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Longitudinal variability of size-fractionated N₂ fixation and DON release rates along 24.5°N in the subtropical North Atlantic

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1. Introduction

[1] Dinitrogen (N₂) fixation and dissolved organic nitrogen (DON) release rates were measured on fractionated samples (>10 μm and <10 μm) along 24.5°N in the subtropical North Atlantic. Net N₂ fixation rates (N₂ assimilation into biomass) ranged from 0.01 to 0.4 nmol N L⁻¹ h⁻¹, and DON release rates ranged from 0.001 to 0.09 nmol N L⁻¹ h⁻¹, DON release represented ~14% and ~23% of >10 μm and <10 μm gross N₂ fixation (assimilation into biomass plus DON release), respectively. This implies that by overlooking DON release, N₂ fixation rates are underestimated. Net N₂ fixation rates were higher in the east and decreased significantly toward the west (rₑ = −0.487, p = 0.002, and rₛ = −0.496, p = 0.001, for the >10 μm and <10 μm fractions, respectively). The sum of both fractions correlated with aerosol optical depth at 550 nm (AOD 550 nm) (rₛ = 0.382, p = 0.017) and phosphate (PO₄³⁻) concentrations (rₛ = 0.453, p = 0.018), suggesting an enhancement of diazotrophy as a response to aerosol inputs and phosphorus availability. In contrast, DON release was constant among size fractions and did not correlate with any of these variables. We also compared N₂ fixation rates obtained using the ¹⁵N₂ dissolved and bubble methods. The first gave average rates 50% (49% ± 39) higher than the latter, which supports the finding that previously published N₂ fixation rates are likely underestimated. We suggest that by combining N₂ fixation and DON release measurements using dissolved ¹⁵N₂, global N₂ fixation rates could increase enough to balance oceanic fixed nitrogen budget disequilibria.


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activity of widespread unicellular diazotrophic cyanobacteria (UCYN) has not been studied directly however [Mulholland, 2007]. This DON flux is generally unaccounted for in N\textsubscript{2} fixation studies, leading to potential underestimations of total N\textsubscript{2} fixation rates.

[5] Finally, global N\textsubscript{2} fixation rates can be miscalculated when regional estimates are extrapolated to larger oceanic basins or the global ocean. The North Atlantic Ocean is the basin where the greatest quantity of N\textsubscript{2} fixation and diazotrophic diversity data are available, and therefore N\textsubscript{2} fixation rates obtained here have been frequently used to calculate global oceanic rates. As we move forward in the investigation of oceanic N disequilibrium, more detailed spatial and temporal variability of diazotroph assemblages and their diazotrophic activity are needed.

[6] With the aim of covering the longitudinal variability of N\textsubscript{2} fixation rates in the subtropical North Atlantic and to avoid underestimations, we measured fractionated N\textsubscript{2} fixation and DON release rates at 40 stations over the 24.5° N parallel using the improved \textsuperscript{15}N\textsubscript{2} tracer technique [Gilbert and Bronk, 1994; Mohr et al., 2010].

2. Materials and Methods

2.1. Hydrographic Measurements and Nutrients

[7] Sampling was performed crossing the North Atlantic Ocean from the northwest (NW) African coast (Cape Jubi) to the Bahamas onboard the R/V Sarmiento de Gamboa from 27 January to 15 March 2011. The major part of the cruise was conducted over 24° 30’ N (World ocean circulation experiment—WOCE—section A05) (Figure 1a). Temperature, salinity, and fluorescence data were recorded with a SeaBird 911 plus conductivity, temperature and depth probe (CTD) equipped with redundant temperature and salinity sensors and a Sea-Tech fluorometer, all mounted on a General Oceanics 24 Niskin bottle rosette sampler. At each station, temperature and salinity were measured from the surface down to 20 m above the seafloor. Water samples were collected at each station in order to calibrate salinity values using a Guildline AUTOSAL model 8400B salinometer with a precision better than 0.002 for single samples.

[8] Samples for nutrient analyses were collected from the rosette Niskin bottles in 15 mL polypropylene tubes and immediately frozen until analysis ashore. The concentrations of nitrate (NO\textsubscript{3}^-), nitrite (NO\textsubscript{2}^-), NH\textsubscript{4}^+, phosphate (PO\textsubscript{4}^{3-}), and silicon (SiO\textsubscript{2}^-) were determined with a Technicon segmented-flow autoanalyzer. Standard methods were modified to obtain a detection limit of 2 nmol L\textsuperscript{-1} [Raimbault et al., 1990; Kerrouel and Aminot, 1997].

