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Population dynamics and environmental conditions affecting *Trichodesmium* spp.
(filamentous cyanobacteria) blooms in the south-west lagoon of New Caledonia

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1 **Abstract.** The present description of *Trichodesmium* spp. population dynamics and associated
2 environmental variables is the first one using a very short sampling interval (2-3 days). Such a
3 strategy allows a fine description of changes involving the population density and characteristics. It
4 also lends itself to interpreting those changes from past environmental conditions. During the two
5 surveys, 2 and 5 month-long, *T. erythraeum* dominated the *Trichodesmium* population, with
6 temporary occurrence of *T. thiebautii* and five blooms could be described. These events appeared at
7 temperatures $> 24^{\circ}\text{C}$ and followed, in all cases, nitrate, soluble reactive phosphorus and chlorophyll
8 *a* enrichments, with a 3-7 day time lag. Low wind speed ($< 4\text{m s}^{-1}$) was not a prerequisite for
9 *Trichodesmium* bloom developments as long as temperatures exceeded 26°C . As abundance
10 increased during the bloom, so did the number of filaments (trichomes) in colonies and their
11 buoyancy, leading to a clear positive biomass gradient from the bottom to the surface. A simple
12 model, using variable growth rates, showed trichome ascent would be responsible for 87-99% of
13 concentrations at 0.5m, with *Trichodesmium* net growth rates ranging from 0.11 to 0.38 d^{-1} . Finally,
14 rapid trichome density declines could be ascribed to nutrient depletion and massive surface death
15 following ascent.

16

17

18 **1. Introduction**

19

20 High surface concentrations of the chain-forming diazotrophic cyanobacteria *Trichodesmium* spp.
21 have drawn attention, not only for their spectacular characteristics, but also for their potential
22 toxicity (Landsberg, 2002,) and contribution to the carbon and nitrogen cycles in oligotrophic
23 regions via diazotrophy, i.e. di-nitrogen (N_2) fixation (Karl et al., 2002). Such natural phenomena,
24 called "blooms" in most references, have often been observed in the south-west Pacific, both in the
25 open ocean (Dupouy et al., 2000) and coastal areas (Jones et al., 1982), as well as in other parts of
26 the tropical belt (Capone et al., 1997). While many causative mechanisms for bloom development

27 have been proposed in the past (Sellner, 1997), we still lack reliable descriptions of past
28 environmental conditions that triggered their increase. The word "bloom" itself remains imprecise
29 and is generally understood as corresponding to rapid and intense growth, as it does for other
30 phytoplankters, although it is now admitted that aggregation may be responsible for many observed
31 surface *Trichodesmium* concentrations. One of the reasons for such uncertainties is that many reports
32 are based on instant observations, no account being provided on past characteristics, biomass and
33 vertical distribution of the *Trichodesmium* population, as stressed by Carpenter et al. (2004).
34 Therefore, the understanding of bloom dynamics and causes would benefit from high frequency
35 observations of both cyanobacteria and environmental variables over the water column.

36 Such a strategy, involving very frequent sampling on a long period of time can virtually be made
37 in coastal areas, only. Present knowledge, however, rely on rather low frequency observations with
38 sampling being made every 7 – 15 days (Lugomela et al., 2002), 1 month (Post et al., 2002) or 10 –
39 15 days (Muslim and Jones, 2003). While correlations between simultaneous data on *Trichodesmium*
40 densities and environmental variables could be evidenced in these studies (Post et al., 2002; Muslim
41 and Jones, 2003), effect of potential bloom triggers and environmental conditions that are required
42 for bloom developments were not clearly proved because no information was collected in the few
43 days preceding the start of the blooming event. For the same reason, causes of high surface
44 aggregations remained hypothetical for lack of high temporal resolution surveys of *Trichodesmium*
45 density along the water column, so that discriminating between effects of growth, vertical migrations
46 and physical aggregation was impossible. Such considerations led to organizing “high” frequency
47 sampling at the entrance of the Bay of Ste Marie (Noumea, New Caledonia) with two surveys made
48 during the austral spring and summer, as both seasons are reported to be favourable for bloom
49 development. Surveys lasted for 2.5 (Survey I: 9 October - 19 December 2003) and 5 months
50 (Survey II: 15 November 2004 - 12 April 2005) with field measurements made at the same station,
51 three times per week.

52 Here, we describe the *Trichodesmium* population, the changes in filament (trichome) density,
53 degree of aggregation in colonies and vertical distribution during bloom development. From such a
54 description and observed biovolume increase rates, we will show that high surface concentrations are
55 due to growth and trichome ascent. Further, potential bloom triggers and favourable environmental
56 conditions, will be identified from monitored environmental variables that prevailed between 2 and 7
57 days before. These variables were selected, taking into account results of the Muslim and Jones
58 (2003) study in a similar shallow coastal site of the Great Barrier Reef (GBR, Australia):
59 meteorology (wind strength and rainfall), hydrology (temperature and salinity), dissolved inorganic
60 nutrients, turbidity, a proxy of suspended sediments, and chlorophyll, in addition to phytoplankton
61 and zooplankton composition. Finally, one of the usual criticism of conclusions based on a solitary
62 station deals with the effect of spatial heterogeneity on time-series data. This point will be
63 considered taking into account observations made simultaneously at the entrance and 2.0 km off the
64 bay.

65

66 **2. Materials and methods**

67

68 *2.1. Sampling sites and general conditions (Fig. 1)*

69 The study site, Sta. SM (22°18.86' S - 166°27.95' E, Fig. 1), was visited every Monday,
70 Wednesday and Friday, between 07:00 and 09:00. While the station is away from inhabited areas
71 and from urban inputs, it is protected from the prevailing SE trade winds and swell. A second
72 station, Sta. O (22°19.80' S – 166°28.48' E) was visited in 2004-2005, weather permitting for
73 comparison. Water column depth is 12 m and 28 m for Stas SM and O, respectively.

74

75 *2.2. Meteorology, hydrology, and water sampling*

76 Meteorological data were recorded at Faubourg Blanchot Meteo-France Station, 4.2 km away
77 from the entrance of the Bay of Ste Marie.

78 CTD profiles (1 m to bottom) were obtained with a SeaBird[®] SBE 19 probe, fitted with pressure,
79 temperature, salinity, turbidity, fluorescence and Photosynthetic Available Radiation (PAR) sensors.

80 Water samples were collected with 5 L-Niskin bottles at 0.5, 4, 7, and 10 m in 2003 and 0.5, 3, 6,
81 and 10 m in 2004-2005. Niskin bottles were siphoned in the laboratory on land for nutrient and
82 pigment analyses and *Trichodesmium* abundance, within one hour of sampling.

83

84 2.3. *Nutrients and pigments*

85 Nutrients were analyzed on HgCl₂-preserved samples from 0.5 and 6-7 m. Nitrate (NO₃)
86 concentrations were determined by colorimetry using a Technicon[®] auto-analyzer and standard
87 techniques (Strickland and Parsons, 1972), except for NO₃ < 0.1 μM, when the “high sensitivity”
88 procedure of Raimbault et al. (1990) was used. Soluble reactive phosphorus (SRP) concentrations
89 were measured with a Cecil[®] CE 1011 (10cm length-cell) spectrophotometer, using the molybdenum
90 blue reaction (Murphy and Riley, 1962).

91 Chlorophyll *a* (Chl *a*) was measured on the < 20 and > 20 μm size fractions at all sampled depths,
92 except for the > 20 μm fraction in 2004-2005, when analyses were only made at 3 and 10 m. For the
93 < 20 μm Chl *a*, 200ml were prefiltered through a 20 μm nylon screen and collected on Whatman[®]
94 GF/F filters. For the > 20 μm fraction, the whole Niskin bottle content was filtered through a 20 μm
95 screen and gently rinsed off with filtered seawater onto GF/F filters. All filters were stored in liquid
96 nitrogen until analysis within less than 2 weeks. Chl *a* was extracted in methanol and measured by
97 fluorometry as described by Le Bouteiller et al. (1992).

98

99 2.4. *Trichodesmium* abundance and biometry

100 *Trichodesmium* were collected on 20 μm nylon screen from the total content of Niskin bottles
101 (mean volume: 5.74 L) and transferred to a “sedimentation cell” with 5% formaldehyde. When
102 trichomes did not sink, a few drops of pure acetic acid were added to the cell, in order to lower
103 solution p_H and break gas vesicles (Carpenter and Carmichael, 1995). Microscopic examinations and

104 enumeration were made 24 h later using a Leitz[®] Fluovert inverted microscope. Enumeration
105 considered free trichomes and fusiform (tuft) and spherical (puff) aggregates, usually called
106 “colonies” in the literature. The number of trichomes per colony was recorded and the volume of
107 each trichome was calculated from its measured length and diameter, assuming a cylindrical shape
108 for the organism. In this paper, *Trichodesmium* abundance has been expressed in terms of total
109 number of trichomes (i.e. free and in colonies) per litre (trich. L⁻¹), or as trichome biovolume per
110 litre (µm³ L⁻¹). Integrated values have been computed by the trapezoidal method. While trichome
111 density and biovolume are closely related (Fig. 2), due to the fact that biovolume data includes
112 trichome numbers, biovolume is often preferred as it gives a better estimate of the biomass.

