Long-Term m5C Methylome Dynamics Parallel Phenotypic Adaptation in the Cyanobacterium *Trichodesmium*

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Abstract

A major challenge in modern biology is understanding how the effects of short-term biological responses influence long-term evolutionary adaptation, defined as a genetically determined increase in fitness to novel environments. This is particularly important in globally important microbes experiencing rapid global change, due to their influence on food webs, biogeochemical cycles, and climate. Epigenetic modifications like methylation have been demonstrated to influence short-term plastic responses, which ultimately impact long-term adaptive responses to environmental change. However, there remains a paucity of empirical research examining long-term methylation dynamics during environmental adaptation in nonmodel, ecologically important microbes. Here, we show the first empirical evidence in a marine prokaryote for long-term m5C methylome modifications correlated with phenotypic adaptation to CO2, using a 7-year evolution experiment (1,000+ generations) with the biogeochemically important marine cyanobacterium *Trichodesmium*. We identify m5C methylated sites that rapidly changed in response to high (750 μatm) CO2 exposure and were maintained for at least 4.5 years of CO2 selection. After 7 years of CO2 selection, however, m5C methylation levels that initially responded to high-CO2 returned to ancestral, ambient CO2 levels. Concurrently, high-CO2 adapted growth and N2 fixation rates remained significantly higher than those of ambient CO2 adapted cell lines irrespective of CO2 concentration, a trend consistent with genetic assimilation theory. These data demonstrate the maintenance of CO2-responsive m5C methylation for 4.5 years alongside phenotypic adaptation before returning to ancestral methylation levels. These observations in a globally distributed marine prokaryote provide critical evolutionary insights into biogeochemically important traits under global change.

Key words: microbial evolution, global change, ocean acidification, methylation, marine microbiology, nitrogen fixation.

Introduction

Epigenetic modifications like methylation have been shown to regulate myriad transcriptional, translational, and phenotypic processes without a change in the underlying DNA sequence (Kronholm et al. 2017). Epigenetic regulation can mediate phenotypic plasticity in response to environmental change (West-Eberhard 2005; Nishikawa and Kinjo 2018). Phenotypic plasticity can be defined as a short-term change in phenotype in response to a perturbation without a change in the underlying genotype. In contrast, evolutionary adaptation results from a selective change in phenotype derived from an underlying change to the genetic (allelic) composition of a population as a result of natural selection (Collins et al. 2013; Lenski 2017). Environmentally induced epigenetic modifications of nucleotides or peptides that drive differentiated plastic phenotypes can serve as molecular proxies for adaptation through different evolutionary processes (Kronholm and Collins 2015; Schaum et al. 2016; Kronholm et al. 2017; Nishikawa and Kinjo 2018). This is especially important because the two processes can be linked, where epigenetics can “lead the way” for genetic adaptation to occur. For example, an environmentally induced, nongenetic change in phenotype can become genetically encoded during the course of selection, accompanied by a loss of environmental sensitivity in the adapted phenotype (Stajic et al. 2019). This evolutionary mechanism is known as genetic assimilation (Ehrenreich and Pfennig 2015; Danchin et al. 2019).

Regardless of mechanism, the selective fixation of beneficial traits in microbial populations is heavily influenced by the strength of selective pressure in a new environment, the exposure time relative to generation time, and the standing genetic variation. Many large-scale environmental sequencing studies examining marine microbial diversity have demonstrated local selection by environmental and ecological
gradients driven by ocean circulation among globally distributed populations (Whittaker and Rynearson 2017; Delmont et al. 2019; Salazar et al. 2019). Understanding how timescales of physicochemical gradients drive evolution is critical to be able to predict how global biogeography will evolve as global change factors increase in speed and frequency (Hutchins and Boyd 2016; Boyd et al. 2018). Although these studies paint an extensive picture of biogeographic patterns relative to the global seascape, they offer little insight into specific evolutionary mechanisms that give rise to these patterns over time. As marine microbial populations can be exposed to one or many selection events through factors like ocean upwelling and/or advection (Doblin and van Sebille 2016), evolutionary processes have the potential to respond either as a relatively isolated incident, or in succession. Furthermore, several long-term evolution studies have revealed that short and long-term microbial responses to global change factors can change in magnitude and sign (Schluter et al. 2016; Schaum et al. 2018). Hence, it is important to study the molecular underpinnings connecting short- and long-term environmental responses to help explain large-scale biogeographic patterns shaped by local environmental gradients and how they may evolve due to global change.

