

Iron conservation by reduction of metalloenzyme inventories in the marine diazotroph *Crocospaera watsonii*

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The marine nitrogen fixing microorganisms (diazotrophs) are a major source of nitrogen to open ocean ecosystems and are predicted to be limited by iron in most marine environments. Here we use global and targeted proteomic analyses on a key unicellular marine diazotroph *Crocospaera watsonii* to reveal large scale diel changes in its proteome, including substantial variations in concentrations of iron metalloproteins involved in nitrogen fixation and photosynthesis, as well as nocturnal flavodoxin production. The daily synthesis and degradation of enzymes in coordination with their utilization results in a lowered cellular metalloenzyme inventory that requires ~40% less iron than if these enzymes were maintained throughout the diel cycle. This strategy is energetically expensive, but appears to serve as an important adaptation for confronting the iron scarcity of the open oceans. A global numerical model of ocean circulation, biogeochemistry and ecosystems suggests that *Crocospaera's* ability to reduce its iron-metalloenzyme inventory provides two advantages: It allows *Crocospaera* to inhabit regions lower in iron and allows the same iron supply to support higher *Crocospaera* biomass and nitrogen fixation than if they did not have this reduced iron requirement.

cyanobacteria | marine iron cycle | nitrogen cycle

The biological fixation of atmospheric dinitrogen into ammonia has a major impact on the extent of marine primary production (1, 2). Only a small number of bacteria are known to contribute to nitrogen fixation in open ocean environments, and they typically comprise a tiny fraction (<0.1%) of the overall microbial community (3, 4). Among these, the unicellular diazotroph *Crocospaera watsonii* is estimated to be a significant contributor to oceanic nitrogen fixation (4–6). Whereas much of the primary productivity in the tropical and subtropical regions of the oceans is predicted to be nitrogen limited, the diazotrophs' nitrogen source, resupplied from the large atmospheric reservoir, is essentially limitless. Instead, iron is considered the critical micronutrient for marine diazotrophs due to their use of the iron-nitrogenase protein complex containing a homodimeric iron protein with a 4Fe:4S metallocluster (NifH) and a heterotetrameric molybdenum-iron protein with an 8Fe:7S P cluster and a 7 Fe and 1 Mo MoFe cofactor (NifDK, α and β subunits) (7, 8) (Table S1). Field experiments and models both predict the distribution of oceanic nitrogen fixation to be primarily constrained by the availability of iron (2, 9–11). Despite this importance of iron on marine nitrogen fixation, there is a limited understanding of how marine diazotrophs have adapted to this low iron environment (12–17).

The coexistence of oxygenic photosynthesis and nitrogen fixation metabolisms presents a unique challenge for diazotrophs due to their high iron demands and the chemical incompatibility of molecular oxygen and the nitrogenase protein complex. Previous elemental studies measuring whole cell iron content in *Crocospaera* found increased iron during the dark period when nitrogen fixation is occurring (15). Whereas this finding was con-

sistent with theoretical studies predicting a large iron requirement in marine diazotrophs (13, 18), it has been somewhat controversial due to the implication of a dynamic yet poorly understood diel cycle for intracellular iron. Several unicellular diazotrophs, including *Crocospaera watsonii*, have been observed to fix nitrogen during the dark period (Fig. 1A), and this is widely viewed as an adaptation for temporal separation of photosynthesis and nitrogen fixation to avoid the oxygen disruption of the nitrogenase complex (6, 19, 20). Global transcriptional studies have also observed large scale changes in the transcriptomes of the unicellular diazotrophs *Crocospaera watsonii* (21), *Cyanothece* (22), *Gloeothece* (23), and a hot-spring *Synechococcus* (24) during the diel cycle, consistent with a temporal separation of photosynthesis and nitrogen fixation. Yet the possibility that diel cycling of the transcriptome or proteome might affect the iron requirement has not been previously discussed, likely because transcriptional studies provide information about gene expression rather than actual enzyme inventories, and immunological protein studies thus far, while having detected diel oscillations of the NifH subunit in *Crocospaera* and *Gloeothece* (23, 25), are limited to a few select proteins with no global or absolute quantitative proteome studies of diazotrophs published as of yet. As a result, it has remained uncertain whether the observed diel transcriptome cycling was needed for the maintenance of a relatively consistent proteome or was in fact causing large changes in the global proteome composition during each diel cycle.

Recent advances in proteomic technologies hold promise for elucidating mechanistic connections between the biochemistry of important microbes and global biogeochemical cycles. There are two broad approaches to mass spectrometry based proteomics: global shotgun proteomic approaches that can semiquantitatively survey the relative abundance of hundreds of proteins simultaneously using spectral counting (26), and targeted approaches that use selected reaction monitoring (SRM) mass spectrometry with isotopically labeled peptide standards for absolute quantification of proteins of interest (27). Though there are several culture-based studies of marine microbes (26, 28) and field assessments of abundant proteins (29, 30), the combined application of global and targeted proteomic approaches to problems of

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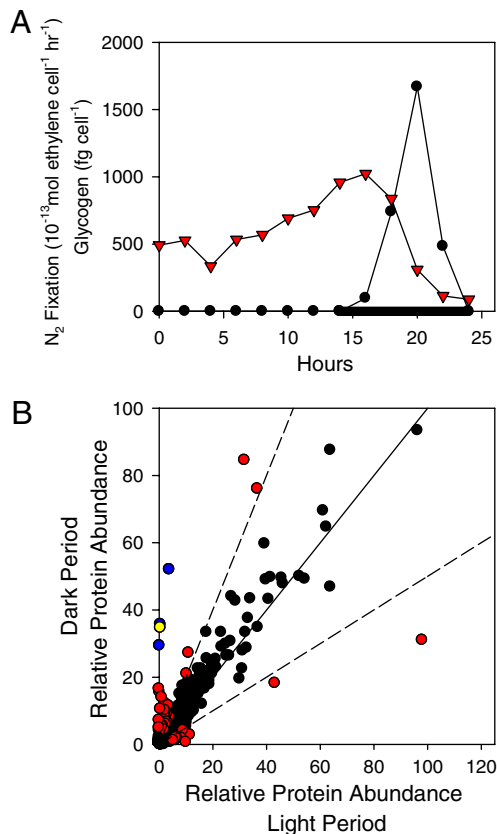


Fig. 1. Variations in the *Crocosphaera* proteome between the light and dark photoperiod. (A) Dark period nitrogen fixation (black) and glycogen storage (red) during a diel cycle (dark, black bar). (B) Relative protein abundance during light and dark periods, where similar expression (solid line, black circles; also see Fig. S1A for comparison of technical replicates) is contrasted by a number of proteins (colors), including the nitrogenase complex metalloenzymes (blue) and flavodoxin (yellow), with greater than 2-fold change in relative abundance (dashed lines) above a threshold of 5 normalized spectral counts in at least one treatment. Data are mean of six technical replicates.

marine biogeochemical relevance, such as the iron limitation of marine diazotrophy, has not yet been reported.

Results and Discussion

The diel cycle and inventory of iron metalloenzymes in *Crocosphaera watsonii* were investigated using global and targeted liquid chromatography mass spectrometry (LC-MS) proteomic techniques from three distinct culture experiments all grown on a 14:10 light:dark cycle: a day-night experiment sampled during the light and dark photoperiod (sampled 10 h from lights on and 7 h from lights off) analyzed for global proteome using both 1D and 2D LC-MS approaches, a time course study (diel hereon) sampled every 2–3 h for 30 h (beginning 4 h from lights on) and analyzed for global proteome using 1D LC-MS and for absolute quantities of targeted proteins, and a set of biological triplicate cultures (sampled 6.5 h into light period, and 3 and 6 h into the dark period) analyzed for targeted protein abundances by mass spectrometry.

Of the 477 unique proteins identified in the global proteomic 1D analysis of the day-night experiment [0.65% false positive rate (FPR) (31)], 37 proteins were found to have a 2-fold or greater change in abundance between the day and night (Fig. 1B). These results were confirmed in the deeper 2D chromatography proteome where 160 of the 1,108 identified proteins (0.11% FPR) showed a greater than 2-fold change (Fig. S1B). A time course of the diel cycle clearly showed a cycling of protein abundance (Fig. 2), where 100 proteins were found to be above a threshold

