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Global distribution and vertical patterns of a prymnesiophyte-cyanobacteria obligate symbiosis

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Running title: Distribution of a prymnesiophyte–UCYN-A symbiosis

Subject Category: Microbial population and community ecology

Abstract

25 A marine symbiosis has been recently discovered between prymnesiophyte species and the unicellular diazotrophic cyanobacterium UCYN-A. At least 2 different UCYN-A phylotypes exist, the clade UCYN-A1 in symbiosis with an uncultured small prymnesiophyte and the clade UCYN-A2 in symbiosis with the larger *Braarudosphaera bigelowii*. We targeted the prymnesiophyte–UCYN-A1 symbiosis by double CARD-FISH (CAlyzed Reporter
30 Deposition-Fluorescence *In Situ* Hybridization) and analyzed its abundance in surface samples from the MALASPINA circumnavigation expedition. Our use of a specific probe for the prymnesiophyte partner allowed us to verify that this algal species virtually always carried the UCYN-A symbiont, indicating that the association was also obligate for the host. The prymnesiophyte–UCYN-A1 symbiosis was detected in all ocean basins, displaying a patchy
35 distribution with abundances (up to 500 cells ml⁻¹) that could vary orders of magnitude. Additional vertical profiles taken at the NE Atlantic showed that this symbiosis occupied the upper water column and disappeared towards the Deep Chlorophyll Maximum, where the biomass of the prymnesiophyte assemblage peaked. Moreover, sequences of both prymnesiophyte partners were searched within a large 18S rDNA metabarcoding dataset from
40 the *Tara*-Oceans expedition around the world. This sequence-based analysis supported the patchy distribution of the UCYN-A1 host observed by CARD-FISH, and highlighted an unexpected homogeneous distribution (at low relative abundance) of *B. bigelowii* in the open ocean. Our results demonstrate that partners are always in symbiosis in nature and show contrasted ecological patterns of the two related lineages.

45 Key words: double CARD-FISH/global abundance/prymnesiophyte host/UCYN-A/vertical distribution

Introduction

Symbiosis between cyanobacteria and eukaryotic organisms is a widespread phenomenon reported both in land and in aquatic systems (Rai *et al.*, 2002). In the marine environment, cyanobacteria appear associated with multicellular organisms, such as ascidians and sponges, and with single-celled organisms, such as diatoms, dinoflagellates, radiolarians, and tintinnids (Carpenter and Foster, 2002; Foster *et al.*, 2006). These findings derive from microscopic observations and often the mutual benefit between the host and the symbiont is poorly understood. Generally, it is assumed that the cyanobacteria provides organic carbon to the host through photosynthesis, while the host provides a stable and secure environment.

Diazotrophic cyanobacteria symbionts can in addition provide nitrogen-derived metabolites through N₂ fixation, as has been demonstrated in some diatom species (Foster *et al.*, 2011).

A related and particular symbiosis has been recently described between two uncultured picoplankters, a small ($\leq 3\mu\text{m}$) prymnesiophyte and the unicellular diazotrophic cyanobacteria UCYN-A (Thompson *et al.*, 2012). Genomics revealed that UCYN-A lacked key metabolic pathways commonly shared by diazotrophic cyanobacteria, such as oxygenic photosynthesis and the Calvin cycle, suggesting it was an obligate symbiont (Zehr *et al.*, 2008; Tripp *et al.*, 2010). Indeed, nanoSIMS demonstrated that the symbiont gave fixed nitrogen to the host and obtained organic carbon in return (Thompson *et al.*, 2012). The 18S rDNA prymnesiophyte sequence, identified by single cell analysis (Thompson *et al.*, 2012), was identical to an environmental sequence from the South Pacific (Shi *et al.*, 2009) and related (98.2%) to *Braarudosphaera bigelowii*, a coastal nanoplanktonic coccolithophore (Hagino *et al.*, 2009). Later, endosymbiotic UCYN-A was also discovered in *B. bigelowii* by TEM observations (Hagino *et al.*, 2013). Phylogenetic analyses using the UCYN-A nitrogenase (*nifH*) gene revealed at least three distinct clades, UCYN-A1, -A2 and -A3 (Thompson *et al.*, 2014), and

highlighted specificity between host and symbiont pairings, suggesting co-evolution between symbionts and hosts (Bombar *et al.*, 2014; Thompson *et al.*, 2014). Thus, the clade UCYN-A1 was associated with an open ocean small prymnesiophyte (hereafter UCYN-A1 host), and the clade UCYN-A2 was associated with the coastal and larger *B. bigelowii* (hereafter UCYN-A2 host), whilst no host has been yet proposed for UCYN-A3.

Environmental surveys of *nifH* genes indicated a rather global distribution of this symbiosis in the oceans (Moisander *et al.*, 2010). Further reports showed that the distinct UCYN-A clades were widespread and often coexisted (Thompson *et al.*, 2014). Nevertheless, microscopic observations of the symbiosis or sequencing data of the prymnesiophytes verifying this widespread distribution are still scarce. Recently, a double CARD-FISH (CAlyzed Reporter Deposition-Fluorescence *In Situ* Hybridization) approach targeting UCYN-A and prymnesiophyte cells, allowed visualization and quantification of this association in the North Atlantic (Krupke *et al.*, 2014a,b). Later, the same double CARD-FISH technique but specifically targeting the two different prymnesiophyte host phlotypes made it possible to distinguish both types of associations and to observe their co-occurrence in the subtropical South Atlantic (Cornejo-Castillo *et al.*, submitted). The emerging picture derived from the above studies was that the prymnesiophyte–UCYN-A association showed a tendency for being more abundant in oligotrophic warm waters, but the environmental drivers explaining its distribution are still poorly understood. In this sense, quantifying the symbiosis by both CARD-FISH and 18S rDNA metabarcoding in a large set of samples covering a wide geographic range help evaluating accurately its distribution in marine ecosystems.

In this work we studied the distribution of UCYN-A1 and UCYN-A2 hosts in samples representative of the World's major oceans. First, we targeted the prymnesiophyte–UCYN-A1 symbiosis by double CARD-FISH using the specific probe for the UCYN-A1 host (Cornejo-Castillo *et al.*, submitted) and the general probe UCYN-A732, targeting all clades of UCYN-

A (Krupke *et al.*, 2013). This FISH setup was used to quantify cell abundances in surface waters tracked during the MALASPINA expedition, to analyze vertical distributions in the NE Atlantic near the Iberian Peninsula, and to evaluate whether the host can appear without the cyanobacteria. Second, we analyzed the global distribution of both hosts within an 18S
100 rDNA metabarcoding dataset from the *Tara-Oceans* expedition sampled at surface and DCM depths (de Vargas *et al.*, 2015). The aim of this work was to confirm the broad distribution of the prymnesiophyte–UCYN-A association, evaluate its putative ecological niche and provide further evidence about the obligatory dependence of both symbiont and host.

105 **Materials and methods**

Oceanographic cruises

We analyzed data from three independent cruises. The MALASPINA circumnavigation expedition took place from December 2010 to July 2011 on board the R/V *Hespérides* and tracked subtropical and tropical latitudes of the Atlantic, Indian and Pacific oceans (see Figure
110 S1a for a map of stations). The *Tara-Oceans* expedition (Karsenti *et al.*, 2011) took place from September 2009 to March 2012 on board the *Tara* schooner and encompassed a different track on the same oceans plus the Mediterranean Sea and subantarctic waters (Figure S1b). The INDEMARES 0710 cruise took place in summer 2010 (28 July - 15 August) on board the R/V *Thalassa* (IFREMER / IEO) in the NE Atlantic near the Iberian peninsula
115 (Figure S1a). For MALASPINA and INDEMARES, sampling and acquisition of hydrological variables are detailed below. Seawater sampling, DNA extract collection and 18S rDNA sequencing for the *Tara-Oceans* expedition is described in detail elsewhere (de Vargas *et al.*, 2015).

Sampling procedures during MALASPINA and INDEMARES cruises

120 During the MALASPINA cruise, surface samples (~3 m) for CARD-FISH and chlorophyll a
analysis were collected with a 30L Niskin bottle at around noon in 59 stations (minimum
bottom depth ~2 000 m). Conductivity-Temperature-Depth (CTD) casts were performed from
surface to mesopelagic depths with a Seabird 911Plus probe mounted on a 24-bottle Niskin
rosette and water was taken at several depths for nutrient analysis. In the INDEMARES
125 cruise, vertical profiles were taken at offshore stations in the Avilés canyon (stations A-B)
and the Galician Bank (stations C-D). Casts were performed after sunset with a CTD profiler
fitted with a Fluorometer (WET labs, ECO-AFL) and a 24-bottle Niskin rosette. Based on the
downcast fluorescence profile, seawater was collected at 7-8 depths from below the Deep
Chlorophyll Maximum (DCM) to surface (~5 m), with higher frequency above the DCM. For
130 chlorophyll a determination, aliquots of 250 ml (MALASPINA) or 2 L (INDEMARES) were
filtered through Whatman GF/F filters (25 mm diameter) and stored at -20°C until extraction
on board or in the lab, respectively. Samples for inorganic nutrients were drawn into
polyethylene vials and kept frozen until analysis. For the CARD-FISH assay, 95 ml aliquots
for MALASPINA or 45 ml aliquots for INDEMARES were fixed with 37% formaldehyde
135 (2% or 4% final concentration, respectively), filtered on 0.6 µm pore size Nucleopore filters
(25 mm diameter) and kept frozen until processed.

