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Periodic and coordinated gene expression between a diazotroph and its diatom host

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Abstract

In the surface ocean, light fuels photosynthetic carbon fixation of phytoplankton, playing a critical role in ecosystem processes including carbon export to the deep sea. In oligotrophic oceans, diatom–diazotroph associations (DDAs) play a keystone role in ecosystem function because diazotrophs can provide otherwise scarce biologically available nitrogen to the diatom host, fueling growth and subsequent carbon sequestration. Despite their importance, relatively little is known about the nature of these associations *in situ*. Here we used metatranscriptomic sequencing of surface samples from the North Pacific Subtropical Gyre (NPSG) to reconstruct patterns of gene expression for the diazotrophic symbiont *Richelia* and we examined how these patterns were integrated with those of the diatom host over day–night transitions. *Richelia* exhibited significant diel signals for genes related to photosynthesis, N₂ fixation, and resource acquisition, among other processes. N₂ fixation genes were significantly co-expressed with host nitrogen uptake and metabolism, as well as potential genes involved in carbon transport, which may underpin the exchange of nitrogen and carbon within this association. Patterns of expression suggested cell division was integrated between the host and symbiont across the diel cycle. Collectively these data suggest that symbiont–host physiological ecology is strongly interconnected in the NPSG.

Introduction

Oligotrophic waters comprise an estimated 30% of the global surface ocean [1]. However, these seeming deserts of primary production are estimated to contribute up to 90% of photosynthetic carbon fixation in the marine environment, of which <15% is exported out of the euphotic zone [2]. In the North Pacific Subtropical Gyre (NPSG), much of this

carbon export has been linked to annual summer increases in biomass of diatom–diazotroph associations (DDAs) [2, 3]. Composition of NPSG DDA blooms varies by year, but blooms are often dominated by *Rhizosolenia* sp. with the symbiotic cyanobacterium *Richelia intracellularis* [3–5]. Fixed N₂ from *Richelia* is transferred to the host and is estimated to provide much of the diatom nitrogen (N) quota in similar associations such as with *Hemiaulus* [6]. Estimates of daytime N₂ fixation rates by *Richelia* associated with *Rhizosolenia* are near 1 pmol heterocyst⁻¹ h⁻¹ [7] and have been shown to provide substantial inputs of new N into the systems where they occur [8, 9].

In oligotrophic waters, access to fixed N₂ provided by a symbiotic partner likely provides distinct advantages to the diatom host. Currently, only a few genera of diatoms are known to form symbiotic relationships with diazotrophs. Further, *Rhizosolenia* is known to exist without *Richelia* [10] suggesting this partnership is facultative. Although the transfer of N from symbionts to diatom hosts has been demonstrated [6, 11], there are no long-term cultures of this symbiosis to study and our current understanding of this association *in situ* is limited. Fundamental questions remain about how the symbiosis is maintained, how resources

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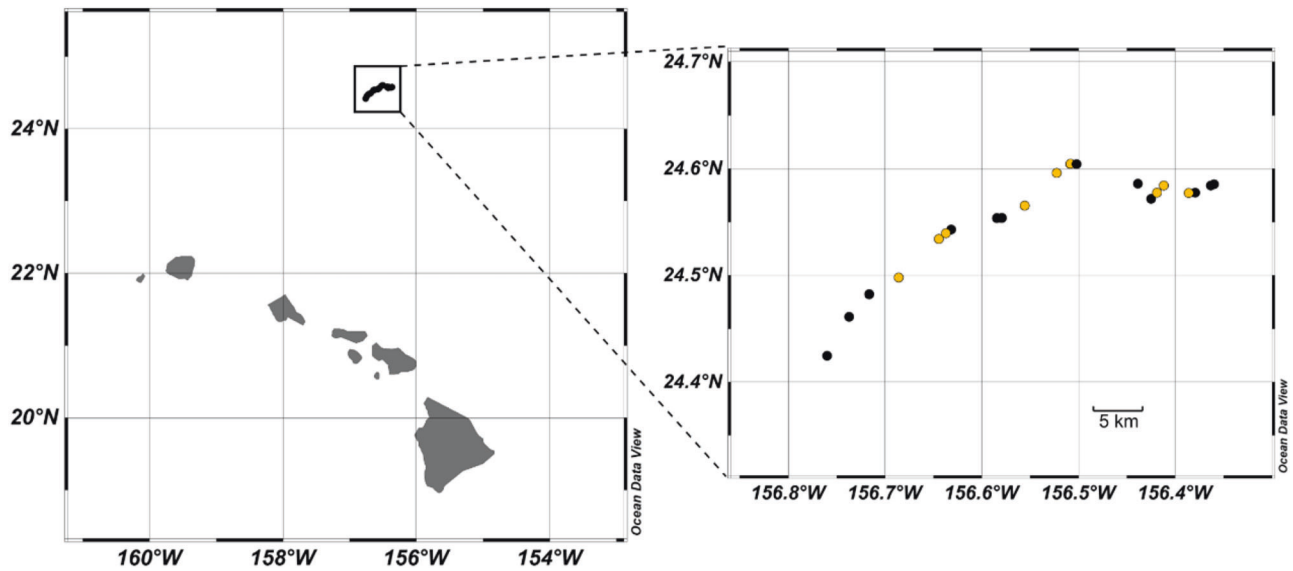


Fig. 1 Map of Lagrangian sampling stations on a July 2015 cruise in the North Pacific Subtropical Gyre. Left panel indicates location relative to the Hawaiian Islands and the right panel displays an

expanded view, with stations (21 in total spanning 4.5 days) sampled during night hours in black and stations sampled during daylight hours in yellow

beyond N might be modulated within the association, and how cell division might be synchronized between host and symbiont in order to propagate this facultative association. To better understand this keystone symbiosis, we reconstructed patterns of gene expression for the diazotroph *Richelia* and examined how these patterns were co-expressed with those of the diatom host over day–night transitions in the NPSG.

Methods

Sample collection

Seawater for diel *in situ* metatranscriptomes was collected aboard the R/V *Kilo Moana* as part of the 2015 Simons Collaboration on Ocean Processes and Ecology (SCOPE) research collaboration HOE-Legacy II Research Cruise near Station ALOHA (Fig. 1; <http://scope.soest.hawaii.edu/data/hoelegacy/>). During this cruise, samples were collected from 15 m depth within an anticyclonic eddy tracking a Lagrangian drifter. Detailed information regarding the sampling regime on the cruise has been described previously [12]. Hydrocasts for sampling were performed using a 24-bottle conductivity–temperature–depth (CTD) rosette sampler at a depth of 15 m every 4 h from 10 p.m. (local time) July 26, 2015 to 6 a.m. (local time) on July 30, 2015 for a total of 21 time points (Fig. 1). Water was collected in acid-washed 20 L carboys and seawater was pre-screened through 200 μm nylon mesh and then filtered onto two 5 μm polycarbonate filters (47 mm) by way of a peristaltic pump, passing $\sim 10\text{L}$ through each filter. This size

fraction was targeted to sample large eukaryotic phytoplankton and associated particle-attached and symbiotic prokaryotes. Samples collected at night were processed in the laboratory under red light. Filters were placed into cryovials and stored in liquid nitrogen until extraction. Total length of time from collection to placement into liquid nitrogen did not exceed 40 min.

