# Massive export of diazotrophs across the tropical south Pacific Ocean

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#### **Abstract**

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Diazotrophs are widespread microorganisms that regulate marine productivity in 60% of our oceans by alleviating nitrogen limitation. Yet, their contribution to organic carbon and nitrogen export fluxes has never been quantified, making an assessment of their impact on the biological carbon pump impossible. Here, we examine species-specific fates of several groups of globally-distributed unicellular (UCYN) and filamentous diazotrophs in the mesopelagic ocean. We used an innovative approach consisting of the combined deployment of surface-tethered drifting sediment traps, Marine Snow Catcher, and Bottlenet, in which we performed nifH sequencing and quantitative PCR on major diazotroph groups across the subtropical South Pacific Ocean. nifH sequencing data from sediment traps deployed at 170 m, 270 m and 1000 m provide clear evidence that cyanobacterial and non-cyanobacterial diazotrophs are systematically present in sinking particles down to 1000 m, with export fluxes being the highest for the UCYN-A1 symbiosis, followed by UCYN-B or Trichodesmium (depending on station and depth), Gamma A and UCYN-C. Specific export turnover rates (a metric similar to the export efficiency adapted to organisms) point to a more efficient export of UCYN groups relative to the filamentous Trichodesmium. This is further confirmed by Marine Snow catcher data showing that the proportion of sinking cells was significantly higher for UCYN compared to Trichodesmium. Phycoerythrin-containing UCYN-B and UCYN-C-like cells were indeed recurrently found embedded in large (> 50 µm) seemingly organic aggregates, or organized into clusters of tens to hundreds of cells linked by an extracellular matrix, facilitating the export. Overall, diazotrophs accounted for 6-13% (170 m) to 45-100% (1000 m) of the total particulate nitrogen export fluxes in our study. We thus conclude that diazotrophs are important contributors to carbon sequestration in the subtropical South Pacific Ocean and need to be considered in future studies to improve the accuracy of current regional and global estimates of export.

# Introduction

Nitrogen (N) availability limits primary productivity throughout much of the surface low-latitude ocean<sup>1</sup>. In such nitrogen (N)-limited waters, microbial dinitrogen (N<sub>2</sub>) fixation by diazotrophic plankton provides the major source of new N to surface waters<sup>2</sup> that on appropriate timescales, equates export production<sup>3</sup>. However, the fate of this production, whether it is exported to the deep ocean or stimulates remineralization in surface waters remains obscure<sup>4</sup>. An increasing number of studies have shown that diazotroph-derived N is quickly translocated to non-diazotrophic plankton such as diatoms<sup>5, 6</sup>, which eventually contribute to secondary export of organic matter out of the photic zone. Yet, the direct gravitational settling of diazotrophs themselves to the deep ocean has rarely been assessed. The few existing observations (see below) conclude that diazotrophs provide a minor contribution to the total amount of N in the exported material.

Diazotrophs may associate with sinking particles to directly contribute to export by different mechanisms. The two most direct include gravitational settling of individual cells/filaments or aggregates. According to Stokes' Law, particle sinking velocity scales with the square of particle size. Therefore, large particles should sink faster and are more likely to reach the deep ocean before being remineralized by bacteria<sup>7</sup>. Aggregation is thus a crucial step for the transport of small phytoplankton species which could export particulate organic carbon (POC) and N (PON) in similar proportion to their production in surface waters8. Diazotrophs have diverse morphologies and their size spans several orders of magnitude. Some types such as the free-living unicellular diazotrophic cyanobacteria (UCYN from groups B and C) are small (2-8 µm in diameter), while others such as Trichodesmium sp. are filamentous and can form large-sized colonies (>100-1000 µm). In addition, some diazotrophs live in symbiosis with calcified (UCYN-A, ~1 µm) or silicified eukaryotes (Richelia sp., Calothrix sp., >20 µm, forming Diatom-Diazotrophic Associations or DDAs). These dense biominerals may provide ballast enhancing the downward export of these symbioses into the deep ocean. Therefore, the presence of different diazotrophs in surface waters may result in drastically different C and N export fluxes via the biological pump. Yet, no field observations have related individual diazotroph groups to the magnitude of downward particles fluxes to date.

Thanks to their inherently ballasted character, DDAs are well known to contribute to particulate matter export<sup>9, 10</sup>, and are involved in seasonal peaks of POC export to the deep sea (4000 m) in the North Pacific subtropical gyre<sup>11</sup>. *Trichodesmium* is one of the major contributors to global N<sub>2</sub> fixation<sup>12</sup>. However, it is thought to have a limited export capacity and to be preferentially remineralized in the

surface layers due to the presence of gas vesicles potentially providing buoyancy<sup>4, 13, 14</sup>. Yet, some studies have reported the presence of *Trichodesmium* in sediment traps material in the Kuroshio Current<sup>15</sup>, the tropical North Pacific<sup>16</sup> and the South Pacific Oceans<sup>17</sup>. Intact filaments and colonies of *Trichodesmium* sp. have also been reported as deep as 3000-4000 m in the tropical Atlantic, Pacific and Indian Oceans<sup>18, 19</sup>, but their contribution to organic matter export has to be quantified.

A priori, UCYN may not contribute importantly to POC export fluxes either due to their small size. Yet, Berthelot et al.  $^{20}$  reported in experimental mesocosms that primary production supported by UCYN was twice as more efficient in promoting POC export than production supported by DDA. In the same experiment, Bonnet et al.  $^{21}$  showed that the sinking of UCYN-B and C was driven by the aggregation of small individual cells (5-6  $\mu$ m) into larger aggregates (100 to >500  $\mu$ m) that were eventually exported and accounted for up to ~20% of the POC flux. Recently, Caffin et al.  $^{17}$  confirmed the presence of UCYN-B in sinking particles collected with sediment traps (330 m) throughout the tropical South Pacific Ocean ( $^{10^3}$  to  $^{10^7}$  nifH gene copies L- $^{1}$  of trap material). Finally, Farnelid et al.  $^{22}$  observed sequenced nifH genes of exported material ( $^{150}$  m) in the North Pacific subtropical gyre and found that all the diazotroph groups above as well as non-cyanobacterial diazotrophs were present in the exported material.

