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Buoynancy and growth characteristics of three positively buoyant marine diatoms

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ABSTRACT The growth rates and buoyancy properties of 3 oceanic diatoms in the genus Rhizosolenia were examined at light levels from 8 to 211 µmol quanta m⁻² s⁻¹. Maximum growth rates ranged from 0.37 to 0.78 divisions d⁻¹ with saturation occurring between 29 and 164 µmol quanta m⁻² s⁻¹. Severe growth rate depressions were noted in R. acuminata and R. formosa at irradiances above 50 to 155 µmol quanta m⁻² s⁻¹. In all 3 species the percentage of positively buoyant cells was inversely related to light intensity. In R. formosa both growth rate and tolerance to high light levels decreased substantially as cell size decreased. Batch culture C:chlorophyll ratios (130 to 261) replicated values found in field Ethmodiscus and Rhizosolenia mats, and suggest that the elevated C:chlorophyll ratios found in buoyant, oceanic phytoplankton are typical of healthy cells. Calculations suggest that carbohydrate ballasting can account for buoyancy changes and that these reserves are adequate to support dark NO₃ uptake. Under steady-state conditions in situ, the observed growth and buoyancy properties would lead to subsurface population maxima in all 3 species. However, the dynamic light-related buoyancy changes probably occur on a shorter time scale than these batch culture experiments. These results indicate that vertical migration is a property basic to these diatoms life history strategy, and, like multispecies Rhizosolenia mats, solitary Rhizosolenia chains transport new nitrogen to the euphotic zone in oligotrophic seas.

KEY WORDS: Rhizosolenia · Buoyancy · Chemical composition · Growth rates · New nitrogen · Suspension · Vertical migration

INTRODUCTION

Rhizosolenia formosa H. Peragallo, R. acuminata (H. Peragallo) H. Peragallo and R. castracanei H. Peragallo are chain-forming marine diatoms which thrive under the low-turbulence, nutrient-poor conditions typical of the open ocean gyres (Karsten 1905, 1907, Sundstrom 1986). These large diatoms (up to 10⁶ µm³) should be at a severe competitive disadvantage in obtaining nutrients relative to small (<5 µm) phytoplankton due to their lower surface/volume ratio (Chisholm 1992). Other large phytoplankton in the same environment, such as Pyrocystis noctiluca and the multispecies Rhizosolenia mats, overcome this size-related disadvantage by obtaining nutrients from below the nutricline through a buoyancy-mediated vertical migration (Rivkin et al. 1984, Hayward 1993, Villareal et al. 1993). This strategy exploits spatially uncoupled light and nutrient fields (Ganf & Oliver 1982) and avoids direct nutrient competition with smaller size fractions. Positive buoyancy is a prerequisite for vertical migration, but has traditionally been considered unlikely or aberrant in diatoms due to the relatively high density of the frustule (Smayda 1970, Walsby & Reynolds 1980, Round et al. 1992). However, numerous diatoms characteristic of the open sea such as Ethmodiscus spp. and solitary Rhizosolenia spp. are capable of positive buoyancy (Villareal 1988, 1992, Villareal & Carpenter 1989), and the evidence strongly suggests that they also migrate vertically (Villareal 1992, Villareal & Carpenter 1994, Villareal & Lipschultz 1995).
This migratory behavior has strong relevance to oceanic biogeochemistry in addition to basic ecological problems. Since vertical migrants directly transport NO$_3^-$ into the upper mixed layer, they circumvent the NO$_3^-$ trap at the base of the euphotic zone (Banse 1987) and represent a source of "new nitrogen" (Dugdale & Goering 1967) to the euphotic zone (Villareal et al. 1993). Calculations suggest that vertically migrating Rhizosolenia mats could transport nitrogen equal to 2–27% of the turbulent upward flux of nitrate in the central Pacific gyre (Villareal et al. 1993). Such inputs would supply some fraction of the new nitrogen required to support the sub-surface oxygen maximum common in oceanic gyres (Hayward 1994); however, quantitative estimates are difficult due to the limited abundance data available. Moreover, the time scale for migration is uncertain but is almost surely less than the carbon specific doubling rates of these taxa in surface waters (Villareal & Carpenter 1994).

