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In-depth characterization of diazotroph activity across the western tropical South Pacific hotspot of N\textsubscript{2} fixation (OUTPACE cruise)

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Abstract. Here we report N\textsubscript{2} fixation rates from a ∼4000 km transect in the western and central tropical South Pacific, a particularly undersampled region in the world ocean. Water samples were collected in the euphotic layer along a west to east transect from 160°E to 160°W that covered contrasting trophic regimes, from oligotrophy in the Melanesian archipelago (MA) waters to ultra-oligotrophy in the South Pacific Gyre (GY) waters. N\textsubscript{2} fixation was detected at all 17 sampled stations with an average depth-integrated rate of 631±286 µmol N m\textsuperscript{−2} d\textsuperscript{−1} (range 196–1153 µmol N m\textsuperscript{−2} d\textsuperscript{−1}) in MA waters and of 85±76 µmol N m\textsuperscript{−2} d\textsuperscript{−1} (range 18–172 µmol N m\textsuperscript{−2} d\textsuperscript{−1}) in GY waters. Two cyanobacteria, the larger colonial filamentous Trichodesmium and the smaller UCYN-B, dominated the enumerated diazotroph community (>80 %) and gene expression of the nifH gene (cDNA >10\textsuperscript{5} nifH copies L\textsuperscript{−1}) in MA waters. Single-cell isotopic analyses performed by nanoscale secondary ion mass spectrometry (nanoSIMS) at selected stations revealed that Trichodesmium was always the major contributor to N\textsubscript{2} fixation in MA waters, accounting for 47.1–83.8 % of bulk N\textsubscript{2} fixation. The most plausible environmental factors explaining such exceptionally high rates of N\textsubscript{2} fixation in MA waters are discussed in detail, emphasizing the role of macro- and micro-nutrient (e.g., iron) availability, seawater temperature and currents.

1 Introduction

In the ocean, nitrogen (N) availability in surface waters controls primary production and the export of organic matter (Dugdale and Goering, 1967; Eppley and Peterson, 1979; Moore et al., 2013). The major external source of new N to the surface ocean is biological di-nitrogen (N\textsubscript{2}) fixation (100–150 Tg N yr\textsuperscript{−1}, Gruber, 2008), the reduction of atmospheric gas (N\textsubscript{2}) dissolved in seawater into ammonia (NH\textsubscript{4}+). The process of N\textsubscript{2} fixation is mediated by diazotrophic organisms that possess the nitrogenase enzyme, which is encoded by a suite of nif genes. These organisms provide new N to the surface ocean and act as natural fertilizers, contributing to sustaining ocean productivity and eventually carbon (C) sequestration through the N\textsubscript{2}-primed prokaryotic C pump (Caffin et al., 2018a; Karl et al., 2003, 2012). This N source is continuously counteracted by N losses, mainly driven by denitrification and anammox, which convert reduced forms of N (nitrate, NO\textsubscript{3}−, nitrite NO\textsubscript{2}−, NH\textsubscript{4}+) into N\textsubscript{2}. Despite the

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critical importance of the N inventory in regulating primary production and export, the spatial distribution of N gains and losses in the ocean is still poorly resolved.

A global-scale modeling study predicted that the highest rates of N\textsubscript{2} fixation would be located in the South Pacific Ocean (Deutsch et al., 2007; Gruber, 2016). These authors also concluded that processes leading to N gains and losses are spatially coupled to oxygen-deficient zones such as in the eastern tropical South Pacific (ETSP), which harbors NO\textsubscript{3}\textsuperscript{-}-poor but phosphate-rich surface waters, i.e., potentially ideal niches for N\textsubscript{2} fixation (Zehr and Turner, 2001). However, recent field studies based on several cruises and independent approaches, including 15N\textsubscript{2} incubation-based measurements and geochemical δ\textsuperscript{15}N budgets, have consistently measured low N\textsubscript{2} fixation rates (average range ∼0–60 μmol N m\textsuperscript{-2} d\textsuperscript{-1}) in the surface ETSP waters (Dekaezemacker et al., 2013; Fernández et al., 2011, 2015; Knapp et al., 2016; Loescher et al., 2014). Low activity in the ETSP has been largely attributed to iron (Fe) limitation (Bonnet et al., 2017; Dekaezemacker et al., 2013), as Fe is a major component of the nitrogenase enzyme complex required for N\textsubscript{2} fixation (Raven, 1988). However, the western tropical South Pacific (WTSP) was recently identified as having high N\textsubscript{2} fixation activity (Bonnet et al., 2017), and collectively these studies plead for a basin-wide spatial decoupling between N\textsubscript{2} fixation and denitrification in the South Pacific Ocean.

The WTSP is a vast oceanic region extending from Australia in the west to the western boundary of the South Pacific Gyre in the east (hereafter referred to as AG waters) (Fig. 1). It has been chronically undersampled (Luo et al., 2012) as compared to the tropical North Atlantic (Benavides and Voss, 2015) and North Pacific (e.g., Böttjer et al., 2017) oceans; however, recent oceanographic surveys performed in the western part of the WTSP, in the Solomon, Bismarck (Berthelot et al., 2017; Bonnet et al., 2009, 2015) and Ararura (Messer et al., 2015; Montoya et al., 2004) seas, report extremely high N\textsubscript{2} fixation rates (> 600 μmol N m\textsuperscript{-2} d\textsuperscript{-1}, i.e., an order of magnitude higher than in the ETSP) throughout the year. In these regions, high N\textsubscript{2} fixation has been attributed to sea surface temperatures > 25 °C and continuous nutrient inputs of terrigenous and volcanic origin (Labatut et al., 2014; Radic et al., 2011). The central and eastern parts of the WTSP, a vast oceanic region bordering Melanesian archipelagoes (New Caledonia, Vanuatu, Fiji) up to the Tonga trench (hereafter referred to as MA waters) have been far less investigated. One study (Shiozaki et al., 2014) reported high surface N\textsubscript{2} fixation rates close to the Melanesian islands in relation to nutrients supplied by land runoff. However, the lack of direct N\textsubscript{2} fixation measurements over the full photic layer impedes accurate N budget estimates in this region. In addition, the reasons for such an ecological success of diazotrophs in the WTSP are still under debate (Bonnet et al., 2017) as the horizontal and vertical distribution of environmental parameters potentially controlling N\textsubscript{2} fixation, in particular measured Fe concentrations, are still scarce in this region.

Recurrent blooms of the filamentous cyanobacterium Trichodesmium, one of the most abundant diazotrophs in our oceans (Luo et al., 2012), have been consistently reported in the WTSP since the James Cook (Cook, 1842) and Charles Darwin expeditions, and later confirmed by satellite observations (Dupouy et al., 2011, 2000) and microscopic enumerations (Shiozaki et al., 2014; Tenório et al., 2018). However, molecular studies based on the nifH gene abundances have shown high densities of unicellular diazotrophic cyanobacteria (UCYN) in the WTSP (Moisander et al., 2010). Three main groups of UCYN (A, B and C) can be distinguished based on nifH gene sequences. In the warm (> 25 °C) waters of the Solomon Sea, UCYN from group B (UCYN-B) co-occur with Trichodesmium at the surface, and together dominate the diazotrophic community (Bonnet et al., 2015), while UCYN-C are also occasionally abundant (Berthelot et al., 2017). Further south in the Coral and Tasman seas, UCYN-A dominates the diazotroph community (Bonnet et al., 2015; Moisander et al., 2010). Both studies reported a transition zone from UCYN-B-dominated communities in warm (> 25 °C) surface waters to UCYN-A-dominated communities in colder (< 25 °C) waters of the western part of the WTSP. Further east in the MA waters, Trichodesmium and UCYN-B co-occur and account for the majority of total nifH genes detected (Stenegren et al., 2018). Although molecular methods greatly enhanced our understanding of the geographical distribution of diazotrophs in the WTSP, DNA-based nifH counts do not equate to metabolic activity. Thus, the contribution of each dominant group to bulk N\textsubscript{2} fixation is still lacking in the WTSP. Previous studies showed that different diazotrophs have different fates in the ocean: some are directly exported, and others release and transfer part of the recently fixed N to the planktonic food web and indirectly fuel export of organic matter (Bonnet et al., 2016a, b; Karl et al., 2012). Consequently assessing the relative contribution of each dominating group of diazotrophs to overall N\textsubscript{2} fixation is critical to assess the biogeochemical impact of N\textsubscript{2} fixation in the WTSP.