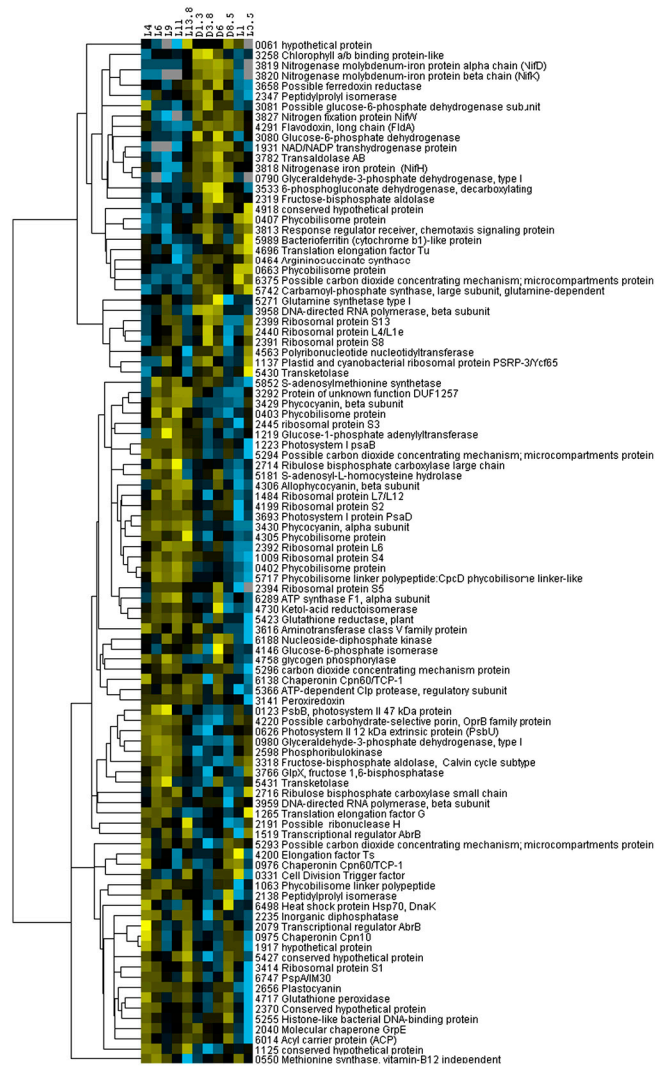


Fig. 2. Cluster analysis of global proteome during the diel cycle. Color indicates higher (yellow) or lower (blue) relative abundance relative to the centered mean value (black). Horizontal axis is hours from onset of light (L) or dark (D) period. Proteome data was filtered for proteins with at least three time points having greater than 10 spectral counts and a difference between maximum and minimum signals ≥ 14 spectral counts, prior to log transformation and normalization. Two major clusters were observed corresponding to the light and dark photoperiods.

spectral count signal (≥ 10) and displayed a diel variation in amplitude (max - min ≥ 14). Cluster analysis revealed two major groups that include key photosystem or nitrogen fixation proteins with increased abundances during the day and night respectively. These experiments demonstrate that the diel cycling observed in the transcriptome of unicellular diazotrophs (21, 22, 32) manifests itself in large scale changes in the global proteome, where more than 20% of the measured proteins in the diel experiment showed diel variation. In both the day-night and diel experiments, the metalloenzymes involved in nitrogen fixation in *Crocosphaera watsonii* were among those with the most pronounced changes in abundance, being largely absent during the day and present at night (Figs 1B and 2, Fig. S2, and Dataset S1).

Based on these global experiments, the absolute abundances of nine protein targets in the diel experiment were determined by triple quadrupole SRM mass spectrometry using isotopically labeled internal standards (Table S2). The abundance of the three nitrogenase metalloproteins ranged from nondetectable levels during the photoperiod to being among the most abundant pro-

teins in the proteome in the dark period (Fig. 3 A–C). At their peak abundances, the ratio of the iron protein to molybdenum-iron protein subunits (α used here) was 3.5:1, consistent with previous observations of a 3:1 cellular ratio (23) and a 2:1 stoichiometry within the nitrogenase protein complex (33). Together these global and targeted proteome measurements provide evidence for the nocturnal synthesis of the metalloenzymes in the nitrogenase complex followed by their complete diurnal degradation.

Flavodoxin was observed to increase in abundance during the dark period (Fig. 3D), similar to the nitrogenase metalloenzymes and in contrast with the increased flavodoxin abundances observed in most phytoplankton under iron deprivation (34). This suggests the ferredoxin-flavodoxin substitution for electron transport in photosynthesis does not occur appreciably in *Crocospaera*. The global proteome of a short-term iron deprivation experiment was sampled during the light photoperiod to allow differentiation of iron effects from the dark expression. No increase in flavodoxin in response to iron stress was observed 24 h after the addition of an exogenous siderophore (Fig. S1 C–F), consistent with the hypothesis that flavodoxin could be functioning as an electron carrier for nitrogenase instead of ferredoxin (35). The use of flavodoxin at night even under iron replete conditions appears to be an adaptation for the overall minimization of the cellular iron demand in this microbe.

In a mirror image of the nitrogenase metalloenzymes and flavodoxin, numerous proteins were more abundant during the photoperiod (Figs. 1B and 2). In particular, several proteins from iron-rich photosystem I decreased during the dark period and increased during light period, including PsaA, PsaB, PsaD, and PsaF (PsaA and PsaB shown in Fig. 3 E and F). The iron-containing cytochromes b6 and c550 were among the most abundant cytochromes detected by spectral counting (f, P450, and b559 and c oxidase were also detected) and showed large decreases in abundance during the dark period as measured by targeted mass spectrometry (Fig. 3 G and H; 2- and 15-fold respectively). The degradation of substantial quantities of cytochrome b6 and c550 should release a number of iron-containing hemes during the dark period. Two heme oxygenases, which are capable of liberating iron from heme, were identified in the proteome, showed diel variations, and may be important in iron recycling (Fig. S2). Iron storage bacterioferritin proteins were also identified in the proteome, one of which showed some diel periodicity (Fig. S2). Diel cycling is not necessarily expected for bacterioferritin given that iron is thought to be added and removed from the ferritin cage without its degradation.

These results portray a *Crocospaera* proteome that is dynamic, shifting its metalloenzyme inventory to suit the alternating temporal needs of nitrogen fixation and photosynthetic biochemical activity. Whereas *Crocospaera*'s distinct nocturnal nitrogenase activity is one of its defining characteristics in both laboratory

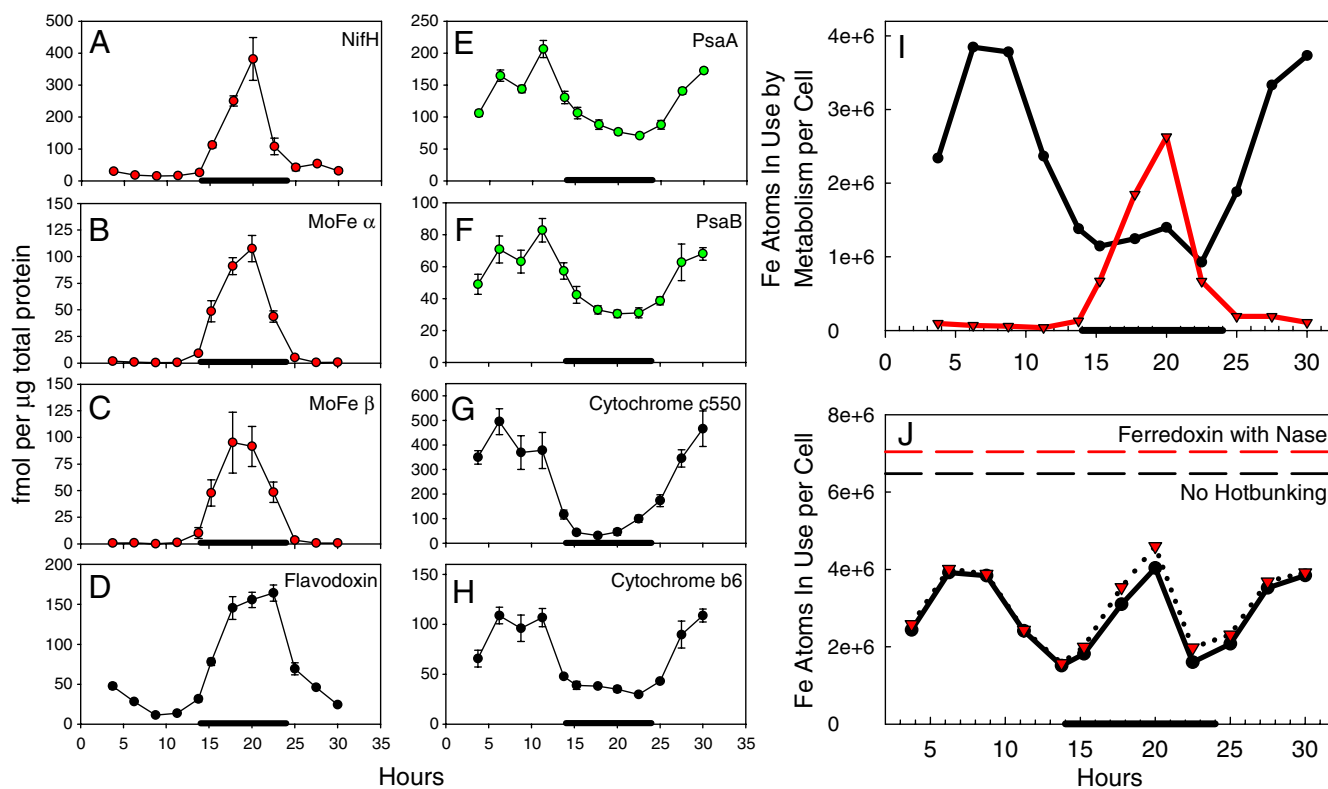


Fig. 3. Absolute quantitation of nitrogenase metalloproteins and selected photosynthesis and electron transport proteins during the diel cycle, and estimates of cellular iron stoichiometry based on absolute protein concentrations. (A–H) Targeted proteomic analysis of iron-nitrogenase (NifH), molybdenum-iron protein subunits (NifDK, MoFe α , MoFe β), and flavodoxin, photosystem I proteins PsaA and PsaB, and cytochromes b6 and c550 during the diel cycle (darkness, black bar) as measured using SRM triple quadrupole liquid chromatography mass spectrometry employing isotopically labeled standards. Data are mean and one standard deviation of technical triplicates. (I) Iron demand associated with nitrogen fixation or photosynthesis estimated from protein abundances in A–H, total protein and cell concentrations, and metal enzyme stoichiometries (Fig. S3 and Table S1) assuming metalloenzymes were populated with iron (darkness, black bar). (J) Sum of photosynthesis and nitrogenase iron estimates in two scenarios (darkness, black bar): (i) the *Crocospaera* hotbunking scenario summing the calculated values in panel I (black circles), and (ii) a hypothetical nonhotbunking scenario (dashed black line), where the maximum iron required for each process [the peak of each line in panel (I) is summed]. Two additional scenarios are shown, where measured flavodoxin is replaced by ferredoxin, assuming a conservative 1:1 substitution (hotbunking, red triangles; nonhotbunking, dashed red line). The hotbunking scenario represents an estimated 40% savings in iron through a reduction of the iron-metalloenzyme inventory involved with nitrogenase and photosynthesis.

