ amplification were independent of the BS conversion protocol used (Additional file 2: Fig. S3a). Interestingly, the amplification-free PBAT also showed a slight Gbias, possibly due to its DNA synthesis and pre-PCR enrichment steps (Fig. 2a, right panel). However, unlike other methods, amplification-free PBAT did not display any significant deviation from control with respect to CG-dinucleotide coverage, where most methylation occurs, suggesting that the amplification-free post-BS approach has the lowest methylation biasing. Amplification-free PBAT datasets also showed the highest consistency in base coverage between individual datasets (Coefficient of Variation, CV = 1.55 %) in comparison to all other methods with amplification (Additional file 2: Fig. S3a). The remaining post-BS methods showed moderate variability (CV = 2.99-3.04 %), whereas most pre-BS protocols were highly variable (CV = 4.52-9.15%), the exception being again 'KAPA', which had a CV of 3.07%., These results link to the reported effect of PCR stochasticity in generating dataset-to-dataset technical variability and biases [21], highlighting that the amplification step generates variability in the quality of WGBS output.

We next asked in more detail how sequencing depth was affected by the % of G or C. These analyses confirmed that KAPA-amplified libraries displayed the lowest amount of G content bias, in contrast to the drastic G-enrichment of Pfu Turbo Cx's 'Heat' and 'Alkaline' pre-BS datasets (Fig. 2b upper panels). The post-BS methods also showed a more balanced G coverage, performing similarly regardless of amplification (Fig. 2b lower panels). With respect to C coverage the post-BS methods strongly outperformed the pre-BS group, where both 'KAPA' and Pfu Turbo Cx's 'Heat' and 'Alkaline' datasets under-represent the C-high (> 25% C) and over-represent the C-low (< 15% C) sequences. This is a key result, suggesting that the BS-degradation bias, characterised by depletion of C-rich sequences, does not affect post-BS approaches to the extent that it affects pre-BS protocols.

Next, we investigated how polymerase bias affected regions with known GC-skew between strands. For both the satellite repeat and mtDNA, PCR amplification exacerbated the strand bias arising from DNA degradation observed in 'PBAT' in Fig. 1b (Additional file 2: Fig. S3b), demonstrating a combined effect of the PCR and degradation biases (lowest for the 'KAPA' datasets). Similarly, the C-rich strand of the telomere repeat was hardly detectable in the raw reads of most datasets (Additional file 2: Fig. S3c).

Finally, to assess the effect of amplification on quantitation of 5mC, we used LC/MS to measure total 5mC levels in mESC within our panel of WGBS datasets (except for 'Am-BS', for which mESC data was not available). The levels in both 'Heat' and 'Alkaline' datasets doubled the LC/MS-measured 5mC value, whilst the overestimation in the 'KAPA' and post-BS datasets was less pronounced (Fig. 2c). Notably, 5mC levels in amplification-free PBAT were not significantly different from those measured in genomic DNA (Fig. 2c) and were comparable to those detected after BS conversion alone (Fig. 1e).

In summary, all WGBS approaches show significant sequence coverage deviations in comparison to conventional WGS. Post-BS approaches, however, show better representation of C-rich and C-low sequences, resulting in a less pronounced overestimation of global methylation values. They also show less technical variability in the quality of sequencing output between datasets.

## **Effect of DNA modifications**

To further evaluate the genome-wide influence of DNA modifications on WGBS biases, we compared gDNA from DNMT-Triple Knockout (TKO) mESCs, which lack DNA methylation, to in vitro methylated TKO gDNA ('meTKO') using the M.CviPI methylase. We sequenced TKO and meTKO WGBS libraries generated using the 'Heat' protocol, which was strongly affected by depletion of C-containing sequences in our previous analyses (Fig. 2a and 2b). The meTKO library displayed ~20% methylation across all cytosine contexts (Additional file 2: Fig. S4a) and this translated into a ~15% higher sequencing coverage at all cytosine-containing dinucleotides when compared to the TKO library (Fig. 3a). This shows that DNA methylation affects sequence coverage, as suggested by our experiments with modified DNA fragments (Fig. 1d). Interestingly, differences in total 5mC levels between meTKO and WT mESC DNA (15% versus 3%, by LC/MS) led to differences in the overestimation of 5mC by WGBS (40% versus 100% overestimation; Fig. 3b and 2c), presumably because meTKO DNA has fewer unmethylated cytosines available for BS-induced degradation. The accuracy of local 5mC measurements by 'Heat' WGBS therefore depends on the extent of methylation at each locus. We also found that DNA methylation affected read coverage depending on the % of C or G content (Fig. 3c). Notably, the even distribution of in vitro deposited 5mC was associated with a more even coverage across C/G content when compared to TKO DNA (Fig. 3c), suggesting that a substantial amount of the coverage bias in WT mESCs (Fig. 2b) is driven by differences in the genomic distribution of 5mC. An increase in coverage, albeit weak, was also observed for the satellite repeat and mtDNA C-rich strands (Additional file 2: Fig. S4b).

These results suggest that highly methylated sequences could be over-represented in WT genomes. To explore how methylation status affected GC-rich regions of interest, we compared averaged CGI coverage in our panel of mESCs WGBS datasets. Low levels of 5mC seen at CGIs have the potential to cause a coverage bias in comparison to entirely unmethylated genomes; therefore we also included unmethylated samples for all methods apart from 'KAPA' and 'EpiGnome' (which were not available). The 'Heat' pre-BS method showed highest coverage bias over WT mESC CGIs, which decreased significantly in the unmethylated sample (Fig. 3d). The post-BS methods showed low ('PBAT') to moderate ('ampPBAT' and 'EpiGnome') coverage bias, which also decreased in the unmethylated samples (Fig. 3d, right panel). No methylation-dependent coverage bias was detected with the 'Alkaline' protocol, and an under-representation of CGIs was observed for the AT-biasing 'KAPA' datasets. These results confirm that particular C/G-rich sequences will be over- or under-represented in WGBS datasets, depending on their methylation status and the chosen library preparation strategy.

## Effect of incomplete BS conversion

It is known that different BS treatment conditions have variable efficiency in converting cytosine to uracil [2,5,14]. Given the variable preference of polymerases towards cytosine and the depletion of successfully converted cytosines through BS-induced degradation, we asked whether unconverted cytosine artefacts could contribute to the observed overestimated methylation values in WGBS datasets (Fig. 2d). To measure conversion efficiency of the BS conversion protocols in Table 1, we BS-treated mESC gDNA and quantified the amount of unconverted cytosines by LC/MS (LC/MS can distinguish unconverted Cs from 5mCs). Our results show that 'Heat' denaturation yielded best conversion, whereas 'Alkaline' denaturation led to 4-fold higher amounts of unconverted cytosine (Fig. 4a). When added to the total 5mC levels measured by LC/MC (Fig. 4b) it is clear that unconverted cytosines make a substantial contribution to the overestimation of 5mC levels by WGBS (Fig. 2c). This effect will be strongest for the milder 'Alkaline' conditions, which was confirmed in our WGBS datasets of the same unmethylated TKO samples prepared with the 'Heat' and 'Alkaline' methods (Fig. 4c). Strikingly, the % of uncoverted cytosines in the 'Alkaline' TKO datasets surpassed the biological value of real 5mC levels in WT mESCs, demonstrating how unconverted cytosines can lead to vastly increased total 5mC estimates.