Several recent empirical studies in model organisms such as yeast have demonstrated direct evidence for epigenetically mediated adaptive responses (Danchin et al. 2019; Stajic et al. 2019), whereas a collection of genomic studies have shown widespread epigenetic potential across diverse microbial phyla including cyanobacteria (Blow et al. 2016; Walworth et al. 2017; Hu et al. 2018; Gärtner et al. 2019). To address the lack of long-term studies examining marine microbial evolutionary epigenetics, we investigated m5C methylation dynamics during CO2 adaptation in a 7-year experimental evolution study of the biogeochemically critical cyanobacterial diazotroph *Trichodesmium erythraeum* IMS101 (hereafter IMS101) under future carbon dioxide (CO2) concentrations (Hutchins et al. 2013; Walworth et al. 2015; Walworth, Lee, et al. 2018). *Trichodesmium* is globally distributed and among the most important contributors to bioavailable nitrogen in the oligotrophic oceans, with some estimates suggesting it makes up as much as half of total N2 fixation in the subtropical gyre regions (Hutchins et al. 2015). It excretes organic metabolites and is a source of the relatively scarce organic coenzyme, cobalamin (Frischkorn et al. 2018; Walworth, Lee, et al. 2018). It can form extensive, recurring blooms and serves as a substantial source of fixed carbon and nitrogen for new and recycled production (Walworth et al. 2015; Lee et al. 2017). Strain IMS101 was originally isolated from the Gulf Stream off North Carolina and has since been maintained in laboratory cultures (Walworth et al. 2015). It remains a challenge to maintain in culture due to its inability to survive cryopreservation, thereby requiring numerous backup cultures to be maintained per biological replicate of each experimental treatment. IMS101, like other *Trichodesmium* isolates, has a large genome relative to other cyanobacteria at 7.75 Mb with only 60% of it coding for proteins (Walworth et al. 2015). It has no detectable plasmids, and large intergenic spaces (~500-bp median length). IMS101 has over 5,000 protein coding genes and is littered with genome-wide transposons, introns, and noncoding RNAs (Pfreundt et al. 2014).

Due to the effects of epigenetically regulated phenotypic variation in response to environmental change in eukaryotes coupled to the widespread conservation of m5C methylation in prokaryotes (Blow et al. 2016), we hypothesized that m5C methylation may play a role in adaptation to a novel environment such as elevated CO2. Indeed, a few short-term studies have pointed to a role for multiple epigenetic controls in the acclimation of model cyanobacteria and diatoms to nitrogen stress and high-CO2, respectively (Hu et al. 2018; Huang et al. 2019). Furthermore, we and others recently demonstrated correlative evidence of m5C methylation potentially impacting transcription in IMS101 (Walworth et al. 2017) and the marine diatom *Phaeodactylum tricornutum* (Veluchamy et al. 2013). However, long-term adaptation and methylation has not yet been addressed. Other recent studies have demonstrated transcriptional and physiological evidence for adaptation (e.g., genetic assimilation) in multiple filamentous cyanobacteria (Walworth, Lee, et al. 2016; Koch et al. 2017). Adaptation is typically inferred through the evolutionary shift of reaction norms that underlie phenotypic expression (Ehrenreich and Pfennig 2015), where reaction norms are defined as short-term trait responses of a given genotype (e.g., N2 fixation and growth) as a function of two or more environments (Walworth, Lee, et al. 2016). Previously, we showed phenotypic evidence of genetic assimilation following high CO2 selection in *Trichodesmium* through evolutionary shifts in reaction norms (Walworth, Lee, et al. 2016), even after the high-CO2 adapted cell lines were grown in the ambient CO2 (ambient) environment for >2 years. Here, we build on this work by investigating the m5C methylome under long-term selective pressure to help elucidate m5C methylation dynamics to long-term high CO2 in the biogeochemically important diazotroph *Trichodesmium*. Theoretical and empirical studies examining epigenetic dynamics underlying adaptation have shown epigenetic modifications including m5C methylation to be influential early in the adaptive process, followed a later reduction in fitness effects (Schmitz et al. 2011; Kronholm and Collins 2015; Stajic et al. 2019; Walworth et al. 2020). Throughout our previous studies demonstrating phenotypic adaptation in IMS101, we hypothesized that epigenetic modifications would exhibit greater variation earlier in the adaptive process. Although various types of epigenetic modifications could influence CO2 adaptation, bisulfite sequencing was the only method that enabled us to examine epigenome-wide methylation dynamics with sufficient sequencing depth using minimal starting material (nanograms of DNA) given the culturing constraints outlined above. Since we demonstrated both robust physiological and transcriptional patterns consistent with genetic assimilation to high CO2, we leveraged bisulfite sequencing to examine if m5C methylation patterns are consistent with these trends including exhibiting more epigenetic activity early in an adaptive walk. We identified differentially methylated (DM) sites that rapidly changed in response to high-CO2 exposure during a short-term, plastic response,
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Results and Discussion

Long-Term Methylation Dynamics

As discussed above, we previously demonstrated physiological and transcriptional evidence of genetic assimilation following long-term high-CO2 exposure (~850 generations) in *Trichodesmium* IMS101 (Hutchins et al. 2015; Walworth, Lee, et al. 2016). We originally started with a single population of IMS101 and split it into an ambient-selected (380 μatm; Amb) and high-selected (750 μatm; High) experimental CO2 treatment with six biological replicates each (six flasks per treatment). At the time of the start of the experiment, 380 μatm CO2 represented the present day (i.e., ambient) concentration in Earth’s atmosphere, whereas 750 μatm CO2 represented the future predicted atmospheric CO2 concentration in the year 2100 under business-as-usual conditions (Hutchins et al. 2007). A diagram of the experimental design can be viewed in supplementary figure S1, Supplementary Material online. Increased growth and N2-fixation rates induced after several weeks of exposure to 750 μatm CO2 (i.e., plastic response; Ambient-High or Amb-High) were maintained 4.5 years later (fig. 1a). When high-CO2 lines were returned to the 380 μatm CO2 condition at this time-point, their increased growth and N2-fixation rates were retained, demonstrating a loss of CO2 sensitivity (fig. 1a; High-Ambient or High-Amb). We then sampled the original experimental cell lines maintained in semicontinuous batch culture under 380 and 750 μatm CO2 at a final ~7-year time point (~1,000+ generations), which showed high-CO2 growth and N2-fixation rates to still be significantly greater than those of ambient cell lines (P < 0.05, Student’s t-test, n = 6; fig. 1b; no reciprocal transplant experiment was carried out at this time point due to loss of the cultures) (Walworth, Fu, et al. 2016, 2018; Lee et al. 2018). Furthermore, fold changes in trait values between the amb-CO2 and high-CO2 conditions remained conserved across 7 years of selection (supplementary fig. S2, Supplementary Material online).

