Myxosporea (Myxozoa, Cnidaria) lack DNA cytosine methylation

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Keywords: Methylome evolution, whole genome bisulfite sequencing (WGBS), Cnidaria, parasite, cytosine methylation.

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ABSTRACT
DNA cytosine methylation is central to many biological processes, including regulation of gene expression, cellular differentiation and development. This DNA modification is conserved across animals, having been found in representatives of sponges, ctenophores, cnidarians and bilaterians, and with very few known instances of secondary loss in animals. Myxozoans are a group of microscopic, obligate endoparasitic cnidarians that have lost many genes over the course of their evolution from free-living ancestors. Here, we investigated the evolution of the key enzymes involved in DNA cytosine methylation in 29 cnidarians, and found that these enzymes were lost in an ancestor of Myxosporea (the most speciose class of Myxozoa). Additionally, using whole genome bisulfite sequencing (WGBS), we confirmed that the genomes of two distant species of myxosporeans, Ceratonova shasta and Henneguya salminicola, completely lack DNA cytosine methylation. Our results add a notable and novel taxonomic group, the Myxosporea, to the very short list of animal taxa lacking DNA cytosine methylation, further illuminating the complex evolutionary history of this epigenetic regulatory mechanism.
INTRODUCTION
DNA methylation is a chemical modification of genomic DNA present in both prokaryotes and eukaryotes that affects gene regulation. DNA methylation of cytosines (cytosine methylation) has been extensively studied. An important functional consequence of DNA cytosine methylation is the suppression of gene expression when methylation is present in promoter regions (Shübeler 2015). This regulation affects a wide variety of key processes in animals including gametogenesis, embryonic development, cellular differentiation, X-chromosome inactivation, and transposon repression (Richa and Sinha 2014). Cytosine methylation also plays an important role in genome evolution. Cytosine methylation is mutagenic, as methylated cytosines can spontaneously deaminate to thymines (Bird 1980; Mendizabal et al. 2014). As a result, cytosine-guanine dinucleotides (CpGs) tend to change to thymine-guanine (TpG) dinucleotides in genomic DNA over the course of evolution (Bird 1980; Mendizabal et al. 2014). Recent methods have allowed researchers to use CpG to TpG conversion rates to reconstruct past DNA methylation patterns (Pedersen et al. 2014; Mendizabal et al. 2014). Methylated genes tend to be more conserved than non-methylated genes, even between distantly related species (Sarda et al. 2012). Such conservation attests to the importance of methylation at the genome level.

Similarly, the fact that cytosine methylation has been described in a wide variety of organisms, including viruses, prokaryotes and eukaryotes, suggests a key biological importance (e.g. Zemach et al. 2010). Most of what we know about DNA cytosine methylation in animals comes from studies in bilaterians. Nevertheless, it has also been described in a number of early branching animal lineages, including sponges, ctenophores and cnidarians (e.g. Zemach et al. 2010; Hassel et al. 2010; Dabe et al. 2015; Li et al. 2018; de Mendoza et al. 2019; Liew et al. 2020), and it is widely assumed that DNA methylation likely existed in the common ancestor of animals (Yi 2012, Zemach and Zilberman 2010). Strikingly, despite the biological importance of cytosine methylation and its high degree of conservation, a handful of animals are known to have secondarily lost this epigenomic modification. For example, cytosine methylation has been completely lost in the free-living nematode Caenorhabditis elegans, indicating that it is not crucial for gene regulation and development in this organism (Greer et al. 2015). It may have been also lost in the helminth parasite Schistosoma mansoni, although data are presently contradictory (Raddatz et al. 2013; Geyer et al. 2013). There is evidence that cytosine methylation has been completely lost in Diptera (flies), and in some hymenopterans (Bewick et al. 2017). However, the loss of cytosine methylation in the model dipteran Drosophila melanogaster is disputed (Raddatz et al. 2013; Capuano et al. 2014; Wang et al. 2018). Some species of lepidopterans, coleopterans, hemipterans and blattodeans also show extremely low levels of methylation, although due to the lack of genome assemblies, whether DNA methyltransferases (DNMTs) exist in these species is currently unresolved (Bewick et al. 2017). Adenine methylation is also known to occur in diverse species, including D. melanogaster and C. elegans, although the function of this type of DNA methylation in eukaryotes remains poorly understood (Low et al. 2001; Ratel et al. 2006; Greer et al. 2015; Zhang et al. 2015; Wu et al. 2016; Liang et al. 2018).