2.2. Sea Surface Height Anomalies and Atmospheric Dust

[9] Daily sea level anomaly (sea surface height anomaly (SSHA)) data (the difference between the total SSH and the average SSH for this time of year) were downloaded from the archiving, validation, and interpretation of
satellite oceanographic remote sensing service (AVISO, http://www.aviso.oceanobs.com/). The daily data were averaged for each three N2 fixation stations.

[10] Aqua-MODIS aerosol optical depth at 550 nm (AOD 550 nm) can be used as a proxy for dust presence in the atmosphere [Kaufman et al., 2005]. In order to assess the effect of atmospheric dust on N2 fixation rates, we used AOD 550 nm data obtained from the National Aeronautics and Space Administration (NASA) Goddard Earth Sciences Data and Information Services Center Giovanni (NASA GES DISC) online database. The spatial distribution of AOD 550 nm during our cruise is plotted in Figure 1b.

2.3. Fractionated Net N2 Fixation and DON Release Rates

[11] Rates of N2 fixation and DON release were measured in the > 10 μm and < 10 μm size fraction of samples collected from the surface (~5 m). The size fractionation was done at the start of the incubation. We sampled one station per day but alternated between day (0900–1200) and night samplings (1900–2100, local time) (Figure 1a). This approach allowed us to study the diazotrophic activity of organisms that fix N2 in the light (e.g., *Trichodesmium*), and organisms that only fix N2 in the dark to avoid oxygen deactivation of the nitrogenase enzyme system (e.g., *Crocosphaera*).

[12] N2 fixation and DON release rates were measured using 15N-labeled N2 gas during 3–4 h incubations. There are two general approaches used to add the labeled gas to a sample: the addition of water that was saturated with 15N2 (e.g., Gilbort and Bronk, 1994; Mohr et al., 2010), and the addition of a bubble [Montoya et al., 1996]. In this study, we used the addition of saturated water in fractionated seawater samples and the bubble method in whole (unfiltered) seawater samples (see below). With the addition of a bubble, the N2 fixation rate is potentially underestimated due to slow dissolution of the gas bubble in water [Mohr et al., 2010].

[13] To prepare the 15N2-saturated water, surface seawater (~5 m) was recovered from the flow-through system of the ship and filtered through a 47 mm GF/F filter. This was done during the upcast of the CTD to ensure that the 15N2-saturated water was the same as the sample water. This filtered seawater was decanted into 0.5 L transparent polycarbonate bottles (Nalgene) and degassed as outlined in Mohr et al. [2010]. Each bottle was filled to overflow, closed with a septum screw-cap and 5 mL of 15N2 (99 at % 15N; Cambridge Isotope Laboratories) were injected using a Hamilton gas-tight syringe. The bottles were vigorously shaken for 10–20 min until the bubble was fully dissolved and then kept in the on-deck incubators until the rosette was back on board (0.5–2.5 h depending on the station depth) to ensure that the 15N2-enriched seawater had the same temperature as the sample at the time of mixing, in order to protect the organisms from thermal shocks.

[14] To check the real 15N enrichment of the 15N2-amended seawater added to the samples, replicate samples of the 15N2-amended seawater prepared on board were taken in 10 mL crimp vials (Chrompack), filled to overflow, and sealed with teflon-lined stoppers and aluminum caps using a seal crimper. These were stored at room temperature in the dark until being analyzed by membrane inlet mass spectrometry (MIMS) in S. Joye’s lab. MIMS analysis showed that the real enrichment of the 15N2-amended seawater was ~100% (99 ± 2%) of that expected from complete dissolution of the 15N2 bubble.

[15] To collect water used to measure fractionated rates, near surface (~5 m) seawater was collected with a 30 L Niskin bottle at each station. The samples were prefractionated by filtering two 2 L replicates through a 10 μm Nitex custom-made sieve. The <10 μm fraction was recovered in 2.4 L transparent polycarbonate bottles (Nalgene). The >10 μm fraction was recovered from the sieve by gently washing and concentrating the biological material using a water sprayer filled with filtered (GF/F) seawater from the same station. Then, 200 mL of 15N2-enriched filtered seawater was added to all bottles, they were filled to the top with filtered seawater and placed in the on-deck incubators for 3–4 h. The incubators were connected to the ship’s flow-through system and covered with neutral density screens (Lee Filters). After the incubation, the samples were filtered through precombusted 25 mm GF/F filters, which were subsequently stored in sterile cryovials (VWR) and frozen. The GF/F filters were then filtered through 0.2 μm polycarbonate filters (GE-Osmotics Porletes) using gentle vacuum pressure (<100 mm Hg) to remove bacteria prior to isolating the dissolved N pools. Rigorous care was taken to avoid light stress in the samples. The filtrates were stored frozen in triplicate 50 mL polypropylene tubes (VWR) and used to measure the concentrations of NO3−, NO2−, NH4+, and total dissolved N (TDN), and the 15N atom % enrichment of NO3−, NH4+, and TDN as described in section 2.5.