and field studies (6, 36), the fate of the nitrogenase protein complex during the light period has remained controversial due to the technical limitations of transcriptional and immunological methods, where transcripts do not measure protein abundance and immunological methods may not detect posttranslationally modified proteins (6, 15, 21, 36, 37). Using mass spectrometry-based analysis of multiple tryptic peptides from each protein, this study demonstrates the near complete degradation of the metalloenzymes from the nitrogenase complex and the partial degradation of certain iron-containing components of photosynthesis. The extent of proteome cycling we observed in *Crocospaera watsonii* was surprising given the potentially significant energetic cost. This proteome cycling likely serves the dual functions of preventing oxygen disruption to the nitrogenase complex as discussed previously (6, 19), as well as providing a mechanism for a significant reduction in metabolic iron demand. We argue that the latter is true based on stoichiometric calculations from the quantitative metalloenzyme data in Fig. 3 A–H and the associated iron stoichiometry for each metalloenzyme (Table S1). By summing the iron associated with major metalloenzymes, we calculated the amount of iron involved in nitrogen fixation and photosynthesis throughout the diel cycle (Fig. 3I). By degrading these metalloenzymes when they are not in use, *Crocospaera* reduces its iron-metalloenzyme inventory for these metabolic functions at any given time by 38%, 40%, and 75% for peak night, peak day, and minimum, respectively, compared to a hypothetical situation where these metalloenzymes are not degraded (Fig. 3J). An additional experiment with targeted protein analyses on biological triplicates was consistent with these estimates of iron conservation from the diel experiment, using $45 \pm 19\%$ less iron during the photoperiod (6.5 h from lights on, $n = 3$), and $38 \pm 14\%$ and $38 \pm 22\%$ less iron during the dark period (sampled at 3 h and 6 h from lights off, $n = 3$ for each). These savings in iron use would be even larger if the iron-free flavodoxin observed during the dark period (Fig. 3D) was replaced by iron-requiring ferredoxin (Fig. 3J). Iron released by the degradation of these metalloenzymes could have several fates including storage in bacterioferritin or chaperones, loss from the cell, and reuse in other metalloenzymes. Given the severe iron limiting conditions of the upper oceans (2, 9), it seems likely that a fraction of this liberated iron inventory is participating in both nitrogen fixation and photosynthesis metabolic processes. This iron conservation strategy is analogous to the maritime practice of hotbunking, referring to ships that sail with more sailors (metalloenzyme requirements) than bunks (iron atoms), where sailors on opposing shifts share the same bunk—keeping the bunks continually hot (or iron atoms in use). Whereas it is methodologically difficult to document the sharing of individual iron atoms between metabolisms, our data clearly demonstrates the reduction in iron-metalloenzyme inventory and concurrent reduction in metabolic iron demand through diel proteome cycling (Fig. 3J).

Iron conservation by reducing the metalloenzyme inventory likely serves as a key component of a low iron marine niche for *Crocospaera*. *Trichodesmium* sp. is another dominant oxygenic marine diazotroph that differs from *Crocospaera* in that it fixes both nitrogen and carbon during the photoperiod (38, 39). As a result *Trichodesmium* sp. likely does not employ this iron conservation strategy to the extent used by *Crocospaera*. This is consistent with reports of an approximately double cellular iron to carbon stoichiometry in *Trichodesmium* compared to *Crocospaera* (>35 and $16 \pm 11 \mu\text{mol Fe mol C}^{-1}$, respectively) (14–16), as well as estimates of a large fraction of cellular iron being necessary for nitrogen fixation (22–50%) in *Trichodesmium* (13). A previous study of cellular iron in *Crocospaera watsonii* observed an increase in iron concentrations per cell during the dark period (15). Our calculated iron per cell attributed to nitrogen fixation and photosynthesis are roughly consistent with measured values (15), and the higher iron content at night likely

results from diel proteome cycling with the added contributions of daytime uptake and accumulation of ferrous iron (21) produced by photochemical reduction (40) and photoperiod cell division (Fig. S4).

By reducing the nitrogen fixation and photosynthetic metalloenzyme inventory when each is not in use, *Crocospaera watsonii* has evolved to require $\sim 40\%$ less iron than if these enzymes were maintained throughout the diel cycle (Fig. 3J). Though difficult to quantify, this process of resynthesizing enzymes every day must have an energetic expense. Based on our quantitative measurements, diel cycling of the three nitrogenase metalloproteins alone contributes $\sim 2.3\%$ of the total protein and all eight targeted proteins in Fig. 3 contribute 5.0% (using the difference between the maximum and minimum expression levels during the diel cycle). With many lower abundance proteins also undergoing diel cycling (Fig. 2), and with protein contributing roughly 40% of cellular biomass (41), these calculations provide some rough sense of the energetic cost for the increased protein synthesis required for this reduction in iron metabolic demand (“hotbunking” hereon). If *Crocospaera watsonii* had evolved hotbunking from an ancestor that maintained its metalloenzymes through the diel, how would the trade-off between lower iron requirement and higher energetic cost affect where *Crocospaera* would be competitive relative to its progenitor and other diazotrophs? A numerical global ocean circulation and ecosystem model (10, 42) is a useful tool to explore the implications of hotbunking on *Crocospaera*'s habitat and nitrogen fixation rates in the global ocean. Our model resolves several phytoplankton types competing for resources, including analogs of several nondiazotrophs types, and unicellular and colonial marine diazotrophs, analogous to *Crocospaera* and *Trichodesmium*, respectively. The modeled unicellular diazotrophs consist of two groups: a hypothetical control group that could not share cellular iron between photosynthesis and nitrogen fixation machinery (analogous to potential *Crocospaera* ancestors that maintained their metalloenzyme inventory), and a hotbunking group (analog of *Crocospaera*, as suggested by this study) that are conferred with a lower cellular iron demand (R) at a cost to their growth rate (μ):

$$\mu_H = \alpha\mu_C, \quad R_H = \gamma R_C \quad \text{where } \alpha, \gamma < 1.$$

The subscript H refers to hotbunkers and C refers to the non-hotbunking control group. We conducted a suite of simulations with different assumptions of the trade-off between reduced growth and iron requirement, the respective values of α and γ . A larger value of α indicates less energetic cost, and a larger value of γ indicates iron requirements closer to the control group.

Iron conservation strategies in *Crocospaera* provided an ecological advantage in the low iron environments of the open ocean (Fig. 4 and Fig. S5). The geographical habitat of the hotbunking-analogs extended further into iron-depleted water than both the hypothetical control group and the *Trichodesmium*-analogs. In simulations with favorable combinations of trade-offs (e.g., Fig. 4), the distribution of *Crocospaera*-analogs is broadly consistent with those suggested by observations (4, 10). These simulation results are consistent with observations suggesting that *Trichodesmium* outcompetes *Crocospaera* in areas with high iron (43). The extent of the hotbunking unicellular diazotroph habitat is strongly regulated by the trade-off assumed (Fig. 5 and Fig. S5). Predictably, if the energetic cost is too high, hotbunking is not a desirable trait. For large cost to growth (small α) hotbunking diazotrophs do not survive (Fig. 5 A and C). On the other hand, for very little cost to growth (large α) and a high reduction in iron requirement (low γ), the hotbunking diazotrophs outcompeted the control group and expand into even more oligotrophic regions (Figs. 4C and 5B and Fig. S5 E and F). Simulations with lower requirements for iron (γ smaller) lead to a larger biomass of unicellular diazotrophs at any location than in a simulation

