Ultra-low-input, tagmentation-based whole-genome bisulfite sequencing

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We have adapted transposase-based in vitro shotgun library construction ("tagmentation") for whole-genome bisulfite sequencing. This method, Tn5mC-seq, enables a >100-fold reduction in starting material relative to conventional protocols, such that we generate highly complex bisulfite sequencing libraries from as little as 10 ng of input DNA, and ample useful sequences from 1 ng of input DNA. We demonstrate Tn5mC-seq by sequencing the methylome of a human lymphoblastoid cell line to ~8.6× high-quality coverage of each strand.

[Supplemental material is available for this article.]

DNA methylation is a widespread epigenetic modification that plays a pivotal role in the regulation of the genomes of diverse organisms. The most prevalent and widely studied form of DNA methylation in mammalian genomes occurs at the five carbon position of cytosine residues, usually in the context of the CpG dinucleotide. Microarrays, and more recently massively parallel sequencing, have enabled the interrogation of cytosine methylation (5mC) on a genome-wide scale (Zilberman and Henikoff 2007). However, the in vivo study of DNA methylation and other epigenetic marks, e.g., in specific cell types or anatomical structures, is sharply limited by the relatively high amount of input material required for contemporary protocols.

Methods for genome-scale interrogation of methylation patterns include several that are preceded by the enrichment of defined subsets of the genome (Meissner et al. 2005; Down et al. 2008; Deng et al. 2009), e.g., reduced representation bisulfite sequencing (RRBS) (Meissner et al. 2005) and anti-methylcytosine DNA immunoprecipitation followed by sequencing (MeDIP-seq) (Down et al. 2008). An advantage of such methods is that they can be performed with limited quantities of starting DNA (Gu et al. 2011). However, they are constrained in that they are not truly comprehensive. For example, the digestion-based RRBS method interrogates only ~12% of CpGs, primarily in CpG islands (Harris et al. 2010), with poor coverage of methylation in gene bodies (Ball et al. 2009) and elsewhere. Furthermore, RRBS does not target cytosines in the CHG or CHH (H = A,C,T) contexts, which have been shown to be methylated at elevated levels in the early stages of mammalian development (Lister et al. 2009). While a small proportion of non-CpG methylation sites can be observed using RRBS, they are restricted to regions within or highly proximal to CpG islands (Ziller et al. 2011).

The most comprehensive, highest resolution method for detecting 5mC is whole-genome bisulfite sequencing (WGBS) (Cokus et al. 2008; Lister et al. 2009; Harris et al. 2010). Treatment of genomic DNA with sodium bisulfite chemically deaminates cytosines much more rapidly than 5mC, preferentially converting them to uracils (Clark et al. 1994). With massively parallel sequencing, these can be detected on a genome-wide scale at single-base-pair resolution. This approach has revealed complex and unexpected methylation patterns and variation, particularly in the CHG and CHH contexts. Furthermore, as the costs of massively parallel sequencing continue to plummet, WGBS is increasingly affordable. However, a key limitation of WGBS is that the current protocols for library construction are based on ligation chemistry and call for 5 μg of genomic DNA as input (Cokus et al. 2008; Lister et al. 2009; Li et al. 2010) which is essentially prohibitive for many samples obtained in vivo.

We recently characterized a transposase-based in vitro shotgun library construction method ("tagmentation") that allows for construction of sequencing libraries from greatly reduced amounts of DNA (Fig. 1A; Adey et al. 2010). Briefly, the method utilizes a hyperactive derivative of the Tn5 transposase loaded with discontinuous synthetic oligonucleotides to simultaneously fragment and append adaptors to genomic DNA. The resulting products are subjected to PCR amplification followed by high-throughput sequencing. The increased efficiency of genomic DNA conversion to viable amplicons and the greatly reduced number of steps allow the construction of low-bias, highly complex libraries from <50 ng of genomic DNA.

