



Nutrient-Colimited *Trichodesmium* as a Nitrogen Source or Sink in a Future Ocean

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ABSTRACT Nitrogen-fixing (N_2) cyanobacteria provide bioavailable nitrogen to vast ocean regions but are in turn limited by iron (Fe) and/or phosphorus (P), which may force them to employ alternative nitrogen acquisition strategies. The adaptive responses of nitrogen fixers to global-change drivers under nutrient-limited conditions could profoundly alter the current ocean nitrogen and carbon cycles. Here, we show that the globally important N_2 fixer *Trichodesmium* fundamentally shifts nitrogen metabolism toward organic-nitrogen scavenging following long-term high- CO_2 adaptation under iron and/or phosphorus (co)limitation. Global shifts in transcripts and proteins under high- CO_2 /Fe-limited and/or P-limited conditions include decreases in the N_2 -fixing nitrogenase enzyme, coupled with major increases in enzymes that oxidize trimethylamine (TMA). TMA is an abundant, biogeochemically important organic nitrogen compound that supports rapid *Trichodesmium* growth while inhibiting N_2 fixation. In a future high- CO_2 ocean, this whole-cell energetic reallocation toward organic nitrogen scavenging and away from N_2 fixation may reduce new-nitrogen inputs by *Trichodesmium* while simultaneously depleting the scarce fixed-nitrogen supplies of nitrogen-limited open-ocean ecosystems.

IMPORTANCE *Trichodesmium* is among the most biogeochemically significant microorganisms in the ocean, since it supplies up to 50% of the new nitrogen supporting open-ocean food webs. We used *Trichodesmium* cultures adapted to high- CO_2 conditions for 7 years, followed by additional exposure to iron and/or phosphorus (co)limitation. We show that “future ocean” conditions of high CO_2 and concurrent nutrient limitation(s) fundamentally shift nitrogen metabolism away from nitrogen fixation and instead toward upregulation of organic nitrogen-scavenging pathways. We show that the responses of *Trichodesmium* to projected future ocean conditions include decreases in the nitrogen-fixing nitrogenase enzymes coupled with major increases in enzymes that oxidize the abundant organic nitrogen source trimethylamine (TMA). Such a shift toward organic nitrogen uptake and away from nitrogen fixation may substantially reduce new-nitrogen inputs by *Trichodesmium* to the rest of the microbial community in the future high- CO_2 ocean, with potential global implications for ocean carbon and nitrogen cycling.

KEYWORDS nutrient colimitation, nitrogen fixation, *Trichodesmium*, global change, ocean acidification, microbial ecology, cyanobacteria, evolution, TMA, trimethylamine, high CO_2 , nutrient limitation, organic nitrogen

Oceanic food webs and climate feedbacks are significantly influenced by carbon dioxide (CO_2)- and nitrogen (N_2)-fixing microbes, thereby contributing to both global productivity and biogeochemistry (1–3). However, only a few studies have investigated the physiological and evolutionary responses of the globally important,

Received 27 September 2017 Accepted 16 November 2017

Accepted manuscript posted online 27 November 2017

Citation Walworth NG, Fu F-X, Lee MD, Cai X, Saito MA, Webb EA, Hutchins DA. 2018. Nutrient-colimited *Trichodesmium* as a nitrogen source or sink in a future ocean. Appl Environ Microbiol 84:e02137-17. <https://doi.org/10.1128/AEM.02137-17>.

Editor Shuang-Jiang Liu, Chinese Academy of Sciences

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photoautotrophic, N₂-fixing cyanobacterium *Trichodesmium erythraeum* strain IMS101 to high-CO₂ conditions under multiple nutrient-limiting regimes (4–6). Simultaneous iron (Fe) and phosphorus (P) limitation of IMS101 under both low- and high-CO₂ conditions has been found to yield higher growth rates, reduced cell sizes, and a unique Fe/P protein complement, compared to the limitation of either Fe or P alone (5, 7). This fitness advantage conferred by Fe/P “balanced limitation” contrasts with the long-standing Liebig limitation model (8) and has implications for global biogeochemical cycles in both the current and future oceans (3).

Oligotrophic populations are predicted to potentially experience longer periods of enhanced nutrient (co)limitation due to intensifying Fe stress under high-CO₂ conditions (5, 9) and reduced vertical P supplies from increased density-driven stratification (3, 10). However, a major unknown is the potential change in future new-nitrogen inputs by globally distributed N₂-fixing microbes (diazotrophs). Indeed, past research has demonstrated significant decreases in the growth and N₂ fixation of marine diazotrophs under Fe limitation, with molecular evidence indicating iron reallocation via reduction in the photosystem I-to-photosystem II (PSI/PSII) ratio, decreases in metalloenzyme inventories, and increases in Fe stress PS antenna abundances (5, 11–13). Additionally, *Trichodesmium* has been demonstrated to take up both inorganic (e.g., nitrate and ammonia) and organic (e.g., amino acids) nitrogen species, thereby inhibiting N₂ fixation (14, 15). However, because much of the low-latitude surface ocean is largely nitrogen limited (16), fixed N sources can be severely depleted, resulting in a marked dependence on Fe and P bioavailability to fuel N₂ fixation (1, 3).

To date, nearly all diazotrophic nitrogen assimilation research has focused on the relationship between N₂ fixation and nitrate, ammonia, and amino acid uptake, resulting in the view that ammonia is the preferred microbial nitrogen source for both diazotrophic and nondiazotrophic microbes (17). Identifying and measuring preferentially scavenged organic nitrogen species by marine microbes has been largely precluded by rapid biochemical turnover *in situ*. Furthermore, microbial consortia aggregating around microbial species of interest can also inhibit discernment of the microbial uptake of specific substrates (18).

In this study, functional genomics and molecular physiology provide strong evidence that following adaptation to high-CO₂ conditions, *Trichodesmium* allocates the greatest biosynthetic investment to the acquisition of the organic nitrogen substrate trimethylamine (TMA) and potentially other organic nitrogen- and sulfur-containing compounds. This shift in nitrogen acquisition pathways under high-CO₂ conditions is predicted to be mediated by large increases in bacterial flavin-containing monooxygenase (FMO) coupled with global shifts in transcription and translation. This indicates a fundamental change in both nitrogen and global cellular strategies whereby iron-rich biosynthetic pathways including N₂ fixation and photosynthesis are significantly reduced under Fe-limited, high-CO₂ conditions, in parallel with increased biosynthesis of the predicted TMA-oxidizing FMO enzyme. Additionally, N₂ fixation is inhibited in the presence of exogenous TMA in a manner similar to that seen with nitrate and ammonia, with TMA simultaneously supporting growth rates equivalent to those seen during N₂ fixation. Methylated amine compounds like TMA are products of protein putrefaction and degradation of quaternary amine osmoregulators (e.g., glycine betaine) (19) and are thus ubiquitous in the marine environment, representing a considerable pool of C and N with reported standing concentrations in the nano- to micromolar range (20, 21). Hence, future CO₂ levels and limiting Fe may exacerbate cellular Fe stress, resulting in a fundamental metabolic shift whereby *Trichodesmium* reallocates resources away from N₂ fixation and toward FMO-mediated organic nitrogen scavenging. The resulting metabolism change observed in *Trichodesmium* necessitates a reassessment of the relationship between new-nitrogen inputs and simultaneous removal by this globally important marine N₂ fixer, as well as a broader characterization of the utilization of methylated amines like TMA by biogeochemically critical marine microbes.

