Trichodesmium spp. physiology and nutrient fluxes in the North Pacific subtropical gyre

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ABSTRACT: The potential role of the diazotrophic cyanobacterium Trichodesmium spp. in nitrogen and phosphorus dynamics of the euphotic zone of the North Pacific subtropical gyre was investigated as one component of the Hawaii Ocean Time-series (HOT) program. Experiments were conducted with natural samples collected at Stn ALOHA (22° 45' N, 158°W) and with isolated cultures in laboratory. In both sets of experiments, we documented aerobic nitrogenase activity in Trichodesmium by acetylene (C2H2) reduction to ethylene (C2H4). Although average C2H4 evolution per unit chlorophyll a (chl a) was lower in naturally occurring single trichomes relative to colonies [3.9 vs 12.5 nmol C2H4 (pg chl a)-1 h-1, respectively], the generally greater biomass of single trichomes in the North Pacific Ocean suggests that trichomes may be important in the oceanic N cycle. Disrupted colonies display the lowest nitrogenase activities, but these activities increase with time in cultures. These observations and the relatively high dark oxygen consumption rates observed for Trichodesmium [0.18 μmol O2 (pg chl a)-1 h-1] suggest that, in nature, this cyanobacterium may be able to protect nitrogenase from oxygen inactivation, and that colony formation enhances, but is not prerequisite for, nitrogenase activity. Trichodesmium spp. collected from different depth strata at Stn ALOHA were also used to study variations in the C:N:P elemental composition of rising and sinking colonies. Although changes in elemental ratios were small, the relative C:N increase in all sinking colonies and the N:P decrease in rising colonies, sampled at approximately 100 m depth, is consistent with the model of Trichodesmium storage of carbohydrate in shallow waters (<20 m) and uptake of P at depth. The active uptake of inorganic phosphorus measured in sinking colonies incubated in the dark combined with a change toward positive buoyancy in colonies during the incubation supports the hypothesis that vertical migrations of Trichodesmium may represent an upward transport of P into the euphotic zone and a potential decoupling of N and P nutrient cycles. However, these results do not explain the large concentration of non-migratory single trichomes observed in the upper water column of Stn ALOHA, unless colony versus free trichome morphology is a transient condition that is under cellular control.

KEY WORDS: Nitrogen fixation - Trichodesmium - Nutrients - North Pacific

INTRODUCTION

The high rates of photoautotrophic productivity in the euphotic zone of the North Pacific subtropical gyre appear to be inconsistent with the relatively low rates of inorganic nutrient input from beneath the permanent thermocline (King 1986, Lewis et al. 1986, Platt et al. 1989). A significant increase in measured primary productivity during years when the frequency of deep mixing events decreases (Karl et al. 1995, Letelier et al. 1996) further exacerbates this condition and demands a closer examination of our basic paradigms of primary production and controls of nutrient fluxes in this oligotrophic ecosystem.

In recent years attention has been focused on other potential sources of inorganic nutrients including atmospheric depositions (Donaghay et al. 1991, Duce et al. 1991), stochastic storm-induced mixing (DiTullio & Laws 1991), dinitrogen fixation (Capone & Carpenter 1982, Karl et al. 1992) and vertical migrations of plant or animal communities (Villareal et al. 1993). In this context, the cyanobacterium Trichodesmium spp. has several characteristics relevant to the study of nutrient
cycling in subtropical pelagic ecosystems. Two of these characteristics, directly related to the input of nutrients into the euphotic zone, are: (1) the faculty of dinitrogen fixation (Goering et al. 1966, Carpenter 1973, 1983, Wada & Hattori 1991) and (2) the capacity of regulating cell density to effect a migration between the sea surface and the nutricline (Walsby 1978, 1992, Gainf & Oliver 1982).

Although nitrogenase activity in *Trichodesmium* is well documented, the quantitative importance of N$_2$ as a source of new nitrogen for the marine pelagic biota is uncertain (Carpenter & Capone 1983, Codispoti & references therein, Carpenter & Romans 1991). Dinitrogen fixation should represent an important fraction of new production in these regions unless growth is limited by nutrients other than nitrogen, as proposed by Martin & Fitzwater (1988), or nitrogenase activity is inhibited by physical or chemical conditions in the environment (Doremus 1982, Reuter 1982, 1988, Howarth & Cole 1985, Paerl & Prufert 1987, Marino et al. 1990).