Cytosine methylation mediated gene regulation involves a network of proteins working to methylate nucleotides, bind to methylated regions, and subsequently alter gene expression. DNMTs are highly
conserved proteins responsible for DNA methylation (Fuks et al. 2000; Jin and Robertson 2013; Edwards et al. 2017). Three DNMTs have been inferred to be present in the common ancestor of animals (DNMT1, DNMT2/TRDMT1 and DNMT3) (Albalat et al. 2012). Studies in mammalian model systems demonstrated that patterns of cytosine methylation are first established by DNMT3, and then maintained by DNMT1 (Goll and Bestor 2005). The third DNMT, DNMT2/TRDMT1, has been implicated in the methylation of tRNA rather than DNA. Despite a very high degree of evolutionary conservation of DNMTs (e.g. Ponger and Li 2005), some DNMTs have been duplicated, or lost, in specific animal lineages (e.g. Yokomine et al. 2006; Barau et al. 2016; Bewick et al. 2017; Alvarez-Ponce et al. 2018). Cytosine methylation can trigger the binding of methyl-CpG-binding domain (MBD) proteins to methylated DNA, resulting in the recruitment of histone methyltransferases and histone deacetylases. These enzymes, in turn, modify the histone tails and affect gene expression (Fuks et al. 2000; Du et al. 2015). Two MBD proteins binding to methylated DNA have been inferred to be present in the common ancestor of animals and in cnidarians: MBD1/2/3 and MBD4/MeCP2 (Albalat et al. 2012). Two additional proteins with an MBD domain, MBDS and MB6, exist in animals but their function is unclear since, in mammals, they do not bind to methylated DNA (Laget et al. 2010); for this reason, they were not considered in our analyses. Methylation of cytosines can be converted into hydroxymethylated cytosines (5hmC), and then to 5-formylcytosine (5fC) and 5-carboxylcytosine (5caC) by proteins known as Ten-eleven translocation methylcytosine dioxygenases (TETs) (Tahiliani et al. 2009; Kriaucionis et al. 2009; Guo et al. 2011). The resulting 5hmC, 5fC and 5caC can be subsequently converted to unmethylated cytosine via base excision repair or replication-dependent dilution (reviewed in Wu and Zhang 2017). Consequently, TETs can actively demethylate methylated cytosines. Since their first discovery, the role of TETs in regulating methylation and demethylation has been shown to be critical in development and gene regulation (Lu et al. 2015). While three copies of TET are present in vertebrates, a single copy of the gene is assumed to have been present in the ancestor of animals (Liu et al. 2020). Similar proteins are involved in adenine methylation, with Methyltransferase like 4 (METTL4) having an analogous function for adenine as DNMTs do for cytosine (Iyer et al. 2016). At present, DNA N6-methyl methyltransferase (DAMT-1), the C. elegans ortholog of METTL4, is the only animal protein that has been confirmed to act as an adenine DNA methyltransferase (Greer et al. 2015; Luo and He 2017). Collectively, these DNA methylation-related proteins lay the foundation for DNA methylation-driven regulation.

DNA methylation enzymes, and methylomes, are relatively understudied in non-model organisms, including cnidarians (see Figure 1) (Fuchs et al. 2014, Li et al. 2018, Putnam et al. 2016; Eirin-Lopez and Putman 2019). Here, we present the first investigation of DNA methylation in a wide variety of cnidarians, including both free-living and parasitic cnidarians. Myxozoans are microscopic and parasitic cnidarians that typically infect fish and annelids (Kent et al. 2001; Chang et al. 2015; Atkinson et al. 2018). They are divided into two classes: Myxosporea, which encompass the vast majority of myxozoans; and the less-studied Malacospora, with less than 10 species (Fiala et al. 2015). Myxosporeans are further divided into a “marine/polychaete-host” and “freshwater/oligochaete-host” clade (Fiala et al. 2015). They exhibit highly reduced genomes (22-175 Mb) compared to other cnidarians (256-1,260 Mb), consistent with the pattern of parasitic organisms having smaller genomes than their closest free-living relatives (Dieterich and Sommer 2009; Chang et al. 2015).
We searched for the presence of known methylation-associated proteins (DNMTs, MBDS, and TETs) in available genomic and transcriptomic data from 29 cnidarians, including 8 myxozoans representative of the myxosporean diversity (representing the “marine/polychaete-host” lineage: *Ceratonova shasta, Kudoa iwatai* and *Enteromyxum leei*; and the “freshwater/oligochaete-host” lineage: *Sphaeromyxa zaharoni, Henneguya salminicola, Thelohanellus kitauei, Myxobolus cerebralis* and *Myxobolus squamalis*; Atkinson et al. 2018; Holzer et al. 2018; Yahalomi et al. 2020). While homologs of genes encoding methylation-associated proteins are present in all non-myxozoan cnidarians with available complete genomes, we did not find any of these genes in any of the studied myxosporean genomes. We thus hypothesized that myxosporeans had lost DNA cytosine methylation. To test this hypothesis, we then performed whole genome bisulfite sequencing (WGBS, a technique that allows determining which DNA cytosines are methylated) on two highly divergent myxosporeans (*C. shasta* and *H. salminicola*) to determine the level of DNA cytosine methylation in these species. Our sequencing results confirmed that these myxosporeans completely lack cytosine methylation. We thus add a new animal group, the Myxosporea, to the very short list of species known to lack DNA cytosine methylation.

**RESULTS**

**DNA Cytosine Methylation Proteins Were Lost in an Ancestor of Myxosporea**

We obtained the sequences of the DNMT, MBD, TET and METTL4 proteins of the cnidarians *Oribicella faveolata* (Anthozoa) and *Hydra vulgaris* (Hydrozoa). Using these sequences, we performed BLAST searches against the genome assemblies, annotated protein sequences, and/or transcriptomes, of 29 cnidarian species. These included 8 myxosporeans, *Polypodium hydriforme* (the most closely related cnidarian species to myxozoans; Chang et al. 2015), and 20 other non-myxozoan cnidarians including both anthozoans (corals and sea anemone) and medusozoans (jellies and hydras). We detected homologs of all these proteins in all of the non-myxozoan cnidarians, except for *P. hydriforme*, which lacked transcripts encoding TET, DNMT3, and METTL4 (Figure 1). We detected METTL4 predicted proteins in all 8 myxosporeans included in our study, but did not detect any DNMT, MBD, or TET protein (Figure 1).