In the present study, we report new bulk and group-specific N\textsubscript{2} fixation rate measurements from a ~4000 km transect in the western and central tropical South Pacific. The goals of the study were (i) to quantify both horizontal and vertical distribution of N\textsubscript{2} fixation rates in the photic layer in relation to environmental parameters, (ii) to quantify the relative contribution of the dominant diazotrophs (Trichodesmium and UCYN-B) to N\textsubscript{2} fixation based on cell-specific measurements, and (iii) to assess the potential biogeochemical impact of N\textsubscript{2} fixation in this region.
2 Methods

Samples were collected during the 45-day OUTPACE (Oliotrophic to UlTra oligotrophic PACific Experiment) cruise (DOI: https://doi.org/10.17600/15000900) onboard the R/V L’Atalante in February–March 2015 (austral summer). The west to east zonal transect along ∼19°S started in Noumea (New Caledonia) and ended in Papeete (French Polynesia) (Fig. 1). It covered a trophic gradient from oligotrophy (deep chlorophyll maximum (DCM) located at ∼100 m) in MA waters around New Caledonia, Vanuatu, Fiji and Tonga, to ultra-oligotrophy (DCM located at 115–150 m) in GY waters located at the western boundary of the South Pacific Gyre (Moutin et al., 2017, for details on this cruise). Data were collected at 17 stations including 14 short-duration (SD; 8 h) stations (SD1 to SD15; note that SD13 was not sampled) and 3 long-duration (LD; 7 days) stations (LDA, LDB and LDC). Vertical (0–200 m) profiles of temperature, salinity, and chlorophyll fluorescence were obtained at all 17 stations using a Seabird 911 plus CTD (conductivity, temperature, depth) equipped with a Wetlabs ECO-AFL/FL fluorometer. Seawater samples were collected by 12 L Niskin bottles mounted on the CTD rosette.

2.1 Macro-nutrient and dissolved Fe concentration analyses

Samples for the quantification of nitrate (NO$_3^-$) and dissolved inorganic phosphorus (DIP) concentrations were collected at 12 depths between 0 and 200 m in acid-washed polyethylene bottles, poisoned with mercuric chloride (HgCl$_2$, final concentration 20 mg L$^{-1}$) and stored at 4 °C until analysis. Concentrations were determined using standard colorimetric techniques (Aminot and Kérouel, 2007) on a Bran Luebbe AA3 autoanalyzer. Detection limits for the procedures were 0.05 µmol L$^{-1}$ for NO$_3^-$ and DIP.

Samples for determining dissolved Fe concentrations were collected and analyzed as described in Guieu et al. (2018). Briefly, samples were collected using a Titane rosette mounted with 24 Teflon-coated 12 L GoFlos deployed with a Kevlar cable. Dissolved Fe concentrations were measured by flow injection with online preconcentration and chemiluminescence detection according to Bonnet and Guieu (2006). The reliability of the method was monitored by analyzing the D1 SAFe seawater standard (Johnson et al., 2007), and an internal acidified seawater standard was measured daily to monitor the stability of the analysis.

The sampling and analytical methods used to analyze the other parameters reported in the correlation table (Table 2) are described in detail in the methods sections for related pa-
pers in this issue (Bock et al., 2018; Fumenia et al., 2018; Stenegren et al., 2018; Van Wambeke et al., 2018).

2.2 Bulk N₂ fixation rate measurements

Whole water (bulk) N₂ fixation rates were measured in triplicate at all 17 stations using the 15N₂ isotopic tracer technique (adapted from Montoya et al., 1996). The 15N₂ bubble technique was intentionally chosen due to the time limitation on making enriched 15N₂ seawater inoculates (e.g., 6–9 depths = 6–9 inoculates) and the larger sample bottles required for making proper estimates of activity in oligotrophic environments. In addition, we aimed to avoid any potential overestimation due to trace metal and dissolved organic matter (DOM) contaminations often associated with the preparation of the 15N₂-enriched seawater (Klawonn et al., 2015; Wilson et al., 2012) in our incubation bottles as Fe and DOM have been found to control N₂ fixation or nifH gene expression in this region (Benavides et al., 2017; Moisander et al., 2011). However, the 15N / 14N ratio of the N₂ pool available for N₂ fixation (the term AN used in Montoya et al., 1996) was measured in all incubation bottles by membrane inlet mass spectrometry (MIMS) to ensure accurate rate calculations (see below).

Table 1. 15N / 14N ratio of suspended particulate nitrogen (PN) (average over the photic layer) across the OUTPACE transect.

<table>
<thead>
<tr>
<th>Station no.</th>
<th>15N / 14N ratio – PNₘₐₛₚ (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>MA waters</td>
<td></td>
</tr>
<tr>
<td>1</td>
<td>2.00</td>
</tr>
<tr>
<td>2</td>
<td>0.78</td>
</tr>
<tr>
<td>3</td>
<td>0.57</td>
</tr>
<tr>
<td>A</td>
<td>−</td>
</tr>
<tr>
<td>4</td>
<td>2.71</td>
</tr>
<tr>
<td>5</td>
<td>1.57</td>
</tr>
<tr>
<td>6</td>
<td>1.91</td>
</tr>
<tr>
<td>7</td>
<td>0.50</td>
</tr>
<tr>
<td>8</td>
<td>−2.45</td>
</tr>
<tr>
<td>9</td>
<td>−2.21</td>
</tr>
<tr>
<td>10</td>
<td>−2.70</td>
</tr>
<tr>
<td>11</td>
<td>−7.05</td>
</tr>
<tr>
<td>12</td>
<td>1.89</td>
</tr>
<tr>
<td>B</td>
<td>−2.88</td>
</tr>
<tr>
<td>Average MA waters</td>
<td>−0.41</td>
</tr>
<tr>
<td>GY waters</td>
<td></td>
</tr>
<tr>
<td>C</td>
<td>7.91</td>
</tr>
<tr>
<td>14</td>
<td>8.72</td>
</tr>
<tr>
<td>15</td>
<td>7.55</td>
</tr>
<tr>
<td>Average GY waters</td>
<td>8.06</td>
</tr>
</tbody>
</table>

Seawater samples were collected from Niskin bottles into 10 % HCl-washed, sample-rinsed (three times) light-transparent polycarbonate (2.3 L) bottles from six depths (75, 50, 20, 10, 1, and 0.1 % surface irradiance levels) at all short-duration stations SD1 to SD15 and nine depths (75, 50, 35, 20, 10, 3, 1, 0.3, and 0.1 % surface irradiance levels) at LD A, LDB and LD C, corresponding to the sub-surface (5 m) down to 80 to 180 m, depending on the station. Bottles were sealed with caps fitted with silicon septa and amended with 2 mL of 98.9 at. % 15N₂ (Cambridge isotopes). The purity of the 15N₂ Cambridge isotope stocks was previously checked by Dabundo et al. (2014) and more recently by Benavides et al. (2015) and Bonnet et al. (2016a). They were found to be lower than 2 × 10⁻⁸ mol : mol of 15N₂, leading to a potential N₂ fixation rate overestimation of < 1 %. Each bottle was shaken 20 times to break the 15N₂ bubble and facilitate its dissolution, and was incubated for 24 h. At SD stations, bottles were incubated in on-deck incubators connected to surface circulating seawater at the specified irradiances using blue screening as the duration of the station (8 h) was too short to deploy in situ mooring lines. At LD stations (7 days), one profile was incubated following the same methodology in on-deck incubators and another replicate profile was incubated in situ for comparison on a drifting mooring line located at the same depth from which the samples were collected. Incubations were stopped by filtering the entire incubation bottle onto pre-combusted (450 °C, 4 h) 25 mm diameter glass fiber filters (GF/F, Whatman, 0.7 µm nominal pore size). Filters were subsequently dried at 60 °C for 24 h before analysis of 15N / 14N ratios and particulate N (PN) determinations using an elemental analyzer coupled to a mass spectrometer (EA-IRMS, Integra CN, SerCon Ltd) as described in Bonnet et al. (2011).