Chlorophyll a and inorganic nutrient measurements

During MALASPINA, pigments were extracted by placing the filters in 5-7 ml of 90%
acetone at 4°C for 24 h and determining the fluorescence of the extract in a Turner Designs
140 fluorometer (Yentsch and Menzel, 1963). A Chlorophyll a standard (Sigma Aldrich) was used
to calibrate the fluorometer and no phaeophytin correction was applied. For INDEMARES
samples, pigments were extracted with 90% acetone, sonicated, kept at -20°C for 24 h and

cleared by filtration through GF/F filters. Total chlorophyll a was determined by HPLC following the procedure described by Latasa (2014).

145 In MALASPINA samples, nitrate (NO_3^-) concentration was measured spectrophotometrically with a Skalar autoanalyzer (Skalar SANplus) following standard procedures (Grasshoff *et al.*, 1999; Moreno-Ostos, 2012), and phosphate (PO_4^{3-}) concentration was measured manually with a Perkin Elmer spectrophotometer. For INDEMARES, both nutrients were measured with a Skalar autoanalyzer. Detection limits were $0.02 \mu\text{M}$ for NO_3^- and $0.01 \mu\text{M}$ for PO_4^{3-} .

150 N/P ratios were calculated from the ratio between significant slope values ($P < 0.05$) of nitrate and phosphate across the nutricline. Dissolved organic nitrogen and phosphorus (DON and DOP) were estimated as the difference between total (TN and TP) and inorganic values. TN and TP concentrations were determined by nitrate and phosphate measures done after alkaline and acidic persulphate oxidation, respectively (Grasshoff *et al.*, 1999; Moreno-Ostos, 2012).

155 *Double CARD-FISH assay*

The double CARD-FISH assay was already used for the same symbiosis (Krupke *et al.*, 2014a) and followed the Multi-color CARD-FISH protocol (Pernthaler *et al.*, 2004). Filters were embedded in 0.1% low gelling point agarose to minimize cell loss and treated with lysozyme (37°C, 1 h) and acromopeptidase (37°C, 0.5 h) to permeabilize cell covers. For the
160 first CARD-FISH step, we used probe UPRYM69 (5'-CACATAGGAACATCCTCC-3') specific for the UCYN-A1 host. Design, specificity and optimization to ensure maximal stringency of the probe are detailed in Cornejo-Castillo *et al.* (submitted). This probe was combined with oligonucleotide helpers (sequences contiguous to the probe region [Helper A-PRYM 5'-GAAAGGTGCTGAAGGAGT-3'; Helper B-PRYM 5'-
165 AATCCCTAGTCGGCATGG-3']) and a competitor (having one mismatch in the probe region [5'-CACATTGGAACATCCTCC-3']). Filter pieces (1/8 of the filter) were covered

with hybridization buffer (40% deionized formamide, 0.9 M NaCl, 20 mM Tris-HCl pH 8, 0.01% sodium dodecyl sulfate [SDS], and 20 mg ml⁻¹ Blocking reagent [Roche Diagnostic Boehringer]) containing a mixture of the HRP-labeled probe, helpers and competitor each at 5
170 ng μl⁻¹. After incubating at 46°C overnight, two washing steps of 10 min were done at 48°C
in a buffer (56 mM NaCl, 5 mM EDTA, 0.01% SDS, 20 mM Tris-HCl pH 8), and filters were
equilibrated in PBS buffer for 15 min at room temperature (RT). Tyramide signal
amplification (TSA) was done for 40 min at RT in the dark in a solution of 1x PBS, 2 M
NaCl, 1 mg ml⁻¹ blocking reagent, 100 mg ml⁻¹ dextran sulfate, 0.0015% H₂O₂ and 4 μg ml⁻¹
175 Alexa 488-labeled tyramide. Filters were transferred to PBS for 20 min, rinsed with distilled
water and air-dried. Before the second hybridization, probe peroxidases were inactivated with
0.01M HCl for 10 min at RT in the dark (Pernthaler *et al.*, 2004) and filters were rinsed twice
with MilliQ water and air-dried. The second CARD-FISH used the probe UCYN-A732,
targeting both UCYN-A1 and -A2 cells, and the hybridization conditions described in Krupke
180 *et al.* (2013). Filters were embedded in the hybridization buffer (50% formamide, 0.9 M
NaCl, 20 mM Tris-HCl pH 8, 0.01% SDS, 10 mg ml⁻¹ Blocking reagent and 100 mg ml⁻¹
dextrane sulfate) with the probe and helpers at 0.16 ng μl⁻¹, incubated at 35°C for 3 h, then
rinsed in a washing buffer (as described above but with 9 mM NaCl) at 37°C, and TSA was at
46°C in the same solution described above but using 1 μg ml⁻¹ Alexa 594-tyramide. Filters
185 were counterstained with 5 μg ml⁻¹ DAPI (4', 6- diamidino-2-phenylindole), mounted in
antifading reagent (77% glycerol, 15% VECTASHIELD, and 8% 20x PBS) and kept frozen.

In a few samples, double CARD-FISH was done targeting *B. bigelowii*, using the specific
UBRADO69 probe (5'-CACATTGGAACATCCTCC-3') together with the same helpers than
before and a different competitor (5'-CACATAGGAACATCCTCC-3') (Cornejo-Castillo *et*
190 *al.*, submitted), and probe UCYN-A732 under the same hybridization and amplification
conditions described above. In INDEMARES samples, the class-specific probe PRYM02

(Simon *et al.*, 2000) was applied to target all prymnesiophytes in addition to the specific probe UPRYM69. Hybridization and TSA conditions for probe PRYM02 were as described for probe UPRYM69 with modified hybridization temperature (35°C) and NaCl concentration
195 in the washing buffer (37 mM).

Epifluorescence microscopy counts

Filters were observed by epifluorescence microscopy (Olympus BX61) at 1000x under UV (DAPI signal of the nucleous), blue light (green labeled host cells with Alexa 488) or green light (red labeled symbionts with Alexa 594) excitations. Counts of hybridized cells were
200 performed in 3-4 transects (~8.0 x 0.1 mm each) across the filter piece, which were analyzed twice. A first inspection was done under blue light to count host cells and examine the presence of associated symbionts by switching to green light. A second analysis was done under green light to detect free symbionts. Detection limit was about 2 cells ml⁻¹ (calculated assuming only one counted cell). Cell counts included four categories: 1) labeled host cells in
205 association with labeled UCYN-A cells, 2) non-labeled host cells with labeled UCYN-A cells, 3) labeled hosts without symbionts (non-associated hosts), and 4) labeled non-associated UCYN-A cells (free symbionts). Prymnesiophyte cells were classified into five size classes (1-2, 2-3, 3-4, 4-5, and >5 µm) by measurements with an ocular micrometer. Micrographs were taken with an Olympus DP72 camera (Olympus America Inc.) attached to the
210 microscope.

Similarity search of prymnesiophyte sequences in the Tara-Oceans expedition

The Tara-Oceans expedition studied protist diversity in several size fractions by 18S rDNA metabarcoding (de Vargas *et al.*, 2015). Here we analyzed the V9 18S rDNA Illumina reads derived from the 0.8-5 µm size fraction taken at surface and DCM in 40 stations. High quality
215 reads were clustered at 100%, derived OTUs were taxonomically classified to remove OTUs

from Archaea, Bacteria and Metazoa. From the final OTU table (73 samples with 87 millions reads), we noted the total number of reads and the number of prymnesiophyte reads per sample. Then, we did BLAST searches to identify the two prymnesiophyte phylotypes using sequences Biosope T60.034 (FJ537341) and *B. bigelowii* Genotype III (AB250784) as seeds.

220 Two OTUs identical to the seeds were identified, represented by 62 423 and 29 803 reads for the UCYN-A1 and UCYN-A2 hosts, respectively. These two phylotypes were also found in the 5-20 μm size fraction but at lower abundance (1 010 and 3 244 reads, respectively), so we decided to use here only the data for the 0.8-5 μm size fraction. V9 rDNA dataset is available at Pangaea under doi <http://doi.pangaea.de/10.1594/PANGAEA.843018>. Identifiers of the

225 OTUs were 07f330a1423bc41c2473a1a93a47f94b for the UCYN-A1 host and c2716f4902d581e7907e3c0a33917c17 for the UCYN-A2 host.