In a prior study, we used bisulfite sequencing to characterize the genome-wide m5C methylation landscape of IMS101 and its biogeographic conservation across spatiotemporally separated isolates (Walworth et al. 2017). This method is specifically designed to investigate a type of modification known as m5C cytosine methylation, defined as the addition of a methyl group to the C5 carbon residue. *Trichodesmium*
and other cyanobacteria contain the genomic capacity for other epigenetic modifications (Zehr et al. 1991; Blow et al. 2016), but this method does not capture those dynamics. To examine if genome-wide, site-specific methylation levels ($S_{\text{met}}$; number of reads with a methylated cytosine/total number of reads mapping to that cytosine) significantly changed in response to both short- (Amb-High; several weeks) and long-term (High; 4.5 years) CO$_2$ exposure, we compared genome-wide methylation changes among experimental treatments using a modified version of the RADMeth algorithm with conservative thresholds (see Materials and Methods) (Dolzhenko and Smith 2014). From this, 967 and 1,136 DM sites were identified in Amb-High and High-CO$_2$ treatments, respectively, relative to those maintained under constant 380 µatm CO$_2$. We also bisulfite-sequenced the original population from which all cultures originated (Amb-ancestral) to examine if methylation levels of environmentally or growth rate-responsive DM cytosines were more similar to those of ambient or high-CO$_2$ conditions at the 4.5-year mark. We note that at this step, we simply performed pairwise differential methylation analysis to detect DM sites but had no knowledge if site-specific methylation levels had increased or decreased in high-CO$_2$ conditions relative to ambient CO$_2$. All DM cytosines resided within the CpG motif with 44% DM on one strand and 56% on both strands (supplementary file 1, Supplementary Material online). Of the DM sites methylated on one strand ($n = 800$), we did not observe a strand preference as 49% and 51% were methylated on the sense and antisense strand, respectively. Overall, methylated cytosines have an average $S_{\text{met}}$ of 74% (median = 84%), and overall DM sites show widespread reductions in $S_{\text{met}}$ levels with an average of 53% (median = 57%).

We also checked for genetic changes shared by the high-CO$_2$ adapted cell lines (High at 4.5 years, High-Amb at 4.5 years, and High at 7 years) using the BS-SNPer pipeline for bisulfite-seq data (Gao et al. 2015) and found 52 mutations (supplementary file 1, Supplementary Material online). Of the 52 mutations, 40 occurred in only 1 high-CO$_2$ adapted sample, 7 occurred in 2 high-CO$_2$ adapted samples, 4 in 3 high-CO$_2$ adapted samples, and 1 in 4 high-CO$_2$ adapted samples. None occurred across all high-CO$_2$ samples ($n = 9$) nor did any occur in the genes or promoter regions of the five identified DNA-cytosine methyltransferases (Walworth et al. 2017). Additionally, no changes in expression levels were observed across treatments for any MTase genes (supplementary file 2, Supplementary Material online). Due to the limitations of short-read, bisulfite-seq data, and the numerous transposable elements found genome-wide in Trichodesmium (Walworth et al. 2015), other larger genomic changes may have occurred in the high-CO$_2$ selected cell lines but were undetected. Additionally, high-CO$_2$ selected cell lines may exhibit convergent phenotypes derived from different underlying genetic changes as previously observed in other microbial experimental evolution studies (Papadopoulos et al. 1999; Fong 2005). Here, we focus on long-term m5C methylome dynamics as they relate to phenotypic and transcriptomic changes.

We wanted to first ensure that comparative analysis among treatments was conducted on those methylation sites that exhibited significant pairwise differences between a high-CO$_2$ treatment and the ambient reference condition (see Materials and Methods). We identified 686 DM cytosines that met this criterion among the high-CO$_2$ treatments (Amb-High and High) and extracted the corresponding methylation levels from the ambient and 380-ancestral conditions. Bray–Curtis dissimilarities were calculated across all experimental treatments and hierarchical clustering with multiscale bootstrap resampling (replicates = 1,000) was performed to explore the possibility of treatment-specific methylation patterns. The 380-ancestral and ambient cultures formed 1 high-confidence cluster that significantly segregated (Approximately Unbiased P-value [AUP] > 0.95) (Suzuki and Shimodaira 2006) from the other cluster comprised the high-
CO₂ phenotypic lines (Amb-High and High replicates) (fig. 2). These analyses provide robust evidence that methylation changes of these DM sites are significantly correlated to changes in CO₂ conditions, considering that the ambient and 380-ancestral cell lines were sampled 4.5 years apart. The formation of these clusters using this clustering method rejects the hypothesis that these clusters do not exist at the 0.05 significance level (i.e., AUP > 0.95). Thus, this analysis provides strong statistical support that specific m5C methylation levels exhibit a significant short-term response (Amb-High) followed by long-term maintenance (High) to either CO₂ growth rate, or both, consistent with previously identified phenotypic and transcriptional patterns (Walworth, Lee, et al. 2016).