With the exception of the experiments conducted by Saino & Hattori (1980), measurements of nitrogenase activity in *Trichodesmium* have been carried out preferentially in samples that selected for the colonial morphology (Dugdale et al. 1964, Mague et al. 1974, 1977, Carpenter & McCarthy 1975, Carpenter & Price 1977, McCarthy & Carpenter 1979, Carpenter et al. 1987, Scranton et al. 1987). The strong bias toward the ecological study of colonies, rather than the native population containing both colonies and free filaments (trichomes), is probably due to the unsupported hypothesis that N$_2$ fixation by *Trichodesmium* in nature is restricted to micro-aerophilic and anaerobic environments found in the center of colonies (Taylor et al. 1973, Carpenter & Price 1976, Bryceson & Fay 1981, Paerl & Bland 1982, Carpenter 1983, Paerl & Bebout 1988, Paerl et al. 1989). This bias may have resulted in an underestimation of the N$_2$ fixation because, as reported in Marumo & Asaoka (1974) and Letelier & Karl (1996), most of *Trichodesmium* biomass at Stn ALOHA and in the North Pacific subtropical gyre is in the form of free filaments. Even though Saino & Hattori (1980) reported significant N$_2$ fixation rates by naturally occurring single trichomes, the remaining evidence that single trichomes fix dinitrogen in natural environments has been only indirect (Saino & Hattori 1982, Ohki & Fujita 1988, Carpenter et al. 1990, Bergman et al. 1993). Furthermore, even if N$_2$ fixation proved to be an important source of new nitrogen for the marine environment, the availability of other nutrients such as phosphorus and iron (Doremus 1982, Fogg 1982, Martin & Fitzwater 1988) would limit the ecological benefits of an essentially unlimited nitrogen supply.

In 1992, Karl et al. proposed a phosphorus transport mechanism (P-Transport model) based on the capacity of *Trichodesmium* to control its buoyancy and migrate between light saturated-nutrient depleted (surface) waters and light limited-nutrient rich (deep) waters. In this model *Trichodesmium* accumulates energy in the form of heavy ballast polycarbohydrates under light saturated conditions near the surface and subsequently sinks into the nutricline (125 to 150 m) where these carbohydrates are consumed to provide energy, in part, for the uptake and intracellular storage of phosphorus. Subsequent migration into surface waters, as a result of the consumption of the ballast, would provide the cells with an opportunity to grow at the expense of N$_2$ and light, using these translocated P stores.

During the first few years of the Hawaii Ocean Time-series (HOT) program we collected evidence suggesting that N$_2$ fixation is an important source of new nitrogen to the biota at Stn ALOHA (22°45'N, 158°00'W; Fig. 1), accounting for approximately 50% of the exported N production (Karl et al. 1995, 1997). Because over this period of time the only identified nitrogen fixers constantly present in our samples were *Trichodesmium* spp., we decided to investigate some basic physiological aspects of natural populations of these cyanobacteria at Stn ALOHA, in an attempt to understand their role in the nitrogen and phosphorus cycles of the North Pacific subtropical gyre. In the present paper, we report results of experiments addressing the importance of aerobic nitrogen fixation by *Trichodesmium*, including the potential contribution by naturally occurring single trichomes. We further assess the rates of oxygen consumption by these cyanobacteria as a mechanism to protect nitrogenase from oxygen evolution and we compare the C:N:P elemental composition of positively and negatively buoyant colonies, as well as the rates of phosphorus uptake under dark conditions, in an attempt to understand the source of nutrients other than nitrogen supporting the *Trichodesmium* spp. populations at Stn ALOHA.

**METHODS**

**Field N$_2$ fixation assay.** Nitrogenase activity of *Trichodesmium* colonies and single filaments collected in the euphotic zone at Stn ALOHA was estimated by C$_2$H$_4$ evolution during selected HOT cruises. Colonies and trichomes were isolated and transferred into filtered seawater as described in Letelier & Karl (1996). Because of the low sensitivity of this assay, a large *Trichodesmium* biomass was required for each sample (>200 filaments per sample). Single trichomes used for these experiments were isolated from a large volume of seawater (>80 l) collected at a single predetermined...
water depth that was predicted from previous observations to have a high *Trichodesmium* filament concentration (25 m). The presence of nitrogenase activity was tested by incubating *Trichodesmium* for 2 h in on-deck incubators that simulated both *in situ* light level and ambient water temperature. *Trichodesmium* aliquots (3 ml) were enclosed in 13.5 ml serum bottles and sealed with a gas-tight stopper. This was followed by the addition of 0.5 ml H₂SO₄ (2N) to each serum bottle. Ethylene (C₂H₂) in the headspace was measured using a Varian 3300 gas chromatograph equipped with a PORAPAK T separation column and a hydrogen flame ionization detector. Nitrogenase activity (reported here as C₂H₄ evolution) was normalized to chl a (fluorometric detection of acetone extract; Chavez et al. 1995) to provide a relative index of N₂ fixation that was suitable for comparison among the individual experiments.

**Laboratory N₂ fixation assay.** In August 1990, a *Trichodesmium* culture was established and was sustained for nearly 2 yr in the Marine Microbiology Laboratory at the University of Hawaii, USA (Letelier 1994). *Trichodesmium* colonies from this culture were used to study the temporal evolution N₂ fixation under aerobic conditions. These experiments were conducted in aquil medium (Morel et al. 1979) without fixed nitrogen. Colonies were disrupted by bubbling with Ar for 30 min (Saino & Hatton 1982) followed by filtration through a 202 μm mesh to obtain a source of single trichomes. Two culture flasks were filled with the resulting *Trichodesmium*. One was kept in microaerophilic conditions by flushing with N₂ gas for 5 min before sealing and the other flask was incubated under aerobic (air) conditions.