RESULTS AND DISCUSSION

Nutrient-limited physiology under increased CO₂. A comprehensive depiction of the experimental design used in this study can be found in Fig. S1 in the supplemental material, and proteomic results and their implications for nutrient limitation theory were presented previously (5). Here, we focus on steady-state, transcriptional/translational changes in nitrogen source-sink metabolism as they relate to other biogeochemically important pathways. Briefly summarizing Fig. S1, a single IMS101 population originally isolated from an individual *Trichodesmium* colony (22) was split into low-CO₂ (380 μatm of CO₂ [380-selected]) and high-CO₂ (750 μatm of CO₂ [750-selected]) treatments of six biological replicates each and grown for ~7 years (~1,000 and ~1,500 generations for the 380- and 750-selected treatments, respectively). Semicontinuous culturing methods were used in modified trace metal-clean Aquil medium (devoid of ammonia or other fixed nitrogen) containing 25 μM EDTA, standard vitamins, and trace metals with 250 nM Fe and 10 μM phosphate (PO₄³⁻). Next, Fe/P-colimited treatments were generated by subculturing three biological replicates from each CO₂ treatment and incubating them in the aforementioned medium except with colimiting levels of Fe (10 nM) and PO₄³⁻ (0.25 μM) for >1 year. To then mimic episodic pulses of Fe or P typically experienced by Fe/P-colimited microbes in the oligotrophic ocean (3, 23, 24), either Fe or P was added at replete concentrations to subcultures of the Fe/P-colimited cell lines to generate P- and Fe-limited lines at both CO₂ levels, respectively. Following a 2-month acclimation period to create steady-state conditions, the lines were sampled for molecular analyses.

Growth and N₂ fixation rates were significantly increased at high-CO₂ levels under both nutrient-replete (r750) and P-limited (750-P) conditions relative to low-CO₂/nutrient replete (r380) and low-CO₂/P-limited (380-P) treatments, respectively (Fig. 1). Conversely, no growth rate differences were observed in high-CO₂/Fe-limited (750-Fe) and high-CO₂/Fe/P-colimited (750-Fe/P) cell lines relative to the corresponding low-CO₂/Fe-limited (380-Fe) and low-CO₂/Fe/P-colimited (380-Fe/P) conditions, respectively. In a recent study, Hong et al. (9) observed a significant decrease in growth and N₂ fixation rates under both replete (r750) and Fe-limited conditions at high CO₂ concentrations (750-Fe) relative to the controls, unlike this study and a number of other previous replete (3–6, 10, 25–28) and high-CO₂/Fe-limited studies (5). Hong et al. (9) attributed these discrepancies to ammonia contamination and metal toxicity in all other past studies, although ammonia was below the detection limit in our prepared Aquil medium and 25 μM EDTA was added to control for metal toxicity (29). One central difference between this study and nearly all others relative to that of Hong et al. (9) is that *Trichodesmium* cell lines were adapted (4–6) or acclimated (10, 26–28) to a low pH as a product of increasing CO₂ partial pressure on seawater carbonate chemistry similar to pH reductions *in situ*. This contrasts with artificially controlling pH with short-term acid/base additions as in the study of Hong et al., which may also result in other unknown and uncontrolled effects on cell physiology. A second fundamental difference is that our Fe single-limitation treatments were generated by pulsing phosphorus into 7-year, high-CO₂-adapted cell lines colimited for iron and phosphorus for ~1 year, similar to persistent colimitation conditions observed in the oligotrophic ocean (23, 24).

Looking ahead, it is predicted that increasing CO₂ will impact ecosystems more rapidly than an increasing sea surface temperature (30), which in turn will affect the adaptive evolution of marine microbes and their contribution to elemental cycling. Hence, following long-term (~7-year) adaptation of IMS101 to high-CO₂ conditions (3, 5, 6), we investigated changes in IMS101 molecular physiology and nitrogen acquisition after ~1 year of exposure to Fe/P colimitation, followed by additions of either Fe or P to create steady-state Fe and P single-limitation treatments.

Variable nitrogen assimilation strategies under nutrient colimitation in a future ocean. Below, we focus on the dynamics of global transcription and translation as they relate to changing nitrogen metabolic strategies under intensifying nutrient

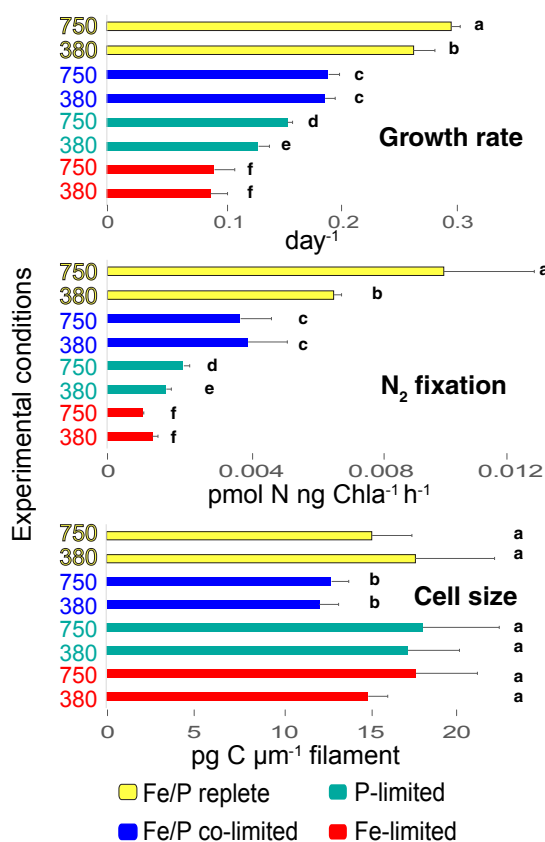


FIG 1 Cell physiology. Cell specific growth rates (day⁻¹, top), N₂ fixation rates (middle), and cell sizes (assessed by using the proxy carbon content per unit of filament length [pg of C/μm]) (bottom) are shown. Experimental conditions are shown to the left of all bars, which are color coded to indicate the nutrients used. Data are shown as means and standard errors (*n* = 6). Different letters denote statistical significance.

(co)limitation as a function of high-CO₂ adaptation. A more in-depth discussion of other molecular changes in broad biochemical pathways can be found in the supplemental material. Genes in 380-Fe that were differentially expressed (DE) compared to those in r380, but not relative to those in 380-Fe/P, showed significantly different transcript levels in both 380-Fe and 380-Fe/P, respectively, relative to r380 (Fig. 2a; see also Fig. S2a and Data Set S1). This pool was enriched in Gene Ontology (GO) pathways involving nitrogen fixation (downregulated), cytochrome *c* oxidase activity (downregulated), copper ion binding (downregulated), nitrate/nitrite transport (upregulated), respiration (downregulated), glutamate synthase (upregulated), and amino acid transport (upregulated). These pathway enrichments indicate a downregulation of Fe-heavy, N₂-fixing enzymes similar to other findings (5, 31) in conjunction with the upregulation of external nitrogen acquisition genes targeting inorganic (e.g., nitrate; Fig. 3) and/or possibly organic (e.g., amino acid transport) sources (Data Set S1).