Interestingly, methylation levels in the plastic (Amb-High) and adaptive (High) treatments mostly exhibited reductions in the high-CO₂ environment (fig. 2; supplementary file 1, Supplementary Material online). We tested the probability of sharing these DM sites between the Amb-High and High treatments by chance through a pairwise hypergeometric test between DM pools. We observed a very low probability (P < 10⁻⁶) that these DM sites were shared by chance alone, suggesting them to be nonrandomly associated with both the plastic and adaptive responses to high-CO₂.

Next, we analyzed these shared DM sites’ methylation levels in steady-state High cell lines transplanted back to 380 µatm CO₂ levels for 2 years (High-Amb). Interestingly, two of the High-Amb biological replicates significantly grouped with the high-CO₂ conditions (Amb-High and High) whereas one replicate clustered with ambient treatments (Ambient and Amb-ancestral; fig. 2). Given that we sampled all selection treatments for methylation after CO₂ adaptation (Hutchins et al. 2015; Walworth, Lee, et al. 2016), significant segregation between the High-Amb replicates suggests that exposing High cell lines to their ancestral CO₂ conditions may have induced the return of CO₂ responsive methylation modifications to their ancestral methylation levels, albeit at different rates. These differing rates may be due to differences in underlying genetic factors interacting with the ambient environment (e.g., genetic drift and/or genetic divergence) after 4+ years of selection at high-CO₂, but future studies are needed to confirm this hypothesis.

This separation may also be due to our time of sampling. The exact timing of adaptation in the High cell lines before our 4-year sampling period is unknown. If adaptation occurred much earlier than our sampling time point, then methylation levels may have been receding for some time at differing rates. Had we performed the High-Amb transplantation experiment just after high-CO₂ adaptation (assuming it occurred at a similar time in all replicates), the High-Amb biological replicates may have still had enough methylation levels to cluster together mirroring their High counterparts, as they are technically the same cell lines. Regardless, if environmentally induced methylation changes are primarily influential early in an adaptive walk, and then regress following adaptation as evidenced here and in other systems (Schmitz et al. 2011; Kronholm and Collins 2015; Danchin et al. 2019; Draghi 2019; Stajic et al. 2019), then methylation levels should eventually return to ancestral conditions. This should occur irrespective of the ambient or high-CO₂ environment (i.e., either in High-Amb or High), with the former occurring at a faster rate, as we observed (fig. 2).

Moreover, the 380-ancestral and ambient cell lines are significantly more similar to each other than those of the short- (several weeks; Amb-High) and long-term (4.5 years; High) high-CO₂ treatments. Taken together with the maintenance of specific High-Amb methylation profiles, these data provide evidence for long-term m5C methylation during adaptation in this cyanobacterium, similar to previous epigenetic dynamics during adaptation in eukaryotic model systems (Danchin et al. 2019; Stajic et al. 2019). Numerous High-Amb DM sites also remained differential methylated even after ambient CO₂ exposure (fig. 2) for several hundred generations, further supporting the notion that DM sites are specific to the phenotype rather than to mere changes in carbonate chemistry. However, more research is needed to determine if methylation plays a causative role in mediating the CO₂ adaptive response in *Trichodesmium*.

The vast majority of the 686 shared DM sites between the Amb-High and High lines (supplementary file 1, Supplementary Material online) were distributed among 196 genes containing varying amounts of DM sites (1–11 DM sites per gene) in both treatments (fig. 3a; supplementary file 2, Supplementary Material online). To examine if gene-bound, shared-DM sites were overrepresented in any particular metabolic pathways, hypergeometric tests with Benjamini and Hochberg’s FDR correction (Benjamini and Hochberg 1995) (FDR < 0.05; see Materials and Methods) were conducted with Gene Ontology (GO) categories (The Gene Ontology Consortium 2018) as previously described (Walworth et al. 2017). Significantly enriched GO categories included pathways involved in select core metabolic processes including cell signaling, protein metabolism, membrane processes, and cobalamin metabolism (fig. 3a). *Trichodesmium* was recently demonstrated to be a potential important source for the often limiting coenzyme cobalamin (vitamin B12) in the open oceans (Frischkorn et al. 2018; Walworth, Lee, et al. 2018). Although methylation changes are enriched in these central cellular processes, more targeted studies are needed to investigate the mechanistic roles of methylation changes within genes in these pathways.