Such life history parameters are difficult to determine from field populations and diatom vertical migration has been studied using internal NO$_3^-$- pools, buoyancy changes and $\delta^{15}$N signature to infer behavior and past history (Villareal et al. 1993, Villareal & Lipschultz 1995). However, mechanistic understandings can only be derived from laboratory studies that evaluate to which nutritional or environmental variables buoyancy is responding. In addition, the relationships between light, chemical composition and growth rate require clarification since the literature suggests these large phytoplankton have unusually high C:chlorophyll ratios (Villareal & Carpenter 1989, 1994). We therefore undertook a series of experiments to examine selected growth, chemical composition and buoyancy properties of Rhizosolenia formosa and R. acuminata, R. castreacanei. We wanted to determine if vertical migration by single-species diatom chains was feasible, and to examine the buoyancy responses to various environmental factors which might permit this migration. To investigate the influence of light on buoyancy, we measured the buoyancy of all 3 species under steady-state, nutrient-replete conditions over a range of light intensities. This experiment was repeated 3 times over a 10 mo period using the R. formosa clone to examine how changing cell size affected buoyancy/light interactions. Compositional characteristics of R. formosa and R. acuminata at different light levels were also examined. The effects of changing light levels on buoyancy were examined in R. formosa over a diel cycle and for a period of several days. All of the Rhizosolenia spp. in this study are capable of positive buoyancy and are sometimes members of Rhizosolenia mats, thus the results may also provide insight into the physiological mechanisms of mat vertical migration (Villareal et al. 1993).

MATERIALS AND METHODS

Culture and analytical methods. Rhizosolenia formosa H. Peragallo (clone B8), R. acuminata (H. Per) H. Peragallo (clone G3) and R. castreacanei H. Peragallo (clone G12) were isolated from the Sargasso Sea near Bermuda in August 1992 (R. formosa) or in September 1993 (others). Cells were hand-collected using SCUBA techniques and isolated in modified MET 44 medium (Schöne & Schöne 1982, Villareal 1991). Cell cultures were unialgal and grown under a 12:12 h light:dark cycle in 2 or 2.5 l polycarbonate bottles at 21°C. Species identification was made according to Sundström (1986).

Cell buoyancy was measured using a petri dish. A homogeneous sample was poured into the bottom half of the petri dish and the inverted lid was gently slid across the top so as to exclude air bubbles. After turbulence subsided (~5 min), most cells rested against the top or bottom dish; their buoyancy was operationally defined as positive or negative, respectively. A small percentage of cells drifting between the top and bottom dish were defined as neutrally buoyant. Cells were counted using an Olympus SZH stereoscope. Cell size was measured with an Olympus BH-2 compound light microscope. Cell volumes were calculated assuming a cylindrical shape.

Nitrate, phosphate, silicate and ammonium concentrations in the medium and cell sap were measured using a TRAACS 800 autoanalyzer. To measure extracellular nutrient concentrations, samples were syringe-filtered (0.8 µm pore size polycarbonate filter) and frozen in an acid-washed plastic scintillation vial for later analysis. For measuring intracellular nutrient concentrations, samples were filtered onto 0.8 µm pore size polycarbonate filters, rinsed with approximately 5 ml of a nitrate-free synthetic seawater and frozen in 5 ml of distilled water in acid-washed plastic scintillation vials. Upon thawing, the vials were placed in a boiling water bath for 5 min to rupture cells and extract intracellular nutrients.

Samples for total protein, carbohydrate, particulate carbon and nitrogen were filtered onto pre-combusted (2 h at 400°C) GF/F glass fiber filters. Chlorophyll samples were collected on GF/F filters. Carbohydrate and chlorophyll were frozen in foil and protein samples were frozen in plastic sample vials. Particulate C and N were frozen in pre-combusted aluminum foil and analyzed using a Perkin-Elmer 2400 CHN Elemental Analyzer. Protein samples were boiled in 0.1 N NaOH for 1 h and then measured using a bicinchoninic acid assay (Smith et al. 1985, Sigma procedure no. TPRO-562). Particulate carbohydrate was measured using an anthrone reagent standardized to glucose (Parsons et al. 1984a). Chlorophyll was measured fluorometrically.
in acetone extracts (Parsons et al. 1984b). Light intensity was measured inside the culture bottles using a Biospherical Instruments, Inc., QSP170B quantum sensor with 4π collector. Cultures were grown under cool white fluorescent light. Blue light culture bottles were wrapped in a blue plastic screening which had a broad wavelength transmission (380 to 550 nm) with a peak transmission (near 80%) at 455 to 465 nm.

Growth rates (μ) were calculated as the slope of the linear portion of ln transformed cell numbers plotted against time. Divisions d⁻¹ were calculated as μ/ln(2). Preliminary studies showed that daily sampling inhibited normal cell division, so the minimum sampling interval was every other day.

**Experimental methods.** Cultures of each species were pre-adapted for 10 d at different light levels ranging from 8 to 211 μmol quanta m⁻² s⁻¹. White light simulated near-surface light quality at sea, and the blue light simulated the light quality that phytoplankton experience lower in the euphotic zone. The transition point varied from 30 to 40 μmol quanta m⁻² s⁻¹ and was based primarily on the maximum irradiance that could be used with blue screening which did not overheat the samples. Buoyancy and cell numbers of all 3 species were measured periodically through the exponential growth phase. Carbon, nitrogen, carbohydrate, protein, chlorophyll, internal nitrate pools and medium nitrate concentrations were measured from *Rhizosolenia formosa* and *R. acuminata* during the exponential growth phase. The apparent production rate (P₈, mg C mg⁻¹ chlorophyll a h⁻¹) was calculated using the observed growth rates, average C and chlorophyll a values, and the photoperiod.