Results

Microscopic observations

230 We targeted the prymnesiophyte–UCYN-A1 symbiosis by double CARD-FISH in 59 surface samples from all major ocean basins taken during the MALASPINA expedition and in 30 water column samples from vertical profiles in the NE Atlantic taken in the INDEMARES cruise (see Figure S1a for a map of stations). Symbiosis was detected in about 50% of these samples. The labelling with probe UPRYM69 allowed us to distinguish the specific host from

235 the other prymnesiophyte cells (Figure 1a). Most cells targeted by probe UPRYM69 (86%, $n=2\ 030$ cells) were associated with UCYN-A. These prymnesiophyte cells were always very small and had a coherent size among samples. About 60% of the cells observed were between 1 and 2 μm in size, 38% within 2-3 μm , and 2% within 3-4 μm , resulting in a typical size of ~ 2.5 μm .

240 The specific prymnesiophyte was associated with one UCYN-A cell in most cases, but some
carried two symbionts (Figure 1b). This was rarely seen in MALASPINA samples, only in 5
out of 24 samples and always at very low abundance (up to 2% of the observed cells). On the
other hand, host cells with two symbionts were more frequent in INDEMARES samples,
being detected in virtually all samples along the four vertical profiles analyzed. On average,
245 25% of the observed host cells in station A carried two UCYN-A cells, with lower values in
the other casts (2.4% in station B, 6.4% in station C, and 5.0% in station D).

We never observed UCYN-A cells associated with non-prymnesiophyte eukaryotes. In a few
cases we detected UCYN-A cells hosted by larger prymnesiophyte-like cells not labeled by
probe UPRYM69 (Figure S2). In some of these samples, the larger host was later positively
250 targeted with the UBRADO69 probe, verifying that it was *B. bigelowii* (Figure 1c). All *B.*
bigelowii cells counted (n=17) were within the 4-5 μm size fraction.

To characterise the dependence of the prymnesiophyte host to UCYN-A1, we categorized the
samples based on the observations of free partners (Table S1). In general, free partners were
more frequent in samples that had high symbiosis abundances ($131 \pm 141 \text{ cells ml}^{-1}$, n=22) than
255 in samples with low symbiosis abundances ($32 \pm 27 \text{ cells ml}^{-1}$, n=14). Samples where free
partners were not detected presented the lowest symbiosis abundances ($15 \pm 14 \text{ cells ml}^{-1}$,
n=11). Moreover, by combining the counts from the 47 samples with symbiosis, we observed
that non-associated host abundances were positively related to free symbiont abundances
(Figure 2a). The similar abundances of both free partners strongly suggests that they derive
260 from the same association, being likely detached due to sample processing (Figure S3). There
was no significant relationship between partners likely detached and symbiotic states (Figure
2b), implying that the extent of the putative disruption depended on how particular samples

were handled. A rough estimate suggests that between 10 and 20% of the original symbiosis could be disrupted by sample manipulation.

265 *Global distribution in surface waters*

The abundance of the prymnesiophyte–UCYN-A1 association in surface waters was analyzed by double CARD-FISH in 59 MALASPINA and 3 INDEMARES samples. CARD-FISH counts showed that the symbiosis was present in all major ocean basins usually in patches of neighbouring stations (Figure 3a). It was detected in 27 out of 62 surface samples collected in
270 geographic sites with different environmental settings (Figure S4). There were some hotspots of high abundance in the NE Atlantic (476 cells ml⁻¹ in station A), near the Hawaiian archipelago (128 and 213 cells ml⁻¹ in stations 100 and 101, respectively) and in the Indian Ocean close to South Africa (140 cells ml⁻¹ in station 45) or Australia (338 and 397 cells ml⁻¹ in stations 69 and 64, respectively). Excluding these six stations, the averaged cell counts for
275 the symbiosis was 28 cells ml⁻¹ (SD= 25; n=21).

We occasionally observed UCYN-A cells associated with larger prymnesiophytes non-targeted by the UPRYM69 probe (Figure S2). In two of the five samples where this was seen, we did an additional hybridization with probe UBRADO69, specific for UCYN-A2 hosts. These larger prymnesiophytes were labeled with this new double CARD-FISH setup, showing
280 that they were the UCYN-A2 hosts. Counts of these larger prymnesiophytes were 8 cells ml⁻¹ in the eastern subtropical North Atlantic (station 144), and 18 cells ml⁻¹ in the southeast Indian Ocean close to Australia (station 68), and in both stations they co-occurred with UCYN-A1 hosts.

The two UCYN-A hosts were then searched in the 18S rDNA metabarcoding dataset obtained
285 from surface samples of the *Tara*-Oceans expedition. Prymnesiophytes were present in all *Tara* samples (n=40) (Figure 5a) in averaged percentages of 3.9% (±3.0) of total community

reads, and maximal values of ~12% in the Southern Ocean (station 85) and minimal of ~0.8% in the eastern South Pacific gyre (station 100). UCYN-A1 and UCYN-A2 hosts appeared to follow different distributions, as inferred by the contribution of their reads to total
290 prymnesiophyte reads (Figure 3b). The UCYN-A1 host appeared in a few spots with a high relative abundance (~30% of prymnesiophytes in two South Atlantic stations), while it was barely detected (< 0.1%) in 10 stations and undetected in 25 additional ones. On the other hand, the UCYN-A2 host appeared in virtually all samples (in 38 out of 40 stations; only undetected in stations 84 and 85 from the Southern Ocean), displaying an averaged
295 percentage of 1.2% (± 0.9).

Ancillary parameters measured during the cruises did not explain clearly the presence and abundance of the prymnesiophyte–UCYN-A1 symbiosis (Figure S4). The symbiosis was detected in a wide range of temperatures (19°C to 29°C), in ultraoligotrophic to eutrophic sites (chl *a* values from 0.035 to 0.65 mg m⁻³), in phosphate ranges from 0.01 to 0.23 μ M, in nitrate
300 ranges from < 0.02 to 1.38 μ M, and in N/P ratios from 6 to >20. Indeed, symbiosis abundances were only weakly correlated to temperature and phosphate, while they were not correlated with chlorophyll *a*, nitrate, N/P ratio, and dissolved organic phosphorus or nitrogen (Figure 4). In order to constrain the occurrence of the symbiosis we took into account all surface samples and compared its abundance in samples inside and outside a given range of
305 the environmental variables (Table 1). Abundances were significantly higher in samples with temperatures lower than 25°C, and phosphate lower than 0.08 μ M. No differences in abundances were found between samples with nitrate concentrations below and above the median value observed in all samples (0.27 μ M) or between samples with N/P ratios below or above 16. Moreover, symbiosis abundances were not significantly different in oligo, meso- or
310 eutrophic waters, although they were significantly lower in ultraoligotrophic stations.

Vertical distribution pattern in the water column

The vertical distribution of the prymnesiophyte–UCYN-A1 symbiosis was studied in detail in four vertical profiles in the NE Atlantic (Figure 6). A stratified water column was observed in all cases with a DCM peak coupled to the beginning of the nutricline, and depths above the
315 DCM having low nitrate and phosphate concentrations ($0.023 \pm 0.025 \mu\text{M NO}_3$; $0.075 \pm 0.013 \mu\text{M PO}_4$). At selected depths, symbiosis counts together with total prymnesiophyte counts were obtained by double and single CARD-FISH assays, respectively. In all profiles, UCYN-A1 host cells occupied the upper water column and decreased towards the DCM, where they disappeared. The decrease in symbiosis abundances appeared more coupled to the
320 beginning of the fluorescence gradient than to the beginning of the nutricline. On the contrary, the other prymnesiophyte cells, not involved in symbiosis, peaked at the DCM and were present throughout the photic water column. Station B lacked the surface samples, but its vertical pattern was consistent with the other three profiles. The abundance and contribution of the UCYN-A1 host varied between the two regions sampled. Highest abundances were
325 seen in the upper water column of station A ($435 \text{ cells ml}^{-1}$ on average) in the Avilés canyon region, where they represented on average 70% of the prymnesiophyte cells (up to 86% at 5 m). In the Galician Bank region, UCYN-A1 host abundances (40 to 80 cells ml^{-1} on average) had a lower contribution to total prymnesiophytes (8-16% on average).