2.4. Whole Seawater Net N2 Fixation Rates

[16] Parallel to seawater collection for fractionation, whole surface seawater was transferred from the 30 L Niskin bottle to 2.4 L transparent polycarbonate bottles. The bottles were completely filled using silicone tubing to prevent the introduction of air bubbles. They were then sealed with septum screw caps before 2 mL of 15N-labeled N2 gas were injected through the septum. The pressure across the septum was equilibrated by allowing the excess water to escape through a sterile syringe tip piercing the septum. The bottles were placed in the on-deck incubator for the same incubation period. After the incubation, samples were also filtered through precombusted GF/F filters, stored in cryovials, and frozen until isotope ratio mass spectrometer (IRMS) analysis ashore. N2 fixation rates measured with the bubble method were calculated as outlined in Montoya et al. [1996].

2.5. Chemical Analyses and Rate Calculations

[17] Nutrient pools in filtrates (surface seawater samples incubated with 15N2 and filtered through GF/F and 0.2 μm, see above) were analyzed in D. Bronk’s lab after the cruise. Concentrations of NH4+ were analyzed in duplicate with the manual phenol hypochlorite method [Hansen and Koroleff, 1983] using a Shimadzu UV-1601 spectrophotometer (detection limit of ~0.03 μmol L−1). Concentrations of NO3− and NO2− were determined with a Lachat QuikChem 8500 autoanalyzer (detection limits of 0.1 and 0.03 μmol L−1, respectively). TDN concentrations were analyzed on a Shimadzu TOC-V equipped with a total nitrogen module (TNM). DON concentrations were calculated by subtracting NO3−, NO2−, and NH4+ concentrations from TDN concentrations; propagation of error calculations were used to estimate errors associated with DON concentrations [Bronk et al., 2000].
was then converted to nitrous oxide (N\textsubscript{2}O, N\textsubscript{2}H\textsubscript{4}/C\textsubscript{2}H\textsubscript{3}], [NH\textsubscript{2}O\textsubscript{2} atom % excess target pool

In contrast, SSS

TDN pool, we used

atom % NO\textsubscript{3} fixation rates. The

atom %NO\textsubscript{3} fixation rates [Dudek et al., 2001]. Isotope ratios of N\textsubscript{2}O were then measured using a ThermoFinnigan GasBench + PreCon trace gas concentration system interfaced to a ThermoScientific Delta V Plus IRMS at the University of California Davis Stable Isotope Facility (Davis, CA). The same approach without the persulfate oxidation step was used to isolate the NO\textsubscript{3} pool. The atom % enrichment of TDN and NO\textsubscript{3} fixation rates obtained from

atom %TDN pool, 15 mL subsamples of the filtrate were oxidized to NO\textsubscript{3}\textsuperscript{−} via persulfate oxidation [Valderrama, 1981]. The NO\textsubscript{3} was then converted to nitrous oxide (N\textsubscript{2}O\textsubscript{4}) using denitrifying bacteria lacking N\textsubscript{2}O-reductase activity [Sigman et al., 2001]. Isotope ratios of N\textsubscript{2}O were then measured using a ThermoFinnigan GasBench + PreCon trace gas concentration system interfaced to a ThermoScientific Delta V Plus IRMS at the University of California Davis Stable Isotope Facility (Davis, CA). The same approach

where atom% excess is the

\begin{equation}
\text{Atom}\%\text{DON} = \frac{\text{(atom}\%\text{TDN} \times [\text{TDN}]) - \left( (\text{atom}\%\text{NO}_3^- \times [\text{NO}_3^-]) + (\text{atom}\%\text{NH}_4^+ \times [\text{NH}_4^+]) \right)}{[\text{DON}]}
\end{equation}

[20] Solving equation (1), the \text{^{15}N} atom % enrichment of the DON pool was calculated as follows:

\begin{equation}
\text{Rate} = \left( \frac{\text{atom}\%\text{excess target pool}}{\text{atom}\%\text{excess source pool} \times \text{time}} \right) \times \text{target pool concentration}
\end{equation}

where atom% excess is the

\text{^{15}N} enrichment over natural abundance.

\text{^{15}N} N\textsubscript{2} fixation and DON release rates were then calculated using standard tracer equations:

were observed near the coastal upwelling off NW Africa (Figure 2). From east to west, SST increased up to 24.26\textdegree C at 54.03 W, and SSS up to 37.54 at 37.57 W. Moving westward, SST presented sharp decreases of 1–2\textdegree C coincident with strong negative SSHA (<−12 cm). In contrast, SSS decreased until 62\textdegree W (station 104), where it reached 36.28. From 62\textdegree W to the west, SST increased in two steps coincident with positive or low values of SSHA, and finally reached ~36.74.

