Abundances of Iron-Binding Photosynthetic and Nitrogen-Fixing Proteins of *Trichodesmium* Both in Culture and *In Situ* from the North Atlantic

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Abstract

Marine cyanobacteria of the genus *Trichodesmium* occur throughout the oligotrophic tropical and subtropical oceans, where they can dominate the diazotrophic community in regions with high inputs of the trace metal iron (Fe). Iron is necessary for the functionality of enzymes involved in the processes of both photosynthesis and nitrogen fixation. We combined laboratory and field-based quantifications of the absolute concentrations of key enzymes involved in both photosynthesis and nitrogen fixation to determine how *Trichodesmium* allocates resources to these processes. We determined that protein level responses of *Trichodesmium* to iron-starvation involve down-regulation of the nitrogen fixation apparatus. In contrast, the photosynthetic apparatus is largely maintained, although re-arrangements do occur, including accumulation of the iron-stress-induced chlorophyll-binding protein IsiA. Data from natural populations of *Trichodesmium* spp. collected in the North Atlantic demonstrated a protein profile similar to iron-starved *Trichodesmium* in culture, suggestive of acclimation towards a minimal iron requirement even within an oceanic region receiving a high iron-flux. Estimates of cellular metabolic iron requirements are consistent with the availability of this trace metal playing a major role in restricting the biomass and activity of *Trichodesmium* throughout much of the subtropical ocean.

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Introduction

Open-ocean diazotrophic cyanobacteria, such as *Trichodesmium* spp., are of particular importance due to their contributions to the carbon (C) cycle through primary production and to the nitrogen (N) cycle through the fixation of atmospheric N₂ [1–3]. *Trichodesmium* spp. are thought to comprise the most abundant diazotrophic cyanobacteria in the oceans [3], often forming widespread blooms over subtropical and tropical regions and potentially contributing a larger fraction of total marine nitrogen fixation than any other organism [3–6]. While marine phytoplankton are N-limited through much of the tropical and subtropical oceans [7–9], diazotrophic growth is likely limited by the availability of other nutrients such as phosphorous (P) and iron (Fe) [10–15].

Within the photosynthetic apparatus, the chlorophyll-binding membrane protein complexes photosystem II (PSII) and photosystem I (PSI) both have an absolute requirement for iron. Photosystem I (PSI) represents the largest photosynthetic requirement for iron, each PSI trimer containing 36 Fe atoms in 9 [4Fe-4S] clusters [16]. Compared with other autotrophs, photosynthetic diazotrophs, including *Trichodesmium*, have a significant further iron requirement due to the abundance of iron-containing enzymes in the nitrogen-fixation apparatus [17–19]. The nitrogenase complex (the key enzyme in N₂ fixation) is composed of two Fe-proteins encoded by the *nifH* gene, each containing 4 Fe atoms [20] and a dimeric MoFe protein encoded by *nifD* and *nifK* genes, which contains a total of 30 Fe atoms [20,21], making it one of the most iron-rich enzymes in nature [17,22]. As a consequence, the availability of iron in marine systems appears to greatly influence N₂ fixation in cyanobacteria [10,23–26].

Models predict that the distribution of nitrogen fixation in the modern ocean may be constrained by the availability of iron [27–29]. Moreover, oceanographic distributions are consistent with the availability of iron affecting N₂ fixation and, consequently, the biogeography of diazotrophic organisms including *Trichodesmium* [14,15]. The North Atlantic Ocean has some of the highest N₂ fixation rates in the global ocean [15], with waters characterized by high dissolved Fe concentrations tightly linked to high atmospheric dust inputs [14,30]. However, evidence for enhanced N fixation rates following Fe addition to North Atlantic waters suggest that enhanced Fe may influence *Trichodesmium* growth even in this ocean basin [12,31].

Diazotrophy is a significant challenge for oxygenic photosynthetic microorganisms because O₂ is inhibitory to the N₂ reduction enzyme nitrogenase [32,33]. Diazotrophs have developed specific molecular and physiological strategies to protect nitrogenase from the O₂ evolved during photosynthesis [28,34,35]. Some diazotrophs have adapted to fix nitrogen during the dark period to avoid photosynthetic oxygen inhibition of the nitrogen-
nase complex (temporal separation) [36–39], while others have terminally differentiated cells, termed heterocysts, with thickened cell walls and reduced photosynthetic activity (spatial separation) [40]. The non-heterocystous *Trichodesmium* uniquely undertakes both CO₂ and N₂ fixation during the day in the same cell through a complex combination of apparently reversible temporal and intracellular-spatial separation of these processes [4,34,35,41]. The simultaneous occurrence of two iron-rich metabolic processes in *Trichodesmium* may also suggest a higher specific iron requirement per cell than for other diazotrophic organisms [38].