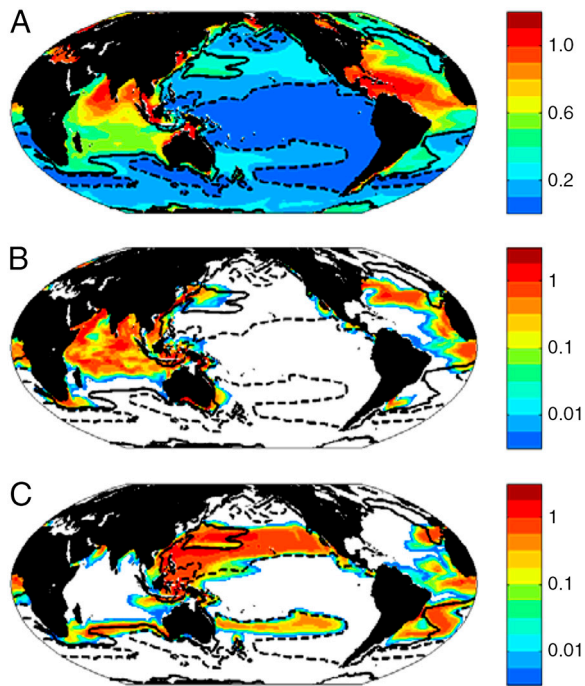


Fig. 4. Hotbunking unicellular diazotrophs inhabit waters with lower iron than the *Trichodesmium*-analogs and the unicellular control group (see Fig. S5) in the numerical simulations. Results are shown here for simulation where the hotbunking group have 90% of the growth rate and 60% the iron requirement of the control group ($\alpha = 0.9$, $\gamma = 0.6$, circle in Fig. 5). In this simulation the hotbunking diazotrophs outcompete the hypothetical control group everywhere. Annual mean model results for 0–50 m: (A) iron concentration (nM); (B) *Trichodesmium*-analogs (mgCm^{-3}); (C) hotbunking unicellular diazotrophs (mgCm^{-3}), analogs of *Crocosphaera watsonii*. Dashed and solid contours indicate 0.1 nM and 0.3 nM Fe respectively, and no shading (in B and C) indicates areas devoid of diazotrophs. The distribution of the diazotrophs is broadly consistent with those suggested by observations (4, 10). The model also includes analogs of several types of nondiazotrophic phytoplankton (diatoms, other large eukaryotes, *Prochlorococcus*, and other small phytoplankton). For less advantageous combinations of α and γ , the hotbunking group outcompete the control group only in the most iron-depleted regions (see Fig. 5 and Fig. S5).

with higher γ ; the same iron supply could support a higher biomass. Sharing cellular iron between photosynthesis and nitrogen fixation machinery allowed the model hotbunking diazotrophs to not only expand their habitat, but also to have higher biomass per mole of available iron. These two benefits of hotbunking lead to increased global unicellular diazotroph biomass than simulations that did not include this adaptation (Fig. 5C). As a consequence, total global nitrogen fixation increases between simulations with α and γ combinations that shifted the population of unicellular diazotrophs from only control types to only hotbunking types (Fig. 5D). For the simulation shown in Fig. 4 (circles in Fig. 5) there was a 10% increase in global autotrophic nitrogen fixation relative to simulation with no hotbunkers (e.g., the white area of Fig. 5A). The overall affect of hotbunking on total global nitrogen fixation would be less than this value, because recent evidence demonstrates that heterotrophic nitrogen fixers also contribute significantly to marine nitrogen fixation (44).

Enzymes are agents for chemical transformations in the cycling of elements on Earth. Whereas genomic and transcriptional techniques can monitor the potential for enzyme synthesis, proteomic methods are now capable of their direct quantitation. Many require metals and hence reside at the intersection of coupled biogeochemical cycles. Our study demonstrates an application of proteomic technologies to study iron conservation in marine nitrogen fixation, where the important diazotroph *Crocosphaera* appears to have evolved a complex diel proteome

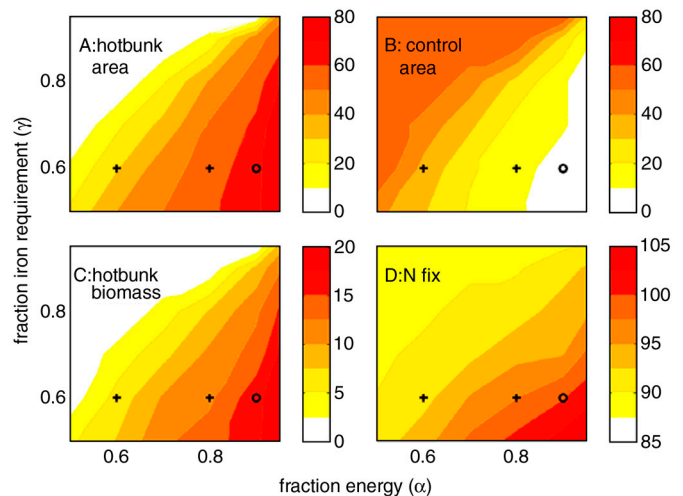


Fig. 5. Results from a series of global circulation/ecosystem model numerical simulations with different assumptions on the trade-off between reduced growth and iron requirement (values of α and γ). (A) Area (10^6 km^2) inhabited by hotbunking unicellular diazotrophs. (B) Area (10^6 km^2) inhabited by the control nonhotbunking unicellular diazotrophs. (C) Global abundance of hotbunking unicellular diazotrophs (TgC/y). (D) Changes in global total nitrogen fixation (TgN/y). With increased α (a smaller cost to growth) and a decreased γ (a lower requirement for iron) there are more regions where diazotrophy is supported, there is higher biomass of unicellular diazotrophs, and consequently higher total global autotrophic nitrogen fixation rates. If the energetic cost is low enough the hotbunking diazotrophs outcompete the control group everywhere (white regions of B). “Area” is calculated as the ocean surface area where the respective unicellular diazotrophs biomass is greater than $10^{-3} \text{ mgCm}^{-3}$. Nitrogen fixation rate is calculated from all three types of diazotrophs included in the simulation (hotbunking unicellular diazotrophs, control group, and *Trichodesmium*-analogs). The circle denotes parameter values used for results shown in Fig. 4 and crosses for those in Fig. S5.

cycle that results in a reduction of iron-metalloenzyme inventory. The process elucidated by the molecular-level measurements described here appears to be of global importance. With rising atmospheric CO_2 suggested to cause increases in marine nitrogen fixation (45), as well as a potential decrease in iron availability to marine phytoplankton in general (46), understanding the biochemical adaptations for iron scarcity in diazotrophs and their implications for marine primary productivity will be of increasing importance.

Methods

Axenic cultures of *Crocosphaera watsonii* were grown in 50 medium (15) on a 14:10 light dark cycle. Frozen pellets were resuspended, sonicated, and centrifuged, and supernatants were solvent precipitated at -20°C . Precipitated protein was resuspended, reduced, alkylated, and trypsin-digested. The whole cell lysate digests were analyzed using liquid chromatography mass spectrometry (LC-MS) using a Paradigm MS4 HPLC system with reverse phase chromatography, a Michrom ADVANCE source, and a Thermo LTQ ion trap mass spectrometer. Two-dimensional chromatography involved adding an offline strong cation exchange separation using a salt gradient prior to reverse phase separation. Nine specific peptides were selected for quantitative analyses via selected reaction monitoring (SRM). Known amounts of heavy labeled versions of each peptide of interest (AQUA peptides; Sigma) were added as internal standards to *Crocosphaera* peptide extracts and analyzed using a Thermo Vantage TSQ Triple Quadrupole Mass Spectrometer with the LC and source as described for the LTQ. Peptide standard sequences and transition information are listed in Table S2. Linear Ion Trap mass spectra were processed using SEQUEST and PeptideProphet, and spectral counts were tabulated in Scaffold 2.0, with a false positive rate of less than 1% (31). Numerical simulations used the MIT general circulation model (MITgcm). The physical circulation flow fields and diffusion, constrained by observations (47), transport the organic and inorganic components of the ecosystem model. The ecosystem model was adapted from Monteiro et al. (10) and includes several types of autotrophs including several nondiazotrophs (analogs of diatoms, other large eukaryotes, *Prochlorococcus*, and other small

phytoplankton) and three types of diazotrophs (analogs of *Trichodesmium*, *Crocospaera*, and a hypothetical control group of unicellulars that could not share cellular iron between photosynthesis and nitrogen fixation machinery). Other abundant groups of marine diazotroph groups (3) such as *Richelia* sp. symbionts found inside the diatom *Rhizosolenia* spp. and the uncultivated unicellular group A (UCYN-A) are not represented in this study due to lack of information on their iron requirements (4, 44).

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Supporting Information

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SI Methods

Culture Technique and Treatment. Axenic cultures of *Crocospaera watsonii* strain WH8501 from the Waterbury laboratory culture collection were grown in nitrogen-free iron-replete SO medium (1) at 28 °C under a 14:10 light-dark cycle at 150 $\mu\text{E m}^{-2} \text{sec}^{-1}$ in polycarbonate vessels, unless otherwise noted. All cultures were verified as axenic prior to each experiment by marine purity broth test (2) and during the experiment by examination of DNA staining microscopy slides used for cell counts (see *Microscopy and Flow Cytometry* below). Three distinct diel experiments were conducted to obtain proteome samples. The day-night experiment consisted of discrete day and night samples collected from a 1 L culture at 10 h into the light period and 7 h into the dark period during exponential growth and analyzed for relative abundance of major proteins (1D chromatography; Fig. 1B) and deeper proteome comparison (2D chromatography; Fig. S1B). The diel experiment consisted of a 1.5 L culture grown to late log phase in a gently stirred magnetic culture vessel (Nalgene) sampled every 2–3 h over a diel cycle (under dim red light at night) for protein, microscopy (Fig. S3A), and flow cytometry (Fig. S3 D–F) analyses. Protein samples were analyzed for relative abundance of major proteins (1D chromatography; Fig. 2) and absolute protein abundances of selected targets (Fig. 3 A–H). A biological triplicate experiment was also conducted using 3 1 L cultures in glass furnbach flasks in midlog growth phase. Protein samples were taken 6.5 h into the light cycle and again at 3 and 6 h from onset of darkness. Two night sampling times were used to capture the transient nature of the nitrogenase expression peak. The triplicate cultures were analyzed for absolute protein abundances of selected targets, and used for an additional estimate of the overall reduction in iron-metalloenzyme inventories at each time point (see main text). An iron stress experiment was conducted with two 250 mL cultures, one with replete iron and one transferred into media with no added iron (Fig. S1 C–E). In late log phase, each culture was split into two flasks and 500 nM of the iron binding ligand desferrioxamine B (DFB, Sigma) was added to one high and one low-iron replicate. This resulted in four treatments: high iron with and without DFB and low-iron with and without DFB. In all experiments, biomass was harvested 24 h after the addition of DFB during the light period, and centrifuged at 12,400 rcf for 20 min at 4 °C, decanted and microcentrifuged at 6,700 rcf for 8 min, decanted and frozen at –80 °C.