\text{^{15}N} The surface concentrations of NO\textsubscript{3}\textsuperscript{−} + NO\textsubscript{2}\textsuperscript{−}, PO\textsubscript{4}\textsuperscript{3−}, and SiO\textsubscript{2} were generally maximum at the eastern end of the transect (coinciding with the NW African coastal upwelling), and decreased toward the west. The lowest concentrations were observed west of ~45° W, coinciding with the oligotrophic Sargasso Sea.

Along with SST and SSS increments and high values for nutrients, the first seven N\textsubscript{2} fixation stations showed positive SSHA values, which then decreased considerably until station 71 (44.5° W). From this position to the west, high and predominantly negative SSHA dominated.

3.1. Hydrographic Features and Nutrient Concentrations

Low values of sea surface temperature (SST) and sea surface salinity (SSS) of ~19\textdegree C and 36.8, respectively,
3.2. Dissolved $^{15}\text{N}_2$ Versus $^{15}\text{N}_2$ Bubble Net $\text{N}_2$ Fixation Rates

The sum of net $\text{N}_2$ fixation rates in the >10 $\mu$m and <10 $\mu$m fractions measured using the dissolved method [Mohr et al., 2010] was compared to net $\text{N}_2$ fixation rates (in whole seawater samples collected at the same stations) using the $^{15}\text{N}_2$ bubble method [Montoya et al., 1996]. The difference between both rates tended to increase toward the east (Figure 3a). Indeed, the percent underestimation of the bubble method (average 49 ± 39%; range 6–93%) was significantly correlated with longitude ($r_s = 0.518$, $p = 0.001$). The concentrations of DON in >10 $\mu$m and <10 $\mu$m samples did not correlate significantly with the percentage underestimation of $\text{N}_2$ fixation rates, but followed a similar longitudinal distribution (Figure 3b).

3.3. $\text{N}_2$ Fixation and DON Release Estimated by the Dissolved $^{15}\text{N}_2$ Method

Both >10 and <10 $\mu$m organisms presented a similar range of $\text{N}_2$ fixation rates (~0.01 to ~0.44 nmol N L$^{-1}$ h$^{-1}$), although >10 $\mu$m rates were statistically different from <10 $\mu$m rates (Wilcoxon test, $p < 0.0001$). On average, net $\text{N}_2$ fixation rates of organisms >10 $\mu$m were slightly higher than those <10 $\mu$m (0.16 and 0.1 nmol N L$^{-1}$ h$^{-1}$, respectively). Both rates showed a tendency to decrease toward the west (Figure 4), showing significant relationships with longitude (Spearman’s correlation rank coefficient $r_s = 0.487$, $p = 0.002$, and $r_s = 0.496$, $p = 0.001$, for the >10 $\mu$m and <10 $\mu$m fractions, respectively). $\text{N}_2$ fixation rates of organisms >10 $\mu$m measured during the day were not significantly different from those measured during the night (Wilcoxon test, $p = 0.936$), while the differences between rates of organisms <10 $\mu$m measured during the day and the night were almost statistically significant (Wilcoxon test, $p = 0.059$). In most stations, $\text{N}_2$ fixation rates measured during the night were higher than those measured during the day for both >10 $\mu$m and <10 $\mu$m fractions (Figure 4).

DON release rates in organisms >10 $\mu$m and <10 $\mu$m ranged from 0.001 to ~0.09 nmol N L$^{-1}$ h$^{-1}$ (Figure 5). Considering gross $\text{N}_2$ fixation as net $\text{N}_2$ fixation plus DON release [Mulholland et al., 2004] DON release represented ~14% of >10 $\mu$m gross $\text{N}_2$ fixation and ~23% of <10 $\mu$m gross $\text{N}_2$ fixation (data not shown). Significant differences in DON release were not found between >10 $\mu$m and <10 $\mu$m fractions, neither between day and night (Wilcoxon test, $p > 0.05$). Also, DON release rates did not show a clear trend with longitude ($r_s = 0.203$, $p = 0.391$, and $r_s = 0.046$, $p = 0.848$, for the >10 $\mu$m and <10 $\mu$m fractions, respectively).