Colonies isolated from the original culture were also incubated under micro-aerophilic and aerobic environments following the same procedure. Nitrogenase activity was measured 3 times each day (07:00–08:00, 13:30–14:30 and 19:00–20:00 h), over a 4 d period. At each sampling period, 3 ml aliquots were withdrawn from each flask and incubated for 1 h under modified natural light conditions in 13.5 ml serum bottles filled with 2.0 ml C₂H₂ (approximately 15% v/v). Transfers were made under a flow of N₂ to maintain micro-aerophilic conditions, as required. The samples were placed into a temperature-controlled (24 to 26°C), light incubator located in the roof of the Marine Science Building on the University of Hawaii campus. A combination of blue and neutral density filters in the incubators reduced the incident irradiance to 23.5% of ambient light. Termination of the incubation and measurement of C₂H₄ evolution were performed as described for the N₂ fixation field assay. The cultures were monitored periodically for the formation of new colonies in single trichome suspensions by microscopic observation.

**Oxygen evolution of colonies.** Changes in the oxygen concentration in glass iodine flasks due to the metabolic activity of *Trichodesmium* colonies were quantified by potentiometric micro Winkler titration (Carpenter 1965) during April 17 and 18, 1992. Colonies were collected from the mixed-layer at Stn ALOHA and transferred into filtered seawater before incubation. Eight replicate samples of seawater collected at each of 4 depths (5, 25, 45 and 75 m) were withdrawn into iodine flasks and capped until the addition of colonies. Of these 8 replicates, 4 randomly selected subsamples were inoculated with 5 colonies each. Immediately after the addition of colonies, the dissolved O₂ content
of 1 sample with added *Trichodesmium* and 1 control bottle (filter seawater only) were fixed by the simultaneous addition of manganese chloride and alkaline iodide. The remaining samples were incubated on-deck at simulated *in situ* irradiance and temperature. One additional set of 8 replicates with seawater collected at 5 m depth was used to estimate oxygen changes in the dark for the *Trichodesmium* treatment only.

Comparison of elemental composition between positively and negatively buoyant colonies. Rising and sinking *Trichodesmium* colonies collected at approximately 5 and 100 m depth, as described by Letelier & Karl (1996), were isolated during a 10 min separation period using 8 settling columns (SETCOL: Bienfang 1981). The segregated colonies were rinsed and transferred into 50 ml of 0.2 μm filtered seawater where they were mechanically disrupted to produce suspensions of single trichomes. From each suspension, 10 ml aliquots were filtered onto combustible 25 mm GF/F Whatman filters and stored frozen for subsequent particulate carbon (PC) and particulate nitrogen (PN) analyses. Ten ml aliquots were also filtered though combusted and acid-washed (HCl) 25 mm GF/F Whatman filters and stored frozen for subsequent particulate phosphorus (PP) analyses. PC and PN were measured using a Perkin-Elmer model 2400 CHN analyzer and PP was quantified spectrophotometrically after combustion (450 to 500°C) and acidification (0.5 N HCl at 90°C for 90 min) according to the method of Karl et al. (1996).

Dark phosphorus uptake. Phosphorus uptake by sinking *Trichodesmium* colonies incubated in filtered seawater was measured using 32P as a tracer (in the form of H232PO4; carrier free, ICN Radiochemicals). Five liters of water collected at 150 m depth at Stn ALOHA were sterile filtered through a 0.2 μm Nalgene filter and stored in acid washed polycarbonate bottles. Subsamples of the filtered water were withdrawn for soluble reactive phosphorus (SRP) and total dissolved phosphorus (TDP) analyses. SRP concentration was estimated by the molybdenum blue reaction (Murphy & Riley 1962). TDP was estimated using the same procedure after liberating the combined phosphorus by ultraviolet photolytic oxidation (Armsstrong et al. 1966). Dissolved organic phosphorus (DOP) concentration was estimated as [TDP] – [SRP] which assumes that all soluble non-reactive P is organic.

Sinking colonies, collected at approximately 5 m depth, were isolated using SETCOL columns and were rinsed before transferring them into polycarbonate incubation bottles containing filtered seawater. Three incubation bottles with *Trichodesmium* colonies and a negative control (filtered seawater without colonies) were prepared for this experiment. Each bottle was filled with 500 ml of 150 m depth filtered seawater and 45 colonies were added to each of 3 bottles. Radiotracer 32P was added to each bottle to yield a final activity of 2.8 MBq 32P l⁻¹. The initial specific activity was 15.5 MBq (μmol SRP)⁻¹ or 7.7 MBq (μmol TDP)⁻¹. All samples were kept in the dark and at simulated *in situ* temperature during the incubation.