Taken together, these studies suggest that all diazotroph groups, small or large, free-living or symbiotic, ballasted or not, are present below the photic layer. However, a detailed examination relating types of diazotrophs to the magnitude of downward particles fluxes and their contribution is needed to refine our understanding of their role in the magnitude and mechanisms controlling the biological pump. This is all the more topical as climate models predict an expansion of the oligotrophic gyres<sup>23, 24</sup>, where diazotrophs represent a major component of the plankton biomass and sustain most new primary production.

Here, we examine species-specific fates of diazotrophs in the mesopelagic ocean. We used an innovative approach consisting of the combined deployment of surface-tethered drifting sediment traps, Marine Snow Catcher (MSC), and Bottle-net, in which we performed *nifH* sequencing and quantitative PCR on major diazotroph groups across the subtropical South Pacific (Fig. 1a). This study builds up on prior investigations that have characterized this region as a hotspot for marine  $N_2$  fixation  $^{25}$ . We show that all globally-significant  $N_2$ -fixing cyanobacteria and non-cyanobacterial diazotrophs are systematically present in sinking particles down to 1000 m. Small size UCYN (1-8  $\mu m$ ) are exported more efficiently than large filamentous diazotrophs as large (>50  $\mu m$ ) aggregates, but both are major contribution to PON export fluxes. We thus conclude that diazotrophs are important contributors to carbon sequestration in the ocean.

# **Results and discussion**

## **Surface conditions**

The study region was characterized by typical oligotrophic waters (chlorophyll concentrations <0.15  $\mu$ g L<sup>-1</sup>, DCM 100-180 m) east and west of Tonga, corresponding to the first group of stations S01, S02, S07 and S08 (Fig. 1a). The second group of stations (S03, S04, S05, S10, S11) was located in mesotrophic waters (chlorophyll >0.15  $\mu$ g L<sup>-1</sup>, DCM 70-90 m) in the vicinity of the Tonga volcanic arc (Fig. 1a). Surface (0-50 m) nitrate concentrations were consistently close or below the detection limit (0.05  $\mu$ mol L<sup>-1</sup>) throughout the transect, while phosphate concentrations were typically 0.1  $\mu$ mol L<sup>-1</sup> at oligotrophic stations and depleted down to detection limit (0.05  $\mu$ mol L<sup>-1</sup>) at mesotrophic stations, likely due to consumption by higher plankton stocks (Fig. S1). Seawater temperature ranged from 23.1 to 27.3 °C in the mixed layer (0-15 m to 0-60 m) determined according to de Boyer Montégut et al. (2004)<sup>26</sup> (Fig. S1).

The abundances of key diazotroph groups indicated that UCYN-A1 were present at high abundances throughout the photic layer (0-130 m) of the transect (average 3.2 x  $10^8$  nifH gene copies L<sup>-1</sup>) (Fig. 1b), peaking at oligotrophic stations (average 1.1 x  $10^9$  nifH gene copies L<sup>-1</sup>). Trichodesmium was the second most abundant group, particularly in the vicinity of Tonga (average 1.3 x  $10^7$  nifH gene copies L<sup>-1</sup>), followed by UCYN-B (5.7 x  $10^5$  nifH gene copies L<sup>-1</sup>). UCYN-C and Gamma-A (a non-cyanobacterial diazotroph) were also detected throughout the transect albeit at lower abundances ( $10^2$  to  $10^4$  nifH gene copies L<sup>-1</sup>).

# How efficient are different diazotrophs groups to export?

We first examined the *nifH* gene community composition of particles collected in sediment traps moored at 170 m, 270 m and 1000 m at stations S05M and S10M (Fig. 1). 32% of the retrieved Amplicon Sequence Variants (ASV) corresponded to cyanobacteria genera on average over both stations (Fig. 2). The most abundant genus was *Trichodesmium* sp., but ASVs related to *Crocosphaera watsonii* (UCYN-B), *Candidatus Atelocyanobacterium thalassa* (UCYN-A) and *Katagnymene* spp. were also

identified (Fig. 2). Genera related to DDAs (*Richelia* spp., *Calothrix* spp.) contributed little to the percentage reads. Besides autotrophic diazotrophs, 68% of ASVs were affiliated to non-cyanobacterial diazotrophs including Alpha-, Beta-, and Gammaproteobacteria at both stations.

Amplicon sequencing provides relative abundances, which are not necessarily indicative of the abundance of the target microorganism in the natural environment<sup>27, 28</sup>. Hence, to assess the export capacity of individual diazotroph groups, we quantified the abundance of five groups spanning different forms, sizes, lifestyle (symbiotic or not), using qPCR assays (UCYN-A1 symbiosis, UCYN-B, UCYN-C, *Trichodesmium* and Gamma-A; see Methods). DDAs were not quantified as they were almost not detected on microscopic filters, pointing to their rarity in those waters at the time of the cruise. All five groups were detected at abundances ranging from  $10^3$  to  $10^8$  *nifH* gene copies L<sup>-1</sup> of trap material, the highest being reported for UCYN-A1 and the lowest for UCYN-C (Fig. 3a). The diazotrophs assemblage exported in sediment traps generally reflected that of the photic layer, although some diazotrophs seem to sink more efficiently than others: the highest diazotroph export fluxes were measured for UCYN-A1 at both sites and all depths of traps deployment  $(9.5 \pm 1.9 \times 10^8$  to  $1.3 \pm 1.0 \times 10^9$  *nifH* gene copies m<sup>-2</sup>d<sup>-1</sup>), followed either by UCYN-B  $(1.3 \pm 0.6 \times 10^7$  to  $4.7 \pm 1.2 \times 10^7$  *nifH* gene copies m<sup>-2</sup>d<sup>-1</sup>) or *Trichodesmium*  $(1.6 \pm 1.2 \times 10^6$  to  $7.6 \pm 1.7 \times 10^8$  *nifH* gene copies m<sup>-2</sup>) depending on station and depth (Fig. 3). Gamma-A and UCYN-C were also exported, albeit at lower rates (~10<sup>5</sup>-10<sup>6</sup> and ~10<sup>5</sup> *nifH* gene copies m<sup>-2</sup>d<sup>-1</sup>), respectively).