**Correlating Methylation to Transcription**

Next, we examined the dynamics between methylation changes and gene expression to identify potential methylated sites correlated to transcriptional changes. We must note here that our data below between methylation and expression levels are strictly correlative and solely identify interesting targets for future study. Although m5C methylation is generally widespread throughout microbial genomes, some evidence suggests that it may only indirectly or directly influence a small fraction of transcription in cyanobacteria, and therefore must be involved in alternative roles like restriction—modification systems (Blow et al. 2016) or those that have yet to be characterized (Walworth et al. 2017; Hu et al. 2018; Gartner et al. 2019). In a previous study, we observed
significantly reduced transcript levels in genes exhibiting relatively higher methylation density ($R_{\text{met}}$; number of methylated cytosines/total base pairs per gene) within the gene body at $R_{\text{met}}$ levels approaching 0.01 (Walworth et al. 2017). We also conducted this analysis for cytosine-specific methylation levels ($R_C$; number of methylated cytosines/number of per-base cytosines), which yielded the same results as $R_{\text{met}}$ and $R_C$ values show strong correlation ($R^2 = 0.89$). This is generally consistent with observations in eukaryotic algae, where transcriptional repression was observed in genes with higher $R_{\text{met}}$ (Veluchamy et al. 2013). To first scan all shared intragenic DM sites, we plotted the length- and depth-normalized mRNA expression levels (see Materials and Methods) of all 196 shared genes containing DM sites as a function of CO$_2$ condition, which yielded no significant difference between the expression pools (fig. 3b; Kruskal–Wallis test; $P > 0.05$)—supporting prior evidence that cyanobacterial methylation may only influence a small fraction of gene expression (Walworth et al. 2017; Hu et al. 2018; Gärtner et al. 2019).

Next, we searched for DM sites within genome-wide promoters ($n = 6,080$) that were experimentally verified in IMS101 (Pfreundt et al. 2014) and found five promoters containing DM sites (supplementary file 3, Supplementary Material online). Every promoter only had one DM site except for the promoter of the Tery_4419 gene annotated as a hypothetical protein, which had two DM sites in the promoter region and one DM site in the gene body. Four out of the five sites within these promoters including those of Tery_4419 were DM in both Amb-High and High-selected conditions (supplementary file 3, Supplementary Material online). However, Tery_4419 was the only one gene that also exhibited differential expression across both of the high-CO$_2$ conditions (fig. 4a). Similar to increasing methylation density correlated with transcriptional changes (e.g., $R_{\text{met}}$), IMS101 genes may need to contain a certain density of m5C DM sites across the promoter and/or the gene body (see below) to influence transcription. More targeted research is needed in this area.

To investigate increasing gene-body methylation, we binned the 196 shared genes by their $R_{\text{met}}$ levels across both ambient and high-CO$_2$ treatments, respectively, and plotted their length-normalized transcription as a function of $R_{\text{met}}$ (fig. 3c). We experimented with different numbers of
bins until a specific bin (bin 5) with a certain \( R_{\text{met}} \) range showed significantly different median expression versus all other bins (Kruskal–Wallis, post hoc Dunn test with Benjamini–Hochberg correction; FDR <0.05). Since it is unknown which of these 196 genes’ expression may at least be partially influenced by methylation, this method allowed us to use a sliding window of methylation density to search for a set of genes that exhibited consistently different expression changes associated with significantly higher methylation density. Consistent with previous findings (Veluchamy et al. 2013; Walworth et al. 2017), we observed gradual transcriptional reduction with increasing methylation density. The significantly reduced median expression level in the final bin (bin 5) with the highest \( R_{\text{met}} \) levels \( (R_{\text{met}} \geq 0.01) \) relative to all other bins suggests a nonrandom population-wide transcriptional reduction for this pool (fig. 3c). Although bin 5 exhibits the strongest statistical evidence for at least partial transcriptional reduction under increased methylation density \( (n = 39; \text{supplementary file 2, Supplementary Material online}) \), it is worth noting that genes in other bins \( (e.g., 2, 3, \) and 4) also display a similar although nonsignificant trend \( (\text{fig. 3c}) \). Here, we focus on representative genes in bin 5 as candidates that demonstrate consistently different expression changes associated with higher methylation density and leave genes in other bins for future studies. Although the significantly increased methylation levels in these genes are significantly correlated to corresponding reductions in their expression, it is of course possible that methylation may play alternative or indirect roles in addition to, or rather than, directly impacting transcription.

**Fig. 4.** Gene-specific transcript profiles containing DM sites in high-CO\(_2\). Methylation levels \( (S_{\text{met}}) \) no. of reads with a methylated cytosine/total no. of reads mapping to that cytosine) of intragenic/promoter DM sites (left axis) and expression level of the corresponding gene (right axis). (a) Methylation (boxplot) and transcriptional (bar plot) profiles of a gene (Tery_4419) containing 2 DM sites in its promoter and 1DM site in the gene body while also exhibiting differential expression across the high-CO\(_2\) conditions. (b) An intragenic DM site exhibiting significantly reduced methylation levels \( (S_{\text{met}}) \) across high-CO\(_2\) phenotypes (yellow bar; left y-axis) in concert with significantly increased transcript levels of the gene Tery_2760 (blue bars, right y-axis). (c) Methylation (boxplot) and transcriptional profiles (bar plot) of a gene (Tery_2481) containing DM sites and upregulated expression. This gene contains 19 intragenic DM sites and the clear boxplots show the distribution of \( S_{\text{met}} \) methylation levels of all sites across experimental conditions demonstrating reduced methylation in high-CO\(_2\) phenotypes (left y-axis). The bar plots show the corresponding gene expression level (right y-axis) exhibiting highest expression in high-CO\(_2\) phenotypes. (d) This plot is the same as in (b) except now there are two intragenic DM sites (light blue and orange bars) plotted adjacent to the corresponding transcript levels of that gene (dark blue bars). In transcript plots (a–d), no transcript data are available for the 380-ancestral. Bars and error bars in bar plots are means and standard deviations of three biological replicates for methylation levels and two biological replicates for RNA-Seq. Genes inside of overrepresented GO pathways are denoted above the plots.
Focusing on genes in bin 5, all intragenic methylation levels exhibited significant reductions going from ambient to high CO$_2$. Hence, we next examined if the pooled transcription of these genes displayed an average increase going from ambient to high CO$_2$. Upon plotting the normalized expression of upregulated genes as a function of treatment within bin 5 (supplementary file 2, Supplementary Material online), we observe a gradual increase in expression going from the ambient environment to the plastic (Amb-High) and then the adapted cell lines (High) (fig. 3d). The High-Amb lines are the High cell lines moved to the ambient environment. Indeed, the High-Amb expression pool exhibited slightly lower average expression values relative to the High treatment (fig. 3d).

