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Divergent responses of Atlantic coastal and oceanic *Synechococcus* to iron limitation

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Marine *Synechococcus* are some of the most diverse and ubiquitous phytoplankton, and iron (Fe) is an essential micronutrient that limits productivity in many parts of the ocean. To investigate how coastal and oceanic Atlantic *Synechococcus* strains acclimate to Fe availability, we compared the growth, photophysiology, and quantitative proteomics of two *Synechococcus* strains from different Fe regimes. *Synechococcus* strain WH8102, from a region in the southern Sargasso Sea that receives substantial dust deposition, showed impaired growth and photophysiology as Fe declined, yet used few acclimation responses. Coastal WH8020, from the dynamic, seasonally variable New England shelf, displayed a multitiered, hierarchical cascade of acclimation responses with different Fe thresholds. The multitiered response included changes in Fe acquisition, storage, and photosynthetic proteins, substitution of flavodoxin for ferredoxin, and modified photophysiology, all while maintaining remarkably stable growth rates over a range of Fe concentrations. Modulation of two distinct ferric uptake regulator (Fur) proteins that coincided with the multitiered proteome response was found, implying the coastal strain has different regulatory threshold responses to low Fe availability. Low nitrogen (N) and phosphorus (P) availability in the open ocean may favor the loss of Fe response genes when Fe availability is consistent over time, whereas these genes are retained in dynamic environments where Fe availability fluctuates and N and P are more abundant.

To explore this acclimation potential in the North Atlantic context, we compared the responses of coastal and oceanic Atlantic *Synechococcus* to Fe limitation. Comparison of strain origins with trace metal and low-level nutrient measurements from the recent US North Atlantic Zonal Transect GEOTRACES expedition demonstrates large differences in nitrate, phosphate, and dissolved Fe between sites in the North Atlantic basin. The coastal strain, WH8102, was isolated from the New England shelf (Fig. 1 A and B), a region with a dynamic nutrient (12) and Fe (13) regime. Oceanic strain WH8102 was isolated from the permanently stratified southern Sargasso Sea, an oligotrophic site that consistently receives Fe from aeolian dust year round, with very high levels in the summer (8). The consistent dust supply results in higher dissolved Fe levels in the open ocean (~0.8 nM within the Sahara dust plume) relative to North American coastal waters (Fig. 1B).

To test their abilities to acclimate, both strains were rendered Fe limited by sequential transfer into no-Fe-added media and were rescued with varying concentrations of Fe’ (Fe’ defined as the sum of inorganic species). Cellular protein was harvested during log-phase growth to characterize Fe responses (Fig. 2 C and H). Protein abundance was measured following cell pellet extractions of single cultures grown at each prescribed Fe concentration, followed by high-resolution LC-MS proteomic analysis for relative abundance and targeted proteomics for absolute abundance (Materials and Methods).

**Significance**

Conventional knowledge suggests that coastal phytoplankton are less able to adapt to Fe limitation than open ocean species. Here we show that in contrast to the established paradigm, coastal *Synechococcus* from the New England shelf is capable of dynamic, multitiered Fe adaptation that allows it to thrive over a broad range of Fe concentrations by partitioning Fe among different uptake and storage proteins. This protein-based response is beneficial in high nitrogen (N) waters with low and variable Fe:N ratios. An oceanic Atlantic *Synechococcus* isolate lacks this adaptive response, suggesting the small yet significant N cost of retaining Fe response proteins offsets the benefit of Fe adaptability in the southern Sargasso Sea, where N is chronically scarce and Fe:N ratios are high.

**Materials and Methods**

The marine *Synechococcus* are a cosmopolitan group of cyanobacteria that contribute ~17% of marine net primary production worldwide (1) and whose diversity is linked to nutrient availability in both coastal and open ocean waters (2). Of the nutrients required by phytoplankton, the micronutrient iron (Fe) limits productivity in ~30% of the world’s oceans (3) and shapes *Synechococcus* genome content (4, 5). Seawater Fe concentrations in the euphotic zone can vary due to inputs from atmospheric, coastal, and upwelled sources. The open ocean, being more removed from these sources, is typically depleted in Fe relative to coastal waters. Consistent with these distributions, experiments on oceanic phytoplankton have found them able to grow at lower metal conditions than related coastal strains, implying a general phenomenon of open ocean phytoplankton being better adapted to low metal availability (6, 7).