4. Discussion

4.1. Dissolved Versus Bubble Methods

Using the dissolved $^{15}\text{N}_2$ method, in this study we have measured $\text{N}_2$ fixation rates on average 50% higher than those estimated using the $^{15}\text{N}_2$ bubble method. This difference between the two methods is at the lower end of the few comparisons available in the literature. Mohr et al. [2010] observed that $\text{N}_2$ fixation rates in cultures of...
Crocosphaera watsonii were 40% higher when the dissolved method was used, compared to the bubble method. Recently, Wilson et al. [2012] compared both methods in natural waters of the North Pacific Ocean (station ALOHA), and obtained rates that were 2–3.5 fold higher with the dissolved method than they were with the bubble method. Großkopf et al. [2012] observed that both methods differ by 62% when the diazotrophic community is dominated by Trichodesmium, and up to 570% when dominated by diazotrophs other than Trichodesmium (symbionts of diatoms, UCYN, heterotrophic diazotrophs). Adding to differences in the diazotrophic community, the factors influencing N$_2$ fixation differences between the dissolved and bubble methods include (1) temperature, which affects gas dissolution into water, (2) the agitation of the incubation bottles, (3) the volume of the incubation bottle, (4) the volume of $^{15}$N$_2$ injected, (5) the duration of the incubation, (6) the time the incubation starts relative to the onset of nitrogenase activity, which differs among diazotroph species, and (7) the DOM coating of the $^{15}$N$_2$ bubble [Mohr et al., 2010]. These factors vary widely among the previously published N$_2$ fixation data measured using the bubble method, which makes the recalculation of rates difficult if not impossible [Großkopf et al., 2012].

[30] This study does not include diazotrophic organisms’ abundance and distribution data, so we cannot analyze how this affected the percentage underestimation of the bubble method with regards to the dissolved method. However, a recent compilation of diazotroph abundance data [Luo et al., 2012] shows that Trichodesmium clearly dominates in the tropical Northwest Atlantic, while UCYN are somewhat more abundant in the eastern than in the western side of the basin and at higher latitudes. This distribution of diazotrophs and percentage underestimation by the bubble method versus the dissolved method agrees with Großkopf et al. [2012], although our percentage underestimation values are much lower. During our cruise, Mompéan et al. [2013] measured the abundance of Trichodesmium using vertical net tows. These authors observed that Trichodesmium was almost absent from the beginning of the transect (NW African coast) until ~25 W, followed by a peak of ~15 trichomes L$^{-1}$ at ~30 W, and decreased toward the west. This longitudinal distribution is contrary to the arguments of Großkopf et al. [2012]. However, the low abundance of Trichodesmium recorded suggests that their role in the diazotrophic activity measured in our experiments was minor.

[31] With regards to the potential effect of DOM in slowing the dissolution of the $^{15}$N$_2$ bubble, we checked for any significant correlations between percentage underestimations and DON concentrations. Correlations between DON and the difference between the dissolved and bubble methods were not statistically significant (data not shown), but their longitudinal trend was similar. We observed that the differences between the dissolved and bubble methods were greater in the eastern part of the transect, coinciding with higher DON concentrations (Figure 3b). The difference between both methods would have probably correlated better with dissolved organic carbon (DOC) since it is a better proxy for the total DOM pool. Oceanic DOC concentrations range from 34 to >90 μM, while DON concentrations range from 2 to 13 μM [Hansell and Carlson, 2002]. Unfortunately, DOC data are not available in this case.

[32] Another factor specific to this study that may have influenced differences between both methods is the prefractionation of samples prior to incubation with $^{15}$N$_2$. The sum of fractionated N$_2$ fixation rates (>10 μm + <10 μm) may be not directly comparable to whole seawater N$_2$ fixation rates, and sample fractionation is not always successful. For example, <10 μm organisms may be retained in the >10 μm fraction due to clogging of the mesh used for fractionation. However, due to the oligotrophic character of the area of study, seawater samples flowed easily through the mesh used for fractionation, and thus significant clogging or cell disruption may have not been significant.

[33] Finally, we must consider that the short incubation period used in our study (3–4 h) diminishes the dissolution of the $^{15}$N$_2$ bubble in the seawater sample in comparison with 24 h incubations conducted in other studies, which together with the large standard deviation found between methods (% underestimation = 49 ± 39%; see section 3) suggests that the underestimation of rates when using the $^{15}$N$_2$ bubble method could have been less severe if a longer incubation period had been chosen. As an alternative to the preparation of $^{15}$N$_2$-enriched seawater to conduct N$_2$ fixation assays and to avoid any uncertainties related to this new method until it is definitely widely proven and a general protocol is established, we recommend to monitor the equilibration of the $^{15}$N$_2$ bubble during the incubation period in order to calculate accurate N$_2$ fixation rates.

4.2. DON Release

[34] DON release rates were not significantly different between fractions, nor were they different between day and night, or significantly correlated with longitude. This lack of spatial and daily variability in both size fractions suggests that the release of a percentage of recently fixed N$_2$ is inherent to natural assemblages of marine diazotrophs, although we cannot discern if this release was active or passive, via predation, viral infection, or cell death.