Conversion artefacts occur mainly in non-CG context (or CH, where H is A, T or C) (Fig. 4d). This is because CH context is over 20-fold more abundant in mammalian genomes than the CG context (Additional file 3). We therefore sought examples of genomic locations that could be particularly affected by this artefact. Mouse major satellites have been shown to have non-CG context methylation through a BS cloning based approach [43]. As targeted BS sequencing relies on primers that select for fully converted fragments [9], these results are more likely to reflect the real non-CG methylation distribution. We therefore produced our own BS cloning data for major satellites to compare against WGBS results (Fig. 4e and Additional file 2: Fig. S5a,b). Analyses of our panel of WT mESC WGBS datasets showed that, unlike what is seen by BS cloning, pre-BS 'Alkaline' datasets had pronounced strand asymmetry in the CH methylation levels, which were more abundant on the C-rich (bottom) than the C-poor (top) strand (Fig. 4e, left panel). The 'Heat' protocol had a reduced bias, whereas the 'KAPA' method did not show strand asymmetry, although certain positions had higher mCH values than those obtained by BS cloning. None of the post-BS methods showed strand asymmetry, even after amplification (Fig. 4e, right panel). These results can be explained by the preferential amplification of poorly converted reads, which are less likely to be degraded, whereas well converted C-rich reads from the bottom strand tend to be degraded, as we have shown. This gives the false perception of asymmetric methylation in such regions. Indeed, asymmetrically methylated regions are commonly reported in BS-seq datasets, especially in CH context [44]. Notably, mCG levels were also asymmetrically elevated in the 'Heat' and 'Alkaline' pre-BS approaches in comparison to the post-BS methods (Additional file 2: Fig. S5c). These results illustrate a direct link and interplay between sequence-specific BS-induced degradation, conversion errors and the amplification of those artefacts by PCR, leading to higher overall bias in pre-BS library preparation protocols.

## Effect on quantitation of methylation

To investigate how the different biases affect the final quantitation of methylation in genomic features we analysed two sets of data: 1) Set 1 included the previously used public mESCs WGBS datasets prepared with six different protocols by different teams [29–32,34,37,38,40,45] and 2) Set 2 included a combination of 'PBAT' and BS-seq 'Heat' datasets from four different biological samples (mESC, blastocyst, oocyte and sperm) prepared by a single team [29].

We have already shown how the different methods in Set 1 performed in estimating global 5mC levels (Fig. 2c) and, apart from 'PBAT', all showed a significant increase in CG coverage (Fig. 2a). To test whether this indeed translates into overestimation of CG methylation, we first looked at the methylation levels of imprinted differentially methylated regions (iDMRs), which are expected to have

around 50 % CG methylation in mESCs. 'PBAT', which showed a global 5mC value nearest to the LC/MS value (Fig. 2c) yielded the lowest DMR methylation (44 %), followed by 'ampPBAT' (48 %), 'Alkaline' and 'KAPA' (both 53 %), and 'Heat' and EpiGnome (both 58 %) (Fig. 5a). 'PBAT' consistently yielded the lowest values of methylation across multiple genomic features such as genes, intergenic regions and repeats, whereas the rest of the methods showed comparably higher mCG values for all but the highly methylated IAP repeats (Additional file 2: Fig. S6a). This prompted us to test whether the methylation increase over the 'PBAT' values was linear across all 5mC values, for which we did a genome-wide comparison of each method against PBAT. This revealed that the largest discrepancies lay in the middle range of CG methylation values, i.e. the moderately or variably methylated genomic regions (Fig. 5b and Additional file 2: Fig. S6b). The same result was obtained when comparing 'Heat' and 'PBAT' data for the four biological samples from Set 2, but not for the individual 'Heat' replicates, validating that the observed non-linear discrepancies are not due to batch effects or lab-to-lab variability (Additional file 2: Fig. S6c). Importantly, moderately methylated regions are commonly studied targets in biological samples, since they indicate areas of variability and heterogeneity within an epigenetic pool and often include enhancers, promoters and transcription factor binding sites [46]. A closer look into such features in Set 1 confirmed that they display larger methylation differences between PBAT and the other methods when compared to genic and repeat regions (Fig. 5c and Additional file 2: Fig. S7a-c). These results further highlight that some regions are more susceptible to biases than others and the overestimation of 5mC across the genome does not follow a linear fashion.

Despite the clear differences in absolute methylation values between the WGBS methods, DNA methylation is often studied as a relative change between conditions, treatments and biological samples. To address whether relative changes in methylation differ when analysed with pre- or post-BS WGBS, we analysed methylation differences between BS-seq and PBAT datasets from sperm and ESC from Set 2. We selected regions with more than 20% methylation difference between the two samples in both directions from one of the methods, and compared the positioning of those regions in the other method (Fig. 5d and Additional File 2: Fig. 8a). More than half of the regions identified with the 'Heat' method were also identified with 'PBAT' but the larger proportion of regions from 'PBAT' were not identified with 'Heat' (Fig. 5e). This is an important finding, showing that researchers can obtain different results and be led to different conclusions depending on which WGBS method they use in their study.

Our results highlight that overestimation of mCG values is a common feature of WGBS protocols with amplification, despite their performance in various bias tests. This is explained by our observations that different sources of bias contribute to this effect in the different protocols, and we have summarised our estimates for some of our datasets in Figure 5f.

## Coping with biases and artefacts

We have shown that many of the biases in WGBS datasets reflect themselves in sequence composition of the libraries. To aid with the evaluation of WGBS biases, we have incorporated, as part of the quality control (QC) package of the Bismark program [47], a 'bam2nuc' module to quantitate the mono- and di-nucleotide composition in a WGBS dataset against the genomic expected values (Fig. 6a). A depletion of cytosine mono- and dinucleotide content is likely to be a result of BS-induced DNA degradation, whereas cytosine enrichment would indicate poor conversion efficiency; G or AT depletion/enrichment could serve as measures of amplification bias and polymerase preferences. For specific sequences, the publically available GC-content tracks on genome viewers could be used as an indication for likelihood for biases. We have also created and made public (see Declarations: Availability of Data and Materials) strand-specific G and C content wiggle tracks for the mouse genome.

Another strategy is related to the way methylation is quantitated in regions of interest. Our results showed that polymerase and PCR biases can enrich the sequence output of WGBS datasets either towards GC-rich or towards AT-rich sequences (Fig. 2a). We experimented with common ways

methylation can be estimated and identified a strategy that is less affected by coverage biases (Fig. 6b). Namely, quantitating methylation for individual cytosines within a region of interest and averaging their values outperforms the alternative practice of pooling all methylation calls within a region. Importantly, the very common practice of selecting for analysis only cytosines with a minimal fold coverage (usually above 5 or 10), results in reinforcement of the coverage bias effects and skews the resulting methylation values (Additional File 2: Fig. S9a).