Here we describe a modified approach, which we call Tn5mC-seq, that retains the advantages of transposase-based library preparation in the context of WGBS. Because the target of the transposition reaction is double-stranded DNA, whereas bisulfite treatment yields single-stranded DNA, the method was extensively modified such that the tagmentation reaction could take place prior to bisulfite treatment (Fig. 1B). First, the adaptors to be incorporated were methylated at all cytosine residues to maintain cytosine identity during bisulfite treatment, with the exception of the 19-bp transposase recognition sequence (in order to minimize differential binding during transposome assembly). Second, an oligonucleotide replacement scheme (Supplemental Fig. S1B; Grunenwald et al. 2011; Gertz et al. 2012) was utilized to ensure that each strand would have adaptors covalently attached to both ends of the molecule. Specifically, this entails initial transposition with a single adaptor in which the double-stranded transposase recognition sequence is truncated to 16 bp (Tm = 36°C), thereby facilitating its post-incorporation removal by denaturation. A second adaptor is then annealed and the gap repaired, resulting in each strand being covalently linked by both a 3’ and 5’ adaptor. The fragmented, adapted, double-stranded genomic DNA is then subjected to standard bisulfite treatment for the conversion of unmethylated cytosine to uracil. Degradation during the conversion process likely remains a primary source of loss, but the in-
increased efficiency of the prior steps and the lack of gel-based size selection result in an overall increase in the fraction of DNA that is converted, PCR-amplified, and sequenced.

Results

Ultra-low-input transposase-based WGBS library performance

We applied Tn5mC-seq to sequence the methylome of a lymphoblastoid cell line (GM20847) using libraries constructed from 1–200 ng of input genomic DNA. Each library was barcoded during PCR amplification and subjected to either a spike-in (5%) or majority (80%–90%) of a lane of sequencing on an Illumina HiSeq2000 (paired-end 100 bp [PE100]; v2 chemistry with custom sequencing primers). These data are summarized in Table 1 and Supplemental Figure S2. In addition, several PCR conditions were investigated to optimize amplification uniformity (Supplemental Fig. S3), as well as a modified protocol (Tn5mC-seq 1.1) (Supplemental Figs. S1D, S4) that eliminates the need for custom sequencing primers and may increase library construction efficiency. Reads were aligned to an in silico converted hg19 (GRCh37) to both the top (C→T) and bottom (G→A) strands using BWA (Li and Durbin 2009) followed by read trimming of unmapped reads and secondary alignment using the same parameters. Unaligned reads typically consisted of low-quality artifacts that likely arose during amplification due to the reduced base complexity of bisulfite converted amplicons.

For each library constructed using ≥10 ng of genomic DNA, over 100 million aligned reads were obtained (60%–75% of total filtered reads; see Methods) of high complexity (90%–97% nonduplicates). Despite the significantly reduced performance of libraries prepared from 1 ng, ~12 million reads were still aligned and the library was of reasonable complexity (78% nonduplicates). Post-alignment reads were merged and quality filtered for a total of 51.7 Gb of aligned, unique sequence. The average read depth was 8.6× per strand with >96% of CpG and >98% of non-CpG cytosines covered genome-wide (Fig. 1C; Supplemental Fig. S2). Because unmethylated nucleotides are incorporated during the gap-repair step (first 9 bp of the second read and last 9 bp before the adaptor as determined by insert size on the first read), the gap-repair regions must be excluded from methylation analysis. However, these bases also serve as an internal control for the conversion rate of the bisulfite treatment. We found this to be >99% for all libraries, and this was independently confirmed using unmethylated lambda DNA spike-ins to two libraries.

For comparison, ligation chemistry–based libraries were constructed using 1000, 100, and 10 ng of GM20847 DNA of the same isolation as the batch used for Tn5mC-seq. These libraries were prepared following the protocols outlined by Lister et al. (2009) with the exception of PCR, which was performed using Kapa Robust due to its higher efficiency over other polymerase choices (Supplemental Fig. S3). During amplification, the 100 and 10 ng preparations did not show significant amplification above a negative control background and were not carried through to sequencing, precluding a comparison of Tn5mC-seq and ligation chemistry–based library construction with identical inputs (a 1000 ng Tn5mC-seq preparation was also not feasible due to the dilute concentration of the commercially available transposase, which would result in a reduced density of transposition events on a high input mass).

Post-alignment, the 1000 ng ligation chemistry–based library consisted of low-quality artifacts that likely arose during amplification due to the reduced base complexity of bisulfite converted amplicons.
We were able to detect approximately 46 million 5mC positions (1% FDR; see Supplemental Fig. S5B, C). We also compared the methylation levels of CpGs well-covered by sequence contexts (Supplemental Fig. S5A). Comparable uniformity was also observed with respect to G+C content as well as for tetramer/pentamer sequence contexts (Supplemental Fig. S5B, C). We also compared the methylation levels of CpGs well-covered by sequencing of libraries corresponding to both methods, and observed good agreement at positions with 5 × or greater coverage ($r^2 = 0.55$) as well as 10 × or greater coverage ($r^2 = 0.82$) (Supplemental Fig. S5D).