The fast rate of evolution of myxozoans might have hindered the detection of the proteins of interest using BLAST searches. To address this possibility, we designed seven Hidden Markov Model (HMM) profiles (see Methods) for the DNMT1, DNMT2/TRDMT1, DNMT3, MBD1/2/3, MBD4/MeCP2, TET and METTL4 proteins to search against predicted protein databases deposited in the NCBI or generated from published transcriptomes of 6 myxosporeans (*C. shasta, H. salminicola, K. iwatai, T. kitauei, M. cerebralis* and *M. squamalis*). We manually evaluated the homology of each HMM hit with reciprocal BLASTp searches against the nr database (NCBI Resource Coordinators 2016). Phylogenetic trees were constructed for each protein dataset including the database of proteins representative of the animal and cnidarian diversity selected for the HMM construction and the hit found in Myxosporea and *P. hydriforme*, with the sponge *Amphimedon queenslandica* as outgroup (Supplementary Figures 1-7). Once again, we detected METTL4 predicted proteins for the myxosporeans included in our study, but did not detect any DNMT, MBD, or TET protein in myxosporeans. Thus, our HMM searches corroborate our BLAST searches.
Mapping the presence or absence of these genes onto a cnidarian phylogenetic tree (Figure 1) allowed us to infer that (1) the most recent common ancestor of cnidarians harbored the same number of methylation genes as the ancestor of animals; (2) DNMT1, DNMT2/TRDMT1 and MBD proteins were lost in a common ancestor of Myxosporea; (3) DNMT3 and TET were lost in a common ancestor of P. hydriforme and Myxozoa, and (4) METTL4 were lost in an ancestor of P. hydriforme. It should be noted, however, that no complete genome is available for P. hydriforme, and because transcriptomic datasets do not represent all genes, it is possible that DNMT, MBD and TET proteins might have been lost in an ancestor of Myxosporea (Figure 1).

**Ceratonova shasta** and **Henneguya salminicola** Lack DNA Cytosine Methylation

The absence of genes related to cytosine methylation in myxosporeans led us to hypothesize that they lacked DNA cytosine methylation. To test this hypothesis, we sequenced the methylomes of two distantly related myxosporeans, *C. shasta* (representative of the “marine/polychaete” lineage) and *H. salminicola* (representative of the “freshwater/oligochaete” lineage). To directly measure cytosine methylation in these myxosporeans, we obtained total genomic DNA from the intestine of a rainbow trout (*Oncorhynchus mykiss*) infected with *C. shasta* and from a muscle cyst of a Chinook salmon (*Oncorhynchus tsawytscha*) infected with *H. salminicola*. We added lambda phage DNA to both samples (as a negative control for bisulfite conversion rate, since it is unmethylated), and then we performed whole genome bisulfite sequencing (WGBS). Therefore, our raw WGBS data for *C. shasta* contained sequences from *C. shasta*, rainbow trout and lambda phage genomic DNA (Supplementary Table 1). Similarly, our raw WGBS data for *H. salminicola* contained *H. salminicola* genomic DNA sequences, as well as trace amounts of Chinook salmon and lambda phage DNA sequences (Supplementary Table 2).

We first aligned both WGBS datasets to the lambda phage genome using Bismark to calculate the bisulfite non-conversion rates (Krueger and Andrews 2011). In addition to the CpG context, we also examined CHG and CHH cytosine methylation (where H represents A, T or C nucleotides), which occurs in eukaryotic genomes (Zemach et al. 2010). For the *C. shasta* WGBS dataset, we determined the bisulfite non-conversion rates to be 0.1%, 0.2%, and 0.1% for the CpG, CHG, and CHH methylation contexts, respectively (Table 1). For the *H. salminicola* WGBS dataset, we determined the bisulfite non-conversion rates to be 0.1% for each of the CpG, CHG, and CHH methylation contexts, respectively (Table 1).

We then aligned the remaining WGBS reads (those that did not align to the lambda phage genome) in the *C. shasta* and *H. salminicola* datasets to the rainbow trout and Chinook salmon genomes (Berthelot et al. 2014; Christensen et al. 2018), respectively. These alignments served as positive control, as fish genomes are known to be highly methylated (Jabbari et al. 1997). We detected 73.8% CpG methylation for the rainbow trout in the *C. shasta* WGBS dataset (Table 1), and 49.0% CpG methylation for the Chinook salmon in the *H. salminicola* WGBS dataset (Table 1). The lower methylation levels measured for the Chinook salmon were most likely due to the low number of sequence reads for this species contained in the *H. salminicola* dataset (Supplementary Table 2).
We also performed deep whole genome sequencing to identify polymorphic sites that occur at cytosines between the reference genome and the sequenced samples. We observed a 2.8% SNP difference between the C. shasta genome assembly and our C. shasta WGS dataset, and a 0.5% SNP difference between the H. salminicola genome assembly and our H. salminicola WGS dataset (Supplementary Table 3). These sites only impacted between 3.5-4.8% and 0.58-0.72% of CpG, CHG, and CHH sites for C. shasta and H. salminicola respectively and we removed them from downstream analyses to avoid errors due to SNPs.

We aligned the remaining WGBS reads (those that did not align to either the lambda phage genome or the fish host genomes) to the genome assemblies of C. shasta and H. salminicola (see Material and Methods). By virtue of extremely high sequencing depth, the average coverages of the mapped cytosines were very high: ~400 X for C. shasta and ~1600 X for H. salminicola (Figures 2 and 3; Supplementary Tables 4 and 5). For both species, we determined the CpG, CHG, and CHH methylation rates to be 0.1% for each of these cytosine contexts (Table 1). These methylation rates were nearly equal to the bisulfite non-conversion rates, indicating that C. shasta and H. salminicola lack DNA cytosine methylation.

