Differential DNA methylation in discrete developmental stages of the parasitic nematode *Trichinella spiralis*

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Abstract

**Background:** DNA methylation plays an essential role in regulating gene expression under a variety of conditions and it has therefore been hypothesized to underlie the transitions between life cycle stages in parasitic nematodes. So far, however, 5'-cytosine methylation has not been detected during any developmental stage of the nematode *Caenorhabditis elegans*. Given the new availability of high-resolution methylation detection methods, an investigation of life cycle methylation in a parasitic nematode can now be carried out.

**Results:** Here, using MethylC-seq, we present the first study to confirm the existence of DNA methylation in the parasitic nematode *Trichinella spiralis*, and we characterize the methylomes of the three life-cycle stages of this food-borne infectious human pathogen. We observe a drastic increase in DNA methylation during the transition from the new born to mature stage, and we further identify parasitism-related genes that show changes in DNA methylation status between life cycle stages.

**Conclusions:** Our data contribute to the understanding of the developmental changes that occur in an important human parasite, and raises the possibility that targeting DNA methylation processes may be a useful strategy in developing therapeutics to impede infection. In addition, our conclusion that DNA methylation is a mechanism for life cycle transition in *T. spiralis* prompts the question of whether this may also be the case in any other metazoans. Finally, our work constitutes the first report, to our knowledge, of DNA methylation in a nematode, prompting a re-evaluation of phyla in which this epigenetic mark was thought to be absent.

**Background**

Developmental regulation of gene expression plays a crucial role in the transitions between significantly differentiated life-history stages, such as is the case in parasitic nematodes; however, the underlying mechanisms of this gene regulation are poorly understood. Although DNA methylation has been established in other organisms as an important method for altering chromatin structure and regulating the expression of genes, its contribution to nematode development has not been adequately assessed given that so far no 5’ cytosine methylation has been identified in any stage of *Caenorhabditis elegans* [1]. Most vertebrate cell types have approximately 60 to 90% of the CpG dinucleotides modified to 5-methylcytosine (5mC) [2], whereas invertebrate genomes vary extensively in the extent of DNA methylation, and some genomes have undetectable levels of methylation [3]. Recently, technological progress has enabled high-resolution detection of 5mC, opening the way for more detailed examination of the role of DNA methylation in a greater variety of eukaryotic genomes [4].

Parasitic nematodes are a good example of the biological importance of developmental regulation of genes, including the principal agent of human trichinellosis, *Trichinella*...
spira
cellular stages. This food-borne agent infects a wide
typhicently (ML). ML are released by the
host's gastric juices, after which they grow substantially
and mature into sexually active adults (Ad) in the host's
intestines. New-born larvae (NBL) are released from
mature females and then disseminate through the blood-
stream, invade skeletal muscles, and encyst in a collagen
capsule to form a new generation of ML [5]. The host
muscle cells proliferate as they are transformed into 'nurse
cells' for the parasite [6]. The major clinical symptoms of
trichinellosis (myopathy) derive from inflammation directed
against the encysted ML. Thus, successful nematode
development entails a series of physically and functionally
distinct stages that require accurate recognition of specific
biological cues. In this way, the life cycle of parasitic
nematodes is distinct from that of free-living nematodes,
such as Caenorhabditis elegans, that live in a more homo-
geneous environment.

Stage-specific expression has been observed for genes in
Trichinella spp. [7]. Differential expression was especially
obvious for genes encoding the excretory-secretory (E-S)
proteins released from the larvae. For example, a gene
encoding a 43-kDa glycoprotein is expressed in precapsular
and postcapsular muscle larvae, but not in adults [8]. E-S
proteins may therefore contribute to capsule formation [9].
Stage-specific gene expression may also assist parasitic eva-
sion or forestalling of immune reactions that would inhibit
continued transmission. Thus, how stage-specific tran-
scriptional regulation is accomplished in these organisms might
prove useful for understanding and preventing infection.

Recent innovations in high-throughput sequencing have
enabled researchers to infer methylation patterns at sin-
gle-base resolution [10]. MethylC-seq enables methylation
analyses with unprecedented precision, and the recently
released draft genome sequence of T. spiralis [11] pro-
vided us the means to evaluate the methylome of its three
distinct stages. Our work here describes the first com-
prehensive study that confirms the existence of DNA methyla-
tion in T. spiralis and characterizes the differential
methyloes of the organism during these life stages. We
further identified sets of genes whose DNA methylation
status varied between the developmental stages. Our data
shed light on the developmental biology of an important
food-borne zoonosis, and our approach opens the way for
future assessment of methylation as a mechanism of de-
velopmental regulation in this and other metazoans that
undergo similar life cycle transitions.