To ensure accurate rate calculations, the 15N / 14N ratio of the N₂ pool in the incubation bottles was measured on each profile from triplicate surface incubation bottles from SD1 to SD14 and at all depths at SD15 and LD stations. Briefly, 12 mL was subsampled after incubation into Exetainers fixed with HgCl₂ (final concentration 20 mg L⁻¹) that were preserved upside down in the dark at 4 °C until analyzed using a MIMS according to Kana et al. (1994). Lastly, we collected time zero samples at each station to determine the natural N isotopic signature of ambient particulate N (PN). The minimum quantifiable rates (quantification limit, QL) calculated using standard propagation of errors via the observed variability between replicate samples measured according to Gradoville et al. (2017) were 0.035 nmol N L⁻¹ d⁻¹.

Discrete N₂ fixation rate measurements were depth integrated over the photic layer using trapezoidal integration procedures. Briefly, the N₂ fixation at each pair of depths is averaged, and then multiplied by the difference between the two depths to get a total N₂ fixation in that depth interval. These depth interval values are then summed over the entire depth range to get the integrated N₂ fixation rate. The rate nearest
Table 2. Summary of relationships between measured N\textsubscript{2} fixation rates and various physical and biogeochemical parameters. Also shown are correlations between measured rates and the several diazotrophic or non-diazotrophic planktonic groups enumerated at the respective stations. The corresponding unit is given for each parameter, and Spearman’s rank correlation (\(n = 102, \alpha = 0.05\)) is provided; significant correlations (\(p < 0.05\)) are indicated by an asterisk (*).

<table>
<thead>
<tr>
<th>Variable</th>
<th>Unit</th>
<th>(\text{N}_2) fixation rate Spearman’s correlation coefficient</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Physical and biogeochemical parameters</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Pressure</td>
<td>dbar</td>
<td>−0.705*</td>
</tr>
<tr>
<td>Temperature</td>
<td>°C</td>
<td>0.658*</td>
</tr>
<tr>
<td>Salinity</td>
<td>psu</td>
<td>−0.701*</td>
</tr>
<tr>
<td>Oxygen</td>
<td>µmol kg(^{-1})</td>
<td>0.151</td>
</tr>
<tr>
<td>PAR</td>
<td>µmol photons m(^{-2}) s(^{-1})</td>
<td>0.319*</td>
</tr>
<tr>
<td>(\text{NO}_3^-)</td>
<td>µmol L(^{-1})</td>
<td>−0.544*</td>
</tr>
<tr>
<td>(\text{NH}_4^+)</td>
<td>µmol L(^{-1})</td>
<td>−0.024</td>
</tr>
<tr>
<td>DIP</td>
<td>µmol L(^{-1})</td>
<td>−0.770*</td>
</tr>
<tr>
<td>Si(OH)(_4^+)</td>
<td>µmol L(^{-1})</td>
<td>−0.724*</td>
</tr>
<tr>
<td>Dissolved Fe</td>
<td>nmol L(^{-1})</td>
<td>0.398*</td>
</tr>
<tr>
<td>DON</td>
<td>µmol L(^{-1})</td>
<td>0.517*</td>
</tr>
<tr>
<td>DOP</td>
<td>µmol L(^{-1})</td>
<td>0.418*</td>
</tr>
<tr>
<td>DOC</td>
<td>µmol L(^{-1})</td>
<td>0.573*</td>
</tr>
<tr>
<td>PON</td>
<td>µmol L(^{-1})</td>
<td>0.721*</td>
</tr>
<tr>
<td>POC</td>
<td>µmol L(^{-1})</td>
<td>0.723*</td>
</tr>
<tr>
<td>Biogenic silica</td>
<td>µmol L(^{-1})</td>
<td>0.274*</td>
</tr>
<tr>
<td>Chl (a)</td>
<td>µg L(^{-1})</td>
<td>0.266*</td>
</tr>
<tr>
<td>Primary production</td>
<td>µg C L(^{-1}) h(^{-1})</td>
<td>0.657*</td>
</tr>
<tr>
<td>Bacterial production</td>
<td>µmol C L(^{-1}) h(^{-1})</td>
<td>0.692*</td>
</tr>
<tr>
<td>(T_{\text{DIP}})</td>
<td>days</td>
<td>−0.721*</td>
</tr>
<tr>
<td><strong>Planktonic groups</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>\textit{Trichodesmium sp.}</td>
<td>\textit{nifH} copies L(^{-1})</td>
<td>0.729*</td>
</tr>
<tr>
<td>UCYN-A1</td>
<td>\textit{nifH} copies L(^{-1})</td>
<td>−0.051</td>
</tr>
<tr>
<td>UCYN-A2</td>
<td>\textit{nifH} copies L(^{-1})</td>
<td>−0.147</td>
</tr>
<tr>
<td>UCYN-B</td>
<td>\textit{nifH} copies L(^{-1})</td>
<td>0.511*</td>
</tr>
<tr>
<td>\text{het-1}</td>
<td>\textit{nifH} copies L(^{-1})</td>
<td>0.538*</td>
</tr>
<tr>
<td>\text{het-2}</td>
<td>\textit{nifH} copies L(^{-1})</td>
<td>0.576*</td>
</tr>
<tr>
<td>\text{het-3}</td>
<td>\textit{nifH} copies L(^{-1})</td>
<td>0.276*</td>
</tr>
<tr>
<td>\textit{Prochlorococcus sp.}</td>
<td>cell mL(^{-1})</td>
<td>0.697*</td>
</tr>
<tr>
<td>\textit{Synechococcus sp.}</td>
<td>cell mL(^{-1})</td>
<td>0.720*</td>
</tr>
<tr>
<td>Pico-eukaryotes</td>
<td>cell mL(^{-1})</td>
<td>−0.450*</td>
</tr>
<tr>
<td>Bacteria</td>
<td>cell mL(^{-1})</td>
<td>0.780*</td>
</tr>
<tr>
<td>Protists</td>
<td>cell mL(^{-1})</td>
<td>0.680*</td>
</tr>
</tbody>
</table>

the surface is assumed to be constant up to 0 m (Knap et al., 1996).

2.3 Statistical analyses

Spearman’s rank correlation was used to examine the potential relationships between \(\text{N}_2\) fixation rates and hydrological, biogeochemical, and biological parameters across the longitudinal transect (\(n = 102, \alpha = 0.05\)). A non-parametric Mann–Whitney test (\(\alpha = 0.05\)) was used to compare the MIMS data obtained following on-deck versus in situ incubations, and to compare nutrient and Chl \(a\) distributions between the western part and the eastern part of the transect.

2.4 Group-specific \(\text{N}_2\) fixation rate measurements

2.4.1 Experimental procedures

At three stations along the transect (SD2, SD6, LDB), where \textit{Trichodesmium} and UCYN-B accounted for > 90% of the total diazotrophic community (see below and Stenegren et al., 2018), eight additional polycarbonate (2.3 L) bottles were collected from the surface (50% light irradiance) to determine \textit{Trichodesmium} and UCYN-B-specific \(\text{N}_2\) fixation rates by nanoSIMS and quantify their contribution to bulk \(\text{N}_2\) fixation. Two of these bottles were amended with \textit{\text{N}}\textsubscript{15}\text{N} as described above for further nanoSIMS analyses on individual cells (the six remaining bottles were used for DNA and RNA
analyses; see below) and were incubated for 24 h with the incubation bottles dedicated to bulk N$_2$ fixation measurements in on-deck incubators as described above. To recover large-size diazotrophs (Trichodesmium) after incubation, 1.5 L was filtered on 10 µm pore size 25 mm diameter polycarbonate filters. The cells were fixed with paraformaldehyde (PFA) (2% final concentration) for 1 h at ambient temperature (∼25 °C) and the filters were then stored at −20 °C until nanoSIMS analyses. To recover small-size diazotrophs (UCYN-B), samples were collected for further cell sorting by flow cytometry prior to nanoSIMS; 1 L of the remaining $^{15}$N$_2$ labelled bottle was filtered onto 0.2 µm pore size 47 mm polycarbonate filters. Filters were quickly placed in a 5 mL cryotube filled with 0.2 µm filtered seawater with PFA (2% final concentration) for 1 h at room temperature in the dark. The cryovials were vortexed for 10 s to detach the cells from the filter (Thompson et al., 2012) and stored at −80 °C until cell sorting. Cell sorting of UCYN-B was performed on a Becton Dickinson Influx Mariner (BD Biosciences, Franklin Lakes, NJ) high-speed cell sorter of the Regional Flow Cytometry Platform for Microbiology (PRECYM), hosted by the Mediterranean Institute of Oceanography, as described in Bonnet et al. (2016a) and Berthelot et al. (2016). After sorting, the cells were dropped onto a 0.2 µm pore size polycarbonate 13 mm diameter polycarbonate filter connected to a low-pressure vacuum pump, and then stored at −80 °C until nanoSIMS analyses. Special care was taken to drop the cells on a surface as small as possible (∼5 mm in diameter) to ensure the highest cell density possible to facilitate subsequent nanoSIMS analyses.