We calculated the fractional methylation (number of methylated reads/total reads per cytosine site) for all three cytosine contexts (CpG, CHG, and CHH) for the final sequence alignments of our WGBS datasets. The histograms of fractional methylation frequency showed only one peak at the zero end of the x-axis for each of the three methylation contexts, thus indicating no cytosine methylation for either C. shasta or H. salminicola (Figures 2 and 3).

DISCUSSION
We searched the genomes, proteomes and/or transcriptomes of 29 cnidarians, including 8 myxosporeans, for the presence of genes encoding core proteins involved in DNA methylation. Our BLAST and HMMER searches revealed that DNMTs, MBDs and TET are present in most of the studied non-myxozoan cnidarians, in agreement with previous reports of DNA cytosine methylation in a wide range of cnidarians (Zemach et al. 2010; Hassel et al. 2010; Putnam et al. 2016; Groves et al. 2016; Liew et al. 2018; Li et al. 2018) (Figure 1). However, we could not find these genes in any of the 8 myxosporeans with available genomes or transcriptomes (Figure 1). Placing this information into a phylogenetic context suggests that proteins encoding these genes were lost in a common ancestor of Myxosporea (Figure 1).

The only non-myxozoan cnidarian that appears to lack some of these proteins is P. hydridiforme, an enigmatic parasite of paddlefish and sturgeon oocytes which is the sister taxon to Myxozoa (Chang et al. 2015). This species appears to lack DNMT3 and TET proteins, which would indicate that these proteins would have been lost in a common ancestor of Myxozoa and P. hydridiforme. It should be noted, however, that our searches against P. hydridiforme relied on transcriptome data, as a complete genome assembly is currently unavailable. As transcriptomes do not capture the whole set of genes of an organism, it is possible that these genes are present in P. hydridiforme, in which case they would have been most likely lost in an ancestor of Myxosporea, subsequent to divergence from the common
ancestor with *P. hydriforme*. Future analyses using a whole genome assembly of this species will allow to draw a definitive conclusion.

Our analyses are based on eight myxozoan species that belong to the subclass Myxosporea (a clade that contains the vast majority of myxozoan species). Myxozoa, however, encompass a second distant subclass, the Malacosporea, which include only two genera (*Buddenbrockia* and *Tetracapsuloides*) and less than 10 species (Fiala et al. 2015). Unfortunately, in public databases, genomic data are limited to less than 1,500 EST for Malacosporea, which precluded the inclusion of this subclass in our analyses. Preliminary BLAST searches and phylogenetic analyses, however, revealed the presence of MBD1/2/3 in *Buddenbrockia plumatellae* (sequence accession: ES599526). This suggests that the loss of cytosine methylation is limited to Myxosporea and not a characteristic shared by all Myxozoa. Future studies should determine if Malacosporea possess a complete set of methylation enzymes or represent an intermediate stage in the loss of cytosine methylation.

In agreement with our observation that myxosporeans lack DNMTs, MBDS and TET proteins, our newly generated WGBS data show that two highly divergent myxosporeans, *C. shasta* and *H. salminicola* (representatives from the fresh-water and the marine lineages, respectively; Fiala et al 2015, Holzer et al 2018), completely lack DNA cytosine methylation (Table 1; Figures 2 and 3). Thus, we have identified a new taxonomic group with no cytosine methylation, placing Myxosporea among the very few animal clades known to lack this form of epigenetic modification.

Given that DNMT1, DNMT2/TRDMT1 and MBDS (and potentially DNMT3 and TET proteins too, as discussed above) were lost in the same branch of the cnidarian tree (the branch predating the diversification of Myxosporea), it is possible that the loss of DNMTs in an ancestor of Myxosporea may have led to a subsequent loss of MBD and TET proteins, as without DNA cytosine methylation, these proteins would have been rendered nonfunctional. Alternatively, MBDS could have been lost first, which would have rendered cytosine methylation (and thus DNMTs) nonfunctional. The sequencing of Malacosporea may help to answer this question.

Given the importance of cytosine methylation in regulation of gene expression, development and genome defense, it is a pressing question to understand how the loss of cytosine methylation impacts these processes. In terms of gene expression, other epigenetic mechanisms, notably histone modification, may play a major role of regulation, as in *C. elegans* and *D. melanogaster*, which also lack cytosine modification (Chang and Liao 2017). Similar molecular mechanisms may exist in myxosporeans. Interestingly, we found METLL4 orthologs in all the studied cnidarians (including all myxosporeans), except *P. hydriforme* (for which, as discussed above, we lack a complete genome). Thus, cnidarians, including myxozoans, might harbor adenine DNA methylation (even though the functional relevance of this modification needs to be resolved; Flusberg et al. 2010).

Our study adds to a growing list of species where DNA methylation has been lost. Evolutionary forces that facilitate loss of DNA methylation remain unclear (Yi 2017, Takuno et al. 2016, Schmitz et al. 2019, Bewick et al. 2019). An interesting observation is that myxozoans generally have very small genomes,
and thus are potentially relatively free from the burden of active genome defense. In plant genomes, there is a strong negative correlation between genome size and the degree of genomic CHG DNA methylation (Niederhuth et al. 2016, Vidalis et al. 2016), which is the primary DNA methylation of transposable elements in plants. In comparison, factors affecting genomic DNA methylation in animal lineages are still relatively little understood. Future studies should determine the significance of genome defense and genomic DNA methylation in cnidarians.