RNA extraction and sequencing

Total RNA was extracted from individual filters using a Qiagen RNeasy Mini Kit (Qiagen, Hilden, Germany) modifying the lysis step with the addition of $\sim 500\ \mu\text{L}$ of 0.5 mm zirconia/silica beads (Biospec, Bartlesville, OK, USA). Briefly, lysis buffer and beads were added to each filter set ($n = 2$) for each time point and vortexed for 1 min, placed on ice for 30 s, and vortexed again for 1 min. Lysate from each filter set ($n = 2$) was removed with a pipette and pooled into a single 5 mL microcentrifuge tube. The rest of the Qiagen RNeasy Mini Kit protocol was then followed according to the manufacturer's instructions, adjusting volumes accordingly and incorporating the on-column DNase digestion step, using a Qiagen RNase-free DNase kit. Resulting total RNA was eluted with RNase-free water and then purified and concentrated with a Qiagen RNeasy MinElute kit according to the manufacturer's instructions. Quantity and quality of extracted total RNA was assessed on an Agilent 2100 Bioanalyzer (Agilent, Santa Clara, CA, USA).

To identify both eukaryotic and prokaryotic gene expression signals, extracted samples were split into two equal volumes of total RNA where one volume was

sequenced after a polyA pull-down step (Illumina Truseq library preparation kit; Illumina, San Diego, CA, USA) to enrich for eukaryotic mRNA (herein referred to as “selected reads”) and one volume was sequenced directly (herein referred to as “unselected reads”). Illumina Truseq libraries were sequenced with an Illumina HiSeq 2000 at the JP Sulzberger Genome Center (CUGC) of Columbia University following Center protocols. Samples targeting eukaryotic signals (selected) were sequenced to produce 90 million 100 bp, paired-end reads while samples targeting prokaryotic signals (unselected) were sequenced to produce 60 million 100 bp, paired-end reads. These environmental sequence data are deposited in the Sequence Read Archive through the National Center for Biotechnology Information under accession no. SRP136571. Raw sequence quality was visualized using FastQC [13] and then cleaned and trimmed using Trimmomatic [14] version 0.27 (paired-end mode, 4 bp-wide sliding window for quality below 15, minimum length of 25 bp).

Mapping and periodic expression analysis

Reads were mapped to a reference database after Alexander et al. [15]. The transcriptome *Rhizosolenia setigera* CCMP 1694 (MMETSP0789; <https://www.imicrobe.us/#/projects/104>) was used as a reference to elucidate transcriptional patterns of selected reads for *Rhizosolenia* sp. and combined with transcriptomes of *Chaetoceros* sp. (MMETSP1336, 200, 149, 751, 752, 753, 754, 1429, 1447, 90, 91, 92, 717, and 718), a species commonly associated with diazotroph endosymbionts [16], to create a dataset large enough to minimize false positive read alignments. For *Richelia* sp. analyses, unselected reads were mapped to the genome of *Richelia intracellularis* RC01 (<https://www.patricbrc.org/view/Genome/1164990.3>), which was isolated from *Rhizosolenia clevei* via flow cytometry sorting and included both vegetative cells as well as heterocysts (http://www.ebi.ac.uk/ena/data/view/GCA_000613065; [17]). As with *Rhizosolenia*, a reference database of common diazotrophs was created from genomes of *Richelia intracellularis* RC01, *Trichodesmium erythraeum* IMS101, *Calothrix rhizosoleniae* SC01, *Crocospaera watsonii* WH8501, UCYN-A1, and UCYN-A2. Mapping of both selected and unselected reads was conducted with the Burrows–Wheeler Aligner (BWA-MEM, parameters `-k 10 -aM`; [18]) and counted using the HTSeq 0.6.1 package (options `-a 0, --m intersection-strict, -s no`; [19]). Sequencing results can be found in Table S1. Alignment of trimmed reads to diatom and diazotroph reference databases resulted in a total of 4,229 (55% of total, Tables S2, S3) *Richelia* and 21,705 (92% of total, Table S2, S3) *Rhizosolenia* features (gene or contig) that were detected at sufficient coverage (≥ 3 read counts for both species) for subsequent analyses.

Annotations for *Rhizosolenia* were taken from the MMETSP assignments using the most abundant annotation hit for each contig (Pfam) while *Richelia* annotations were obtained from Patric assignments (<https://www.patricbrc.org/view/Genome/1164990.3>). Kyoto Encyclopedia of Genes and Genomes (KEGG) biochemical pathways for each genetic feature (contig or gene) were identified with KEGG Automatic Annotation Server (KAAS) using the partial genome single-directional best-hit method [20, 21]. Features (gene or contig) receiving KEGG assignments were binned by KEGG module. Analysis of periodic expression of count data was performed using the R package RAIN [22] by first filtering low abundance reads (see above) after Wilson et al. [12], and then normalizing with the DESeq2 “varianceStabilizingTransformation” command [23]. Resulting p -values from this analysis were corrected for multiple testing using the false-discovery rate method [24], with corrected p -values ≤ 0.05 considered to have significant diel periodicity. Peak times were calculated with a harmonic regression, fitting the expression data to a sine curve, as has been used in several recent studies [25, 26]. These estimates of peak expression time were congruent with the normalized expression patterns across the time series and were used as a means of estimating at what time the highest relative expression was occurring.

Network analysis

To examine co-expression patterns between *Rhizosolenia* and *Richelia*, a weighted gene co-expression network analysis was conducted using the R package WGCNA [27] on normalized counts (DESeq2 “varianceStabilizingTransformation” command; [23]) using methods described previously [25]. WGCNA uses a correlation network to identify clusters (modules) of highly correlated genes, which can have a co-expression pattern that is either synchronous or orthogonal. This type of analysis has been used in previous studies comparing host–symbiont dynamics [28–30]. Briefly, a soft-threshold of 7 was chosen based upon a scale-free network topology test (Figure S1) and transcriptional modules were identified using the “blockwiseModules” command (`minModuleSize = 30, mergeCutHeight = 0.25`). The TOMsimilarityFromExpr command was used to calculate topological overlap between transcripts. To evaluate variability in core metabolic processes for both *Rhizosolenia* and *Richelia* in each WGCNA module, normalized read counts for features (gene or contig) in each species with KEGG assignments were summed by KEGG module for WGCNA modules containing >200 genes or contigs (nodes). The relative abundance by species was then determined to view species-specific distribution of KEGG modules within each WGCNA module.