Results
The presence of DNA methylation in the T. spiralis
genome
To understand whether T. spiralis possesses the ability
to methylate DNA, we conducted reciprocal Blast searches
to identify genes that might be related to known DNA
(cytosine-5)-methyltransferases. Our data revealed the
existence of several relevant orthologous genes annotated
in the draft T. spiralis genome [11] (Table S1 in Addi-
tional data file 1). We found that EFV54759.1 and
EFV58204.1 were homologous to dmnt3 de novo methyl-
transferases and to the maintenance methyltransferase
dmnt1, respectively, in species that are known to have
DNA methylation, such as human and mouse. Of addi-
tional interest, T. spiralis appeared to be the only nema-
tode, compared to 11 other nematodes, that possessed de
ovo methylation machinery (dmnt3). The other nem-
todes only contained orthologs to maintenance methyl-
transferase dmnt1, including Caenorhabditis elegans. We
also identified an ortholog to dmnt2 (EFV60295.1), but it
was more similar to a previously identified tRNA methyla-
lase [12-14], which suggests the potential existence of
RNA methylation in T. spiralis. We used the sequences of
des dmnt-like proteins to reconstruct a phylogenetic
tree (Figure 1). This analysis indicated that T. spiralis
dmnt3 was not a close relative of orthologs in its host
mammals, suggesting that T. spiralis dmnt3 did not origi-
nate from its host through horizontal gene transfer.
We performed PCR with reverse transcription (RT-PCR)
and found that T. spiralis dmnt2 and dmnt3 genes were
differentially expressed among three life stages, but that
dmnt1 expression remained at about the same level (Fig-
ure S1a in Additional data file 2). Correspondingly, enzy-
matic data using nuclear protein extracts also showed
differential catalytic activity of the T. spiralis dmnts
(Figure S1b in Additional data file 2). We also carried out
ultra-performance liquid chromatography-tandem mass
spectrometry (UPLC-MS/MS), which further confirmed
the existence of DNA methylation in T. spiralis, showing
that the total amount of DNA methylation in the Ad
stage was significantly higher than in the NBL stage (Fig-
ure S2 in Additional data file 2) [15].

Given these results, we assessed the genome-wide
DNA methylation profiles in the three life stages of
T. spiralis (Ad, ML, and NBL) using MethylC-Seq. We
generated 61.65, 23.52 and 55.77 million raw reads,
respectively. We aligned the reads to the T. spiralis
reference sequence [16] and mapped approximately
96.36% of the reads to Ad, 91.30% to ML and 99.27% to
NBL, yielding 2.91, 1.05 and 2.71 Gb of DNA sequence
to Ad, ML, and NBL, respectively. The average read
depth was 21.36, 10.80 and 26.21 per strand, respec-
tively. On average, over 81.6% of each strand of the 64
Mb T. spiralis reference sequence was covered by at
least one sequence read in each of the three stages.
Because of the potential for the occurrence of non-
conversion and thymidine-cytosine sequencing errors,
we estimated the false-positive rate as the percentage of
cytosines sequenced at cytosine reference positions in
the Lambda genome, which are normally unmethylated (Materials and methods). We then applied the error rates for each stage (0.0060, 0.0064 and 0.0025 for Ad, ML, and NBL, respectively) to correct mC identification according to a method described by Lister et al. [10] that is based on a binomial test and false discovery rate constraints. Corrected estimates resulted in approximately 0.31 million and 0.24 million mCs in the Ad and ML genomes (comprising 1.59% and 1.22% of their sequenced cytosines, respectively). In contrast, methylation
was nearly undetectable in NBL (0.002 million; 0.01%; Table S2 in Additional data file 2). We validated the results using two different methods: (1) bisulfite-PCR (BSP), cloning, and conventional sequencing by the Sanger method; and (2) methylated DNA immunoprecipitation (MeDIP) combined with quantitative PCR (QPCR). For BSP, we assessed six randomly selected genomic regions that varied in their estimated amount of methylation, and obtained strong agreement between the two experimental results (P-value < 0.05 using double t-test; Figure S3 in Additional data file 2; Table S3 in Additional data file 1) [15]. For MeDIP with QPCR, we assessed three randomly selected genomic regions and confirmed the existence of DNA methylation in all three regions (Figure S4 in Additional data file 2).

Characterization of overall methylation patterns in the three life stages
We further characterized the global patterns of DNA methylation in the genomes of the different *T. spiralis* stages. The highest amount of detected mCs (82.25% and 89.06%) in Ad and ML were located in CG regions, indicating a dominant role of CpG methylation in these stages. Due to the very low level of DNA methylation in NBL, the distribution of CG and non-CG methylation was very similar to the background (Figure 2a). The average methylation level of specific cytosine residues could be estimated from the fraction of methylated sequence reads at that site. Here we found that the average methylation level of specific cytosine residues was estimated from the fraction of methylated sequence reads at that site (Figure 2c).