Incomplete conversion artefacts affect CG methylation and are a subject of over-amplification during PCR, but their most dramatic effect is on non-CG methylation. Spiking of unmethylated DNA of foreign origin during library preparation, such as Lambda or M13 phage DNA, is a useful way to monitor the global conversion efficiency per sample (Additional file 2: Fig. S10a). However, BS conversion resistance is very sequence specific [3,11] (Additional file 2: Fig. 10b) and thus not fully represented by a control of different origin and composition. It is also common to use genomic non-CG context methylation as an indication for conversion efficiency, which may be appropriate for some samples but also risky, given the well documented presence of mCH in mammalian, insect and plant genomes [9,26,27,29,30,39,40,43,48,49]. Different approaches can be applied to distinguish real mCH from artefacts and reduce the weight of false positive methylation calls. Here we compared three strategies to cope with conversion errors, tested on the major satellite consensus and our M13 spike-in controls: 1) removal of CH calls below a threshold methylation value (usually set between 3 and 20 %), 2) filtering of reads with 3 or more consecutive unconverted CH bases (the 3xC filter), and 3) a new approach for normalisation of each cytosine's methylation against an unmethylated WGBS control of the same genome. Although the first approach is the most commonly used [48,50,51], our results show that it is the least efficient one, since a large number of false positive calls remain above the threshold, which at the same time is likely to remove real methylation calls (Fig. 6c and Additional file 2: Fig. 10b). The second approach is much more efficient in removing the background noise, although a number of conversion resistant GC-rich sites (such as CCWGGs) can pass the filter (Fig. 6d and Additional file 2: Fig. 10c). For the third approach, we used our unmethylated TKO mESC WGBS datasets prepared with 'Heat' and 'Alkaline' protocols (Additional file 2: Fig. 10d) as background noise controls to subtract from the corresponding WT mES datasets. This substantially reduced the noise from CH context positive calls (shown in Fig. 4e) to levels comparable to those achieved with the classic BS cloning and the 3xC bioinformatic filter (Fig. 6e). The main benefit of this approach is the ability to deal with conversion resistant sequences, such as the ones we observe in the M13 spike-in, which pass through the 3xC filter (Additional file 2: Fig. 10b). Our results demonstrate that WGBS is not a noise-free technique, therefore studies interpreting non-CG methylation should be accompanied with robust controls and have clear strategies for coping with conversion artefacts.

## **Discussion**

Here we present a comparative analysis between five BS conversion methods and seven WGBS library preparation protocols, revealing the most common sources of bias. We have evaluated their performance and summarize our results in Table 3.

Our findings reveal that WGBS protocols suffer from multiple biases and have a highly variable performance, a fact that has not received due attention to date. The biases lead to overestimated absolute levels of both CG and CH context methylation, skewed relative methylation differences between samples and under- or over-representation of vulnerable genomic regions. These unwanted effects can be controlled by a careful selection of the library preparation strategy and specific conditions during key steps, but are best avoided with an amplification-free post-BS approach (Table 3).

Table 3.Summary of biases affecting the pre- and post-BS WGBS library preparation strategies. For simplicity, pre-BS methods have been divided into GC-biasing ('Heat', 'Alkaline' and 'Am-BS') and AT-biasing ('KAPA'), and post-BS methods into amplification-free ('PBAT') and with amplification ('ampPBAT' and 'EpiGnome'). As 'ampPBAT' uses KAPA HiFi polymerase, we have noted this contribution where necessary. (+) indicates low bias, (++) medium and (+++) high bias.

BIAS		PRE-BISULFITE		POST-BISULFITE	
		GC-biasing	AT-biasing (KAPA)	Amplification- free	With amplification
Degradation bias (C depletion)		+++	+++	+	+
PCR (polymerase) bias		+++	++	+	+++ (KAPA ++)
Conversion artefacts		++	++	+	+
Modified C bias		++	++ / NA	+	++
5mC overestimation	global	+++	++	+	++
	CG	+++	+++	+	+++ (KAPA ++)
Variability in output		+++	++	+	++

Our results show that BS-mediated DNA degradation plays a major role in generating biases in WGBS data, affecting the sequence composition and methylation output through depletion of unmethylated C-rich regions. This effect is especially strong in pre-BS approaches, where DNA is fragmented prior to BS conversion. Thus, the adapter-tagged library undergoes a second fragmentation step, where C-rich unmethylated fragments get excluded from the library pool before they undergo amplification, which introduces a sequence bias. As a result, the uneven representation of strands and sequences increases stochasticity in the first cycles during PCR, where certain sequences end up under- or over-represented, irrespective of polymerase GC bias [21]. Post-BS methods, to the contrary, harness the BS-induced fragmentation directly starting from high molecular weight DNA, which conveniently yields the desired fragment size [8,28] thus decreasing the loss of Chigh content and allowing for a more accurate estimation of methylation levels even after amplification (Fig. 2a-c). Notably, post-BS methods can also afford to use harsher BS conversion conditions such as heat denaturation of DNA and higher BS incubation temperatures (65-70°C), which yield a better and more consistent BS conversion efficiency, especially if combined with high molarity of BS (>4 M) and short incubation times (30-90 min) (Fig. 4a and Additional File 2: Fig.10a) [5,8,14]. Longer incubation times have been shown to lead to higher degradation and accumulation of inappropriate conversion (false negatives), without necessarily contributing to conversion efficiency [5,8]. The choice of reliable conversion conditions is particularly important for studying non-CG methylation, where the alkaline denaturation, low BS molarity and lower temperatures are likely to yield false positives, which outnumber the real 5mCH signal in a sample (Fig. 4c and 5e).

PCR bias was found to add up to the over-representation of methylated sequences and conversion artefacts. Indeed, the best results were observed for the amplification-free PBAT approach, where, in addition to its lower degradation bias, it showed insignificant CG-context coverage bias and better matched the 5mC levels measured by LC/MS (Fig. 2a-c). The amplification-free output was also least affected by the underlying methylation status (Fig. 3d) and had the lowest cross-dataset variability in output quality (Additional File 2: Fig 3a). Given that a main advantage has been its use with very low DNA input [28,29], the amplification-free method should be feasible for most standard applications. Whilst the original PBAT approach suffers from lower mapping efficiency due to chimeric reads, these are easily dealt with during the bioinformatic processing [52]. It has been also reported that PBAT is

affected by different versions of the Illumina HiSeq base calling software, which can lead to a drop of up to 5% in global mCG values [53]. These problems, however, are not common for other post-BS methods such as 'EpiGnome' and others [42,53]. Importantly, an alternative amplification-free WGBS approach called ReBuilT has been published more recently, which also shows an improvement in GC-bias [54]. Interestingly, amplification-free protocols are also the least biased solution for NGS microbiome analysis, where sequence diversity and (mis)-representation is of high importance, as in WGBS [55].