The experiment described above was repeated 3 times over a 10 mo period at 3 different cell sizes to examine the influence of cell size on *Rhizosolenia formosa* buoyancy patterns. Volume declined by 54% from the largest (1.18 × 10⁷ μm³) to the smallest cell size (5.69 × 10⁶ μm³). In another experiment, a *R. formosa* culture growing under white light at 143 μmol quanta m⁻² s⁻¹ was divided equally between 2 polycarbonate bottles. A control remained at the relatively high light level at which the original culture had been growing; the other bottle was moved to a low-light environment at 42 μmol quanta m⁻² s⁻¹. Buoyancy and nutrient concentrations in the cell sap and medium were followed for 6 d. To investigate diel patterns, a *R. formosa* culture grown in white light at 154 μmol quanta m⁻² s⁻¹ was sampled every 3 h for a 24 h period for buoyancy, cell sap nitrate, population and the number of cells recently divided. Recently divided cells were defined as cells in a chain with nuclei less than 1 cell diameter apart. This method allowed recently divided cells to be counted using the stereoscope at the same time as buoyancy measurements were made; thus, sample volume was minimized and the buoyancy of the recently divided cells could be recorded. To prevent cell damage from frequent sampling, the culture was divided equally among 9 polycarbonate bottles 12 h before the experiment began, and 1 bottle was sampled at the end of each 3 h period.

**RESULTS**

The growth rate of *Rhizosolenia formosa* (cell diameter = 129 μm, average cell volume = 6.58 × 10⁶ μm³) was saturated from 23 to 58 μmol quanta m⁻² s⁻¹ with an observed maximum of 0.37 divisions d⁻¹ at the relatively low light level of 29 μmol quanta m⁻² s⁻¹ (Fig. 1A). Cultures at 156 and 211 μmol quanta m⁻² s⁻¹ died at low population densities (<500 cells l⁻¹) apparently due to photoinhibition. The culture growing at 106 μmol quanta m⁻² s⁻¹ grew slowly (0.13 divisions d⁻¹) and also died before nutrients became limiting (at ~4500 cells l⁻¹). The cultures at 156 and 211 μmol quanta m⁻² s⁻¹ died after approximately 20 d, and the culture at 106 μmol quanta m⁻² s⁻¹ died after about 30 d. In all 3 high-light cultures cells underwent several divisions and appeared healthy during the first 10 d of the experiment. This light-induced mortality

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![Diagram](image-url)

**Fig 1** Growth rates of 3 *Rhizosolenia* spp. in cultures grown at different light levels. Growth rates were calculated from the average of duplicate cell counts.
buoyant cells were observed at the 2 highest light levels (156 and 211 µmol quanta m\(^{-2}\) s\(^{-1}\)). This relationship between light level and buoyancy (Fig. 2A) had the best fit linear equation, \(\% \text{POS} = -4.36 \times 1 + 76.2, r^2 = 0.88\), where \(\% \text{POS}\) is the percentage of positively buoyant cells and \(I\) is light intensity (µmol quanta m\(^{-2}\) s\(^{-1}\)). The percentage of \(R. \text{acuminata}\) cells positively buoyant under nutrient-replete conditions was fairly constant (between 45 and 58%; Fig. 2B) at light levels up to 110 µmol quanta m\(^{-2}\) s\(^{-1}\); however, it declined to less than 30% at the 2 highest light levels (155 and 212 µmol quanta m\(^{-2}\) s\(^{-1}\)). The average percentage positively buoyant at 155 and 212 µmol quanta m\(^{-2}\) s\(^{-1}\) was significantly lower (1-sided t-test, \(p < 0.05\)) than in the cultures at 17, 38, and 39 µmol quanta m\(^{-2}\) s\(^{-1}\). Note that buoyancy patterns were similar where blue and white light cultures overlapped. The best fit regression of buoyancy and light intensity in \(R. \text{acuminata}\) had the equation \(\% \text{POS} = -0.18 \times 1 + 61.7, r^2 = 0.84\). In \(R. \text{castacanei}\) cultures, the highest percentages of positively buoyant cells were also seen in light-limited cultures (71% at 11 µmol quanta m\(^{-2}\) s\(^{-1}\); Fig. 2C). The 2 high-light cultures had 38 and 33% positively buoyant at 164 and 210 µmol quanta m\(^{-2}\) s\(^{-1}\), respectively. The best fit equation of this relationship (Fig. 2C) had the equation \(\% \text{POS} = 0.15 \times 1 + 63.1, r^2 = 0.76\).