Since the mCs in the *T. spiralis* genome are relatively sparse compared to vertebrate genomes, we identified methylation regions (MRs) of the genome using relatively dense mCs (Materials and methods). Different CG and non-CG methylation might be subject to distinct forms of genetic control; therefore, MR identification was performed independently for CG and non-CG contexts. Across the genome, we observed an increase in CG methylation as the parasites matured from the NBL to the ML stage and, to a lesser extent, in the transition from ML to Ad. In addition, CG methylation levels fluctuated drastically across the genome, indicating a mosaic methylation pattern where relatively dense methylated domains are interspersed with regions that are not methylated (Figure 2b). Such a pattern has been observed in previous studies on other invertebrates [3]. In contrast, we identified only a small number of non-CG MRs (Table S4 in Additional data file 1).

In all types of genomic elements, we saw a methylation increase from NBL to ML and from ML to Ad as well as the global pattern. We then examined patterns of methylation in distinct genomic elements, including genes, tandem repeats, and transposable elements. Genes were methylated more frequently than the genome average (Figure 3a, b). Within genes, the coding sequences were more methylated than the flanking DNA or promoter regions, while introns were the least methylated (Figure 3c, d). Notably, repeat elements, including tandem repeats and transposable elements, exhibited much higher DNA methylation than the genome average (Figure 3a, b). Previous studies have indicated that the methylation level of transposons across different phylogenetic units may vary. It has been reported that transposable elements are highly methylated in mammals, plants and zebrafish (*Danio rerio*), and moderately methylated in *Ciona intestinalis*, but are usually unmethylated in the honey bee *Apis mellifera* and silkworm *Bombyx mori* [17,18]. In *T. spiralis*, we observed higher methylation on transposons relative to the immediate flanking regions as well (Figure S5 in Additional file 2), which is similar to what is seen in *Ciona intestinalis* [17].

The relationship between stage-dependant methylation and gene expression
We evaluated differential gene expression among the three life stages using Illumina high-throughput RNA-seq technology. Most of the raw reads (numbering 28,662,704, 26,128,346 and 28,962,820, respectively, for the Ad, ML and NBL stages) could be uniquely mapped to previously annotated genes (62.26%, 64.38%, and 64.34%). We detected 12,675, 12,683 and 12,909 annotated genes out of the total 16,379 with at least one unique read. The majority of these genes (11,636) were expressed in all three life stages, and we saw 234 Ad-stage specific, 183 ML-specific and 445 NBL-specific genes. Of note, we also detected stage-dependent expression of methyltransferases that were concordant with prior RT-PCR results (Figure S2 in Additional data file 2). Finally, among genes that were expressed in more than one stage, we identified differential expression in 1,752 pair-wise comparisons (Table S5 in Additional data file 1).

We characterized the changes in DNA methylation among the three distinct life stage methylomes and the relationship between methylation and differential gene expression. For this, we divided expressed genes with at least one sequencing read into quartiles of expression levels, and examined the expressed genes together with another category composed of genes exhibiting no expression. We found that DNA methylation levels of gene upstream regions had a negative correlation with gene expression levels, and non-expressed genes in particular had different patterns of DNA methylation as the methylation levels in their upstream regulatory regions were higher than in the coding sequences (Figure 4a, b). Based on this, it is likely that methylated promoters induce silencing in *T. spiralis*, akin to the widely accepted
role for hypermethylation of promoters as a means of repressing gene expression in plants and mammals [19,20]. In contrast, the gene-body methylation levels in our analysis showed a bell-shaped, rather than monotonic, relationship with gene expression levels. Generally in the gene body, the non-expressed and most highly expressed genes had the lowest DNA methylation levels, whereas the mid-level expressed genes had the highest percentage of DNA methylation (Figure 4a-c). A bell-shaped relationship between gene-body methylation and expression levels has been observed previously in plants (Arabidopsis thaliana and Oryza sativa), invertebrates (Ciona intestinalis and Nematostella vectensis), and humans as well [4,21,22], indicating conservation of the role of methylation across phylogenetically diverse species.
Biological implications of stage-dependant methylation in *T. spiralis*