Classically PCR bias has been associated with enrichment of GC-rich sequences due to polymerase preferences [16–20,23]. The KAPA HiFi family of polymerases have been shown to have low GC-bias and best AT-tolerance, however our results demonstrate an AT-bias for KAPA HiFi Uracil+ (Fig. 2a,b and Fig. 3d). Importantly, we show that the overall low bias of KAPA Uracil+ cannot overcome the underlying degradation bias in the pre-BS approach and still leads to an overestimation of methylation levels (Fig. 5a-c and Additional File 2: Fig 7). Thus the post-BS methods with amplification perform in overall better than the pre-BS protocols (Table 3), recently reported also for post-BS approaches not tested here [42]. Investing efforts into optimising polymerase use both during DNA synthesis and amplification could further improve their performance in WGBS. It is necessary to stress, however, that all of our compared pre-BS datasets were generated with a relatively high number of PCR cycles (>10), and it is very likely that applying fewer cycles as advised in Urich et al, would lead to a less biased performance [56].

A growing number of WGBS datasets are currently available in the public domain and often datasets generated by different labs get used together in one study. Given the presented differences in the methods' absolute and relative methylation estimates, analysing and comparing data generated by different protocols should be avoided or done with caution, and the biases must be accounted for during the interpretation of results. The newly presented 'bam2nuc' module in the QC package of the Bismark software will hopefully help avoid the interpretation of purely technical methylation differences as biological, as our results show that changes up to 20% in DNA methylation can be purely technical.

Our results show the existence of conversion artefacts 'noise' that is particularly relevant in the context of non-CG methylation, especially as the latter becomes increasingly a focus of biological interest [29,30,39,40,43,48,49]. We show that bioinformatic filtering of reads with three or more consecutive unconverted CH cytosines is necessary and more efficient than setting a cut-off threshold value, and should become a standard even in datasets with high overall conversion rates. This tool has now been integrated as an optional 'filter\_non\_conversion' module in the Bismark methylation caller [47]. Alternatively, sequencing a whole genome amplified (WGA) unmethylated sample in parallel with samples of interest can be used to normalise false discovery rates with single base resolution. The latter approach is particularly important for studies of model organisms with very low or questionable methylation levels near the detection limit, such as insects. Such studies, especially reporting methylation in C-rich regions or non-CG context [44], should be backed up by unmethylated genome controls and validated with non-BS methods such as LC/MS [39].

## **Conclusions**

Our findings establish basic principles for minimising biases when designing and optimising WGBS strategies. We envisage that, in the current state-of-the-art, the gold standard for WGBS library preparation should evolve towards an amplification-free or amplification-minimal post-BS approach with optimised BS treatment conditions and bias-free DNA polymerases. Such benchmark method would be of great value for the research community and enable researchers from outside fields to always generate methylation data with minimal biases. It would also encourage the development of newer and better amplification-free protocols. New sequencing technologies, not dependent on BS treatment, will also push the field forward and help obtain degradation bias-free and conversion error-free maps of DNA methylation.

## **Methods**

## ES cell culture and DNA preparation

The J1 ES cell line (129S4/SvJae) was purchased from ATCC (Cat. SCRC-1010) and the Dnmt1-/,3a-/-,3b-/- TKO line is a kind gift from Masaki Okano [57]. The TKO was cultured on gelatine without feeders and the J1 on a  $\gamma$ -irradiated pMEF feeder layer, at 37°C and 5% CO<sub>2</sub> in complete ES medium (DMEM 4500 mg/L glucose, 4 mM L-glutamine and 110 mg/L sodium pyruvate, 15% foetal bovine serum, 100 U of penicillin/100  $\mu$ g of streptomycin in 100 mL medium, 0.1mM non-essential amino acids, 50  $\mu$ M  $\beta$ -mercaptoethanol, 10³U/ml LIF ESGRO®). Mycoplasma tests on cell lines are routinely performed in the lab. Genomic DNA was extracted with DNeasy Blood and Tissue Kit (Qiagen) following manufacturer's instructions and quantitated via Quant-iT<sup>TM</sup> PicoGreen dsDNA Assay Kit (Invitrogen). *In vitro* DNA methylation with M.CviPI (New England Biolabs, NEB) was performed on 0.5 – 1.0  $\mu$ g of TKO mESC genomic DNA, incubated for 2 hours at 37°C, purified with GeneJet PCR Purification kit (Thermo) and quantitated by Quant-iT<sup>TM</sup> PicoGreen® dsDNA Assay Kit (Invitrogen).

## M13 fragments analysis

M13mp18 (NEB) was used as a template. PCR of C-poor and C-rich fragments (Additional file 2: Table S1) was performed using either a standard dNTP mix (Bioline), or substituting the dCTPs with modified dm5CTPs (10 mM, NEB) or d5hmCTPs (100 mM, Bioline). The PCR was performed with Dream Taq DNA Polymerase (Thermo Scientific) (50 μl volume, 200 nM primer, 200 μM dNTPs, 1.25 units enzyme) with an initial step at 95°C for 2 minutes, 35 cycles of 30 seconds at 95°C, 20 seconds at 57°C, and 30 seconds (C, hmC) or 5 minutes (mC) at 72°C, with a 7 minute final step at 72°C (primers are listed in Additional file 2: Table S2). All PCR products were verified on a DNA resolving 2% agarose gel, purified with GeneJet PCR Purification kit (Thermo Scientific) and quantitated by both Quant-iT<sup>TM</sup> PicoGreen dsDNA Assay Kit (Invitrogen) and Agilent 2100 Bioanalyzer. Identical aliquots were prepared from each fragment for BS treatment with the different protocols and an aliquot each was kept as an input control. BS treated fragments were eluted in the same final volume as the input and quantitated for recovery 3 to 4 times for each sample. Each BS treatment was repeated twice.

# **Bisulfite conversion of DNA**

Genomic DNA and purified M13-derived fragments were treated with sodium bisulfite using all of the following kits: EpiTect Bisulfite kit from Qiagen (FFTP protocol), Imprint DNA Modification kit from Sigma-Aldrich (1-step and 2-step) and EZ DNA Methylation Kit (Zymo Research) according to manufacturer's instructions. The *in vitro* M.CviPI-methylated TKO DNA was converted with the EpiTect Bisulfite kit. A minor modification was applied for all samples treated with the EpiTect kit: the 5 hour incubation programme was run twice (10 hours) following a commonly accepted practice [26,49,58,59]. Conversion with 9 M ammonium bisulfite was performed at 70°C for 30 minutes as in Hayatsu et al. [25]; 50% Ammonium Hydrogen Sulfite Solution was purchased from Wako Chemicals GmbH, the rest of the reagents were supplied by Sigma-Aldrich. Half of the samples converted with the 1-step ('heat') and 2-step ('alkaline') Imprint DNA Modification kit and the 9M ammonium bisulfite procedure were purified with Amicon Ultra 0.5mL Ultracel 30k filters (Millipore), with the clean-up reagents and following the manufacturer's purification instructions of the TrueMethyl kit v1 (Cambridge Epigenetix).