The percentage of positively buoyant \(R. \text{formosa}\) cells decreased over time in batch cultures (Fig. 3A). Two low-light cultures (15 and 19 µmol quanta m\(^{-2}\) s\(^{-1}\)) which did not reach stationary phase during the experiment had consistently high (47 to 83%) percentages of positively buoyant cells. In the cultures at 17, 23, 29, and 58 µmol quanta m\(^{-2}\) s\(^{-1}\), the percentage of positively buoyant cells decreased from 61-83% initially to <25% at the onset of stationary phase (cell densities between 12 000 and 15 000 cells l\(^{-1}\); Fig. 3A). Cultures grown at high light intensities (106, 156, and 211 µmol quanta m\(^{-2}\) s\(^{-1}\)) had consistently low percentages of positively buoyant cells (0 to 19%; Fig. 3A; data not shown for 156 and 211 µmol quanta m\(^{-2}\) s\(^{-1}\) light levels where no positively buoyant cells were observed). Buoyancy patterns over time in \(R. \text{acuminata}\) batch cultures also varied with light level (Fig. 3B). In the cultures at 17 and 39 µmol quanta m\(^{-2}\) s\(^{-1}\), the percentage positively buoyant remained above 50% throughout the experiment; these 2 low-light, slow-growing cultures did not reach stationary phase. In faster-growing cultures (>39 µmol quanta m\(^{-2}\) s\(^{-1}\)) the percentage of positively buoyant cells declined over time. High-light cultures had consistently lower percentages of cells positively buoyant. As cultures approached stationary phase, the percentage of positively buoyant cells decreased to 32% or less (Fig. 3B). The percentage of \(R. \text{castacanei}\) cells positively buoyant also tended to decrease over time in low-light cul-

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Fig. 2. Percentage of \(R. \text{formosa}\) cells positively buoyant in nutrient-replete, exponentially growing cultures at different light levels. All cultures had at least 10 d to photoacclimate before buoyancy measurement. Error bars indicate the standard deviation from 2 or 3 buoyancy measurements. \(R. \text{formosa}\) symbols are from a single measurement.

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was not observed in 2 earlier experiments when cell diameter was 176 µm and average cell volume was 1.18 × 10\(^{7}\) µm\(^{3}\). At this larger cell size a growth rate of 0.45 divisions d\(^{-1}\) was observed at 150 µmol quanta m\(^{-2}\) s\(^{-1}\), and a lower rate of 0.32 divisions d\(^{-1}\) was observed in a culture at 209 µmol quanta m\(^{-2}\) s\(^{-1}\). The growth rate of \(R. \text{acuminata}\) (cell diameter = 122 µm, cell volume = 6.83 × 10\(^{6}\) µm\(^{3}\)) increased steadily with increasing light level, to a maximum of 0.78 divisions d\(^{-1}\) at 155 µmol quanta m\(^{-2}\) s\(^{-1}\), and then declined to 0.2 divisions d\(^{-1}\) at 212 µmol quanta m\(^{-2}\) s\(^{-1}\) (Fig. 1B). The maximum \(R. \text{castacanei}\) growth rate observed was 0.38 divisions d\(^{-1}\) at 164 µmol quanta m\(^{-2}\) s\(^{-1}\) (Fig. 1C). No significant photoinhibition of growth was observed at the highest light level of 210 µmol quanta m\(^{-2}\) s\(^{-1}\).

All 3 species had an inverse relationship between positive buoyancy and light intensity under nutrient-replete conditions (Fig 2). This inverse relationship was most apparent in \(R. \text{formosa}\) (Fig. 2A), where more than 80% of the cells were positively buoyant at the 2 lowest light levels (15 and 17 µmol quanta m\(^{-2}\) s\(^{-1}\)). As light level increased, the percentage of positively buoyant cells decreased; no positively buoyant cells were observed at the 2 highest light levels (156 and 211 µmol quanta m\(^{-2}\) s\(^{-1}\)).
Note that under nutrient replete conditions (Fig. 4), positive buoyancy was observed in both high-light cultures at the largest cell size; at the intermediate cell size some positive buoyancy was observed at light levels up to 106 μmol quanta m⁻² s⁻¹ and at the smallest cell size no positive buoyancy was observed at light levels above 32 μmol quanta m⁻² s⁻¹. There was an inverse relationship between irradiance level and positive buoyancy at all 3 cell sizes (Fig. 4).