While the impacts of iron on growth, C and N₂ fixation, O₂ production and dark respiration in *Trichodesmium* in culture have been well documented [24,25,42–44], relatively little is known of the molecular adaptation of this organism to iron availability in the environment. Although transcriptomic and protein level responses to iron stress have been observed in culture [19,45–47], information from *in situ* natural populations is limited [48–50]. Here we report the responses of photosynthetic and nitrogen fixing proteins to iron availability in laboratory cultures of *Trichodesmium* IMS101 and compare these with natural populations of *Trichodesmium* from the subtropical North Atlantic. We quantified peptides indicative of the abundance of the major iron-binding proteins involved in nitrogen fixation and photosynthesis, including the iron protein of nitrogenase (NifH) and the major chlorophyll-binding proteins in photosynthesis, the D₁ protein of PSII (PsbA) and a core subunit of PSI (PsaC). In addition we assessed the presence and accumulation of the iron-stress-induced chlorophyll-binding protein IsiA [51,52]. Through absolute quantification of these enzymes we characterized the molecular acclimation strategy of *Trichodesmium* to iron availability and hence estimate the iron supplies required to sustain natural populations.

**Materials and Methods**

**Sample Collection**

**Culture.** *Trichodesmium* IMS101 (originally isolated by [53]) was grown as batch-cultures in YBC-II medium [54], made from artificial seawater filtered through 1.0- and 0.2-µm Millipore membrane filters. Culture conditions included a 12/12-h light-dark cycle using cool white lamps at 50 µmol photons m⁻² s⁻¹ and temperature of 25°C. Small volume cultures (50 ml and 200 ml) were grown in polycarbonate flasks (Nalgene) with gentle agitation on an orbital shaker. Cells from a late-log-phase culture (as estimated by chlorophyll concentration measured daily) were used as an inoculum (1/50 dilution) for five parallel cultures grown in YBC-11 supplemented with 20, 40, 120 and 200 nM Fe (cultures grown in YBC-11 medium with no added iron did not grow). Our culture conditions (2 mM EDTA) would not have buffered iron concentrations. Consequently biological iron uptake resulted in the development of starvation within the cultures. Samples were harvested after a 7 to 9-day incubation period, when the 20 nM Fe culture first showed evidence of physiological iron-stress as indicated by reduced photosynthetic efficiencies. Measurements of Fv/Fm were made 2 h after the start of the light period. All nutrient stress experiments were performed as biological triplicates.

**Field.** Samples were collected during three cruises in the tropical and subtropical Atlantic (Figure 1). D326, a SOLAS (Surface Ocean Low Atmosphere Study) research cruise South of the Canary Islands, occurred during January-February 2008. FeAST-6, a cruise south of Bermuda, took place during summer (July) 2008, and AMT19 (Atlantic Meridional Transect), took place during Autumn (Oct.-Dec.) 2009. During these cruises, *Trichodesmium* spp. colonies were collected using a drift-plankton net (50 µm on AMT-19, 60 µm on D326 and FeAST-6) deployed at 5 m depth. Samples were collected at midday (AMT-19 and D326) and pre-dawn (FeAST-6). For protein analysis, colonies were picked individually, vacuum filtered with a hand pump onto glass fiber filters (0.3 µm nominal pore size, Advantec, Southampton, United Kingdom), snap-frozen in liquid nitrogen and stored at -80°C. The time from collection to freezing was always less than 30 min to avoid significant changes in the protein profile of the samples. Protein samples were collected from 1, 5 and 10 locations as part of the D326, FeAST-6 and AMT19 cruises, respectively. During D326, samples were also collected from an additional 7 stations to measure colony chlorophyll, particulate organic carbon and N₂ fixation rates.

**Analytical Methods: N₂ Fixation**

During D326, N₂ fixation was measured on samples of 20 picked *Trichodesmium* colonies following [35], acknowledging the potential for measured rates to be lower bounds due to incomplete equilibrium of the injected ¹⁵N₂ gas with the sample [39]. Samples were analyzed by isotope ratio mass spectrometry calibrated...
against known carbon and nitrogen standards to derive colony particulate carbon, nitrogen and N₂ fixation rates and hence, assuming steady state, estimate growth rates.

Chlorophyll-α Measurement

For culture studies, 2 ml of cell cultures were filtered onto Whatman GF/F filters and snap frozen in liquid nitrogen pending analysis. For in situ samples, 5–20 colonies were incubated onto Whatman GF/F filters and immediately extracted. Filters were incubated into 4–6 ml 90% acetone overnight at 4°C in the dark and fluorescence was then measured according to [56].

