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Combined Effects of CO2 and Light on Large and Small Isolates of the Unicellular N2-Fixing Cyanobacterium Crocosphaera watsonii From the Western Tropical Atlantic Ocean

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Combined effects of CO₂ and light on large and small isolates of the unicellular N₂-fixing cyanobacterium *Crocosphaera watsonii* from the western tropical Atlantic Ocean

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We examined the combined effects of light and pCO₂ on growth, CO₂-fixation and N₂-fixation rates by strains of the unicellular marine N₂-fixing cyanobacterium *Crocosphaera watsonii* with small (WH0401) and large (WH0402) cells that were isolated from the western tropical Atlantic Ocean. In low-pCO₂-acclimated cultures (190 ppm) of WH0401, growth, CO₂-fixation and N₂-fixation rates were significantly lower than those in cultures acclimated to higher (present-day ~385 ppm, or future ~750 ppm) pCO₂ treatments. Growth rates were not significantly different, however, in low-pCO₂-acclimated cultures of WH0402 in comparison with higher pCO₂ treatments. Unlike previous reports for *C. watsonii* (strain WH8501), N₂-fixation rates did not increase further in cultures of WH0401 or WH0402 when acclimated to 750 ppm relative to those maintained at present-day pCO₂. Both light and pCO₂ had a significant negative effect on gross : net N₂-fixation rates in WH0402 and trends were similar in WH0401, implying that retention of fixed N was enhanced under elevated light and pCO₂. These data, along with previously reported results, suggest that *C. watsonii* may have wide-ranging, strain-specific responses to changing light and pCO₂, emphasizing the need for examining the effects of global change on a range of isolates within this biogeochemically important genus. In general, however, our data suggest that cellular N retention and CO₂-fixation rates of *C. watsonii* may be positively affected by elevated light and pCO₂ within the next 100 years, potentially increasing trophic transfer efficiency of C and N and thereby facilitating uptake of atmospheric carbon by the marine biota.

**Key words:** carbon dioxide, *Crocosphaera*, cyanobacteria, diazotroph, light, nitrogen fixation, ocean global change, unicellular

**Introduction**

Within the past two decades, emerging data have suggested that the magnitude of marine N₂ fixation has been grossly underestimated (Deutsch et al., 2007; Capone, 2008; Mulholland et al., 2012). Traditionally, *Trichodesmium* has been widely accepted to be a major contributor to oceanic N₂ fixation; however, estimates of N₂ fixation by unicellular diazotrophs continue to increase and their calculated N inputs to marine systems may narrow the gap in the global N budget (Zehr et al., 2001; Montoya et al., 2004; Church et al., 2008; Moisander et al., 2010). Understanding how these key components of the marine N cycle will respond to rapid global change is essential to predict how the carbon cycle will change. In the next 100 years, anthropogenic inputs of CO₂ to the atmosphere will likely double the present-day partial pressure of CO₂ (pCO₂). At the same time, the average global mixed layer depth is also expected to decrease, thereby contributing to higher mean light intensity experienced by phytoplankton (Sarmiento et al., 2004; Behrenfeld et al., 2006; Boyd et al., 2010). In addition to their individual effects, we now recognize that interactive effects of these and other environmental factors must be considered to realistically predict the net impacts of global change (Hutchins et al., 2007, 2009; Fu et al., 2008; Kranz et al., 2010; Levitan et al., 2010; Garcia et al., 2011). *Crocosphaera watsonii* has been widely studied in the literature in an effort to understand the physiology of unicellular photosynthetic N₂ fixers and biogeochemical implications for models of oceanic biological N₂ fixation (Zehr et al., 2001, 2007; Goebel et al., 2007, 2008). In this study, we examined how two isolates of this genus might respond to global change. Previous studies suggest that elevated pCO₂ acts to enhance gross N₂-fixation rates by the oceanic diazotrophs *Trichodesmium erythraeum* and *C. watsonii* (Barcelos e Ramos et al., 2007; Hutchins et al.,...
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CO₂ experiments

Recently, however, several experiments have indicated that light influences the effect of elevated pCO₂ on gross N₂ fixation by T. erythropus (Kranz et al., 2010; Garcia et al., 2011). Elevated pCO₂ acts to enhance N₂-fixation rates under low light but this stimulatory effect is lower at high light, which may be caused by an enhanced ability to retain newly fixed cellular N at high light (Garcia et al., 2011).