Nitrogenase Activity and Glycogen Content. Nitrogenase activity and glycogen content were measured as previously described (3) on a separate culture experiment in 14:10 light-dark cycle. Briefly, nitrogen fixation was measured using acetylene-reduction assay on a Shimadzu gas chromatograph. Glycogen abundance was measured by digestion of amyloglucosidase and amylase followed by a hexokinase/glucose-6-phosphate dehydrogenase assay coupled to NADP⁺ reduction.

Microscopy and Flow Cytometry. Cells were preserved in 0.125% final concentration ultrapure transmission electron microscopy (TEM) grade glutaraldehyde (Tousimis Research Corporation), incubated at room temperature in the dark for 15 min, flash frozen in liquid nitrogen and stored at –80 °C. 200 μL of thawed cells were collected on 0.22 μm Poretics polycarbonate filters (Osmonics Inc.) and mounted on glass slides using Vectashield hard set mounting media with DAPI (Vector Laboratories, Inc.). *Crocospaera* cells were counted by epifluorescence microscopy and cultures were examined for bacterial contamination on

a Zeiss Axioplan 2 microscope (Carl Zeiss, Inc.). For flow cytometry preserved cells were thawed, diluted with 0.22 μm filtered sterile seawater and stained with SYBR Green I fluorescent dye (Invitrogen) at a final concentration of 10 \times . Sodium citrate was added to the mixture at a final concentration of 10^{–5} M. A FACS-Calibur flow cytometer (Becton Dickinson) was used with filtered seawater for the sheath fluid and 2.5 μm fluorescent beads for standardization (Calibrite, Becton Dickinson). Data was analyzed with WinMDI 2.8 software (J. Trotter, the Scripps Institute).

Protein Extraction and Digestion. Pellets were resuspended in cold 100 mM ammonium bicarbonate (pH 8.0, ambic hereon) and sonicated on ice with a microtip twice for 4 min on 70% duty cycle with a 5 min pause between sonication steps. Samples were centrifuged for 30 min at 6,700 rcf and 4 °C, and supernatants were precipitated overnight in 100% acetone at –20 °C, with the exception of the diel and iron experiments that were precipitated overnight in 50% acetone-50% methanol and 0.5 mM HCl at –20 °C. Precipitated protein was collected by centrifugation at 6,700 rcf for 30 min at 4 °C and dried by speed vacuum. Protein was resuspended in 0.1 mL of 6 M urea 0.1 M ambic and shaken at 400 rpm for 20 min at room temperature, then at 95 °C for 15 min. Aliquots were taken for protein determination by DC assay using bovine serum albumin as a protein standard (BioRad Inc.). Samples were reduced with 10 mM dithiothreitol (DTT) at 56 °C for 1 h, alkylated with 30 mM iodoacetamide for 1 h, and incubated with 40 mM of DTT for 1 h, prior to dilution in 1 mL ambic and digestion with trypsin for 16 h at 37 °C (1:50 ratio with total protein, Promega Gold Mass Spectrometry Grade, Promega Inc.). The sample was concentrated by speed vacuum for 3 h and stored at –80 °C. Aliquots for liquid chromatography mass spectrometry (LC-MS) were mixed 2:1 with ambic and injection volumes were adjusted to achieve equal protein loading in each experiment.

Shotgun Mass Spectrometry. The whole cell lysate digests were analyzed using a peptide Cap Trap in-line with a reversed phase Magic C18 AQ column (0.2 \times 50 mm, 3 μm particle size, 200 Å pore size, Michrom Bioresources Inc.) on a Paradigm MS4 HPLC system (Michrom Bioresources Inc.) at a flow rate of 4 $\mu\text{L min}^{-1}$. A LTQ linear ion trap mass spectrometer (Thermo Scientific Inc.) was used with an ADVANCE nano-capillary electrospray source (Michrom Bioresources Inc.). The chromatography consisted of a hyperbolic gradient from 5% buffer A to 95% buffer B for 300 min, where A was 0.1% formic acid (Michrom) in water (Fisher Optima) and B was 0.1% formic acid in acetonitrile (Fisher Optima). The mass spectrometer was set to perform MS/MS on the top 7 ions using data-dependent settings and a dynamic exclusion window of 30 s. Ions were monitored over the range of 400–2000 m/z. 2D chromatography consisted of strong cation exchange (SCX) chromatography offline, followed by reverse phase chromatography of the fractions. Strong cation exchange was run on a 1 \times 150 mm 5 μm particle size 200 Å pore size strong cation exchange column (Michrom Bioresources Inc.). A gradient was run from Buffer C (95% water, 5% acetonitrile, 0.1% formic acid), to filtered Buffer D (80% water, 20% acetonitrile, 0.1% formic acid, 1M NaCl) at 50 $\mu\text{L min}^{-1}$, 5 min at 2% D, ramping hyperbolically to 90% by 64 min, followed by 10 min of reequilibration at 2% D. 19–21 fractions were collected and analyzed by reverse phase chromatography using the 1D protocol described above except with a shorter 140 min gradient.

Shotgun Mass Spectra Data Processing. The 1,404,663 LTQ mass spectra collected in this study were searched using SEQUEST (Bioworks version 3.3, Thermo Inc.). An amino acid database for *Crocospaera watsoni* from the Joint Genome Institute Integrated Microbial Genomics server (downloaded in November 2008) was combined with a reverse sequence genome for false positive analysis for all subsequent analyses (labels with prefix CwatDraft are shortened to Cwat). The genome has not been manually annotated or closed; as a result we manually annotated the 100 most abundant proteins with diel periodicity found in Fig. 2, whereas JGI-IMG annotations were used otherwise. Database search results were further processed using the PeptideProphet statistical model (4) within Scaffold 2.0 (Proteome Software Inc.). Relative protein abundance was determined using Scaffold 2.0 for normalized spectral counting software operating on a 64-bit Ubuntu Linux workstation. Spectral counts are normalized across samples in each experiment in Scaffold, including technical replicates, to allow comparison of relative protein abundance. For the 2D experiment the spectral counts in each treatment (19 and 21 fractions each) were summed prior to normalization across samples. For the 2D day-night dataset protein identifications results from SEQUEST used filters of $\Delta\text{CN} > 0.1$, $>30\%$ ions, Xcorr vs charge state of 1.9, 2.4, and 2.9 for +1, +2, +3 charges respectively, and peptide probability $<1e-3$ for protein identification (Dataset S1). For all other datasets (Dataset S1), Scaffold was used for protein identifications with protein and peptide probability settings of 99.9 and 95%, respectively, with one tryptic peptide required for identification. Both approaches (SEQUEST and Scaffold) resulted in false positive rates of less than 1% (5). For cluster analysis data were log transformed, centered around the mean, and normalized with the sum of the squares equaling 1, in Cluster 3.0 (6). Analysis was performed using Kendall's tau nonparametric distance metric cluster and the dendrogram was displayed using self-organizing mapping with Pearson's correlation metric (6, 7).

Targeted Mass Spectrometry. Peptides were designed as internal standards for quantitative analyses by selected reaction monitoring (SRM) using the global shotgun mass spectrometry results. Peptides were in the mass range of 800–2400 Da and lacked methionine or cysteine. SRM reactions for monitoring each peptide were selected based on successful peptide identification in the global proteomic analyses as well as those theoretically predicted by Protein Prospector MS Digest (<http://prospector.ucsf.edu/prospector>). Heavy isotope-labeled versions of the specific peptides (8) were obtained from Sigma-Aldrich in defined concentrations calibrated by amino acid analysis (AQUA Peptides) and handled according to the manufacturer's instructions. Heavy and native sequences of peptides, their SRMs, and collision energies are shown in Table S2. SRM analyses were performed on a Thermo Vantage TSQ Triple Quadrupole Mass Spectrometer with a Michrom Advance ion source. The SRM reactions were validated through tuning the mass spectrometer on heavy labeled reference peptides, introduced by direct infusion, to select the most abundant transitions for monitoring and to optimize the collision energies. Reverse phase chromatographic separation consisted of a peptide Cap Trap in-line with a reversed phase Magic C18 AQ column (0.2 × 50 mm, 3 mm particle size, 200 Å pore size) on a Paradigm MS4 HPLC system at a flow rate of 4 $\mu\text{L min}^{-1}$ with a gradient from 5% buffer A then to 45% buffer B over 25 min and to 95% buffer B in 10 min. Buffer A and B are the same as Shotgun LC-MS analysis above. Q1 operated in 0.2 FWHM resolution and Q3 operated in 0.7 FWHM resolution. SRM reaction monitoring was scheduled based on individual peptide pair chromatographic retention times and the cycle time was 1.6 s (Table S2). Linear behavior of each peptide was validated by creating a standard curve from 1.5 to 1,650 fmol of the heavy labeled reference peptides in 1 μg *Crocospaera* tryptic

peptide sample matrix. For quantitative analysis, 150 fmols of each heavy labeled peptide was added as an internal standard to *Crocospaera* peptide samples (1 μg protein per injection on the mass spectrometer). The amount of each target peptide of interest was calculated by ratio to the corresponding internal standard peptide abundance measured via SRM. Each biological sample was analyzed by mass spectrometry in triplicate.