The vertical distribution of the symbiosis could also be inferred by the contribution of UCYN-
330 A host reads in surface versus DCM samples of the *Tara-Oceans* cruise. Considering only the stations with high UCYN-A1 host abundance in surface samples (stations 76, 78 in the South Atlantic; station 4 in the Gibraltar Strait; and station 32 in the Red Sea), the contribution at the DCM was about 150 times lower (Figure 5b). In the other stations where UCYN-A1 host was scarce at surface, it was equally scarce at the DCM. By contrast, UCYN-A2 host was detected
335 at similar abundance in surface and DCM samples in all stations where it appeared (averaged

surface/DCM ratio of 1.38 ± 1.19) (Figure 5c). The contribution of UCYN-A2 host was only significantly lower at the DCM (as compared with surface) in stations where the DCM was deeper than 60 m (Mann-Whitney Test; $P = 0.001$, $n = 29$), a difference not seen for total prymnesiophytes (Figure 5a).

340

Discussion

Characterization of the partnership

Microscopic observations revealed a population size spectra and phenotype consistent with previous reports, with small prymnesiophyte host cells of 2-3 μm typically with one
345 cyanobacterial UCYN-A1 symbiont (Thompson *et al.*, 2012; Krupke *et al.*, 2014a). On the other hand, the cell size of the UCYN-A2 host in open ocean samples was 4-5 μm , clearly smaller than that described in coastal sites, about 7-10 μm (Thompson *et al.*, 2014; Hagino *et al.*, 2013). This is in agreement with a greater contribution of *B. bigelowii* reads in the 0.8-5 μm size fraction (as compared with the 5-20 μm fraction) in the *Tara*-Oceans metabarcoding
350 data analysed here, as well as of UCYN-A2 reads in the same smaller size fraction in metagenomic samples of the South Atlantic (Cornejo-Castillo *et al.*, submitted).

The hypothesis about the obligatory dependence of UCYN-A with its host (Tripp *et al.*, 2010; Thompson *et al.*, 2012; Bombar *et al.*, 2014) has been recently reinforced with the strong coupling observed in the transfer of carbon and nitrogen metabolites between partner cells
355 (Krupke *et al.*, 2014b). On the other hand, the possibility of a free-living population of the host could not be evaluated since the FISH probe used in previous studies (Krupke *et al.*, 2014a) targeted the whole prymnesiophyte assemblage and not the specific host phylotypes. Our use of a specific probe for the UCYN-A1 host allowed us to validate that this association

was also obligatory for the host since i) we never observed a free living population of targeted
360 prymnesiophytes in the absence of UCYN-A and ii) counts of free partner cells, when
detected, were correlated, suggesting a disruption of the association due to sample
manipulation (Thompson *et al.*, 2012; Krupke *et al.*, 2013). To our knowledge this is one of
the few cases showing an obligatory dependency of a phytoplankton species and its symbiont.
The high degree of specificity between partners and the reductive evolution of the UCYN-A
365 genomes (Bombar *et al.*, 2014) are features analogous to those observed in some freshwater
diatom species, which harbor N₂-fixing cyanobacteria as endosymbionts that are inseparable
from the host and transferred to daughter cells during host cell division (Nakayama *et al.*,
2014). The UCYN-A symbiont of *B. bigelowii* is also an endosymbiont as observed by
transmission electron microscopy (Hagino *et al.*, 2013), and it is likely that the
370 prymnesiophyte–UCYN-A1 association presents analogous structural properties.

In our study, about 97% of the hosts carried a single UCYN-A1 cell. This supports previous
observations from the North Pacific (Thompson *et al.*, 2012) and the North Atlantic (Krupke
et al., 2014a). Therefore we consider that the 1:1 UCYN-A1/host ratio is the most plausible
assumption for estimating N host requirements and metabolite exchanges between partner
375 cells and evaluating the contribution of this partnership to N₂ fixation. In addition, we suggest
that the presence of two UCYN-A1 cells per host may derive from repartition of symbionts
before host cell division. We made punctual captures of host cells likely displaying different
stages of cell division (Figure S5). It is known from field (Jacquet *et al.*, 1998) and laboratory
studies (Jacquet *et al.*, 2001) that photosynthetic picoeukaryotes divide during a very short
380 time window just before or during the dark period. We detected two cells per host mostly in
INDEMARES samples, which were collected after sunset and thus the symbiotic population
could be in active cell division. In the future, microscopic observations during diel cycles

coupled with transcriptomic analyses would shed light on the division mechanisms within this symbiosis.

385 *A widespread distribution of the symbiosis in the marine ecosystem*

The distribution obtained based on CARD-FISH counts, together with the 18S rDNA metabarcoding from the circumglobal expedition *Tara-Oceans*, expands the geographic distribution of the symbiosis and presents contrasted ecological patterns of both types. The UCYN-A1 host appeared in few spots of high abundance and was undetected in ~56% of
390 CARD-FISH surface samples. The distribution of this host based on V9 reads supported this patchy distribution. Although the metabarcoding method has a much lower detection limit than the CARD-FISH method, we could not detect the presence of UCYN-A1 host reads in a substantial number of stations (~62%). The UCYN-A2 host was detected only occasionally by
395 CARD-FISH, likely because its abundance was below the detection threshold, consistent with reported cell abundances of *B. bigelowii* of up to 1 cells ml⁻¹ in marine systems (Konno *et al.*, 2007). This limitation to obtain a reliable signal of the UCYN-A2 host was offset with the metabarcoding approach. Thus, the UCYN-A2 host showed an unexpected homogeneous distribution at low relative abundance in the open ocean, with targeted reads detected throughout *Tara-Oceans* samples.

400 Our CARD-FISH observations expanded the geographic coverage of UCYN-A1 in symbiosis to the South Atlantic gyre and the southern Indian Ocean. In these regions, no records of the prymnesiophyte phylotypes or UCYN-A were documented previously, except occasional detections of UCYN-A *nifH* sequences (no clade distinction) in the Benguela upwelling system, in the Arabian Sea and near Madagascar (Mazard *et al.*, 2004; Moisaner *et al.*, 2010;
405 Sohm *et al.*, 2011a). Moreover, our metabarcoding analysis represented the first detection of both hosts in the Red Sea, and widened the presence of the UCYN-A2 host in the South

Pacific, the Arabian Sea and the Mediterranean Sea, where only few *nifH* clones of UCYN-A2 were reported (Man-Aharonovich *et al.*, 2007). Previous reports showed that *B. bigelowii* usually occurred in low salinity, nutrient rich coastal waters (Konno *et al.*, 2007; Hagino *et al.*, 2013) although it was also reported in the Sargasso Sea (Gaarder, 1954; Hulburt, 1962). Our study reinforces the presence of *B. bigelowii* in symbiosis with UCYN-A in open ocean stations, consistent with the finding of *nifH* copies of the UCYN-A2 phylotype in the open sea (Thompson *et al.*, 2014).

In the ocean, cyanobacterial diazotrophs generally show a consistent pattern in depth with higher abundances in the upper euphotic zone (Goebel *et al.*, 2010; Moisander *et al.*, 2010). Recurring summer blooms of these diazotrophs are reported in the North Pacific subtropical gyre (Dore *et al.*, 2008) usually related to shallow mixing conditions that ensure the solar energy needed for nitrogen fixation. A recent study showed that the UCYN-A symbiosis appeared largely restricted to the upper water column, where N/P ratios were below 16 (Krupke *et al.*, 2014a). Our highly resolved vertical profiles show that i) the prymnesiophyte–UCYN-A1 symbiosis occupies nutrient depleted surface waters, and ii) its abundance decreases in the shallow DCM. This pattern suggests a dependence on light intensity, and/or an out-competition of the symbiotic population by other non-diazotrophic species that benefit from the increased nutrient availability at the nutricline. This vertical pattern is comparable to that observed for UCYN-A1 *nifH* gene abundances in station ALOHA (Church *et al.*, 2005, 2009). In these vertical profiles, between 10^4 - 10^6 *nifH* copies per L (roughly equivalent to 10-1 000 cells ml^{-1}) were quantified in the well-lit, nutrient depleted upper waters, and abundances decreased in the DCM to 10^2 - 10^3 *nifH* copies per L (~ 1 cell ml^{-1}).

Environmental controls of cyanobacterial diazotrophs in the ocean

In our study, no environmental factor explained unambiguously the prymnesiophyte–UCYN-

A1 distribution in surface waters. Presence or absence of the symbiosis occurred under similar environmental conditions (Figure S4). Previous reports point to temperature as a driver of the distribution of cyanobacterial diazotrophs (Church *et al.*, 2008; Moisander *et al.*, 2010).

Particularly for UCYN-A, *nifH* gene abundances have been reported to be higher in

435 temperature ranges from 19 to 24°C (Langlois *et al.*, 2008; Church *et al.*, 2008). Our study, with detections at a thermal range of 18-30°C, revealed a weak negative relationship between symbiosis abundance and temperature, agreeing with previous reports. Limiting nutrients such as iron or phosphate might also control the distribution of cyanobacterial diazotrophs (Moore *et al.*, 2009; Sohm *et al.*, 2011b). Indeed, higher abundances of diazotrophs have been linked
440 to oceanic areas where these nutrients are supplied from dust deposition of adjacent desert areas (Ridame and Guieu 2002; Tyrrell *et al.*, 2003; Capone *et al.*, 2005; Mahowald *et al.*, 2009). The patches of symbiosis observed in this study in the eastern North Atlantic, the western coast of Australia or even near Hawaii, could be explained by deposition events, since high dissolved Fe concentrations have been reported in these areas (Brown *et al.*, 2005;
445 Langlois *et al.*, 2012). Patchiness seems to be a common feature of diazotrophic populations (Church *et al.*, 2009; Goebel *et al.*, 2010) and UCYN-A is regarded as very dynamic over small spatio-temporal scales (Robidart *et al.*, 2014).