2.4.2 Abundance of diazotrophs by microscopy and qPCR methods

The abundance of Trichodesmium filaments and the average number of cells per filament were determined microscopically: 1 to 2.2 L was filtered on 2 µm polycarbonate filters. The cells were fixed with PFA prepared with filtered seawater (2% final concentration) for 1 h at 4 °C and stored at −20 °C until counting using an epifluorescence microscope (Zeiss Axioplan, Jena, Germany) fitted with a green (510–560 nm) excitation filter. The whole filter was counted and the number of cells per trichome were counted on at least 10 filaments per station.

Four other diazotrophic phyotypes were quantified using quantitative PCR (qPCR) as they were not easily quantifiable by standard epifluorescence microscopy: UCYN-A1, UCYN-B and two heterocystous symbionts of diatom–diazotroph associations (DDAs): Richelia intracellularis associated with Rhizosolenia spp. (het-1) and R. intracellularis associated with Hemiaulus spp. (het-2). Triplicate 2.3 L bottles were filtered onto 25 mm diameter 0.2 µm Supor filters with a 0.2 µm pore size at each station using a peristaltic pump. The DNA extraction and TaqMAN qPCR assays are fully described in Stenegren et al. (2018). To quantitify nifH gene expression, additional triplicate 2.3 L bottles were filtered as described above. The filters were placed into pre-sterilized head-beater tubes (Biospec Products Inc., Bartlesville, OK, USA) containing a 250 µL Lysis buffer (Qiagen RNaseasy) amended with 1% β-mercaptoethanol and 30 µL of 0.1 mm glass beads (Biospec Products Inc.). The time of filtering for RNA varied between stations (17:00–21:00). Filters were flash frozen in liquid nitrogen and stored at −80 °C until RNA extraction. The RNA extraction and reverse transcription (RT) were performed as previously described using a Super-Script III first-strand cDNA synthesis kit (Invitrogen Corp., Carlsbad, CA, USA) including the appropriate negative controls (water and No-RT) (Foster et al., 2010). The nifH gene expression for het-1, het-2, Trichodesmium, UCYN-A1, and UCYN-B was as described previously (Foster et al., 2010).

2.4.3 nanoSIMS analyses, data processing and group-specific rate calculations

NanoSIMS analyses were performed using an N50 nanoSIMS instrument ( Cameca, Gennevilliers, France) at the French National Ion MicroProbe Facility according to Bonnet et al. (2016a, b) and Berthelot et al. (2016). Briefly, a −1.3 pA cesium (16 KeV) primary beam focused onto a ∼100 nm spot diameter was scanned across a 256 × 256 or 512 × 512 pixel raster (depending on the image size) with a counting time of 1 ms per pixel. Samples were pre-sputtered prior to analyses with a current of −10 pA for at least 2 min to achieve sputtering equilibrium and ensure a consistent implantation and analysis of the cell interior by removing the cell surface. Negative secondary ions ($^{12}$C$^{14}$N$^-$, $^{12}$C$^{15}$N$^-$) were collected by electron multiplier detectors, and secondary electrons were also imaged simultaneously. A total of 10–50 serial quantitative secondary ion images were generated that were combined to create the final image. Mass resolving power was ∼ 8000 in order to resolve isobaric interferences; 20 to 100 planes were generated for each cell analyzed. NanoSIMS runs are time-intensive and not designed for routine analysis, but a minimum of 250 cells of UCYN-B per station and 30 Trichodesmium filament portions were analyzed to take into account the variability of activity among the population.

Data were processed using the LIMAGE software. Briefly, all scans were corrected for any drift of the beam and sample stage during acquisition. Isotope ratio images were created by adding the secondary ion counts for each recorded secondary ion for each pixel over all recorded planes and dividing the total counts by the total counts of a selected reference mass. Individual Trichodesmium filaments and UCYN-B cells were easily identified on nanoSIMS images that were used to define regions of interest (ROIs). For each ROI, the $^{15}$N / $^{14}$N ratio was calculated.

Trichodesmium and UCYN-B cellular biovolume was calculated from cell-diameter measurements performed on ∼ 50
cells or trichomes per station using an epifluorescence microscope (Zeiss Axioplan, Jena, Germany) fitted with a green (510–560 nm) excitation filter. UCYN-B had a spherical shape and Trichodesmium cells were assumed to have a cylindrical shape. The carbon content per cell was estimated from the biovolume according to Verity et al. (1992) and the N content was calculated based on C:N ratios of 6 for Trichodesmium (Carpenter et al., 2004) and 5 for UCYN-B (Dekaezemacker and Bonnet, 2011; Knapp et al., 2012). 15N assimilation rates were expressed “per cell” and calculated as follows (Foster et al., 2011, 2013): assimilation (mol N cell−1 d−1) = (15Nex × Ncon)/ NNr, where 15Nex is the excess 15N enrichment of the individual cells measured by nanoSIMS after 24 h of incubation relative to the time zero value, Ncon is the N content of each cell determined as described above, and NNr is the excess 15N enrichment of the source pool (N2) in the experimental bottles determined by MIMS (see above). Standard deviations were calculated using the variability of N isotopic signatures measured by nanoSIMS on replicate cells. The relative contribution of Trichodesmium and UCYN-B to bulk N2 fixation was calculated by multiplying cell-specific N assimilation by the cell abundance of each group, relative to bulk N2 fixation determined at the same time.

3 Results

3.1 Environmental conditions

Seawater temperature ranged from 21.4 to 30.0 °C in the sampled photic layer (0 to ~80–180 m) over the cruise transect (Fig. 2a). The mixed layer depth (MLD) calculated according to the de Boyer Montégut et al. (2004) method was located around 20–40 m throughout the zonal transect: maximum temperatures were measured in the surface mixed layer (~0–20/40 m) and remained almost constant along the longitudinal transect with 29.1 ± 0.3 °C in MA waters and 29.5 ± 0.4 °C in GY waters.

Based on other hydrographic measurements (dissolved nutrients, dissolved Fe and Chl a concentrations), the longitudinal transect was divided into two sub-regions: (1) the MA region from station SD1 (160°E) to LDB (165° W), and (2) the GY sub-region from station LDB (165° W) to SD15 (160° W). Chl a concentration in the upper 50 m was significantly (p < 0.05) higher in MA waters (0.17 µg L−1 on average) than in GY waters (0.06 µg L−1 on average) (Figs. 1 and 2b). The DCM was located around 80–100 m in MA waters and deepened to ~150 m in GY waters, indicating higher oligotrophy in the GY region. Surface NO3− concentrations (Fig. 2c) were consistently close to or below the detection limit (0.05 µmol L−1) in the upper water column (0–50 m) throughout the transect and the depth of the nitracline gradually deepened from ~75–100 m in MA waters to ~115 m in GY waters. DIP concentrations were slightly higher than or close to detection limits (0.05 µmol L−1) in MA surface waters (0–50 m), and the phosphacline was shallower (20–45 m) than the nitracline and DIP concentrations increased significantly (p < 0.05) in GY waters and ranged from 0.13 to 0.17 µmol L−1 (Fig. 2d).

3.2 N isotopic signature of the N2 pool after incubation

The 15N enrichment of the N2 pool after 24 h of incubation with the 15N2 tracer was on average 6.145 ± 0.798 at. % (n = 54) in bottles incubated in on-deck incubators and significantly higher (p < 0.05) in bottles incubated on the mooring line (7.548 ± 0.557 at. % (n = 44), Fig. 3a). However, the depth of incubation on the mooring line (between 5 and 180 m) did not have any significant effect (p > 0.05) on the isotopic signature of the N2 pool at LDB and LDC, which remained constant over the water column (Fig. 3b).