The current understanding of the evolutionary loss of DNA methylation in animals comes primarily from studies using model species and human parasites. Our work offers an example of a DNA methylation survey of a group of non-model organisms using straightforward and highly replicable molecular and computational techniques. Given that it is becoming less prohibitive to generate whole genome sequencing data and whole genome bisulfite sequencing data of non-model organisms, an investigative framework similar to the method used here could be applied to a variety of eukaryotic species in a near future. Such data will allow evolutionary biologists to generate a more complete understanding of the intricate evolutionary dynamics of DNA methylation. On one hand, the pervasiveness of DNA methylation across a broad variety of taxa, as well as the high level of conservation of the methylation machinery, are a testament to the biological importance of this epigenetic modification in genomic regulation. On the other hand, despite this paradigm of regulation, our findings suggest that a speciose group of parasitic cnidarians – the Myxosporea – are among a short but notable list of animal taxa that have lost DNA cytosine methylation and the enzymatic machinery responsible for it. With future data on genomic DNA methylation surveys from diverse species, we will soon be able to illuminate the factors that determine the tradeoffs between conservation and loss of DNA methylation across the tree of life.

MATERIALS AND METHODS
BLAST Searches
We performed online BLASTp searches (Altschul et al. 1997) using as query the O. faveolata and H. vulgaris DNMT1, DNMT2/TRDMT1, DNMT3, MBD1/2/3, MBD4/MeCP2, TET and METTL4 protein sequences against all cnidarians with protein sequences in the NCBI’s nr database and with a scaffold N50 greater than 90,000 on the NCBI Genome database (NCBI Resource Coordinators 2016). The latter criterion was chosen to ensure that the absence of a protein could not be explained by a low assembly quality (low N50 values result in many genes being spread across multiple scaffolds, which may lead to incomplete and/or erroneous protein sequence predictions). The list of cnidarian species meeting these criteria is provided in bold in Supplementary Table 6. In addition, we performed tBLASTn searches using the same query sequences against the unannotated genomes and transcriptomes of additional non-myxozoans representative of the cnidarian diversity; all cnidarians with an available genome assembly in the NCBI Genome database (as per June 2019) were included in our analyses (Figure 1, Supplementary Table 6). For each cnidarian species considered, the orthology of the first five hits was assessed by performing reciprocal blast searches against the H. vulgaris proteome and reconstructing phylogenetic trees (see below).

We next performed local tBLASTn searches (BLAST+ v2.9.0) (Camacho et al. 2009), using all the non-myxozoan cnidarian protein sequences identified in Supplementary Tables 7-13, against the genome
assemblies or transcriptomes of 8 myxosporeans (C. shasta, H. salminicola, K. iwatai, E. leei, S. zaharoni, T. kitauei, M. cerebralis and M. squamalis), and the transcriptome of P. hydriforme. Accession numbers of the assemblies used are provided in Supplementary Table 6. For each myxosporean hit obtained, a reciprocal BLASTx search was performed against the NCBI’s nr database (last accessed in February 2020) to verify the identity of the protein sequences identified by our local tBLASTn. These BLAST searches ensured that the myxosporean hits were indeed cnidarian sequences rather than host contamination or erroneous BLAST hits.

All BLAST searches were conducted using default parameters. The accession numbers for all protein sequences used for and found during our BLAST searches can be found in Supplementary Tables 7-13.

**HMM Profile Construction and Searches**

For HMM profile construction, all cnidarian sequences identified in our previous BLAST searches were included together with representatives of the main animal lineages (see Supplementary Tables 7-13). Multiple sequence alignments of each group of proteins were created using MAFFT v7.450 with default parameters (“auto”) (Katoh and Standley 2013), and the alignments were manually inspected in Geneious Prime software version 2019.2.3 (Kearse, et al. 2012). The accession numbers of all proteins used in HMM profile construction can be found in Supplementary Tables 7-13. HMM profiles were built based on the MAFFT alignments using hmmbuild from the HMMER package v3.1b2 (Eddy, 2011). Alignments and HMM profiles are provided in Supplementary Files 1-15.

The T. kitauei, M. squamalis and H. salminicola proteomes were retrieved from the NCBI Database (Supplementary Table 6). Proteins sequences of the other three myxosporean species (C. shasta, K. iwatai and M. cerebralis) and P. hydriforme were predicted from transcriptome sequences using Transdecoder version v5.5.0 ([https://github.com/TransDecoder](https://github.com/TransDecoder)) with default parameters. For each species, the protein database generated or downloaded from NCBI was searched using hmmsearch from the HMMER package v3.1b2 under the HMM built as described above (Eddy, 2011). Because there is no transcriptome data available for E. leei or S. zaharoni, these two species were excluded from our HMM analyses.

Finally, to confirm the orthology of the proteins identified using BLAST and HMM searches, phylogenetic trees were reconstructed for each gene considered in this study (DNMT1, DNMT2/TRDMT1, DNMT3, MBD2, MBD4, TET, METTL4) using the amino acid alignments used previously for HMM build. Specifically, the GUIDANCE2 server version 2.02 (Sela, et al. 2015) was used to remove ambiguously aligned positions, with the following parameters: MAFFT program, 100 bootstrap, “--maxiterate 1000”, and “--localpair”. Unreliable column positions with a reliability score below 0.5 were excluded, as well as columns which included more than 50% of gaps. The different alignments are provided in Supplementary File 1. Maximum likelihood phylogenies were inferred using IQ-TREE version 1.6.12 (Nguyen et al. 2015) with automatic model detection using ModelFinder Plus and 1000 bootstrap replicates (Supplementary Figures 1-7).