In the experiment where half of a *Rhizosolenia formosa* culture was moved to a lower light level (Day 1), the low-light culture had a sharp increase in positively buoyant cells after 5 d from 24% to 54%, which then declined to 29% as the culture approached stationary growth phase (Fig. 5). The high-light culture had consistently low percentages of positively buoyant cells (12 to 15%). It is unclear why growth was inhibited in the low-light culture between Days 3 and 5. Both cultures had maximum average cell sap nitrate concentrations on Day 5 (18 and 11 mM for the high- and low-light cultures, respectively). Nitrate concentrations in the medium of both cultures was always greater than 40 μM.

There was a strong diel pattern in the cell division of *Rhizosolenia formosa* (Fig. 6). The highest percentages (29 to 35%) of recently divided cells were observed early in the light period; the minimum (5%) was halfway through the dark period. On average, 89% of the recently divided cells were negatively buoyant. Positive buoyancy remained fairly constant (19 to
of nitrate uptake in the dark. The minimum average intracellular nitrate concentration was 3.3 mM; thus, intracellular nitrate concentrations can vary at least 5 fold during a diel cycle.

Most chemical composition values of Rhizosolenia acuminata and R. formosa were close to the values predicted by previous size-dependent studies of diatoms (Strathmann 1967, Hitchcock 1982; see Table 1). One notable exception was the higher-than-predicted particulate C values in R. formosa. These elevated C values are consistent with the observed high C:N ratios (9.0 to 16.4, mol/mol) and greater than predicted carbohydrate levels (Table 1).

Chlorophyll a values were lower than predicted, especially in R. acuminata. The highest average chlorophyll a value observed in R. acuminata was 0.70 ng cell⁻¹ (in a low-light culture), while Hitchcock’s (1982) size relationship would predict 2.04 ng cell⁻¹. The Rhizosolenia spp. in this study were within the size range considered in the 2 size-composition studies (Strathmann 1967, Hitchcock 1982). The maximum assimilation numbers calculated
for \textit{R. formosa} and \textit{R. acuminata} were 2.9 and 6.9 mg C mg\(^{-1}\) chlorophyll \(a\) h\(^{-1}\), respectively (see Table 3). There was a negative correlation between carbohydrate per cell and positive buoyancy when all values for \textit{R. acuminata} were examined (Fig. 7).

**DISCUSSION**

\textit{Rhizosolenia formosa}, \textit{R. acuminata} and \textit{R. castracanei} had similar buoyancy properties (Table 2). All 3 species exhibited an inverse relationship between light level and positive buoyancy under nutrient-replete conditions, and all had low percentages of positively buoyant cells in stationary phase cultures. \textit{R. formosa} and \textit{R. acuminata} also experienced growth inhibition at relatively low light intensities. Photoinhibition is typically exhibited at light intensities >500 \(\mu\)mol quanta m\(^{-2}\) s\(^{-1}\) (Platt et al. 1980, Neale & Richerson 1987). Light-induced cell mortality was observed in \textit{R. formosa} at light levels as low as 105 \(\mu\)mol quanta m\(^{-2}\) s\(^{-1}\). This cell mortality did not occur until cells had been exposed to these light levels for a considerable time period of 20 to 30 d. Initially (up to 10 d), cells appeared healthy and some cell division occurred. A similar photoinhibition-induced mortality has been reported for \textit{Pyrocystis noctiluca}. Kahn & Swift (1978) reported that nutrient-depleted cells died within a few days at 60 \(\mu\)mol quanta m\(^{-2}\) s\(^{-1}\), but not at lower light levels. In another experiment \textit{P. noctiluca} cells under both nutrient-replete and nutrient-deplete conditions died when suspended \textit{in situ} (depth of 10 m) at \(-312\) \(\mu\)mol quanta m\(^{-2}\) s\(^{-1}\) (Ballek & Swift 1986). Like \textit{P. noctiluca}, \textit{R. formosa} (and probably the others) became negatively buoyant during cell division (Swift et al. 1976, Kahn & Swift 1978, Rivkin et al. 1984). Based on these similarities, the once-in-a-generation migration pattern noted in \textit{P. noctiluca} appears to be a valid model for solitary \textit{Rhizosolenia} as well.

\textit{Rhizosolenia formosa} seems well adapted for growth at low light levels as evidenced by the steep slope of the light-limited portion of the growth-irradiance curve (Fig. 1A). The maximum growth rate for \textit{R. formosa} in Fig. 1 was 0.37 divisions d\(^{-1}\) at the very low light level of 29 \(\mu\)mol quanta m\(^{-2}\) s\(^{-1}\). This irradiance level is equivalent to \(-1\) % of surface irradiance levels in the Sargasso Sea (at noon during spring–summer) (Goldman 1993). Thus maximum growth rates were observed in environmental conditions similar to those found near the nutricline.

There were similarities observed between the chemical composition of \textit{Rhizosolenia acuminata} and \textit{R. formosa} in this study under nutrient-replete, light-saturated conditions and that of field populations of \textit{Rhizosolenia} mats, Ethmodiscus rex and \textit{Pyrocystis}.