While the 750-Fe pool of DE transcripts relative to r380 was enriched in GO pathways (Fig. S2) involved in nitrogen fixation (downregulated) and cytochrome *c* oxidase activity (downregulated), similar to 380-Fe relative to r380, genes involved in nitrate/nitrite transport had drastically lower expression levels in 750-Fe (gray bar) and 750-Fe/P (blue bar) than those in 380-Fe (orange bar) and 380-Fe/P (green bar), respectively (Fig. 3). This massive transcriptional reduction in inorganic nitrogen transport going from low- to high-CO₂ conditions under Fe limitation regardless of the P concentration also co-occurred with ~50% fewer genes being DE overall under high-CO₂/low-Fe conditions, including a considerable reduction of protein synthesis transcripts (Fig. 2b; Fig. S2b and Data Set S1).

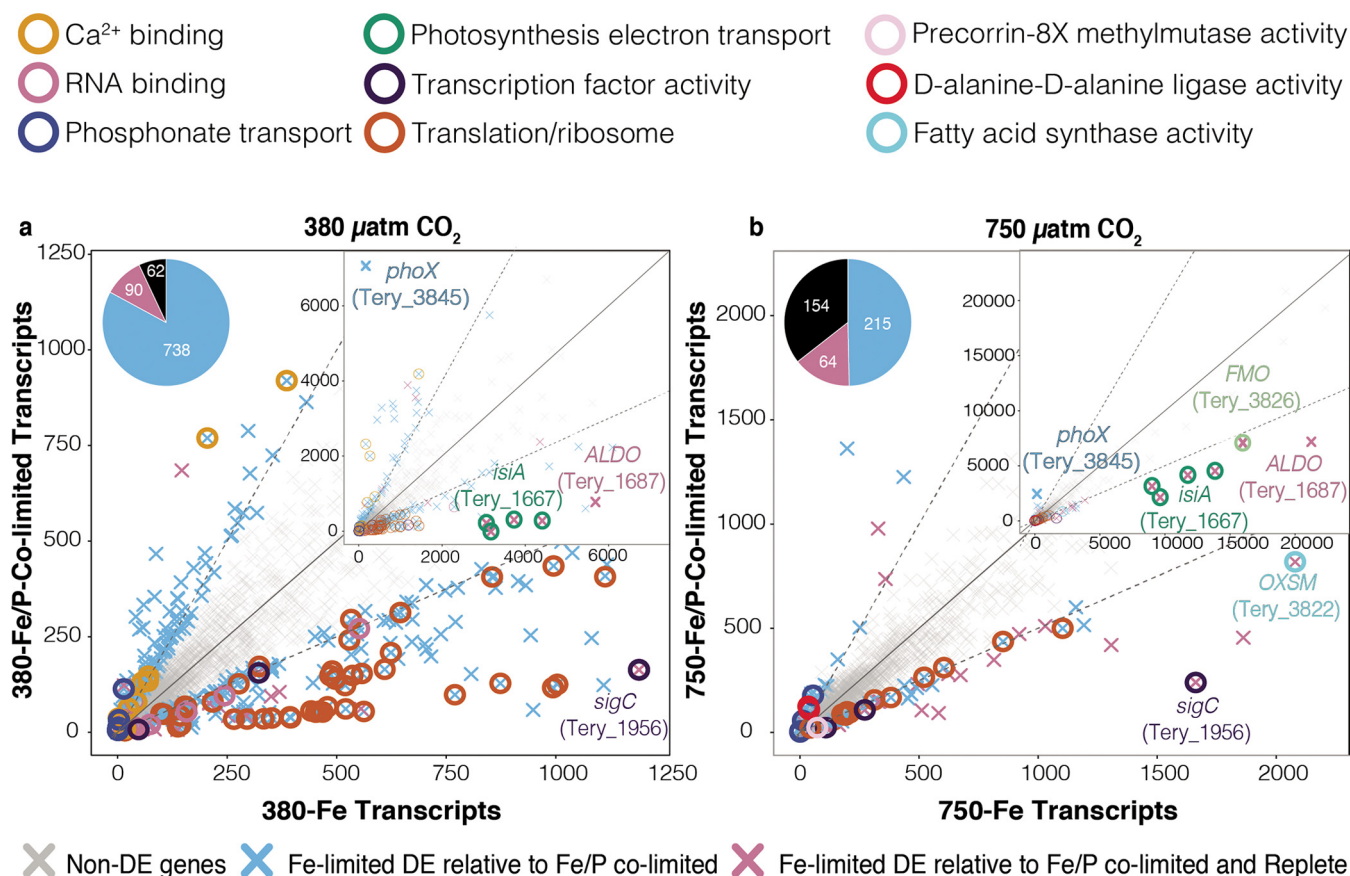


FIG 2 Fe-limited versus Fe/P-co-limited transcripts under 380 and 750 μatm of CO₂. (a) Scatterplot of 380-Fe DE genes relative to 380-Fe/P (blue) and 380-Fe DE genes relative to both r380 and 380-Fe/P (magenta). The center diagonal is a 1:1 line, while the others are 2:1 and 1:2 lines, respectively. Insets are the same plot zoomed out to display highly expressed genes. (b) Similar scatterplot of 750-Fe DE genes relative to either 750-Fe/P (blue) or both r750 and 750-Fe/P (magenta) DE genes. Select genes are labeled. The differently colored circles represent genes within significantly enriched GO pathways under the two conditions.

Interestingly, both transcript and protein abundances of a predicted TMA-oxidizing FMO (Tery_3826) harboring the FMO-specific motif (32) and a predicted transmembrane helix (according to the Integrated Microbial Genomes [IMG] annotation) sharply increased in 750-Fe (Fig. 2b and 3), along with the flavin biosynthesis gene *ribD* (Tery_1661; Data Set S1), relative to all other treatments. In fact, the FMO-encoding gene was one of the most highly expressed transcripts detected in the high-CO₂, Fe-limited treatment. Furthermore, its general expression level was orders of magnitude greater than that of other nitrogen acquisition transcripts across all treatments (Fig. 2b and 3a). Transcripts of an annotated ammonia transporter gene (*amT1*; Tery_4477) remained unchanged across treatments, but it was consistently expressed in the top ~6% of detected transcripts. Notably, although 380-Fe and 750-Fe growth rates were similar, *aldO* (Tery_1687; fructose-bisphosphate aldolase) transcript levels also drastically increased in 750-Fe along with the ferric uptake regulator (Tery_3404, *fur*) and Fe stress/photosynthesis antenna genes (Fig. 2b; Fig. S2b).