**Table 1. Summary of TnSmC-seq libraries and sequencing**

| Name        | Input DNA (ng) | Percentage aligning (%) | Percentage unique (%) | Unique aligned reads | Mean insert size (bp) |
|-------------|----------------|--------------------------|-----------------------|---------------------|-----------------------|
| TnSmC-C     | 200            | 68                       | 93                    | 127,098,152         | 198                   |
| TnSmC-D     | 50             | 75                       | 90                    | 133,382,834         | 254                   |
| TnSmC-E     | 1              | 12                       | 76                    | 11,181,960          | 134                   |
| TnSmC-F     | 10             | 65                       | 95                    | 118,170,302         | 168                   |
| TnSmC-G     | 50             | 61                       | 97                    | 87,294,793          | 180                   |
| TnSmC-H     | 1              | 11                       | 78                    | 12,393,357          | 126                   |
| TnSmC-I     | 10             | 62                       | n/a                   | 29,546,077          | n/a                   |
| TnSmC-J     | 50             | 71                       | 95                    | 132,144,644         | 196                   |
| TOTAL       |                |                          |                       | 651,213,119         |                       |

Raw reads were initially filtered for instrument valve failures at specific locations of reads and then removal of reads containing over three Ns or extremely low-quality bases (phred score ≤ 2) in the first 50 bases. Alignment was then performed using BWA (Li and Durbin 2009) to in silico converted top and bottom strand references of hg19 (GRCh37) followed by trimming and realignment. Duplicate reads were identified and removed according to their start position and insert size. The percentage of post-filtering reads that align for each library is shown, as is the percentage of these that are nonduplicates.

* Valve failures in read 2 resulted in extensive read trimming (50–70 bp).
* Complete valve failure on read 2.

(Supplemental Fig. S1D) libraries constructed from 10 ng, particularly at the lower CpG densities that represent the majority of the genome (Supplemental Fig. SSA). Comparable uniformity was also observed with respect to G+C content as well as for tetramer/pentamer sequence contexts (Supplemental Fig. SSB, C). We also compared the methylation levels of CpGs well-covered by sequencing of libraries corresponding to both methods, and observed good agreement at positions with 5 × or greater coverage ($r^2 = 0.55$) as well as 10 × or greater coverage ($r^2 = 0.82$) (Supplemental Fig. SSD).

**Lymphoblastoid cell line methylation**

We were able to detect ~46 million SmC positions (1% FDR; see Methods), accounting for 4.2% of total cytosines with 5mC. The majority of methylation observed was in the CpG context (97.1%), and the global CpG methylation level was 69.1%. This level is similar to that of the fetal fibroblast cell line IMR90 sequenced by Lister et al. (2009; 67.7%) and is consistent with the observation that CpG methylation levels are reduced in differentiated cell types. Additionally, CHG and CHH methylation levels were substantially lower than in ES cells, at 0.36% and 0.37%, respectively, again consistent with the differentiated cell type. On the chromosome scale, the methylation density correlated with banding patterns and increasing levels were observed extending distally through subtelomeric regions (Fig. 1D). An analysis of functionally annotated genic regions revealed a sharp decrease in CpG methylation through the promoter region followed by a minor increase in the 5′ UTR and then elevated levels of methylation throughout the gene body, particularly at introns (Fig. 1E,F), consistent with previously described CpG methylation profiles (Lister et al. 2009).

**Discussion**

We developed TnSmC-seq as a novel method for rapidly preparing complex, shotgun bisulfite sequencing libraries for WGBS. In brief, the method utilizes a hyperactive Tn5 transposase derivative to fragment genomic DNA and append adaptors in a single step, as previously characterized for the construction of DNA-seq libraries (Adey et al. 2010). In order for library molecules to withstand bisulfite treatment, the adaptors are methylated at all cytosine residues, and an oligonucleotide replacement strategy is employed to make each single-strand covalently flanked by adaptors. The high efficiency of the transposase and overall reduction in loss-associated steps permits construction of high-quality bisulfite sequencing libraries from as little as 10 ng that are comparable to ligation chemistry–based libraries generated from 100× more DNA, as well as useful sequence from 1 ng of input DNA. Additionally, the increased efficiency of transposase-mediated library construction may allow for preparation of WGBS libraries from poor-quality or degraded DNA samples.