PSII Variable Chlorophyll Fluorescence

The photosynthetic physiology of natural Trichodesmium colonies was measured using active chlorophyll fluorescence on a Sadtantic Fluorescence Induction and Relaxation (FIRE) [57], to determine the apparent photochemical quantum efficiency (Fv/Fm). For culture studies, PSII fluorescence of Trichodesmium IMS101 was measured once a day, 2 h after the onset of photoperiod, using a Fasttracka™ Mk II Fast Repetition Rate fluorometer (FRRf) integrated with a FastAct™ laboratory system (Chelsea Technologies Group LTD, West Molesey, Surrey, United Kingdom). Fv/Fm was taken as an estimate of the apparent PSII photochemical quantum efficiency [58]. For cultures, all parameters were measured directly after sampling, without dark adaptation under increasing background irradiance levels from 5 to 30 μmol photons m⁻² s⁻¹, which initially caused an increase in quantum yield. Data presented thus correspond to a background irradiance of 21 μmol photons m⁻² s⁻¹ (PAR).

Total Protein Extraction and Quantification

Filters with 100–150 colonies (field samples) or 40–45 ml of culture (∼5000 trichomes) (culture samples) were resuspended in 1 ml denaturing extraction buffer containing 140 mM Tris base, 105 mM Tris-HCl, 0.5 mM EDTA, 2% lithium dodecyl sulfate, 10% glycerol, and 0.1 mg ml⁻¹ PefaBloc SC protease inhibitor (Sigma-Aldrich, Pool, United Kingdom). The proteins were extracted according to the protocol described by [48]. Samples were centrifuged for 10 min at 15,000 g, pooled and resuspended in a solution of 140 mM Tris base, 105 mM Tris-HCl, 0.5 mM EDTA, 2% lithium dodecyl sulfate, 1% sodium dodecyl sulfate, 10% glycerol, and 0.1 mg ml⁻¹ PefaBloc SC protease inhibitor (Sigma-Aldrich, Pool, United Kingdom) for 1 ml to 250 μl. Total protein concentrations were measured using a modified Lowry assay (Bio-Rad, Hemel Hempstead, United Kingdom) with bovine γ-globulin as a comparative protein standard.

Target Protein Quantification

Key protein quantification was performed using standards (AgriSera, Vannas, Sweden) and followed the procedure described by [48] and [59]. Primary antibodies (AgriSera, Vannas, Sweden) were used in 2% ECL advance blocking reagent in Tris-buffered saline plus Tween 20 for NifH (Fe protein of nitrogenase), PsbA (D1 protein of PSII), PsAC (core subunit of PSII) and CP43 (IsiA: iron-stress-induced protein A) for 1 h incubation. Blots were incubated for 1 h with horseradish peroxidase-conjugated chicken anti-rabbit secondary antibody (Abcam, Cambridge, United Kingdom) for PsbA, PsAC and CP43 primary antibodies and with horseradish peroxidase-conjugated rabbit anti-chicken secondary antibody (Abcam, Cambridge, United Kingdom) for NifH. Blots were developed using ECL Advance detection reagent (Amersham Biosciences, GE Healthcare, Little Chalfont, United Kingdom) and a CCD imager (VersaDoc™, Bio-Rad Laboratories Ltd, Hemel Hempstead, United Kingdom). For estimating the amounts of protein in experimental samples, protein levels on Millipore immunoblots were quantified using QuantityOne™ and Image Lab™ software and calculated from standard curves (for each blot, after [48]). Example protein detection images are shown in Figure 2.

Redox kinetics of P700 Measured on Natural Trichodesmium Populations

During the D326 cruise, redox kinetics of the P700 reaction center chlorophyll of PSI were also measured on natural populations of Trichodesmium collected during net tows. To obtain sufficient biomass for measurements of absorption changes at 830 nm relative to 870 nm, samples were further concentrated by gentle filtration onto 10 μm polycarbonate filters followed by careful resuspension in the filtrate. Absorption changes at 830 nm (∆A830) were measured alongside variable chlorophyll fluorescence in a Walz Dual-PAM 100™ (Heinz Walz GmbH, Germany). Comparison of Fv/Fm values measured on concentrates and hand-picked colonies confirmed that the concentration step had no observable adverse effect on photochemical processes. A range of dilutions of concentrates were prepared for parallel measurement of ∆A830 and chlorophyll concentration to enable absolute quantification of the P700:Chl ratio for these natural populations (Figure 3A).