113 Two species of *Trichodesmium* were identified according to Carpenter and Carmichael (1995)
114 and Janson et al. (1995). Individual cells of *T. thiebautii* Gomont are more long than wide and the
115 trichome has a convex apex whereas *T. erythraeum* Ehrenberg cells are more wide than long and the
116 trichomes have a flat apex. *T. erythraeum* colonies are usually darker than *T. thiebautii* colonies
117 (Carpenter et al., 1993). Such morphological differences between the two species have also been
118 confirmed by genetic finger prints of *Trichodesmium* spp. sampled in different areas of New
119 Caledonia, considering the *hetR* and *rrs* encoding for 16S rDNA genes, the internal transcribed
120 spacer (ITS) of the 16S-23S rDNA region and a highly iterated palindrome, HIP1 (Trottet, 2003).

121

122 2.5. *Trichodesmium* C, N and chlorophyll a contents

123 Plankton nets (35-µm mesh), towed at 1 m below the surface, were the source of 18-75 colonies
124 hand-picked, using a plastic inoculation loop, and transferred onto 25 mm Whatman GF/F filters for
125 Chl *a* measurements, or on pre-combusted filters for carbon (C) and nitrogen (N) analyses.

126 Filters for Chl *a* measurements were stored in liquid nitrogen and analyzed as described above.

127 For C, N determinations, filters were dried at 60°C and kept in a dessicator until analysis with a

128 Integra-CN PDZ EUROPA mass spectrometer. Single colonies were also selected for carbon

129 assimilation and nitrogen fixation experiments using ^{13}C and ^{15}N techniques (data not shown here,
130 but reported in Le Borgne et al., 2006).

131 Carbon, N and Chl *a* contents are expressed per colony (col^{-1}) and per trichome (trich.^{-1}). Since
132 analyses were made on colonies originating from plankton nets, the mean number of trichomes per
133 colony was obtained from microscopic observations of aliquots originating from the same catches
134 and processed as described before.

135

136 2.6. *Microphytoplankton (>20 μm) and Macrosetella abundance*

137 Counts of the major phytoplankton taxa were conducted on samples from 2004-2005 survey.
138 Selected taxa were diatoms, dinoflagellates and *Trichodesmium* spp. The two specific
139 *Trichodesmium* spp. grazers, copepods *Macro-* and *Microsetella* spp. (Roman, 1978; O'Neil and
140 Roman, 1994) were all enumerated on all samples. They were grouped into the "*Macrosetella*"
141 taxon, since *Macrosetella gracilis* is the most abundant species in both our samples and lagoons of
142 the west coast of New Caledonia (Binet, 1984).

143

144 2.7. *Trichodesmium biovolume net increase and carbon-specific growth rates*

145 Using observed *Trichodesmium* biovolume increases, ΔV_i , exponential net increase rates, k_i (in
146 d^{-1}), can be calculated at depth, z_i :

147

$$148 \quad \Delta V_i = V_{f,i} - V_{o,i} \text{ with } V_{f,i} = V_{o,i} e^{k_i t} \Rightarrow k_i = \ln(V_{f,i}/V_{o,i})/(t_f - t_o) \quad (1)$$

149

150 where $V_{f,i}$ is the maximum biovolume of a peak period at time t_f and $V_{o,i}$, at the initial time t_o .

151 Biovolume-based doubling time, $d'_{A,i}$ (in d), is equal to:

152

$$153 \quad d'_{A,i} = \ln(2)/k_i = \ln(2) (t_f - t_o)/(\ln(V_{f,i}/V_{o,i})) \quad (2)$$

154

155 The carbon-specific growth rate for 0.5m depth, $g_{C,0.5}$, was obtained from ^{13}C primary
 156 productivity and elemental composition measurements, made on colonies in 2004-2005 (data
 157 presented in Le Borgne et al., 2006). Calculations of $g_{C,0.5}$ used the following equation:

158

$$159 \quad g_{C,0.5} = \ln((C_0 + \rho_C)/C_0) \quad (3)$$

160

161 where C_0 is the initial carbon concentration, ρ_C is the daily rate of carbon fixation, and time is
 162 understood as one day.

163

164 2.8. Data processing

165 Non-parametric and Gaussian statistics were used, following Snedecor and Cochran (1967).
 166 Bravais-Pearson and Spearman's rank correlation coefficients are r and r_s , respectively, with level of
 167 significance shown by * and ** for $p < 0.05$ and $p < 0.01$ significance, respectively; n is the number of
 168 data per series, s , the standard deviation and p , the probability of rejection in the Wilcoxon and sign
 169 tests.

170

171 3. Results

172

173 3.1. *Trichodesmium* population dynamics: abundance, vertical distribution and net increase rates

174 *Trichodesmium* spp. were always present, at least at one of the sampled depths and made almost
 175 the totality of the pelagic filamentous cyanobacteria, *Katagnymene* spp. being quasi-absent. *T.*
 176 *erythraeum* was the dominant species in 2003 (99.0% of the 5,513 counts) and 2004-2005 (83% of
 177 the 10,650 counts) with *T. thiebautii* making the remainder.

178 Amidst a background of low trichome concentration, several periods of abrupt increase and
 179 decline were observed during the two surveys (Fig. 3 and 4). When the abundance exceeded 5
 180 trich.L^{-1} for at least two consecutive sampling times (i.e. 2-3 days), trichomes were considered as

181 thriving. Given the sharp variations, the term “bloom” will be used hereafter even though it is not
182 quite appropriate, in the sense that the *Trichodesmium* never dominated the microphytoplankton,
183 contributing a maximum of 27.7% of microphytoplankton counts.

184 Five blooms were observed with various features summarized in Table 1. Each bloom was
185 characterized by one peak abundance, except in 2003, when the 3 consecutive maxima were
186 interpreted as belonging to the same bloom (see Discussion). Peak amplitudes were variable from
187 one event to another and that of February 2005, although low, was considered as a bloom because
188 trichome densities were above 5 L^{-1} for 3 weeks. *T. erythraeum* was responsible for most of the
189 blooms, except in December 2004 and February 2005 when contribution of *T. thiebautii* was
190 significant. A striking point was the vertical distribution during the blooms, with a clear positive
191 gradient of trichome densities from deep to shallow depths (Fig. 3 and 4).

192 During the peak periods, the exponential net increase rate, k_i , (equation 1), ranged from 0.19 to
193 1.51 d^{-1} at 6-7 and 10m and from 0.52 to 3.29 d^{-1} at 3-4 and 0.5m (Table 2). These k_i values were
194 higher than carbon-specific growth rates, $g_{C,0.5}$, calculated from ^{13}C uptake rates at 0.5m: 0.12-0.17
195 d^{-1} (Le Borgne et al., 2006). Exponential net increase rate calculations from biovolume variations
196 were not possible outside the peak periods, while carbon-specific growth rates, $g_{C,0.5}$, provided by
197 the ^{13}C uptake method were low, 0.022 d^{-1} on average (range: $0.017 - 0.050 \text{ d}^{-1}$).

198 Finally, density variations were synchronized and of the same magnitude at stations SM and O
199 (Fig. 4) except from February 2005 on, when higher *Trichodesmium* densities were often found at
200 the latter site. Growth rates, estimated from the ^{13}C experiments at both stations were similar (Le
201 Borgne et al., 2006).

202

203 3.2. *Trichodesmium* population features: trichome size, degree of aggregation and chemical 204 constituents

205 For both surveys, biovolume per trichome displayed quite a large range (Table 3), their length
206 variability was greater than the width one and *T. thiebautii* trichomes were significantly longer than

207 those of *T. erythraeum* ($p < 0.001$). There was no significant size difference between trichomes from
208 0.5 and 6 m, as well as from Sta. SM and O (not shown) and between 2003 and 2004-2005 for both
209 species.

210 During the two surveys, *Trichodesmium* population consisted mostly of free trichomes (65 ± 22
211 and 63 ± 15 % in 2003 and 2004-2005, respectively) and colonies in tuft morphology prevailed
212 (98% of the total number of colonies). Tufts were typically small, made of 2 to 34 trich. col^{-1} while
213 puffs of *T. thiebautii* were made up of 7 to 123 trich. col^{-1} . As the number of free trichomes
214 increased during the peak periods, so did the number of trichomes in colonies, leading to a
215 significant correlation between the two forms ($p < 0.01$; Fig. 5). Regression curves between densities
216 of trichomes in colonies and free trichomes were calculated for each of the two periods, using a
217 semi-log scale in order to take the anormal distribution of the free trichomes into account (Fig. 5).

218 *Trichodesmium* carbon, nitrogen and Chl *a* content data, presented in Table 4, were quite variable,
219 likely due to different physiological or trophic states of filamentous cyanobacteria. Based on
220 trichome densities and Chl *a* content, *Trichodesmium* Chl *a* was estimated as accounting for no more
221 than 21% of total Chl *a*. Atomic C/N ratio averaged 6.2 (range: 4.6 - 7.4), a value close to the slope
222 (6.75) of the C versus N regression line ($r^2 = 0.96$, $n=10$) and to the Redfield ratio.