Our results illustrate how derivatives of the transposase-based method for DNA-seq library preparation can enable key applications of next-generation sequencing where its advantages are perhaps even more relevant. The ability to generate such libraries from very low amounts of input genomic DNA substantially improves the practicality of whole methylome sequencing and removes a key advantage of less encompassing methods such as RRBS (Meissner et al. 2005; Harris et al. 2010). Specifically, low-input WGBS with TnSmC-seq may make possible the comprehensive interrogation of methylation in many contexts where DNA quantity is a bottleneck, e.g., developing anatomical structures, microdissected tissues, or pathologies such as cancer, where the epigenetic landscape is of interest but tissue quantity limits high-resolution WGBS.

**Methods**

**TnSmC-seq library construction and sequencing**

Transposome complexes were generated by incubating 2.5 μL of 10 μM TnSmC-A1 (TnSmC-A1top: 5'-GAT [5mC] TA [5mC] A[5mC] G [5mC] [5mC] T [5mC] [5mC] [5mC] T [5mC] G [5mC] AGAGATGTGTATAAGAGACAG-3'), 10 μM TnSmC-A1bot: 5'-[Phos]-CTGTCTCTTTACATAAC-3', IDT, annealed to TnSmC-A1bot: 5'-[Phos]-CTGTCTCTTTACATAAC-3', IDT, by incubating 10 μL of each oligo at 100 μM and 80 μL of EB [QIAGen] for 2 min at 95°C and then cooling to room temperature at 0.1°C/sec with 2.5× 100% glycerol and 5 μL EZ-Tn5 transposase (Epicentre – Illumina) for 20 min at room temperature.

Genomic DNA prepared from GM20847 cell lines was used at respective input quantities with 4 μL Nextera HMW Buffer (Epicentre-Illumina), nuclease-free water (Ambion) to 17.5 μL and 2.5 μL prepared TnSmC transposomes (regardless of the quantity of DNA used). Reactions were incubated for 9 min at 35°C in a thermocycler followed by SPRI bead cleanup (AMPure) using 36 μL of beads and the recommended protocol with elution in 14 μL nuclease-free water (Ambion). Adapter 2 annealing was then carried out by adding 2 μL of 10× Amplification Reaction Buffer (Epicentre-Illumina), 2 μL 10 mM dNTPs (2.5 mM each; Invitrogen), and 2 μL 10 μM TnSmC-A2top (5'-5Phos/-SPhos/CGTCTCTTTACATAACAT [5mC] TGGG [5mC] GGG [5mC] TGG [5mC] AAGG [5mC] AGA [5mC] [5mC] GAT [5mC]-3'); IDT to each reaction and incubating for 50 min at 95°C and then cooling at 0.1°C/sec to 37°C and subsequent incubation for 10 min. Gap-repair was then performed by adding 3 μL of Amplify at 5U/μL (Epicentre-Illumina) and 1 μL of either T4 DNA Polymerase (TnSmC libraries A-G, NEB) or Sulfobolus DNA Polymerase IV (TnSmC libraries H-J, NEB) and additional incubation for 30 min at 37°C. Reactions were then cleaned up using SPRI beads (AMPure) according to recommended protocol using 36 μL beads and elution in 50 μL nuclease-free water (Ambion). Bisulfite treatment was performed using an EZ.
DNA Methylation Kit (Zymo) according to recommended protocols with a 14-h 50°C incubation and 10 μl elution. Elute was then used as the template for PCR using 12.5 μl Kapa 2G Robust HotStart ReadyMix (Kapa Biosystems), 1 μl 10 μM Ts5Mc-C1 (5'-AATGATACGGGCGACCCGAGATCTACACGCTCCCTCCGCTCGGCGCAGTACAGCAAGG-3'; IDT); index read, Ts5Mc-Cx (5'-TTGTGTITTTTATATA TATTTCTGAGCGGGCTGGCAAGGC-3'; IDT); and read 2, Ts5Mc-C2 (5'-GCTCTGGCCACCGCCGCTGAAATATAAAAAACAA-3'; IDT). Read lengths were either single-read at 36 bp with a 9-bp index (PE101, libraries C–J). Libraries were only read; 10 sec at 72° C; 40 sec at 72°C; Plate Read; 10 sec at 72°C) x 99. Reactions were monitored and removed from thermocycler as soon as plateau was reached (12–15 cycles).