Prior to the inoculation with 32P, 5 *Trichodesmium* colonies from each bottle were removed by Pasteur pipette to measure the initial chl a:PP ratio. These colonies were resuspended in 50 ml filtered seawater and vortexed until only single trichomes were observed in order to obtain a homogeneous suspension of filaments. Twenty-five ml of each suspension was concentrated onto a combusted acid washed 25 mm GF/F filter and rinsed twice with 10 ml filtered seawater prior to PP analyses. Chl a was extracted from the remaining 25 ml by filtering the sample through a 25 mm GF/F Whatman filter and storing the sample in 100 % acetone at −20°C. The extracted chl a concentrations were measured by fluorometry.

Samples, consisting of either 5 colonies per bottle and 5 ml of the negative control (filter seawater without *Trichodesmium*), were withdrawn approximately every 6 h during the first 24 h. The colonies were rinsed twice, resuspended in 50 ml filtered seawater and vortexed until obtaining an homogeneous filament suspension. A set of samples was collected immediately after the addition of 32P (t = 0 h) and a final set was collected after a 46 h incubation period. Twenty-five ml of suspension was used for chl a determination and the remaining 25 ml was filtered through a 25 mm GF/F Whatman filter, rinsed twice with filtered seawater and the filters stored at 4°C in polyethylene scintillation vials for determination of inorganic phosphorus uptake. Specific activity in each incubation bottle was estimated from triplicate 1 ml samples withdrawn at the beginning of the incubation. In addition, triplicate 1 ml samples were withdrawn at each experimental time-point to assess the remaining 32P in solution for mass balance determination. Radioactivity was measured as soon as we returned to our shore based laboratories by the Cerenkov technique (Kamp & Blanchard 1971) using a Packard Tri-Carb scintillation counter. Counting efficiency was 43%.

RESULTS

Nitrogen fixation

All *Trichodesmium* samples collected at Stn ALOHA had measurable nitrogenase activity (Table 1, Fig. 2). The activity appeared to display limited photoinhibi-
Table 1. Comparison of the aerobic evolution of ethylene (C$_2$H$_4$) by single trichomes, intact colonies, and disrupted colonies for a sample collected at Stn ALOHA. Values are means (±1 standard deviation) following a 1 h incubation.

<table>
<thead>
<tr>
<th>Collection depth (m)</th>
<th>Nitrogenase activity [nmol C$_2$H$_4$ (µg chl a)$^{-1}$ h$^{-1}$]</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Single trichomes Intact colonies Disrupted colonies</td>
</tr>
<tr>
<td>5</td>
<td>2.3 (0.4) 7.6 (1.1) 0.3 (0.2)</td>
</tr>
<tr>
<td>25</td>
<td>3.6 (0.9) 12.3 (1.6) 0.7 (0.3)</td>
</tr>
<tr>
<td>45</td>
<td>3.1 (0.7) 11.7 (2.6) 0.5 (0.4)</td>
</tr>
<tr>
<td>75</td>
<td>0.3 (0.3) 3.1 (1.8) 0.1 (0.2)</td>
</tr>
</tbody>
</table>

Fig. 2. (A) Depth profile of aerobic acetylene reduction (C$_2$H$_4$ evolution) by Trichodesmium colonies, disrupted colonies, and naturally occurring single trichomes collected at Stn ALOHA and incubated under simulated in situ irradiance and temperature. (B) Depth profile of aerobic acetylene reduction (C$_2$H$_4$ evolution) by colonies collected during different cruises. Error bars indicate ±1 standard deviation.

Oxygen evolution

Trichodesmium colonies display an unusually shallow compensation depth (Fig. 4A) relative to other pelagic photoautotrophs. Although net oxygen evolution reaches 0.2 mol O$_2$ (µg chl a)$^{-1}$ h$^{-1}$ at 550 µE m$^{-2}$ s$^{-1}$ (~10 m; Fig. 4B), at 25 m depth it is not significantly different from zero ($p > 0.1$). At 75 m the net oxygen evolution in the light is not statistically different from the oxygen consumption measured in samples incubated.

Table 2. Daily variations in nitrogenase activity measured as nmol C$_2$H$_4$ (µg chl a)$^{-1}$ h$^{-1}$ in Trichodesmium cultures incubated under aerobic and micro-aerophilic conditions. Values are means ±1 standard deviation) following a 1 h incubation.