223 To sum up, present surveys evidenced a shallower vertical distribution, higher growth rates and
224 more trichome aggregation in colonies during *Trichodesmium* blooms. These are the result of
225 environmental changes that are now considered.

226

227 3.3. General environmental conditions and causes of bloom developments (Fig. 6 and 7)

228 Both surveys started in austral spring, a season characterized by settled SE trade winds ($> 6 \text{ m}\cdot\text{s}^{-1}$)
229 and low rainfall. Sea temperature increased gradually until the beginning of February (Survey II),
230 reaching 29°C. Amidst this increasing trend, windy periods induced lower sea temperature and
231 higher salinity in spring. In summer and autumn, however, effect of wind on salinity was the
232 opposite, wind being associated to heavy rainfalls brought by tropical depressions, Kerry being one

233 of them in January 2005. Their major effect was high rainfall on the mainland and resulting lower
234 lagoon salinity (<35.00).

235 Nutrient concentration increases followed local rainfall, as recorded at Noumea Met station in
236 most instances. When they did not, they could be due, either to regional precipitation, not necessarily
237 recorded by coastal stations, or to wind events leading to resuspension of shallow sediments and
238 associated benthic nutrients (Muslim and Jones, 2003). The latter interpretation might indeed explain
239 the nitrate peaks of November 2003 (Fig. 6) and March 2005 (Fig. 7) and the peak of SRP of
240 January 2005, that took place during windy periods with no rainfall.

241 These increases in nutrient concentrations were not necessarily followed by total Chl *a* peaks for
242 reasons probably linked to vertical mixing or trace metal inhibitions, as discussed later for
243 *Trichodesmium*. Microphytoplankton contributed 15.9% ($s = 8.0$; $n = 119$) to total Chl *a* (>20 μm +
244 < 20 μm) in 2003 and 8.0% ($s = 7.2$; $n = 130$) in 2004-2005. Their main peak period occurred in
245 February 2005, following January nutrient inputs (Fig. 7). Diatoms dominated the
246 microphytoplankton, contributing 82% of the counts and dinoflagellates made most of the remainder
247 (11%).

248 *Trichodesmium sp.* started thriving at a water temperature, ranging from 24.2 to 28.6°C (Fig. 6
249 and 7). The vertical temperature gradient was variable from one bloom event to the other (0.05 to
250 0.84°C) and, therefore, did not seem to affect *Trichodesmium* development. The same conclusion
251 may be drawn for the vertical salinity gradient which ranged between 0.00 and 1.00.

252 A calm period (mean wind speed < 4 m s^{-1}) preceded development of *Trichodesmium* blooms,
253 except in January 2005 (Fig. 7) when the bloom started and continued during a windy period (mean
254 wind speed before the bloom commenced: 7.3 m s^{-1}). In this case, however, temperature was > 26°C.
255 In the other instances, windy periods did prevent *Trichodesmium* growth, as observed in November-
256 December 2003, for example. Higher turbidity was not systematically associated with wind speed
257 and rather low turbidity was the rule most of the time. We conclude turbidity, as measured by the
258 CTD sensor, is not a good descriptor of blooming conditions.

259 *Trichodesmium* always (5/5 cases) thrive after nutrient (NO₃, SRP) and Chl *a* concentration
260 increases, that occurred within 3-7 days before the peak abundance (Fig. 6 and 7). Since it happened
261 in all 5 cases, its occurrence was significant (sign test at $p < 0.10$). In addition, amplitude of the peak
262 abundance at 0.5 m was significantly correlated ($r_s = 0.879^*$) with NO₃ concentration at the start of
263 the bloom (Fig. 8), but not with SRP.

264 In order to test the nutrient and Chl *a* trigger hypothesis, we have considered whether all of their
265 increases were followed by blooms. During the two surveys, there were only two exceptions (5-7
266 November 2003 and 10-14 January 2005) when nutrient enrichments did not lead to any
267 development. In those two cases, the nutrient-Chl *a* increases were followed by windy periods (Fig.
268 6 and 7) which could, therefore, prevent bloom development. However, as seen previously, the wind
269 factor alone may not explain the second exception (January 2005) since temperature was $>26^\circ\text{C}$. A
270 lower than usual salinity may be the direct or indirect reason for the lack of *Trichodesmium* bloom as
271 it was also for the lack of microphytoplankton biomass increase at the same period (Fig. 7).

272 In summary, according to the present observations, development of *Trichodesmium* blooms
273 would follow nutrient and Chl *a* concentration increases with a 3-7 day lag, provided temperature is
274 $>26^\circ\text{C}$ and no heavy and sustained rainfall occurs. The same conditions apply to $> 24^\circ\text{C}$
275 temperatures, as long as wind speed is low ($< 4 \text{ m s}^{-1}$).

276

277 4. Discussion

278

279 4.1. Sampling strategy: are conclusions drawn from one station valid?

280 Study of *Trichodesmium* population dynamics employed high frequency sampling at Sta. SM,
281 assuming the same watermass and planktonic population were sampled for a long enough period.
282 This implies a rather long residence time, which is supported by the study of Jouon et al. (2006).
283 Thus, the "e-flushing time", a proxy of the water residence time, is 15-25 days at the entrance of the
284 bay, for a $\sim 8 \text{ m s}^{-1}$ (15.5 knots) SE wind velocity, and longer, of course, for lower wind speeds.

285 Besides, no significant tide effect was evidenced, especially in terms of salinity at 1m depth, because
286 water exchanges proceed from both the south and the east of Sta. SM (Fernandez et al., 2006). Note
287 also that *Trichodesmium* density and environmental parameter variations at Sta. SM displayed no
288 erratic trends, at least at a period of a week. It appears, therefore, that this station lent itself to a
289 satisfactory temporal description.

290 Moreover, present observations were representative of the surrounding lagoon waters, as seen for
291 the following parameters: (1) *Trichodesmium* spp. densities and microplankton abundance at 0.5 m
292 were in a fair agreement at the two stations even though *Trichodesmium* densities became higher out
293 of the bay from February 2005 on. (2) Proportions of the two *Trichodesmium* species or of other
294 microphytoplankton taxa were not significantly different at Sta. SM and O and (3) *Trichodesmium*
295 spp. growth rates, as provided by the ^{13}C method, were similar at the two stations. All these
296 arguments support the view of a reasonable spatial homogeneity in the sampled area, which is the
297 prerequisite to temporal variation descriptions.

298

299 4.2. *Trichodesmium* population characteristics

300 *Density as compared to other studies.* Trichomes have been observed systematically during the
301 two surveys and appeared to be a component of the usual phytoplankton population, as happens in
302 the GBR where they would occur in 83% of the samples (Jones, 1992). However, they never made
303 the bulk of the phytoplankton at Sta. SM, with their Chl *a* accounting for no more than 21% of the
304 total. Similar contributions were reported by Letelier and Karl (1996) and Dupouy et al. (2000) for
305 the open ocean of the tropical Pacific although higher values (11- 62%) were found in the Atlantic
306 by Carpenter et al. (2004).

307 Maximum *Trichodesmium* abundances (240 and 244 trich. L^{-1} in 2003 and 2004, respectively)
308 reported in this study, are rather modest compared to those found in the literature (Table 5) and this
309 may be explained by differences in both methods and environmental conditions. Use of a 20 μm
310 mesh, trichome transfer from the silk into the sedimentation cell, insufficient trichome sinking

311 (Lugomela et al., 2002) or use of a 5.74 L sampling bottle instead of a net (Chang, 2000) may be
312 reasons linked to the methodology. In addition, present results refer to samples at 0.5m while some
313 of the reported densities in the literature refer to the very near-surface (i.e. 0m), where much higher
314 concentrations could be associated with discoloured waters and accumulations along fronts or
315 Langmuir cells. It remains some of the high densities found in literature refer to subsurface maxima
316 (e.g. 12-24 m of Carpenter et al., 2004) and not to the very top surface, leading us to look for
317 environmental-related reasons. Thus, an element may limit growth of the filamentous cyanobacteria
318 and it is probably not dissolved iron, given its very significant concentrations in the SW lagoon,
319 ranging between 2 and 5 nM (unpublished data). These values are much above 1 nM, the
320 concentration given by Sanudo-Wilhelmy et al. (2003) for areas of non Fe-limited diazotrophy in the
321 Atlantic ocean. Phosphorus, however, could be the limiting element of primary production,
322 considering that its two possible sources are phosphorus depleted: runoffs from lateritic soils of this
323 part of New Caledonia (Tenorio *et al.*, 2005) and exchanges with the surrounding oligotrophic ocean
324 (Van Den Broeck et al., 2004). This limitation could explain both the modest trichome density
325 maxima and the low Chl *a* ($< 1 \text{ mg m}^{-3}$) concentrations (Fig. 6 and 7). Lastly, the observed positive
326 and significant correlation found between trichome and diatom densities ($r_s = 0.395^{**}$; $n = 41$)
327 would suggest no negative effect of the main component of the microphytoplankton on
328 *Trichodesmium*. In conclusion, modest trichome density maxima of the present study might result
329 from a combination of different factors, the main one being possibly phosphorus limitation.