The regional dominance of different diazotrophic groups can be related to distinct temperature and nutrient requirements (Church *et al.*, 2008; Langlois *et al.*, 2008). For instance, in the
450 tropical North Atlantic, *Trichodesmium* was dominant in the warmer western side whereas UCYN-A was more abundant in the cooler eastern side near Cape Verde Islands (Goebel *et al.*, 2010). During the MALASPINA expedition, a similar picture for the distribution of UCYN-A and *Trichodesmium* within this basin was observed, with these two diazotrophic groups dominating in different regions (see Fernández-Castro *et al.* in press for a

455 *Trichodesmium* distribution map). In addition, N₂ fixation rates measured during the

MALASPINA expedition did not correlate with *Trichodesmium* abundances (Fernández-Castro *et al.*, in press), but we noticed that regions of enhanced N₂ fixation rates as in the western South Atlantic or the eastern Indian ocean were coupled to the presence of the prymnesiophyte–UCYN-A association. A recent latitudinal study along the eastern Australian coast also described a community shift from *Trichodesmium* in the north to unicellular diazotrophs in the south where the highest N₂ fixation rates were reported (Raes *et al.*, 2014).

Different ecological strategies between the two related lineages

Our results obtained by FISH counts and metabarcoding sequences in a large set of marine samples demonstrated that the two types of prymnesiophyte–UCYN-A symbiosis were widespread in the photic layer of the world ocean. We observed that UCYN-A2 host was homogeneously distributed at both surface and DCM but in low abundance, whereas UCYN-A1 host was only detected in some hotspots of very high abundance at surface, suggesting different ecological strategies. We hypothesize that UCYN-A1 host could follow the r strategy, with fast growth under favourable conditions, such as unpredictable events of dust deposition or the onset of thermal stratification, using surface light for the energy-demanding process of nitrogen fixation. On the other hand, the UCYN-A2 host could follow the k strategy (also known as stress-tolerant species by Reynolds, 1997), persisting at low abundances and being more competitive in stable and low resource conditions. In this sense, the small UCYN-A1 host seems excluded in the DCM likely because other species are more efficient in waters with higher nutrients and dimmer light, while this depth limitation could be less severe in the UCYN-A2 host, which seems to be less affected by changes in the light regime (Thompson *et al.*, 2014). A better understanding of the role of UCYN-A diazotrophy along the vertical profile may certainly benefit from transcriptional profiles of these populations, including the mechanisms of the host to regulate UCYN-A activity. Overall,

480 further studies on the biogeochemical role of the UCYN-A diazotrophy may need to include
contrasted ecological differences among the different lineages.

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Conflict of Interest

The authors declare no conflict of interest.

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Supplementary information is available at the ISME Journal website.

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640 **Titles and legends to figures**

Figure 1 Epifluorescence microscopy images of the double CARD-FISH assay. Left panels correspond to DAPI signal (blue-labeled nucleous). Right panels correspond to the combined signal of the prymnesiophyte specific probes (green-labeled host under blue light excitation) and the cyanobacteria (red-labeled symbiont under green light excitation). **(a)** Two
645 prymnesiophyte host cells carrying the symbiont close to two non-host prymnesiophyte cells from MALASPINA station 64. **(b)** A prymnesiophyte host harboring two UCYN-A cells from surface of station A (INDEMARES cruise) where this type of association was frequently observed. **(c)** A *B. bigelowii* cell targeted by the UBRADO69 probe from MALASPINA station 68.

650 **Figure 2** Relationship between cell counts categories in samples where the prymnesiophyte-UCYN-A1 symbiosis was detected. **(a)** Abundances of non-associated hosts versus free symbionts. **(b)** Abundances of symbiosis versus free partner cells.

Figure 3 Global distribution of prymnesiophyte hosts in surface waters. **(a)** Cell abundances of the prymnesiophyte–UCYN-A1 symbiosis along the MALASPINA and INDEMARES
655 cruises determined by double CARD-FISH using UPRYM69 and UCYN-A732 probes. **(b)** Relative abundances of V9 reads of UCYN-A1 host (upper panel) and UCYN-A2 host (lower panel) to total prymnesiophytes along the *Tara*-Oceans expedition.

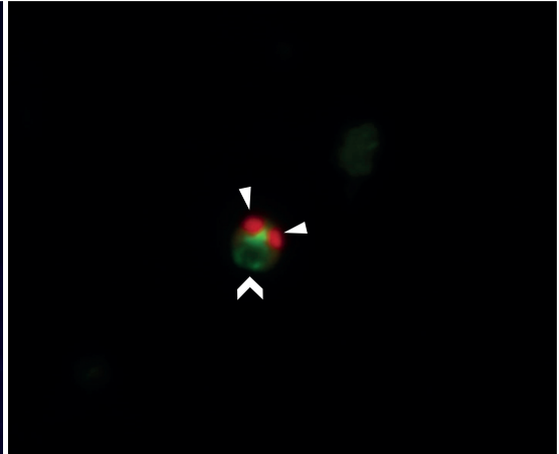
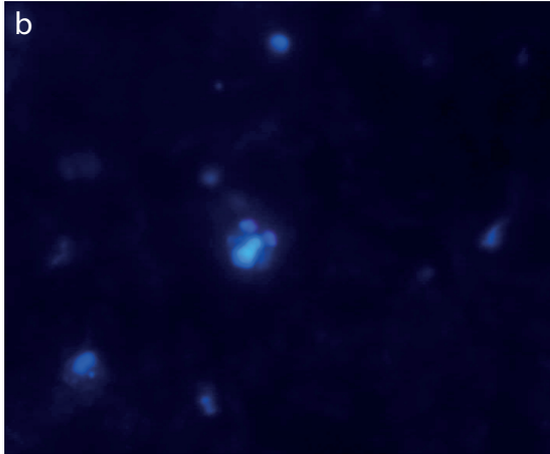
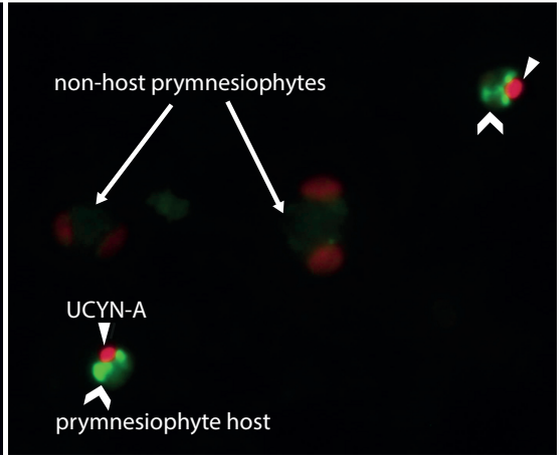
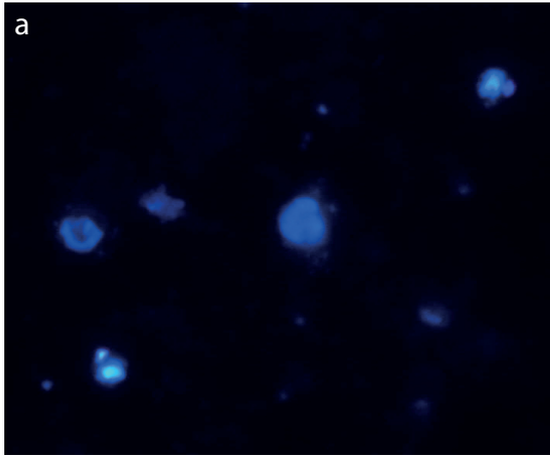
Figure 4 Cell counts of the prymnesiophyte–UCYN-A1 symbiosis versus **(a)** total chlorophyll, **(b)** temperature, **(c)** nitrate, **(d)** phosphate, **(e)** N/P ratio, **(f)** DOP and **(g)** DON
660 measured in surface waters during the MALASPINA (white dots) and INDEMARES (black dots) cruises. Regression lines are plotted in cases displaying a significant relationship ($P < 0.05$).