# Purification and BS-sequencing of the major satellite repeat

J1 (WT) and TKO ES [57] genomic DNA was bisulfite converted using EpiTect Bisulfite Kit (Qiagen) as explained previously. The major satellite was amplified with HotStart Taq (Qiagen) in a mixture of 200 nM primer, 200 µM dNTPs, 2 mM MgCl<sub>2</sub>, 1.0 unit of enzyme at 94°C for 15 minutes, 35 cycles of

20 seconds at 94°C, 20 seconds at 55°C, and 20 seconds at 72°C, and a final step at 72°C for 3 minutes. DNA fragments spanning over one repeat (370bp) were excised from 2% agarose gels and purified with a MinElute Gel Extraction kit (Qiagen) following the kit protocol. The fragments were cloned into pGEM-T using the pGEM-T Easy Vector Kit (Promega) and transformed into Invitrogen's Subcloning Efficiency DH5α Competent Cells according to manufacturer's instructions. Positive clones were selected on LB plates containing 100 μg/ml ampicillin and covered with X-gal (40 mg/ml). Colonies were screened with Roche's Taq DNA Polymerase (25 μl volume, 300 nM primer, 200 μM dNTPs, 1.25 units enzyme) at 94°C for 10 minutes, 35 cycles of 30 seconds at 94°C, 30 seconds at 55°C, and 30 seconds at 72°C, with a final 72°C for 10 minutes and sent for Sanger sequencing at Beckman Coulter Genomics. All oligonucleotides are listed in Additional file 2: Table S2. Methylation of the sequenced clones was analysed with QUMA [60] and plotted with a custom made R script.

# Mass spectrometry

Untreated or BS treated genomic DNA (0.3-1 µg) was digested with a DNA Degradase Plus™ (Zymo Research) for 3 hours at 37°C according to manufacturer's instructions. Approximately 50-100 pg per sample were analysed by LC-MS/MS on a Thermo Q-Exactive mass spectrometer coupled to a Proxeon nanoLC. Three replicates of each sample were analysed and the amounts of C, 5mC, 5hmC and U and T were quantified relative to external standards. Recovery of BS treated genomic DNA for the different BS conversion protocols and clean-up procedures was assessed in the same way as for the M13 fragments, but quantitated with the LC/MS.

# Library preparation and next generation sequencing

Approximately 250 ng genomic DNA was fragmented via sonication with a Covaris E220 instrument with the 300 bp programme, and spiked in 1:10,000 with a 2 kb unmethylated PCR fragment from M13mp18 (New England Biolabs). Early Access Methylation Adaptor Oligos (Illumina) were ligated to the fragmented DNA with the NEB Next DNA Library Prep Master Mix Set for Illumina (E6040), according to the manufacturer's instructions and purified after each step with Agencourt® AMPure® XP beads. Bisulfite treatment was performed as described above in 'Bisulfite conversion of DNA', using all listed methods, except for the ammonium bisulfite protocol. The BS converted libraries were amplified using PfuTurbo Cx Hotstart DNA Polymerase (Agilent Technologies): 300 µM dNTPs, 400 nM indexed adaptor-specific primers[19], 2.5 units enzyme, with an initial step at 98°C for 30 seconds, 15 cycles of 98°C for 10 seconds, 65°C for 30 seconds, and 72°C for 30 seconds, with a final elongation step at 72°C for 5 minutes. Library quality control was performed with an Agilent 2100 Bioanalyzer and quantity determined via KAPA Library Quantification Kit (KAPA Biosystems). For the unmethylated DNA controls, library preparation was performed in the same way, with the following modifications: 1.0 µg of DNA was sonicated and adaptor-ligated with Illumina TruSeq indexed adaptors, no BS conversion was performed, and amplification was done with the NEB Next 2x Phusion mix for 6 cycles, following manufacturer's instructions.

Paired-end 100 bp NGS was performed on an Illumina HiSeq 2000 system at the Bespoke Facility at the Wellcome Trust Sanger Institute.

# WGBS data mapping and quality analysis

Data from both BS converted and non-BS converted datasets, was trimmed with Trim Galore and raw data quality analysis performed with FastQC [61]. Mapping was carried out with Bismark [47] to NCBIM37 and GRCm38 builds for the mouse genome, GRCh38 for the human genome, HS3.3 for the ant *Harpegnathos saltator*, GCA\_000297895.1 for the Pacific oyster *Crassostrea gigas*, Chinese hamster reference sequence in Ensembl for the CHO cells, and assembly scaffolds for the ant *Dinoponera quadriceps*. For consistency and to reduce error, the non BS converted datasets were also mapped with Bismark. All alignments were performed with high stringency allowing for only one base

mismatch (n=1) and mapped data was deduplicated before analyses. For PBAT libraries, all mapping errors from chimeric reads and M-bias were taken into consideration upon processing and first 4 bases from each read were excluded from the analysis [52]. All datasets were deduplicated, consistent with common analyses pipelines, although this decreased the sequence and methylation bias, normally stronger in the raw duplicated data (unpublished observation). Some of the analyses, however, such as telomere and major satellite, were performed on raw (duplicated) data, since those sequences are not mappable.

## Read coverage analysis

After processing with Bismark, read coverage depth was analysed with SeqMonk [62]. Custom genomes were created for mtDNA, satellite repeat and M13, and the reads were aligned preserving the strand information. Because of its repetitive nature, the data for major satellite was not deduplicated following alignment. Read coverage was assessed from the total read count for forward or reverse strands in each custom-built genome.

For the mouse CGI coverage, regions with unusually high read coverage (1kb tiled genomic probes with more than 1000 reads), likely to represent alignment artefacts were excluded from the analysis. The CGI coverage analysis was performed in SeqMonk via a relative trend plot over CGIs (coordinates from Illingworth et al. 2010 [63]), including all reads, 'forced to be relative' option chosen and allowing for 1kb flanks.

## Methylation analysis of WGBS

CG methylation quantitation was performed in SeqMonk with the integrated bisulfite analysis pipeline. Regions likely to attract alignment artefacts (having more than 1000 reads over 1kb tiled genomic probes) were excluded from the analyses. CG methylation analysis on genic and regulatory features was quantitated on the individual replicate datasets via probes created over each feature, without setting cytosine coverage thresholds but requiring minimum 3 observations in order to include a probe. This analysis was undertaken only for mouse ESC datasets, for which public datasets were available for all compared protocols, except for Am-BS (see Additional file 1 for details). Promoters were defined as -1000 + 200bp from the TSS, DMR coordinates were obtained from Seisenberger et al. [38], CGI coordinates from Illingworth et al. 2010 [63], active, poised and primed enhancers from Creyghton et al. 2010 [64], super-enhancers from Whyte et al. 2013 [65], ES-specific LMRs from Stadler et al. 2011 [46], Yamanaka factor binding sites from Chen et al. 2008 [66] and transcription factor binding sites (TFBS) from UCSC (Caltech annotation). Exons and introns were defined with Ensembl-derived coordinates integrated in SeqMonk. IAP, LINE, LTR, Satellite and SINE coordinates were derived from UCSC [67].

The genome-wide analysis (scatter plots) was performed for two groups of datasets: 1) the panel of published mouse ESC datasets from six WGBS methods generated by different studies or labs and used for the feature analyses (see Additional file 1), and 2) datasets generated by the same lab with PBAT and Heat BS-seq from four different biological samples (mESC, sperm, blastocyst and oocyte) [29]. The first group provides comparison between all protocols (except for Am-BS due to unavailability), while the second set serves to validate the observations from the first group with the difference that it should not be affected by potential sample strain or batch differences. For the first group probes were made over non-overlapping 50-cytosine containing tiles over the PBAT mESC datasets and quantitated for all remaining datasets as pooled replicates. For the second set of datasets 150C probes were made over each corresponding PBAT dataset and quantitated over pooled or individual replicates for the PBAT vs Heat BS-seq comparison or for the inter-replicate comparison, respectively.