We examined the correspondence of methylation status to divergent gene expression in different stages. Due to the overall low level of genome methylation in general, we limited this analysis to MRs exhibiting high levels of methylation where we had at least 5× depth of coverage. Using this criteria, we found a total of 652 ML and 767 Ad MRs enriched for methylation in CG regions, but MRs in non-CG regions were rare. In contrast, we found no MRs in the NBL stage. As shown in Figure 5a, 389 MRs were shared between Ad and ML. These MRs were located in 486 and 551 genes in the ML and Ad stages, respectively, with the majority located in gene-body regions (Table S4 in Additional data file 1). We carried out a Gene Ontology analysis to functionally characterize those genes with CG MRs in Ad and ML using Gostat [23]. Enrichment of GO terms defined by a significant false discovery rate-corrected *P*-value (≤ 0.01) in the 'molecular function' category was indicated in 'DNA integration', 'DNA metabolic process' and so on. In the 'biological process' category, 'nucleic acid metabolic process' and 'endopeptidase activity' and so on were enriched. Of note, we found that many genes were shared among different molecular pathways and constituted a central focus of study in parasitic nematodes (Table S6 in Additional data file 1). Given this, we explored the potential for DNA methylation regulating genes that are related to parasitic activities. For example, the protein EFV53263.1 is encoded by a DNase II gene in the ‘DNA
**Figure 4** Relationship between DNA methylation and expression levels of genes in Ad and ML stages of *T. spiralis* (a, b) Average density (a) or percentage of methylation levels (b) within genes that were classified based on expression levels. The first class includes silent genes with no sequencing read detected, and the second to fifth classes cover expressed genes from the lowest 25% to highest 25%. In the curve graph, 2-kb regions upstream and downstream of each gene are divided into 100-bp intervals, and each gene was divided into 20 intervals (5% per interval). In the histogram graph, overall average (± standard error) methylation levels for genes are indicated.

**Figure 5** Analysis of highly enriched MRs and annotation of the genes containing MRs (a) Venn diagram of shared and stage-specific MRs in different sequence contexts of Ad and ML stages. (b) Venn diagram of shared and disparate genes containing MRs in Ad and ML stages, expressed (red) and silent (blue) genes are separated.
metabolic process’ category that is expressed in a stage-specific manner and important for *T. spiralis* parasitism [24], and we found that this gene was uniquely transcribed in the NBL stage, whereas it was not expressed and had hypermethylated promoter regions in the Ad and ML stages. Considering the globally monotonic and negative correlation between promoter methylation and gene expression, this finding strongly suggested that promoter methylation plays a role in the regulation of stage-specific expression of certain DNase II genes in *T. spiralis* (Figure 6a).

In addition to a role for promoter methylation in expression, recent studies have suggested that gene-body methylation is involved in alternative splicing regulation. Our RNA-seq results indicated that many *T. spiralis* genes were alternatively spliced (Figure S6 in Additional data file 2), and in relation to this, we saw a steep change in the methylation frequency across splice junctions on both sense and antisense strands in the MR genes (Figure 6b). Moreover, there were considerable methylation differences between skipped and constitutive exons or between retained and spliced introns (Figure 6c). Particularly prominent was also the ‘infiltration of methylation’ in intronic sequences of neighboring splice junctions (Figure 6b, c). These results are in agreement with previous studies [25-27]. Taken together, our data suggest that DNA methylation status has a relationship with the donor/acceptor sequence context around the splice junctions, indicating the potential influence of DNA methylation on alternative splicing of MR genes.

**Discussion**

The status of DNA methylation in nematode genomes has been unclear. Research on *Caenorhabditis elegans*, which reportedly lacked mC in age-synchronous senescent populations, indicated it had negligible influence [1]. With regard to *Caenorhabditis elegans*, our computational searches here indicate that it has dnmt1, but not dnmt2 [12] or dnmt3. As dnmt3 is essential for de novo methylation [28], the lack of dnmt3 in *Caenorhabditis elegans* might explain the absence of de novo methylation in this nematode. In contrast, we identified three methyltransferase genes in the *T. spiralis* genome that were orthologs to known dnmts in vertebrates, including dnmt1, dnmt2 and dnmt3. Intriguingly, of 11 species of nematodes tested, *T. spiralis* appeared to be the only nematode possessing de novo methylation machinery (dnmt3). Moreover, we found that expression levels of dnmt2 and dnmt3 increased during development from larva to adulthood, but that dnmt1 did not. We note...
that our analysis indicated dnmt2 was more similar to a tRNA methylase instead of a DNA methyltransferase [12-14], which might indicate that RNA methylation also plays a role in *T. spiralis* development, and is therefore worth more detailed analysis in the future.