330 *Specific composition.* *T. erythraeum* was the dominant species with temporary occurrence of *T.*
331 *thiebautii*. The two species have distinct morphometric characteristics and they displayed much
332 higher length variations at Sta. SM (Table 3) than those reported by Post et al. (2002) in the Gulf of
333 Aqaba: 300 - 800 μm for *T. erythraeum* and 1000 - 2000 μm for *T. thiebautii*.

334 *T. erythraeum* dominance at Sta. SM, as in the GBR lagoon (Jones, 1992; Muslin and Jones,
335 2003) or the Tanzanian coast (Lugomela et al., 2002 ; Bryceson and Fay, 1981), the Mississippi
336 plume (Eleuterius et al., 1981) or the Bresilian coasts (Satô et al., 1963) might lead to the conclusion

337 it would be found more often in coastal areas than *T. thiebautii*. This seems to be corroborated by
338 observations made in the open ocean off the east coast of New Caledonia, where *T. erythraeum*
339 makes only 20-25% of the *Trichodesmium* filaments, while *T. thiebautii* and *T. tenue* would make
340 40-45% and 30% of the total, respectively (Tenorio, 2006). Actually, each of the two species is able
341 to prevail in the open ocean. It is *T. erythraeum* in the Coral Sea and *T. thiebautii* in the Caribbean
342 Sea (O'Neil et al., 1996), *T. thiebautii* in the Kuroshio area (Saino and Hattori, 1980) and the
343 Atlantic (Carpenter et al., 2004), but *T. erythraeum* in the north Indian Ocean (Capone et al., 1998).
344 Therefore, it seems unlikely that dominance of the two species would rely on the "hemisphere" as
345 suggested by Capone and Carpenter (1999), but rather on physiological and/or environmental
346 conditions, such as those prevailing in rather closed coastal areas, that seem to be more appropriate
347 to *T. erythraeum*. Such a feature may be explained by a lower diazotrophic capacity of this species
348 (Carpenter et al., 1993), which would imply its preference for less oligotrophic areas than *T.*
349 *thiebautii*, on the one hand, and by less resistance of its vacuoles to pressure, which means less
350 adaptation to deep environments, on the other hand (Carpenter et al., 1993).

351 *Contribution of trichomes in colonies to total.* Most (98%) of the colonies were of the tuft type
352 and made of < 20 trichomes, on average. Similar numbers were found for *T. erythraeum* by
353 Bryceson and Fay (1981) and Capone et al. (1998) with respectively, 5.5-10 and 10-30 trich. col⁻¹.
354 But Carpenter et al. (2004), Letelier and Karl (1996) and Post et al. (2002) reported numbers > 100
355 for both *T. erythraeum* and *T. thiebautii*.

356 Also variable in the literature is the proportion of the number of aggregated trichomes to total
357 number of trichomes, or "bundleness" (Bryceson and Fay, 1981): ratios <20% were observed in the
358 North Pacific (Saino and Hattori, 1980 ; Letelier and Karl, 1996) and in the Atlantic (Orcutt et al.,
359 2001 ; Tyrell et al., 2003) while bundleness ranged from 25 to 90% off the Tanzanian coast
360 (Bryceson and Fay, 1981) and in the tropical Atlantic (Carpenter et al., 2004). Such variability is
361 discussed by the latter authors and could originate from damages to the colonies during the sampling
362 and sorting processes, too small a sampled volume, and/or a "fundamental difference in the state of

363 trichomes" between the different regions. Further, Orcutt et al. (2001), Fu and Bell (2003b) and Bell
364 et al. (2005) suggest there would be more of free trichomes during the optimum growth phase, and
365 more aggregation during the non optimum phases. Effect of wind speed and related water turbulence
366 on bundleness is invoked by Bryceson and Fay (1981) and would be negative, although the present
367 work does not support such a view, with bundleness not lower than usual during the windy January
368 2005 peak. More simply, present observations (Fig. 5) suggest bundleness would follow encounter
369 probability law, i.e. the higher the trichome density, the higher the probability, thus agreeing with the
370 view of Carpenter et al. (2004) of density-related bundleness. But, whatever the factors involved in
371 colony formation, high bundleness seems to be typical of *Trichodesmium* blooms and responsible for
372 higher nitrogen fixation rates (Saino and Hattori 1980; Bryceson and Fay, 1981; Letelier and Karl,
373 1998; Capone, 2001).

374 375 4.3. *Trichodesmium* bloom features: respective roles of growth and ascent

376 A striking result in the present study and many other publications, is the intensity of
377 *Trichodesmium* density increases, which appear to be inversely related to depth (Table 2): the
378 shallower the level, the greater the increase. Moreover, doubling times d'_{A} , calculated on biovolume
379 increases at 0.5, 3 - 4 and 6-7 m (Table 2), are much shorter than those reported in the literature for
380 growth (Table 6), while $d'_{A,10}$ at 10 m are in good agreement. From these observations, we conclude
381 $d'_{A,10}$ mainly result from growth and $d'_{A,i}$ of shallower depths, result from growth plus another
382 process. The latter is very likely the trichome ascent due to their positive buoyancy, a process which
383 delivers them more or less rapidly to the surface as they grow and which would explain the observed
384 vertical density gradient from the bottom to the surface (Fig. 3 and 4). Inversely, this gradient cannot
385 be ascribed to trichome sinking because microscopic observations showed healthy trichomes at all
386 depths and no or few trichomes were retrieved in sediment traps of the 2003 survey (Le Borgne et
387 al., 2004).

388 *Trichodesmium* positive buoyancy is well known and *T. erythraeum* has been reported more
 389 buoyant than *T. thiebautii* (Carpenter et al., 1993; Lugomela et al., 2002), which would explain the
 390 modest increases of February 2005 at 0.5 m when *T. thiebautii* contribution was significant. Such
 391 positive buoyancy, linked to the presence of gas vesicles (Capone et al., 1997), is different from that
 392 governing daily up and down vertical motion, with diel variations of the carbohydrate to protein ratio
 393 (Villareal and Carpenter, 2003).

394 In order to test whether trichome biovolume increases (ΔV_i) observed during the bloom periods,
 395 resulted from both their net growth (G_i) and ascent (A_i) from deeper levels and in order to estimate
 396 the respective contributions of the two processes, we used a simple model. Provided spatial
 397 heterogeneity is negligible, as compared to temporal variability (see 4.1.), observed ΔV_i at depth z_i ,
 398 between t_0 and t_f (see eqn (1)) may be ascribed to the sum of G_i and A_i :

$$400 \quad \Delta V_i = V_{f,i} - V_{o,i} = G_i + A_i \leftrightarrow \Delta V_i - (G_i + A_i) = 0 \quad (4)$$

401

402 with $G_i = \sum_{t=t_0}^{t_f} V_{o,i} e^{g_i t} \Delta t$, g_i , being the exponential net growth rate, Δt , the time interval.

403 Corresponding doubling time, $d_{A,i}$, is equal to: $d_{A,i} = \ln(2)/g_i$. Now, assuming trichome ascent, A_i ,
 404 would proceed at the same pace as growth and g_i be the same along the water column, A_i will be the

405 integral of G_i with depth: $A_i = \sum_{z=12}^{z_i} G_i \Delta z = \sum_{t=t_0}^{t_f} \sum_{z=12}^{z_i} V_{o,i} e^{g_i t} \Delta t \Delta z$, where Δz is the difference

406 between two sampled depths and 12 m, the Sta. SM depth.

407 Calculations of A_i and G_i have been made for $z_i = 0.5$ m, with the following Δz : 12-10 m, 10-7
 408 m, 7-4 m, 4-0.5 m in 2003 and 12-6 m and 6-0.5 m in 2004-2005. We solved equation (4), and
 409 considered the realism of computed doubling times, $d_{A,0.5}$ (Table 7). Except for the December 2004
 410 event, $d_{A,0.5}$ values (1.82 - 6.45 d) are within the range of the doubling times in the literature (Table
 411 6) and include the December 2003 biovolume doubling time at 10m, $d'_{A,10} = 2.9$ d (Table 2). The
 412 December 2004 low $d_{A,0.5}$ value (0.53 d) may be ascribed to an underestimated initial biovolume,

413 $V_{0,0.5}$, which is used in equation 4. Estimates of $A_{0.5}$ and $G_{0.5}$ show that most (87 to 99%) of
414 biovolume increase observed at 0.5m ($\Delta V_{0.5}$), would be due to $A_{0.5}$. Ascent contribution variability
415 might be linked to the mixing intensity of the water column, as illustrated by the windy January 2005
416 bloom, which had the lowest ascent contribution (Table 7).

417 In this type of calculation applied to peak periods, the growth rates (g_i) varied from one peak
418 abundance to another ($g_{0.5} = 0.11 - 0.38 \text{ d}^{-1}$). They varied also between non bloom and bloom
419 periods, as illustrated by the carbon-specific growth rates, provided by ^{13}C uptake measurements.
420 Indeed, a ratio of 5.1 - 7.4 (see Results) between bloom and "normal" periods may be computed,
421 leading us to the next question: what is the origin of g_i variations leading to bloom developments?