The relative methylation analysis was performed with PBAT and Heat BS-seq datasets from Kobayashi et al. [29], used for the whole genome analysis above. We made probes over 150-cytosine containing tiles over the whole genome for the mESC PBAT dataset (due to lowest coverage) and

quantitated CG methylation for these probes in PBAT sperm, as well as in the BS-seq datasets for both samples. We then selected regions with 20% CG methylation difference between sperm and ESC in both the PBAT and BS-seq datasets and plotted the regions obtained with PBAT onto the BS-seq plot and vice versa. Overlapping differentially methylation probes between the PBAT and BS-seq lists were quantitated in SeqMonk and plotted as venn diagrams with R.

CH filtering of major satellite and M13 was done by removing every read containing more than three unconverted CH cytosines. The script can be found in the provided Github deposition and also integrated as 'filter\_non\_conversion' module in the Bismark package v0.17.0 [47]. All analyzed datasets and the number of replicates per protocol are listed in Additional file 1.

# Sequence composition analyses

All described sequence composition analyses were performed with SeqMonk and custom-made Perl and R scripts. All scripts can be accessed from the Github deposition directory provided under Declarations.

Dinucleotide coverage was assessed through the quantitation of the total number of dinucleotide instances within the mapped data of each dataset. It was plotted against the expected occurrence of each dinucleotide derived from the relevant annotated genome (see Additional file 3 for genomic references). This analysis is integrated as 'bam2nuc' module in Bismark v0.16.0 [47].

For the telomere analyses we used only raw data reads, since the tandem hexamer units of the telomere are not mappable. We quantitated the number of occurrences of each hexamer (or heptamer for *A. thaliana*) per read, as follows: TTAGGG (G-strand hexamer) or TTTTAA (BS converted version of the CCCTAA C-strand hexamer) for BS-seq and EpiGnome datasets; the reversed sequences CCCTAA(A) or (T)TTAAAA for PBAT and ampPBAT datasets (heptamers, in parentheses, were used for *A. thaliana*); TTAGGG, CCCTAA or TTTTAA for the non-BS converted control. The TTTTAA hexamer was quantitated in the non-BS converted control in order to assess the original genomic occurrence of this non-telomere sequence prior to BS conversion. Read lengths varied between 30-100bp for BS-seq datasets, 44bp, 76bp or 121bp for PBAT and 100bp for the non-BS control, hence the variation in number of units in Fig. 1c and S1b. Our quantitation revealed that TTTTAA and (T)TTAAAA occurred mostly in one to five units per read before BS conversion, indicating that the majority of those reads are native to the genome and do not derive from the telomere repeat. The telomere hexamers (T)TTAGGG and CCCTAA(A), however, were present in higher numbers per read. In order to exclude the non-telomere derived TTTTAA and (T)TTAAAA hexa/hepta-mers from the BS treated data, therefore, all reads containing less than 5 units per read we removed from the analyses.

For the correlation of read coverage with C and G content, we generated non-overlapping 100bp running window tiles over the whole mouse genome. Regions likely to attract alignment artefacts (defined previously as having more than 1000 reads over 1kb tiled genomic probes) were excluded from the analysis. Read count of each 100bp-tile was quantitated and base composition was extracted from the genomic sequence and not the actual read sequence, where cytosines are BS converted to thymines. The tiles were grouped in 100 bins by their G or C content and the mean read count per tile, normalised to the total read count per dataset, was plotted against the % G or C.

# Statistics and sample size

Statistical analyses were performed with GraphPad Prism 6.0. Error bars represent standard deviation (s.d.) or standard error of the mean (s.e.m), as described in each figure legend, and indicate whether biological or technical replicates are compared, respectively. Where possible, exact p-values are stated in the figures or figure legends, otherwise star-symbols are used with the corresponding p-value ranges indicated. Where applicable, value distribution was tested with D'Agostino-Pearson normality test and sample variance estimated with Coefficient of Variation (CV). All LC/MS analyses were performed on two separate BS conversion experiments for each method, with 3 replicate samples each (i.e. 6 in total), and a minimum of three LC/MS measurements per sample. For WGBS,

we performed our analyses on a panel of datasets either sequenced for this study (2 to 3 replicates each) or publically available, generated for different studies or by different laboratories (see details in Additional file 1). Where possible, information on number of samples is provided in the figures or figure legends.

#### List of abbreviations

mC: methylcytosine; hmC: hydroxymethylcytosine; BS: bisulfite; WGBS: whole genome bisulfite sequencing; RRBS: reduced representation bisulfite sequencing; DNA: deoxy-ribonucleic acid; NGS: next generation sequencing; PCR: polymerase chain reaction; PBAT: post-bisulfite adaptor-tagging; C: cytosine; LC/MS: liquid chromatography with mass-spectrometry; mESC: mouse embryonic stem cells; TMAC: tetramethylammonium chloride; SNP: single n ucleotide polymorphism; indel: insertion and deletion-caused mutations; CV: coefficient of variation; s.d.: standard deviation; s.e.m.: standard error of the mean; TKO: triple-Dnmt knock-out cell line; CGI: CG-island; (i)DMR: (imprinted) differentially methylated region; QC: quality control; WGA: whole genome amplification.

#### **Declarations**

## Ethics approval and consent to participate

Not applicable

## **Consent for publication**

Not applicable

# Availability of data and materials

All datasets generated in this study are listed in Additional file 1 and deposited in NCBI's GEO under the accession number <u>GSE77961</u>.

All Perl and R scripts used, as well as wiggle tracks are deposited in GitHub: <a href="https://github.com/NellyOlova/BS\_bias">https://github.com/NellyOlova/BS\_bias</a>.

Modules 'bam2nuc' and 'filter\_non\_conversion' have been added to the Bismark v0.16.0 and v0.17.0 packages respectively, <a href="https://github.com/FelixKrueger/Bismark">https://github.com/FelixKrueger/Bismark</a>.

## **Competing interests**

The authors declare no competing financial interests. W.R. is a consultant and shareholder at Cambridge Epigenetix Ltd.

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## **Author's contributions**

N.O. and M.R.B. conceived the study and designed experiments. N.O. performed experiments and data interpretation, N.O., F.K. and S.A. conceived and performed the bioinformatic analyses, D.O.

performed mass spectrometry. N.O., M.R.B. and W.R. wrote the manuscript. All authors read and approved the final manuscript.

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#### **Additional files**

Additional file 1: an Excel spreadsheet with three tabs listing 1) datasets generated in this study, 2) datasets used from the publically domain, together with key parameters, and 3) number of datasets used per analysis.

Additional file 2: a PDF file with supplementary Tables S1-3 and supplementary Figures S1-9.

Additional file 3: an Excel spreadsheet with reference genome compositions.