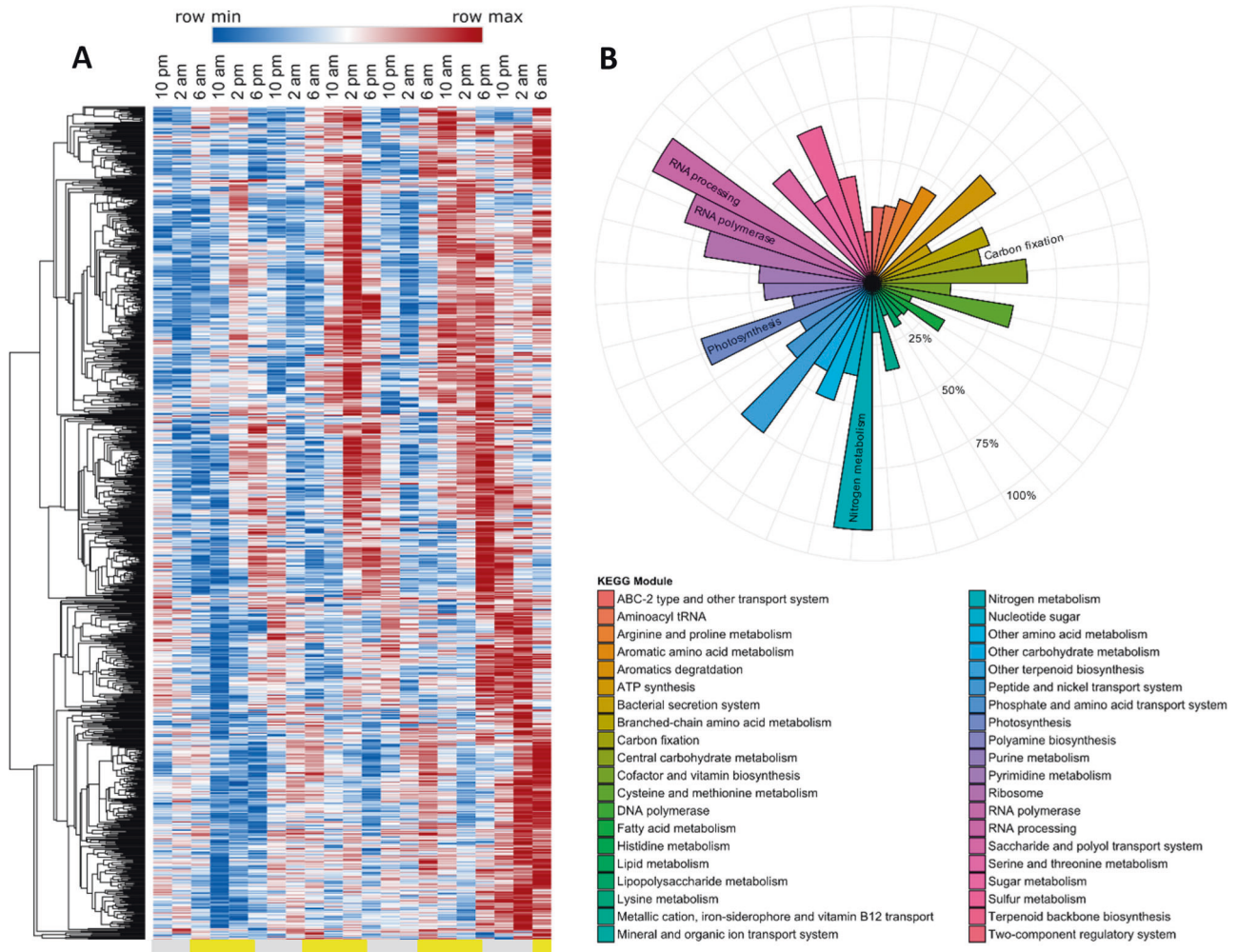


Fig. 2 Significantly periodic gene expression and metabolic profiles for *Richelia*. **a** Hierarchically clustered heat map of *Richelia* genes that were significantly periodic (RAIN, FDR < 0.05) where red indicates the maximum normalized expression (row max) and blue indicates the minimum normalized expression (row min) for that row (gene). Gray bars below the figure indicate dark periods (7 p.m. to 6 a.m.) while

yellow indicates light periods (6 a.m. to 7 p.m.). **b** The percent of significantly periodic *Richelia* genes with KEGG annotations clustered by KEGG module. KEGG modules mentioned in the text are labeled in the circle plot as a guide to the color legend. Each ring is labeled as percent increases from the center of the plot

Results and discussion

The oscillation from day to night is one of the strongest and most predictable perturbations imposed on organisms, selecting for pronounced diel patterns in both metabolism and growth [31]. Although it is expected that the diazotroph *Richelia* exhibits daily oscillations in N₂ fixation rates based upon diel patterns of *nifH* gene transcript copy abundance [32], little is known about diel patterns across other aspects of metabolism for *Richelia* and its diatom host. Here we mined high frequency (21 time points over 4.5 days) environmental metatranscriptomes for signals of a common bloom forming DDA *Rhizosolenia–Richelia* in order to investigate the diel rhythms in *Richelia* and identify

significantly co-expressed transcripts between the diazotroph symbiont and its diatom host.

The *Rhizosolenia–Richelia* DDA has been observed in a wide variety of environments and in the NPSG it is often the dominant organism in summer phytoplankton blooms [3]. During the sampling period, the presence of *Richelia* (associated with *Rhizosolenia*) was observed in micrographs taken from seawater filtered from a shipboard uncontaminated seawater line [33]. The presence of the DDA was further confirmed with quantitative PCR (qPCR) of *nifH* gene copy abundance [12], being the second most abundant diazotroph at 15 m after *UCYN-B* (Figure S2). Although *Richelia* was not the most abundant diazotroph, these results are consistent with the importance of the *Rhizosolenia–*