Specific export turnover rates provide information on the rate of diazotrophs "lost" from the photic layer per day due to export. This metric is similar to the export efficiency (e-ratio=POC export/POC produced by Primary Production), but adapted to organisms. The range of calculated rates  $(0.8 \pm 0.6 \times 10^{-5} \text{ to } 7.0 \pm 1.2 \times 10^{-3} \text{ d}^{-1})$  spans through two orders of magnitude and vary depending on the diazotroph group, depth and station considered. On average (over all stations and depths), the export turnover rate of UCYN  $(3.5 \pm 1.9 \times 10^{-3} \text{ d}^{-1})$  was ca. four times higher than that of *Trichodesmium*  $(0.8 \pm 1.3 \times 10^{-3} \text{ d}^{-1})$  (Fig. 3c), pointing to preferential export of UCYN groups relative to the filamentous *Trichodesmium*. Among UCYN groups, the highest export turnover rate was measured for UCYN-B, followed by UCYN-C and UCYN-A1, although differences were not significant between groups (Mann-Whitney test, p<0.05). The export turn-over rate of Gamma-A (average  $1.1 \pm 0.9 \times 10^{-3} \text{ d}^{-1}$ ) was intermediate between that of *Trichodesmium* and UCYN groups.

Epifluorescence microscopy confirmed the presence of phycoerythrin-containing UCYN-B and UCYN-C-like cells in sediment trap samples (Fig. 4a-f). Scanning electron microscopy revealed that they were recurrently found embedded in large seemingly organic aggregates, or organized into clusters of tens to hundreds of cells linked by an extracellular matrix (Fig. 4g-l), which was further confirmed by alcian blue straining (Fig. S2). *Trichodesmium* was also observed in all samples, mostly as free filaments, but intact colonies were observed in sediment trap samples especially at 1000 m at both stations. Gamma-A and UCYN-A1 cannot be visualized by these techniques and are thus assessed solely on the basis of gPCR counts (above).

Pigment concentrations measured in sediment trap samples at both stations indicate that total pigment concentrations and diversity decreased with depth. Apart from the degradation pigments phaeophorbide and phaeophytin, zeaxanthin, a biomarker of cyanobacteria predominated at all depths (Fig. S2). The Chlorophyll *a*: pheopigments ratios were elevated (average 1.8, range 0.8-3.), indicative of fresh organic matter in sediment trap samples.

## Contribution of diazotrophs to PN export fluxes

Sediment trap-based export fluxes of PN at 170 m, 270 m and 1000 m ranged between 1.6  $\pm$  0.2 and 6.1  $\pm$  1.2 mg N m<sup>-2</sup> d<sup>-1</sup> at S05M and 5.2  $\pm$  0.1 and 13.4  $\pm$  1.3 mg N m<sup>-2</sup> d<sup>-1</sup> at S10M (Table 1). Fluxes attenuated with depth, with transfer efficiencies of 0.02 and 0.03 between 170 m and 1000 m at both stations. The N export efficiency (the amount of PN export at the base of the euphotic zone over the integrated N<sub>2</sub> fixation rate) was 18% and 53% at stations S05M and S10M. The C:N ratios in exported material was lower (5.2) than that of the surface (7.5, integrated value over the photic layer) at station S10M, pointing to a preferential export of N relative to C.

The proportion of PN attributed to diazotrophs and collected by sediment traps was evaluated based on qPCR results and estimated PN content per diazotroph (see Methods). The PN flux attributed to diazotrophs ranged from 0.6-12.3 mg N m<sup>-2</sup> d<sup>-1</sup> at S05M and 2.3-7.8 mg N m<sup>-2</sup> d<sup>-1</sup> at S10M, accounting for 13% (170 m) to 100% (1000 m) and 6% (170 m) to 45% (1000 m) of the total PN export flux, respectively at stations S05M and S10M (Table 1). *Trichodesmium* was the major contributor of the PN attributed to diazotrophs at 1000 m at both stations, whereas UCYN dominated at shallower depths (Table S1). Despite non-cyanobacterial diazotrophs were present in exported material (68% of sequences), their C and N composition remains unknown due to the poor availability of cultured representatives. Thus, quantifying their contribution to export fluxes is not possible at this stage. In

addition, sequencing data also revealed the presence of Katagnymene spp., which are large (>100-1000  $\mu$ m) and likely have high C and N content. Hence the contribution of diazotrophs to exported PN reported here represent a lower-end estimate.

# The fate of diazotrophs in the mesopelagic ocean

We further assessed the fate of the exported diazotroph community by collecting fresh particles in mesopelagic waters using the Marine Snow Catcher (MSC, see Methods). MSC samples were collected at the same depths as those of sediment traps (170, 270, 1000 m) at stations S05M and S10M and in addition at seven other stations (Fig. 1). In essence, the MSC allows the separating non-sinking particles from slow sinking and fast sinking particles<sup>29</sup> (see Methods). These different fractions can hence be studied individually.

The *nifH* composition in the various fractions generally mirrored that of traps at stations S05M and S10M with 32-54% of sequences affiliated with cyanobacteria and a prevalence of *Trichodesmium* (Fig. 5). This was also the case at stations S03, S04, S07 and S11, but not at stations S01, S02 and S08, where sequences assigned to *Candidatus Atelocyanobacterium thalassa* and *Crocosphaera watsonii* were generally more abundant than *Trichodesmium* sequences. As in sediment traps, a large number of *nifH* sequences (46-58%) were affiliated to non-cyanobacterial diazotrophs, which composition was overall consistent with that of the traps, although some classes not detected in sediment traps (Epsilonproteobacteria and Clostridia) were present in all MSC fractions. In general, the total number of reads was consistently 10-25% higher in the slow and fast sinking fractions compared to the suspended fraction at all stations. This trend was mainly driven by non-cyanobacterial sequences that are suspected to be attached to particles<sup>30</sup>, and to a lesser extent by sequences affiliated with cyanobacteria.