422
423 *4.4. Causes of bloom developments: favorable conditions and triggers*

424 These may be shared into necessary conditions which allow blooms to occur, and triggers that
425 effect g_i . The latter have been identified as nutrient increases during the 3-7 preceding days for all
426 the blooms, which supports observations made by Carpenter and Price (1977), Bell et al. (1999),
427 Muslim and Jones (2003) regarding a phosphate effect and Lugomela et al. (2002) about nitrate-
428 related blooms. Because diazotrophy is phosphorus-dependent (Sanudo-Wilhelmy et al., 2001; Fu
429 and Bell, 2003a; Mills et al., 2004), the role played by SRP in bloom developments seems to be
430 obvious. Less straightforward, however, may look the NO_3 increase effect on diazotrophic
431 cyanobacteria growth. In fact, di-nitrogen fixation is usually low or absent when other nitrogenous
432 compounds are available since it is high energy demanding and repressed by NH_4 (Mulholland and
433 Capone, 2000; Karl et al., 2002). At Sta SM, very low $^{15}\text{N}_2$ fixation rates were indeed measured on
434 *Trichodesmium* colonies during the March-April 2005 bloom (Le Borgne et al., 2006), suggesting
435 most of their nitrogen needs were fulfilled by other compounds. Ammonium and very small
436 dissolved organic nitrogen (DON) molecules, which are released or produced by the microbial loop,
437 seem to be the best candidates, although such a statement cannot be proved for a lack of
438 measurements.

439 Then we get to the following scenario which fits observations at Sta SM. Observed Chl *a*
440 increases are very quick responses to NO₃ inputs. Part of the Chl *a* increases may be due to diatoms,
441 which have a higher NO₃ uptake capacity than most phytoplankters (Sarhou et al., 2005), and likely
442 respond immediately to nutrient inputs and lead to higher concentrations of ammonium and small
443 DON molecules through bacterial remineralization and microzooplankton grazing. These are taken
444 up eventually by filamentous cyanobacteria with a 3-7 d time-lag between NO₃ increases and
445 *Trichodesmium* bloom initiation. The direct relation between *Trichodesmium* and diatom densities
446 reported before may therefore be interpreted in this way.

447 The nutrient-related scenario suffers a troubling exception: the most important nutrient increase
448 of the two surveys, in January 2005, was followed nor by *Trichodesmium* nor by
449 microphytoplankton development, while the whole area was invaded by waters of lower than usual
450 salinity. The salinity may not be incriminated *per se*, considering maximum *Trichodesmium* growth
451 occurs in the 30 - 37 psu range (Fu and Bell, 2003b; Bell et al., 2005). It may rather indicate
452 important terrigenous inputs of suspended sediments and heavy metals and their possible inhibition
453 of microphytoplankton (including *Trichodesmium*) growth. Inhibition could proceed through light
454 limitation due to a heavy load of suspended particles. No significant increase in turbidity (data not
455 shown) was observed, however. Different could be the heavy metals inhibitory effect and Fernandez
456 et al. (2006) have shown Fe, Ni, Cr, Co, Mn, Cu and Sr are issued from laterite weathering, and
457 carried along from several estuaries, south of Noumea (e.g., La coulee, Fig. 1). There may be also an
458 impact of the amount and type of organic matter from terrigenous origin, on the microphytoplankton,
459 although this topic has not been documented, yet.

460 Bloom triggers work, provided necessary conditions are met, *i.e.* appropriate iron concentration,
461 temperature and wind velocity. As seen above, iron is probably non limiting in the present study, but
462 sunny and calm weather conditions may be necessary ones as invoked in the past by many authors
463 (Eleuterius et al., 1981; Jones et al., 1992; Sellner, 1997; Lugomela et al. (2002) ; Muslim and
464 Jones, 2003 ; Carpenter et al., 2004). However, calm weather is not sufficient a condition because

465 there may be blooms during long periods of wind, as illustrated by the January 2005 event. Based on
466 this example, when temperature was $\sim 26^{\circ}\text{C}$, we conclude that blooms may occur during windy
467 periods, only if sea temperature is above this threshold value, thus confirming the Carpenter and
468 Capone (1992) analysis on bloom conditions. Inversely, below 26°C , a low wind velocity appears to
469 be necessary for bloom development. The minimum temperature for them to occur is 24°C in the
470 present study, which is less than the 25°C threshold of Chen et al. (2003). Combination of wind and
471 temperature effects can be interpreted as the result of two antagonist actions on growth rates,
472 positively correlated to temperature and negatively to wind induced-mixing. It follows that, when
473 temperature reaches a threshold value, its effect on growth will overcome the negative effect of
474 mixing.

475

476 4.5. Decline origin

477 One of the striking points in the two surveys, was the brevity of the bloom decline phase, as for
478 the growing one. Decline may be ascribed to mortality and to a recruitment diminution due to a
479 growth rate slowdown as for any population, even though they might have common origins. In the
480 present study, most blooms ended with very low nutrient concentrations ($<0.030\ \mu\text{M NO}_3$ at the end
481 of the December 2003 and February 2005 events and $<0.030\ \mu\text{M SRP}$ for those of February and
482 April 2005), which suggests nutrient exhaustion represents one of the decline causes, as mentioned
483 already by Lenés et al. (2005) and Moutin et al. (2005). This process could reduce growth rates, in a
484 way opposite to the nutrient increase effect described above, or induce mortality through a
485 "Programmed Cell Death" (PCD) pathway (Berman-Frank et al., 2004, 2007). Viral lysis (Ohki,
486 1999) and PCD, caused by other factors than nutrient depletion, as listed by Berman-Franck et al.
487 (2004), are other possible processes. Inversely, intense grazing is likely excluded, considering the
488 low *Macrosetella* densities that were observed and their lack of relationship with trichome
489 concentrations. But the present observations on trichome ascent and surface accumulation during
490 bloom periods with no obvious sinking or downward migration do not exclude massive destruction

491 by solar radiation as infra-red or ultra-violet rays, independently of other processes. The solar effect
492 is likely linked to the sea surface agitation, with a more rapid destruction during calm sea and low
493 wind conditions. On the whole, apart from trichome viral lysis or PCD, abrupt decay could be the
494 result of ascent to the very top surface, solar destruction and lack of new trichome formation for
495 nutrient exhaustion.

496

497 *4.6. Summary of Trichodesmium bloom dynamics as illustrated by the November - December 2003* 498 *event*

499 A summary of present conclusions on the bloom triggering factors (nutrient concentration
500 increases), necessary conditions (combination of wind and temperature effects) and causes of bloom
501 decay can be illustrated by the Nov-Dec 2003 event. Although it was made of three peak
502 abundances, the following description shows it was only one event (Fig. 6).

503 An increase in nutrient concentrations was observed on 19 - 21 November for NO_3 and 17
504 November for SRP. At that time, water temperature was around 24°C and development could not
505 start before the wind velocity diminished. This occurred on the 23rd and was likely responsible for
506 the observed trichome density and biovolume increase between 21 and 28 November. On 29
507 November, while temperature was now 25°C , but still $< 26^\circ\text{C}$, the wind started blowing again, which
508 resulted in the first trichome density diminution between 28 November and 3 December. The wind
509 slowed down between 4 and 5 December, allowing another *Trichodesmium* increase with a peak on
510 the 5th. Calm weather conditions lasted until the 11th, allowing trichome ascent and mortality at 0m
511 with the resulting biovolume decrease at all depths between 5 and 8 December and a small
512 development between 10 and 12 December, once temperature had reached 26°C . The bloom ended
513 when NO_3 became exhausted ($< 0.005 \mu\text{M}$ at 0.5m), i.e. from the 10th, on.

514

515 **5. Conclusion**

516 Photosynthetic filamentous cyanobacteria of the *Trichodesmium* genus, display the same
517 physiological processes as the other co-occurring phytoplankton and have the same requirements. In
518 particular, dissolved nutrients are primarily involved in their growth processes. But the present
519 surveys have shown major differences, with very steep density variations and changes in the
520 cyanobacterial vertical distribution during the blooms, both being the result of the trichome positive
521 buoyancy. This characteristic has two consequences: (1) most of the biomass accumulation will
522 happen at 0m and not in the water column; (2) once they have reached the very top surface,
523 trichomes will remain there, and eventually be destroyed. In addition, because of their toxicity, the
524 grazing control of the *Trichodesmium* population appears to be restricted to a few species, like
525 harpacticoid copepods (Hawser et al., 1992) and these were not abundant in the studied area. These
526 two features of the *Trichodesmium* population dynamics, i.e. effects of positive buoyancy and low
527 grazing losses, make it quite different from what happens to the rest of the phytoplankton
528 community. Such a statement, however, is based on observations of a *T. erythraeum* dominated
529 population, which is known to be more buoyant and less diazotrophic than *T. thiebautii*. Therefore,
530 bloom causes and characteristics of *T. thiebautii* populations might be different to some extent and
531 would deserve a similar study, related to the toxicity issue, this species being considered as more
532 toxic than *T. erythraeum* (Sellner, 1997; Landsberg, 2002).

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Tables

Table 1

Main characteristics of the five *Trichodesmium erythraeum* blooms (* denotes presence of *T. thiebautii*).