Metabolic Iron Use Estimates. Absolute protein abundances were measured by targeted mass spectrometry (Fig. 3 A–H) and normalized to total protein extracted (Fig. S3B). These values were used to estimate the iron use by metabolism (Fig. 3I) using quantities of iron nitrogenase, molybdenum-iron β subunit, PsaA, cytochrome b6, and cytochrome c550 proteins to estimate iron involved in nitrogen fixation (former 2) and photosynthesis (latter 3) using the stoichiometries for each protein complex (Table S1). Assumptions of a 1:1:1 ratio of photosystems I, II, and ferredoxin were used here as in previous studies (9, 10), with the modification that only two iron atoms were attributed to PSII in our study (for 1 nonheme iron in PSII and 1 heme iron in cytochrome b-559), because we have direct measurements of the abundant cytochrome c550 [Fig. 3G; c550 can be found as part of the PSII complex and in soluble form (11)]. Because photosystem I, which we measure directly, contains far more iron than photosystem II and ferredoxin (Table S1), these assumptions should have a small effect on the calculations. Due to a potential inability to detect the dominant photosynthesis-associated ferredoxin (see *SI Results and Discussion*), the 4Fe4S stoichiometry was conservatively used. For the hypothetical situation, calculated in Fig. 3I, where nitrogenase substitutes ferredoxin for flavodoxin as its primary electron carrier, we used the 2Fe2S stoichiometry based on observations that a 2Fe2S ferredoxin increased slightly in abundance at night, and a conservative 1:1 stoichiometric substitution (ferredoxins are known to be more efficient than flavodoxins). Protein abundances were converted to per cell concentrations by multiplying by the measured total protein from extracted cells and dividing by the cell concentration at each time point. Whereas the measured peptide signals associated with targeted peptides appeared to be robust during the diel experiment (Fig. 3) suggesting reproducible results, we interpret these as lower estimates due to the potential for inefficiencies associated with the protein extraction. The extraction protocol was chosen for minimal processing to improve quantitative reproducibility, yet some proteins such as membrane associated proteins were likely not completely extracted. These calculations assume that the metalloproteins are populated with iron. These iron use by metabolism estimates should be lower than total cellular iron due to use in other unconstrained iron reservoirs such as other metalloenzymes (e.g., aconitase), bacterioferritin storage, and chaperones. As a result, we use the terminology “iron-metalloenzyme inventory” to describe the total number of iron atoms that could be associated with each metabolism based on our measurements of peptide abundance.

Global Circulation Ecosystem Modeling. We used the global three-dimensional physical-biogeochemical-ecosystem model of Follows et al. (12) to investigate the potential global importance of hotbunking trophs. The model used here had various adaptations discussed in Dutkiewicz et al. (13) and Monteiro et al. (14), in particular to explicitly include nitrogen-fixing phytoplankton (diazotrophs) and a representation of denitrification. We use the Massachusetts Institute of Technology (MIT) general circulation model (MITgcm) and the physical circulation flow fields and diffusion are state estimates as produced by ECCO-GODAE and have been constrained by observations (15). The biogeochemical cycling of nitrogen, phosphorus, silicon and iron is represented and the ecosystem model includes several types of autotrophs consisting of nondiazotrophs and diazotrophs. The model re-

solves a diatom-analog, an analog of other large eukaryotes, high and low light *Prochlorococcus*-analogs, an analog of other small phytoplankton, *Trichodesmium*-analog, hotbunking unicellular diazotrophs (analogs of *Crocospaera watsonii*) and nonhotbunking unicellular diazotrophs (a hypothetical control group). Non-diazotrophs' growth could be limited by phosphorus, silica (for diatoms), iron or nitrogen; diazotrophs growth is limited by phosphorus or iron. Growth rates are parameterized as a function of light, temperature and nutrient concentration. Simple allometric trade-offs are imposed (see refs. 12 and 13 for more discussion), in particular smaller phytoplankton have lower nutrient half-saturations (more efficient at acquiring nutrients) than large phytoplankton. The model also resolves explicit grazing and sinking and all organic and inorganic components were advected and diffused by the physical fields. Parameter values for the different phytoplankton analogs are taken from those found most fit in a previous simulation representing 10s of phytoplankton types (14). Here, given the number of simulations required, we only resolve the above 8 functional types. Additional modifications over (14) are to use the aeolian iron source fields as modeled by (16) that includes estimation of the variable solubility of the iron in seawater, as well as to include a sedimentary source of iron [following Elrod et al. (17)]. These modifications to the iron cycling parameterization used in Monterio et al. (14) provided more iron in some regions of the ocean (e.g., Equatorial Pacific) that had been potentially too low in Monterio et al., though this also leads to concentrations too high relative to observations in other regions (e.g., North Atlantic).

The hotbunking diazotrophs were modeled with a lower cellular iron to nitrogen ratio (R) relative to the control group as a consequence of their ability to share iron between photosynthesis and nitrogen fixation processes. But because this is an energetically expensive process, we also assumed that the hotbunking diazotrophs had a lower growth rate (μ) than the control group:

$$\mu_H = \alpha\mu_C, \quad R_H = \gamma R_C \quad \text{where } \alpha, \gamma < 1.$$

The subscript H refers to the hotbunking group and C refers to the control nonhotbunking group. The trade-off between reduced growth and iron requirement is therefore represented by the values of α and γ . A larger value of α indicates less energetic cost, and a larger value of γ indicates iron requirements closer to the control group. We conduct several simulations with different combinations of α and γ . The laboratory studies described here suggest a decrease of iron requirement by as much as 40% ($\gamma = 0.6$), thus we consider a range of $\gamma = [0.5, 0.6, 0.7, 0.8, 0.9, 0.95]$. Crude back of the envelope calculations on the energetic cost suggest that it is at least 5%, but potentially several fold higher. Thus we consider the cost to growth $\alpha = [0.5, 0.6, 0.7, 0.8, 0.9, 0.95]$. Fig. 5 is constructed from results from the 36 simulations with the combinations of these values of α and γ . The simulations were run for 10 years and results from the annual mean for the tenth year are shown in Fig. 4 and Fig. S5. In all the experiments we set $\mu_C = 0.7 \text{ d}^{-1}$, $K_{\text{FeC}} = 0.36 \text{ nM}$, and $R_C = 7.5 \times 10^{-4} \text{ mol Fe/mol N}$. We relate the change in the iron cellular demand (R) to the half saturation coefficient of growth (K_{FeC}) following Verdy et al. (18).

SI Results and Discussion

Global Proteome Results. Relative quantitation results obtained using 1D chromatography LC-MS analysis of the day-night experiment resulted in 477 unique protein and 3,800 unique tryptic peptides identifications with a false positive rate (FPR) of 0.65% (5), based on 20,506 total peptide identifications from 12 LC-MS injections (6 technical replicates per sample) and a total of 380,015 mass spectra (Dataset S1). Additional 2D chromatography analyses of the day-night experiment identified 1,108

unique proteins and 3,540 unique tryptic peptides (Dataset S1) for 18% coverage of the genome open reading frames and a 0.11% FPR, based on 21,620 total peptide identifications over 40 LC-MS fraction injections (~20 fractions per sample) for a total of 388,076 mass spectra. The diel experiment and iron stress experiments also identified 552 and 589 proteins, respectively, for a combined total of 724 proteins identified from the 1D global proteome experiments (day-night, diel, and iron stress experiments, see Dataset S1 for complete list of protein identifications, spectral count values of technical replicates, and fold change calculations). As is typical of proteomic experiments, each individual injection provided a large number of protein identifications, whereas repeat injections and 2D chromatographic separation provided confidence in the robustness of the results and increased proteome coverage incrementally with discovery of lower abundance tryptic peptides.

Examination of Flavodoxin and Ferredoxin Expression by Iron Stress Experiment. Whereas ferredoxin and flavodoxin are typically involved in electron transport in photosynthesis, an electron donor is also needed for the nitrogenase enzyme in the diazotrophic cyanobacterial species. Ferredoxins contain iron-sulfur centers and have a higher catalytic efficiency than the iron-free flavodoxin (19). In marine phytoplankton flavodoxin is generally considered a low-iron alternative to ferredoxin that is upregulated under iron stress in eukaryotic algae. Based on this expression behavior, flavodoxin has been applied as an iron nutrition diagnostic (20). Yet, as described in the main text, the expression of flavodoxin in *Crocospaera* diverged from this behavior and was clearly present and abundant at night in *Crocospaera* under iron-replete conditions (Fig. 3D).