This global shift in transcript profiles, in conjunction with the downregulation of nitrate transport and the concurrent upregulation of FMO and *ribD*, suggests that high-CO₂ conditions intensify iron limitation (5, 9), resulting in fundamental resource (e.g., riboflavin) reallocation toward increased biosynthetic investment in FMO-mediated, iron-lean organic nitrogen assimilation strategies. The large increase in FMO transcripts and proteins observed in 750-Fe suggests that *Trichodesmium* preferentially scavenges TMA and possibly other organic nitrogen and sulfur compounds under severe iron stress. Furthermore, FMO transcript and protein abundances are in the top

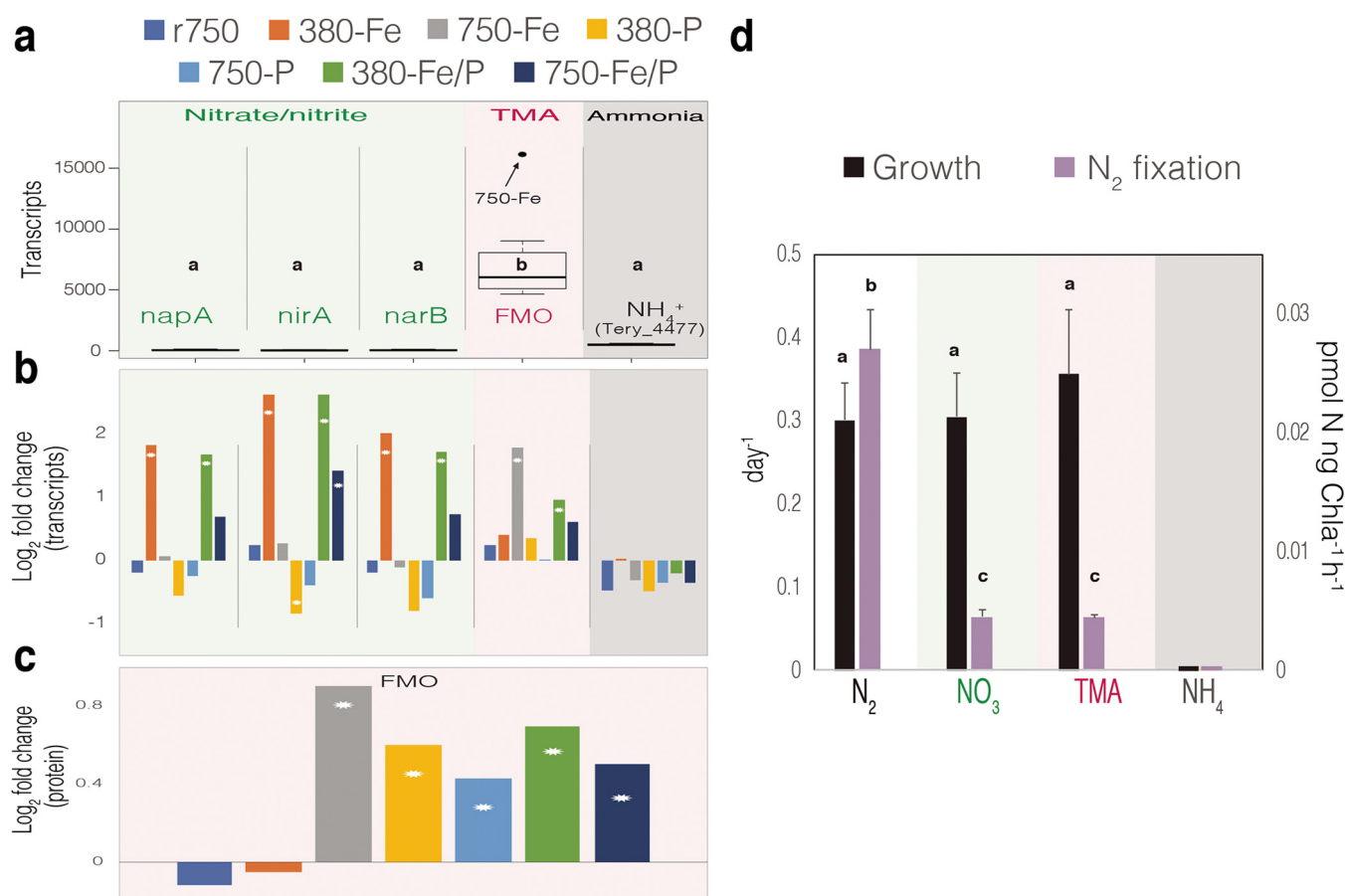


FIG 3 Distributions of nitrogen acquisition transcripts across treatments and log₂ changes in transcript and protein abundance per treatment and physiological data of IMS101 grown on several nitrogen species. (a) The distribution of transcripts per gene across all treatments. Different letters signify significantly different mean values. (b) Log₂ fold changes in gene transcript abundance relative to the r380 condition across experimental treatments. (c) Log₂ fold changes in normalized FMO protein spectral counts relative to r380. For panels b and c, stars indicate statistically significant differences from r380. (d) Growth and N₂ fixation rates of IMS101 grown without fixed nitrogen (N₂) and in the presence of NO₃, TMA, or NH₄ at 20 μM. Different letters signify significantly different mean values.

~0.5% and ~10% of all detected gene transcripts and protein products, respectively, irrespective of treatment, which indicates significant energetic investment in persistent FMO biosynthesis. Conversely, nitrate transport gene transcripts are only in the top ~50 to 70% of the total transcripts, with no corresponding protein products detected in the proteome. Upon calculation of the total mean expression of nitrogen acquisition transcripts across all treatments, the gene for FMO was consistently expressed at significantly higher levels ($P < 0.0001$) than other nitrogen acquisition genes (Fig. 3a), providing further evidence of persistent and enriched biosynthesis of FMO. Taken together with the ammonia transporter expression in the top 6% of overall transcripts, these data support the notion that FMO-acquired compounds and ammonia are preferred relative to nitrate acquisition, as previously noted for ammonia (17). The TMA protein was significantly more abundant under high-CO₂/low-Fe, low-P, and Fe/P colimitation conditions than under r380, r750, and 380-Fe conditions, suggesting enhanced energetic investment in TMA enzyme biosynthesis across various CO₂/nutrient-limited global change scenarios (Fig. 3c). It is interesting that FMO transcript and protein levels considerably increased in 750-Fe but not 380-Fe, although Fe stress genes signaled iron limitation under both conditions. Strain IMS101 N₂-fixing metabolism may be better adapted to low-CO₂/low-Fe conditions because of prevalent Fe limitation *in situ* (24), thereby reducing the need to scavenge for nitrogen, while molecular metabolic restructuring induced by high-CO₂/low-Fe conditions may require increased exogenous nitrogen for growth in a high-CO₂ ocean. Since heterotrophs have

been shown to grow on TMA as a sole C and N source (32), *Trichodesmium* may utilize TMA for both C and N (33, 34). It was also recently shown that an associated heterotrophic *Alteromonas macleodii* strain widely conserved in *Trichodesmium* metagenomic consortia has the genetic potential to cleave dimethylsulfoniopropionate to produce dimethylsulfide (DMS) (35), which may be further oxidized by the *Trichodesmium* FMO, as DMS has been shown to have nearly as high an affinity for FMO as that of TMA (32). Hence, TMA- and other FMO-mediated assimilation may be a more general compensatory strategy utilized by *Trichodesmium* for physiological maintenance in the face of fluctuating nutrients, which is evidenced by the significant increase in FMO protein abundance across nutrient-limited treatments (Fig. 3c, bottom).