Period	Number of peaks	Difference in time evolution at 6 and 0.5 m	Total duration (d)	Maximum at 0.5 m (trich. L ⁻¹)	Maximum at 6-7 m (trich. L ⁻¹)
24 Nov - 14 Dec 2003	3	No	18	238.7	47.4
13 - 20 Dec 2004*	1	at 6m, 2 d before	7	244.6	20.7
3 - 14 Jan 2005	1	No	11	81.5	14.8
4 - 18 Feb 2005*		No	14	10.1	19.4
29 Mar - 7 Apr 2005	1	at 6m, 4 d before	9	156.5	44.3

Table 2

Exponential net increase rates (k_i in d^{-1}) and corresponding doubling times ($d'_{A,i}$ in d) in parentheses, from changes in *Trichodesmium* population biovolume, at different depths (equations 1 and 2).

2003	<i>T. erythraeum</i>				2004-2005	<i>T. erythraeum</i>		<i>T. thiebautii</i>	
	10m	7m	4m	0.5m		6m	0.5m	6m	0.5m
24-30 Nov		0.36 (1.9)	0.43 (1.6)	0.36 (1.9)	13-12 Dec	0.39 (1.8)	3.29 (0.2)	1.51 (0.5)	2.20 (0.3)
3-9 Dec	0.24 (2.9)	0.68 (1.0)	0.61 (1.1)	0.72 (1.0)	3-14 Jan	0.98 (0.7)	1.13 (0.6)		
11-14 Dec			0.15 (4.6)	0.40 (1.7)	4-18 Feb	0.19 (3.7)	0.52 (1.3)	0.23 (2.9)	0.94 (0.7)
					29 Mar - 4 Apr	0.67 (1.0)	0.59 (1.2)		

Table 3

Sizes of *T. erythraeum* and *T. thiebautii* during the 2003 and 2004-2005 surveys at Sta SM. Means \pm standard deviations, and range (in parenthesis) of size parameters for n observations.

		Width (μm)	Length (μm)	Volume ($10^3 \mu\text{m}^3$)
<i>T. erythraeum</i>				
2003	0.5m - 10 m	11.0 ± 0.6	477 ± 190	45.3 ± 20.7
	$n = 2612$	(5.5 - 22.0)	(56-1489)	(5.3-426.1)
2004-2005	0.5m - 6 m	10.5 ± 1.4	550 ± 344	48.4 ± 34.8
	$n = 6362$	(5.5 - 22.0)	(60 - 2675)	(4.1 - 448.6)
<i>T. thiebautii</i>				
2003	0.5m - 10m	11.3 ± 2.3	822 ± 499	67.1 ± 26.0
	$n = 8$	(8.2 - 16.5)	(440 - 2031)	(25.6 - 96.9)
2004-2005	0.5m - 6m	5.7 ± 1.0	847 ± 440	22.0 ± 16.4
	$n = 1618$	(5.5 - 11.0)	(57 - 3386)	(1.3 - 186.4)

Table 4

Trichodesmium carbon, nitrogen and chlorophyll *a* content of the 2004-2005 survey: means, standard deviations (sd) and ranges.

	Per colony			Per trichome			Ratio	
	Chl <i>a</i>	C	N	Chl <i>a</i>	C	N	C/N	C/Chl <i>a</i>
	(ng)	(μ g)	(μ g)	(ng)	(ng)	(ng)	(mol/mol)	(g/g)
Mean \pm sd	7.84 \pm 5.16	1.25 \pm 0.29	0.20 \pm 0.12	0.44 \pm 0.26	103.1 \pm 36.1	16.8 \pm 5.7	6.2 \pm 0.8	265.1 \pm 161.5
Range	2.41 – 14.9	0.79 – 1.57	0.15 – 0.39	0.17 – 0.92	61.5 – 191.0	10.7 – 28.0	4.6 – 7.4	87.0 – 486.5

Table 5

Trichodesmium spp. densities (as trich. L⁻¹) literature review (Note: references considering colony abundance are not presented. Specific difference between *T. erythraeum* (§) and *T. thiebautii* (*) are reported when available. m: mean value; max: maximum value.

Region	Depth	Density	Author(s)
Sargasso and Caribbean seas	0 m (range)	0-49*	Carpenter and Price (1977)
	15-200 m (range)	0-294*	
Kuroshio (Japan)	0-100 m (max)	40 – 50000	Saino and Hattori (1980)
Great Barrier Reef lagoon	bloom (max)	50,000	Revelante and Gilmartin (1982)
Cleveland Bay (Australia)	inshore "blooms" (m)	8515 [§]	Jones (1992)
	offshore "blooms" (m)	3300 [§]	
HOT Station (Hawaii)	0-45 m (range)	11 – 84	Letelier and Karl (1996)
Southwest tropical Pacific	0 m - Fiji (max)	10,000	Dupouy et al. (2000)
	0 m New Caledonia (max)	1000	
Tanzania coast	0 m "bloom" (range)	38,000 - 120,000 [§]	Lugomela et al. (2002)
	0 - 20 m (range)	0 - 63,000 [§]	
Gulf of Aqaba	0-90 m (range)	0.05 - 2	Post et al. (2002)
South China Sea	0 m (m, range)	77 (0 - 962)	Chen et al. (2003)
Magnetic island (Australia)	surface (range)	9 - 102,000	Muslim and Jones (2003)
	15 m (m)	13,000	
North and South Atlantic	7 m (m)	300	Tyrell et al. 2003
	(max)	2200	
Dampier Archipelago (Australia)	0-20 m (m)	1800 [§]	Negri et al. (2004)
North tropical Atlantic	"surface" (m)	222 – 292*	Carpenter et al. (2004)
	"surface", peak period (m)	2250 (up to 10867)*	
Bay of Ouinné (New Caledonia)	0 - 45 m (m)	0.7 - 17 [§] ; 0.6 – 29*	Tenório et al. (2005)
Loyalty channel (New Caledonia)	0 - 60 m (range)	0 - 1011 [§] ; 4 – 2450*	Tenorio (2006)
Bay of Sainte Marie (New Caledonia)	0.5 m (range)	0 - 240 [§] ; 0 – 34*	Present study
	6 m (range)	0 - 47 [§] ; 0 – 18*	

Table 6

Literature review of *Trichodesmium* carbon-specific doubling times (d).

Reference	Conditions	Temperature (°C)	Doubling time (d)
Carpenter and Romans (1991)	natural populations, tropical -subtropical Atlantic	26.5 - 28	1.8 - 18
Carpenter and Capone (1992)	Review	>27	3 - 6
Carpenter et al. (1993)	natural populations <i>T. erythraeum</i>	26.5 - 28	3.8
	<i>T. thiebautii</i>		3.0
Prufert-Bebout et al. (1993)	Cultures		3.0*
Mulholland and Capone (2000)	Cultures		2.2
	natural populations		3.8 - 198
Orcutt et al. (2001)	natural populations, Bermuda		2
Bell et al. (2005)	Cultures	25 ± 3	2.3 – 3.5
Mulholland and Bernhardt (2005)	continuous culture	28	3 - 10
Le Borgne et al. (2006)	natural populations, bloom, Bay of Ste Marie,	27 - 29	4.1 - 5.8
	<i>idem</i> , non-bloom conditions	27 - 29	13.9 - 40.8

* inferred from their 0.23 division d⁻¹ growth rate

Table 7

Values of the trichome exponential net growth rate ($g_{0.5}$) and their corresponding doubling time ($d_{A,0.5}$) satisfying equation 4, for $z = 0.5$ m. Percent contributions of net growth ($G_{0.5} / \Delta V_{0.5}$) and ascent from deeper levels ($A_{0.5} / \Delta V_{0.5}$) to observed biovolume increases at 0.5 m, $\Delta V_{0.5}$.

Period	$g_{0.5}$ (d^{-1})	$d_{A,0.5}$ (d)	$A_{0.5} / \Delta V_{0.5}$ (%)	$G_{0.5} / \Delta V_{0.5}$ (%)
24-28 Nov 2003	0.18	3.86	87.3	12.5
3-5 Dec 2003	0.38	1.82	97.6	2.4
10-12 Dec 2003	0.12	5.57	87.3	12.6
13-15 Dec 2004	1.32	0.53	98.2	1.8
3-5 Jan 2005	0.11	6.45	87.0	13.0
25 Mar - 4 Apr 2005	0.17	4.14	98.9	1.1

Figure captions

Fig. 1. Sampling station locations at the entrance (Sta. SM) and off (Sta. O) the Bay of Ste Marie.

Fig. 2. *Trichodesmium* abundance as trichome numbers versus biovolume during the 2003 and 2004-2005 surveys. Slopes (b) of the linear regressions are : $b = 44240$ in 2003 and $b = 45084$ in 2004-2005, respectively.

Fig. 3. Temporal variations in *T. erythraeum* biovolume at Sta. SM four sampled depths, during Survey I (2003). Note : *T. thiebautii* densities, contributing for <1% of the total, have not been represented on the figure.

Fig. 4. Temporal variations in *T. erythraeum* and *T. thiebautii* biovolume at Sta SM two sampled depths, during Survey II (2004-2005). Less frequent observations were made at Sta. O, 0.5 m

Fig. 5. Relationships between trichome in colony (y) and free trichome (x) densities during the 2003 and 2004-2005 surveys.