The North Atlantic Ocean presents an extreme endmember with which to examine the coastal–open ocean phytoplankton adaptation phenomenon due to its large Fe flux associated with seasonal atmospheric dust deposition from the Saharan desert (8, 9). Moreover, there has been increasing evidence that Fe availability may be dynamic in coastal regions, with observations of Fe limitation of phytoplankton in eastern boundary upwelling regions upon upwelling of macronutrient-rich and Fe-depleted waters (10, 11). Whereas the diversity and distribution of *Synechococcus* strains have been linked to macronutrient availability (2), the considerable range of Fe levels across which *Synechococcus* thrive suggests they may also possess unique and diverse regulatory mechanisms for Fe acclimation.

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Ferredoxin
We observed that the growth rate and photophysiology (Fv/Fm) of oceanic Synechococcus WH8102 was proportional to Fe concentration (Fig. 2 A–D). In contrast, coastal Synechococcus WH8020 tailored its response to Fe limitation through the differential expression of Fe response proteins in a multilayered, hierarchical cascade by which cells partitioned and conserved Fe, and so thrived over a broad range of Fe concentrations. Fe levels had a minimal effect on growth rate (Fig. 2E), although a sharp decline in photosynthetic protein abundance was observed below 1 nM (Fig. 2F), followed by a decline in photosynthetic efficiency below 0.3 nM (Fig. 2F). Thresholds for photosynthetic efficiency and photosynthetic protein abundance were similar between strains (Fig. 2D, E, I, and J).

The global proteome of coastal Synechococcus WH8020 revealed large changes in the abundance of Fe sensing, acquisition, and storage proteins (Fig. 3A). Fe levels ≤1 nM led to induction of the Fe transport protein, IdiA, that assists in Fe uptake during periods of low Fe availability (14) and substitution of Fe-free flavodoxin for Fe-containing ferritin (15, 16). At Fe levels above 1 nM, coastal WH8020 produced ferritin, which stores Fe when it is abundant for later use when levels decline. Several additional Fe response systems were present in WH8020 that are also shared in the genome of coastal strain CC9311 isolated from the California Coastal Current, although the latter has additional genes including a ferrous Fe uptake transporter (feo) (Fig. 3B and SI Appendix, Table S1). The strong proteome response of coastal WH8020 to low Fe implies that coastal Atlantic waters may share aspects of the “patchwork mosaic” dynamics of Fe limitation and excess similar to California coastal waters (10, 11).

The multilayered response in coastal WH8020 appears to be regulated by two isoforms of the ferric uptake regulator protein (Fur), which acts as a transcriptional repressor when it binds intracellular Fe(II) (17). Interestingly, the two Fur isoforms quantified here appeared to respond to different Fe concentrations and may govern “Fe-limited” and “Fe-stressed” conditions, respectively (Fig. 3A).

In contrast to the coastal strain, oceanic WH8102 did not show a multilayered proteome response to Fe limitation and lacked many of the Fe response genes discussed above. Oceanic WH8102 lacks flavodoxin, although at least one of its ferredoxin proteins was still regulated by Fe levels (SI Appendix, Fig. S1). Both WH8102 and WH8020 genomes have more than one ferredoxin isoform that may have different physiological or regulatory roles (SI Appendix, Table S1). Additionally, the coastal WH8020 genome contains both ferritin (GenBank accession no. WB44_04810) and heme-based bacterioferritin (GenBank accession no. WB44_10870) (18, 19), whereas oceanic WH8102 lacks both.