As in sediment traps, five diazotroph groups were quantified in the suspended, slow and fast sinking fractions to more finely assess their sinking dynamics over short time scales (h). Overall, UCYN-A1 and *Trichodesmium* were the most abundant groups in the mesopelagic MSC samples (170 m to 1000 m), and their numbers varied in parallel with the abundance of their population in the photic layer (Fig. 6). UCYN-A1 were the most abundant at oligotrophic stations (S01, S02, S07, S08, chlorophyll <0.15  $\mu$ g L<sup>-1</sup>), while *Trichodesmium* peaked at mesotrophic stations (Chlorophyll a >0.15  $\mu$ g L<sup>-1</sup>, Fig. 1). Conversely, the abundance of UCYN-B, UCYN-C and Gamma-A in mesopelagic waters was poorly correlated with their photic layer abundance.

The MSC suspended fraction accounted for the most abundant pool of diazotrophs at all stations, followed by either the slow or the fast sinking pools, depending on groups (Fig. 6a-e). At the mooring stations S05M and S10M, the total concentration of diazotrophs and the diazotroph community composition remained generally constant at the three sampled depths (170 m, 270 m and 1000 m), consistent with the results observed in sediment traps. To evaluate the export potential of each individual diazotroph group, we calculated the ratio of sinking versus non-sinking cells (the sum of the slow and fast sinking fractions over the total). The proportion of sinking cells varied widely across the transect, but was the highest for UCYN-A1 (32 ± 33 %), UCYN-C (31 ± 38 %), Gamma-A (29 ± 19 %) and UCYN-B (27 ± 11 %), and the lowest for *Trichodesmium* (18 ± 20 %) (Fig. 6f). Among the sinking fractions, UCYN-B and Gamma-A were generally equally distributed among the slow and fast sinking fractions, whereas the majority of nifH gene copies were found in the fast sinking fraction for UCYN-A1 (59%), UCYN-C (62%) and Trichodesmium (67%). Taken together, these results indicate that over short time scales (2 h, the conventional settling of particles following MSC deployment<sup>29</sup>): i) small UCYN (few µm) generally sink more efficiently than large Trichodesmium (>100 µm), ii) when sinking, Trichodesmium sinks fast, iii) UCYN-A1 and UCYN-C sink faster than UCYN-B and Gamma-A. A detailed imaging study performed on 170 m MSC samples at the mooring stations S05M and S10M indicates that the 50-80% of phycoerythrin-containing UCYN in the fast sinking fraction were organized into aggregates of tens to >250 cells measuring 30 to >100 µm, while the majority (60-95%) of UCYN were free living in the suspended fraction (Figure S3). This indicates that under minimum turbulent agitation, UCYN quickly (within 2 h) form aggregates large/dense enough to sink.

Concentrations of PN averaged across all stations 1.85  $\pm$  0.49  $\mu$ g L<sup>-1</sup>, 0.25  $\pm$  0.13  $\mu$ g L<sup>-1</sup> and 0.26  $\pm$  0.08  $\mu$ g L<sup>-1</sup> within the suspended, slow sinking and fast sinking fractions (Fig. S4). PON in the fast sinking fraction contributed 11  $\pm$  2 %, while the slow sinking and suspended fractions contributed 11  $\pm$  4 % and 79  $\pm$  5 % in terms of total PON. Thus, ~22% of the PON was sinking out of the upper part of the MSC within 2h during our study. We converted transparent expolymeric particles into C (TEP-C), that revealed higher concentrations in the suspended fraction compared to the fast sinking fraction at all stations (TEP-C concentrations were null in the SS fraction) (Fig. S4). The TEP-C:POC ratio was generally higher in the suspended fraction compared to the fast sinking, indicating a larger contribution of TEP to the POC pool.

Finally, we quantified diazotrophs on vertical profiles spanning the water column between 200 m and 2000 m by using a Bottle-net mounted on the CTD rosette frame at six stations across the transect (Fig. 1). The bottle-net consists of a 20-μm conical plankton net housed in a cylindrical PVC pipe<sup>18</sup>. It is lowered with the top cover closed, which is opened at the desired bottom depth (2000 m) of the tow, remains opened during the ascent of the rosette, and closed again at the upper depth (200 m) of the water column to be sampled. This results in one integrated sample of 200 m to 2000 m per deployment. The concentration of diazotrophs in this deep layer averaged 1.4 x 10<sup>7</sup> *nifH* gene copies m<sup>-2</sup> and was primarily dominated by UCYN-A1 (63% on average over all sampled stations) and *Trichodesmium* (27%), and secondarily by UCYN-B (9%) (Fig. 7), generally mirroring the diazotroph community in surface water (these numbers are conservative and may be underestimated as some individual UCYN pass through the 20 μm mesh net of the bottle-net). The 2000-200 m stock was relatively constant among stations, except at the most oligotrophic station S08, where it was lower by two orders of magnitude than that of other stations.

# **Biogeochemical implications**

Apart from silicified DDAs<sup>11</sup>, diazotrophs have been seldom regarded as important contributors to organic matter export. Yet, our results provide clear evidence that they are present and ubiquitous in the mesopelagic ocean and they contribute significantly to the PN export fluxes in the sub-tropical south Pacific Ocean. They have different behaviors however.

We showed that diazotroph cell sizes is not necessarily a key variable controlling their ability to sink out. Small UCYN (1-8 µm) displayed the highest export turn-over rates ~10<sup>-3</sup> d<sup>-1</sup>, in the range or higher than those reported for diatoms and coccolithophores<sup>31</sup>. Within UCYN, UCYN-A1 was clearly dominating the diazotroph community targeted by qPCR in mesopelagic waters. Although the UCYN-A1 symbiosis has been detected once in mesopelagic waters of the South Atlantic ocean<sup>32, 33</sup>, their capacity to leave the photic layer and sink throughout the mesopelagic zone down to 1000 m (Fig. 3) has not been thoroughly explored before. As UCYN-A rank among the most abundant diazotrophs in the global ocean<sup>12</sup> and spans from tropical to polar waters<sup>34, 35</sup>, we suggest that they potentially play a significant contribution to ocean organic matter sequestration. In future studies, better constraining UCYN-A sedimentation rates and aggregation processes would be of primary importance to assess their role in particle fluxes and cycling.