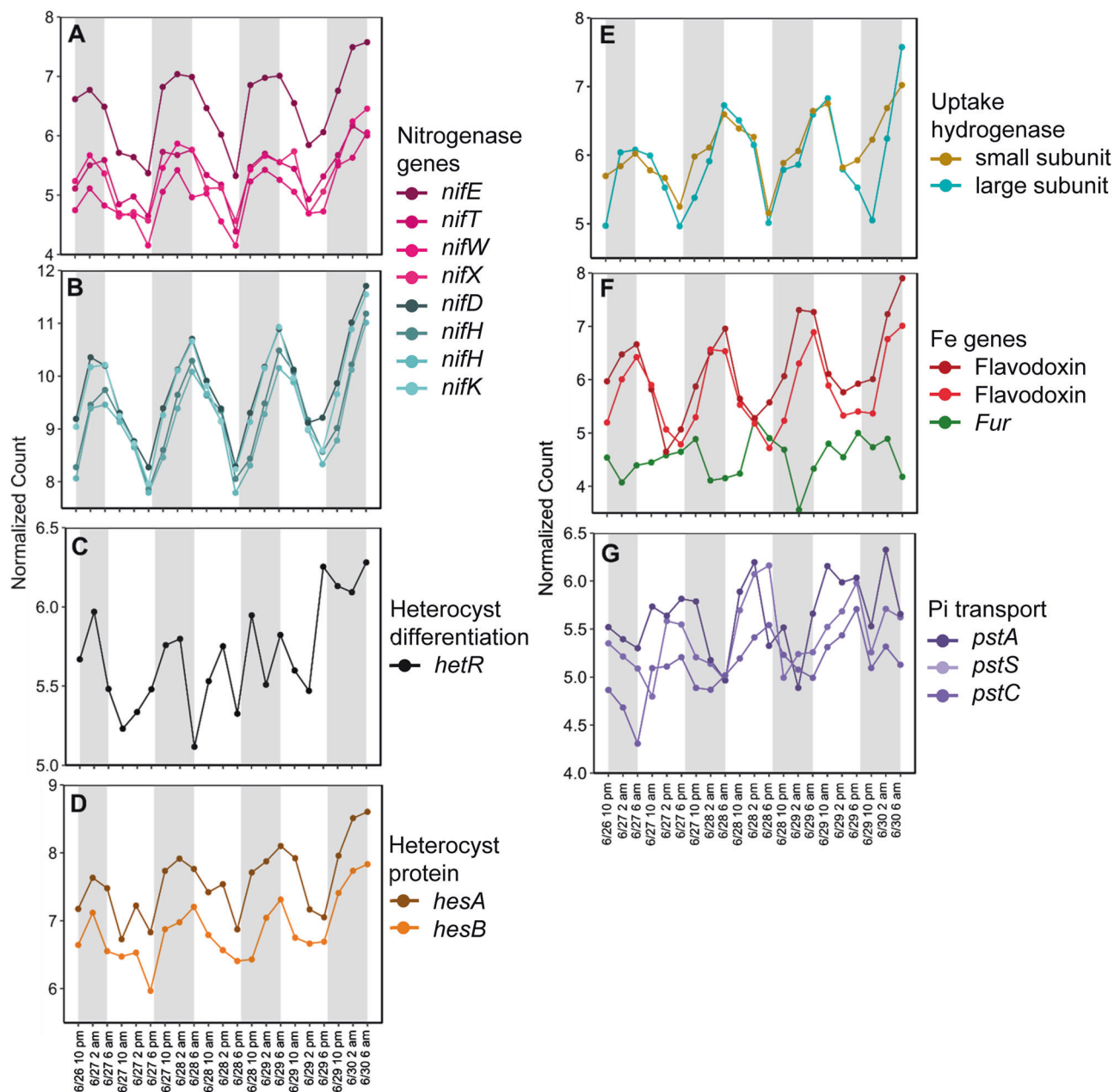


Fig. 3 Normalized expression patterns for genes in *Richelia* with significant periodic expression (RAIN, FDR < 0.05). **a** Nitrogen fixation genes with peak expression ~4 a.m. **b** Nitrogen fixation genes with peak expression ~6 a.m. **c** The heterocyst differentiation regulator *hetR*. **d** Heterocyst-associated genes with peak expression ~4 a.m.

e Uptake hydrogenase subunits with peak expression ~6 a.m. **f** Genes related to iron (Fe) homeostasis, and **g**) genes involved in phosphorus (Pi) acquisition. Gray shading highlights the dark periods (7 p.m. to 6 a.m.) while light periods are white (6 a.m. to 7 p.m.)

Richelia DDA in the NPSG during the summer export period [2, 3].

Patterns of diel gene expression in *Richelia* from the NPSG

RAIN analysis identified 1,053 genes with significant periodicity in *Richelia*, representing 25% of the analyzed transcriptome (Fig. 2a, Table S2). A similar magnitude of

the transcriptome (34%) was identified as significantly periodic in NPSG field populations of the free-living diazotroph *Crocospaera* [12] suggesting that although *Richelia* typically occurs in association with other organisms, it may (at least when in association with *Rhizosolenia*) be equally subject to diel forcing as other free-living diazotrophs. As with free-living diazotrophs, N₂ and photosynthetic carbon fixation processes exhibited significant periodicity [12, 34], in addition to RNA synthesis and

processing (Fig. 2b). Diel patterns in nitrogenase (*nif*) gene expression are also common in diazotrophs [34–36] and here a total of 16 genes were identified as encoding nitrogenase components in *Richelia*, of which 12 had significant periodicity (Table S2). The KEGG module for N metabolism contained the *nifHDK* nitrogenase genes with significantly periodic expression (Fig. 2b). Nitrogenase genes involved in assembly and incorporation of iron and molybdenum into nitrogenase (*nifE*, *nifT*, *nifW*, and *nifX*) [37] had peak expression around 4 a.m. (Fig. 3a), followed by expression of the core components of nitrogenase (the heterotetrameric core encoded by *nifD* and *nifK*, as well as the subunits encoded by *nifH*) [38], with peak times at 6 a.m. (Fig. 3b). This cascade in gene expression is consistent with a coordinated assembly of the functional enzyme components, and the timing mirrored prior investigations in the region focused solely on *Richelia nifH* gene expression [32].

In *Richelia*, N_2 fixation occurs in heterocysts and the heterocyst differentiation regulator, *hetR* (Rich_326, peak expression at 1 a.m.), heterocyst-associated genes *hesA* (Rich_5163, peak expression at 4 a.m.) and *hesB*, (Rich_5165, peak expression at 4 a.m.), as well as genes encoding an uptake hydrogenase system (Rich_4671 and Rich_4672, peak expression 7 a.m. and 6 a.m., respectively) were significantly periodic (Fig. 3c–e). Although the function of *hesA* and *hesB* are uncharacterized, they are thought to be linked to N_2 fixation, as insertional inactivation of *hesA* impairs N_2 fixation [39] and both are upregulated during N depletion [40]. Here, peak expression of these genes occurred prior to dawn (Fig. 3d), mirroring the pattern that is observed in other heterocystous cyanobacteria [41] and the pattern observed for the *Richelia nif* genes (Fig. 3a, b). Uptake hydrogenases, located in the heterocysts [42], are thought to efficiently recycle H_2 (consuming O_2 in the process) which is produced during N_2 fixation [43]. They may also act to provide ATP via the oxyhydrogen (knallgas) reaction or provide reducing equivalents to nitrogenase, further enhancing N_2 fixation [44–46]. In prokaryotes, transcription and translation are closely coupled in time and space with many genes translated shortly after transcription [47]. The diel oscillations in gene expression for N_2 fixation genes in *Richelia* therefore likely represent the biosynthesis and assembly of nitrogenase in the early morning in preparation for daytime N_2 fixation.

In the NPSG, seasonal fluctuations in phosphorus (P), iron (Fe), and light are thought to drive annual blooms of DDAs in the region [48], and Fe and P in diazotrophs is important for fueling maximal N_2 fixation [49]. Given the Fe demand associated with N_2 fixation, Fe homeostasis may be tightly controlled over the diel cycle. Consistent with this hypothesis a ferric uptake regulatory (Fur) protein, typically involved in maintaining Fe homeostasis [50, 51], was

significantly periodic in *Richelia* with peak expression at 4 p.m. (Fig. 3f, Table S2). Fur typically acts as a negative regulator of Fe stress genes, however, it can both induce and repress a suite of genes in other bacteria [52, 53]. When Fe is plentiful, Fur binds to Fe^{2+} acquiring a configuration that enables it to bind to target Fur boxes on the genome, inhibiting transcription. When iron is scarce, Fe^{2+} is released from Fur and transcription of Fe-related functions commences [54]. The expression pattern suggests that *Richelia* Fe quota and Fe demand are tightly modulated over the diel cycle.