Results

PSII Photophysiology in Trichodesmium IMS101 and Natural Populations

After a 7- to 9-day incubation period, Trichodesmium cultures grown at the lowest levels of added Fe (20 nM Fe) exhibited decreased Fv/Fm when compared with cultures grown at higher iron concentrations (Figure 4). Although we cannot quantify the relative iron availability within the cultures at the point of harvest, over the range of added Fe levels physiologies would have varied from iron-replete to iron-starved. As mentioned above, cultures with no added Fe did not grow. The maximal Fv/Fm value was observed at 120 nM Fe. Data for cultures grown at the highest added Fe concentration (200 nM) indicated slightly suppressed Fv/Fm compared to 120 nM.
Maximal daily values of $F_v/F_m$ measured on natural populations ranged from around 0.47 to 0.37 (data from cruise D326 and FeAST-6). As previously observed [34], $F_v/F_m$ measured in Trichodesmium spp. natural populations displayed a distinct diel cycle, with minimal values observed around 1 h after local noon (Figure 3B). In contrast, the quantum yield of PSI photochemistry displayed no clear diel cycle (Figure 3C,D). Consequently light response curves for PSII and PSI electron transfer rates (ETR) differed (Figure 3D), with relative PSI ETR measured at high saturating irradiance remaining constant throughout the photoperiod, while PSII electron transport reduced around midday (Figure 3D), coincident with likely maximal rates of $N_2$ fixation [34].
Abundance of Major Iron-binding Proteins in *Trichodesmium* IMS101 and Natural *Trichodesmium* spp. Populations

Absolute protein concentrations obtained from quantitative Western-blot analysis were used to determine the photosynthetic/nitrogen-fixation protein profiles of *Trichodesmium* IMS101 harvested after 7 to 9 days growth. The level of iron starvation had little effect on the abundance of PsbA (D1 protein of PSII; Figure 5A); PsbA amounts decreased from 0.041 ± 0.001 pmol (μg total protein)^-1^ for the iron-replete culture to 0.031 ± 0.005 pmol (μg total protein)^-1^ under the most iron-starved condition. PsaC, a subunit of PSI, was more affected by Fe availability, decreasing from 0.037 ± 0.01 pmol (μg total protein)^-1^ under iron-replete conditions to 0.02 ± 0.01 pmol (μg total protein)^-1^ under iron-starvation (Figure 5A). The relative abundance of the two photosystems, PSI:PSII, estimated from the ratio of PsaC:PsbA, thus decreased with decreasing Fe availability across 4 culture conditions (one-way ANOVA p<0.01, Table 1). The Fe protein subunit of nitrogenase (NifH) was also influenced by Fe availability (Figure 5B). The NifH
chlorophyll protein in natural communities, ranged from 0.02–0.3 pmol (µg total protein)$^{-1}$ (Figure 5C), similar to that measured on iron-starved cultures of Trichodesmium IMS101 (0.49 ± 0.1 pmol. µg total protein)$^{-1}$ and never approached the concentrations per unit protein observed in iron-replete cultures (Figure 5B).

Chlorophyll Budget in Culture and Natural Populations of Trichodesmium spp

Quantification of all the major chlorophyll-containing complexes enabled us to derive a cellular chlorophyll budget. The presence of phycobilisomes as the major light-harvesting complex in Trichodesmium, results in the majority of chlorophyll in cyanobacteria being bound to the reaction centers PSI and PSII or, under iron-stress, potentially to Isa [52]. The number of chlorophyll molecules in each of the complexes has been determined through high-resolution structural analysis of the photosynthetic reaction centers [16,60,61] [ChlPSII = 36; ChlPSI = 100; ChlIsa = 12]. These values were thus used to derive ratios of total cellular chlorophyll to each complex, and hence also estimate the relative contribution (expressed as a %) of the total cellular chlorophyll bound within each of the complex pools [52]. For example, assuming 1 PsbA per PSII, and 1 PsaC per PSI, the ratio of total cellular chlorophyll to the cellular concentration of PSI (Chl:PSI) can be estimated from:

$$\text{Chl : PSI} = \text{Chl}_{\text{PSII}} + \text{Chl}_{\text{PSI}} \times \text{PSI} : \text{PSI} + \text{Chl}_{\text{Isa}} \times \text{Isa} : \text{PSI} = 100 + 36\text{PSII} : \text{PSI} + 12\text{Isa} : \text{PSI}. $$

The % of total cellular chlorophyll bound to PSI can further be calculated as:

$$\% \text{Chl bound to PSI} = 100(1 + 0.36\text{PSII} : \text{PSI} + 0.12\text{Isa} : \text{PSI})^{-1} ,$$

with similar equations derivable for PSII and Isa:

$$\% \text{Chl bound to PSII} = 100(2.778\text{PSII} : \text{PSII} + 1 + 0.333\text{Isa} : \text{PSII})^{-1} ,$$

$$\% \text{Chl bound to Isa} = 100(8.3333\text{PSI} : \text{Isa} + 3\text{PSII} : \text{Isa} + 1)^{-1} .$$

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**Table 1.** Selected subunit ratios for Trichodesmium IMS101 grown under different concentrations of added Fe.

| Trichodesmium IMS101 [culture] | [200 nM] Fe | [120 nM] Fe | [40 nM] Fe | [20 nM] Fe |
|-------------------------------|------------|------------|------------|------------|
| PSI:PSII | 0.9 ± 0.5  | 1.1 ± 0.3  | 0.6 ± 0.2  | 0.6 ± 0.3  |
| Isa:PSI  | u.d.       | u.d.       | 2.6 ± 1.6  | 6.4 ± 3.3  |
| Isa:PSII | u.d.       | u.d.       | 1.33 ± 0.25| 3.9 ± 1.1  |

u.d. indicates that Isa was undetectable at the two lowest concentrations of iron.