We further carried out the first comprehensive, high-resolution analysis of methylation in *T. spiralis*, to assess the intriguing possibility, given the presence of *de novo* DNA methyltransferase orthologs, that epigenetic control might help govern the development of its distinct life history stages via temporally regulated gene expression. Methylome sequencing revealed a mosaic methylation pattern in *T. spiralis*, typical of other invertebrates [29,30]. DNA methylation increased drastically during maturation from NBL to ML, and adults exhibited the highest observed DNA methylation level. This finding contrasts with a trend seen in some other species where methylation patterns remain stable throughout the life cycle [29]. For instance, in the sea urchin, which also has distinct life stages, methylated and non-methylated regions in its genome retain the same overall methylation composition throughout all tested stages [31]. The relative overall constancy of methylation patterns is also a feature of vertebrate genomes. However, the extent of DNA methylation may reflect changes in both intrinsic and environmental exposure [32]. For instance, studies in humans indicated that total genomic 5-methylcytosine has been found to typically decrease during aging [33,34], in concordance with declining Dnmt1 activity with age [35]. Parasites such as *T. spiralis* certainly undergo more drastic lifespan changes in response to environmental cues, that is, metamorphosis critical to their survival and reproduction. Our findings here provide evidence to indicate that these DNA methylation changes might play an important role in regulating such transformations in *T. spiralis*.

Previous studies have indicated that methylation may be an evolutionarily ancient means of transcriptional control as it is maintained in phylogenetically diverse lineages. In both plants and vertebrates, the notion that methylation in promoters primarily represses genes by impeding transcriptional initiation has been widely accepted [19,20], whereas intermediate levels of expression have been associated with genes experiencing the greatest extent of methylation in the gene body, indicating a bell-shaped relationship [4,21,22,36]. However, in the fungus *Neurospora crassa* [37] and the silkworm *Bombyx mori* [18], transcription initiation is unaffected. Thus, DNA methylation shows remarkable diversity in its extent and function across eukaryotic evolution. Here, our results indicate that the presence of promoter methylation correlates with reduced gene expression levels. Promoter hypermethylation may regulate a portion of stage-specific genes by repressing their transcription initiation in non-expressed stages, as exemplified by a NBL-specific DNase II gene (Figure 6a). Our assessment of gene-body methylation, which had a bell-shaped relationship between methylation and gene expression, indicates there was no overt relationship between expression and methylation levels. We did, however, see evidence for a relationship between methylation within the gene-body and alternative splicing of these genes in *T. spiralis*, indicating these regulatory mechanisms of gene and protein activity are an area of interest for future study as well; and the presence of a tRNA methylase ortholog makes further study of RNA regulation in general in the organism of interest.

In relation to the notion that complex regulatory machinery of DNA methylation has been developed in *T. spiralis* with species-specific characteristics, probably in response to environmental cues, we found that many of the MR genes are enriched in pathways that are functionally important for parasitic nematodes. Such genes modulate the interaction between the parasite and its host so as to protect the parasite against host immune responses. There was also enrichment in pathways that are important to parasitic activity, including previously reported catalytically active E-S proteins. Of note, hydrolases are among the most abundant proteins secreted by parasites and facilitate host tissue invasion [38]. Also important to *T. spiralis* is the conversion of muscle cells to nurse cells, and DNA-binding proteins [39], which are often affected by methylation changes, are believed to interfere with host cell signaling in ways that promote this conversion. Additionally, many such proteins are encoded by large, developmentally regulated gene families and assume different isoforms, which is also relevant to our findings that MRs were primarily distributed in gene bodies rather than promoter regions and the DNA methylation status was related to the donor/acceptor sequence context around their splice junctions.

**Conclusions**

We describe the first comprehensive study confirming the existence of DNA methylation in three life stages of *T. spiralis*. Our data also provide support for DNA methylation being associated with the regulation of genes that are closely related to the parasitism of the organism. In this context, in *T. spiralis* and in other organisms that experience discrete and highly specialized development forms, further consideration should be given to mechanisms where DNA methylation is involved in suppression of spurious transcriptional initiation of infrequently transcribed genes, promotes transcriptional termination, or mediates alternative splicing, as has been shown for other model organism systems.
Materials and methods

Collection of *T. spiralis* muscle larvae, adults and newborn larvae

Infected *T. spiralis* ML were obtained from infected mice at 35 days post-infection by digestion of minced skeletal muscle in 1% pepsin and 1% HCl for 45 minutes at 42°C with agitation, as previously described [40]. Seventy male 6-week-old Wistar rats were then orally inoculated with a dose of 8,000 infective ML. Adult worms (Ad1) were obtained from the intestine of ten rats at 30 h post-infection. The remaining 60 rats were sacrificed at 6 days post-infection, and the adult worms (Ad6) were recovered and incubated in Iscove’s Modified Dulbecco’s Medium (IMDM) in 75-cm² cell culture plates at 37°C. Newborn larvae were harvested every 6 h. All experiments were performed in accordance with the Guide for the Care and Use of Laboratory Animals published by the National Institutes of Health (publication no. 85-23, revised 1996). The protocol was approved by the Ethical Committee of the Institute of Zoonosis, Jilin University, China (reference number 20080106).