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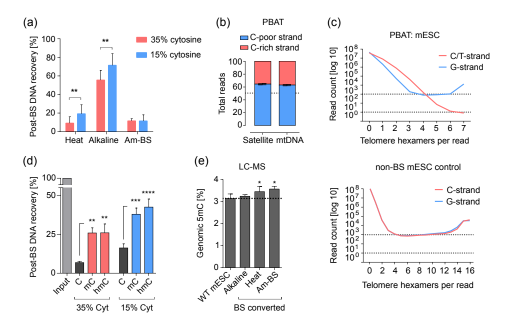


Figure 1. Biased degradation of unmethylated C-rich DNA after bisulfite treatment. (a) Postbisulfite recovery of C-rich and C-poor DNA fragments treated with different BS conversion protocols. Fragment sequences originate from the M13 phage sequence (Additional file 2:Table S1). Statistical analysis was performed with a two-way ANOVA, p = 0.0034 for cytosine content ('Heat' and 'Alkaline' only) and p < 0.0001 for the method. (b) Asymmetric C-rich (23-24% C) and C-poor (12-14% C) strand representation in the mouse major satellite repeat and mtDNA in amplification-free PBAT datasets. The total read count per strand is represented as a proportion of 100%.(c) Telomere repeat count per read in amplification-free PBAT and non-BS converted NGS control. To assess the fragmentation rate, reads containing G-strand tandems [TTAGGG] were quantitated separately from C-strand tandems [CCCTAA] n. In PBAT the C-strand is converted to a 'T-strand', [TTTTAA] n. (d) Post-bisulfite recovery of unmethylated, fully methylated and hydroxymehtylated C-rich and C-poor DNA fragments treated with different BS conversion protocols. One-way ANOVA was performed with Tukey's multiple comparisons test. (e) LC/MS measurement of total genomic 5mC levels in mouse ESC gDNA before ('WT mESC') and after treatment with different BS-conversion protocols ('Alkaline', 'Heat', 'Am-BS'). 5mC is represented as a percentage of all cytosines; the converted cytosines were measured as uracils in the BS converted samples. Individual two-tailed paired t-tests were performed within matched sample-control pairs. Details on number of WGBS datasets used for each analysis are presented in Additional File 1. \*p<0.05, \*\*p<0.01; error bars represent s.d.

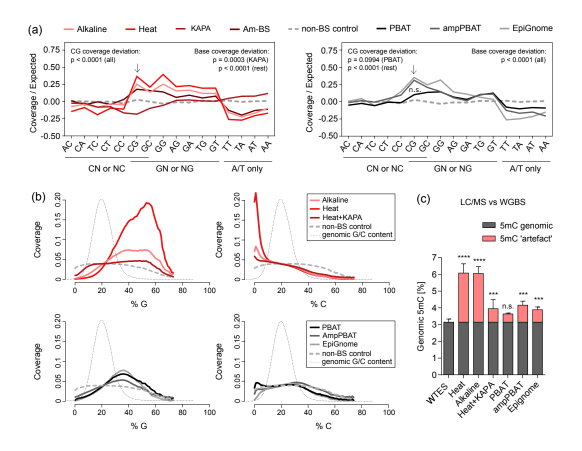


Figure 2. Effect of PCR and polymerase bias on sequence coverage and methylation estimates. (a) Coverage of dinucleotides in WGBS datasets generated with four pre-bisulfite (left panel) and three post-bisulfite (right panel) library preparation protocols. The coverage is expressed as log2 difference from the genomic expected and compared to a non-BS-treated control. For clarity, the dinucleotides are underlined as derived from C, G, or A/T only. Statistical analysis of overall dinucleotide (base) coverage was performed on the average absolute deviations from control with one sample two-tailed t-tests followed by Bonferroni correction. Statistical analysis on CG coverage was performed with one-way ANOVA with Bonferroni correction. The number of datasets used is detailed in Additional File 1 and Additional file 2: Fig. S3a. (b) Read coverage dependence on the G/C composition in 100 bp tiles. Cytosine content of reads was calculated from the corresponding genomic sequence and not the actual read sequence, where unmethylated cytosines appear as thymines. The tile distribution per G/C content is plotted in the background along the x-axis for reference. 'Am-BS' was omitted from this analysis due to unavailability of same species datasets. (c) Global methylation levels of mouse ES cells as measured by LC/MS and a panel of WGBS datasets. The LC/MS value is an average for J1 and E14 lines from different passages and studies to account for lineage and tissue culturing variances. The differences between the LC/MS values and WGBS measurements are marked as a methylation 'artefact' within each WGBS method. Significance is calculated by one-way ANOVA with Dunnett's multiple comparisons test on the absolute WGBS values ('genomic' + 'artefact') against the LC/MS value; \*\*\*\*p < 0.0001, error bars represent s.d.

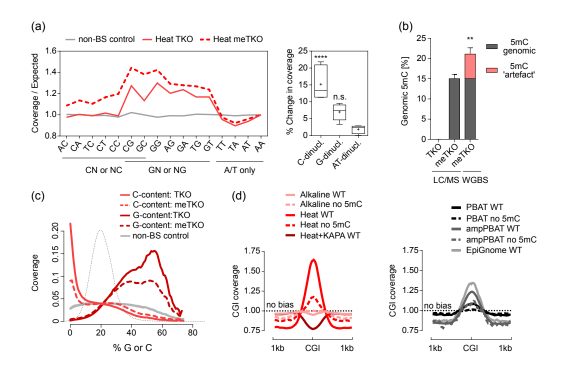


Figure 3. Effect of DNA methylation status on the degradation and amplification biases. (a) Coverage of dinucleotides in WGBS datasets from unmethylated and in vitro M.CviPI-methylated TKO DNA prepared with the 'Heat' BS-seq protocol. For direct comparison, the increase in coverage is expressed as fold difference from the genomic average and normalised to the AA dinucleotide. The dinucleotides are grouped as derived from C, G, or A/T only and presented in the box-plot panel (right) as total % increase in coverage; crosses mark mean values and error bars represent minimum and maximum values. Statistical analysis was performed with one-way ANOVA with Dunnett's multiple comparisons test against the AT-only dinucleotides. (b) Global methylation levels of the in vitro M.CviPI -methylated TKO DNA. The difference between the LC/MS and WGBS values is marked as a methylation 'artefact' as in Fig. 2. Significance between the two values was assessed with a twotailed unpaired t-test, p=0.001, error bars represent s.d. (c) Read coverage dependence on the G/C composition before and after in vitro M.CviPI methylation. Cytosine content of reads was calculated over 100 bp tiles and matched to the corresponding genomic sequence and not the actual read sequence, where unmethylated cytosines are converted to thymines. Dotted black line represents the tile count distribution in G/C content. (d) Coverage of CGIs in WGBS datasets of mouse WT ESCs (i.e. with similar level of methylation) compared to unmethylated DNA generated with the same library preparation protocols. 'Heat+KAPA' and 'Epi Gnome' protocols are included only as WT values for reference, due to unavailability of corresponding unmethylated DNA datasets. Values are expressed as fold-difference from a coverage 'no bias' line.