3.3 Natural isotopic signature of suspended particles and N2 fixation rates

The natural N isotopic signature of suspended particles measured over the photic layer was on average ~0.41 ‰ in MA waters and 8.06 ‰ in GY waters (Table 1). Those isotopic values were used as time zero samples to calculate N2 fixation rates.

N2 fixation was detected at all 17 sampled stations and the average measured N2 fixation rates in the two previously defined sub-regions were (1) 8.9 ± 10 nmol N L−1 d−1 (range QL-48 nmol N L−1 d−1) over the photic layer in MA waters, and (2) 0.5 ± 0.6 nmol N L−1 d−1 (range QL-4.0 nmol N L−1 d−1) in GY waters (Fig. 2e). In MA waters, N2 fixation was largely restricted to the mixed layer, where average rates were 15 nmol N L−1 d−1, with local maxima (> 20 nmol N L−1 d−1) at stations SD1, SD6 and LDB and local minima (< 5 nmol N L−1 d−1) at SD8 and SD10. In GY waters, maximum rates reached 1–2 nmol N L−1 d−1 and were located deeper in the water column (~50 m). When integrated over the photic layer, N2 fixation represented an average net N addition of 631 ± 286 µmol N m−2 d−1 (range 196–1153 µmol N m−2 d−1) in MA waters and of 85 ± 79 µmol N m−2 d−1 (range 18–172 µmol N m−2 d−1) in GY waters.

N2 fixation rates were significantly positively correlated with seawater temperature and photosynthetically active radiation (PAR) (p < 0.05), significantly negatively correlated with depth and salinity (p < 0.05) and not significantly correlated with dissolved oxygen concentrations (p > 0.05) (Table 2). N2 fixation rates were significantly positively correlated with dissolved Fe, dissolved organic N (DON), phosphorus (DOP), carbon (DOC), particulate organic N (PON), particulate organic carbon (POC), biogenic silica (BSi), Chl a concentrations, and primary and bacterial production (p < 0.05), and significantly negatively correlated with concentration of dissolved nutrients: NO3−, NH4+, DIP and sil-
icate \( (p < 0.05) \). \( \text{N}_2 \) fixation rates were significantly positively correlated with \( \text{nifH} \) abundances for \emph{Trichodesmium} spp., UCYN-B and the symbionts of DDAs (het-1, het-2) \( (p < 0.05) \) and not significantly correlated with UCYN-A1 and UCYN-A2 abundances \( (p > 0.05) \) (Table 2). Regarding non-diazotrophic plankton, \( \text{N}_2 \) fixation rates were significantly positively correlated with \emph{Prochlorococcus} spp., \emph{Synechococcus} spp., heterotrophic bacteria and protist abundances \( (p < 0.05) \) and significantly negatively correlated with picoeukaryotes \( (p < 0.05) \).

\section*{3.4 Contribution of \textit{Trichodesmium} and UCYN-B to \textit{N}_2 fixation and nitrogenase gene expression}

At the three stations where cell-specific \( \text{N}_2 \) fixation rates were estimated by nanoSIMS (SD2, SD6 and LDB), the most abundant diazotroph phylotype was \emph{Trichodesmium} with \( 1.3 \times 10^5, 3.3 \times 10^5 \) and \( 1.2 \times 10^5 \) cells L\(^{-1}\), respectively, followed by UCYN-B, whose abundances were \( 2.0 \times 10^4, 1.5 \times 10^5 \) and \( 3.8 \times 10^2 \) \text{nifH} copies L\(^{-1}\), respectively. Het-1 and het-2 combined were 1 to 2 orders of magnitude lower, ranging from 1.0 to 9.9 \times 10^3 \text{nifH} copies L\(^{-1}\), and UCYN-
Figure 3. (a) The average measured $^{15}$N / $^{14}$N ratio of the N$_2$ pool in the incubation bottles incubated either in on-deck incubators ($n = 54$) or in situ (mooring line) ($n = 44$). The dashed lines represent the theoretical value ($\sim 8.2$ at. %) calculated assuming complete isotopic equilibration between the gas bubble and the seawater based on gas constants. Error bars represent the standard deviation. (b) Depth profiles of the $^{15}$N / $^{14}$N ratio of the N$_2$ pool in the incubation bottles incubated either in on-deck incubators (filled symbols) or on an in situ mooring line (open symbols).

A1 were below detection at the three stations. In summary, *Trichodesmium* and UCYN-B accounted for 98.2, 99.8 and 92.1 % of the total diazotroph community (based on the phylootypes targeted here) at SD2, SD6 and LDB, respectively (Table 3).

The $^{15}$N / $^{14}$N ratios of individual cells/trichomes of UCYN-B and *Trichodesmium* were measured via nanoSIMS analyses and used to estimate single-cell N$_2$ fixation rates. A summary of the enrichment values and cell-specific N$_2$ fixation is provided in Table 3. Individual trichomes exhibited significant $^{15}$N enrichments ($0.610 \pm 0.269$, $0.637 \pm 0.355$ and $0.981 \pm 0.466$ at. % at stations SD2, SD6 and LDB, respectively) compared with time zero samples ($0.369 \pm 0.002$ at. %). UCYN-B were also significantly $^{15}$N-enriched, with $1.163 \pm 0.531$ and $0.517 \pm 0.237$ at. % at SD2 and SD6, respectively (note that no UCYN-B could be sorted and analyzed by nanoSIMS at LDB as they accounted for only 0.3 % of the diazotroph community). Cell-specific N$_2$ fixation rates of *Trichodesmium* were $38.9 \pm 8.1$, $29.3 \pm 5.4$ and $123.8 \pm 24.8$ fmol N cell$^{-1}$ d$^{-1}$ at SD2, SD6 and LDB. Cell-specific N$_2$ fixation of UCYN-B was $30.0 \pm 6.4$ and $6.1 \pm 1.2$ fmol N cell$^{-1}$ d$^{-1}$ at SD1 and SD6. The contribution of *Trichodesmium* to bulk N$_2$ fixation was 83.8, 47.1 and 52.9 % at stations SD2, SD6 and LDB, respectively. The contribution of UCYN-B was 10.1 and 6.1 % at SD2 and SD6, respectively (Table 3).

The in situ *nifH* expression for all diazotroph groups targeted by qPCR was estimated using a TaqMAN quantitative reverse transcription PCR (RT-qPCR) (Table 4). The sampling and filtering time (17:00–21:00 h) was not optimal for quantifying the *nifH* gene expression for all diazotrophs; however, it provides useful information about which diazotrophs were potentially active during the experiment and complements the nanoSIMS analysis which measures the in situ activity. Both *Trichodesmium* and UCYN-B dominated the biomass (Stenegren et al., 2018), as did their *nifH* gene expression at all three stations, especially SD2 and SD6. Of the two DDAs, het-1 had a higher *nifH* gene expression, which was consistent with its higher *nifH* abundance by DNA qPCR (Stenegren et al., 2018). UCYN-A1 was consistently below detection for the *nifH* gene expression and was also the least detected diazotroph by *nifH* qPCR (Stenegren et al., 2018).

4 Discussion

4.1 Methodological considerations: the importance of measuring the $^{15}$N / $^{14}$N ratio of the N$_2$ pool

Our understanding of the marine N cycle relies on accurate estimates of N fluxes to and from the ocean. Here we decided to use the “bubble addition method” to minimize potential trace metal and organic matter contaminations, which may have resulted in overestimating rates (Klawonn et al.,...
fixation rates were measured. Our MIMS results measured a significantly ($p < 0.05$) lower $^{15}$N enrichment of the N$_2$ pool (6.145 ± 0.798 at.%) when bottles were incubated in on-deck incubators compared to when bottles were incubated on the mooring line (7.548 ± 0.557 at.%). This suggests that the $^{15}$N$_2$ dissolution is much more efficient when bottles are incubated in situ, probably due to the higher pressure in seawater at the depth of incubation (1.5 to 19 bars between 5 and 180 m) compared to the pressure in the on-deck incubators (1 bar). The seawater temperature checked regularly in the on-deck incubators was equivalent to ambient surface temperature and likely did not explain the differences observed. This result highlights the need to perform routine MIMS measurements to use the most accurate N$_2$ value for rate calculations, independently of the $^{15}$N$_2$ approach used (gas or dissolved). In our study, the theoretical AN$_2$ value based on gas constant calculations (Weiss, 1970) was ~ 8.2 at.%, so the deviation from this value is more important when bottles are incubated in on-deck incubators as compared to when they are incubated in situ. This suggests that the use of the bubble addition method without MIMS measurement potentially leads to higher underestimations when bottles are incubated in on-deck incubators, which is the case in the great majority of marine N$_2$ fixation studies published so far (Luo et al., 2012). We are aware that the dissolution kinetics of $^{15}$N$_2$ in the incubation bottles may have been progressive along the 24 h of incubation (Mohr et al., 2010); therefore, the N$_2$ fixation rates provided here represent conservative values.