**Table 2. Buoyancy and growth characteristics summary for \textit{Rhizosolenia formosa} (at 2 cell sizes), \textit{R. acuminata} and \textit{R. castracanei}**

<table>
<thead>
<tr>
<th></th>
<th>\textit{R. formosa}</th>
<th>\textit{R. acuminata}</th>
<th>\textit{R. castracanei}</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cell diameter ((\mu)m)</td>
<td>129</td>
<td>176</td>
<td>122</td>
</tr>
<tr>
<td>Average cell volume ((\times 10^6) (\mu)m(^3))</td>
<td>6.60</td>
<td>11.8</td>
<td>6.83</td>
</tr>
<tr>
<td>Max. growth rate (divisions d(^{-1}))</td>
<td>0.37</td>
<td>0.51</td>
<td>0.78</td>
</tr>
<tr>
<td>Light level of max. growth rate ((\mu)mol quanta m(^{-2}) s(^{-1}))</td>
<td>29</td>
<td>150</td>
<td>155</td>
</tr>
<tr>
<td>Lowest light level with photoinhibition of growth</td>
<td>105</td>
<td>200</td>
<td>210</td>
</tr>
<tr>
<td>% of cells positively buoyant:</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>(1) under nutrient replete, low-light conditions (&lt;40 (\mu)mol quanta m(^{-2}) s(^{-1}))</td>
<td>61–83 %</td>
<td>50–56 %</td>
<td>48–71 %</td>
</tr>
<tr>
<td>(2) under nutrient replete, high-light conditions (&gt;100 (\mu)mol quanta m(^{-2}) s(^{-1}))</td>
<td>0–14 %</td>
<td>21–40 %</td>
<td>22–51 %</td>
</tr>
<tr>
<td>(3) in stationary phase cultures</td>
<td>&lt;25 %</td>
<td>&lt;25 %</td>
<td>&lt;32 %</td>
</tr>
</tbody>
</table>
in situ (Table 3), including high C:N and C:chlorophyll ratios and similar cell sap nitrate concentrations and assimilation numbers. Villareal & Carpenter (1994) reported that E. rex cells contained only ~40% of the predicted amounts of chlorophyll a and 1.2 to 2.5 times as much C cell⁻¹ as predicted by the same size-composition equations as in Table 1 (Strathmann 1967, Hitchcock 1982). Typically C:chlorophyll ratios in cultures growing at high relative growth rates range from 25 to 50 (wt/wt) and C:N ratios from 5 to 7 (mol/mol) (Goldman 1980). The high C:N ratios seen in R. formosa in this study under nutrient-replete conditions strongly support the hypothesis that the high C:N ratios seen in field populations of the buoyant phytoplankton species listed in Table 3 do not indicate N limitation (Villareal & Carpenter 1989, 1994). Rivkin et al. (1984) suggest that Pyrocystis accumulates excess carbon at high (near-surface) light intensities which it catabolizes at depth during nutrient uptake, thus maintaining a relatively high growth rate throughout the water column. Rhizosolenia and Ethmodiscus spp. appear to have similar utilization patterns. The maximum assimilation number of R. acuminata was quite close to the assimilation numbers seen in Rhizosolenia mats in situ (Table 3), and suggests that the mats were growing at high relative growth rates. Cell sap nitrate concentrations were consistently an order of magnitude higher than medium nitrate concentrations in both Rhizosolenia acuminata and R. formosa. R. formosa had average concentrations as high as 17 to 18 mM and a maximum of 26 mM, seen in several samples. R. acuminata had lower average (2.2 to 6.4 mM) and maximum (8.4 mM) concentrations. These concentrations are at the lower end of the intracellular nitrate concentrations seen in other diatoms under N-replete conditions (up to 94 mM); however, the percent of particulate N accounted for by intracellular nitrate is near the upper end (5.4%) of the range observed in other diatoms (Dortch et al. 1984). In general, the cell sap nitrate concentrations observed in R. acuminata and R. formosa were comparable with the concentrations seen in field populations of Rhizosolenia mats and Ethmodiscus rex (Table 3). Individual Rhizosolenia chains collected in the Sargasso Sea had average intracellular nitrate concentrations of 4.2 mM and the positively buoyant chains had significantly higher internal nitrate concentrations than sinking chains (Villareal & Lipschultz 1995). This is further evidence that these field populations are not N-limited, because N-starved diatoms do not accumulate nitrate internally (Dortch 1982, Dortch et al. 1984).