N₂ fixation is strongly inhibited in the presence of 20 μ M TMA, just as it is with 20 μ M nitrate (Fig. 3d). However, *Trichodesmium* growth rates are virtually the same in the presence of 20 μ M TMA as growth rates supported by N₂ fixation, comparable to results obtained in the presence of 20 μ M nitrate, indicating that both fixed nitrogen sources can support growth as efficiently as N₂. In contrast, the presence of 20 μ M ammonia was toxic, and the cells were unable to grow or fix N₂ at all (Fig. 3d). One recent study observed enigmatically high growth and N₂ fixation rates in the presence of 20 μ M ammonia (9), contrasting with all other previous studies demonstrating drastic repression of N₂ fixation at exogenous ammonia concentrations of >10 μ M (15, 17, 36, 37).

Although Hong et al. (9) observed decreases in the PSI/PSII ratio under 750-Fe conditions similar to those of this study (Fig. 4a) and others (5, 31, 38), they also observed increases in the iron-rich NifH protein under high-CO₂/Fe-limiting conditions (750-Fe), contrasting with our observed decreases in multiple nitrogenase transcripts and protein subunits in this study (Fig. 4b) and another *Trichodesmium* iron-limited proteome (31). Although Hong et al. (9) suggested that more NifH is necessary in 750-Fe because of decreased enzyme efficiency, we observed a contrasting trend via a fundamental shift in nitrogen metabolic strategy, whereby iron-rich gene abundances (e.g., PSI and *nif* subunits) along with nitrate transport transcripts are significantly reduced under severe iron stress concomitantly with increases in TMA-scavenging FMO and riboflavin biosynthesis gene expression (750-Fe; Fig. 4; Fig. S2).

Among the various forms of available exogenous nitrogen (e.g., amino acids, nitrate, TMA, ammonia), one possible reason why FMO may be preferred is that it can oxidize both nitrogen- and sulfur-containing compounds, including TMA, which may also serve as potential organic carbon sources. This may enable *Trichodesmium* to assimilate several macronutrients by minimizing energetic investment. Additionally, since *Trichodesmium* is physically colonized by a diverse microbial consortium (35, 39) and methylated amine compounds like TMA are ubiquitous as end products of, for example, protein putrefaction (32), *Trichodesmium* may have consistent access to organic nitrogen/carbon sources as a product of the biochemical decomposition of cellular components by members of its consortium. This contrasts with the lifestyles of other planktonic unicellular marine bacteria, which are typically associated with much less abundant and diverse consortia. When searching *Trichodesmium* consortium metatranscriptome sequences (35) for FMO via BLASTP (40), no significant hits containing the FMO-specific motif were observed, suggesting that it may not be widespread or expressed highly in consortia.

Although FMO homologs have been detected across global ocean data sets, including within the genomes of the ubiquitous SAR11 clade (32), only a few robust cyanobacterial and bacterial homologs to IMS101 were detected, with many of the remaining top high-scoring pairs deriving from metazoans (Fig. S3; high-scoring pair, $\geq 70\%$ of the original *Trichodesmium* FMO sequence length; E value, $\leq 10^{-5}$), yielding an average identity of 27.9% (median, 27.8%) over all pairs. We did detect the FMO genes in another *T. erythraeum* genome (strain 2175) and a *Trichodesmium* metagenome from hand-picked natural colonies (41), demonstrating widespread retention and conservation of FMO. Interestingly, no homologs of other biogeochemically relevant marine cyanobacteria were detected. Taken together, the findings show that FMO-mediated scavenging may be ecologically advantageous for *Trichodesmium* in the

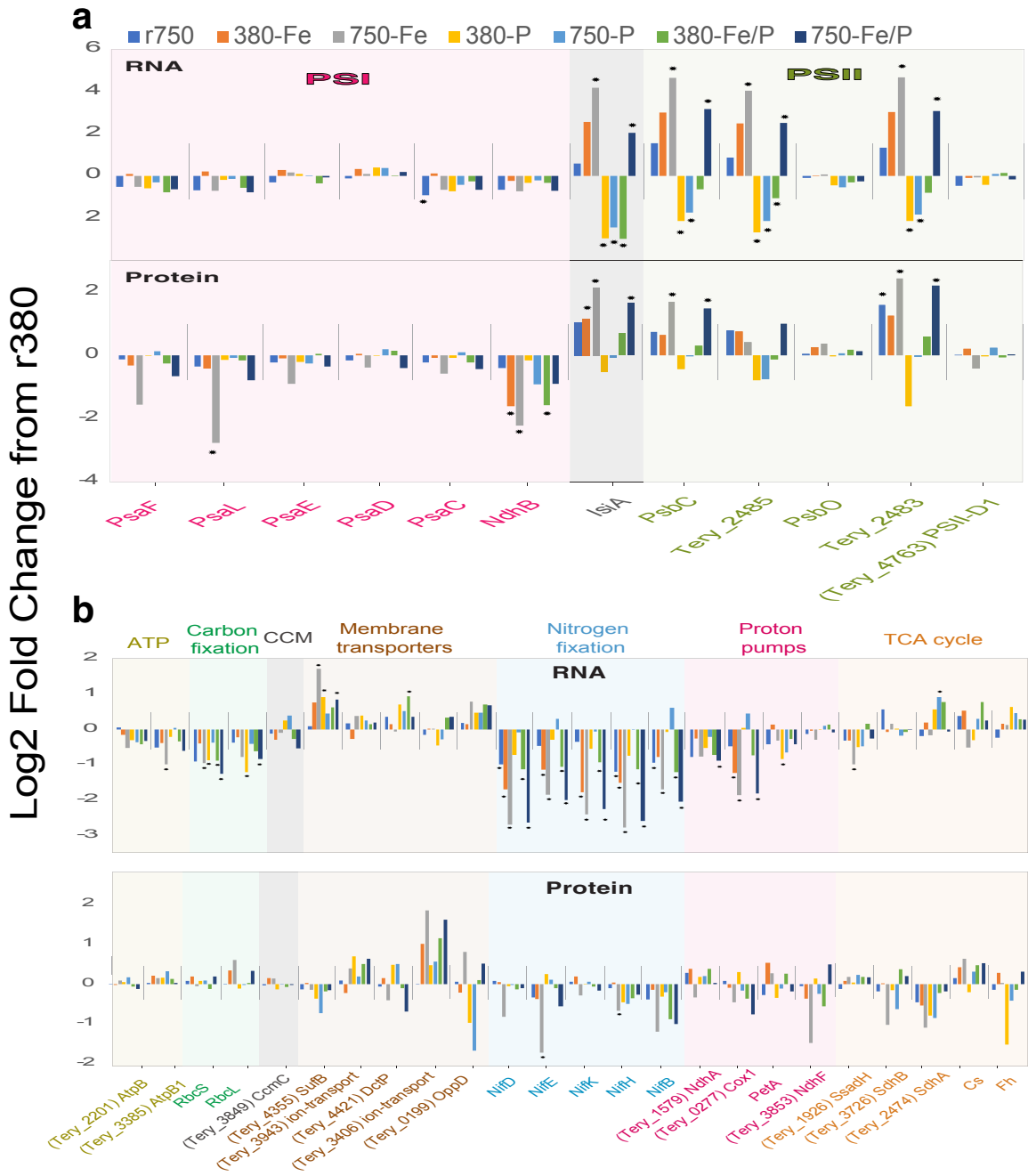


FIG 4 Log₂ fold changes in the normalized transcript and protein abundance of photosystem genes and genes from various other pathways. (a) Log₂ fold changes in gene transcript abundance relative to the r380 condition across experimental treatments. (b) Log₂ fold changes in normalized FMO protein spectral counts relative to r380.

oligotrophic ocean, given its potentially persistent availability via biochemical degradation and its relative scarcity in its physically attached consortia, as well as other sympatric cyanobacteria.