Enzymatic activity analysis of Dnmts

To test the dnmt enzymatic activity of *T. spiralis*, 11 μg of nuclear extracts for each assay were incubated in 37°C for 2 h using a EpiQuik™ DNMT Activity/Inhibition Assay Ultra Kit (Epigentek, Farmingdale, NY, USA) according to the manufacturer’s instructions.

BlastP searches and phylogenetic analysis of Dnmts

Reciprocal BlastP comparisons were first performed to identify dnmt orthologs. Significant hits were defined as those satisfying the following criteria: E-value < 10⁻⁵ and the aligned segments covering at least 30% of the sequence length of the hit. For phylogenetic analysis, multiple sequence alignment was performed by ClusterW [41]. The ClusterW with the neighbor-joining method [42] based on JTT + G (Jones-Taylor-Thornton and Gamma Distribution) model was applied to reconstruct the phylogenetic tree.

UPLC-MS/MS analysis of global DNA methylation

UPLC-MS/MS analysis was performed according to a previously published method [43]. Genomic DNA (0.2 μg) extracted from Ad and NBL was digested with 1U DNase I, 2U Alkaline Phophatase, Calf Intestinal (CIP) and 0.005U snake venom phosphodiesterase I at 37°C for 24 h. A microcon centrifugal filter device with a 3,000 D cutoff membrane was used to remove protein from the digested DNA samples by centrifuging at 12,000 rpm for 60 minutes. The mobile phase, consisting of 5.0% methanol and 95% water (plus 0.1% formic acid), was used for UPLC separation of the nucleotides at a flow rate of 0.25 ml/minute. Enzymatically digested DNA samples (10 μl each) were injected for UPLC-MS/MS analysis and each run took 10 minutes. Mass spectrometry conditions were as follows: ionization mode, ESI-positive; capillary voltage, 3,500 V; nitrogen drying gas temperature, 300°C; drying gas flow, 9 L/min; nebulizer, 40 psi. For MS/MS analysis of nucleotides, the fragmentor voltage was 90 V, collision energy was performed at 5 eV and scan time was 100 ms. Multiple-reaction monitoring (MRM) mode was used for the UPLC-MS/MS analysis by monitoring transition pairs of m/z 242.1/126.0 corresponding to 5mdC. The isotope labeled internal standard (5mdC-d3) was used to quantify genomic DNA methylation level, whose m/z was 245.4/129.0.

MethylC-seq library construction and sequencing

Prior to library construction, 5 μg of genomic DNA spiked with 25 ng unmethylated Lambda DNA (Promega, Madison, WI, USA) was fragmented using a Covaris sonication system to a mean size of approximately 200 bp. After fragmentation, libraries were constructed according to the Illumina Pair-End protocol with some modifications. Briefly, purified randomly fragmented DNA was treated with a mix of T4 DNA polymerase, Klenow fragment and T4 polynucleotide kinase to repair, blunt and phosphorylate the ends. The blunt DNA fragments were subsequently 3’ adenylation using Klenow fragment (3’-5’ exo-), followed by ligation to adaptors synthesized with 5’-methylcytosine instead of cytosine using T4 DNA ligase. After each step, DNA was purified using the QIAquick PCR purification kit (Qiagen, Shanghai, China). Next, a ZYMO EZ DNA Methylation-Gold Kit™ (ZYMO Research, Irvine, CA, USA) was employed to convert unmethylated cytosine into uracil, according to the manufacturer’s instructions, and 220 to 250 bp converted products were size selected. Finally, PCR was carried out in a final reaction volume of 50 μl, consisting of 20 μl of size-selected fractions, 4 μl of 2.5 mM dNTP, 5 μl of 10x buffer, 0.5 μl of JumpStart™ Taq DNA Polymerase, 2 μl of PCR primers and 18.5 μl water. The thermal cycling program was 94°C for 1 minute, 10 cycles of 94°C for 10 s, 62°C for 30 s, 72°C for 30 s, and then a 5-minute incubation at 72°C, before holding the products at 12°C. The PCR products were purified using the QIAquick gel extraction kit (Qiagen). Before analysis with Illumina Hiseq2000, the purified products were analyzed by the Bioanalyser analysis system (Agilent, Santa Clara, CA, USA) and quantified by real time PCR. Raw sequencing data were processed using the Illumina base-calling pipeline (Illumina Pipeline v1.3.1). The sodium bisulfite non-conversion rate was calculated as the percentage of cytosines sequenced at cytosine reference positions in the Lambda genome.
RNA-sequencing and real-time PCR validation