This study belies the expectation that oceanic phytoplankton would tend to experience greater Fe stress than coastal phytoplankton (6) for two reasons. First, certain open ocean regions like the southern Sargasso Sea can have relatively high and stable dissolved Fe levels depending on their location (Fig. 1B). Second, as we show here, the underlying factor governing Fe acclimation capacity is not Fe availability alone, but the simultaneous pressure exerted by multiple scarce nutrients, in particular N, P, and Fe. The Fe adaptations of coastal WH8020 (relative to WH8102) require P for genome maintenance and N for protein synthesis. These costs can be estimated by examining genomic content and by using quantitative proteomic measurements. Overall, oceanic WH8102 has a ~10% smaller genome than either coastal strain (WH8020 and CC9311), consistent with the loss of Fe response genes like those encoding ferritin and flavodoxin (Fig. 3B and SI Appendix, Table S1). This smaller genome likely imparts a selective advantage during P-limited periods in the south Sargasso Sea (20, 21). However, the cost of maintaining a single Fe-adaptive gene within the Synechococcus WH8102 genome is only ~0.04% of the genome content (1 gene in 2,526 genes total), implying that maintaining each individual gene would incur only a small ecological cost.

The N cost associated with the multilayered Fe-adaptive response in coastal WH8020 was examined using quantitative proteomic methods, where isotopically labeled peptide internal standards were synthesized for ferritin, IdiA, ferredoxin, and flavodoxin, and quantified using multiple reaction monitoring on a quadrupole-orbitrap mass spectrometer (22). N requirements were calculated based on the absolute concentration and chemical formula of each protein (Materials and Methods). Ferritin and IdiA synthesis requires ~3.3–4.4 pmol N per microgram of total protein (Fig. 3C). Because the cells must either synthesize ferritin or IdiA depending on ambient Fe concentrations, these Fe-adaptive proteins incur an N cost for the cell regardless of whether Fe levels are high (for Fe storage) or low (for Fe uptake). Flavodoxin has the largest N cost (14.7 pmol N per microgram of total protein; Fig. 3C) of the Fe-adaptive proteins targeted here. The ~100-fold greater abundance of flavodoxin (98.9 fmol per microgram of total protein) compared with ferredoxin (0.73 fmol per microgram of total protein) in WH8020 is consistent with reports of flavodoxin’s lower reaction efficiency compared with ferredoxin (23). Whereas flavodoxin only contributes to ~0.2% of total protein N at its most abundant (0.0019 μg flavodoxin per microgram of total protein), this observation suggests that even the small N costs associated with adaptive responses to nutrient scarcity are ecologically significant, and this is consistent with previous findings of small N cost for adaptive responses to vitamin B12 scarcity in diatoms (24). The benefit that flavodoxin imparts in the dynamic coastal ocean likely outweighs the N cost of its synthesis. Moreover, the small N cost of synthesizing Fe-responsive proteins like flavodoxin is still higher than the P cost of maintaining the corresponding genes.

Our results suggest that the capacity of Synechococcus to adapt to Fe limitation therefore depends not on Fe alone, but on the balance between availability of Fe, N, and P. The combination of relatively high dissolved Fe levels (25) and scarcity of N and P in the permanently stratified southern Sargasso Sea may place stronger selective pressure on oceanic WH8102 to eliminate Fe-adaptive genes. A recent North Atlantic transect that sampled near both Synechococcus isolation environments illustrates this: oceanic surface waters had elevated Fe:nitrate ratios (11.4:1 μM:nM on average between 64–66°W, the longitude of isolation for WH8102),
Fig. 2. Physiological and photosynthetic responses of oceanic WH8102 (Left, A–E) and coastal WH8020 (Right, F–J) to [Fe]. Vertical dashed lines in A and F and bars in B and G indicate the concentration of Fe’ plotted as log[Fe’] at which a given physiological parameter showed a threshold response (Materials and Methods). Growth rate (circles) and photosynthetic efficiency (F/Fm measured at the time the protein samples were collected, shaded regions) are shown in A and F. ETC, membrane proteins in the photosynthetic electron transport chain. Growth curves are shown in C and H and photosynthetic efficiency (F/Fm) values collected each day are shown in D and I. Legend for C, D, H, and I is shown below J. (E and J) Heatmap of photosynthetic electron transport chain proteins for Synechococcus sp. strains WH8102 (E) and WH8020 (J). Normalized spectral counts are given in SI Appendix, Table S4. Color indicates higher (yellow) or lower (purple) abundance relative to the centered mean value (black).