Dark nitrate uptake and storage occurred in the Rhizosolenia formosa diel experiment (Fig. 6) where maximum cell sap nitrate concentrations (17.0 mM) were observed at the end of the dark period. Cullen (1985) calculated a rough stoichiometry whereby for each nitrate ion taken up and reduced nocturnally, about 6 CH₂O units were utilized. The observed average nocturnal uptake by R. formosa (0.15 nmol NO₃⁻ cell⁻¹) would utilize 0.02 nmol CH₂O or roughly 27.7 ng carbohydrate if fully reduced. Cells equal in size to those in the diel experiment contained up to 149 ng carbohydrate. This was enough carbohydrate to extend the uptake and full reduction of nitrate in the dark for up to 5.3 times the length of the dark period (149 ng/27.7 ng = 5.3) or approximately 2.7 d. Based on this conservative calculation, cells below the nutricline for more than a day might accumulate considerably higher internal nitrate concentrations. For example, assuming a constant uptake rate, a cell below the nutricline for 36 h would increase its intracellular nitrate concentration from 4.7 to 43.7 mM. Since all other sampling was done during the light period, cell sap nitrate concentrations in other experiments were likely not at their maximum values.

The inverse relationship between positive buoyancy and irradiance level in all 3 Rhizosolenia spp. under nutrient-replete conditions implies that physiological changes in cells at supersaturating light intensities result in negative buoyancy. Accumulation of excess carbohydrates at high light intensities seems a logical cause, especially since this type of carbohydrate ballasting has been widely documented for buoyant cyanobacteria (van Rijn & Shilo 1985, Ibelings et al. 1991, Oliver 1994).

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Table 3: Comparison of the chemical composition of Rhizosolenia acuminata and R. formosa in light-saturated, nutrient-replete cultures with field populations of Rhizosolenia mats, Ethmodiscus rex and Pyrocystis noctiluca

<table>
<thead>
<tr>
<th></th>
<th>C:N (mol/mol)</th>
<th>C:chl (wt/wt)</th>
<th>Assimilation number (mg C mg⁻¹ chl a h⁻¹)</th>
<th>Intracellular nitrate (mM)</th>
</tr>
</thead>
<tbody>
<tr>
<td>R. acuminata</td>
<td>6.2</td>
<td>130</td>
<td>6.9</td>
<td>2.3</td>
</tr>
<tr>
<td>R. formosa</td>
<td>9.1 - 16.4</td>
<td>261</td>
<td>2.9</td>
<td>3.3 - 26.5</td>
</tr>
<tr>
<td>R. mats</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Atlantic</td>
<td>9.7</td>
<td>590</td>
<td>6.6</td>
<td>-</td>
</tr>
<tr>
<td>Pacific</td>
<td>7.4</td>
<td>150</td>
<td>6.3</td>
<td>0.3 - 14.1</td>
</tr>
<tr>
<td>Pacific</td>
<td>5.8 - 13.1</td>
<td>-</td>
<td>6</td>
<td>-</td>
</tr>
<tr>
<td>E. rex</td>
<td>14.4</td>
<td>129</td>
<td>1.8 - 4.7</td>
<td>14.4 - 27</td>
</tr>
<tr>
<td>P. noctiluca</td>
<td>10</td>
<td>121 - 165</td>
<td>3.3</td>
<td>-</td>
</tr>
</tbody>
</table>

*This study, †Carpenter et al. (1977), ‡Villareal & Carpenter (1989), §Allredge & Silver (1982), †Villareal et al. (1993), †Villareal et al. (in press), ‡Villareal & Carpenter (1994), §Rivkin et al. (1994)
There was a negative correlation ($r^2 = 0.67$) between ng carbohydrate cell$^{-1}$ and positive buoyancy in $R.\, acuminata$ (Fig. 7). Theoretical calculations demonstrate the feasibility of the carbohydrate ballasting hypothesis for $Rhizosolenia$. Oliver (1994) described how the ‘excess’ density contributed by a cellular component such as carbohydrate can be calculated as the difference in density between its mass and the mass of the volume of water it displaces. Given a carbohydrate density of 1.6 g ml$^{-1}$ (Oliver 1994) and that the density of water is approximately 1.0 g ml$^{-1}$, the excess density contributed by carbohydrates is 1.6 – 1.0 = 0.6 g ml$^{-1}$.