Our goal in this study was to understand how light influences the effect of elevated CO₂ on growth, CO₂-fixation and N₂-fixation rates of two isolates of C. watsonii from the western tropical Atlantic Ocean in laboratory culture experiments. To date, most published works investigating the physiological responses of C. watsonii have focused on one strain (WH8501) and physiological studies of other strains are currently lacking. Recently, Webb et al. (2009) compared N₂-fixation rates by strains of C. watsonii differing in cell size. Chl a-normalized N₂-fixation rates in a large strain isolated from the North Pacific Ocean (WH0003) were twice as high as those in a smaller strain isolated from the South Atlantic Ocean (WH8501). To consider differences in responses between strains of C. watsonii, we examined one isolate with small cells (2–3 µm diameter; WH0401) and another isolate with large cells (4–6 µm diameter; WH0402).

Materials and methods

Culturing and experimental design

Stock cultures of the two Atlantic C. watsonii isolates used in this study were provided courtesy of Dr. Eric Webb. Both isolates were collected in March 2002, WH0401 from 6°58.78′N, 49°19.70′W and WH0402 from 11°42.12′S, 32° 00.64′W. An outline of all experiments with both isolates, including experimental analyses used for each, is presented in Table 1. In all experiments, triplicate cultures were grown using a semi-continuous culturing technique (Garcia et al., 2011) at 28°C in an artificial seawater medium (Chen et al., 1996). Nutrients were added to autoclaved seawater at the concentrations listed in the AQUIL recipe (Morel et al., 1979), except for nitrate, which was omitted. The growth rates of cultures were measured over 2–3 day intervals and were used to determine the dilution rate. Culture cell density was kept low (cells ml⁻¹ = 50–500 × 10⁶) for experiments with WH0401 and 5.0–30 × 10⁶ for WH0402; Table 1) to prevent light limitation of photosynthesis and deviation from the expected pH values for respective pCO₂ culture treatments. Light was supplied with cool-white fluorescent lamps on a 12 : 12 h light : dark cycle and measured with a LI-250A light meter (LiCor Biosciences, light sensor serial# SPQA 4020).

Because of large differences in cell size between WH0401 and WH0402, we cultured WH0401 at higher cell densities to maintain relatively equivalent levels of total culture biomass (0.1–2.5 mM particulate C for cultures of WH0401; 0.1–1.3 mM particulate C for WH0402). For CO₂ experiments, media and cultures were bubbled with filtered air from the room (0.2 µm filtered, present-day pCO₂ concentration of ~385 ppm) or premixed air prepared by Gilmore Liquid Air Company with certified values of 190 ppm pCO₂ (last glacial maximum levels: Petit et al., 1999) and 750 or 761 ppm pCO₂ (within the range predicted for the year 2100: Alley et al., 2007) for the entire term of the experiment (Table 1). Cells were considered fully acclimated to treatment conditions after cultures had remained at steady-state growth for seven generations or more (unless stated otherwise). Fast-growing cultures (i.e. high light cultures) were acclimated for more than 10 generations while slow growing cultures (i.e. low light and low pCO₂ cultures) were acclimatized over 2 months but for fewer generations. Cultures were sampled over the period between 24 and 48 h after the preceding dilution to measure growth rates, gross and net N₂-fixation rates, CO₂-fixation rates, particulate elemental composition, and carbonate system measurements (for CO₂ experiments).

Light experiments

In order to quantify differences in growth and in the CO₂- and N₂-fixation rate capacities of these two isolates of C. watsonii, we measured growth, CO₂-fixation and gross and net N₂-fixation rates, and particulate carbon and nitrogen composition in response to a range of light intensities (labelled experiments 1 and 2 in Table 1).
CO₂ experiments

To investigate variability in the effects of CO₂ on growth and N₂-fixation rates between strains of C. watsonii, we conducted experiments with cultures of WH0401 and WH0402. We measured growth and gross and net N₂-fixation rates (see N₂-fixation rates) in response to three levels of CO₂ (190, air and 750 ppm) at a light intensity of 155 µmol quanta m⁻² s⁻¹ (labelled experiments 3 and 4 in Table 1). We chose this light intensity because we did not want growth rates in these cultures to be limited by light.