Non-Fe requiring flavodoxins can substitute for ferredoxin to help mitigate Fe demand, and their induction can be an indicator of Fe stress in cyanobacteria [55]. Two flavodoxin transcripts (Rich_2552, Rich_7201) were significantly periodic (Fig. 3f), with peak expression near dawn (4 and 5 a.m., respectively), mirroring N_2 fixation genes. Flavodoxin genes have been shown to track *nif* gene expression patterns in other diazotrophs [34], consistent with the patterns observed here. Flavodoxins in cyanobacteria can be under the transcriptional control of Fur [50, 51]. In other bacteria, Fur regulates a cascade of genes starting with Fe transport, siderophore production, and then genes related to Fe substitution and sparing-related functions such as flavodoxin [53]. This potential cascade may explain the offset in peak time observed for Fur and the flavodoxin genes (Fig. 3f). Regardless of the expression pattern, these day–night patterns in Fur and flavodoxin expression suggest oscillations in Fe demand across the 24 h cycle. These apparent oscillations in Fe demand have also been observed in free-living diazotrophs like *Crocosphaera* [34, 56] and *Trichodesmium* [36] and may be a common feature of these genera.

As with the apparent diel oscillations in demand for Fe, P demand appeared to also fluctuate over a diel cycle. Peak expression of genes within the inorganic phosphate (Pi) transport system (*pstACS*; Rich_4810, 4809, and 1959) occurred between 1 and 3 p.m. and were significantly periodic (Fig. 3g), suggesting variation in P demand and homeostasis over the diel cycle. Similar diel modulations of *pstS* and other genes involved in P acquisition were reported in free-living diazotrophs [12, 57], suggesting this apparent modulation of P acquisition may be a common feature. Unlike the choreography observed between Fe demand and *nifH* expression (Fig. 3b), peak expression time for the *pst* genes (Fig. 3g) was offset relative to that of *nifH* (Fig. 3b). Given the direct role (e.g. enzyme co-factors) Fe has in N_2 fixation and photosynthesis, these apparent fluctuations in Fe demand may be more tightly linked to these processes than apparent fluctuations in P demand. The increased expression of *pstS* is often used as a marker of P stress [58–60]. In studies of P physiology, the diel modulations of *pstS* observed here for *Richelia* further emphasizes the

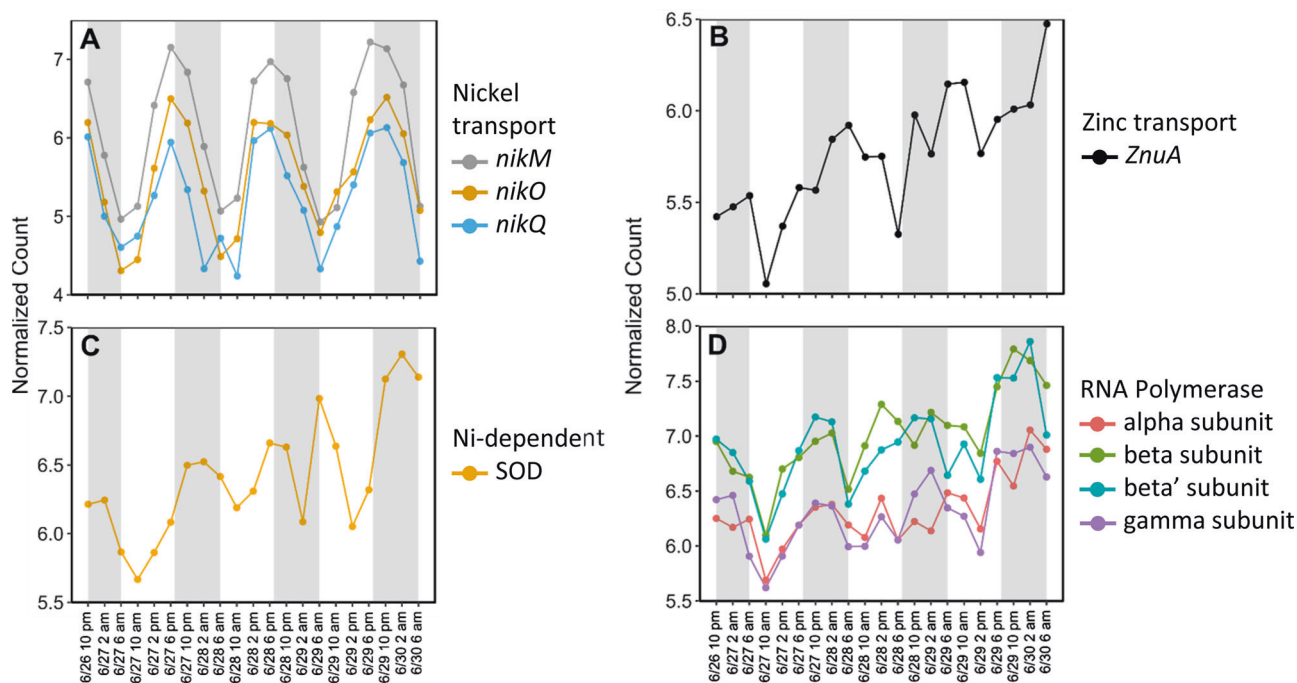


Fig. 4 Normalized expression patterns for genes in *Richelia* with significant periodic expression (RAIN, FDR < 0.05). **a** Nickel transport genes, **b** Ni-dependent superoxide dismutase (SOD), **c** zinc transport,

and **d** RNA polymerase subunits. Gray shading highlights the dark periods (7 p.m. to 6 a.m.) while light periods are white (6 a.m. to 7 p.m.)

importance of comparing samples from the same time of day to avoid diel variability which may confound interpretation of P stress signals observed in field populations [57].

Diel oscillations were apparent for additional resources in *Richelia*. For example, a set of Nickel (Ni) transport genes (Rich_1479, 1480, 1481, *nikMOQ*, 7–8 p.m. peak time) exhibited significant periodicity (Fig. 4a). Ni is an essential transition metal, present at low levels in oceanic environments [61] and is a co-factor in metalloenzymes like uptake hydrogenase and Ni-superoxide dismutase (Ni-SOD) [62]. Ni transport appears to be linked to the periodic expression of metalloenzymes as the Ni-dependent SOD (Rich_4190, 2 a.m. peak time, Fig. 4b) and Ni-dependent uptake hydrogenase subunits (Rich_4671, 4672, 6–7 a.m. peak time, Fig. 3e) were also significantly periodic (Table S2). In the diazotroph *Trichodesmium*, increased Ni availability is correlated with increased N₂ fixation where Ni-dependent enzymes like SOD and uptake hydrogenase reduce oxidative stress by removing oxygen radicals and utilizing released H₂ [63]. As such, daily modulation of Ni demand might be common in both symbiotic and free-living diazotrophs.