<table>
<thead>
<tr>
<th>Date-time (mo/d/yr-h)</th>
<th>Colonies Aerobic Micro-aerophilic</th>
<th>Filaments Aerobic Micro-aerophilic</th>
</tr>
</thead>
<tbody>
<tr>
<td>11/12/90-07:30</td>
<td>3.3 (2.1) 4.3 (2.1)</td>
<td>0.9 (1.1) 2.0 (0.4)</td>
</tr>
<tr>
<td>11/12/90-14:00</td>
<td>26.9 (1.3) 33.2 (1.4)</td>
<td>1.6 (1.7) 26.6 (1.2)</td>
</tr>
<tr>
<td>11/12/90-19:30</td>
<td>4.9 (1.3) 9.8 (1.5)</td>
<td>1.2 (0.8) 7.4 (1.9)</td>
</tr>
<tr>
<td>11/13/90-07:30</td>
<td>10.2 (1.1) 15.6 (1.0)</td>
<td>1.1 (0.8) 9.7 (0.7)</td>
</tr>
<tr>
<td>11/13/90-14:00</td>
<td>34.5 (1.0) 42.8 (0.8)</td>
<td>8.5 (0.5) 32.6 (1.6)</td>
</tr>
<tr>
<td>11/13/90-19:30</td>
<td>9.4 (1.6) 11.6 (1.1)</td>
<td>2.7 (0.6) 6.0 (0.9)</td>
</tr>
<tr>
<td>11/14/90-07:30</td>
<td>13.1 (3.7) 17.6 (0.7)</td>
<td>4.1 (3.6) 11.4 (0.8)</td>
</tr>
<tr>
<td>11/14/90-14:00</td>
<td>42.5 (1.8) 50.9 (2.7)</td>
<td>15.4 (0.7) 35.7 (0.7)</td>
</tr>
<tr>
<td>11/14/90-19:30</td>
<td>8.6 (0.5) 13.4 (2.4)</td>
<td>3.0 (0.4) 9.6 (0.5)</td>
</tr>
<tr>
<td>11/15/90-07:30</td>
<td>11.7 (1.2) 14.7 (0.7)</td>
<td>4.3 (1.3) 9.3 (0.4)</td>
</tr>
<tr>
<td>11/15/90-15:00</td>
<td>45.5 (1.4) 47.3 (1.2)</td>
<td>16.3 (1.3) 33.7 (1.4)</td>
</tr>
</tbody>
</table>
Fig. 3. (A) Temporal changes in acetylene reduction (C$_2$H$_4$ evolution) measured in disrupted and intact *Trichodesmium* colonies grown in culture (solid symbols = micro-aerobic; open symbols = aerobic; squares = disrupted colonies; circles = intact colonies). (B) Temporal aerobic trend of acetylene reduction (C$_2$H$_4$ evolution) by disrupted *Trichodesmium* colonies under aerobic conditions expressed as the fraction of acetylene reduction measured in intact colonies. Error bars indicate ±1 standard deviation.

Fig. 4. Net oxygen changes during shipboard incubation of *Trichodesmium* colonies collected on 2 consecutive days at Stn ALOHA. (A) Depth profile. (B) Oxygen plotted against irradiance. Error bars indicate ±1 standard deviation.

in the dark ($p > 0.1$), suggesting that photosynthetic carbon assimilation at this depth is negligible. Over the light range of our incubations net oxygen evolution may be described by a geometric mean (model II) linear function of the photon flux $y = -0.18 + 6.7 	imes 10^{-4} x$, where $y =$ moles O$_2$ (g chl a)$^{-1}$ h$^{-1}$ and $x =$ μE m$^{-2}$ s$^{-1}$. $R^2 = 0.873, n = 29$.

**Elemental composition**

The C:N:P elemental ratios obtained from positively and negatively buoyant colonies collected at different depth strata suggest only subtle changes in the elemental composition of *Trichodesmium* (Table 3). Sinking colonies have, on average, a PC:PN ratio higher than the Redfield stoichiometry (Redfield et al. 1963), while rising colonies are, on average, below this ratio. The lowest average concentration of PC per unit PN is found in rising colonies collected at 100 m depth, suggesting either a net consumption of cellular carbon per unit nitrogen or active assimilation of nitrogen relative to carbon at depth, or both.

The PP content of *Trichodesmium* colonies is consistently lower, with respect to PC and PN, than Redfield stoichiometry of 106C:16N:1P (Table 3). The lowest mean PN:PP measured (34.2 ± 2.8) was found in rising colonies collected at 100 m depth. This value is 2.2 times greater than the Redfield ratio of 16:1 and about 30% lower than that measured for colonies sampled at 5 m depth.
Table 3. Comparison of elemental ratios (particulate carbon:particulate nitrogen, PC:PN, and particulate nitrogen:particulate phosphorus, PN:PP) in sinking and rising colonies collected at surface (5 m) and depth (100 m) at Stn ALOHA. Values are means (±1 standard deviation)

<table>
<thead>
<tr>
<th>Buoyancy</th>
<th>PC:PN 5 m</th>
<th>PC:PN 100 m</th>
<th>PN:PP 5 m</th>
<th>PN:PP 100 m</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sinking</td>
<td>7.32 (0.36)</td>
<td>6.61 (0.27)</td>
<td>42.76 (1.71)</td>
<td>44.80 (1.12)</td>
</tr>
<tr>
<td>Rising</td>
<td>6.32 (0.32)</td>
<td>5.86 (0.34)</td>
<td>43.75 (1.64)</td>
<td>34.19 (2.83)</td>
</tr>
</tbody>
</table>

**Dark phosphorus uptake**

Sinking colonies collected near the sea surface display active uptake of inorganic phosphorus in the dark during the first 12 to 24 h of incubation (Fig. 5). Based on the SRP concentration in the incubation bottles at the beginning of the incubation period (0.18 μM) and the phosphorus specific activity [15.5 MBq (μmol SRP)-1] it is possible to calculate the absolute inorganic phosphorus uptake by *Trichodesmium* in this experiment. During the first 12 h, the uptake rate per unit chl a appears to be constant [approximately 6 × 10^-2 ng P (ng chl a)^-1 h^-1]. The largest accumulation of 32P_2O_5^- [1.29 ng P (ng chl a)^-1] was measured in the 24 h time point samples. Nevertheless, it should be noted that values measured at 13, 19, and 48 h after the beginning of the incubation are not statistically different (p < 0.05).