Figure 5 Contribution of prymnesiophytes to total community reads (**a**) and contribution of UCYN-A1 host (**b**) and UCYN-A2 host (**c**) to total prymnesiophyte reads at surface and
665 DCM depths of *Tara*-Oceans stations (V9 rDNA metabarcoding data). DCM depth was plotted in panel (**c**). Only stations having both surface and DCM samples are shown (29 out of 40). In the 11 stations not shown here, the percentage of prymnesiophytes to total community reads at surface was similar ($4.6\% \pm 4.3$ on average).

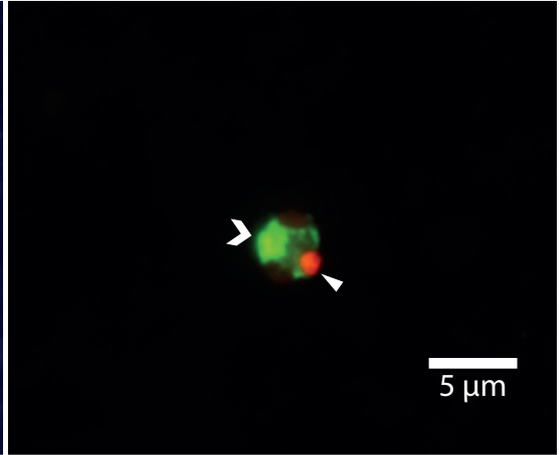
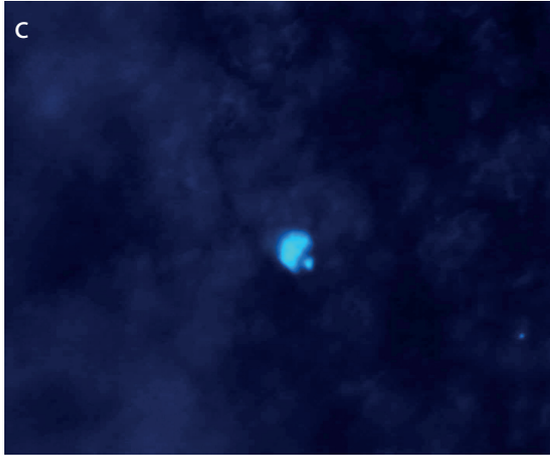
Figure 6 Depth profiles of hydrological variables (right) and distribution in depth of the prymnesiophyte–UCYN-A1 symbiosis (black dots) and the rest of the prymnesiophyte
670 community (white dots) (left) for the INDEMARES stations in the NE Atlantic. Stations A and B correspond to the Avilés canyon region; stations C and D correspond to the Galician Bank region. The grey shaded area represents the fluorescence profile recorded at each station. Error bars represent standard errors.

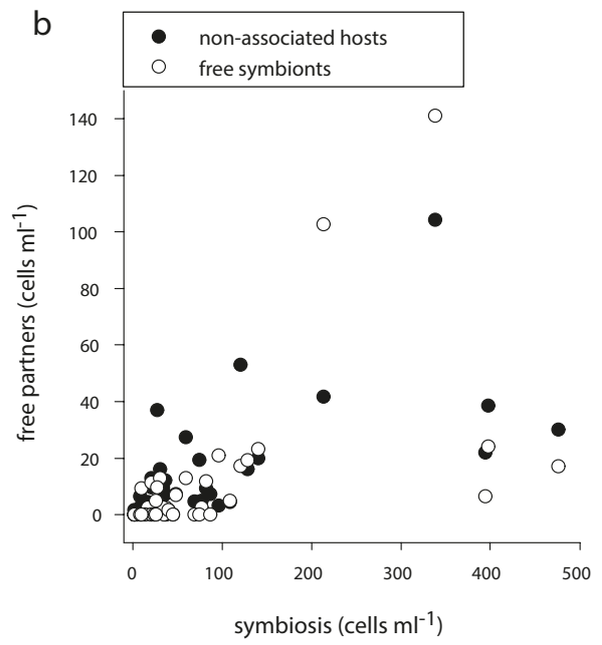
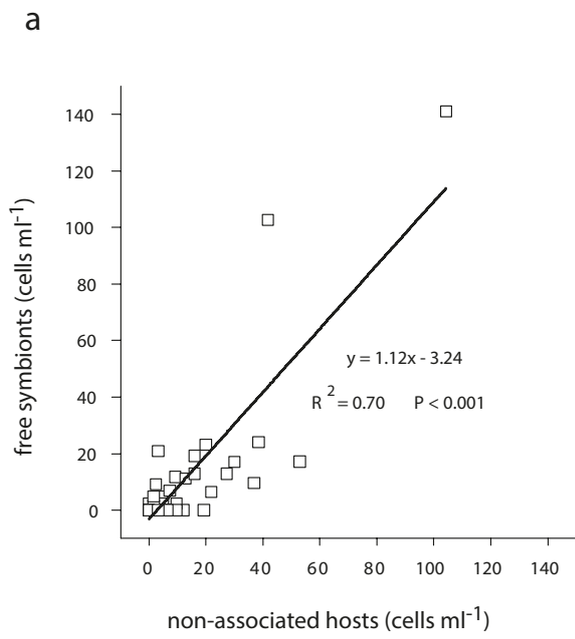
675