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**Table 2.** Selected subunit absolute quantification and ratios for Trichodesmium spp.

| Cruise | Trichodesmium spp. [natural populations] | AMT-19 | FeAST-6 | D326 | all |
|--------|-----------------------------------------|--------|---------|------|-----|
| PSI (µmol of protein)$^{-1}$ | 0.005 ± 0.005 | 0.004 ± 0.005 | 0.049 ± 0.02 | 0.019 ± 0.011 |
| PSII (µmol of protein)$^{-1}$ | 0.007 ± 0.005 | 0.008 ± 0.004 | 0.004 ± 0.003 | 0.007 ± 0.001 |
| IsiA (µmol of protein)$^{-1}$ | 0.034 ± 0.02 | 0.032 ± 0.04 | 0.12 ± 0.1 | 0.06 ± 0.05 |
| NifH (µmol of protein)$^{-1}$ | N/A | 0.132 ± 0.1 | 0.054 ± 0.002 | 0.09 ± 0.1 |
| PSI/PSII | 1.0 (0.3 – 1.7) | 1.4 (0.6 – 2.2) | 5.9 (4.6 – 7.3) | 2.0 (0.3 – 7.3) |
| IsiA/PSII | 4.1 (2.3 – 8.0) | 7.5 (1.7 – 19.4) | 1.97 (1.6 – 2.4) | 5.0 (1.7 – 19.4) |
| IsiA/PSII | 3.9 (2.2 – 6.7) | 7.7 (0.7 – 23.6) | 16.2 (11.0 – 22.5) | 8.5 (0.7 – 23.6) |

Colonies collected from 1, 5 and 10 locations as part of the D326, FeAST-6 and AMT19 cruises, respectively. For proteins absolute values, SD correspond to n = 16 samples. For ratios, average means are presented along with the corresponding range into brackets.

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Thus, for example, estimated ratios of Chl:PSI ranged from 150 mol:mol for iron-replete cultures to 800 mol:mol for iron-starved cultures where chlorophyll bound to IsiA significantly contributed to the total cellular budget, while the calculated value for Chl:PSI was 180 mol:mol for average field populations. As an independent check, during the D326 cruise spectrophotometric measurements averaged 93 ± 30 Chl:P700 (Figure 3A and 3C), compared to 130 Chl:PSaC as derived from protein ratios.

From the above equations, chlorophyll budgets for *Trichodesmium* in culture and in the field were thus compared (Figure 6A). Chlorophyll partitioning between the main chlorophyll-binding complexes in *Trichodesmium* IMS101 (e.g. PSI, PSII and IsiA) was significantly modified by iron starvation, partly due to a reduced PSI:PSII ratio, but principally due to the shift from two (PSI and PSII) to three (PSI, PSII and IsiA) main chlorophyll complexes (Figure 6A). The percentage of chlorophyll associated with PSII complexes remained relatively unchanged (25-33%); consequently, we effectively observed a re-allocation of total cellular chlorophyll from PSI (decreasing from 75% to 34%) to the IsiA complex (increasing from 0% up to 38%). Interestingly, the average chlorophyll % associated with each of the complexes in natural populations of *Trichodesmium* collected from the North Atlantic approximated that within iron-starved cultures, with 17%, 51% and 32% chlorophyll allocated to PSII, PSI and IsiA, respectively.

Partitioning of Fe Atoms between the Major Iron-binding Proteins in Culture and Natural Colonies of *Trichodesmium* spp