Moreover, a recent extensive meta-analysis (13 studies, 368 observations) between bubble and enriched amendment experiments to measure $^{15}$N$_2$ rates reported that underestimation of N$_2$ fixation is negligible in experiments that last 12–24 h (e.g., error is ~ 0.2%); hence our 24 h based experiments should be within a small amount of error (Wannicke et al., 2018). However, we paid careful attention to accurately measure the term AN$_2$ to avoid any potential underestimation and reveal that the way bottles are incubated (on-deck versus in situ) has a great influence on the AN$_2$ value, and thus on N$_2$ fixation rates.

Our MIMS results measured a significantly ($p < 0.05$) lower $^{15}$N enrichment of the N$_2$ pool (6.145 ± 0.798 at.%) when bottles were incubated in on-deck incubators compared to when bottles were incubated on the mooring line (7.548 ± 0.557 at.%). This suggests that the $^{15}$N$_2$ dissolution is much more efficient when bottles are incubated in situ, probably due to the higher pressure in seawater at the depth of incubation (1.5 to 19 bars between 5 and 180 m) compared to the pressure in the on-deck incubators (1 bar). The seawater temperature checked regularly in the on-deck incubators was equivalent to ambient surface temperature and likely did not explain the differences observed. This result highlights the need to perform routine MIMS measurements to use the most accurate AN$_2$ value for rate calculations, independently of the $^{15}$N$_2$ approach used (gas or dissolved). In our study, the theoretical AN$_2$ value based on gas constant calculations (Weiss, 1970) was ~ 8.2 at.%, so the deviation from this value is more important when bottles are incubated in on-deck incubators as compared to when they are incubated in situ. This suggests that the use of the bubble addition method without MIMS measurement potentially leads to higher underestimations when bottles are incubated in on-deck incubators, which is the case in the great majority of marine N$_2$ fixation studies published so far (Luo et al., 2012). We are aware that the dissolution kinetics of $^{15}$N$_2$ in the incubation bottles may have been progressive along the 24 h of incubation (Mohr et al., 2010); therefore, the N$_2$ fixation rates provided here represent conservative values.

Despite the AN$_2$ value being different according to the incubation mode, it did not change with the depth of incubation on the mooring line, indicating that a slightly higher pressure than atmospheric pressure (1.5 bar at 5 m depth) is enough to promote the $^{15}$N$_2$ dissolution. It also indicates that the slightly lower seawater temperature (22–24 °C) recorded at ~ 100–180 m where the deepest samples were incubated likely did not affect the solubilization of the $^{15}$N$_2$ gas. In our study, the vertical profiles performed at LD stations and incubated either on-deck in triplicate or in situ in triplicate reveal identical ($p > 0.05$) N$_2$ fixation rates regardless of the incubation method used (Caffin et al., 2018b). This indicates that in situ incubations and on-deck incubations that simulate appropriate light levels are a valid methodology for $^{15}$N$_2$ fixation rate measurements on cruises during which mooring

### Table 3. Summary of diazotroph abundances and nanoSIMS analyses at SD2, SD6 and LDB.

<table>
<thead>
<tr>
<th>Station no.</th>
<th>Trichodesmium abundance (cells L$^{-1}$)</th>
<th>Contribution to diazotroph community (%)</th>
<th>Contribution to N$_2$ fixation rates (fmol cell$^{-1}$ d$^{-1}$)</th>
<th>Contribution to bulk N$_2$ fixation (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>SD2</td>
<td>1.3 × 10$^5$</td>
<td>84.9 ± 0.610</td>
<td>38.9 ± 8.1</td>
<td>83.8</td>
</tr>
<tr>
<td>SD6</td>
<td>3.3 × 10$^5$</td>
<td>68.0 ± 0.637</td>
<td>29.3 ± 5.4</td>
<td>47.1</td>
</tr>
<tr>
<td>LDB</td>
<td>1.2 × 10$^5$</td>
<td>91.8 ± 0.981</td>
<td>123.8 ± 24.7</td>
<td>52.9</td>
</tr>
</tbody>
</table>

### Table 4. Summary of nifH gene expression data determined by qRT-PCR at selected stations (SD2, SD6, LDB), where the cell-specific N$_2$ fixation rates were measured.

<table>
<thead>
<tr>
<th>Diazotroph</th>
<th>Station SD2 cDNA nifH (gene copies L$^{-1}$)</th>
<th>Station SD6 cDNA nifH (gene copies L$^{-1}$)</th>
<th>Station LDB cDNA nifH (gene copies L$^{-1}$)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Trichodesmium</td>
<td>1.1 × 10$^5$</td>
<td>5.1 × 10$^5$</td>
<td>5.78 × 10$^4$</td>
</tr>
<tr>
<td>UCYN-B</td>
<td>1.9 × 10$^5$</td>
<td>1.5 × 10$^5$</td>
<td>1.03 × 10$^2$</td>
</tr>
<tr>
<td>het-1</td>
<td>6.83 × 10$^2$</td>
<td>1.56 × 10$^3$</td>
<td>2.04 × 10$^2$</td>
</tr>
<tr>
<td>het-2</td>
<td>5.44 × 10$^2$</td>
<td>2.14 × 10$^2$</td>
<td>bd</td>
</tr>
<tr>
<td>UCYN-A1</td>
<td>bd</td>
<td>bd</td>
<td>bd</td>
</tr>
<tr>
<td></td>
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<td></td>
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</tr>
</tbody>
</table>
lines cannot be deployed, as long as routine measurements of the isotopic ratio of the N$_2$ pool are performed in incubation bottles.

### 4.2 Drivers of high N$_2$ fixation rates in the WTSP?

N$_2$ fixation rates measured in MA waters (average 631 ± 286 µmol N m$^{-2}$ d$^{-1}$) are 3 to 4 times higher than model predictions for this area (150–200 µmol N m$^{-2}$ d$^{-1}$, Gruber, 2016). They are in the upper range of the higher category (100–1000 µmol N m$^{-2}$ d$^{-1}$) of rates defined by Luo et al. (2012) in the N$_2$ fixation MAREDAT database for the global ocean and thus identify the WTSP as an area for high N$_2$ fixation in the global ocean. Recent studies performed in the western part of the WTSP, i.e., in the Solomon, Bismarck (Berthelot et al., 2017; Bonnet et al., 2009, 2015) and Arafura (Messer et al., 2015; Montoya et al., 2004) seas, also reveal extremely high rates (> 600 µmol N m$^{-2}$ d$^{-1}$), indicating that this high N$_2$ fixation activity extends geographically west–east from Australia to Tonga and north–south from the Equator to 25–30° S, covering a vast ocean area of ~13 × 106 km$^2$ (i.e., ~20% of the South Pacific Ocean area). However, the driver(s) for diazotrophy in this region is(are) still poorly resolved and raise(s) the question of which factors influence the distribution and activity of N$_2$ fixation in the ocean. In a global-scale study conducted by Luo et al. (2014), which investigated the correlations between N$_2$ fixation and a variety of environmental parameters commonly accepted to control this process, they concluded that SST (or surface solar radiation) was the best predictor to explain the spatial distribution of N$_2$ fixation in the surface ocean. Below we highlight the most plausible factors explaining such high N$_2$ fixation rates in our study area.

**Seawater temperature.** Seawater temperature was unlikely to be the factor explaining the differences in N$_2$ fixation rates observed between MA and GY waters, as it was consistently high (> 28°C in the surface mixed layer) and optimal for the growth and nitrogenase activity of most diazotrophs (Breithardt et al., 2007; Nübel et al., 1997) all along the cruise transect. This indicates that other factors such as nutrient availability may explain the distribution of N$_2$ fixation.