whereas the Atlantic coast had relatively lower Fe-nitrate (5.3:1 pM:nM at 69°W close to the WH8020 isolation site; Fig. 1C). Additionally, oceanic (64–66°W) seawater nitrate:phosphate ratios were low (~2.1:1 nM:nM) relative to both the Redfield ratio (26) (16:1) and WH8102 cell quotas (10.9:1 and 43.8:1 in P-replete and P-limited cultures, respectively) (27). These nutrient availability patterns are consistent with a selection pressure against expending additional N within Fe-adaptive proteins in the oligotrophic southern Sargasso Sea. Specifically, low N levels in the open ocean may make flavodoxin less beneficial for oceanic WH8102, given that
less Fe adaptability would be needed under its relatively stable Fe regime (Fig. 1 B and C). Oceanic WH8102 is likely well suited to these high dust waters, whereas other oceanic strains with greater Fe acclimation capacity may exist in regions with lower Fe availability, such as the Equatorial Pacific (28) and Costa Rica Dome (29), or even seasonally in the northern Sargasso Sea near the Bermuda Atlantic Time Series (BATS; Fig. L4) where dust supply wanes seasonally (9) and Fe is depleted within the deep chlorophyll maximum (30).

In comparison with Synechococcus, the N cost of Fe adaptability is apparently not sufficient to deter the highly streamlined Prochlorococcus MED4 and MIT9301 genomes from retaining flavodoxin, yet ecotypes adapted to chronic Fe limitation in the Pacific Ocean have eliminated certain Fe-containing genes from their genomes (28). The selective loss of Fe-containing proteins (but not Fe-free Fe response proteins like flavodoxin) from Prochlorococcus genomes implies a particular importance of adapting to Fe scarcity in Prochlorococcus. Prochlorococcus has approximately twofold lower cellular N requirements than Synechococcus (27), and obtains N from urea (31, 32). As a result, Synechococcus may experience greater selective pressure to conserve N, given that smaller Prochlorococcus, with its larger surface area-to-volume ratio, would likely outcompete Synechococcus for Fe in either case.

These results demonstrate distinct responses of marine Synechococcus from coastal and open Atlantic Ocean environments. The surprising dynamic, multitiered Fe response of coastal Synechococcus WH8102 implies that periods of Fe scarcity likely extend well into the coastal zones of continental shelf regions, in addition to the open ocean and coastal upwelling systems. The limited response of oceanic Synechococcus WH8102 demonstrates the challenge of multinutrient scarcity in the open ocean, where the advantages of maintaining Fe adaptability are offset by even small nutritional costs. These responses suggest that the need to conserve Fe plays an important role in species evolution in the coastal and open ocean, but the outcome rests on a complex interplay between the availability of multiple scarce nutrients and an organism’s complement of biochemical responses.

Materials and Methods

Strains and Culturing Conditions. Synechococcus sp. strain WH8102 was isolated from surface waters in the oligotrophic southern Sargasso Sea (22.495°N, 65.6°W by F. Valois and J. Waterbury) north of Puerto Rico, where the waters are permanently stratified, and belongs to Synechococcus clade III (2). Strain WH8020 was isolated from surface waters near Woods Hole, MA (38.68°N, 69.32°W by