In a similar manner to the derived cellular chlorophyll budgets, the proportions of Fe in different cellular metabolic pools were calculated based on estimates of Fe atoms associated with each complex from structural studies. PSI and PSII have 3 and 12 Fe atoms per complex respectively [48,19]. Further, to more fully account for additional Fe within the photosynthetic electron transport chain, we assume a ratio of 1:1 for the Cytochrome b₅f:PSI ratio, with 6 Fe atoms per Cytochrome b₅f (Cytb₅f) complex [18]. Consequently we assume a total of 18 Fe atoms per Cytb₅f+PSI. The total Fe content of the nitrogenase complex depends on the ratio of Fe to MoFe protein (coded for by *nifH* and *nifDK* respectively) which has not to our knowledge been measured in *Trichodesmium*. Maximal specific activity for nitrogenase appears to occur at a ratio near 5 Fe:MoFe [62,63], with structural analysis of the protein [20] and *nifH/nifDK* transcript products [64] indicating a range of ratios from 2-5. This corresponds to a possible Fe content of 38–50 mol Fe (mol nitrogenase complex⁻¹), i.e. a factor of 2 variability in Fe content. As we measure the NifH complex, we thus estimate a potential range of total Fe in nitrogenase: NifH of 10–19 mol Fe (mol NifH⁻¹). We consequently perform calculations of Fe allocation across the different metabolic complexes for the upper and lower end of the ranges indicated above. For example, taking $\frac{PsbA_{Fe}}{PSI_{Fe}} = 3$; $\frac{Cytb_{6f} + PSI_{Fe}}{PSI} = 18$; $\frac{NifH_{Fe}}{PSI} = 19$, we can derive the following:

\[
\% \text{ Fe associated with PSII} = 100(6PSI : PSII + 1 + 6.333NifH : PSII)^{-1}
\]

\[
\% \text{ Fe associated with PSI} = 100(1 + 0.166PSI : PSI + 1.06NifH : PSI)^{-1}
\]

\[
\% \text{ Fe associated with NifH} = 100(0.95PSI : NifH + 0.1579PSII : NifH + 1)^{-1}
\]

Clearly there are likely to be other unaccounted for cellular Fe requirements, such as Fe superoxide-dismutase [18]. Our
estimates thus represent lower bounds for the total cellular Fe budget for *Trichodesmium*. In a manner similar to [28], we thus consider such estimates to represent the Fe metalloenzyme or metabolic inventory associated with the cellular processes of N₂ fixation and photosynthesis, although, as argued below, this potentially accounts for the majority of cellular Fe.

The percentage of Fe in the major iron-binding complexes was compared between culture and natural populations of *Trichodesmium* spp. (Figure 6B). Based upon the relative abundance of PsAC, PsbA and NifH, the major metabolic iron inventory (83–95% of Fe) under all culture conditions was estimated to be associated with the nitrogenase enzyme complex (Figure 6B). An inverse relationship was also observed between iron stress and iron allocation to PSI. The same analysis performed on average values derived from natural populations of *Trichodesmium* spp. indicated a similar partitioning of Fe between the resolved metabolic activities, with 9–15% Fe bound to the PSI complex versus 83–90% for the nitrogenase complex.

**Comparison of Theoretical and Prior in vitro/in vivo Data**

Combining our derived Fe and Chl budgets, we further calculate the cellular metabolic Fe:Chl ratio. The combined influence of IsA accumulation and a reduction in nitrogenase (Figure 3) resulted in estimated cellular metabolic Fe:Chl reducing from 1.7–3 mol mol⁻¹ under iron-replete conditions to 0.2–0.34 mol mol⁻¹ under iron-starved conditions, with all quoted ranges again indicating the sensitivity of calculations to the assumed Fe:MoFe protein (nifH/nifDK transcript product) ratio. For comparison, the average value estimated from our measurements of natural populations was 0.9–0.5 mol mol⁻¹. Although significant variability in cellular Chl:C needs to be acknowledged, assuming Chl:C ratios of 6–60 ng Chl colony⁻¹ (16 ± 14, mean ± 1 s.d., n = 33 samples, 7 stations, Table 3). Particulate organic carbon contents measured at a more limited number of stations ranged from 3–6 μg C colony⁻¹ (4.8 ± 1.4, n = 4), which, combined with an average of 37 ± 11 ng Chl colony⁻¹ at these same locations, resulted in an estimated Chl:C ratio of 107 ± 20 μmol:μmol. These values are consistent with previous observations of Chl:C ratios for natural *Trichodesmium* colonies [67] and comparable, although slightly lower than, values for diazotrophically growing cultures [44], the lower values in situ potentially reflecting a contribution from other microbes within the colony matrix [68–70].

Combining our estimates for metabolic Fe:Chl ratios of 0.3–0.5 mol mol⁻¹ with measured colony Chl contents, we estimate a conservative range for the metabolic Fe content of *Trichodesmium* of 2–35 pmol Fe colony⁻¹ (average 7 pmol Fe colony⁻¹) [11]. Measured Chl:C ratios further suggest 15–75 μmol metabolic Fe (mol C)⁻¹ for natural *Trichodesmium* populations, which is again comparable with previous direct elemental ratios measured on both natural populations and iron limited cultures [44]. Overall estimates of the metabolic iron demand were thus consistent with the majority of the cellular pool being associated with the metalloenzymes of the photosynthetic and nitrogen fixation apparatus (Table 3).