This is possibly due to divergence of the High-Amb biological group with the ambient CO$_2$ cell lines (fig. 2). Despite this, the Amb-High, High, and High-Amb treatments share statistically indistinguishable phenotypes (i.e., growth and N$_2$ fixation; fig. 1) (Walworth, Lee, et al. 2016). Interestingly, High average transcription levels are typically highest for these 196 genes (fig. 3d) and lowest for intragenic DM sites relative to other treatments (e.g., fig. 4). However, with population-wide transcriptomics alone, only correlation and not causation can be inferred to link these methylation and transcriptional changes.

Accordingly, our primary aim was to identify gene targets and enriched pathways exhibiting significant transcriptional shifts that robustly correlate to significant intragenic methylation shifts for further targeted studies. To do this, we looked for significant differences in both expression and methylation through independent pairwise analyses. We screened for genes in bin 5 exhibiting significant upregulated expression (see Materials and Methods) in at least one high-CO$_2$ condition (Amb-High or High) while also retaining the same sign in the other corresponding high-CO$_2$ condition relative to the ambient treatment. We also ensured that selected genes harbored both significantly DM sites and $R_{net} \geq 0.01$. From these criteria, 17 candidates emerged (supplementary table S2 and file 2, Supplementary Material online). Roughly half of these ($n = 8$; supplementary table S2, Supplementary Material online) appeared within overrepresented GO pathways containing significant methylation shifts (see above) including protein binding (GO: 0005515), membrane processes (GO: 0016021), and proteolysis (GO: 0006508). These genes, and particularly the eight in GO-enriched pathways, are thus strong candidates for more targeted studies as significant molecular activity was detected from independent methods (i.e., transcriptomics and metatranscriptomics) and several independent statistical tests (supplementary file 2, Supplementary Material online). Figure 4 shows four of these genes exhibiting strikingly consistent methylation and transcriptional patterns.

For example, we observed a gene annotated as a tetratricopeptide repeat (TPR) that harbored 19 DM sites with reduced methylation levels and upregulated mRNA expression (fig. 4c; supplementary files 1 and 2, Supplementary Material online). TPR domains are found across all domains of life and participate in a variety of functions, ranging from cell division to RNA metabolism to protein transport. They have garnered recent interest for the discovery of their role in photosynthetic metabolism, particularly in the assembly and stability of photosystems I and II (Klinkert et al. 2004). In the cyanobacterium Synechocystis, a membrane-bound TPR protein was demonstrated to be involved in the processing of the D1 reaction center (Klinkert et al. 2004) and additionally was subsequently found to play a role in photosynthetic performance and light-dependent chlorophyll synthesis (Schottkowski et al. 2009). Other genes involved in oxidative stress and photosynthetic maintenance displayed similar profiles (fig. 4b and d; see supplementary note S1, Supplementary Material online). These results are generally supportive of prior observations in Trichodesmium under both short- and long-term CO$_2$ exposure that showed significant shifts in PSI:PSII ratios and photosynthetic electron transport transcripts (Levitan et al. 2007; Walworth, Lee, et al. 2016). These data may show molecular evidence for the association of redox changes and protein turnover that potentially result in previously observed changes in PSI:PSII ratios under high-CO$_2$, which may in turn liberate iron (Fe) from the Fe—heavy PSI complex for use in N$_2$ fixation.

**Return of high-CO$_2$ Induced Methylation to Ancestral Levels**

At the 7-year mark (1,000+ generations) of this evolution experiment, we again bisulfite-sequenced the ambient and High cell lines (hereafter “ambient” and “High, respectively) and compared methylation levels with those of the Amb-ancestral and CO$_2$-selected conditions at the 4.5-year mark. Upon hierarchical clustering of Bray—Curtis dissimilarities calculated from methylation levels from the same 686 shared genes identified above across 7 years of selection, the 7-year “ambient” replicates clustered with the Amb-ancestral and 4.5-year ambient conditions as expected. However, the 7-year “High” replicates also clustered with all ambient treatments, indicating the return of methylation levels in initially responsive DM sites back to ambient ancestral levels (fig. 5).