Total RNA was extracted using the Invitrogen TRizol® Reagent and then treated with RNase-free DNase I (Ambion, Guangzhou, China) for 30 minutes. The integrity of total RNA was checked using an Agilent 2100 Bioanalyzer. cDNA libraries were prepared according to the manufacturer’s instructions (Illumina). The poly(A)-containing mRNA molecules were purified using Oligo (dT) Beads (Illumina) from 20 μg of total RNA from each sample. Tris-HCl (10 mM) was used to elute the mRNA from the magnetic beads. To avoid priming bias when synthesizing cDNA, mRNA was fragmented before the cDNA synthesis. Fragmentation was performed using divalent cations at an elevated temperature. The cleaved mRNA fragments were converted into double-stranded cDNA using SuperScript II, RNaseH and DNA Pol I, primed by random primers. The resulting cDNA was purified using the QIAquick PCR Purification Kit (Qiagen). Then, cDNA was subjected to end repair and phosphorylation using T4 DNA polymerase, Klenow DNA polymerase and T4 Polynucleotide Kinase (PNK). Subsequent purifications were performed using the QIAquick PCR Purification Kit (Qiagen). These repaired cDNA fragments were 3’-adenylated using Klenow Exo- (Illumina) and purified using the MinElute PCR Purification Kit (Qiagen), producing cDNA fragments with a single ‘A’ base overhang at the 3’ end for subsequent ligation to the adapters. Illumina PE adapters were ligated to the ends of these 3’-adenylated cDNA fragments and then purified using the MinElute PCR Purification Kit (Qiagen). To select a size range of templates for downstream enrichment, the products of the ligation reaction were purified on a 2% TAE- Certified Low-Range Ultra Agarose (Bio-Rad, Hercules, CA, USA). cDNA fragments (200 ± 20 bp) were excised from the gel and extracted using the QIAquick Gel Extraction Kit (Qiagen). Fifteen rounds of PCR amplification were performed to enrich the adapter-modified cDNA library using primers complementary to the ends of the adapters (PCR Primer PE 1.0 and PCR Primer PE 2.0, Illumina). The 200 ± 20 bp PCR products were purified using QIAquick Gel Extraction Kit (Qiagen), using the MinElute spin columns (Qiagen). Finally, after detection on an Agilent Technologies 2100 Bioanalyzer using the Agilent DNA 1000 chip kit and quantification on a StepOne plus qPCR (ABI, Woodlands, Singapore), the cDNA library products were sequenced using the Illumina Genome Analyser. Real-time PCR validation was conducted using the Maxima® SYBR® Green qPCR Master Mix kit (Fermentas, Beijing, China), according to the manufacturer’s instructions, in an ABI Prism 7500 Sequence Detection System machine (Applied Biosystems Inc., CA, USA). All real-time RT-PCR data were normalized to the NBL stage (see Additional data file 1 for primer information).

Sequence alignment of MethylC-seq

The reads generated by Illumina sequencing were aligned onto the T. spiralis reference genome [11]. The Lambda genome was also included in the reference sequence as an extra chromosome so that reads originating from the unmethylated control DNA could be aligned. Because DNA methylation has strand specificity, the plus strand and the minus strand of the T. spiralis genome were separated to form alignment target sequences. To do this, each cytosine in the genome was converted to a thymine, termed the T-genome, which represented the plus strand. Meanwhile, each guanine in genome sequences was converted to adenosine, termed the A-genome, which represented the minus strand. Additionally, the original forms of the reads were also transformed to deal with the bisulfite treatment nucleotide conversion in the alignment process. First, the observed cytosines on the forward read of each read pair were replaced in silico by thymines, and secondly, the observed guanines on the reverse read of each read pair were replaced in silico by adenosines. We then mapped the ‘alignment form’ reads to the ‘alignment form’ target sequence using SOAPaligner with default parameters [16]. Every hit of a single placement with a minimum number of mismatches and a clear strand assignment was defined as an unambiguous alignment, and each alignment was used for mC ascertainment.

Gene annotation and functional analysis

For gene annotation, the BLAST algorithm was applied to further annotate the genes defined in the available T. spiralis genome annotation because the current annotation is incomplete. All the predicted protein sequences of T. spiralis genes were aligned using BLAST with known annotated protein sequences from three databases, including SWISS-Prot, TrEMBL and InterPro. A cutoff E-value < 1e-05 was applied for annotation, and a best alignment term for each query protein sequence was selected if more than one query sequence was aligned based on this cutoff E-value from BLAST.

For function analysis, GO analysis was performed based on the annotated genes by Gostat software [23]; 8,286 genes with annotation out of 16,380 genes were used as background, and 287 and 242 genes with annotation out of the 540 and 454 genes containing MRs in Ad and ML, respectively, were used as input genes. Fisher’s exact test was performed and the P-values generated for each GO category were adjusted according to the Benjamini and Hochberg correction method.