CO₂-light experiments

To determine if light influences the effect of elevated pCO₂ on growth, CO₂-fixation and N₂-fixation rates of C. watsonii, we first grew WH0402 with two concentrations of CO₂ (air and 750 ppm) at five light intensities (18–300 µmol quanta m⁻² s⁻¹; labelled experiment 6 in Table 1). In this experiment we measured similar growth and N₂-fixation rates at the two CO₂ concentrations. Therefore, when examining responses of WH0401 with this experimental design, we added a low CO₂ treatment (190 ppm) under the same range of light intensities (labelled experiment 5 in Table 1). Despite several attempts, we were not able to acclimate WH0401 to any of these CO₂ concentrations at 18 or 50 µmol quanta m⁻² s⁻¹ for unknown reasons.

Growth rate and cell density estimates

Growth rate was determined as an increase in culture cell density over time with the equation NT = N₀eµT, where N₀ and N_T are the initial and final culture cell densities, respectively, T is the time in days between culture cell density estimates, and µ is the specific growth rate. Culture cell density was determined using a haemocytometer and an Olympus BX51 microscope. Cell diameter was measured using an ocular micrometer calibrated with the same microscope. Growth rates were fitted to a Monod linear hyperbolic function of light (Monod, 1949) using Sigma Plot 10 software program. The hyperbola was fit to the data without including the origin to yield the highest r² value. With Sigma Plot 10, we did this by calculating the Kₘ and maximum rate values after aligning the data to include the origin. The point of alignment was determined by achieving the highest r² value. We then realigned the data to their original values along with the best-fit functions. This method yields more realistic Monod parameters with critical threshold values.

Carbonate system measurements

Culture pH was measured intermittently during the CO₂ experiments with a pH meter using the NBS seawater scale (Orion 5 star Thermo Scientific, Beverly, MA, USA). We preserved samples for total CO₂ (TCO₂) measurements in unfiltered water collected from cultures (5–70 ml; stored at 4°C) with a 5% HgCl₂ solution (0.5% final concentration) until later analysis with a carbon coulomb meter (CM 140, UIC, Joliet, IL, USA). We measured TCO₂ by acidifying a 5 ml sample with phosphoric acid (1–2% final concentration) and quantifying the CO₂ trapped in an acid sparging column as described in Garcia et al. (2011). TCO₂ analyses were not available in our preliminary CO₂ experiments. We calculated pCO₂ with the CO₂sys program (Lewis & Wallace, 1998) using the NBS pH scale and K₁ and K₂ constants from Mehrbach et al. (1973), refit by Dickson & Millero (1987).

N₂ fixation

For all experiments we used the acetylene reduction assay described by Capone (1993) to estimate the gross N₂-fixation rate. All rate measurements in the light and CO₂-light experiments were initiated at the beginning of the 12-h dark period, when C. watsonii is known to fix N₂ (Mohr et al., 2010a; Saito et al., 2011). For the CO₂ experiments the acetylene assay was initiated during the seventh hour of the 12-h dark period and continued for 4 h. For this assay, two 50 ml (light and CO₂-light experiments) or 60 ml (CO₂ experiments) culture samples were collected from each replicate and incubated in 80-ml polycarbonate bottles at 28°C. Four millilitres of acetylene were injected into the headspace ~1 h after the beginning of the dark period and samples were withdrawn from the headspace every 2–3 h to measure acetylene reduction. In the CO₂-light experiment with WH0401, we measured rates throughout the dark period and continued to measure them during the early portion of the light cycle, when samples were exposed to treatment light levels (Table 1). In this experiment, we gently agitated incubation bottles to equilibrate ethylene in the seawater with ethylene in the headspace. Gross N₂-fixation rates were calculated in the same way as described in Garcia et al. (2011), using a Bunsen coefficient for ethylene of 0.082 (Breithbarth et al., 2004) and an ethylene production : N₂-fixation ratio of 3 : 1.