UCYN-B and -C are free-living unicellular cyanobacteria and their export pathway therefore does not rely on mineral ballasted hosts. Our results indicate that the sinking of such small cells is made possible thanks to aggregation of UCYN into large-size aggregates (30->100 µm) of tens to hundreds of cells, large enough to sink, in accordance with a previous mesocosm study<sup>21</sup>. The presence of UCYN-B in sediment traps is also in accordance with previous reports of *Crocospheara watsonii* sequences at station ALOHA either just below the euphotic zone<sup>22</sup>, or as deep as 4000 m<sup>36</sup>. The visualization of our sediment trap samples shows that they can be subsequently embedded in mixed aggregates with other cells and debris (Fig. 4, S2). The UCYN-B large (>4 µm) ecotype, which was dominating in this study (data not shown), produces large amounts of C-rich transparent exopolymeric particles (TEP) at rates one to two order of magnitude higher than that of diatoms and coccolithophorids<sup>37</sup>, probably in response to nutrient limitation and excess light<sup>38</sup>. In cultures of *Crocosphaera watsonii*, TEP account for ~22% of the particulate C pool<sup>37</sup>. Hence, TEP produced by UCYN-B do not only provide a matrix for the formation of large aggregates, but may also account for a significant fraction of C export. TEP are indeed greatly enriched in C relative to N (C:N ~25<sup>37</sup>), which is confirmed by the high TEP-C:PC ratios in the fast sinking fraction of the MSC.

Trichodesmium was generally the second contributor to the diazotroph community targeted by qPCR in mesopelagic waters after UCYN-A1, which contradicts the common assertion that they are entirely remineralized in the euphotic layer<sup>4, 13</sup>. Several potential mechanisms could explain the presence of Trichodesmium's in mesopelagic waters. Trichodesmium colonies are able to perform vertical migrations for exploiting the deep phosphate stock (Villareal and Carpenter 2003) (the phosphacline was around 200 m in our study region at the time of the cruise). According to this theory, they overcome their positive buoyancy by fixing C that result in carbohydrate ballasting. However, Walsby (1978)<sup>39</sup> observed that 100% of the gas vacuoles of Trichodesmium erythraeum (the most abundant species in this study) collapse at depths between 105 m and 120 m, resulting in a loss of their buoyancy. Hence, Trichodesmium could be locked in a persistent and irreversible downward trajectory. Alternatively, Berman-Frank et al.<sup>40, 41</sup> have shown that programmed cell death induces internal cellular degradation, vacuole loss and increased production of TEP, resulting in an increase in the vertical flux of Trichodesmium<sup>42</sup>. Whatever the mechanism behind the vertical flux of Trichodesmium, we report here higher abundances at 1000 m compared to those at shallower depths at both stations, consistent with microscopic observations showing intact colonies at 1000 m. This likely results from a spatio-temporal

decoupling between production and export<sup>43</sup>. Despite sinking less efficiently than UCYN in the aggregates, they are much larger and contain more C and N per filament than UCYN<sup>12</sup>, and accounted for a significant fraction of the diazotroph-attributed PN export in this study, especially at 1000 m (Table S1), which is in accordance with previous studies reported intact *Trichodesmium* colonies down to 3000-4000m<sup>18, 19</sup>.

More than 90% of the organic matter sinking below the euphotic zone is respired before it reaches a depth of 1000 m<sup>7</sup>. Fast sinking particles will therefore theoretically make a greater contribution to the deep ocean flux than slow sinking particles, since the latter will be rapidly recycled at shallow depths. The relative similarity of the taxonomic diazotroph community composition in mesopelagic waters sampled by three independent sampling approaches (traps, bottle-net, and MSC) compared to the diazotrophic composition of the euphotic layer suggests a rapid export mode of diazotrophs. This is further confirmed by i) the high proportion of diazotroph groups quantified in the MSC fast sinking fraction, ii) the whole cells and colonies and high Chlorophyll a: pheopigments ratios in sediment traps, indicative of healthy phytoplankton (Fig. 4; S2). Finally, iii) while PN export fluxes were attenuated with depth in our study, which is a classical feature in the oligotrophic ocean<sup>44</sup>, the diazotroph export fluxes were not, suggesting a high transfer efficiency to the deep ocean, and thus sinking velocities high enough to escape short-term remineralization. Indeed, a parallel study performed during the same cruise<sup>45</sup> reveals significant N<sub>2</sub> fixation rates in sediment traps, MSC and bottle-net samples, suggesting that part of the diazotroph community in mesopelagic waters sunk fast enough to remain alive at mesopelagic depths. Bar-Zeev et al. 42 reported sinking velocities of *Trichodesmium erythraeum* of ~200 m d<sup>-1</sup>. More recently, Ababou et al. <sup>46</sup> reported sinking velocities of *Trichodesmium erythraeum* and UCYN aggregates formed in roller tanks of 92 ± 37 m d<sup>-1</sup> and 333 ± 176 m d<sup>-1</sup>, respectively. We calculated that Trichodesmium sinking at those rates (92-200 m d<sup>-1</sup>) would take 5 to 10 days to reach 1000 m, whereas UCYN would take 3 days, which would be compatible with finding active cells observed at those depths<sup>45</sup>.