Identification of methylcytosines and methylation regions and determination of methylation level

For mC identification, we transformed each aligned read and the two strands of the T. spiralis genome back to
their original forms to build an alignment. In the unique part of the genome, cytosines that were covered by cytosines from reads on the same strand or guanines from those on the opposite strand (hereafter, referred to as ascertainment bases) were called as potential methylated sites. To account for the inefficient conversion of the bisulfite treatment and for sequencing errors, a correction method based on binomial tests and false discovery rate constraints [10] was applied to the data to build a high-quality methylome for each of the three stages. The probability \( p \) in the binomial distribution \( B(n, p) \) was estimated from the number of cytosine bases sequenced in reference cytosine positions in the unmethylated Lambda genome (referred to as the error rate: non-conversion plus sequencing error frequency). The bisulfite conversion rates for all samples were over 99%, and the error rates were as follows: Ad, 0.0060; ML, 0.0064; NBL, 0.0025. Then, the mCs from the binomial distribution analysis were selected for determination of MRs, which were defined as being under a threshold of more than five continuous mCs (either CpG or non-CpGs in at least one strand) and having a distance between adjacent mCs of less than the median value (18 bp) of all distance values. Stage-specific MRs were defined as containing more than five continuous mCs and no overlap between two samples.

Percentage methylation was computed as the fraction of reads number of ‘C’ in the total reads number of ‘C’ and ‘T’ for each covered CpG site, and herein average percentage methylation of all cytosine residues for any genomic region covered was computed as the fraction of reads number of ‘C’ in the total reads number of ‘C’ and ‘T’ for each genomic region. Density methylation (mC/C) was determined as the number of mCs divided by total number of C sites in any genomic region.

Validation of DNA methylation
Two strategies were applied to validate the methylation status of randomly selected genomic regions. BSP combined with cloning Sanger sequencing. BSP primers were designed by the online MethPrimer software (Additional data file 1). Genomic DNA (500 ng) was converted using the ZYMO EZ DNA Methylation-Gold Kit™ according to the manufacturer’s instructions. PCR amplification was carried out with a thermal cycling program of 94°C for 1 minute, 30 cycles of 94°C for 10 s, 58°C for 30 s, 72°C for 30 s, and then 5 minutes at 72°C. The products were then held at 12°C. Following amplification, PCR products were gel selected and purified using the QIAquick gel extraction kit (Qiagen), and the purified PCR products were subcloned. The colonies from each region were sequenced on a 3730 genetic analyzer (Applied Biosystems) to determine the methylated cytosine levels.

MeDIP followed by QPCR (see Additional data file 1 for primer information) was performed on 300 to 400 ng of original genomic DNA for each sample, which was randomly sheared to an average length of 200 to 500 bp by sonication. A MeDIP assay was then performed using the Magnetic Methylated DNA Immunoprecipitation Kit (Diagenode, Liege, Belgium) according to the instructions. The immunoprecipitated products and 10% amount of original input DNA were purified with ZYMO DNA Clean & Concentrator-5 kit (ZYMO) in parallel. The purified DNA was analyzed by QPCR on an ABI StepOne Plus Real Time PCR System (Applied Biosystems Inc.) using Eva Green (Biotium, Shanghai, China). The relative methylation levels of particular genomic loci among samples were compared by measuring the amount of immunoprecipitated DNA after normalization to the 10% of input DNA: % (MeDNA-IP/Total input) = 2^[Ct(10%input) - 3.32 - Ct(MeDNA-IP)] × 100%.

Data availability
Sequence data and processed data are available under the Gene Expression Omnibus accession GSE39328. UPLC-MS/MS and BS-PCR data have been deposited in GigaDB, the GigaScience database, with the unique identifier doi:10.5524/100043 [15].

Additional material

Additional file 1: Supplemental Tables S1 to S6 and corresponding captions
Additional file 2: Information on all primer sequences, and Supplemental Figures S1 to S6 and corresponding legends

Abbreviations
Ad: adult; BSP: bisulfite-PCR; ES: excretory-secretory; Gb: gigabase; GO: Gene Ontology; mC: methylcytosine; MeDIP: methylated DNA immunoprecipitation; ML: muscle larvae; MR: methylation region; NBL: newborn larvae; QPCR: quantitative PCR; UPLC-MS/MS: ultra-performance liquid chromatography-tandem mass spectrometry.

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Authors' contributions

FG, ML, QJ, XZ and HY conceived the project. FG designed experiments and interpreted data. XL, XPF, XLW, YS, JW, JD, XH, and YT performed experiments. FD, DL, YL and SL conducted bioinformatic analysis. FG prepared and wrote the manuscript. All authors have read and approved the manuscript for publication.

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