Coincident with oscillations in Ni was a significantly periodic *Richelia* Zn ABC-transporter (Rich_242, *znuA*), peaking in expression at 5 a.m. (Table S2, Fig. 4c). The modulation of this Zn transporter suggests diel oscillations in internal Zn inventory. In the NPSG, zinc (Zn) is depleted

in surface waters [64] but, like Ni, is required for many metalloenzymes critical to processes, such as carbon fixation (carbonic anhydrase), nucleic acid replication (RNA polymerase, reverse transcriptase), and phosphorus metabolism (alkaline phosphatase) [65]. Although the Zn-dependent alkaline phosphatase (*phoA*) and carbonic anhydrases were not significantly periodic, a group of alpha, beta, and gamma RNA polymerases (RNAP) were significantly periodic with peak expression after midnight (Rich_1744, 3195, 3196, 3197; Fig. 4d). Zinc appears to play a structural role in promoting RNAP assembly and additions of Zn promote production of RNAP [66, 67]. As RNA polymerase is transcribed, Zn inventories may become depleted prompting the cell to express genes for Zn transport.

Taken together, these diel patterns in resource-related gene targets suggests that both macro and micronutrient demand is tightly modulated in *Richelia* over diel cycles. These diel patterns could influence the cycling of these resources within the host–symbiont association, and the surrounding water column. More diel studies of field populations are required to know to what extent patterns in resource demand are consistent across other diazotroph associations and phytoplankton in general, and to what extent synchronous competition for resources might drive competitive outcomes and resulting bloom dynamics in oligotrophic regions.

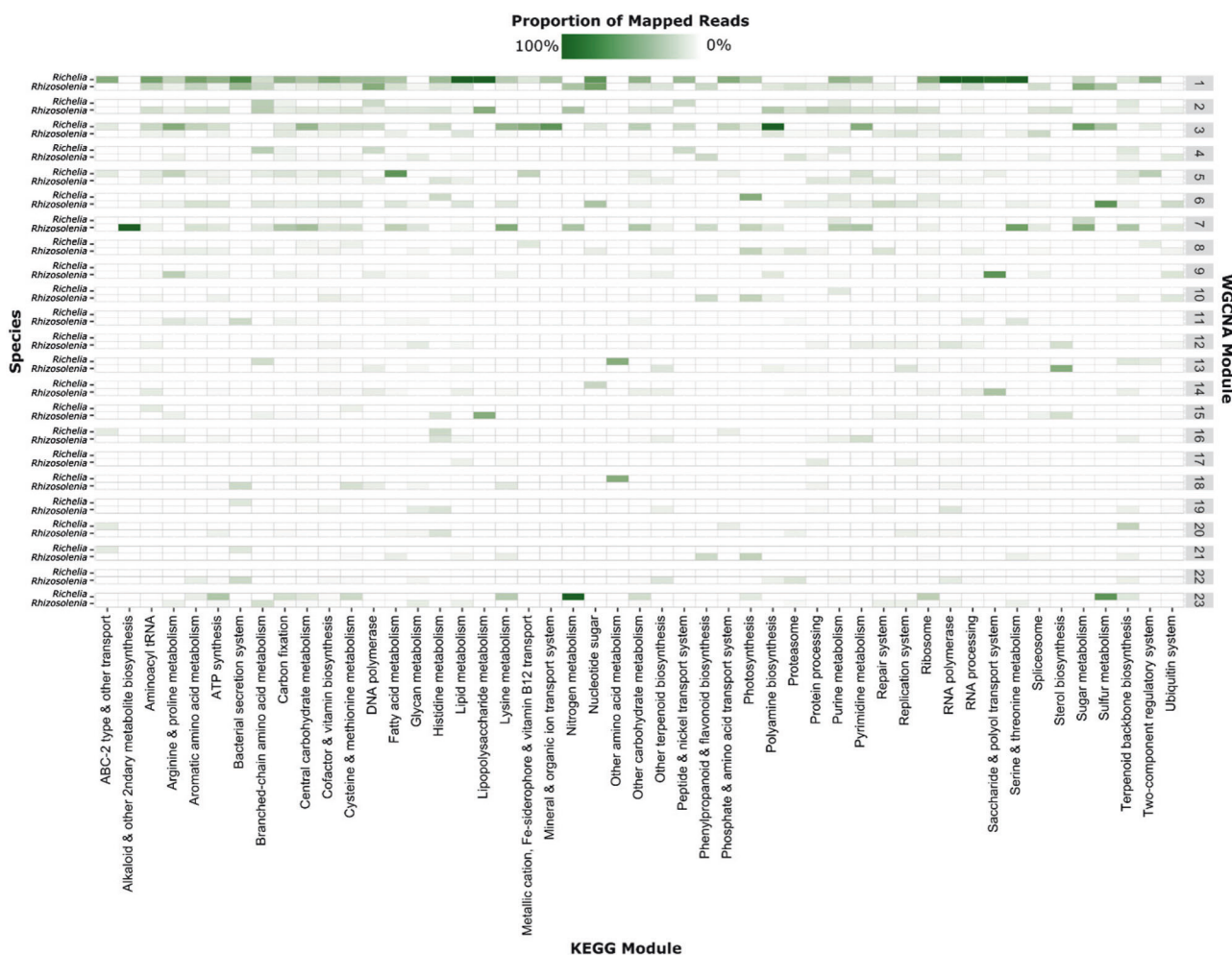


Fig. 5 The species-specific proportion of mapped reads per KEGG module and their distribution across WGCNA modules containing >200 nodes (*Richelia* genes or *Rhizosolenia* contigs). Color

corresponds to the proportion of mapped reads in each KEGG module (column) for each species visualized for WGCNA modules

Patterns of diel gene expression in *Rhizosolenia* from the NPSG

The analysis of selected reads from the same pool of RNA allowed us to also identify diel patterns in the diatom host *Rhizosolenia* and evaluate coordinated patterns in gene expression within the symbiosis. RAIN identified 398 *Rhizosolenia* contigs with significant periodicity in the NPSG, representing 2% of the analyzed transcriptome (Figure S3, Table S2). Many of these periodically expressed genes were related to photosynthesis, carbon fixation and related processes (Figure S4, Table S2). For example, 43 chlorophyll *alb* binding proteins were periodically expressed (Figure S4A) and significant periodicity was also observed in violaxanthin de-epoxidase (VDE), an enzyme that converts diadinoxanthin into diatoxanthin, protecting against photooxidative damage [68]. Peak expression of VDE in *Rhizosolenia* (1 p.m.; Figure S4B) was later than that observed in cultures of *Thalassiosira pseudonana*

(dawn) [69] perhaps suggesting differing strategies or responses to this ubiquitous problem in photosynthetic organisms. Lastly, phosphoglycerate kinase, which catalyzes an essential step in glycolysis was found to have peak expression in *Rhizosolenia* at 8 a.m. (Figure S4C) while in *T. pseudonana* this gene peaked in expression near dusk [69]. It is clear that further investigation is warranted to determine if differences between *Rhizosolenia* and *Thalassiosira* are intrinsic to different diatom genera or due to inherent differences between culture studies and observations of field populations.