The effect of reduced iron availability on the flavodoxin expression in the *Crocospaera* proteome was examined through the growth of cultures under high and low total iron concentration, as well as with and without the addition of the strong iron chelator desferrioxamine B (DFB; Fig. S1 C–F). Cell counts showed a cessation of growth with the addition of DFB in both iron conditions, whereas the no iron added treatment did not show a physiological effect, as evidenced by growth similar to that of the standard (+Fe) media. This similar growth was expected based on the lack of multiple transfers in low-iron media, and allowed a low-iron treatment under similar growth rates for comparison with the replete iron. DFB is an exogenous siderophore and is known to greatly reduce the iron available to phytoplankton by depressing the abundances of the free ionic form and inorganic chemical species of the metal (21, 22). Flavodoxin protein expression under iron stress was examined during the light period because the influence of iron stress would potentially be difficult to differentiate from the high dark period expression observed (Fig. 3D). Flavodoxin (Cwat4291) was more abundant under high iron relative to no iron added (Fig. S1D), although both of these light period treatments were >5-fold less than the large peak in flavodoxin observed during the dark period in the day-night experiment (Fig. S2 and Fig. 1B in spectral count units; also Fig. 3D). Flavodoxin during the day period did not appear affected by DFB. Additionally, ferredoxin (Cwat3830) was 3-fold more abundant under high iron conditions relative to low-iron conditions (Fig. S1D), regardless of whether DFB was added. The absence of a light period increase in flavodoxin abundance under these two iron stress treatments (–Fe, and +DFB) suggests that flavodoxin's primary role is as the major electron donor protein for nitrogen fixation, rather than as a low-iron substitute for ferredoxin associated with photosynthesis as observed in other phytoplankton (19, 20, 23). The dark period expression of flavodoxin even under iron-replete conditions may be an adaptation for the minimization of *Crocospaera*'s cellular iron demand.

Analysis of other putative ferredoxins in the genome revealed that three ferredoxin-like proteins lack more than one suitable

tryptic peptide for shotgun mass spectrometry detection. Because confidence in protein identification increases greatly with the identification of more than one tryptic peptide for a given protein, some of these small putative ferredoxin proteins may have been present but were not detected. Many open reading frames are annotated as ferredoxin in the current *Crocospaera watsonii* genome, but of these only four were identified in our proteome samples. Cwat3830 is a 2Fe-2S ferredoxin, whereas Cwat5974 is misannotated and is actually *psaC*. Cwat5441 and Cwat1598 are barely detected and are likely too large to be real ferredoxins. We suspect that due to the small size of this protein, the major cellular ferredoxin has gone undetected in our study due to the lack of a suitable tryptic peptide within the sequence. Cwat1273, Cwat1274, and Cwat0916 are possible candidates for the major ferredoxin because all are lacking more than one suitable internal tryptic peptide needed for the shotgun proteomic techniques applied here.

Several other notable changes in global protein expression under iron stress were observed as shown in Fig. S1 *E* and *F* and Dataset S1, including a tripling in relative abundance of a possible carbon concentrating mechanism/microcompartments protein (Cwat5294) under low-iron without the addition of DFB. Ribulose-bisphosphate carboxylase (Cwat2714) was two to three-fold more highly expressed under low-iron, and fructose-bisphosphate aldolase (Cwat1319), which is involved in carbohydrate synthesis, was 20-fold higher under iron stress. The upregulation of these proteins suggests that reduced iron availability results in cellular demand for more efficient inorganic carbon fixation. *Crocospaera* strains have been observed to secrete significant amounts of extracellular organic material (24). The increases in carbon fixation proteins under iron stress observed here may be connected to these organic secretions, perhaps to promote bacterial associations that produce siderophores and aid iron scavenging and acquisition (25).

Other Proteins of Interest. Cytochrome c550 is a monoheme cytochrome containing 1 Fe atom present as a component of the PSII complex of cyanobacteria and involved in the stabilization of the PSII Mn cluster (26). It has also been isolated as a soluble protein with a possible additional role in anaerobic carbon and hydrogen metabolism (11). This protein had a particularly strong diel pattern decreasing 15-fold during the dark from the daytime maximum as measured by targeted proteomic analysis (Fig. 3*G*). This decreased abundance during the dark period should reduce the oxygen production capability of PS II, which would be beneficial to the preventing damage to the nitrogenase complex. Cytochrome b6 also decreases in abundance by more than 2-fold during the dark period in our targeted proteomic analyses (Fig. 3*H*). This protein is a component of the b6f complex, which contains four iron containing heme groups and an Fe₂S₂ cluster (27) and is reported to have a dual function in photosynthesis and respiration in cyanobacteria (28).

Two heme oxygenases (Cwat4859, Cwat2459) were identified in the proteome, one with a dark period maxima (Cwat2459) and the other with a day period maxima (Cwat4859) in relative abundance (Fig. S2). Heme oxygenases are capable of releasing iron from hemes, for example by pathogenic bacteria to gain nutritional iron from host cells (29). This enzyme has also been shown to be involved in phycobilin synthesis in cyanobacteria through the conversion of hemin to ferric-biliverdin IX_α (29). We hypothesize that one of these heme oxygenases (Cwat2459) is involved in releasing the iron from cytochrome heme groups at night for use in the nitrogenase complex, whereas the other (Cwat4859) is involved in the conversion of hemin to ferric biliverdin IX_α phycobilin. Peptidase S41A also showed strong diel cycling with a presence during the photoperiod (Fig. S2), suggestive of its involvement in the changing proteome composition throughout the diel cycle (e.g., Fig. 2). Measurements of the

ribosomal protein L3 (Cwat2439) are consistent with two more highly active periods of protein synthesis per diel cycle (Fig. S3*C*), and is consistent with the two peaks of ribosomal protein activity observed in the diel global proteome in Fig. 2. Plastocyanin, the copper containing water soluble electron carrier, was identified in the *Crocospaera* proteome (Dataset S1). In addition, the iron-manganese superoxide dismutase present in the genome (Cwat6299) was not found in any of the proteome analyses here, whereas a nickel superoxide dismutase was detected in the 2D proteome (Cwat2983; Dataset S1), consistent with the theme of iron conservation in *Crocospaera*.

Connecting Cellular Iron Estimates with Iron Biogeochemistry, Uptake, and Cell Division. Cellular iron was estimated during a diel cycle by combining light period ferrous iron uptake (30), our proteomic measurements, and allowing cell division after dawn. As mentioned in the main text, the motivation for this effort was to reconcile our diel protein cycling observations with the observation of increased cellular iron during the dark period in Tuit et al.'s study (31) by overlaying the two processes of cell division and ferrous iron uptake (and their temporal variability) onto our iron-metalloenzyme inventories. Before describing the results of this calculation below, we describe what is known about each of these processes with respect to *Crocospaera*.

Cell division is known to occur solely during the photoperiod in *Crocospaera* (e.g., Fig. S3*A*), and division must divide the cellular iron between two daughter cells, perhaps also involving some cellular leakage in the process. With growth rates of ~0.5 d⁻¹ observed in Tuit et al. (31), 65% of cells are dividing daily, contributing to a 40% decrease in iron per cell when averaged across the population. Hence, cell division alone can account for a significant amount of the lower photoperiod cellular iron content observed by Tuit et al. In addition, a potential iron uptake pathway for *Crocospaera* involves a combination of photochemical reduction of the natural iron ligands that dominate seawater iron chemistry or Fe-EDTA complexes in laboratory media and uptake of the resultant ferrous iron (30, 32, 33). This scenario would cause production of Fe(II) during the light period followed by rapid reoxidation and complexation by strong ligands in darkness at tropical and subtropical temperatures as modeled by Weber et al. (30) (Fig. S4*A*). In addition, the *Crocospaera watsonii* genome contains genes for the ferrous iron transporter, which a recent microarray transcriptional study showed are greatly upregulated during the photoperiod (34), consistent with light period photochemical ferrous iron source. Ferrous iron is very rapidly oxidized to ferric iron in seawater (35), and hence only exists in appreciable amounts during the photoperiod under the tropical temperature conditions of *Crocospaera*'s niche (30). *Crocospaera*'s genome does not appear to contain genes for other important iron acquisition mechanisms such as an iron reductase system, siderophore biosynthesis, or siderophore transport (36). These facts taken together suggest a scenario where *Crocospaera* acquires ferrous iron during the light period, resulting in iron accumulating through the light period as shown in the schematic in Figs. S4*A* and *B*.

Combining the influence of the temporal processes of cell division and ferrous iron uptake, with our diel iron-metalloenzyme inventory calculations allows a schematic of diel cellular iron budget to be estimated that might be expected in natural environments. Ferrous iron uptake was calculated to be proportional to the light period ferrous iron availability from Weber et al. (30) (Fig. S4*A*), where the timing of iron uptake was inferred from chemical modeling of ferrous iron abundance (30), whereas the magnitude of uptake was tuned to minimize storage category (see below) to emphasize the extent of intracellular trafficking and to allow a steady-state cellular iron abundance. The iron estimated to be involved in nitrogen fixation and photosynthesis (Fig. 3*I* and *J*) was then subtracted to make a rough estimate of

the remaining cellular iron that might be found in storage (bacterioferritin), other enzymes (e.g., aconitase), and chaperones (storage category from hereon). This storage/other reservoir is estimated to be quite dynamic during the diel cycle as shown in red in Fig. S4B. No efflux term is needed in these calculations to approximate steady-state growth. Increasing the ferrous iron production and resultant uptake rate in this calculation increases this reservoir as well as cellular iron in subsequent generations, as might be expected during transient pulses caused by dust input or increased light fluxes, presumably with iron storage in bacterioferritin. Our estimates of cellular iron associated with iron metalloenzymes involved in photosynthesis and nitrogen fixation are consistent with previous quota studies (10, 31, 37) differing by a factor of less than 2, which is within the error observed in those cellular iron measurements. This schematic also highlights the possibility of apo forms of nitrogen fixation and photosynthetic metalloenzymes existing, a likely scenario given the challenge of coordinating the movement of iron between intracellular reservoirs as estimated here. Future research on how chaperones and metal trafficking are involved in the diel metalloenzyme cycling we have observed would be particularly interesting in *Crocospaera watsonii*.