# Conclusion

These findings challenge the common assumption that the fate of diazotrophic new nitrogen is constrained to the surface layer. We show that diazotrophs other than DDAs get exported below the euphotic zone through direct gravitational settling, and that small diazotrophs are the most efficiently exported thanks to aggregation processes in large particle. However, direct export through gravitational settling of diazotrophs is most likely only the tip of the iceberg and diazotrophs can potentially be exported through secondary pathways: diazotrophs release in seawater 10–50% of recently fixed N<sub>2</sub> (referred to as Diazotroph Derived, DDN) as NH4+ and dissolved organic N (DON)<sup>47, 48, 49</sup>. This DDN is potentially available for assimilation by the surrounding phytoplanktonic communities, supporting their growth and leading a potential secondary (indirect) export pathway of diazotroph-derived OC<sup>6, 21, 50</sup>. Moreover, DDN is also transferred to zooplankton, which produces dejections termed fecal pellets, that sink fast and play a major role in OC export to the deep ocean<sup>51</sup>. In future studies, there is an urgent need to develop appropriate approaches to decipher diazotroph export pathways if we are to understand the role of diazotroph of the biological carbon pump. This is an especially pressing question given that current climate models predict an expansion of the oligotrophic gyres (60% of our oceans), where diazotrophs thrive and sustain most new primary productivity<sup>23, 24</sup>.

#### Methods

Samples were collected during the TONGA cruise (Fig. 1, DOI: 10.17600/18000884) in the tropical South Pacific Ocean onboard the R/V *L'Atalante* from November 1 to December 5 2019. We collected suspended and sinking particles in the mesopelagic zone using three complementary devices: surface-tethered drifting sediment traps<sup>52</sup>, MSC and Bottle-net. Additionally, water column samples were collected from the euphotic layer using Go-Flo bottles mounted on a trace metal clean rosette to quantify the stock of the major groups of diazotrophs in the photic layer.

**Satellite products and photic layer sampling**. Satellite-derived surface chlorophyll *a* concentration during the TONGA cruise (November 2019) was accessed at <a href="http://oceancolor.gsfc.nasa.gov">http://oceancolor.gsfc.nasa.gov</a> (MODIS Aqua, 4 km, 8-days composite, level 3 product). Vertical 0-150 m depth profiles were performed at each station using a trace metal clean titanium rosette of Go-Flo bottles equipped with a fluorometer and temperature, conductivity and oxygen sensors. Seawater samples were collected from 5 depths (75, 50, 20, 10, 1% surface irradiance levels) to quantify the stock of major groups of diazotrophs in the photic layer by quantitative PCR (qPCR) as described below. Additionally, primary production and N<sub>2</sub> fixation rates were measured in triplicates at the same six depths at the locations of the traps deployments (S05M and S10M), as described in <sup>53, 54</sup>. N<sub>2</sub> fixation rates were measured at two occasions (on day 1

and day 3 of the traps deployment). The average value between both profiles was used to calculate the e-ratios (POC export:primary production) and the contribution of N<sub>2</sub> fixation to primary production rates.

Sediment trap deployment and sample analyses. A surface tethered mooring line (~1000 m long) was deployed at stations S05M (21.157°S;175.153°W, 5 days) and S10M (19.423°S;175.133°W, 4 days). The line was equipped with sediment traps (KC Denmark®) at 3 depths: 170 m (corresponding to the base of the photic layer), 270 m and 1000 m. Each trap was composed of four particle interceptor tubes (PITs) mounted on a cross frame (collecting area of 0.0085 m<sup>2</sup>, aspect ratio of 6.7). Two of the tubes were used for this study: one tube was used for biogeochemical analyses (hereafter referred to as 'Biogeo' tube), and one for microbiological analyses ('Microbio' tube). Prior to deployment, the tubes were filled with 0.2 µm filtered seawater with added saline brine (50 g L<sup>-1</sup>). Borate-buffered formalin (5%) was also added to the Biogeo tube to prevent in situ microbial decomposition<sup>52</sup>. After recovery, the density gradient was visually verified, and the PITs were allowed to settle for 2 h before the supernatant seawater was carefully removed with a peristaltic pump. The remaining water containing the sinking material was transferred to a chlorohydric acid-washed container, while being screened with a 500 µm mesh to remove swimmers (zooplankton that actively entered the traps)<sup>55</sup>. Subsequently, samples were split into 12 aliquots. A triplicate set of aliquots were filtered onto 25-mm diameter Supor filters for nifH sequencing and nifH qPCR as described below. Another triplicate set of aliquots were filtered onto 25mm diameter combusted (4h, 450°C) glass microfiber filters (Whatman GF/F), which were subsequently dried for 24 h at 60°C, pelleted and from which particulate N (PN) and C (PC) were analyzed by EA-IRMS (Elemental Analyzer-Isotope Ratio Mass Spectrometry) using an Integra-CN (Sercon) mass spectrometer. Lastly, a triplicate set of aliquots were filtered on GF/F filters for further pigment analyses<sup>56</sup>, and another triplicate filtered on 1 µm polycarbonate filters for microscopic analyses (see below for methods). The turnover rate of diazotrophs in the photic layer due to vertical export was determined by dividing cell fluxes (nifH gene copies m<sup>-2</sup> d<sup>-1</sup>) by their integrated abundance in the photic later (*nifH* gene copies m<sup>-2</sup>).

# **MSC** deployment

 Suspended and sinking particles were sampled using a MSC at 3 depths at the mooring stations S05M and S10M (170 m, 270 m and 1000 m), and at 200 m, 400 m and 1000 m at S8. Additionally, the MSC was deployed at 200 m at S01, S02, S03, S04, S05V, S07, S10V and S11 (Fig. 1). The MSC is a large volume water sampler (100 L) that collects sinking particles with minimal turbulent agitation<sup>57</sup>. Upon recovery, the MSC is placed on deck for 2 h while any particles present settle onto the base of the bottom 7 L chamber. After each recovery, the MSC is conventionally left on deck for 2 h allowing particles to settle<sup>29</sup>. Fast sinking particles were thereby deposited collected in a dedicated plate at the bottom of the MSC, slow sinking particles were collected from the 7-litre compartment above the plate, and the non-sinking (or suspended) fraction was collected from the upper part of the MSC<sup>29</sup>. All three fractions were collected separately (4 L for the suspended, 1.2 L for the slow sinking). For the fast sinking fraction, the entire volume of the plate was sampled (280 to 310 mL). Subsamples from each fraction were filtered for PN and PC quantification, TEP quantification according to Passow and Alldredge (1995)<sup>58</sup>, microscopic observations, and *nifH* sequencing, *nifH* qPCR as described below. We used the formula described in Riley et al., (2012)<sup>29</sup> to obtain values associated with fast or slow sinking particles only, quantitatively removing the contribution of other fractions.