Fe-limited and Fe/P-colimited metabolism in the future ocean. Our data suggest that interaction of limiting Fe with high-CO₂ adaptation will intensify iron stress in Fe/P-colimited molecular metabolism. For example, known P stress genes (e.g., *phoX*, *sphX*, *phnG*), but not Fe stress genes (e.g., *isiA*), were significantly upregulated in 380-Fe/P compared to r380, suggesting that P stress may be more prominent than Fe stress under low-CO₂/colimiting conditions (Fig. 4 and 5; Fig. S4 and S6 and Data Sets

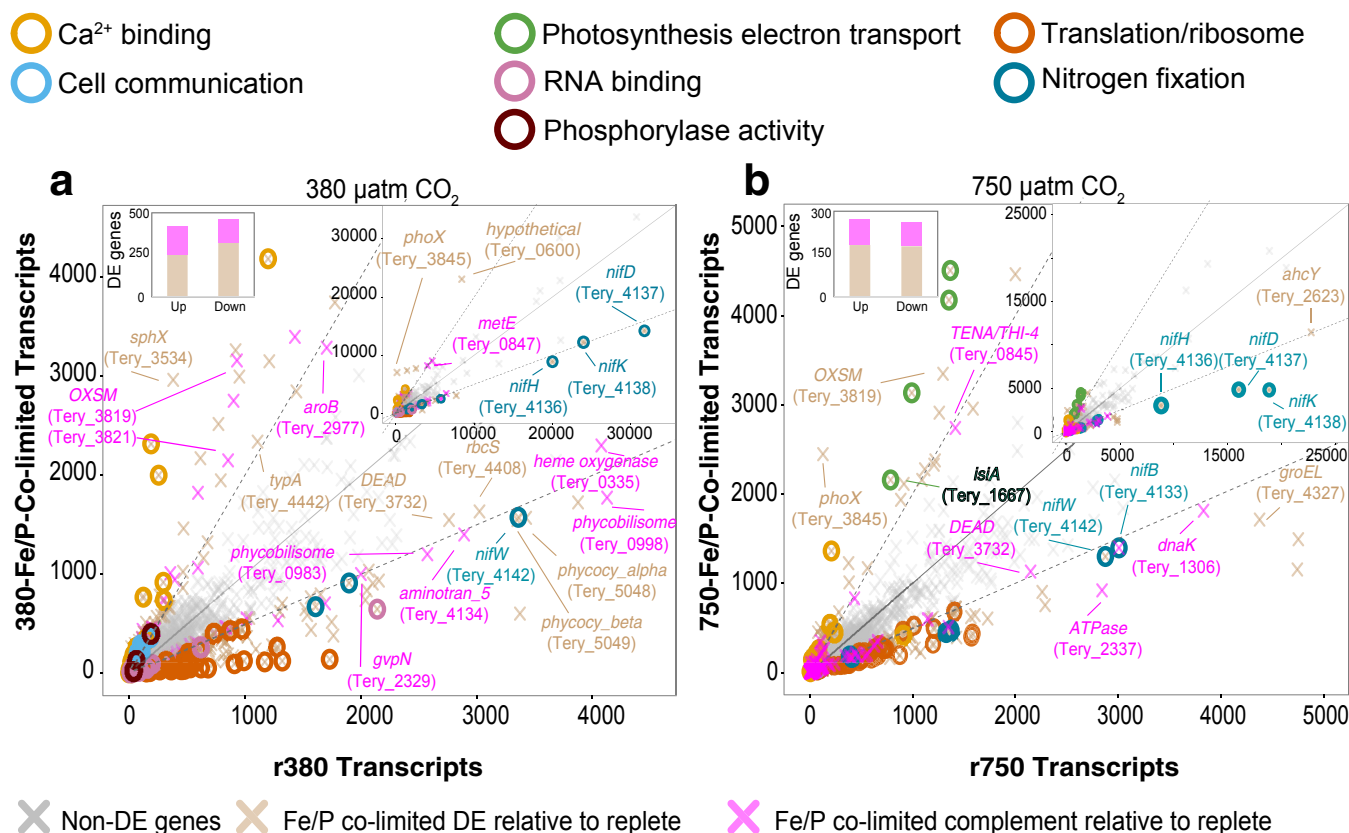


FIG 5 Replete versus Fe/P-colimited transcripts under 380 and 750 $\mu\text{atm CO}_2$. (a) Scatterplot of 380-Fe/P DE genes relative to r380 (tan) and 380-Fe/P DE genes relative to only r380 and no other 380 nutrient limitations (magenta). The center diagonal is a 1:1 line, while the others are 2:1 and 1:2 lines, respectively. The upper right insets are the same plots zoomed out to display highly expressed genes. (b) Similar scatterplot of 750-Fe/P DE genes relative to r750 (tan) and 750-Fe/P relative to only r750 and no other 750 nutrient limitations (magenta). Selected genes are labeled. The differently colored circles represent genes within significantly enriched GO pathways under the two conditions.

S2 and S3). In fact, *isiA* transcripts were significantly downregulated under both P-limited conditions, irrespective of CO₂, as well as during 380-Fe/P colimitation (Fig. 4a). However, transcripts of Fe stress genes, including *isiA*, were significantly increased in 750-Fe/P relative to those in 380-Fe/P and this was accompanied by a significant decrease in the PSI/PSII ratio, indicating iron stress under high-CO₂ conditions with colimitation (Fig. 4 and 5). Finally, this shift toward Fe stress is further evidenced by the high-confidence grouping of 750-Fe with the 380- and 750-Fe/P-colimited treatments (approximately unbiased [AU] bootstrap *P* value, >0.95) resulting from hierarchical clustering of genes exclusively DE (i.e., transcriptional complement; Data Set S3) in both 380- and 750-Fe/P (Fig. 6). Collectively, these data provide strong evidence of the intensification of Fe stress on cellular metabolism upon interaction with a high CO₂ concentration, regardless of fluctuating P concentrations. Hence, iron-rich metabolic pathways that generate energy and fixed nitrogen will likely be impacted as cells utilize alternative iron- and energy-conserving strategies to acquire nutrients, with consequent reductions in new-nitrogen inputs.