Fig. 3. (A) Abundance of Fe-related proteins in coastal Synechococcus sp. strain WH8020 in response to [Fe] (gene IDs correspond to genes in SI Appendix, Table S1). (B) Venn diagram of Fe response genes (SI Appendix, Table S1) in the genomes of oceanic strain WH8102 and coastal strains WH8020 and CC9311. (C) N costs associated with Fe response proteins in coastal Synechococcus WH8020 under high (>1 nM) and low (≤1 nM) Fe conditions (the expression threshold for these proteins). Flavodoxin (GenBank accession no. WB44_08620) was ~100-fold less abundant than flavodoxin (GenBank accession no. WB44_06685). Although ferredoxin is more abundant under high Fe (see A), the effect is masked in C due to the scale used for flavodoxin.
F. Valois and J. Waterbury), both strains were generously provided by the Waterbury laboratory. The WH8102 genome is available at www.ncbi.nlm.nih.gov/Taxonomy/Browser/wwwtax.cgi?id=845988. The WH8200 genome reported in this paper has been deposited in the GenBank BioProject database (accession no. PRJNA278997).

Synechococcus spp. WH8102 and WH8020 were cultured at 23 °C under a light intensity of 0.039, based on empirical determination of the photosynthetic efficiency in the dark-adapted state ($\mu_{\text{P}}$; Mackey et al., see below) was monitored daily. Cultures for protein extraction were harvested during late log phase by gentle vacuum filtration onto 0.2-μm Supor filters and frozen at −80 °C until processing.

Protein Extraction and Digestion. Frozen cell samples for protein extraction were rinsed from the filters in 100 mM ammonium bicarbonate, sonicated on ice a minimum of 10 min to lyse cells and break up cellular membranes, and centrifuged to pellet cell debris. Proteins in the supernatant were precipitated overnight in 100% (vol/vol) acetone at −20 °C. The pellet was resuspended in 6 M urea with 0.1 mM ammonium bicarbonate, reduced with 10 mM DTT in a thermomixer (56 °C, 450 rpm, 1 h), and alkylated with 30 mM iodoacetamide (20 °C on bench for 1 h). Proteins were then digested with trypsin (Promega Trypsin Gold; 37 °C, 400 rpm overnight in a thermomixer) at an enzyme:protein ratio of 1:50. The tryptic peptides were concentrated by evaporation and resuspended in 2% (vol/vol) acetonitrile with 0.2% formic acid.

Global Proteome Analysis. Chromatography and tandem mass spectrometry (LC-MS/MS) of the tryptic peptides was performed on a Q Exactive (QE) Orbitrap mass spectrometer (Thermo Scientific). Chromatography was performed using a 180-min nonlinear gradient of 0.1% formic acid in water and 0.1% formic acid in acetonitrile on a 2.1 mm × 150 mm column (New Objective, Acquity UPLC HSS T3) packed with 3 μm C18 silica packing material (Repsol-Gold 120 C18, 3 μm; Dr. Maisch GmbH). Tandem mass spectrometry was performed on the top 15 ions. Full MS scans were performed at 70k resolution with a scan range of 380-2,000 m/z and 1e5 AGC target. Data-dependent MS2 scans were monitored at 17.5k resolution with 1e5 AGC target, 100 ms maximum injection time, and a 2.0 m/z isolation window.

Proteins were identified against the genome for each strain using the SEQUEST algorithm within Proteome Discoverer (Thermo) and normalized spectral counts tabulated in Scaffold software (version 4.3.2 Proteome Software). Protein identification criteria included a protein identification probability of 99%, a peptide identification probability of 95%, and identification of two or more peptides from the protein’s sequence using the Peptide Prophet algorithm (37). Relative protein abundance data were normalized to total spectral counts. The resulting data matrices were then imported into Cluster 3.0 (38), log transformed, centered on mean, and normalized across treatments to create a heatmap (Fig. 2). Proteins were clustered using the correlation (uncentered) similarity metric and centroid linkage options.

The global proteome values show the spectral counts for each protein, which are uncalibrated measurements of relative abundance for which comparisons within a protein across treatments is the best application. In contrast, the targeted calibrated measurements for which isotopically labeled peptide standards are used (see below), are the appropriate measurement for comparisons of concentrations of different proteins.