Fig. 6. Temporal variations of environmental variables and *Trichodesmium* spp. densities at Sta. SM during Survey I. Temperature and salinity refer to the 1 m depth. Nitrate (NO_3), Soluble Reactive Phosphorus (SRP) and chlorophyll *a* (Chl *a*) concentrations are averaged through the 0-12 m water column, wind velocities, on 24 h. Significant rainfall ($>10\text{mm d}^{-1}$) are indicated by vertical arrows. *Trichodesmium* spp. abundances are integrated over 10m.

Fig.7. Same as Fig. 3, but for Survey II.

Fig. 8. *Trichodesmium* peak amplitude versus nitrate concentration at the beginning of the five blooms.

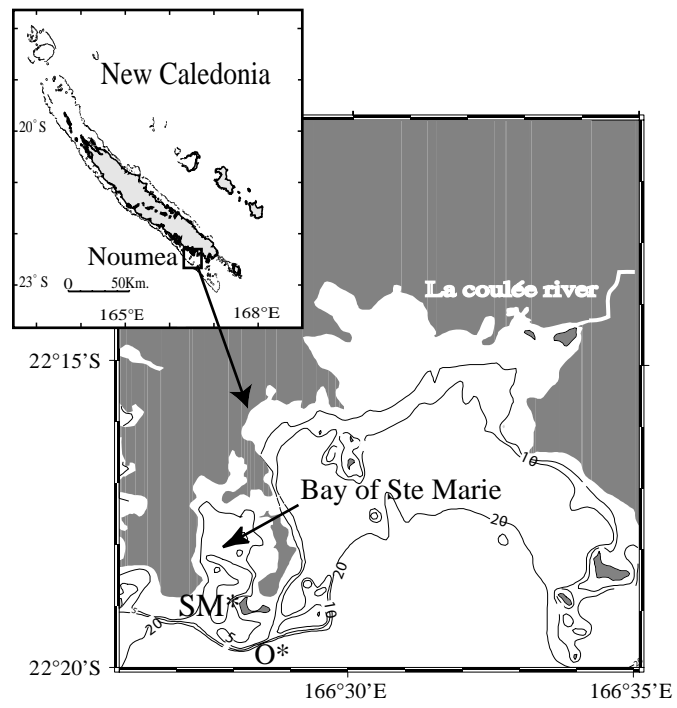


Fig. 1

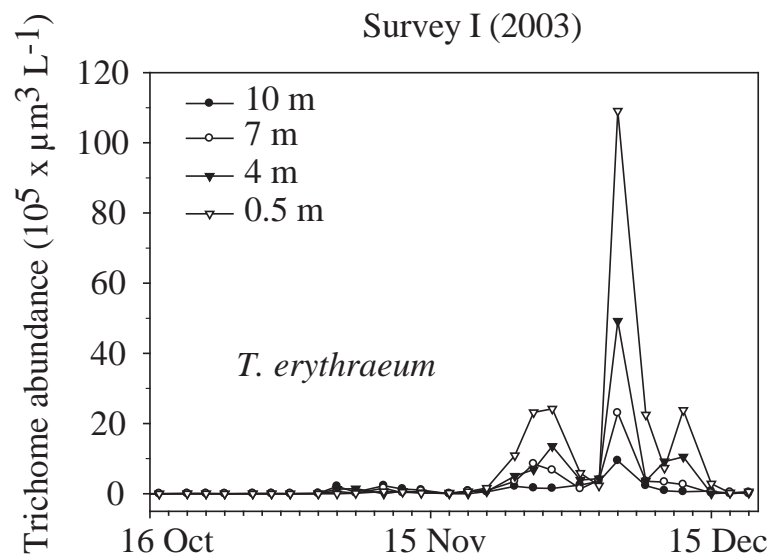


Fig. 3

Survey II (2004-2005)