**Genome Assemblies**
The *H. salminicola* genome assembly was obtained from Yahalomi et al. (2020). We used our draft *C. shasta* genome assembly (Version Velvet2015-93; Atkinson, unpublished data; 15,423 sequences, average contig length: 6823, N50: 47,028, assembled size: 105.2 Mb, average coverage: ~250x, filtered of rainbow trout and bacterial contamination). The genome assembly IDs for all genomes used in our study are listed in Supplementary Table 6.

**Biological Material**

Fresh *H. salminicola* was obtained in September 2016 from the skeletal muscle of an adult Chinook salmon (*O. tschawytscha*) from the Willamette River, Oregon, USA. Parasite material, predominantly mature myxospores, was aspirated from a single pseudocyst, and then dried in a SpeedVac (Savant), before being stored at -20 °C until processing for sequencing. Fresh *C. shasta* was obtained in February 2014 from the intestine of a juvenile rainbow trout (*O. mykiss*) that had been infected during a cage exposure in the Willamette River, Oregon. Fresh intestine was dissected from an infected fish, macerated with scissors, and suspended parasite spores washed through a 70 µm cell sieve using phosphate-buffered saline (PBS). The filtered material was washed twice, by pelleting by centrifugation (~2000 x G, 10 min), with the supernatant exchanged with fresh PBS after each centrifugation. The semi-pure myxospores were then layered on top of a Percoll (Sigma-Aldrich) gradient (layers with concentrations of 25%, 50% and 75%) and centrifuged for ~2500 x G for 15 min. The almost pure myxospores formed a band on top of the 75% layer. The spores were aspirated from the gradient, then again washed by centrifugation in PBS to remove Percol. The spores were then dried in a SpeedVac (Savant), before being stored at -20 °C until processing for sequencing.

**Whole Genome Sequencing**

Whole genome sequencing libraries were generated from DNA extracted using the DNeasy Blood and Tissue DNA kit (Qiagen). From each sample, 500 ng-1 µg of DNA was sheared on a Covaris ultrasonicator (Covaris, Woburn, MA) to 200-600 bp at the Emory Integrated Genomics Core. The DNA fragment ends were repaired with the End-It DNA End-Repair Kit (#ER81050, Epicentre, Madison, WI) and A-overhangs were added (#M0202, New England Biolabs, Ipswich, MA) before Nextera barcode adaptors were ligated to the DNA fragments overnight. Finally, the libraries were PCR amplified to increase concentration and enrich for adaptor-ligated DNA fragments. WGS libraries were sequenced using Illumina HiSeq X 150 with paired-end reads at the Macrogen Clinical Laboratory.

**Variant Calling for Whole Genome Sequences**

We used FastQC (v0.11.7) and Trim Galore (v0.5.0) to examine the quality and remove the adaptor sequences in our raw, paired-end whole genome sequencing (WGS) data for *C. shasta* and *H. salminicola*. The raw WGS data for *C. shasta* contained *C. shasta* and rainbow trout genomic DNA sequences. The raw WGS data for *H. salminicola* contained *H. salminicola* and Chinook salmon genomic DNA sequences. We used BBSPsplit, part of the BBMap package (v38.62) (Bushnell, B. 2014), to remove all fish contamination from both sets of WGS data. Next, we aligned the WGS reads to the respective *C. shasta* and *H. salminicola* reference genome assemblies using BWA (v0.7.17) (Li and Durbin 2009) under default parameters. We then used bamtools (v1.9) to convert the SAM files produced by the alignments to BAM files. Then, we used samtools (v1.9) again to extract correctly paired reads, remove duplicate
reads, and extract all reads with a map quality of 30 or higher, and acquire statistics to calculate the average genome coverage for the *C. shasta* and *H. salminicola* WGS data.

To perform variant calling, we used bcftools (v1.9) (Li et al. 2009). We then used SnpSift (v4.0) (Cingolani et al. 2012) to extract all SNPs where the sequence depth was greater than or equal to 20. These high quality SNPs were used for a comprehensive variant calling analysis performed with bcftools for the *C. shasta* and *H. salminicola* WGS data. The SNP density was computed with vcftools (v1.9) (Danecek et al. 2011), which allowed to calculate average numbers of SNPs per Kb. Afterwards, we used ANGSD (v0.928) (Korneliussen et al. 2014) to convert the BAM files to consensus FASTA files. Finally, we used Mauve (2015-02-13 build) (Darling et al. 2004) to compare the reference genome assembly FASTA files and consensus FASTA files for *C. shasta* and *H. salminicola* and the outputted statistics to calculate percent differences between the genome assembly and consensus FASTA files. All the relevant statistical information obtained from our analysis are presented in Supplementary Table 3.

**Whole Genome Bisulfite Sequencing**

To generate whole genome bisulfite sequencing Libraries, genomic DNA was pooled with 1-5% lambda phage DNA as a test to control for bisulfite reaction efficiency. The DNA samples were then sheared on a Covaris ultrasonicator to 200-600 bp. The DNA fragment ends were repaired and A-overhangs were added before bisulfite compatible adaptors were ligated to the DNA fragments overnight. Next, the DNA fragments were bisulfite-converted using the MethylCode Bisulfite Conversion Kit (#MECOV50, ThermoFisher). Purified gDNA was treated with CT conversion reagent in a thermocycler for 10 min at 98 °C, followed by 2.5 h at 640 °C. Bisulfite-treated DNA fragments remain single-stranded as they are no longer complementary. Low-cycle (4-8) PCR amplification was performed with Kapa HiFi Uracil Hotstart polymerase enzyme (#KK2801, KAPA Biosystems, Wilmington, MA), which can tolerate Uracil residues. The final library fragments contain thymines and cytosines in place of the original unmethylated cytosine and methylated cytosines, respectively. WGBS libraries were sequenced using Illumina HiSeq X 150 with paired-end reads at the Macrogen Clinical Laboratory.