Total *Trichodesmium* chlorophyll standing stocks ranged from <0.0001 μg Chl m⁻² to maximal values of 0.07 μg Chl m⁻² in the subtropical North Atlantic during February-March 2008, with direct estimates of chlorophyll in the >10 μm fraction measured on a 10 L sample being of a similar magnitude. Bulk community in situ chlorophyll concentrations (i.e. >0.2 μm measured on 200 ml of a whole water sample) thus indicated that *Trichodesmium* accounted for up to 20% of the total chlorophyll standing stock at some stations. Over the observed ~50 m mixed layers, maximal *Trichodesmium* concentrations of 0.07 μg Chl m⁻² during D326 thus represented standing stocks of 3.5 mg *Trichodesmium* Chl m⁻². When combined with our estimates of cellular metabolic Fe requirements, this was thus the equivalent of a metabolic Fe standing stock of 1–2 μmol m⁻² (Table 4).

**Discussion**

In cultures of *Trichodesmium* IMS101, the onset of iron starvation was indicated by a decline in photochemical efficiency (Fv/Fm), as previously shown [25,41]. Although comparison of absolute values
of $F_r/F_o$ will be complicated by a range of other growth factors, in situ values measured on natural populations of *Trichodesmium* were consistent with those previously reported in the literature [34] and most comparable to iron-starved cultures.

The photosynthetic/nitrogen-fixation protein profiles of *Trichodesmium* either in culture or in the field show a high ratio of NifH to PsbA and PsAC proteins (Figure 5B and 5C), potentially reflecting a lower enzymatic rate for nitrogenase compared to photosynthetic proteins [48]. Our results reveal a small decrease in abundance of photosystem I and II complexes, but a much more significant decline in nitrogenase in iron-starved cultures. These results corroborate the trends observed at a gene-expression level by [19] who documented an early decline in nifH transcripts during the development of iron stress followed by a later decline in transcripts encoding the photosynthetic apparatus. Consequently it appears that the metabolic process of N$_2$ fixation may be more sensitive to iron limitation than photosynthesis [19]. In addition, PSII appears to be less sensitive to iron limitation than PSI at both gene and protein levels [19,48]. Our results indicate that the nitrogenase enzyme is the major metabolic sink for iron in *Trichodesmium* cells. Transfer of iron from the energetically and iron-expensive processes of nitrogen fixation to maintain energy production via the photosynthetic apparatus [19,47] could thus be speculated to act as a mechanism for surviving in oligotrophic subtropical and tropical areas characterized by fluctuations of iron supply [71].

Nitrogen fixation has been studied extensively in the tropical and subtropical North Atlantic [4,5,14,67]. Due to the proximity of African deserts, most significantly the Sahara, the area is subject to high rates of dust deposition [71–75], which likely result in the observed high surface iron concentrations [14]. Indeed, the low latitude North Atlantic has the highest known Fe concentrations of any of the global subtropical basins [13]. Despite this relatively high iron availability, we still found potential protein level evidence of reduced iron requirements within natural *Trichodesmium* populations based on comparisons with the cultured strain IMS101.

Significant caveats clearly need to be acknowledged when comparing culture results from a starvation experiment, run on a specific strain, under a limited set of other culture conditions, to field populations experiencing variable additional environmental forcings. Thus, although N$_2$-fixing enzyme abundances were in similar range in both *Trichodesmium* natural populations and in iron-starved cultures, phosphate is also severely depleted in the subtropical North Atlantic gyre, likely as a result of enhanced N$_2$ fixation due to dust deposition [14,76]. Consequently, natural *Trichodesmium* colonies from the region frequently display evidence of phosphate stress [13,50,77–78], which may thus contribute to any reduced capacity for N$_2$ fixation [11].

The IsiA protein was also present in all the natural populations sampled (Figure 5C) and is expressed [19] or accumulates (Figure 5A) under development of iron stress in *Trichodesmium* cultures. Although IsiA may be expressed under other growth conditions, including high light [79], acclimation to iron stress appears to be the primary functional role for this chlorophyll binding complex [47,80–83], which can act as a light-harvesting antenna for PSI [52,82].

In the present study, the IsiA protein constitutes a significant portion of the chlorophyll-binding protein in iron-starved cultures of *Trichodesmium* ISM101 and in natural populations (Figure 6A), with up to 4 and 6 times more IsiA than PSI and PSII proteins in starved cultures and natural populations, respectively.

The observed maintenance of PSI electron transport throughout the photoperiod in natural communities, in contrast to the down-regulation of electron transport through oxygen-evolving PSI (Figure 3D), supports the suggestion that PSI electron transport may act to consume cellular O$_2$ in order to protect nitrogenase from inactivation in *Trichodesmium* [33]. A subsequent requirement for maintenance of cellular PSI concentrations may further enhance the iron requirements of this organism compared with other diazotrophs [41]. For example, *Crocosphaera watsonii* separates N$_2$ fixation and oxygenic photosynthesis over the diel period [34] and hence can effectively share cellular Fe between these molecular processes [28]. Maintenance of PSI electron transport may also provide a rationale for increasing the light-harvesting cross-section of PSI in *Trichodesmium* under conditions of reduced iron availability through the expression of IsiA and the synthesis of IsiA-PSI supercomplexes [52,84].