P-limited and Fe/P-colimited metabolism in present and future oceans. Upon adaptation to high-CO₂ conditions, both 750-P growth and N₂ fixation increased relative to those in 380-P (Fig. 1), while most 750-P-limited transcript levels remained statistically unchanged between 380-P and 750-P. This is evidenced by the much smaller number (*n* = 4) of genes DE between 380-P and 750-P than those DE between 380-Fe and 750-Fe (*n* = 284; Fig. S5 and S6). Translation dominated the GO-enriched pathway pools for downregulated genes relative to both r750 only and 750-Fe/P (Fig. S4 and S6 and Data Set S2), as in 380-P. It is possible that increased N₂ fixation rates of

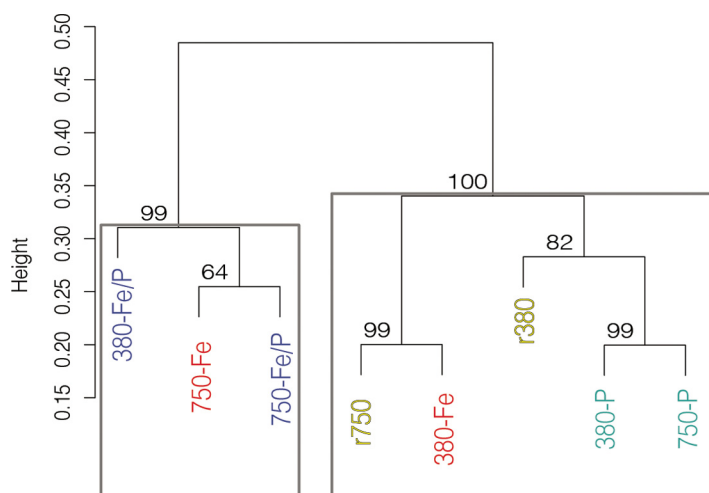


FIG 6 Hierarchical clustering of 380- and 750-Fe/P gene transcript complements. Hierarchical clustering of Bray-Curtis dissimilarities with multiscale bootstrap resampling calculated from normalized transcripts of all DE genes in both 380- and 750-Fe/P complements. Values at nodes are AU *P* values. Boxes highlight high-confidence clusters with AU *P* values of >0.95.

equivalently sized IMS101 cells under various CO₂ conditions (e.g., r380 versus r750 and 380-P versus 750-P) are driven by factors contributing to altered enzymatic rates rather than by increases in enzyme abundance, as has been observed for temperature (42). This possibility is supported by either no change or decreases in N₂ fixation transcripts and/or protein abundances (e.g., *nifH*, *nifB*, *nifK*) in replete, 7-year high-CO₂-adapted IMS101 (r750) relative to replete, low-CO₂ (r380) cells, despite higher N₂ fixation rates in cells selected under high-CO₂ conditions (Fig. 4) (5). For some N₂ fixation genes, transcript increases were mirrored by increases in protein abundance (e.g., those encoded by *nifB* [Tery_4133] and *nifK* [Tery_4138]), though most remained unchanged in the proteomic data (e.g., those encoded by *nifH* [Tery_4136] and *nifD* [Tery_4137]). These trends suggest that different nitrogenase genes may be independently regulated at the transcriptional level or have different mRNA half-lives. Taken together, the results show that other regulatory mechanisms impacting enzymatic rates, but not abundances, may be more generally associated with changes in CO₂-impacted physiology under P limitation.

Fe/P-colimited transcription in present and future oceans. Photosynthesis-related (e.g., Tery_0983, Tery_0998, Tery_5048, Tery_5049, Tery_0728), ferredoxin (e.g., Tery_0916, Tery_4539, Tery_5051), heme-containing (e.g., Tery_1714, Tery_0335), and N₂-fixing genes (Fig. 2, 4, and 5; Fig. S7) were all downregulated in 380-Fe/P. This indicates a potential reduction and/or reallocation of Fe away from these pathways in the colimited cells coinciding with reduced cell size (and therefore cellular elemental quotas), growth, and N₂ fixation relative to those seen under replete conditions. Additionally, downregulation of GO-enriched pathways in 380-Fe/P involved in translational machinery and RNA binding (Fig. 5; Fig. S7) are consistent with observations in yeast and bacteria (i.e., stringent response), where proteins involved in cell growth and division (e.g., ribosomal proteins) have been observed to be downregulated in minimal (e.g., 380-Fe/P) versus rich (e.g., r380) medium (43, 44). Moreover, the upregulated 380-Fe/P (relative to r380) transcriptional complement also contained many genes involved in precursor metabolism, potentially allowing for greater cellular plasticity and metabolic flexibility under low-nutrient regimes (5). Accordingly, we identified a set of upregulated proteins specific to 380-Fe/P colimitation that included many involved in precursor metabolisms (i.e., isoprenoid biosynthesis), a finding consistent with prior observations in yeast maintained in minimal medium (44). For example, 3-dehydroquinate synthase (Tery_2977, *aroB*) is upregulated solely in 380-Fe/P and is part of the shikimate pathway that produces

chorismate, a precursor of aromatic amino acids, including phenylalanine, tyrosine, and tryptophan (45) (Fig. S7).

Following 7 years of adaptation to high-CO₂ conditions and 1 year of growth under high-CO₂ Fe/P colimitation conditions (750-Fe/P), transcript levels had shifted across broad metabolic pathways, presumably orchestrating the maintenance of growth and N₂ fixation similar to those seen under 380-Fe/P conditions (Fig. 5; Fig. S7). Of those genes that were exclusively DE during Fe/P colimitation, only ~5% ($n = 26$; Data Set S3) from the 380-Fe/P complement ($n = 313$) remained in the 750-Fe/P complement ($n = 174$) (Data Set S3). In other words, these genes were either upregulated ($n = 15$) or downregulated ($n = 11$) only during 380-Fe/P and 750-Fe/P colimitation, relative to the respective replete reference conditions (r380 and r750). Shared upregulated genes include the universal stress global response regulator (Tery_2353, *uspA*), the aforementioned GTP-binding gene (Tery_1904), and a carbohydrate selective porin gene (Tery_0838, *oprB*). The consistent upregulation of these genes suggests that 380- and 750-Fe/P-colimited cellular physiology responds to environmental flux by consistently transcribing genes involved in general stress, biotimer regulation, and potential organic carbon uptake, as has been observed in other cyanobacteria (33, 34). Shared downregulated genes include that for a 5',5''-P₁P₄-tetraphosphate phosphorylase (Tery_0108, *ap₄A*), which has been implicated in bacterial stress responses and gene regulation (46), and the mutator gene *mutT* (Tery_4056), which has been demonstrated to suppress transversion mutations (A-T to C-G) (47). Accordingly, *Escherichia coli* Δ *mutT* deletion mutants exhibited a 1,000-fold increase in unidirectional A-T-to-C-G transversion frequencies. Other studies have postulated that mutators can accelerate adaptive evolution under certain circumstances via increased mutation rates (48). Hence, the significant reduction of *mutT* transcription in long-term 380-Fe/P and 750-Fe/P treatments may help enable adaptive mutation by possibly derepressing transversion mutations in low-nutrient, stress-inducing environments.