Thus, the observed average carbohydrate increase (75.9 ng carbohydrate cell$^{-1}$) in $R.\, acuminata$ from the lowest to the highest light level would displace approximately $47.4 \times 10^{-9}$ ml of water ([75.9 $\times 10^{-9}$ g]/1.6 g ml$^{-1}$) = $47.4 \times 10^{-9}$ ml). Since the density of water is 1.0 g ml$^{-1}$, the weight of this volume of water would be 47.4 $\times 10^{-9}$ g, or 47.4 ng. Thus, the excess density contributed by the carbohydrate increase would be 75.9 – 47.4 = 28.5 ng cell$^{-1}$. Assuming a constant seawater density of 1.0235 g cm$^{-3}$ (a typical density in Sargasso Sea surface waters; Villareal 1988), and subtracting the average density difference between positively buoyant cells and seawater of 2.3 mg ml$^{-1}$ calculated for $R.\, acuminata$ (Moore 1994), the average cell density would be 1.0212 g cm$^{-3}$. Given an average cell volume of 6.245 $\times 10^{-6}$ µm$^3$, the average cell’s weight would be 6377.4 ng. Adding the 28.5 ng cell$^{-1}$ of excess density from carbohydrates to this cell weight would increase cell density to 1.0258 g cm$^{-3}$ (an increase of 4.6 mg ml$^{-1}$). Thus the average increase in carbohydrate cell$^{-1}$ between the lowest and highest light levels observed in $R.\, acuminata$ would be more than sufficient to cause the average positively buoyant cell to become negatively buoyant. This calculation assumes that other cellular constituents remain the same at the 2 light levels, but it does indicate that carbohydrate ballasting alone could account for the observed buoyancy patterns. Lowering carbohydrate content by the same amount would restore positive buoyancy. The increase in positive buoyancy seen in the $R.\, formosa$ culture moved from high to low light (Fig. 5) is consistent with this hypothesis. Further evidence for the carbohydrate ballasting hypothesis is seen in recent data from $Rhizosolenia$ mats in the central North Pacific where negatively buoyant mats had very high carbohydrate/protein ratios (average 2.2) relative to positively buoyant mats (average 0.7) (Villareal et al. in press).

Under steady-state conditions, the buoyancy properties summarized in Table 1, if replicated in situ, would lead to sub-surface population maxima in all 3 $Rhizosolenia$ spp. similar to what has been documented for $Pyrocystis\, noctiluca$ (Swift et al. 1976, Rivkin et al. 1984). Cells would not maintain a single preferred depth but would oscillate around a preferred depth or light level; on average, cells below the preferred depth would be positively buoyant and cells above that depth would be negatively buoyant. Examining Fig. 2A, it appears that nutrient-replete $R.\, formosa$ would likely be positively buoyant at light levels <20–40 µmol quanta m$^{-2}$ s$^{-1}$ and negatively buoyant at light levels >20–40 µmol quanta m$^{-2}$ s$^{-1}$. Thus, the apparent preferred depth was where light levels were approximately 20 to 40 µmol quanta m$^{-2}$ s$^{-1}$. Maximum growth rates were observed (Fig 1A) in this same light range. The relationship between light level and buoyancy (and inferred preferred depth) was also a function of cell size (Fig. 4). This oscillating depth regulation is consistent with vertical migration if the lower end of the oscillation brings cells below the nutrient-sufficiently frequently (see the model of Kromkamp & Walsby 1990, 1992), an event that clearly occurs based on field populations of solitary $Rhizosolenia$ (Villareal & Lipschultz 1995). The growth and buoyancy characteristics of all 3 species are consistent with being part of the ‘shade flora’, a flora which occurs preferentially or exclusively at some depth below the surface, usually around 100 m depth near the nutricline, and includes $R.\, acuminata$ and $R.\, castracanei$ (Sournia 1982, Venrick 1982). However, the steady-state assumptions of cultures may not be valid for open-ocean populations. These cultures were pre-adapted to a single light intensity for 10 d, an unlikely occurrence in the open sea, where cell buoyancy appears to respond to nutrient depletion on the order of a few days (Villareal & Lipschultz 1995). Nonetheless, these culture studies indicate that even under conservative steady-state conditions, $R.\, acuminata$, $R.\, castracanei$ and $R.\, formosa$ are capable of positive buoyancy and under laboratory conditions exhibit buoyancy and growth characteristics consistent with vertical migration. $R.\, acuminata$ and $R.\, formosa$ had chemical compositions similar to those observed in vertically migrating populations of $Rhizosolenia$ mats, $Pyrocystis\, noctiluca$ and $Ethmodiscus\, rex$ in oligotrophic seas (Rivkin et al. 1984, Villareal et al. 1993, Villareal & Carpenter 1994). In addition, all 3 species in this study are capable of ascent rates on the order of cm to m h$^{-1}$ (Moore 1994) which permit the euphotic zone to be traversed in as little as 24 h. Thus, it seems likely that short chains of $R.\, formosa$, $R.\, castracanei$ and $R.\, acuminata$ are capable of a buoyancy-mediated vertical migration and, thus, are a source of new nitrogen to the euphotic zone. Unfortunately, quantitative abundance data for these species are not available. While $Rhizosolenia$ mats are found mainly in the Pacific, single-species chains of large $Rhizosolenia$ are commonly found in all warm water seas, particularly in the Sargasso Sea (Sundström 1986,
Villareal 1988). Thus, vertical migration by single-species chains of Rhizosolenia may be a significant source of new nitrogen on a global scale.

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