**DIP availability.** The ~4000 km transect was divided into two main sub-regions: (1) the MA waters, harboring typical oligotrophic conditions with surface NO$_3^-$ and DIP concentrations close to detection limits (0.05 µmol L$^{-1}$), a nitracline located at 75–100 m, moderate surface Chl a concentrations (~0.17 µg L$^{-1}$), a DCM located at ~80–100 m and very high N$_2$ fixation rates (631 ± 286 µmol N m$^{-2}$ d$^{-1}$ on average), and (2) the GY waters harboring ultra-oligotrophic conditions with undetectable NO$_3^-$, a deeper nitracline (115 m), and comparatively high DIP concentrations (0.15 µmol L$^{-1}$), very low Chl a concentrations (0.06 µg L$^{-1}$, DCM ~150 m) and low N$_2$ fixation rates (85 ± 79 µmol N m$^{-2}$ d$^{-1}$).

In the NO$_3^-$-depleted MA waters, low DIP concentrations are indicative of the consumption of DIP by the planktonic community, including diazotrophs. This is consistent with the negative correlation found between N$_2$ fixation and DIP turnover time (the ratio between DIP concentrations and DIP uptake rates) (Table 2) and suggests a higher DIP limitation when N$_2$ fixation is high and consumes DIP. The high DIP concentrations (> 0.1 µmol L$^{-1}$) in GY surface waters compared to MA waters are consistent with former studies that consider the South Pacific Gyre as a high phosphate, low chlorophyll ecosystem (Moutin et al., 2008), in which DIP accumulates in the absence of NO$_3^-$ and low N$_2$ fixation activity. In the high phosphate, low chlorophyll scenario, the community is limited by temperature and/or Fe availability (Moutin et al., 2008; Bonnet et al., 2008). During the OUTPACE cruise, the DIP turnover time was variable but close to or below 2 days in MA waters (Moutin et al., 2018), indicating a potential limitation by DIP at some stations. *Trichodesmium*, the most abundant and major contributor to N$_2$ fixation during the cruise, is known to synthesize hydrolytic enzymes in order to acquire P from the dissolved organic phosphorus pool (DOP) (Sohm and Capone, 2006). Moreover, *Trichodesmium* spp. differs from the other major diazotroph UCYN-B enumerated on the cruise in the forms of organic P it can synthesize. It is thus likely that DOP species that favored *Trichodesmium* over UCYN-B played a role in maintaining high *Trichodesmium* biomass in MA waters. It has to be noted that average DIP turnover times in MA waters were always much higher than those typically measured in severely DIP-limited environments such as the Mediterranean and Sargasso seas (e.g., Moutin et al., 2008), suggesting that DIP concentrations are generally favorable for the development of certain diazotrophs in the WTSP, and do not alone explain why N$_2$ fixation is high in MA waters and low in GY waters. However, it is likely that the depletion of DIP stocks at the end of the austral summer season forces the decline of diazotrophic blooms in the WTSP (Moutin et al., 2005), concomitantly with the decline of SST.

**Fe availability.** Before OUTPACE, our knowledge on Fe sources and concentrations in the WTSP was limited, especially in MA waters. During OUTPACE, Guieu et al. (2018) reported high dissolved Fe concentrations in MA waters (range 0.2–66.2 nmol L$^{-1}$, 1.7 nmol L$^{-1}$ on average over the photic layer), i.e., significantly ($p < 0.05$) higher than those reported in GY waters (range 0.2–0.6 nmol L$^{-1}$, 0.3 nm on average over the photic layer). The low dissolved Fe concentrations measured in the GY waters agree well with previous reports for the same region (Blain et al., 2008; Fitzsimmons et al., 2014). However, the high dissolved Fe concentrations measured in MA waters were previously undocumented and reveal several maxima (~50 nmol L$^{-1}$) between stations SD7 and SD11, indicative of intense fertilization processes taking place in this region. Guieu et al. (2018) found that atmospheric deposition measured during the cruise in this region was too low to explain the observed dissolved Fe concentrations in the surface water column. The seafloor of the WTSP hosts the Tonga–Kermadec subduc-
tion zone which stretches 2500 km from New Zealand to the Tonga archipelago (Fig. 1). It has among the highest densities of submarine volcanoes associated with hydrothermal vents recorded in the ocean (2.6 vents/100 km, Massoth et al., 2007), which discharge large quantities of material into the water column, including biogeochemically relevant elements such as Fe and Mn. Guieu et al. (2018) used hydrological data recorded by Argo float in situ measurements, atlas data and simulations from a general ocean circulation model to argue that the high dissolved Fe concentrations may be sustained by a submarine source. They show that such Fe inputs could spread throughout the WTSP through mesoscale activity mainly westward through the South Equatorial Current, SEC (Fig. 1). Guieu et al. (2018) hypothesize that the high dissolved Fe concentrations in MA waters are due to shallow inputs of hydrothermal origin together with potential Fe input from islands themselves (Shiozaki et al., 2014). In our study, dissolved Fe concentrations were significantly positively correlated with N2 fixation and help to explain the distribution of N2 fixation rates measured across the OUTPACE transect.

In summary, our hypothesis to explain the spatial distribution of N2 fixation in MA waters is the following: when high DIP waters flow westward from the ETSP through the SEC and cross the South Pacific Gyre, N2-fixing organisms do not develop despite optimal SST (> 25 °C), likely because Fe concentrations are Fe-depleted (Moutin et al., 2008; Bonnet et al., 2008). When the high DIP, low DIN (dissolved inorganic N) waters from the gyre are advected west of the Tonga trench in Fe-rich and warm (> 25 °C) waters, all environmental conditions are fulfilled for diazotrophs to bloom extensively. According to Moutin et al. (2018), the strong depth difference between the nitracline and the phosphacline in MA waters associated with winter mixing allows a seasonal replenishment of DIP, which creates an excess of P relative to N and thus also favors N2 fixation in this region. Further investigations are required to better quantify Fe input from both islands and shallow volcanoes and associated hydrothermal activity along the Tonga volcanic arc for the upper mixed layer, study the fate of hydrothermal plumes in the water column at the local and regional scales, and investigate the potential impact of such hydrothermal inputs on diazotrophic communities at the scale of the whole WTSP.

N2 fixation rates were significantly negatively correlated with NO3 concentrations, consistent with the high energetic cost of N2 fixation compared to NO3 assimilation (Falkowski, 1983). They were also negatively correlated with depth and logically positively correlated with PAR and seawater temperature, two parameters which are depth dependent. Most N2 fixation took place in the surface mixed layer and rates were ~15 nmol N L−1 d−1 in MA waters with local maxima at stations SD1, SD6 and LDB and local minima at SD8 and SD10. *Trichodesmium*, the most abundant diazotroph enumerated at those stations (Stenegren et al., 2018), is buoyant, and horizontal advection is well known to result in patchy distributions of *Trichodesmium* in the surface ocean (Dandonneau et al., 2003), with huge surface accumulations named slicks, as observed during OUTPACE (Stenegren et al., 2018), surrounded by areas with lower accumulations. These physical processes may explain the differences between stations rather than local enrichments of nutrients due to islands as those three stations where the highest rates were measured are not located close to islands. However, the huge surface bloom observed at LDB (Fig. 1) and extensively studied by de Verneil et al. (2017) was mainly sustained by N2 fixation (secondary fueling picoplankton and diatoms, Caffin et al., 2018a), rather than deep nutrient inputs (de Verneil et al., 2017). This bloom had been drifting eastwards for several months and initially originated from Fiji and Tonga archipelagoes (https://outpace.mio.univ-amu.fr/spip.php?article160, last access: February 2016), which may have provided sufficient Fe to alleviate limitation and trigger this exceptional diazotroph bloom.

### 4.3 *Trichodesmium*: the major contributor to N2 fixation in the WTSP

In MA waters, the dominant diazotroph phyotypes quantified using nifH quantitative PCR assays were *Trichodesmium* spp. and UCYN-B (Stenegren et al., 2018), which commonly peaked at >10^6 nifH copies L−1 in surface (0–50 m) waters. DDAs (mainly het-1, but het-2 and het-3 were also detected) were the next most abundant diazotrophs (Stenegren et al., 2018). This result is consistent with the fact that abundances of those phyotypes co-varied and were significantly positively correlated with N2 fixation rates (Table 2). The two UCYN-A lineages (UCYN-A1 and UCYN-A2) were less abundant (<1.0–1.5% of total nifH copies, Stenegren et al., 2018) and not significantly correlated with N2 fixation rates (Table 2).