Conclusions. Here, we show a marked shift in nitrogen metabolism going from low-CO₂/Fe-limited to high-CO₂/Fe-limited regimes whereby iron-heavy pathways are significantly reduced, and cellular investment is reallocated toward a predicted FMO-mediated organic nitrogen scavenging, relative to acquisition pathways for inorganic nitrogen substrates. This fundamental iron-saving strategy may also enable simultaneous assimilation of several other required elements, including C and S from methylated amines like TMA. Our data also suggest that intensifying Fe stress under high-CO₂ conditions may shift Fe/P-colimited metabolism into a more Fe-limited metabolic state. This trend further highlights the need for iron-saving metabolic strategies for nutrient limitation in a future high-CO₂ ocean. Additionally, N₂ fixation but not growth was inhibited in the presence of TMA. If interactive global change factors intensify nutrient limitation, leading to enhanced organic nitrogen scavenging and reduced N₂ fixation by *Trichodesmium*, future work must consider the relationship between N₂ fixation-mediated new-nitrogen inputs and simultaneous removal from the fixed nitrogen pool. The shifting balance between these two processes may have global implications for the role of *Trichodesmium* in future ocean biomes. Finally, transcript patterns in Fe/P-colimited metabolism also suggest reductions in iron-heavy pathways in exchange for increases in precursor-related genes that may aid cellular plasticity in response to various nutrient concentrations. Although limitations in this long-term experiment and others (49) include a steady-state environment lacking natural variability, as well as the use of a single strain of *Trichodesmium*, we believe that our results demonstrating molecular reallocations of nitrogen acquisition and iron-sparing expression provide robust indicators of potential *in situ* strategies some *Trichodesmium* populations may employ while under covarying nutrient-limited regimes. We highlight consistent molecular patterns in genes under Fe and/or P limitation while covarying CO₂, which strengthens the broad trends outlined above. Future field work investigating colimitation should aim to compare *in situ* molecular patterns with those that we have

identified in culture to try to help elucidate universal indicators of colimitation in the future ocean.

MATERIALS AND METHODS

Culturing methods. Details of the nutrient-limited culturing methods used in this study can be found in reference 5 and in the supplemental material. For the growth experiment comparing TMA with other N sources, IMS101 was grown in four N treatments, (i) N_2 , (ii) 20 μM $NaNO_3$, (iii) 20 μM TMA (C_3H_9N), and (iv) 20 μM NH_4Cl . For all treatments, triplicate cultures were maintained semicontinuously at 26°C under a light irradiance intensity of 150 μmol of photons $m^{-2} s^{-1}$ (12-h light/12-h dark cycle), cultures were diluted every other day, and 20 μM N sources were added after every dilution. Growth and nitrogen fixation rates were measured after 10 generations as described above and below.

Aquil medium was bubbled with 0.2- μm -filtered prepared air- CO_2 mixtures (Praxair), and in-line high-efficiency particulate air filters were employed to avoid Fe contamination from particles in the gas tanks or lines. The pH was monitored daily, and dissolved inorganic carbon concentrations were measured at the final sampling. The acetylene reduction method was used to measure *Trichodesmium* N_2 fixation rates (4, 14, 25). Samples from each experimental triplicate (20 ml) were incubated in gas-tight vials for 1 h after being injected with 2 ml of acetylene in 23 ml of headspace. The amount of ethylene produced was then analyzed by injecting a 200- μl aliquot of headspace gas into a gas chromatograph (model GC-8A; Shimadzu Scientific Instruments, Columbia, MD, USA). An ethylene accumulation-to- N_2 fixation conversion ratio of 3 was used to calculate N_2 fixation rates. Upon steady-state growth, *Trichodesmium* filament abundance and length were measured in a 1-ml phytoplankton counting chamber by epifluorescence microscopy and significant differences between nutrient conditions were calculated by two-way analysis of variance along with the Tukey test. For RNA sampling, cultures were swiftly and gently filtered onto 5- μm polycarbonate filters (Whatman) during the middle of the photoperiod, immediately flash frozen, and stored in liquid nitrogen until RNA extraction.

RNA isolation and extraction for Illumina sequencing. For each replicate, samples for RNA extraction were taken concomitantly with proteome samples during the middle of the photoperiod, as described in reference 5. Briefly, cells were swiftly and gently filtered at 11 a.m. with 5- μm polycarbonate filters (Whatman), immediately flash frozen, and stored in liquid nitrogen until RNA extraction. RNA was extracted from two randomly chosen biological replicates per treatment with the Ambion MirVana microRNA isolation kit (Thermo Fisher Scientific) in an RNase-free environment in accordance with the manufacturer's instructions, followed by two incubations with the Ambion Turbo DNA-free kit to degrade trace amounts of DNA. RNA was then submitted to the University of California San Diego Institute for Genomic Medicine core for library preparation and sequencing (<http://igm.ucsd.edu/genomics/services.shtml>). Briefly, rRNA removal and library construction were done with the Ribo-Zero rRNA removal kit (Illumina) and the TruSeq Stranded RNA Library Prep kit (Illumina), respectively, and multiplexed libraries were sequenced by using Illumina Hi-Seq, yielding single-end 50-bp read libraries. All of the protein spectral data used in the above-described analyses can be found in Supplementary Data 4 of reference 5.

Expression analysis. Differential expression was done as previously described (6), and detailed methods can be found in the supplemental material.

GO enrichment analysis. As in reference 6, GO annotations for *Trichodesmium* were downloaded from the Genome2D web server (<http://genome2d.molgenrug.nl>). Next, the phyper function in R (R Core Team 2014) was used to test for significant enrichment of GO categories among the treatments and *P* values were corrected by the Benjamini-Hochberg method (50) with the p.adjust function ($P \leq 0.1$). Finally, genes in enriched GO categories were manually checked.

Phylogenetic analysis. FMO sequences for *T. erythraeum* strains IMS101 and 2175 were taken from the IMG website (<https://img.jgi.doe.gov/>), and *Trichodesmium* environmental metagenomic sequence data were downloaded from reference 41. The BLASTP algorithm was used to search sequences against the RefSeq protein database (51), in which all high-scoring pairs were retained if the aligned portion spanned >70% of the original query length with an E value of $<10^{-5}$. Sequences were then aligned with MUSCLE v3.8.31 by using the default settings (52), and spurious sequences and poorly aligned regions were removed with trimAl 1.2rev59 (53). RAXML (54) was used for all maximum-likelihood phylogenetic analyses with the following settings: -f a -p 12345 -m PROTCATLG -N 100 -x 12345.

Availability of data. All of the transcriptome sequencing data used in this study have been deposited as raw fastq files in the NCBI Gene Expression Omnibus (55) and are accessible through GEO Series accession number [GSE94951](https://www.ncbi.nlm.nih.gov/geo/query/acc.cgi?acc=GSE94951). All of the protein spectral data used in the above-described analyses can be found in Supplementary Data 4 of reference 5. Physiological and proteome data are archived through the U.S. National Science Foundation Biological and Chemical Oceanography Data Management Office (<http://www.bco-dmo.org/dataset/649904>).

SUPPLEMENTAL MATERIAL

Supplemental material for this article may be found at <https://doi.org/10.1128/AEM.02137-17>.

SUPPLEMENTAL FILE 1, PDF file, 2.6 MB.

ACKNOWLEDGMENTS

This work was supported by U.S. National Science Foundation grants OCE-1260233, OCE 1260490, and OCE 1657757 to D.A.H., E.A.W., and F.-X.F. and grant OCE OA 1220484 to D.A.H. as well as G. B. Moore Foundation grants 3782 and 3934 to M.A.S.

We have no competing interests to declare.

F.-X.F., D.A.H., and E.A.W. designed the research; N.G.W., M.D.L., X.C., and F.-X.F. performed the research; N.G.W., M.D.L., X.C., F.-X.F., D.A.H., M.A.S., and E.A.W. analyzed the data; and N.G.W., M.D.L., F.-X.F., D.A.H., M.A.S., and E.A.W. wrote the paper.

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