[35] We must note that the DON release rates presented here may be biased by the methodological approach used. The fractionation of seawater samples into >10 μm and <10 μm fractions prior to incubation with $^{15}$N$_2$ could have disrupted cells, artificially inflating DON release rates. However, the differences between prefractionation and postfractionation NH$_4^+$ concentration—an indicator of cell breakage [Bronk and Gilbert, 1993]—were not significant for either fraction (t-test, p = 0.92 and p = 0.13 for the >10 μm and <10 μm fractions, respectively, data not shown), suggesting that the DON release rates presented here are robust. Another factor potentially influencing the DON release rates presented here is the disruption of trophic interactions between different sized microorganisms occasioned when incubated separately [Havens, 2001]. If the separate incubation of >10 μm and <10 μm diazotrophs had any effect on their DON release rates, we could not quantify it. Prefractionation has been previously used in a number of studies [e.g., Bronk and Gilbert, 1993; Ohlendiek et al., 2000; Benavides et al., 2011] and has been used here because it is the only possible approach to measure fractionated DON release rates.
Previous studies of diazotrophic DON release have focused on the filamentous cyanobacterium *Trichodesmium* [Gilbert and Bronk, 1994]. These authors found that *Trichodesmium* released 50% of their recently fixed N\textsubscript{2} as DON, implying that true N\textsubscript{2} fixation rates were considerably underestimated if DON release was not taken into account. The release of DON by other diazotrophs has not been studied directly in the field and therefore it is difficult to put the results obtained here into context. However, when our hourly surface DON release rates are integrated to 100 m and scaled up to days, the average integrated DON release rates obtained (up to 0.75 µmol N m\textsuperscript{-2} d\textsuperscript{-1}) are in the range of the N\textsubscript{2} fixation in the <0.7 µm fraction (seawater filtered through a GF/F) obtained by Konno et al. [2010]. Nonetheless, we must note that the DON release rates measured here result from the quantification of the 15N atom\% enrichment of the DON pool (in filtrates of samples passed through GF/F filters and 0.2 µm PC filters) and not of any particles (organic or inorganic) <0.7 µm. In another attempt to measure DON release rates using axenic cultures of *Cyanothecapsa* sp. Miami BG 043511 grown in optimum conditions, we found that this organism releases only ~1% of their recently fixed N\textsubscript{2} as DON [Benavides et al., 2013]. This is much lower than the ~23% DON release obtained in this study for the <10 µm fraction. However, *Cyanothecapsa* or other closely related UCYN belonging to group C (UCYN-C) diazotrophs are less frequent in the North Atlantic Ocean, and therefore it can be expected that the <10 µm diazotroph community in this transect was more likely composed of *Crocophaspha* (UCYN group B—UCYN-B), UCYN of group A (UCYN-A), and heterotrophic diazotrophs [Langlois et al., 2008]. The DON release activity of those diazotrophs is unknown [Mulholland, 2007], but some evidence is accumulating. The difference between gross N\textsubscript{2} fixation rates obtained using the acetylene reduction assay and net N\textsubscript{2} fixation rates obtained using the 15N\textsubscript{2} method has been used as a proxy to estimate the release of recently fixed N\textsubscript{2} [Mulholland et al., 2004]. Benavides et al. [2011] measured both rates on fractionated samples (>10 µm and <10 µm) over the Canary Current and observed that the <10 µm fraction potentially released ~60% of their recently fixed N\textsubscript{2}. In this study, we measured the 15N enrichment of the DON pool and found that both the >10 µm and <10 µm fractions DON release rates represent a significant percentage of gross N\textsubscript{2} fixation. Now that the correction of N\textsubscript{2} fixation rates using the dissolved 15N\textsubscript{2} method is in the spotlight, we call attention to the inclusion of DON release measurements in routine field samplings. The application of the dissolved method combined with DON release rates could perhaps raise global N\textsubscript{2} fixation rates enough to balance denitrification, balancing the oceanic fixed N budget.