These results are in line with prior modeling and empirical studies examining genetic assimilation theory where phenotypic fixation is observed irrespective of treatment while initial, environmentally induced nongenetic changes return to ancestral levels prior to selection (Madlung et al. 2002; Pigliucci et al. 2006; Schlichting and Wund 2014; Ehrenreich and Pfennig 2015; Kronholm and Collins 2015). The two overarching clusters are strongly statistically supported (AUP > 0.95), which rejects the null hypothesis that these clusters do not exist at the 0.05 significance level. Furthermore, no significantly DM sites were detected between the “ambient” and “High” conditions, which serves as robust evidence for statistically analogous global methylation states at the 7-year mark. This contrasts strongly with observations of significantly different methylated sites at 4.5 years of selection (fig. 2), and with the significant phenotypic differences in growth and N$_2$-fixation rates still observed after 7 years between the “ambient” and “High treatments (fig. 1). Hence, at this 7-year time point any previous phenotypic differences
Fig. 5. Hierarchical clustering (HC) with multiscale bootstrap resampling of DM sites. HC of Bray—Curtis dissimilarities calculated from shared (n = 686) DM methylation levels in figure 2. Values at nodes are AUPs. The ancestral population is labeled in bold and the 7-year time points are labeled in green. Both the plastic (Amb-High) and adaptive (High) high-CO₂ conditions at the 4.5-year mark maintain segregation from the ancestral (Amb-ancestral) and ambient conditions as in plots in figure 2. Also consistent with plots in figure 2, two of the High-Amb replicates still cluster with the 4.5-year high-CO₂ conditions, whereas one replicate clusters with the ambient conditions. This demonstrates CO₂-responsive methylation levels to be returning to ancestral conditions once placed back into the ancestral environment. Upon adding *ambient and *High selected conditions at the 7-year time point (green), the *ambient replicates cluster with the Amb-ancestral and 4.5-year ambient conditions as expected, but the *High replicates also cluster with all ambient conditions, demonstrating the return of methylation levels originally triggered by high-CO₂ back to their ancestral ambient levels.
that may have been associated with nongenetic changes are presumably now decoupled from them, potentially implicating a genetic change.

**Conclusion**

In summary, we identified m5C methylated sites that rapidly changed in response to high-CO2 exposure during a short-term, plastic response (Amb-High) relative to their levels in both the ancestral population (Amb-ancestral) and ambient selected conditions. High-CO2 methylation levels at these sites were then maintained for at least 4.5 years, during which time adaptive shifts in reaction norms were observed (High cell lines). This is evidenced by the fixation of multiple traits (i.e., growth and N2 fixation) that were retained in the High lines after being returned to a ambient environment (i.e., High-Amb). The vast majority of these DM sites were intragenic and overrepresented in select core metabolic pathways, thereby implicating m5C methylation changes to be significantly associated with these genes during high-CO2 selection. Additionally, we identified specific genes that both harbored DM sites and exhibited differential mRNA expression as candidates for potential mechanistic studies under long-term high-CO2. Their annotated functions suggest central roles in cellular redox changes and protein turnover potentially associated with photosynthetic activity. Multiple lines of evidence from other independent analyses examining *Trichodesmium* metabolism under high-CO2 have also indicated significant cellular investment toward energetic and photosystem rearrangement, which seem to facilitate increased growth and N2 fixation. Placing the high-CO2 cell lines back in the ambient environment (High-Amb) appears to influence the return of high-CO2 methylation levels back to their ancestral state with different rates among the High-Amb replicates. Upon resampling the ambient and High cell lines at the 7-year mark, no DM sites were observed between conditions, indicating that methylation levels of initially responsive DM sites had returned to their ancestral levels. This is strongly supported by the clustering of all sampled replicates from across 7 years of selection. Though more work needs to be done to determine if methylation played a causative role in the phenotype, these long-term dynamics are consistent with genetic assimilation theory wherein the adapted phenotype is maintained through the loss of environmental sensitivity while initial molecular changes return to ancestral levels in the selection environment. Further study examining methylation during adaptation in marine photosynthetic microbes will help to elucidate the importance of short-term microevolutionary processes on long-term evolutionary divergence to interacting ocean and global change dynamics. Here, we provide a first step to investigating long-term m5C methylation in conjunction with high-CO2 selection for a globally distributed marine microbe. Importantly, the long-term dynamics of these methylated sites were highly consistent with both prior observations in other systems, and those predicted by genetic assimilation theory. This work presents a framework for examining long-term roles of epigenomes as potential mechanisms in biogeochemically important trait evolution under global change.

**Materials and Methods**

**Culturing and Physiology**

Detailed culturing and physiology methods and results can be found in previously published studies (Hutchins et al. 2015; Walworth, Lee, et al. 2016; Lee et al. 2018). Briefly, stock cultures of *Trichodesmium* strain IMS101 were maintained at the University of Southern California (Los Angeles, CA) in a modified Aquil medium devoid of added combined nitrogen with standard vitamins and trace metals, which included 500 nM iron and 20 μM phosphate. Cultures were grown at 26°C under a light–dark cycle of 12:12 (LD) and a light intensity of 120 mmol photons per m2 s−1 incident irradiance. A diagram of the experimental design can be viewed in supplementary figure S1, Supplementary Material online. At the beginning of the experiment, we started with a single population of IMS101 and split it into an ambient-selected (380 μatm; Amb) and high-selected (750 μatm; High) experimental CO2 treatment. Six biological replicates per treatment were examined using semi-continuous culturing methods where each replicate was diluted individually based on the growth rate calculated for the respective replicate. For all phenotypic assays, six biological replicates were used to calculate means and standard deviations. Each replicate bottle was kept optically thin to avoid self-shading, nutrient limitation, and perturbations to targeted CO2 levels. Cultures were constantly bubbled with prepared air/CO2 mixtures (Praxair, Los Angeles, CA) to sustain stable CO2 concentrations of 380 (ambient) and 750 (high-CO2) μatm CO2 for a total of 7 years. Growth and N2 fixation rates reported in this study were measured during the middle of the photoperiod at 4.5 and 7 years, respectively, as previously described (Walworth, Fu, et al. 2016; Walworth, Lee, et al. 2016). To measure the CO2 plastic response following 4.5 years of growth at 380 μatm CO2, subcultures of ambient cell lines were transferred to 750 μatm CO2 for 2 weeks (Amb-High) and growth and N2-fixation rates were measured during the middle of the photoperiod. The correlated response (High-Amb) was measured in the same manner, except that High cell lines were transferred to 380 μatm CO2 for 2 weeks before assaying and were then maintained under the changed CO2 conditions for over 2 years. The phenotypic trends of these published experiments are presented for readers in figure 1 and supplementary figure S1, Supplementary Material online, in order to facilitate interpretation of the epigenetic results presented here.