## **Bottle-net deployment**

Vertical 2000-200 m profiles were done using a bottle-net mounted on the CTD rosette frame at stations S03, S04, S05M, S07, S08, S10M, S11 and S12. The bottle-Net consists of a 20-µm conical plankton net housed in a cylindrical PVC pipe<sup>18</sup>. It is lowered with the top cover closed, which is opened at the desired bottom depth (Db, 2000 m) of the tow, remains opened during the ascension of the rosette, and closed again at the upper depth (Du, 200 m) of the water column to be sampled. This results in one integrated sample of 2000 to 200 m per deployment. Once on deck, the bottle-nets were gently rinsed with filtered seawater before retrieving the sample from the collector. At each station, samples were split into aliquots that were processed for were analyzed for *nifH* qPCR. Sampled volume was estimated as the product between the cross-sectional area of the mouth of the bottle-Net (7.5 cm, aspect ratio of 4) and the vertical distance covered by the device from the start of the ascension to the closure of the top cover (1800 m). Blank casts were performed with the bottle-net closed during the entire cast to assess for potential contaminations, and blanks were subtracted.

## nifH gene sequencing and bioinformatics

The nifH gene was sequenced from a total of 71 samples: 18 samples collected from sediment traps deployed at 170 m, 270 m and 1000 m at S05M and S10M, and 53 samples collected from the suspended, slow sinking and fast sinking fractions of particles collected with the MSC. To that end, DNA was extracted using the DNeasy Plant Mini Kit (Qiagen, Courtaboeuf, France) with additional freezethaw bead beating and proteinase K steps before the kit purification<sup>59</sup>. Triplicate nested PCR reactions were conducted using degenerate *nifH* primers<sup>60</sup>. The PCR mix was composed of 5 μL of 5X MyTaq red PCR buffer (Bioline), 1.25 μL of 25 mM MgCl<sub>2</sub>, 0.5 μl of 20 μM forward and reverse primers, 0.25 μL Platinum Taq and 5 µL of DNA extract (1 µL on second round). The reaction volume was adjusted to 25 µL with PCR grade water. Triplicate PCR products were pooled and purified using the Geneclean Turbo kit (MP Biomedicals). Partial adapters were added by ligation at the sequencing facility (Genewiz) and llumina MiSeq 2x300 paired end sequenced. Demultiplexed paired-end sequences were dereplicated, denoised, assembled and chimeras discarded using the DADA2 pipeline<sup>61</sup>. Fifty-thousand to 60,000 reads were obtained per sample. In total, >2.9 millions of high quality nifH sequences were obtained resulting in 14925 ASVs (submitted to NCBI with accession numbers xx-xx). ASVs were annotated down DADA2 nifH the genus usina а formatted gene database (https://github.com/moyn413/nifHdada2). Sequences were grouped into 17 genus according to the database, and sequences not identified to the genus level were grouped as "others". The nifH gene was successfully amplified from all samples.

# Abundance of diazotrophs et contribution to N export fluxes

The abundance of diazotrophs was determined using TaqMan qPCR assays and previously published primer-probe sets for *Trichodesmium*, UCYN-A1, UCYN-B, UCYN-C and γ-24474A11 (the latter hereafter referred to as Gamma-A) targeting the *nifH* gene<sup>62, 63, 64</sup>. The qPCR was run in 25 μL reactions consisting of 12.25 μL TaqMan PCR Master Mix (Applied Biosystems, Villebon Sur Yvette, France), 1 μL of the forward and reverse primers at 10 μM (HPLC purified, Eurofins, Nantes, France), 0.25 μL probe at 10 μM, 8.25 μL PCR grade water, 0.25 μL bovine serum albumin at 10.08 μg μL<sup>-1</sup>, and 2 μL standard or template sample. The qPCR program was run on a CFX96 Real-Time System thermal cycler (BioRad, Marnes-la-Coquette, France) and consisted of 2 min at 50°C, 10 min at 95°C continued by 45 cycles of 15 s at 95°C and 1 min at 64°C. The annealing temperature was changed to 60°C for UCYN-A1 qPCR runs<sup>64</sup>. Standard dilutions (10<sup>7</sup>-10<sup>1</sup> gene copies) were run in duplicate, and samples and notemplate controls (NTCs) in triplicate. NTCs did not show any amplification. The efficiency was 98-113%. Inhibition tests were carried out on all samples and each primer-probe set by adding the 2 μL of the 10<sup>5</sup> copies standard to each sample. No inhibition was observed. The limit of detection and detected but not quantifiable limits were 1 and 8 gene copies per reaction, respectively.

The diazotroph turnover rate representing the fraction of surface diazotrophs exported out of the photic layer per day was calculated as follows: Turnover rate (d<sup>-1</sup>) = export flux/abundance in the photic layer. The proportion of PN collected by sediment traps attributed to diazotrophs was calculated based on estimated PN content per cell. For nanoplanktonic UCYN, cell dimensions were directly measured and their biovolume calculated<sup>65</sup>. The C content per cell was estimated from the biovolume according to Verity et al. (1992)<sup>66</sup> and the N content calculated based on C:N ratios of 5 for UCYN-B <sup>67,68</sup> and 8.5 for UCYN-C<sup>47</sup>. For the UCYN-A1-host symbiosis, we considered a size of 1 µm for the UCYN-1 and 2 µm for the host and a C:N ratio of 6.3<sup>69</sup>. For *Trichodesmium*, we considered the lower-end value of 10 ng N per *Trichodesmium* filament from the same region<sup>70</sup>. To circumvent the polyploidy of *Trichodesmium*<sup>71</sup>, we divided the qPCR-based abundances by 12. This ratio was determined for this study by dividing the qPCR-based abundances by the number of *Trichodesmium* filaments counted at S05M and S10M stations at 6 depths in the photic layer (we considered 100 cells per filament of *Trichodesmium*).