Global Circulation Ecosystem Modeling. The simulations produced nutrient and biomass fields that compare well with observations. Total global primary production was between 38.2 and 38.8 GtC/y and total global nitrogen fixation rates were between 90 and 107 TgN/y (depending on the values of α and γ). The distribution of the different phytoplankton types compare well to observations, as shown in Follows et al. (12), Dutkiewicz et al. (13) and Monteiro et al. (14). Though never dominant in the model, diazotrophs could coexist with nondiazotrophs in regions where nitrate concentrations are low and there is sufficient iron and phosphorus for the diazotrophs to exist (38). Unicellular diazotrophs, due to their small size, can out-compete the *Trichodesmium*-analogs in more oligotrophic regions and their distribution and abundance are broadly consistent with those suggested by observations (14, 39). Observations of *Crocospaera* as summarized in ref. 14 suggest higher abundances in the tropical North Atlantic (40, 41), eastern subtropical North Pacific (42, 43) and Arabian Sea (44). *Crocospaera* have also been detected in the eastern subtropical North Atlantic (40) and eastern tropical South Atlantic (41), eastern subtropical and tropical North Pacific (42) and western South Pacific (45). They were however not detected in the western subtropical North Atlantic (40), eastern subarctic North Pacific, and eastern subtropical South Pacific (46). The pattern of presence and absence of *Crocospaera*-analogs is in general captured by the numerical model (Fig. 4), with concentration levels consistent in most regions with the observations (see ref. 14 for details). The iron concentrations in the Caribbean and eastern tropical North Atlantic are too high in this version of the model, leading to strong phosphorous limitation there. This discrepancy leads to less than ideal conditions for diazotrophs,

and concentrations there do not compare as well as in other parts of the oceans. A similar problem was found by Moore et al. (47). Presence, absence and abundances of *Trichodesmium*-analogs are also, in general, captured by the model. In particular the geographic patterns of *Trichodesmium* limited to regions with higher iron and *Crocospaera* existing in regions with lower iron matches patterns suggested by the observations of Campbell et al. (45), who found that in the southwest Pacific *Trichodesmium* appeared constrained to coastal areas with high iron, whereas *Crocospaera* dominated in offshore waters where Fe was low.

A series of experiments were conducted with varying degrees of trade-off between cellular iron to nitrogen reduction (γ) and cost to growth rate (α). These results (Fig. 5) indicate that the cost to growth cannot be too high (small α) or the hotbunkers do not survive. However, for lower cost to growth, hotbunking can confer an advantage in low-iron regions (Fig. 4 and Fig. S5). With the right combinations of α and γ the hotbunking diazotrophs could inhabit some model regions that would not support the control group (Fig. S5 B and D): The maximum area covered by unicellular diazotrophs increases by over 30% (from 59×10^6 km² to more than 79×10^6 km²; Fig. 5 A and B) from simulations where only the control group survived to ones where hotbunkers are the only unicellular group surviving. However, the region of parameter space (α , γ) where hotbunkers completely out-compete the control group (white area of Fig. 5B) is quite small, suggesting that the energetic cost cannot be too high and there must be a significant reduction in iron needs. With more advantageous values of α and γ , the hotbunking diazotrophs could even out-compete *Trichodesmium* in some specific areas of the model ocean, leading to a reduced global biomass of *Trichodesmium* in those simulations (compare Fig. 4A and Fig. S5 A and B).

The value of γ also affected the local concentration of hotbunking diazotrophs. Simulations with lower requirement for iron (γ smaller) leads to a larger biomass of unicellular diazotrophs at any location compared to simulations with higher γ : the same iron supply could support a higher biomass. This leads to a globally increased biomass of hotbunking unicellular diazotrophs (Fig. 5C) with lower γ , and consequently an increase in total global nitrogen fixation (Fig. 5D) by as much as 20% (from 90 GtN/y for simulations with no hotbunkers to 107 GtN/y for most extreme case of α and γ , bottom right Fig. 5D). We note that more reasonable values of the trade-off would suggest a 5 to 10% increase. Because over many regions of the ocean the non-diazotrophic phytoplankton are nitrogen limited, this increased source of fixed nitrogen led to increased primary production. Globally this increase was as much as 1.5% between a simulation with no hotbunking unicellular diazotrophs (poor combination of α and γ) and one where hotbunkers outcompeted the control group everywhere (advantageous α and γ).

Other Supporting Information Files

Dataset S1 (XLSX)

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Table S2. Heavy and native sequences of peptides, their SRMs, and collision energies

Protein description	Peptide name	Peptide sequence	Parent m/z	Product m/z	Collision energy	Start time (min)	Stop time (min)
Photosystem 1 protein PsaB (Cwat 1223)	psaB L	TPLANLVR	442.272	501.314	18	12.9	15.1
				572.351	21	12.9	15.1
	psaB H	TPLAN[L_C13N15]VR	445.78	685.436	21	12.9	15.1
				508.331	18	12.9	15.1
				579.368	21	12.9	15.1
Nitrogenase molybdenum-iron protein alpha subunit (Cwat3819)	MoFeA2 L	ILLEEIGLR	528.327	345.22	18	18	21
				716.39	19	18	21
	MoFeA2 H	I[L_C13N15]LEEIGLR	531.84	829.48	17	18	21
				345.22	18	18	21
				716.39	19	18	21
Nitrogenase NifH protein (Cwat 3818)	nifH2 L	LNTQMIHFVPR	678.37	655.368	27	15.5	19
				768.452	23	15.5	19
	nifH2 H	[L_C13N15]NTQMIHFVPR	681.88	899.492	22	15.5	19
				655.37	27	15.5	19
				768.45	23	15.5	19
Nitrogenase molybdenum-iron protein alpha subunit (Cwat 3819)	MoFeA1 L	LIADVLSTYPEK	674.872	373.208	29	15.2	17.6
				724.351	17	15.2	17.6
	MoFeA1 H	[L_C13N15]IADVLSTYPEK	678.38	837.435	20	15.2	17.6
				1122.567	21	15.2	17.6
				373.208	29	15.2	17.6
Cytochrome b6 of the b6f complex; PetB (Cwat 4239)	petB1 L	LEIQAISDDITTK	723.888	724.351	17	15.2	17.6
				779.378	26	15.2	17.6
	petB1 H	[L_C13N15]EIQAISDDITTK	727.397	837.435	20	15.2	17.6
				1122.567	21	15.2	17.6
				373.208	29	15.2	17.6
Photosystem 1 protein PsaA (Cwat 1222)	psaA1 L	VAVDVDPVPTSFEK	751.89	724.351	17	15.2	17.6
				779.378	26	15.2	17.6
	psaA1 H	VA[V_C13N15]DVDPVPTSFEK	754.897	892.462	28	15.4	17.9
				963.499	28	15.4	17.9
				779.378	26	15.4	17.9
Flavodoxin, long chain, PetE (Cwat 4291)	petE1 L	FVGLPIDEINQSDK	787.907	892.462	28	15.4	17.9
				963.499	28	15.4	17.9
	petE1 H	FVG[L_C13N15]PIDEINQSDK	791.415	963.499	28	15.4	17.9
				892.462	28	15.4	17.9
				963.499	28	15.4	17.9
Nitrogenase molybdenum-iron protein beta subunit (Cwat 3829)	MoFeB1 L	INFIPGFETYIGNLR	877.467	708.356	33	14.9	17.1
				904.477	27	14.9	17.1
				1019.504	24	14.9	17.1
				708.356	33	14.9	17.1
				904.477	27	14.9	17.1
	MoFeB1 H	INFIPGFETYIGN[L_C13N15]R	880.976	1019.504	24	14.9	17.1
				591.273	32	16.7	19
				948.426	31	16.7	19
				1158.563	22	16.7	19
				591.273	32	16.7	19
Nitrogenase molybdenum-iron protein beta subunit (Cwat 3829)	MoFeB1 L	INFIPGFETYIGNLR	877.467	948.426	31	16.7	19
				1158.563	22	16.7	19
				591.273	32	16.7	19
				948.426	31	16.7	19
				1158.563	22	16.7	19
Nitrogenase molybdenum-iron protein beta subunit (Cwat 3829)	MoFeB1 H	INFIPGFETYIGN[L_C13N15]R	880.976	402.246	11	22.5	25.5
				459.267	20	22.5	25.5
				572.351	20	22.5	25.5
				836.462	8	22.5	25.5
				1266.648	30	22.5	25.5
Nitrogenase molybdenum-iron protein beta subunit (Cwat 3829)	MoFeB1 H	INFIPGFETYIGN[L_C13N15]R	880.976	1379.732	20	22.5	25.5
				409.263	11	22.5	25.5
				466.285	20	22.5	25.5
				579.369	20	22.5	25.5
				843.479	8	22.5	25.5
Nitrogenase molybdenum-iron protein beta subunit (Cwat 3829)	MoFeB1 H	INFIPGFETYIGN[L_C13N15]R	880.976	1273.664	30	22.5	25.5
				1386.749	20	22.5	25.5