The concentration of chl a in colonies does not vary significantly over time (Fig. 5A), suggesting that dark photoadaptation is negligible in colonies during the incubation. Using the PP:chl a ratio (2.2:1 w:w) measured at the beginning of this experiment it is possible to normalize phosphorus uptake to initial phosphorus content. Based on this calculation, we find that sinking *Trichodesmium* colonies were able to assimilate the equivalent of 35 to 57% of their phosphorus content from inorganic phosphorus over the first 12 h of dark incubation during HOT-49 (September 1993; Fig. 5).

**DISCUSSION**

In 1980 and 1982, Saino & Hattori presented strong evidence that *Trichodesmium* is able to fix nitrogen under aerobic conditions. However, until recently nitrogenase activity was still thought to be restricted to low oxygen environments within *Trichodesmium* colonies (Paerl et al. 1989). A re-evaluation of *Trichodesmium* nitrogenase activity under aerobic conditions has been conducted recently (Capone et al. 1990, Carpenter et al. 1990, Bergman et al. 1993, Prufert-Bebout et al. 1993). As part of this reassessment, Carpenter et al. (1990) found an even distribution of photosystems I and II in the colonial trichomes. This observation indicates that nitrogenase activity is not protected from photosynthetic O_2 evolution as a result of the absence of photosystems II in the central region of the colony, as initially proposed by Carpenter & Price (1976). Furthermore, nitrogenase activity in *Trichodesmium* appears to be strongly dependent on light (Fig. 3A; Saino & Hattori 1978, Ohki & Fujita 1988), suggesting that there is no temporal separation between photosynthesis and nitrogen fixation.

Field and laboratory data obtained during our study at Stn ALOHA support the results of Saino & Hattori (1980) and reinforce the notion that, under ambient oxygen levels, N_2 fixation is reduced but not completely suppressed. These data also suggest that *Tri-
*Trichodesmium* possesses an intracellular oxygen scavenging mechanism that is capable of protecting nitrogenase activity from O$_2$ inactivation. When single trichomes obtained from the disruption of colonies grown in culture without combined nitrogen were incubated under aerobic conditions, there was a lag period of approximately 40 h before nitrogenase activity was significantly different than seawater controls (Fig. 3A). This lag period could not be attributed to the handling of the samples during the disruption process because single trichomes obtained in the same manner but incubated under micro-aerophilic conditions displayed nitrogenase activity close to the activity measured in colonies. Also, field sample results indicate that naturally occurring free trichomes incubated under aerobic conditions have higher nitrogenase activity than filaments obtained from colonies by mechanical disruption (Fig. 2A). Furthermore, *Trichodesmium* colonies at Stn ALOHA display high consumption of O$_2$ under light conditions (Fig. 4B). This phenomenon has also been observed by Kana (1992) who attributed the high O$_2$ consumption to the Mehler reaction (production of O$_2^-$ by photosystem I followed by the conversion of O$_2^-$ into H$_2$O$_2$ by a superoxide dismutase) and to a high basal respiration rate.

Cunningham & Capone (1992) have reported the presence of a Fe superoxide dismutase in *Trichodesmium*, and Bergman et al. (1993) observed a strong correlation between the expression of nitrogenase and an aa$_3$-type cytochrome oxidase in *Trichodesmium thiebautii* cells. The lag period measured for N$_2$ fixation after the disruption of colonies grown in culture suggests that these biochemical processes are not always present at a level sufficient to protect nitrogenase activity from oxygen evolution in our *Trichodesmium* cultures. However, the existence of this lag period is absent in the Saino & Hattori (1982) report, as well as in the results of field experiments with disrupted colonies developed during this study. Colonies collected from natural populations that had been disrupted and incubated under aerobic conditions display highly reduced but still detectable nitrogenase activity (Fig. 2; Saino & Hattori 1982).

The light intensity used for the laboratory culture experiments (15 to 30 $\mu$E m$^{-2}$ s$^{-1}$) may help explain this discrepancy. Under low irradiance, respiration rates must decrease in order to sustain net growth rate. Because N$_2$ fixation has a higher demand for adenosine triphosphate (ATP) relative to reduced nicotinamide adenine dinucleotide phosphate (NADPH), the Mehler reaction may play an important role, not only by removing oxygen, but also by controlling the supply of ATP for nitrogenase activity (Kana 1992). Hence, when grown under dim light, it may be to the advantage of *Trichodesmium* to couple the removal of oxygen with light processes (Mehler activity) rather than to maintain an elevated respiration rate.