UPRYM69 + UCYN-A732

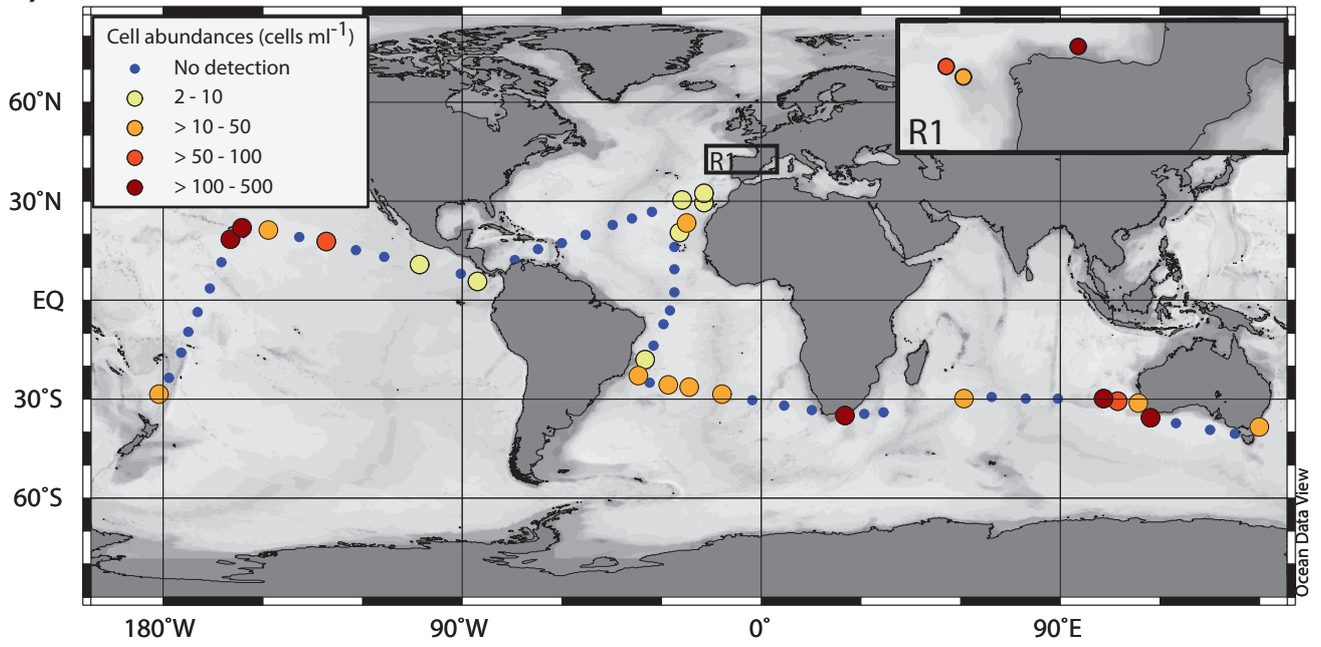


UBRADO69 + UCYN-A732

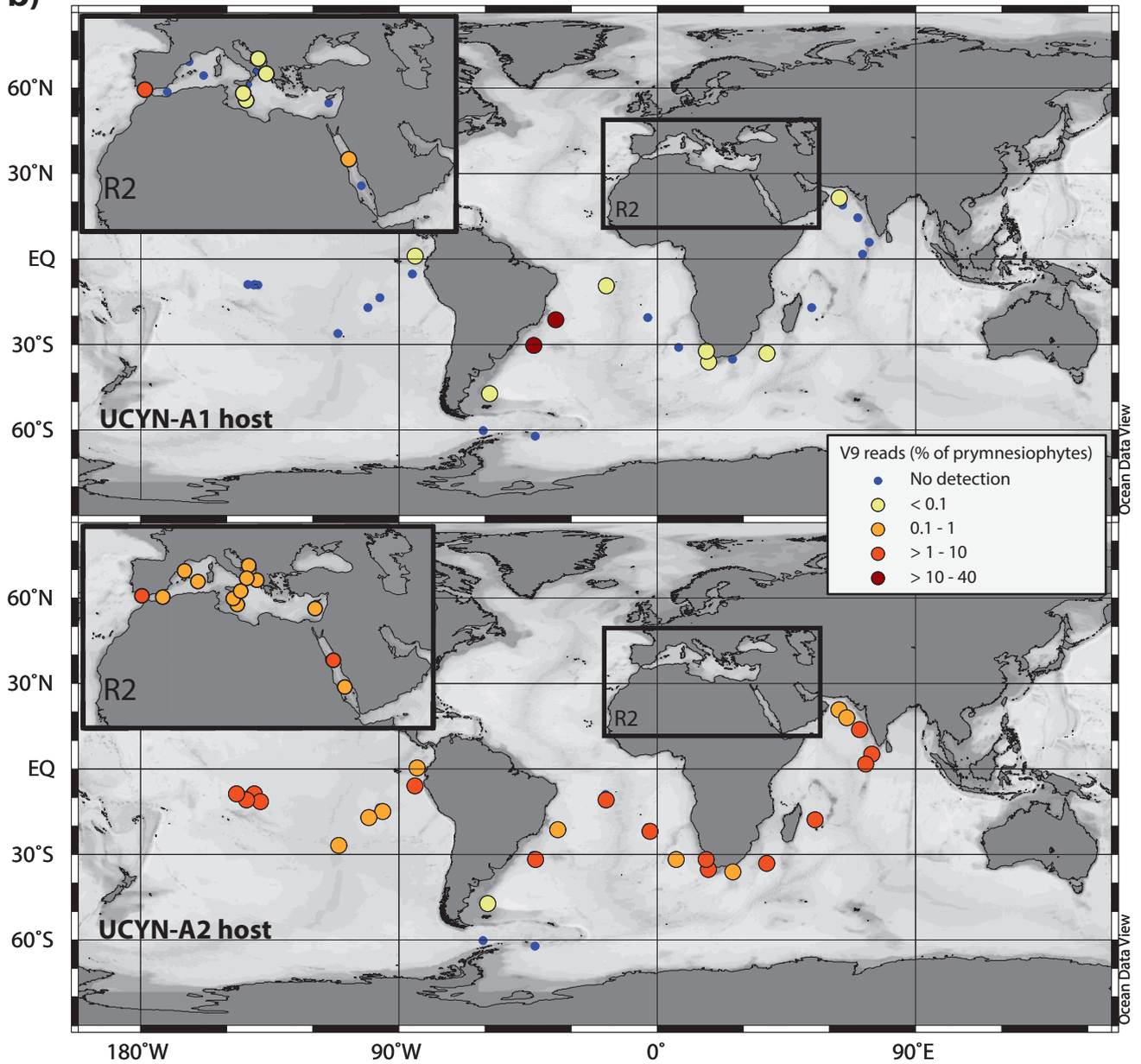


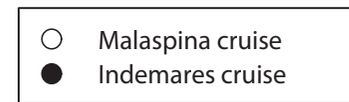
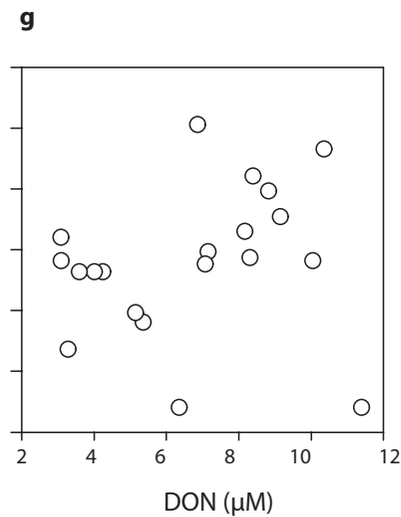
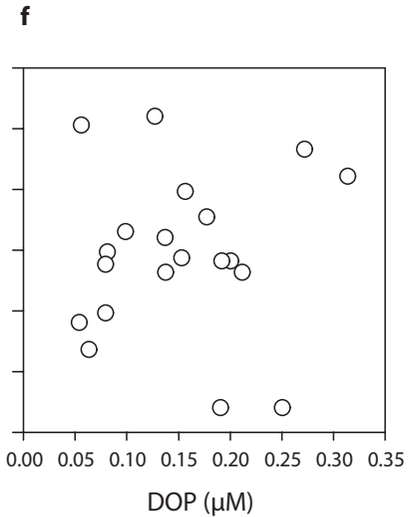
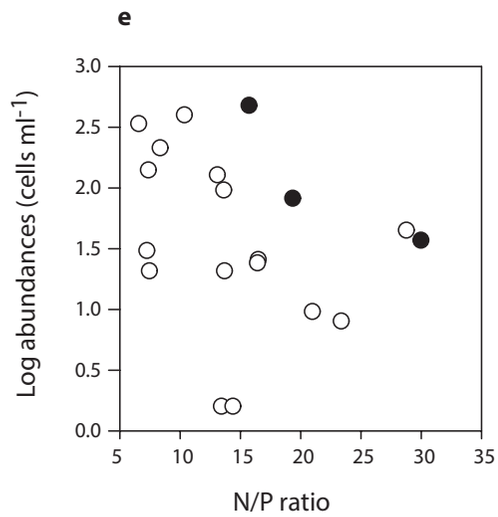
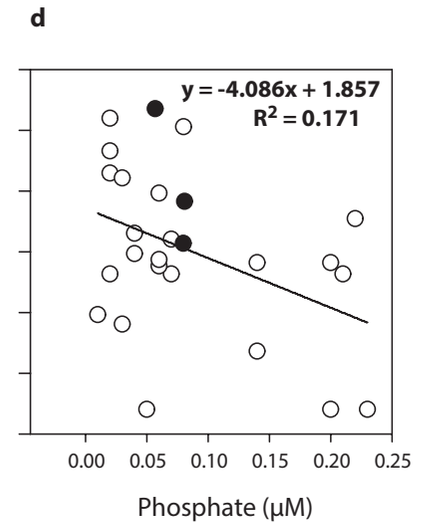
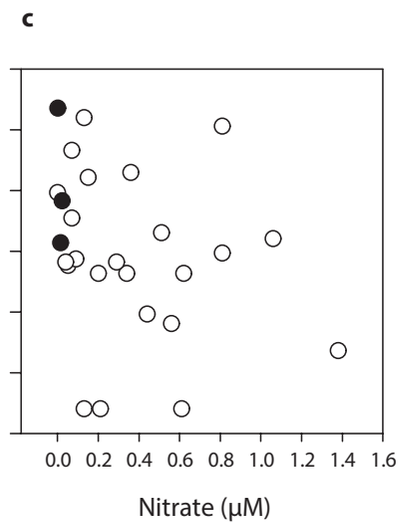
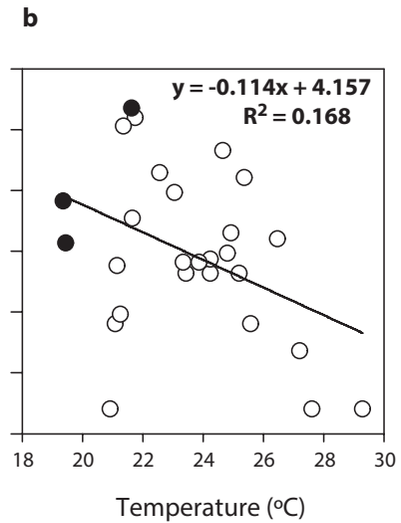
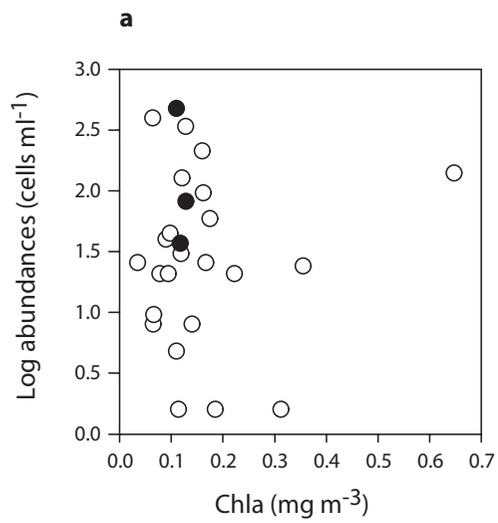