**Nucleic Acid Sampling and Isolation for Illumina Sequencing**

Three independent biological replicates per treatment (three randomly selected separate flasks out of six total flasks per treatment) were gently filtered during the middle of the photoperiod onto 5 mm polycarbonate filters (Sigma Aldrich, St.
Trimmomatic (v0.32) (Bolger et al. 2014) using the following settings: LEADING: 3 TRAILING: 3 SLIDINGWINDOW: 4:15 MINLEN: 35. Reads were then mapped onto the IMS101 genome using Bowtie2 v2.2.5 (Langmead and Salzberg 2012) with default settings. Genes containing an average of $<10$ counts across libraries were removed. Because we compared the distribution of bins containing numerous genes, transcripts were first normalized by gene length via a “Loess regression” in the EDASeq package (Risso et al. 2011) followed by library-depth normalization using calculated size factors per library in the DESeq2 package. Finally, differential expression analysis was conducted using DESeq2 (Love et al. 2014). Genes with a Benjamini and Hochberg’s FDR $<0.05$ were deemed differentially expressed. Normalized expression levels for all detected genes can be found in supplementary file 2, Supplementary Material online. A flowchart of the analysis pipeline can be found in supplementary figure S3, Supplementary Material online.

Statistical Analysis

Hypergeometric tests were conducted as previously described (in Walworth, Fu, et al. 2016; Walworth, Lee, et al. 2016; Walworth, Fu, et al. 2018). GO annotations for *Trichodesmium* were downloaded from the Genome2D web server (http://genome2d.molgenrug.nl). The phyper function in R was used to test for significance and P values were corrected by the Benjamini–Hochberg false discovery rate (FDR) $<0.001$ (Benjamini and Hochberg 1995). A flowchart of the analysis pipeline can be found in supplementary figure S2, Supplementary Material online. Methylation levels for all detected mSC residues can be found in supplementary file 1, Supplementary Material online, along with methylation and RNA sequencing statistics. Methylation and RNA sequencing statistics can also be found in supplementary table S1, Supplementary Material online.

SNP Analysis

Quality processed reads (see above) were first aligned to the *T. erythraeum* IMS101 reference genome with Bismark software (Krueger and Andrews 2011). SNP analysis was conducted using the BS-NPPer pipeline with default settings. BEDTools was used for genome-wide comparative analysis of genetic features (Quinlan and Hall 2010) including experimentally confirmed promoter regions from Pfreundt et al. (2014).

Differential Expression Analysis

Transcript normalization and differential expression was conducted as previously described (Walworth et al. 2017). Briefly, Trimmomatic version 0.35 (Bolger et al. 2014) was used to quality trim and filter with the following settings: SE -threads 35 -phred33 LEADING: 3 TRAILING: 3 SLIDINGWINDOW: 4:15 MINLEN: 35. Processed fastq files were next mapped onto IMS101, IMG-called genes (Chen et al. 2019) using Bowtie2 v2.2.5 (Langmead and Salzberg 2012) with default settings. Genes containing an average of $<10$ counts across libraries were removed. Because we compared the distribution of bins containing numerous genes, transcripts were first normalized by gene length via a “Loess regression” in the EDASeq package (Risso et al. 2011) followed by library-depth normalization using calculated size factors per library in the DESeq2 package. Finally, differential expression analysis was conducted using DESeq2 (Love et al. 2014). Genes with a Benjamini and Hochberg’s FDR $<0.05$ were deemed differentially expressed. Normalized expression levels for all detected genes can be found in supplementary file 2, Supplementary Material online. A flowchart of the analysis pipeline can be found in supplementary figure S3, Supplementary Material online.

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discovery method (Benjamini and Hochberg 1995) with the p.adjust function ($P < 0.05$).

Bray–Curtis dissimilarities were calculated from growth- and depth-normalized methylation levels with the vegan package (Oksanen et al. 2013) and hierarchical clustering with multiscale bootstrapping ($n = 1,000$) was carried out with the pvclust package (Suzuki and Shimodaira 2006). Heatmaps were generated using the Heatplus package (Ploner 2015).

### Supplementary Material

Supplementary data are available at Molecular Biology and Evolution online.

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### Data Availability

Raw read files for Whole-genome bisulfite-sequenced libraries (WGBS) have been deposited in the NCBI’s Gene Expression Omnibus (Edgar et al. 2002) and are accessible through GEO Series accession number GSE138176. The ambient and Amb-High Raw RNA raw reads can be found in another GEO project (Ploner 2015).

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