### 4.3. Longitudinal Variability of N\textsubscript{2} Fixation

In contrast with the general trend of higher N\textsubscript{2} fixation rates in the western Atlantic than in the eastern Atlantic observed in previous studies [Montoya et al., 2007; Liu et al., 2012], in this study both >10 µm and <10 µm N\textsubscript{2} fixation rates were highest close to the NW African coast and decreased westward (Figure 4). This longitudinal variability could be partly affected by nutrient availability in the upper ocean controlled by diffusion of nutrient-rich deep waters through the thermocline. In the Northern hemisphere, anticyclonic eddies promote positive SSAH, deepening the thermocline and hence enhancing organic matter accumulation and bacterial production within their cores [e.g., Balart et al., 2010]. On the contrary, cyclonic eddies foster negative SSAH, which uplifts the thermocline and enhances the upwelling of deeper nutrient-rich cold waters, enhancing primary production and chlorophyll concentrations [e.g., Aristeigui et al., 1997]. In contrast to autotrophic phytoplankton species, which depend on NO\textsubscript{3} upwelled from the deep sea or NH\textsubscript{4} regenerated in situ for their growth, diazotrophic organisms are capable of growing on N\textsubscript{2} as the only source of N. Theoretically, this sets out a different scenario for diazotrophs, which would grow better in the core of warm and nutrient-poor anticyclonic eddies, than in that of cold and nutrient-rich cyclonic eddies. Church et al. [2009] studied a three-year series of SSAH values at station ALOHA and compared it to N\textsubscript{2} fixation rates and nifH gene diversity. These authors observed that pronounced positive SSAH values coincided with enhanced N\textsubscript{2} fixation rates and higher temperatures. However, these authors also pointed out that negative SSAH could promote the upwelling of waters enriched in PO\textsubscript{4}\textsuperscript{3–} and SiO\textsubscript{2} relative to fixed N, which would enhance the growth of phosphorus-limited diazotrophs and diatom-diazotroph associations (DDAs) limited by SiO\textsubscript{2}. In order to test the hypothesis by Church et al. [2009], we operationally divided our transect into two broad areas: an area dominated by weak positive or nearly neutral SSAH values (termed +SSAH zone, east of ~45°W), and an area where strong negative SSAH values predominated (termed −SSAH zone, west of ~45°W) (Figure 2a). Similarly to Church et al. [2009], N\textsubscript{2} fixation rates were higher where positive SSAH values predominated (+SSAH zone), and lower coinciding with negative SSAH values (−SSAH zone). Indeed, net N\textsubscript{2} fixation rates in >10 µm and <10 µm organisms were significantly different in the two zones (Wilcoxon test, p = 0.01 and p = 0.005, respectively). Net N\textsubscript{2} fixation rates of the <10 µm fraction were approximately twofold higher in the +SSAH zone than in the −SSAH zone (Table 1).

#### Table 1. Fractionated Net N\textsubscript{2} Fixation Rates, DON Release Rates, and the Percent Contribution of DON Release to Gross N\textsubscript{2} Fixation

<table>
<thead>
<tr>
<th>Zone</th>
<th>Net N\textsubscript{2} Fixation</th>
<th>DON Release</th>
<th>% DON Release</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>&gt;10 µm</td>
<td>&lt;10 µm</td>
<td>&gt;10 µm</td>
</tr>
<tr>
<td>+SSHA</td>
<td>0.19 ± 0.10</td>
<td>0.14 ± 0.10</td>
<td>0.02 ± 0.02</td>
</tr>
<tr>
<td>−SSHA</td>
<td>0.12 ± 0.07</td>
<td>0.06 ± 0.03</td>
<td>0.02 ± 0.01</td>
</tr>
</tbody>
</table>

*Rates are in nmol L\textsuperscript{−1} h\textsuperscript{−1} Values represent the average and standard deviation (average ± SD). The +SSHA zone comprises stations 1–71, or longitude 13.34–44.5° W. The −SSHA zone comprises stations 77–139, longitude 44.5–76.15° W.*
suggesting that small diazotrophs predominated in the eastern North Atlantic during our sampling. Notwithstanding, the longitudinal tendency of nutrient concentrations observed is opposite to that expected from nutrient diffusion as a consequence of rising or deepening of the thermocline, that is, greater concentrations were observed in the +SSHA zone than in the −SSHA zone (Figure 2c), which covers the oligotrophic Sargasso Sea. Differences in the response of diazotrophic activity to SSHA variability and consequent nutrient inputs to the upper ocean layers between the work of Church et al. [2009] and this study might stem from differential atmospheric inputs in the Pacific and Atlantic Oceans. Voss et al. [2004] also obtained greater N2 fixation rates toward the west over 10 N in the north Atlantic. These authors performed their cruise in autumn, when the intertropical convergence zone (ITCZ) reaches its northernmost position enhancing Saharan dust deposition. Our cruise took place in the late winter (January–March 2011), when dust inputs are also known to be maximum [Torres-Padrón et al., 2002]. Similar to Fernández et al. [2010], we found that total net N2 fixation rates (>10 μm plus <10 μm) correlated significantly with AOD 550 nm (r = 0.382, p = 0.017), suggesting that nutrients made available through Saharan dust deposition enhance N2 fixation rates.[38] Latitudinal gradients in N2 fixation rates seem to be more abrupt than longitudinal ones. Moore et al. [2009] and Fernández et al. [2010] observed N2 fixation rates up to ~5 and ~10 nmol L−1 d−1 (respectively) in the North Atlantic, which decreased to values ~0 in the South Atlantic. Both authors attributed these latitudinal gradients to differences in dissolved iron availability between the North and South Atlantic basins. Longitudinal gradients in N2 fixation rates in the North Atlantic are smoother, ranging from ~0 to 2.5 nmol N L−1 d−1 [Voss et al., 2004; Montoya et al., 2007]. Alternative to the role of iron in shaping spatial trends in N2 fixation rates, Fernández et al. [2012] found that under similar dust deposition regimes, the longitudinal variability of N2 fixation rates may be controlled by phosphorus availability instead. This pattern agrees with the longitudinal trend of PO43− concentrations observed here (Figure 2c) and, indeed, PO43− concentrations were significantly correlated with total (>10 μm + <10 μm) N2 fixation rates (r = 0.453, p = 0.013). The agreement between the longitudinal distribution of N2 fixation rates, AOD 550 nm values and PO43− concentrations supports the iron and phosphorus colimitation hypothesis proposed by Mills et al. [2004] for the North Atlantic Ocean.