The relative contribution of different diazotroph phyotypes to bulk N2 fixation has been largely investigated through bulk and size fractionation measurements (usually comparing > and <10 µm size fraction N2 fixation rates), which may be misleading since some small-size diazotrophs are attached to large-size particles (Benavides et al., 2016; Bonnet et al., 2009) or form colonies or symbioses with diatoms (e.g., UCYN-B, Foster et al., 2011, 2013) and some diazotrophic-derived N released by diazotrophs is assimilated by small and large non-diazotrophic plankton (e.g., Bonnet et al., 2016a). Here we directly measured the in situ cell-specific N2 fixation activity of the two dominating diazotroph groups in MA waters: *Trichodesmium* and UCYN-B.

At all three studied stations, *Trichodesmium* dominated, accounting for 68.0–91.8% of the diazotroph community, followed by UCYN-B, accounting for 0.3–31.7%. In addition, *Trichodesmium* and UCYN-B had the highest measured gene expression (10^2–10^5 cDNA nifH copies L−1). It was not surprising that UCYN-B had a high gene expression given that the sampling time occurred later in the day (17:00–
21:00); however, both *Trichodesmium* and het-1 (which typically reduce N$_2$ and express *nifH* highest during the day, Church et al., 2005) had a detectable and often equally as high expression as UCYN-B. Cell-specific N$_2$ fixation rates reported here are on the same order of magnitude as those reported for field populations of *Trichodesmium* (Berthelot et al., 2016; Stenegren et al., 2018) and UCYN-B (Foster et al., 2013). *Trichodesmium* was always the major contributor to N$_2$ fixation, accounting for 47.1–83.8 % of bulk N$_2$ fixation, while UCYN-B never exceeded 6.1–10.1 %, despite accounting for $>30\ %$ of the diazotroph community at SD6. This may be linked with the lower $^{15}$N enrichment at SD6 (0.517 ± 0.237 at. %), which is due to a high proportion of inactive cells (at. % close to natural abundance) compared to SD2, where the majority of cells were active and highly $^{15}$N-enriched (1.163 ± 0.531 at. %). Such heterogeneity in N$_2$ fixation rates among UCYN-B-like cells has already been reported by Foster et al. (2013). Overall, these results show that the most abundant phylotype (*Trichodesmium*) accounts for the majority of N$_2$ fixation, but not in the same proportion, further indicating that the abundance of micro-organisms in seawater cannot be equated to activity, which has already been reported for other functional groups such as bacteria (Boutrif et al., 2011). In the North Pacific Gyre (station ALOHA), Foster et al. (2013) report a higher contribution of UCYN-B to daily bulk N$_2$ fixation (24–63 %) during the summer season, indicating that this group likely contributes more to the N budget at station ALOHA than in the WTSP, where *Trichodesmium* seems to be the major player.

5 Ecological relevance of N$_2$ fixation in the WTSP and conclusions

N$_2$ fixation was significantly positively correlated with Chl $\alpha$, PON, POC and BSi concentrations, as well as with primary production, suggesting a tight coupling between N$_2$ fixation, primary production and biomass accumulation in the water column. Based on our measured C : N ratios at each depth, the computation of the N demand derived from primary production measured during OUTPACE (Johnson et al., 2007) indicates that N$_2$ fixation fueled on average 8.2 ± 1.9 % (range 5.9 to 11.5 %) of total primary production in the WTSP. This contribution is higher than in other oligotrophic regions such as the northwestern Pacific (Shiozaki et al., 2013), ETSP (Raimbault and Garcia, 2008), northeastern Atlantic (Benavides et al., 2013), or the Mediterranean Sea (Bonnet et al., 2011; Ridame et al., 2014), where it is generally $<5\ %$. However, it is comparable to results found further north in the Solomon Sea (N$_2$ fixation fueled 9.4 % of primary production, Berthelot et al., 2017), which is part of the WTSP “hotspot” for N$_2$ fixation (Bonnet et al., 2017). Caffin et al. (2018b, a) show that N$_2$ fixation represents the major source (> 90 %) of new N to the upper photic (productive) layer during the OUTPACE cruise, before atmospheric inputs and nitrate diffusion across the thermocline, indicating that N$_2$ fixation supported nearly all new production in this region during austral summer conditions.

The large amount of N provided by N$_2$ fixation likely stimulated the growth of non-diazotrophic plankton as suggested by significant positive correlations between N$_2$ fixation rates and the abundance of *Prochlorococcus* spp., *Synechococcus* spp., heterotrophic bacteria and protists. $^{15}$N$_2$-based transfer experiments coupled with nanoSIMS experiments designed to trace the transfer of $^{15}$N in the planktonic food web demonstrated that $\sim10\ %$ of diazotroph-derived N is rapidly (24–48 h) transferred to non-diazotrophic phytoplankton (mainly diatoms and bacteria) in coastal waters of the WTSP (Bonnet et al., 2016a, b; Berthelot et al., 2016). The same experiments performed in offshore waters during the present cruise confirm that $\sim10\ %$ of recently fixed N$_2$ is also transferred to picophytoplankton and bacteria after 48 h (Caffin et al., 2018a). This is in accordance with Van Wambeke et al. (2018), who report that N$_2$ fixation fuels 40 to 70 % of the bacteria N demand in MA waters. This further demonstrates that N$_2$ fixation acts as an efficient natural N fertilization in the WTSP, potentially fueling subsequent export of organic material below the photic layer. Caffin et al. (2018a, b) estimated that the e-ratio, which quantifies the efficiency of a system in exporting particulate carbon relative to primary production (e-ratio = POC export/PP), was 3 times higher ($p < 0.05$) in MA waters compared to GY waters. Moreover, e-ratio values were as high as 9.7 % in MA waters, i.e., higher than the e-ratios in most studied oligotrophic regions (Karl et al., 2012; Raimbault and Garcia, 2008), where they rarely exceed 1 %, indicating that production sustained by N$_2$ fixation is efficiently exported in the WTSP. Diazotrophs were recovered in sediment traps during the cruise (Caffin et al., 2018a, b), but their biomass only accounted for $\sim5\ %$ (locally 30 % at LDA) of the N biomass in the traps, indicating that most of the export was indirect, i.e., after transfer of diazotroph-derived N to the surrounding planktonic communities that were subsequently exported. A $\delta^{15}$N budget performed during the OUTPACE cruise reveals that N$_2$ fixation supports exceptionally high (> 50 % and locally > 80 %) export production in MA waters (Knapp et al., 2018). Together these results suggest that N$_2$ fixation plays a critical role in export in this globally important region for elevated N$_2$ fixation.

The magnitude and geographic distribution of N$_2$ fixation control the rate of primary productivity and vertical export of carbon in the oligotrophic ocean; thus, accurate estimates of N$_2$ fixation are of primary importance for oceanographers to constrain and predict the evolution of marine biogeochemical carbon and N cycles. The number of N$_2$ fixation estimates have increased dramatically at the global scale over the past 3 decades (Luo et al., 2012). The results reported here show that some poorly sampled areas such as the WTSP provide unique conditions for diazotrophs to fix at high rates and contribute to the need to update current N$_2$ fixation estimates for
the Pacific Ocean. Further studies would be required to assess the seasonal variability of N₂ fixation in this region and perform accurate N budgets. Nonetheless, such high N₂ fixation rates question whether or not these high N inputs can balance the N losses in the ETSP. A recent study based on the N* (the excess of N relative to P) at the whole South Pacific scale Fumenia et al. (2018) reveals a strong positive N* anomaly (indicative of N₂ fixation) in the surface and thermocline waters of the WTSP, which potentially influences the geochemical signature of the thermocline waters further east in the South Pacific through the regional circulation. However, the WTSP is chronically undersampled, and a better description of the mesoscale and general circulation would be necessary to assess how N sources and sinks are coupled at the South Pacific scale.

**Data availability.** All data and metadata are available at the following web address: http://www.obs-vlfr.fr/proof/php/outpace/outpace. php, last access: June 2018.

**Author contributions.** SB designed the experiments; SB, MC, HB and MB carried them out at sea. MC, HB, RAF, SHN, CG and OG analyzed the samples; MC and SB analyzed the data. SB prepared the manuscript with contributions from all co-authors.

**Competing interests.** The authors declare that they have no conflict of interest.

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