Similar to previous calculations [25,66], estimation of iron within the molecular mechanisms of N$_2$ fixation and photosynthesis provides a means of extrapolating to the natural environment. Our estimate of 80–90% of the cellular Fe pool being associated with nitrogenase is consistent with theoretical calculations suggesting that 35–78% of the cellular Fe pool would be associated with N$_2$ fixation under optimal catalytic conditions [18]. Kustka et al. [18] estimated that cellular Fe:C ratios of 28–
40 μmol·mol⁻¹ would be required to maintain a moderately iron-limited growth rate of around 0.1 d⁻¹, consistent with both laboratory studies [41] and our observations of natural populations (Table 4). Combining observed nitrogen specific N₂ fixation rates with metabolic iron standing stocks, natural populations of Trichodesmium in the region sampled during D3Z6 would hence require 50–100 nmol Fe m⁻² d⁻¹ (Table 4). Cautious comparison with culture data (Table 3) further indicates that iron requirements could potentially be an order of magnitude higher under fully iron replete conditions. Observed ten-fold higher standing stocks of Trichodesmium in other regions of the North Atlantic [67], would also require correspondingly higher Fe turnover.

Although iron is known to be readily recycled in the upper ocean, knowledge of the differential cycling of nutrients (i.e. N or Fe) in oligotrophic systems is incomplete [85]. Given that the N₂ fixed by diazotrophs represents a source of ‘new’ nitrogen [sensu [86]] to first order it is reasonable to postulate that new iron inputs [85] may be required to balance much of the daily diazotrophic requirement. Given our estimated metabolic Fe demands, atmospheric dust related inputs of dissolved iron to the subtropical North Atlantic would thus be sufficient to satisfy the requirements of the large standing stocks of Trichodesmium observed in this region (Table 4). However, inputs would potentially be insufficient to satisfy the requirements of a metaloenzyme composition equivalent to that of an iron-replete culture.

Simple ecological theory predicts that the rate of supply of a limiting resource will control the organism standing stock, while the ecophysiological characteristics of these organisms will determine the ambient concentration of the resource [29,87,88]. We thus suggest that the response of Trichodesmium to atmospheric dust inputs to the North Atlantic [14] may lead to population growth and subsequent iron uptake reducing bioavailable levels to the point necessitating some reduction of cellular requirements. Such biological control of iron availability by the diazotrophic population would, however, be unlikely to result in severe stress and heavily reduced growth, a scenario which is consistent with the lack of expression of biomarkers of high iron stress in the region [46,50]. Other factors will clearly complicate this simple scenario, including non-biological influences on surface iron bioavailability [84] and, particularly in the North Atlantic, depletion of phosphorous [11,12,50,76]. However, consistent with previous experimental work [12,50], it appears that a limited degree of diazotrophic iron stress may potentially develop even in some regions of high input.

In contrast to the North Atlantic, other subtropical oceanic regions receive much lower iron inputs, which may thus represent a severe constraint on the accumulation of high standing stocks of Trichodesmium [15,25,71]. For example, Trichodesmium biomass [14] and hence estimated iron requirements are 3 orders of magnitude lower in the subtropical South Atlantic (Table 4). Our molecular characterization of the metabolic pools within natural populations of Trichodesmium hence provides further evidence of an important role for iron in dictating the large scale biogeography of this important diazotrophic taxon [15,25,27,29].

Conclusion

Relative to other diazotrophs [28,41], enhanced iron demand resulting from coordination of photosynthesis and nitrogen fixation may contribute to Trichodesmium being particularly sensitive to iron availability. In turn, high iron requirements have likely led to the evolution of both mechanisms for acclimating to reduced availability [19] and novel acquisition strategies [89]. Using a combination of laboratory and field experiments we suggest that natural Trichodesmium populations from the subtropical North Atlantic display molecular characteristics representative of a reduction in metabolic Fe-metaloenzyme requirements relative to iron-replete cultures. It thus appears that iron may influence Trichodesmium ecophysiology even in some regions receiving relatively high dust inputs. The current study adds further molecular level understanding to a growing body of evidence supporting the role of iron as a control on oceanic N₂ fixation.

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Author Contributions

Conceived and designed the experiments: SR TSB CMM. Performed the experiments: SR NJP DJH. Analyzed the data: SR NJP CMM TSB. Contributed reagents/materials/analysis tools: SR AIM DJH CMM TSB. Wrote the paper: SR CMM TSB.

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