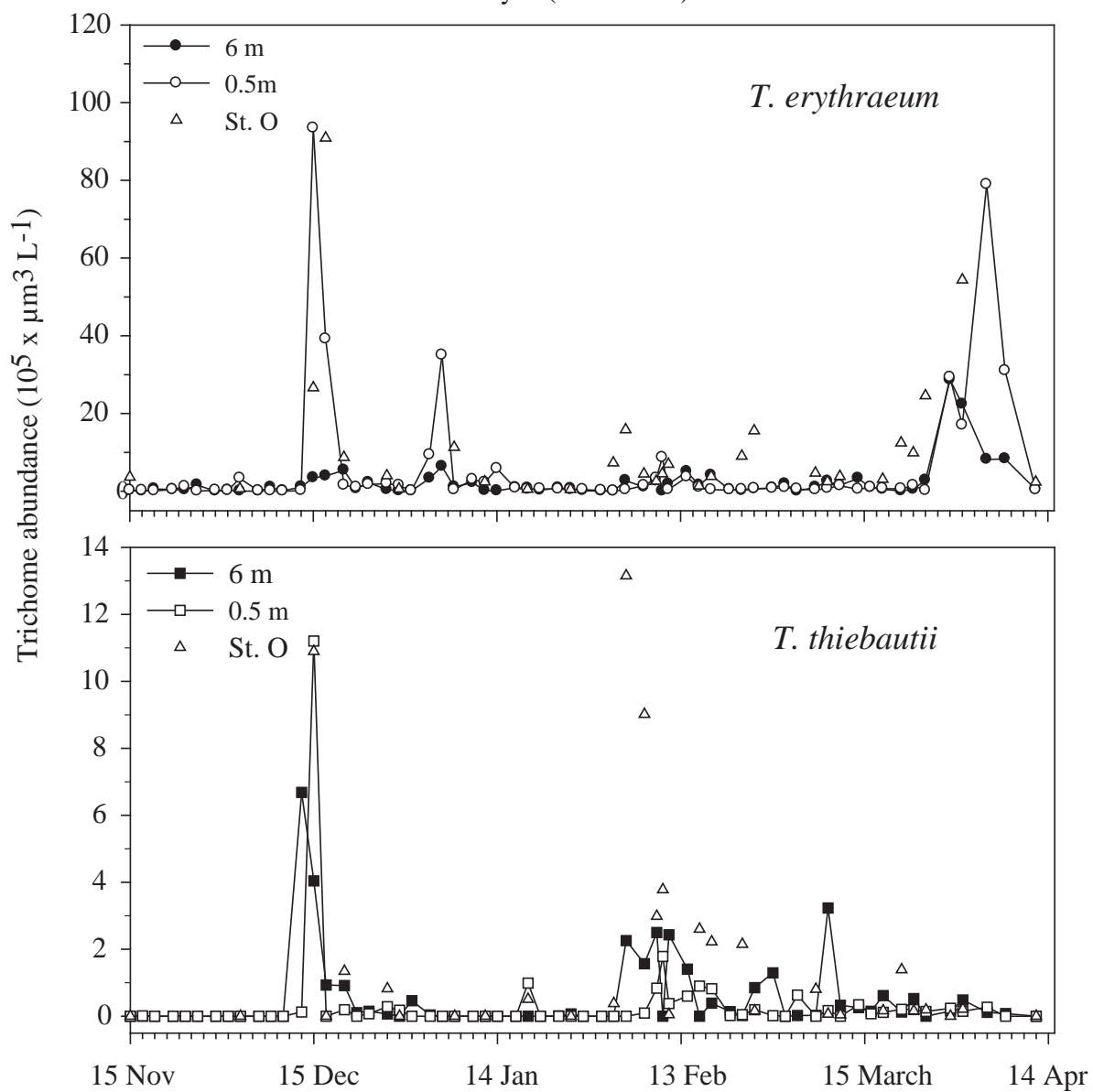


Fig. 4

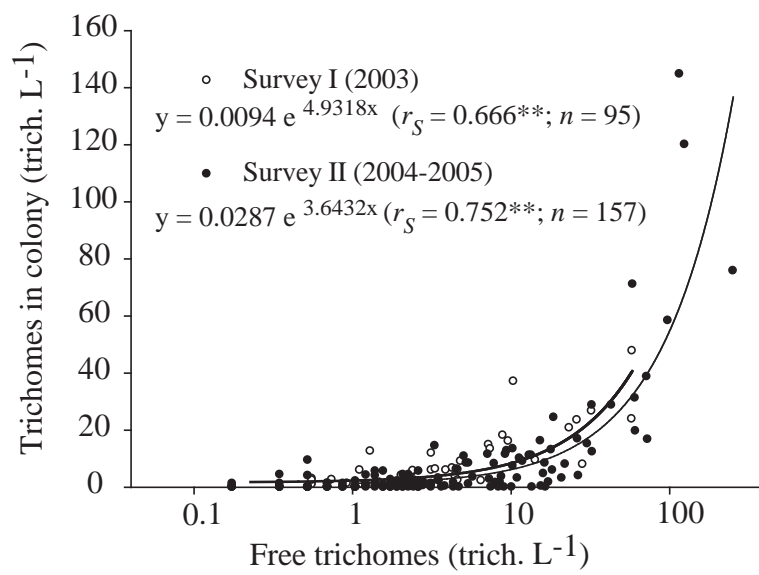


Fig. 5

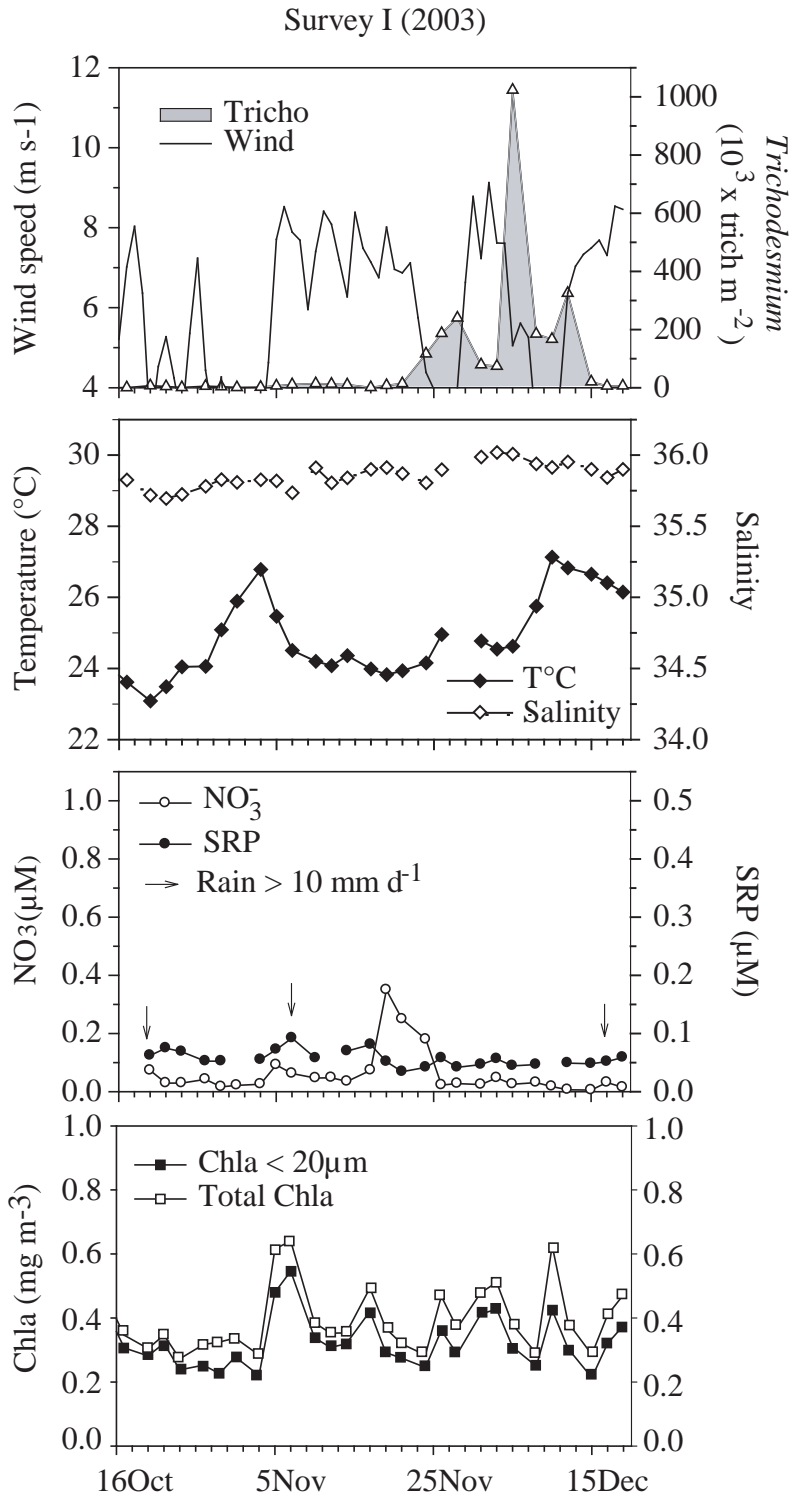


Fig.6

Survey II (2004-2005)

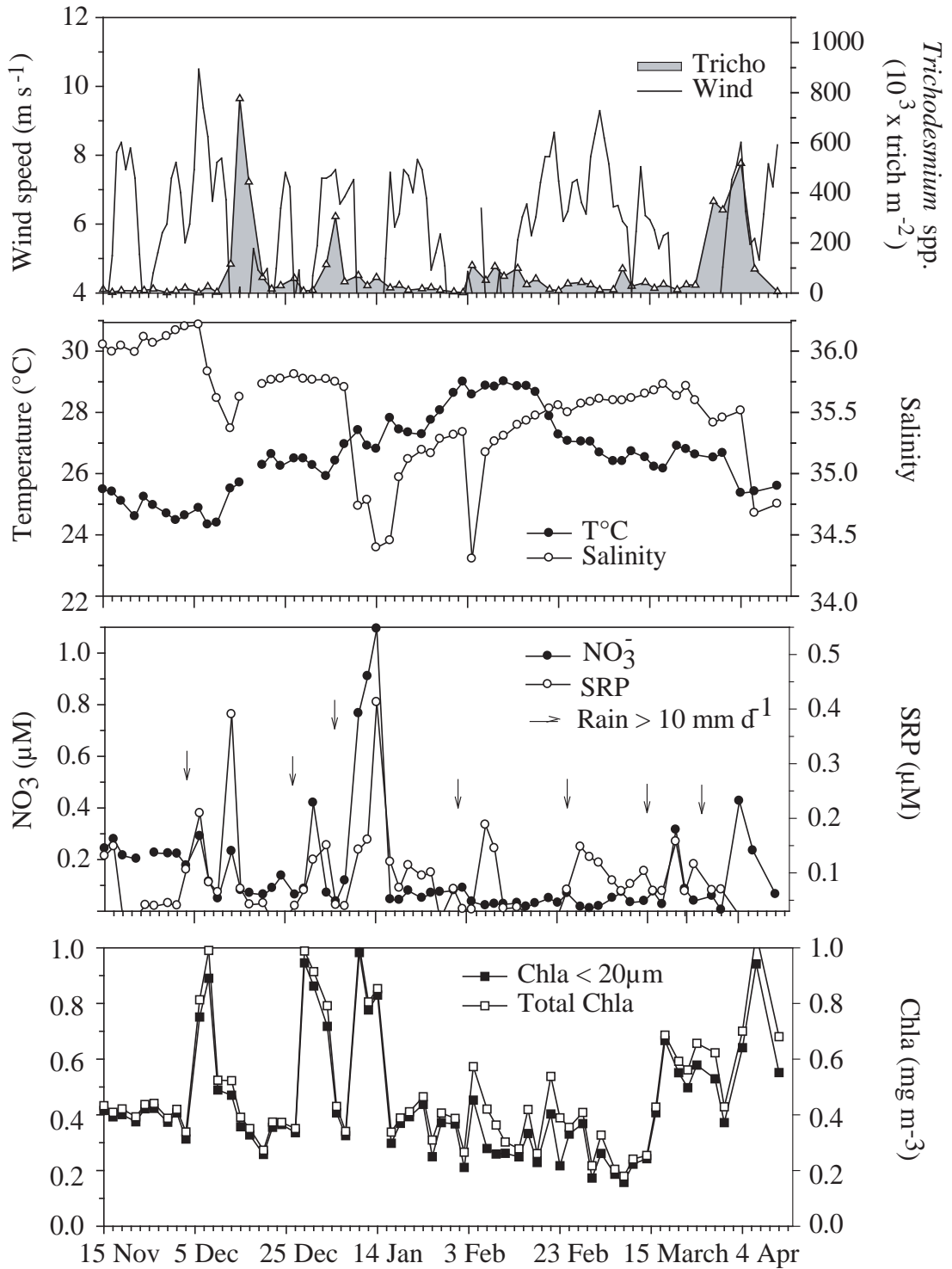


Fig. 7

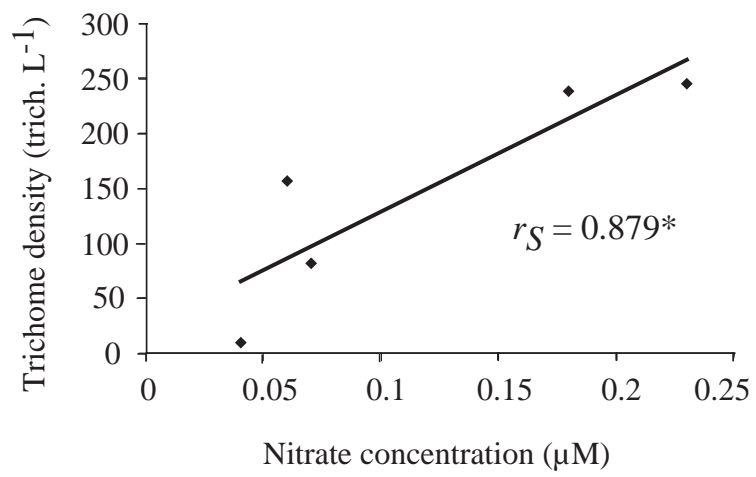


Fig. 8