A sudden increase of oxygen and light upon the disruption of colonies may also change the balance of enzymatic processes controlling the intracellular oxygen. Although this is also true for colonies collected from natural environments, the average radiant flux penetrating down to 45 m during a summer day at Stn ALOHA is approximately 150 $\mu$E m$^{-2}$ s$^{-1}$, 5 to 10 times the irradiance of cultures studies (at 5 m depth the light is 50 times higher). Hence, it is possible that colonies collected at Stn ALOHA, having a higher respiratory oxygen uptake relative to colonies grown in culture, are able to control more effectively the increase of intracellular oxygen tension. Furthermore, *Trichodesmium* grown under dim light would not be expected to accumulate polysaccharides to the extent that they do when incubated under high light (Li et al. 1980). This reserve of reduced carbon may be important for respiration and the removal of oxygen.

It is interesting to note that the short term (1 to 2 h) nitrogenase activity measured at Stn ALOHA appears to be partially inhibited when samples are incubated at 5 m depth light irradiance (Fig. 2). This observation, combined with results from the net oxygen evolution by *Trichodesmium* colonies (Fig. 4), suggests that the optimum light depth for this cyanobacterium to grow and fix N$_2$ is probably close to 25 m depth. This is also the depth at which a *Trichodesmium* biomass maximum is measured under stratified water column conditions (mixed layer <25 m) and non-bloom conditions at Stn ALOHA (Letelier & Karl 1996).

Although sinking and rising colonies isolated from different depth strata at Stn ALOHA display only minor changes in their bulk C:N:P composition (most of them are not statistically different at $\alpha = 0.05$), the changes follow a consistent pattern (Table 3) and support the phosphorus-transport model postulated by Karl et al. (1992; see also Kromkamp & Mur 1984, Fogg 1987). Throughout the water column, *Trichodesmium* has elevated C:P and N:P ratios, relative to the classic Redfield ratio, and this may be a specific cellular adaptation for growth in low phosphorus environments. This P-sparing effect has been reported previously, but the biochemical basis of the effect is not known.

Sinking colonies are enriched in carbon and depleted in phosphorus relative to nitrogen when compared to rising colonies. The increase in carbon may be explained by a preferential carbon assimilation into polysaccharides by *Trichodesmium* cells (Roman et al. 1994) as observed by Li et al. (1980) in samples collected at shallow depths and incubated under high irradiance (100 to 1500 $\mu$E m$^{-2}$ s$^{-1}$). This relative increase in carbon is also in agreement with the hypothesis that *Trichodesmium* cell density is regulated by
the synthesis and degradation of polycarbohydrate reserves (Villareal & Carpenter 1990). Nevertheless, we should be careful with the interpretation of cellular C:N ratios, and how it relates to buoyancy. During an August 1989 *Trichodesmium* bloom at Stn ALOHA the C:N ratio of floating colonies was 7.1:1 (mol:mol). A similar ratio (7:1) was measured by Lewis et al. (1988) during a summer bloom in the North Atlantic. The mean ratio of sinking colonies collected from 5 m depth at Stn ALOHA under non-bloom conditions is only 6.9:1 (±0.3:1).

The exact reason why cyanobacteria lose the capacity to control the cellular density under bloom conditions is not known. In freshwater systems the collapse of gas vacuoles due to a decrease in the availability of inorganic carbon has been shown to be an important factor (Klemer et al. 1982). This is probably not the case for *Trichodesmium* because of the high critical pressure for the collapse of its gas vesicules (Walsby 1978) and the high, non-limiting, concentration of dissolved inorganic carbon in seawater.

An alternative hypothesis states that cell metabolism and the physiological state of an organism is regulated by the source and availability of nutrients (Konopka 1984). Under non-bloom conditions *Trichodesmium* growth in shallow waters is probably limited by the availability of elements other than carbon, hydrogen, oxygen and nitrogen. It is also possible that limitation by phosphorus (or other limiting nutrient) increases the allocation of carbon assimilated into high density carboxysomes. *Trichodesmium* colonies appear to have high degradation rates during bloom events, increasing the availability of nutrients where large ocean surface accumulations are found (Devassy et al. 1978, Fogg et al. 1987). If the growth rate of *Trichodesmium* is balanced by the degradation rate, nutrient limitation will not take place and probably less carbon assimilation will go into carbohydrate ballast. However, inorganic carbon uptake will have to remain large relative to nitrogen uptake in order to produce C:N ratios equal to 7:1 or greater. Under this hypothesized growth condition, cells close to the surface will not sink (they will only grow, degrade or be grazed) and the net increase in *Trichodesmium* biomass will be mainly the result of colonies growing and transporting nutrients to the surface by active vertical migration.