**Whole Genome Bisulfite Sequence (WGBS) Quality Control**

The raw WGBS data for *C. shasta* contained *C. shasta*, rainbow trout, and lambda phage genomic DNA sequences. The raw WGBS data for *H. salminicola* contained *H. salminicola*, Chinook salmon, and lambda phage genomic DNA sequences. We first used FastQC (v0.11.7) (Andrews, S. 2010) to analyze the quality of the reads. Next, we used Trim Galore (v0.5.0) (Martin, M. 2011; Krueger and Andrews 2012; Krueger, F. 2015) to remove the Illuma adaptor sequences and very low quality reads. Finally, we used FastQC again to analyze the quality of the trimmed reads.

**Whole Genome Bisulfite Sequence (WGBS) Alignment with Bismark**

We aligned and analyzed the trimmed reads for the *C. shasta* and *H. salminicola* datasets using Bismark (v0.20.0) (Krueger and Andrews 2011; Krueger and Andrews 2012) with Bowtie 1 (v1.0.0) (Langmead et al. 2009). First, we aligned the reads to the lambda phage genome to calculate the bisulfite conversion rate (Wreczycka et al. 2017). As part of this step, we used Bismark’s “—un” flag to retain the myxosporean and host fish reads, as these reads do not align to the lambda phage genome. Note that...
the term “host fish” refers to the vertebrate hosts of C. shasta and H. salminicola, which are the rainbow trout and the chinook salmon, respectively. Next, we aligned the unmapped reads (myxosporean and host fish sequences) from the previous step to the host fish genomes, while using the “–un” flag to retain all myxosporean reads. Finally, we aligned the unmapped reads (myxozoan sequences) from the previous step to the respective C. shasta and H. salminicola reference genome assemblies. Afterwards, we used deduplicate_bismark on the BAM output files produced in the previous steps to remove PCR duplication bias. Then, we used bismark_methylation_extractor on these deduplicated BAM files to acquire the final methylation data (Krueger and Andrews 2012; Wreczycka et al. 2017). Additionally, we used bam2nuc on the deduplicated BAM output files produced from the final C. shasta and H. salminicola alignments to assess the nucleotide coverage of the alignments (Supplementary Tables 4 and 5). Finally, we used bismark2report to create six HTML data summary report files from the six sets of files created from the previous steps. All code used for the quality control and sequence analysis is available upon request. The methylation data we obtained from each of the six alignments was used to construct Table 1.

DNA Methylation Analysis
We utilized Bismark’s bismark_methylation_extractor, bismark2bedGraph, and coverage2cytosine in series (v0.20.0) (Krueger and Andrews 2011; Krueger and Andrews 2012) to generate reports for cytosine in three nucleotide contexts, CpG, CHG, and CHH. To counteract inaccurate methylation calls that can arise due to polymorphisms between the reference genomes and WGBS samples (i.e. C-to-T polymorphisms would always be designated unmethylated irrespective of reference methylation status), all cytosine sites overlapping identified variants were removed. We calculated the fractional methylation (ratio of the number of methylated cytosine reads to the total number of methylated and unmethylated reads) for each analyzed site. Finally, we plotted the frequencies of fractional methylation in all three contexts for both C. shasta and H. salminicola.

Acknowledgements: The authors are grateful to Tom Parchman for providing access to his server and Richard Tillett for helpful advice on data analysis. RK, ALN, SP and DAP were supported by a grant from the National Science Foundation (MCB 1818288) and a Pilot Grant from the Smooth Muscle Plasticity COBRE of the University of Nevada, Reno, funded by the National Institutes of Health (grant 5P30GM110767-04). TL, DS and SVY were supported by a grant from the National Science Foundation (MCB 1615664). DH was supported by a grant from the Tel-Aviv University Vice President’s fund for research support (grant number 30003072000). SA and JB were supported in part by the Bureau of Reclamation, US Department of Interior through Inter-agency Agreement #R15PG00065, as part of its mission to manage, develop, and protect water and related resources in an environmentally and economically sound manner in the interest of the American public. The views in this report are the authors’ and do not necessarily represent the views of Bureau of Reclamation. The funders had no role in study design, data collection and analysis, decision to publish, or preparation of the manuscript. Mention of trade names does not imply U.S. Government endorsement.

Conflict of Interest Statement: The authors declare that they have no conflicts of interest.
Data availability: Sequencing data have been deposited to the BioProject database (accession numbers PRJNA623035 and PRJNA623156).
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Table 1: Methylation analysis with Bismark showed that *Ceratonova shasta* and *Henneguya salminicola* have no DNA cytosine methylation

| Bisulfite Sequence Data | Reference Genome Assembly | C methylated in CpG context | C methylated in CHG context | C methylated in CHH context |
|------------------------|---------------------------|-----------------------------|----------------------------|-----------------------------|
| *C. shasta* + rainbow trout + lambda phage | lambda phage | 0.1% | 0.2% | 0.1% |
| *C. shasta* + rainbow trout | rainbow trout | 73.8% | 0.2% | 0.2% |
| *C. shasta* | *C. shasta* | 0.1% | 0.1% | 0.1% |
| *H. salminicola* + Chinook salmon + lambda phage | lambda phage | 0.1% | 0.1% | 0.1% |
| *H. salminicola* + Chinook salmon | Chinook salmon | 49.0% | 11.9% | 16.2% |
| *H. salminicola* | *H. salminicola* | 0.1% | 0.1% | 0.1% |