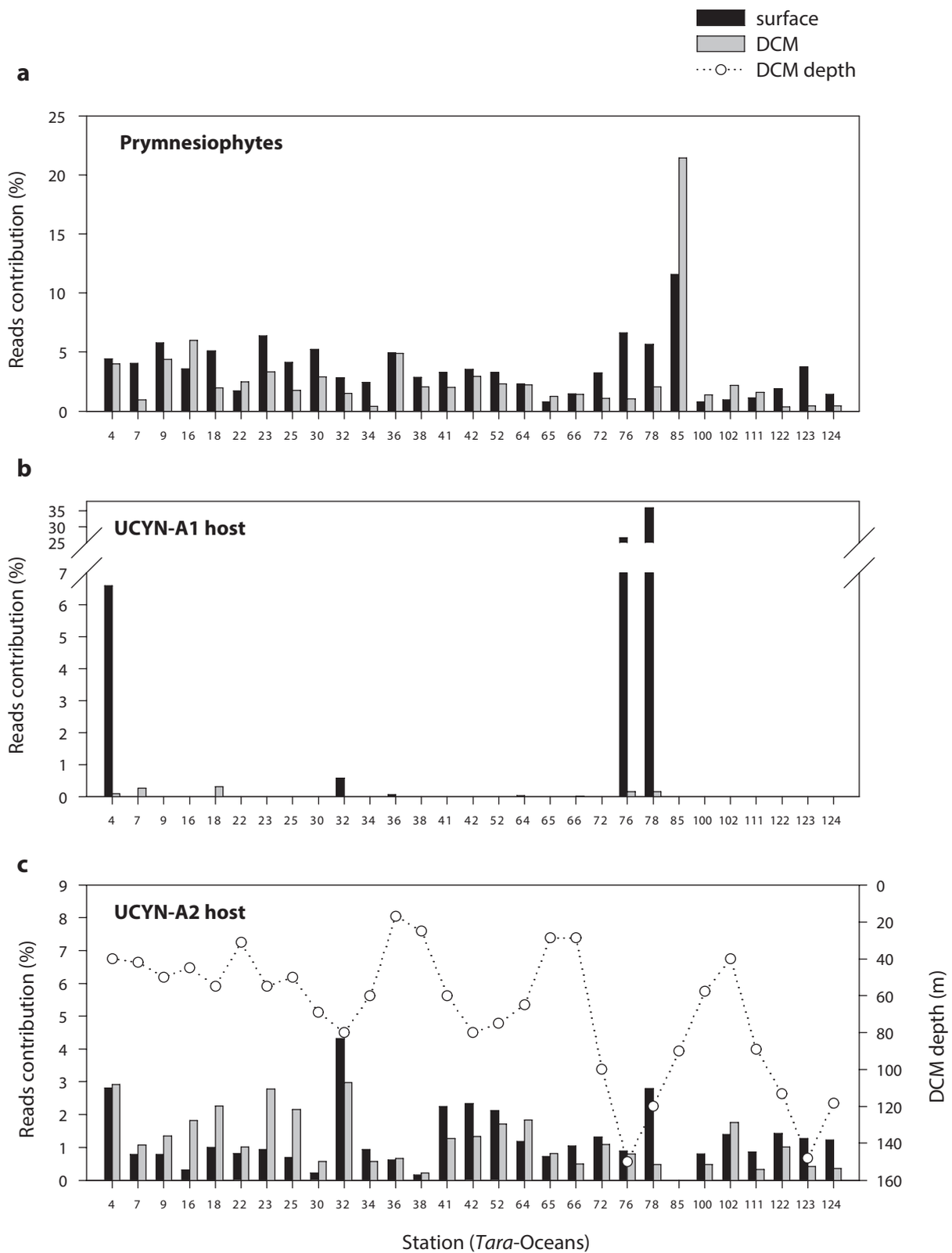
a)



b)







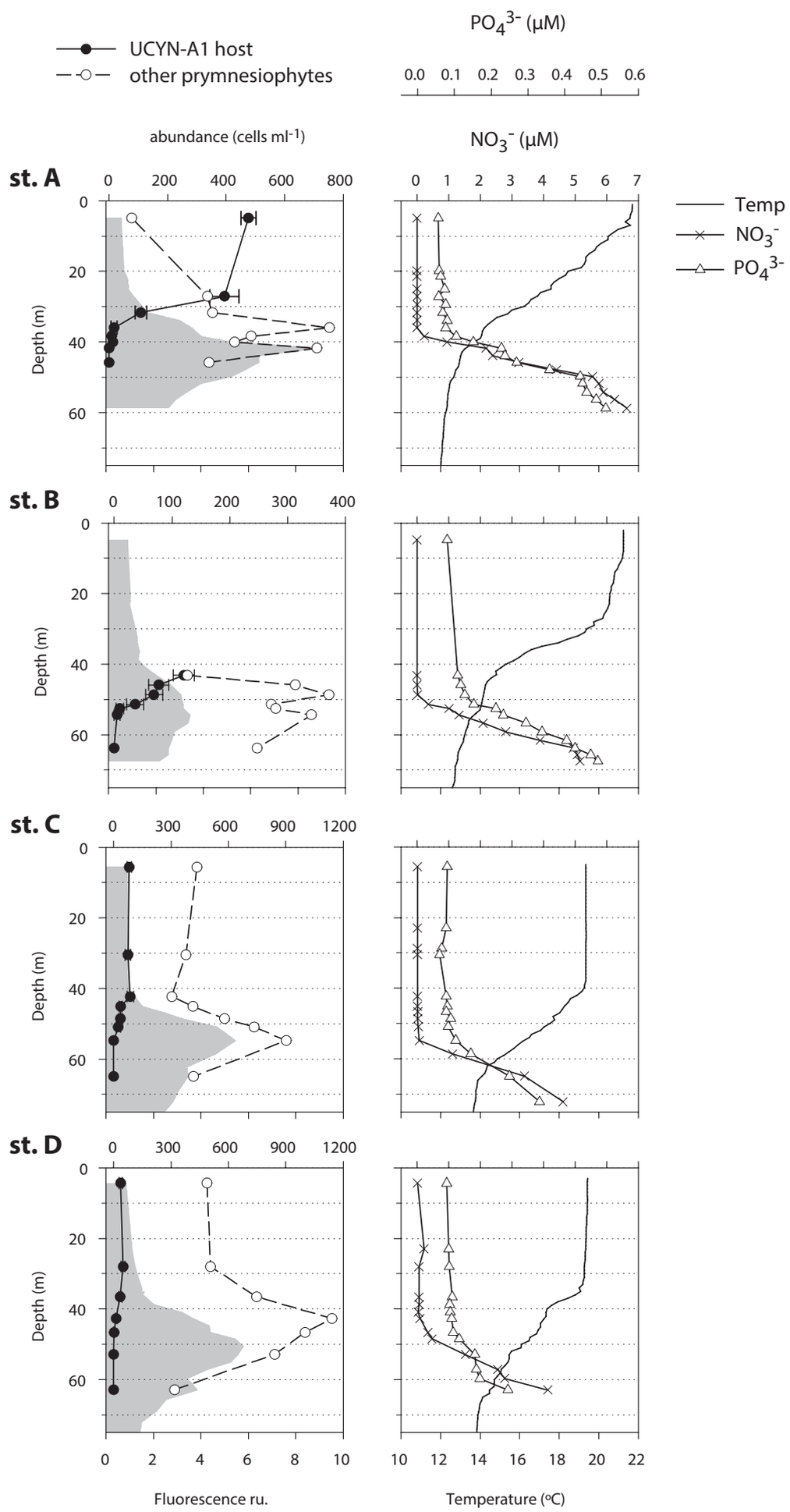


Table 1: Averaged abundances of symbiosis inside and outside of a given range of values for temperature, phosphate, nitrate, N/P ratio and chlorophyll *a*. Thresholds for temperature and inorganic nutrients were around the median value in the 62 MALASPINA and INDEMARES surface samples in order to obtain an equal number of samples in each category. Chlorophyll *a* ranges were established according to the trophic water classification of Shushkina *et al.*, 1997. (*) indicates when abundances values were significantly different ($P < 0.05$; Mann-Whitney test) inside and outside the corresponding range.

Variable	Range		<i>abundances (cells ml⁻¹)</i>				<i>number of samples</i>	
			<i>in range</i>		<i>out range</i>		<i>in range</i>	<i>out range</i>
			Average	SD	Average	SD	n	n
Temperature	< 25°C	(*)	21	120	7	25	33	29
PO ₄ ⁻³	< 0.08 μM	(*)	61	118	6	14	30	26
NO ₃ ⁻	< 0.27 μM		57	119	24	67	28	29
N/P ratio	< 16		72	135	13	22	26	18
TChla ⁽¹⁾	< 0.06 mg m ⁻³	(*)	3	17	43	18	9	52
TChla ⁽²⁾	0.06-0.1 mg m ⁻³		64	20	27	18	16	45
TChla ⁽³⁾	0.1-0.3 mg m ⁻³		35	19	39	18	30	31
TChla ⁽⁴⁾	> 0.3 mg m ⁻³		28	11	38	19	6	55

(¹) *Ultraoligotrophic*, (²) *Oligotrophic*, (³) *Mesotrophic*, (⁴) *Eutrophic*