The evidence for an active uptake of inorganic phosphorus by *Trichodesmium* in the dark is supported by the variation of N:P between sinking and rising colonies collected at depth (Table 3), as well as by the results from the dark incubations of sinking colonies in nutrient-rich waters (Fig. 5). If we assume no absolute changes in the *Trichodesmium* nitrogen content between sinking and rising colonies, then the decrease in the N:P ratio from 45:1 to 34:1 reflects an increase in cellular phosphorus content of 32%. This result is similar to the increase calculated from the inorganic phosphorus uptake during dark incubations. Nevertheless, we should be aware that the dark incubation results may be biased for several reasons. Sinking colonies collected close to the surface at mid morning were suddenly transferred into a medium rich in HPO₄²⁻ and kept in the dark for 48 h. This procedure may have produced an overestimation of the *in situ* phosphorus uptake because, in nature, colonies must expend cellular energy to migrate down to the nutricline. Furthermore, as soon as the cells become positively buoyant colonies will start migrating away from the nutricline.

During our experiment approximately 15% of the colonies were observed to be positively buoyant after a 6 h incubation and 100% after 18 h. Because phosphorus uptake is an active process (it requires energy) a potentially high cellular energy reserve at the beginning of the experiment (colonies did not respire carbohydrate reserves during their migration toward the nutricline) could produce an overestimation of the natural phosphorus uptake. This effect is probably further aggravated by maintaining the colonies in an artificially high inorganic phosphorus environment once buoyancy has changed. Nevertheless, the results of the *³²P* uptake experiments, although not conclusive with respect to the maximum absolute uptake of inorganic phosphorus by *Trichodesmium*, suggest that active uptake only occurs during the first 12 to 24 h. Because most of the colonies were still negative or neutrally buoyant after 6 h incubation, we believe that the artificial confinement of *Trichodesmium* does not contribute significantly to an overestimation of the absolute uptake of inorganic phosphorus when 12 h incubation results are used.

On the other hand our experiment may have underestimated the total assimilation of phosphorus because *Trichodesmium* displays high alkaline phosphatase activity (Yentsch & Yentsch 1972, McCarthy & Carpenter 1979, Elardo et al. 1994). On the date of the *³²P* uptake experiment the concentration of dissolved organic phosphorus (DOP) in the first 100 m depth at Stn ALOHA was consistently above 150 nM, a value that was 5 to 6 times higher than the concentration of inorganic phosphorus measured in the same depth range (Fig. 6; also see Karl & Tien 1997). If we assume that *Trichodesmium* uses both pools at the same rate then the amount of phosphorus assimilated doubles. Our experiment did not discriminate between SRP and DOP uptake and, because DOP may be an equally important source of phosphorus, the total uptake measured by *³²P* incorporation using SRP concentrations to calculate *P*-specific activity gives only a conservative estimate of the total phosphorus uptake. Nevertheless, if the *Trichodesmium* P-transport model
represents a source of phosphorus and other nutrients to the upper euphotic zone. *Trichodesmium* vertical migrations must be regarded as a sink of energy and a source of inorganic nutrients for a stratified euphotic zone as suggested by our results. Uptake of DOP, even if coupled with vertical migrations, represents a recycling of organic matter rather than the input of new phosphorus into the upper euphotic zone.

In conclusion, although the formation of colonies in *Trichodesmium* has been seen mainly as an ecological adaptation for the protection of nitrogenase activity, our results support the notion that this morphological adaptation only enhances N$_2$ fixation. If we consider that the velocity of vertical migration of a particle is proportional, not only to the difference of density with its surrounding medium, but to the square of its radius (i.e. Stoke's law; McCave 1984, Walsby 1992), the formation of colonies may be interpreted as a morphological adaptation to increase the frequency of vertical displacements of *Trichodesmium* in the water column. This suggestion is supported by the active uptake of inorganic phosphorus in the dark by sinking colonies and the accumulation of carbon under high light intensities. But, if *Trichodesmium* obtains its phosphorus for growth by active migration, then it is difficult to explain the high abundance of single trichomes in the water column of Stn ALOHA. One possible explanation comes from the observation by Ohki & Fujita (1982) that, under certain conditions, single trichomes aggregate to form colonies. However, these authors observed de novo aggregations only when the density of trichomes was 2 orders of magnitude higher than the average density observed at Stn ALOHA. Furthermore, because DOP may also be considered a potential source of phosphorus for the single trichomes, the colonial morphology and the P-transport model may not represent a real advantage in terms of phosphorus uptake in the North Pacific subtropical gyre.

Finally, the carbohydrate ballast hypothesis, when considered in conjunction with the high critical tugor pressure of vacuoles observed in *Trichodesmium* spp. (Walsby 1992), suggests that intact filaments of these cyanobacteria are not significant contributors to the sinking flux of organic matter from the euphotic zone. If sinking *Trichodesmium* cells, unable to collapse their vacuoles, use carbohydrate ballast as an energy source, they will become positively buoyant under limiting conditions. Under this scenario, the only possibility for these cells to be exported permanently below the euphotic zone is by association with dense particles such as marine snow or discarded appendicularian houses (Alldredge 1976). Hence, we speculate that most elements incorporated into *Trichodesmium* biomass in the euphotic zone need to be recycled within the euphotic zone before being exported.

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