[39] The data compilation of Luo et al. [2012] shows that small diazotrophs belonging to UCYN groups A, B, and C are more abundant in the eastern than in the western basin of the North Atlantic and consequently their contribution to total N2 fixation rates increase to the east, where they dominate over larger diazotrophs such as Trichodesmium [Voss et al., 2004; Montoya et al., 2007; Benavides et al., 2011]. Instead, Trichodesmium blooms are recurrent in the western North Atlantic where the highest N2 fixation rates in this basin are found [Luo et al., 2012]. In contrast, during this cruise the abundance of Trichodesmium was higher in the eastern side of the basin and decreased toward the west [Mompeán et al., 2013]. However, the maximum Trichodesmium abundances observed are between one and three orders of magnitude lower than observed in previous studies conducted at lower latitudes in the North Atlantic [Luo et al., 2012]. The high rates of >10 μm net N2 fixation observed between the African coast and 20 W during our study (Figure 4a) could not have been supported by the low Trichodesmium abundances (~0) measured by Mompeán et al. [2013] in the same longitudinal range, and therefore we infer a low contribution of this genus to the observed N2 fixation rates in the eastern part of the transect. The contribution of Trichodesmium to total N2 fixation in the rest of the transect (from 20 W to the Bahamas) is uncertain, but given their low abundance we do not believe that this cyanobacterium played an important role in the N2 fixation rates measured here. In contrast, Mompeán et al. [2013] argue that N2 fixation by Trichodesmium largely supported the δ15N low values observed in different size fractions of the planktonic community during our cruise. However, low δ15N signals in the planktonic biomass associated to N2 fixation may be detected in time scales of weeks to months after the N2 was actually biologically fixed, and thus the results obtained by Mompeán et al. [2013] cover a time scale larger than ours (3–4 h incubations). Moreover, the low δ15N values observed could have been influenced by the deposition of anthropogenic N, a flux which is rapidly approaching the magnitude of oceanic N2 fixation rates [Duce et al., 2008].

[40] Finally, we note that the magnitude of the N2 fixation rates measured here is difficult to compare with other studies due to the 3–4 h incubation period used (compared to the usual 12–24 h incubations). Moreover, our measurements correspond only to the surface and therefore are not directly comparable to depth-integrated N2 fixation rates in other studies. The vertical variability of the composition of the diazotrophic community associated with the vertical gradients of light, oxygen, temperature and nutrient availability translates into a vertical variability in N2 fixation rates. It is generally observed that N2 fixation rates are highest at the surface and decrease with depth [e.g., Church et al., 2009; Luo et al., 2012]. Therefore, although the surface N2 fixation rates presented here are not representative of the whole upper ocean layer, they are still valuable to study the longitudinal variability of diazotrophic activity at a large scale.

5. Conclusions

[41] The results obtained here suggest that the longitudinal variability of N2 fixation rates in the North Atlantic is affected by atmospheric nutrient inputs and PO43− availability, which are maximum in its eastern basin and decrease toward the west. In contrast to net N2 fixation rates, the release of recently fixed N2 as DON did not show any longitudinal trend or variability among size fractions. This suggests that this process is inherent of natural assemblages of diazotrophic organisms and should not be overlooked in order to avoid underestimation of true N2 fixation rates. Moreover, we have added to the body of evidence that N2 fixation rates are substantially underestimated when the bubble method is used. Measurements of N2 fixation rates using the dissolved method combined with measurements of DON release in future samplings will probably diminish global N budget unbalances.
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