## Particle imaging

Seawater samples from traps, MSC and bottle-nets were gently filtered on  $0.2~\mu m$  (for scanning electron microscopy, SEM) and  $2~\mu m$  polycarbonate filters (for epifluorescence microscopy) at very low pressure (xxx) to preserve the particle structure. For epifluorescence microscopy, filters were fixed with paraformaldehyde (2% prepared in filtered seawater) for 10 minutes at ambient temperature and stored at-80°C until visualized using a Zeiss Axioplan (Zeiss, Jena, Germany) microscope fitted with a green (510–560 nm) excitation filter, which targeted the phycoerythrin-rich cells. For SEM, samples were fixed with 2.5% glutaraldehyde and 1.6% PFA for 1 h at room temperature. Subsequently, the filters were rinsed twice in 0.2  $\mu$ m filtered in seawater during 15 min, rinsed in osmium during 30 min, and rinsed thrice in filtered seawater to eliminate excess osmium. Next, the filters passed through a series of ethanol drying solutions (50, 70, 95 and 100%, 10 min each), and a series of HDMS solutions (30, 50,

80 and 100%, 10 min each). Finally, the filters were air-dried and stored at room temperature until SEM analyses. The filters were visualized onshore using a Phenom-Pro benchtop scanning electron microscope at 10 kV.

Table 1. Particulate nitrogen (PN) export fluxes at stations S05M and S10M at 170 m, 270 m and 1000 m, and contribution of diazotrophs to total PN export.

Station		S05M			S10M	
Depth of deployment	PN export flux	Diazotroph-PN	%Contribution diazotrophs	PN export flux	Diazotroph-PN	%Contribution diazotrophs
	mg N m <sup>-2</sup> d <sup>-1</sup>	mg N m <sup>-2</sup> d <sup>-1</sup>	%	mg N m <sup>-2</sup> d <sup>-1</sup>	mg N m <sup>-2</sup> d <sup>-1</sup>	%
170 m	4.9±0.6	0.6	13	13.4±1.3	7.6	6
270 m	6.1±1.2	1.9	30	9.7±0.6	7.8	8
1000 m	1.6±0.2	12.3	100	5.2±0.1	2.3	45

Figure 1. a. Surface chlorophyll MODIS composite averaged over the time period corresponding to the TONGA cruise (1 November-6 December 2019) at a resolution of 4 km. Black triangles correspond to stations where surface-tethered drifting sediment traps were deployed (170 m, 270 m, 1000 m). Grey dots correspond to stations where Marine Snow Catcher (MSC) casts were performed at three depths (see Methods), and white dots to MSC casts performed at one depth (200 m). Black circles correspond to stations where the bottlenet profiles were performed between 2000 m and 200 m. b. Abundances of the five *nifH* phylotypes (Log10 *nifH* gene copies L<sup>-1</sup>) studied during the cruise over the transect (averages over the photic layer, ~0-130 m).

Figure 2. Relative abundance of diazotroph genera on sediment trap samples. The dendrogram corresponds to the clustering of samples generated by Bray-Curtis similarity. The legend indicates the *nifH* genus affiliation.

Figure 3. a. Export flux (*nifH* gene copies m<sup>-2</sup> d<sup>-1</sup>) of the five diazotroph groups targeted by qPCR (UCYN-A1 symbiosis, UCYN-B, UCYN-C, *Trichodesmium* and Gamma-A) in sediment trap samples at 170 m, 270 m and 1000 m at stations S05M and b. at S10M. Error bars represent strandard deviations from triplicate aliquote analyzed in duplicates. c. Export turnover rates (d<sup>-1</sup>) of the same diazotroph groups (average of the three depths and the two stations for each group).

Figure 4. Microscopy images showing examples of phycoerythrin-containing UCYN-like cells and *Trichodesmium* in sediment trap samples collected at 170 m, 270 m, and 1000 m at stations S05M and S10M. a-f. Images taken by epifluorescence microscopy (green excitation 510–560 nm, scale bar: 50 µm). k-l. Image taken by scanning electron microscopy (SEM).

Figure 5. Relative abundance of diazotroph genera from marine snow catcher samples. The dendrogram corresponds to the clustering of samples generated by Bray-Curtis similarity. The legend indicates the *nifH* genus affiliation.

Figure 6. a-e. Quantification of the five diazotroph groups targeted by qPCR in Marine Snw catcher samples: UCYN-A1 symbiosis: *Trichodesmium*, UCYN-B, UCYN-C, and Gamma-A in the suspended (dark color), slow sinking (medium color) and fast sinking (light color) pools of the MSC samples across the transect (left Y axis, *nifH* gene copies L<sup>-1</sup>). Dots represent the integrated pool of each phylotype (right Y axix, *nifH* gene copies m<sup>-2</sup>) in the photic (~0-130 m) layer. f. Percentage of sinking versus non-sinking diazotrophs for each group.

Figure 7. Abundance (*nifH* gene copies m<sup>-2</sup>) of the five diazotroph groups targeted by qPCR (UCYN-A1 symbiosis, *Trichodesmium*, UCYN-B, UCYN-C, and Gamma-A) in the bottle-net samples across the transect, representing an integrated sample from 200 m to 2000 m.

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# **Author contributions**

SB designed the experiments and SB, MB and IBF carried them out at sea, with advice from FLM; SB, MB, MC, OG, AT, DS analyzed the samples; SB analyzed the data with the help of MB for the sequencing data. SB prepared the manuscript with contributions of all co-authors.

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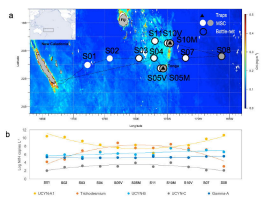
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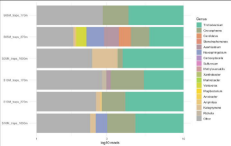
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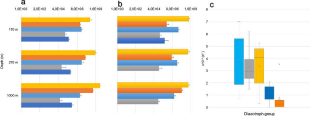
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# Log10 nifH gene copies m² d¹ b



■UCYN-A ■ Trichodesmium ■ UCYN-B ■ UCYN-C ■ Gamma-A