Targeted Protein Quantitation. Peptide target sequences representing proteins of interest were identified from peptides in the global proteome. Two or more peptides per protein of interest were synthesized using SpikeTides Synthetic Proteotypic Peptides (UPT Peptide Technologies GmbH). These peptide standards were then mixed with proteins and heavy isotope-labeled heavy spike-in peptides, allowing precise quantitation of the peptide. Experimental samples were spiked with known amounts of each standard to determine the amount of the respective protein in the sample on a femtomole peptide/microgram total protein basis. Samples were analyzed using parallel reaction monitoring (PRM) (see SI Appendix, Table S4 for settings) (39, 40), where each precursor ion (light and heavy masses) was selected by the quadrupole, fragmented, and then all fragment ions were then introduced into Cluster 3.0 (38). The accurate mass spectrum of each fragment ion intensities was calculated in Skyline (MacCoss Lab Software version 3.1) and used to estimate peptide signal intensity. Peptide concentration was calculated based on the ratio to the heavy peptide standards that were added in known quantity.

To compare the amount of protein under Fe replete and deplete conditions, all PRM values above (replete) and below (deplete) the Fe threshold for that protein were averaged and compared for both peptides per targeted protein (SI Appendix, Table S2). For example, for flavodoxin the threshold for WH8102 occurred for cells grown between 1 and 10 nM, so all protein concentrations for cells grown at ≥10 nM were averaged to obtain the replete, or “high” protein concentration, and all values for cells grown at ≤1 nM were averaged to obtain the deplete, or “low” protein concentration. The replete and deplete values were subtracted to determine the mass of extra protein produced by the cell when these genes are expressed. The N cost associated with each protein was calculated based on the number of N atoms per protein for each target. Fe Response Gene Identification. The WH8102 and CC9311 genome annotations were searched using iron-related key words and for known iron-sparing proteins. The identified genes were used to BLAST the WH8020 genome to identify orthologous genes.

Photosystem II Fluorescence. Photosystem II fluorescence was measured using a F8 fluorometer with FView software (Santificial) and blue excitation light (450 nm with 30 nm bandwidth). Due to the small cross section of Synechococcus for blue light, it is necessary to increase the duration of the saturating flash beyond the default value to ensure that a steady state fluorescence plateau is reached; in this study, a single turnover flash duration of 200 μs was used. Default values were used for all other parameters. The reference excitation profile was recorded for these timing parameters using rose bengal dye. The fluorescence emission parameters $F_{\text{Fus}}$ and $F_{\text{Fv}}$ were measured using the orbitrap (22). The sum of the top five fragment ion intensities was calculated in Cluster 3.0 (38) to determine protein concentrations for which isotopically labeled peptide standards are used (see below), are the appropriate measurement for comparisons of concentrations of different proteins.

Growth Curves. Doubling times in Fig. 2 A and F were calculated from fluorescence growth curves shown in Fig. 2 C and H using the equation $t_d = \ln(2)/[\ln(\text{growth rate})]$, where $t_d$ is the time for the population to double, $t$ is final time, $t_i$ is initial time, and $c_i$ is the initial concentration. Fluorescence trends were confirmed against flow cytometry measurements of samples taken at the final time point of each experiment (SI Appendix, Fig. S2). Formalin fixed samples were flash frozen in liquid nitrogen and stored frozen at −80 °C until analysis on
an Accuri flow cytometer. Counts were triggered on chlorophyll fluorescence. These data are shown in SI Appendix, Fig. S2, and F0/F of 0.94. Threshold response calculated. For Fig. 2, protein thresholds were determined as the concentration at which the protein declined to 50% of its maximal value. Photophysiological thresholds were taken as the concentration above which the F/F0 response diverged from the “no iron added” condition. Nutrient and dissolved Fe concentrations. Sea samples for Fe concentration analysis were collected on US GEOTRACES section GA03 using trace-metal clean sampling protocols (41). Fe concentrations were analyzed on a Nepe- tune multiloupe ICP-MS at the University of South Carolina by isotope dilution, after concentrating and purifying Fe from a 1-L seawater samples