The table lists the percent cytosine methylation for the CpG, CHG, and CHH methylation contexts for each of the six bisulfite sequence alignments performed using Bismark. The first row summarizes the methylation results for the alignment of the raw WGBS sequence reads for *C. shasta* to the lambda phage genome. The second row summarizes the results for the alignment of the *C. shasta* filtered reads from the previous alignment to the rainbow trout genome. The third row summarizes the results for the alignment of the *C. shasta* final filtered reads from the previous alignment to the *C. shasta* genome excluding cytosine sites overlapping SNPs from the reference genome. The fourth row summarizes the methylation results for the alignment of the raw WGBS sequence reads for *H. salminicola* to the lambda phage genome. The fifth row summarizes the results for the alignment of the *H. salminicola* filtered reads from the previous alignment to the *H. salminicola* genome. The sixth row summarizes the results for the alignment of the *H. salminicola* final filtered reads from the previous alignment to the *H. salminicola* genome excluding cytosine sites overlapping SNPs from the reference genome. The percent methylation data shown here are from after deduplication and extraction using deduplicate_bismark and bismark_methylation_extractor.
Figure 1: Presence/absence of methylation related genes across cnidarians.

The tree was assembled from published trees (Kayal et al. 2018; Yahalomi et al. 2020). Black and white squares represent the presence and absence of each gene, respectively. Predicted MBD proteins were characterized as part of our phylogenetic analyses. According to this tree, (1) DNMT1, DNMT2/TRDMT1, and MBD4/MeCP2 would have been lost in an ancestor of Myxozoa (red circle); (2) DNMT3 and TET proteins would have been lost in a common ancestor of Myxozoa and Polypodium hydriiforme (orange
circle); (3) MBD1/2/3 would have been lost in a common ancestor of Myxosporea (black circle); and (4) METLL4 would have been lost in an ancestor of *P. hydriforme* and in an ancestor of *Buddenbrockia plumatellae*. It should be noted, however, that only transcriptomic data are available for *P. hydriforme*, and that only sparse EST data are available for Malacosporea (asterisks represent species with only transcriptomic or EST data), and thus it is possible that all relevant genes may be present in these species and that DNMTs, MBDs and TETs would have been lost in the branch preceding the diversification of Myxosporea. Question marks indicate uncertainties regarding the absence of genes in that species due to incomplete genome data. The “2” indicates that the gene encoding MBD1/2/3 is duplicated in *Dendronephthya gigantea*. The last column corresponds to the presence and absence of DNA cytosine methylation in each species (or in another species of the same genus); the absence of a square indicates that neither a methylome nor the absence thereof has been reported. Cytosine methylation data were obtained from Zemach et al. 2010 (*Nematostella vectensis*), Hassel et al. 2010 (*Hydra*), Putnam et al. 2016 (*Pocillopora damicornis*), Groves et al. 2016 (*Acropora millepora*), Liew et al. 2018 (*Stylophora pistillata*), Li et al. 2018 (*Exaiptasia pallida*), and the current study (*Ceratonova shasta* and *Henneguya salminicola*).
Figure 2: Cytosine methylation in Ceratonova shasta.
Histograms of fractional methylation (ratio of the number of methylated reads to the total number of methylated and unmethylated reads) for the CpG, CHG, and CHH methylation contexts for the alignment of the deduplicated final filtered C. shasta reads to the C. shasta assembly (top row). Histograms of read coverage per base for three different methylation contexts for the alignment of the final filtered C. shasta reads to the C. shasta assembly (bottom row).
Figure 3: Cytosine methylation in *Henneguya salminicola*.
Histograms of fractional methylation (ratio of the number of methylated reads to the total number of methylated and unmethylated reads) for the CpG, CHG, and CHH methylation contexts for the alignment of the deduplicated final filtered *H. salminicola* reads to the *H. salminicola* assembly (top row). Histograms of read coverage per base for three different methylation contexts for the alignment of the final filtered *H. salminicola* reads to the *H. salminicola* assembly (bottom row).
SUPPLEMENTARY FILES
Supplementary File 1. DNMT1 alignment (FASTA format).
Supplementary File 2. DNMT1 alignment filtered using GUIDANCE2 (FASTA format).
Supplementary File 3. DNMT3A alignment (FASTA format).
Supplementary File 4. DNMT3A alignment filtered using GUIDANCE2 (FASTA format).
Supplementary File 5. MBD2 alignment (FASTA format).
Supplementary File 6. MBD2 alignment filtered using GUIDANCE2 (FASTA format).
Supplementary File 7. MBD4 alignment (FASTA format).
Supplementary File 8. MBD4 alignment filtered using GUIDANCE2 (FASTA format).
Supplementary File 9. METTL4 alignment (FASTA format).
Supplementary File 10. METTL4 alignment filtered using GUIDANCE2 (FASTA format).
Supplementary File 11. TET alignment (FASTA format).
Supplementary File 12. TET alignment filtered using GUIDANCE2 (FASTA format).
Supplementary File 13. TRDMT1 alignment (FASTA format).
Supplementary File 14. TRDMT1 alignment filtered using GUIDANCE2 (FASTA format).
Supplementary File 15. HMM profiles (text file).