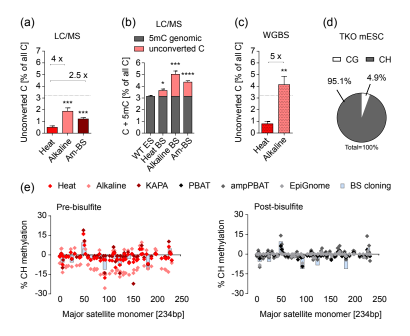


Figure 4. Effect of conversion artefacts on the biases in WGBS. (a) Presence of unconverted cytosines as percent of total cytosine content, measured by LC/MS for three different BS-conversion protocols. The three protocols differ by denaturation method ('Heat' or 'Alkaline') or molarity of bisulfite (4.5 M vs 9 M for 'Am-BS') but not by BS incubation temperature (65-70°C). Averaged folddifferences in quantity are shown above horizontal brackets, and a dotted line shows the usual level of genomic 5mC for reference of scale. For conversion differences between methods with 50°C and 65°C incubation temperatures, see Additional file: Fig. S10a. (b) A theoretical sum of 5mC and unconverted C as measured by LC/MS for J1 WT mESCs for three BS conversion protocols. Both 5mC and unconverted C will be interpreted as 5mC after amplification of WGBS libraries, boosting the overall levels of methylation, depending on the BS treatment protocol. (c) Absolute quantitation of unconverted cytosines in the unmethylated TKO mESC line, as measured by 'Heat' and 'Alkaline' BS seq. (d) Context distribution of BS conversion artefacts; the value is the same for 'Heat' and 'Alkaline' and therefore plotted as an average. (e) CH methylation on both strands of the mouse major satellite repeat as measured by pre- and post-bisulfite WGBS methods. 5mC percentage from the BS cloning from Additional File: Fig. S5a is plotted in both panels for reference. Positive y-axis values indicate the top strand and negative - the bottom strand. Statistical analyses in (a), (b) and (c) were performed for matched experimental pairs with unpaired two-tailed t-tests against 'Heat' in (a) and (c), and 'WT ES' in (b). Error bars in (a), (b) and (c) represent s.e.m.,\*p < 0.05, \*\*p<0.01, \*\*\*p<0.001, \*\*\*p<0.0001.

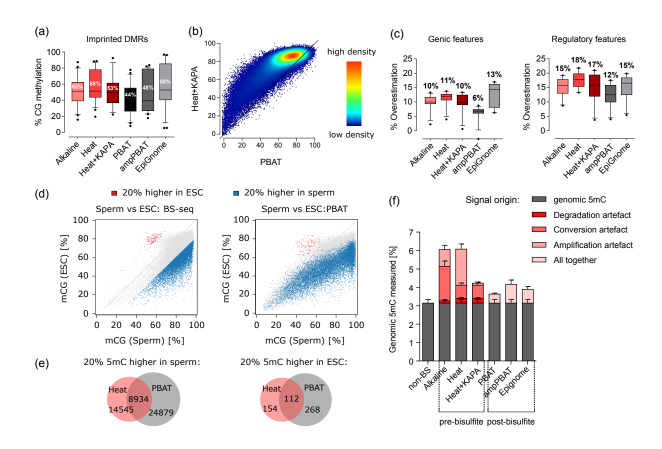
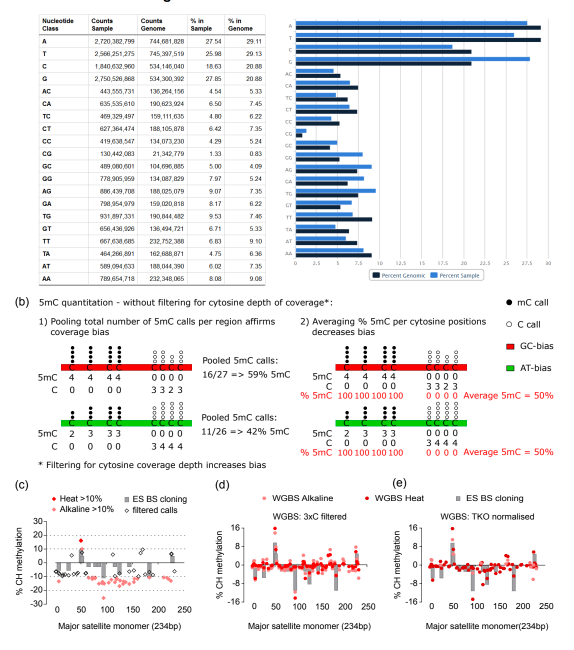


Figure 5. Effect of biases and artefacts on the output of 5mC quantitation. (a) Average methylation values of iDMRs. Numbers indicate the mean value, error bars span the 10-90 percentile. (b) Genome-wide comparison of absolute methylation levels for the amplification-free PBAT approach and the 'Heat+KAPA' BS-seq. (c) Differences in the absolute quantitation of genomic and regulatory features between the amplification-free PBAT (at value 0) and amplified WGBS datasets. Numbers indicate the mean value, error bars show the 10-90 percentile. (d) Comparison of relative methylation differences between sperm and ESC sequenced with either amplification-free PBAT or 'Heat' BS-seq. Each dot represents a probe over 150 consecutive cytosines from the same genomic region in ESC and sperm. The plotted over 20% mCG differences are generated from the BS-seg method (left panel) and visualised with same colour onto the PBAT data (right panel). (e) Venn diagrams showing how many of the over 20% mCG regions from (d) overlap between BS-seq and PBAT. (f) A breakdown of relative contribution of biases for the BS-seq protocols as measured by LC/MS and WGBS. For post-bisulfite protocols, the overall combined bias is shown, sinc4 individual contributions are less trivial to dissect. The non-BS 5mC measurement averages LC/MS measurements for mES from different studies and passages to account for culturing and lineage variances. Significance is lines calculated by one-way ANOVA with Dunnett's multiple comparisons test on the absolute WGBS values ('genomic' + 'artefact') against the LC/MS value; \*\*\*\*p < 0.0001, error bars represent s.d.

## (a) Nucleotide Coverage



**Figure 6. Dealing with biases and artefacts**. (a) A screenshot from the Bismark 'bam2nuc' module output, showing base and dinucleotide content in a 'Heat' dataset against the genomic expected value. The C base indicates the extent of degradation-caused bias (negative correlation), the G-base and its derivative dinucleotides as well as the A/T bases and dinucleotides show the extent and direction of amplification bias – positive or negative, G(C)- or AT-biasing. (b) Comparison of two methylation quantitation strategies to overcome or decrease the effects of GC- or AT-biasing protocols. Counting a total number of methylation calls within a probe (region), irrespective of their position and depth, is compared to calculating methylation values of individual CGs and averaging those for the whole probe. None of those approaches applies initial coverage depth filtering, which is shown in Additional File 2: Figure S9. (c, d, e) Non-CG methylation in the major satellite after removing conversion noise by (c) setting a 10% or 20% 5mC cut-off threshold value, (d) after a bioinformatic filter was applied to remove every read with three or more methylated CH cytosines, and (e) after subtracting the background values from an unmethylated genome control (TKO mESC). Positive y-axis values indicate the top strand and negative – the bottom strand.