Significantly coordinated patterns of expression in the *Rhizosolenia*–*Richelia* association

Chronobiological observations of symbiotic partnerships have revealed metabolic interdependence between host and symbiont, largely centered around the exchange of resources and cell division processes [70, 71]. To explore this

interdependence in the *Rhizosolenia*–*Richelia* partnership, a weighted gene co-expression network analysis (WGCNA) was used to examine patterns of co-expression between the partners over the time series. This analysis was independent of the RAIN analysis and was used to identify clusters (modules) of transcripts that were significantly correlated with either a synchronous or orthogonal pattern. Of the 126 modules constructed, *Richelia* genes were represented in all but five (Figure S5), suggesting that many cellular processes between the host and symbiont were significantly co-expressed. A number of WGCNA modules had patterns of day–night periodicity and contained features with significantly periodic expression (Figure S6). Some of this co-expression was due in part to the coordinated day–night expression of overlapping metabolic pathways like carbon fixation and nucleotide sugar metabolism (e.g. WGCNA module 1) (Fig. 5) and may reflect the shared response that phytoplankton have to light. However, the majority of the co-expression modules contained different KEGG pathways for each species (Fig. 5), and this variability may reflect each organism being individually influenced by day–night transitions. For example, N metabolism was observed in WGCNA modules 1, 2, and 7 for *Rhizosolenia* but only in module 23 for *Richelia* (Fig. 5). As another example, *Richelia* serine and threonine metabolism was predominately in WGCNA module 1, whereas *Rhizosolenia* serine and threonine metabolism was predominately in WGCNA module 7, reflecting different expression patterns for the host and symbiont (Fig. 5). The unique distribution of metabolic processes in the co-expression network analysis suggests that these organisms operate on different clocks in the same environment, where shared aspects of metabolism are offset. As with many host–symbiont relationships, it is assumed resource exchange is the primary function of this DDA partnership, and this metabolic offset may reflect this. However, it may also reflect inherent differences between diatoms and cyanobacteria. Additional surveys of diel patterns in field populations of diatoms and cyanobacteria would help resolve the extent to which these patterns reflect intrinsic differences between different photosynthetic taxa, versus patterns driven by the two organisms living in association. Despite these uncertainties, each species clearly has distinct metabolic patterns, and their metabolisms also appear to be tightly coordinated with each other. This significant coordination is likely important to maintaining the symbiosis and driving the exchange of resources between host and symbiont.

Potential exchange of resources between host and symbiont over light–dark cycles

The transfer of N between symbiont and host has been shown in DDA partnerships [6, 11] and to explore this

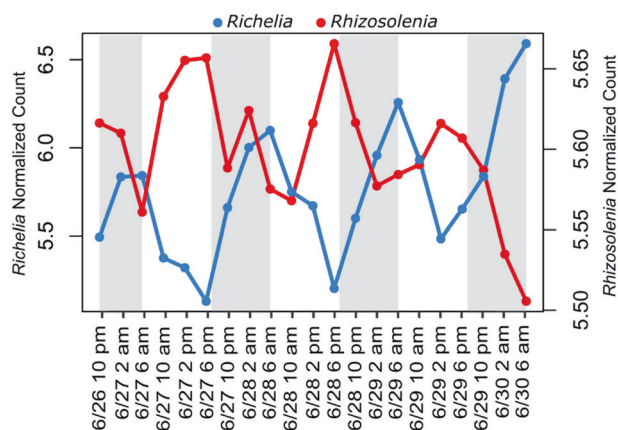


Fig. 6 Average normalized expression patterns of significantly co-expressed features within the WGCNA module 23 for *Richelia* (103 genes, $SE \pm 0.21$) and *Rhizosolenia* (105 contigs, $SE \pm 0.15$). Key features within this module relating to potential exchange of resources are a TRAP = tripartite ATP-independent periplasmic and nitrogenase gene (*nifHDKETWX*) in *Richelia* and transporters for ammonium, amino acids, major facilitator superfamily (MFS) and efflux and EAMA-like transporters in *Rhizosolenia*. Gray shading highlights dark hours (7 p.m. to 6 a.m.) while light periods are white (6 a.m. to 7 p.m.)

function we searched for WGCNA modules containing *nif* genes from *Richelia*. Module 23 of the WGCNA analysis contained 10 of the 12 significantly periodic nitrogenase genes (Fig. 6, Figure S8A) and these were significantly co-expressed with a *Rhizosolenia* ammonium transporter (Rhizo_7354), an amino acid permease (Rhizo_10488), and two major facilitator superfamily (MFS) proteins (Rhizo_156 and Rhizo_18969); a large gene family involved in the transport of amino acids among other potential substrates [72] (Fig. 6, Figure S8A). Although there are many factors regulating expression of N transport functions [73], these significantly co-expressed genes between the host and symbiont may, in part, underpin the known transfer of newly fixed nitrogen from *Richelia* to the diatom [11].

Diatoms in the NPSG are limited by N [74] and *Richelia* is thought to provide a distinct advantage to the diatom host by providing N via N_2 fixation. What *Richelia* gets back in return remains an open question. In legumes (as well as other plant/bacteria symbiosis), reduced carbon such as dicarboxylates, sugars, and amino acids are thought to be exchanged for fixed N_2 [75, 76]. Although *Richelia* expresses genes for photosynthesis (Fig. 2, Table S2), it may still import carbon substrates from the diatom host. In Module 23, *Rhizosolenia* genes for the transport of carbon substrates, including an EAMA-like transporter (Rhizo_1833) and an efflux transporter (Rhizo_14884), were significantly co-expressed with the *Richelia* nitrogenase genes (Fig. 6, Figure S8A). In the diatom *Phaeodactylum tricorutum*, EAMA-like transporters (named after the O-acetyl-serine/cysteine export gene in *E. coli*) were associated with triose-phosphate translocators (TPTs),

which are thought to export photosynthesis-derived carbohydrates [77]. These transporters may be involved in the export of substrates to *Richelia* in exchange for fixed N₂. In support of this hypothesis, a *Richelia* tripartite ATP-independent periplasmic (TRAP) solute receptor gene (Rich_1143) was also co-expressed in the same module (Fig. 6, Figure S8B). TRAP transporters are unique to bacteria and archaea and are involved in the uptake of a number of organic substrates, including sugars [78], and in this case their co-expression may be indicative of *Richelia* actively transporting substrates produced by the host.