The hg19 reference genome was first bisulfite-converted in silico for both the top (C changed to T, C2T) and bottom (G changed to A, G2A) strands. Prior to alignment, reads were filtered based on the run metrics, as several libraries were run on lanes in which instrument valve failures resulted in poor quality or reads consisting primarily of N bases. Filtering was carried out by first calculating the base compositions as well as mean base quality scores at each position in the read. The mean of the lanes had significantly reduced quality scores at the start and/or end of the read and were globally trimmed to remove any start or end positions that had a mean phred score of less than or equal to 10. The start and ends of the reads were additionally globally trimmed if a position within the first or last 25 bases of the read had a mean composition of 10% Ns, which generally corresponded to the quality-based trimming. Additionally, reads that contained three or more Ns were also removed. It is important to note that the reduced qualities in the runs were “flowcell-wide” regardless of the library that was run and not isolated to Ts5Mc-seq libraries. Subsequent runs for the Ts5Mc-seq 1.1 and polymerase testing experiments did not suffer instrument failures, and no trimming of the reads was necessary. Next, reads were aligned to both the C2T and G2A strands using BWA with default parameters. Reads that aligned to both strands were removed. Read pairs in which neither aligned to either strand were then pulled and trimmed to 76 bp (except for SE36 runs) and again aligned to both C2T and G2A strands. Duplicate reads (pairs sharing the same start positions for both reads 1 and 2) were removed and complexity determined. Reads with an alignment score less than 10 were then filtered out prior to secondary analysis. Total fold coverage was calculated using the total bases aligned from unique reads over the total alignable bases of the genome (slightly below 3 Gb per strand).

5MeC Calling

Methylated cytosines were called using a binomial distribution as in the method described by Lister et al. (2009), whereby a probability mass function is calculated for each methylation context (CpG, CHG, CHH) using the number of reads covering the position as the number of trials and reads maintaining cytosine status as successes with a probability of success based on the total error rates that were determined by the combined nonconversion rate and sequencing error rate. The total error rate was initially determined by unmethylated lambda DNA spike-ins; however, we found that the error rate estimation from the gap-repair portion of reads (as described in the main text) gave a more comprehensive estimate, which was slightly higher than that of the lambda estimate. Therefore to be conservative, we used the highest determined error rate at 0.009. If the probability was below the value of M, where M × (number of total unmethylated CpG) < 0.01 × (number of total methylated CpG), the position was called as being methylated, thus enforcing that no more than 1% of positions would be due to the error rate.

Data access

The sequence data presented in this study have been submitted to the NCBI Sequence Read Archive (SRA) (http://www.ncbi.nlm.nih.gov/sra) under accession no. SRP011746.

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Ligation chemistry WGBS library construction and sequencing

We subjected 1000, 100, and 10 ng of genomic DNA prepared from GM18507 cell lines to ligation chemistry–based library preparation according to methods described by Lister et al. (2009) with several minor exceptions: (1) Bisulfite conversion was carried out using an EZ DNA Methylation Kit (Zymo), and (2) PCR was carried out using Kapa 2G Robust Hot Start ReadyMix (Kapa Biosystems). The change in PCR enzyme was due to several unpublished experiments demonstrating a much higher efficiency with Kapa Robust as opposed to PfuTurbo Cx used according to the method described by Lister et al. (2009). Sequencing was performed on an Illumina MiSeq instrument using a single-end 100-bp sequence read run.

Read filtering and alignment

The hg19 reference genome was first bisulfite-converted in silico for both the top (C changed to T, C2T) and bottom (G changed to A, G2A) strands. Prior to alignment, reads were filtered based on the run metrics, as several libraries were run on lanes in which instrument valve failures resulted in poor quality or reads consisting primarily of N bases. Filtering was carried out by first calculating the base compositions as well as mean base quality scores at each position in the read. The mean of the lanes had significantly reduced quality scores at the start and/or end of the read and were globally trimmed to remove any start or end positions that had a mean phred score of less than or equal to 10. The start and ends of the reads were additionally globally trimmed if a position within the first or last 25 bases of the read had a mean composition of 10% Ns, which generally corresponded to the quality-based trimming. Additionally, reads that contained three or more Ns were also removed. It is important to note that the reduced qualities in the runs were “flowcell-wide” regardless of the library that was run and not isolated to Ts5Mc-seq libraries. Subsequent runs for the Ts5Mc-seq 1.1 and polymerase testing experiments did not suffer instrument failures, and no trimming of the reads was necessary. Next, reads were aligned to both the C2T and G2A strands using BWA with default parameters. Reads that aligned to both strands were removed. Read pairs in which neither aligned to either strand were then pulled and trimmed to 76 bp (except for SE36 runs) and again aligned to both C2T and G2A strands. Duplicate reads (pairs sharing the same start positions for both reads 1 and 2) were removed and complexity determined. Reads with an alignment score less than 10 were then filtered out prior to secondary analysis. Total fold coverage was calculated using the total bases aligned from unique reads over the total alignable bases of the genome (slightly below 3 Gb per strand).

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