In addition to N, diatoms can also be limited by trace nutrients such as vitamins, especially during bloom conditions [74]. As they are not able to synthesize vitamin B₁₂, diatoms in DDA partnerships may benefit from their cyanobacteria symbiont as a source for required vitamins such as B₁₂ [79]. Like other cyanobacteria, *Richelia* likely produces a form of B₁₂ (pseudocobalamin) that is unusable by most eukaryotic phytoplankton [80]. Notably, *Rhizosolenia* has genetic machinery to alter pseudocobalamin to a useable form (*cobST*; Rhizo_5562; Table S2), by replacing the axial ligand adenine with exogenous sources of 5,6-dimethylbenzimidazole (DMB) [80]. It appears both genes are present within the assembled contig. These genes are expressed in the NPSG but are not significantly diel (Table S2). In *Richelia*, a gene for cobalamin (B₁₂) synthase (Rich_4005; Table S3) was expressed at too low a level to pass our stringent analysis threshold and portions of the cobalamin biosynthesis pathway *cbiX* (Rich_4021,3078), *cobF* (Rich_5476,5860), and *cobN* (Rich_155) were not significantly periodic (Table S2), suggesting perhaps constitutive low-level exchange of this essential resource. It may be that other vitamins are also exchanged within the association. For example, *Richelia* biotin synthase (Rich_1540) expression is significantly periodic, as are two *Rhizosolenia* biotin ligases (Rhizo_7013 and Rhizo_9114) (Table S2), the latter of which activates biotin and transfers biotin to biotin-accepting proteins [81]. Diatoms are known to produce biotin, but may rely on exogenous sources of biotin in Fe-limited conditions [82]. Although the *Richelia* and *Rhizosolenia* biotin genes do not fall into the same WGCNA module, these diel patterns may still underpin host–symbiont biotin exchange in the low Fe NPSG. Taken together, these data are consistent with the exchange of vitamins within the association. If this is confirmed with further study, then the vitamins from *Richelia* may free the host from competition for vitamins like B₁₂ in addition to N.

Potential coordination of *Rhizosolenia*–*Richelia* growth

In *Rhizosolenia*, *Richelia intracellularis* is an extracellular endosymbiont located in the periplasmic space between the

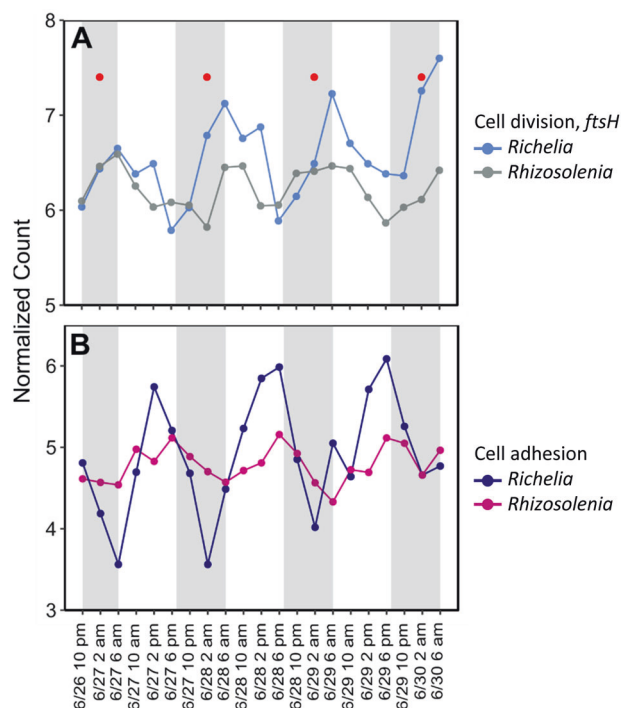


Fig. 7 Normalized expression patterns of **a** the putative cell division marker *ftsH* for both *Richelia* and *Rhizosolenia* and **b** cell adhesion gene forming pili in *Richelia* and a fasciclin domain contig in *Rhizosolenia*. All genes/contigs were significantly periodic (RAIN, FDR < 0.05). Red dots in plot A correspond to peak expression timing of silicic acid transporters (Figure S7) in *Rhizosolenia*. Gray shading highlights dark hours (7 p.m. to 6 a.m.) while light periods are white (6 a.m. to 7 p.m.)

plasmalemma and silica wall [83, 84]. Early culture work suggested asynchronous division frequency between host and symbiont despite similar growth rates in N-limited media [85]. However, little is known regarding natural population division or how the host and symbiont maintain this extracellular association. RAIN analysis identified a putative *ftsH* cell division marker with significant periodicity for both the *Rhizosolenia* host (Rhizo_8174) and the *Richelia* symbiont (Rich_842) with both having peak expression at 7 a.m. (Fig. 7a). Although the genes did not fall into the same WGCNA module, their peak times are consistent, suggesting similar division timing. Uptake and incorporation of silicic acid appears to be linked to cell division as well [86, 87] and so the timing in expression of these transporters in *Rhizosolenia* may be an effective proxy for cell division processes in natural populations of *Rhizosolenia*. Significantly periodic *Rhizosolenia* silicic acid transporters (Rhizo_8532 and Rhizo_9964) had peak expression at ~2 a.m. (Fig. 7a, Figure S7) slightly before the peak expression of the *ftsH* gene cell division markers observed here (Fig. 7a). Taken together, these expression patterns are consistent with a pattern of early morning

division that is coordinated between the host and symbiont. If this pattern is supported with additional observations, then the association may be maintained, at least in part, by a coordinated pattern of cell division which is synchronized over the diel cycle in field populations.

Fasciclin domain proteins in symbioses are thought to be related to adhesion between host and symbiont and they have been found in a wide variety of symbiotic associations [88], where increased expression of fasciclin domain proteins in the host has been linked to the presence of the symbiont [89]. WGCNA Module 3 contained a significantly periodic *Rhizosolenia* fasciclin domain protein (Rhizo_11521) with peak expression at 5 p.m. (Fig. 7b). This fasciclin domain protein was significantly co-expressed with a *Richelia* gene related to pilus formation (Rich_7610) (Fig. 7b). Filamentous surface appendages called pili mediate pilus-dependent adhesion in other associations, such as those found in N₂-fixing bacteria associated with plant roots [90]. Given these coordinated expression patterns, we hypothesize that pili formation and related processes in the diatom host may help maintain the symbiosis over the day–night cycle as each species divides. Collectively, these gene expression patterns suggest that host and symbiont ecology is tightly integrated over the diel cycle and these data provide gene targets with which to further evaluate the ecology of this DDA.

Conclusions

DDAs such as the *Rhizosolenia*–*Richelia* symbiosis are known to exert profound control over the cycling of carbon and nitrogen, particularly in oligotrophic regions like the NPSG, yet there are still major knowledge gaps regarding DDA physiological ecology. In surface ocean prokaryotic communities, diel patterns of synchronous gene expression suggest there is cross-species coordination driving carbon metabolism and cycling in the community [25]. Here, we observed offsets in the expression of core metabolic processes in a photosynthetic host–symbiont pair. These differences may underpin the ability of these two species to live together by providing a temporal compartmentalization of metabolism. Temporal compartmentalization could minimize competition for key resources, while facilitating exchange of resources, like N, and maintain the processes that coordinate growth and stabilize the association over light–dark transitions. Given that oligotrophic waters are expanding due to climate change, with the NPSG expanding most rapidly [91, 92], DDAs may be increasingly important in the future ocean, underscoring the importance of understanding the physiological ecology of this keystone symbiosis.

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Compliance with ethical standards

Conflict of interest The authors declare that they have no conflict of interest.

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