We also measured net N₂-fixation rates using the ¹⁵N₂ isotope tracer method (Mulholland et al., 2004; Mulholland & Bernhardt, 2005). Samples were prepared the same way as described in Garcia et al. (2011). Briefly, 169 ml of each experimental replicate was inoculated with 169 µl of 99% doubly labelled ¹⁵N₂ gas and incubated at 28°C in complete darkness for 12 h during the dark period. The incubation was then terminated by filtering the entire volume onto precombusted (450°C, 4 h) GF/F filters for the analysis of particulate ¹⁵N, total particulate N and total particulate C. Filters were dried at 80–90°C, pelleted, and combusted in a quartz column with chromium oxide and silver wool at 1000°C. For this analysis we used ammonium sulphate and sucrose as standards. At the time we conducted these experiments, we were not aware of the criticisms of the ¹⁵N₂ uptake method that have been discussed by Mohr et al. (2010b). Thus, for another independent estimate of net N₂ fixation, we calculated a particulate N (PN) accumulation rate in cultures over time (ΔPN = PNfinal – PNinitial) by using our estimates of particulate N. Particulate N was measured in subsamples of experimental replicates that were incubated with ¹⁵N₂ at the end of the dark period and used as the end-period PN measurement (PNfinal). Because only one sample of PN was collected, we back-calculated an estimate of PN_initial based on our measurements of cellular growth rate using the equation: growth rate (d⁻¹) = [ln(PNfinal)−ln(PN_initial)]/(t₂−t₁), where t₁ is the initial time and t₂ is the final time. Based on our measurements of growth rates, we assumed that PN cell⁻¹ was in a daily steady state. We then calculated the gross N₂-fixation rate : PN-accumulation rate ratio (hereafter the gross : PN accumulation ratio) and compared it with the ratio of gross N₂-fixation rate : net ¹⁵N₂-fixation rate ratio.
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CO2 fixation

The rate of CO2 fixation was determined as described in Garcia et al. (2011) using the H14CO3 incorporation method. CO2-fixation rates were determined by first calculating the ratio of the radioactivity of 14C incorporated into cells during 24 h to the total radioactivity of H14CO3. This ratio was then multiplied by the total CO2 concentration (TCO2). TCO2 concentrations were measured in our CO2-light experiments and were applied to all experiments to calculate CO2-fixation rates for corresponding CO2 treatments and were applied to all experiments to calculate CO2-fixation rates for corresponding CO2 treatments. For the light experiments, we used a TCO2 value that was measured in the present-day pCO2 treatments of the CO2-light experiments (2053 μM TCO2).

Particulate C and N

Culture samples from each experimental replicate (100 ml) were filtered onto precombusted (450°C, 4 h) GF/F filters for the analysis of cellular N and C. Filters were then dried at 80–90°C and compressed into pellets, and the amounts of C and N were determined using an elemental analyser (Costech Instruments, model 4010).

Statistics

We used a one-way analysis of variance (ANOVA) test (with the light experiment and CO2 experiment data) and a two-way ANOVA test (with the CO2–light experiment data) combined with a Tukey analysis of multiple comparisons to determine statistical differences (P < 0.05) between treatments. For these analyses, we used data from all three replicates from each treatment.

Results

Light experiments (experiments 1 and 2)

Mean specific growth rates of WH0402 were higher than those of WH0401 at all light levels investigated (P < 0.05; Fig. 1a). Cells of WH0401 were considerably smaller than cells of WH0402 and average cell diameters were ~20% larger in high-light acclimated cells compared to low-light acclimated cells in both strains (P < 0.05) (Fig. 1b). The Monod fit of growth as a function of light yielded a theoretical maximum growth rate of 0.95 d⁻¹ (r² = 0.99) for WH0402 and 0.68 d⁻¹ (r² = 0.99) for WH0401. However, the half-saturation constant for light (Kₘ) with respect to light and the light compensation point for growth (Eᵥ, where net growth is zero) were similar between strains (WH0401, Kₘ = 61 μmol quanta m⁻² s⁻¹, Eᵥ = 11 μmol quanta m⁻² s⁻¹; WH0402, Kₘ = 59 μmol quanta m⁻² s⁻¹, Eᵥ = 13 μmol quanta m⁻² s⁻¹). Because of the large differences in cell size between strains, we compared C-specific CO2-fixation rates and N-specific N2-fixation rates. We determined these rates by normalizing N2-fixation rates to particulate organic nitrogen measurements and CO2 fixation rates to particulate organic carbon measurements. Both C-specific CO2-fixation rates (Fig. 1c) and N-specific gross N2-fixation rates (Fig. 1d) were consistently higher in the strain with large cells (WH0402) than in the strain with small cells (WH0401), except at the lowest light level, similar to the pattern of their specific growth rates. Mean growth rates were highly correlated with mean N-specific 15N2-fixation rates (r = 0.85, n = 5 for WH0401; r = 0.99, n = 5 for WH0402) (Fig. 1a, e). Mean gross : net N2-fixation rate ratios declined with increasing light intensity by 72% in WH0401 (from 300 to 50 μmol quanta m⁻² s⁻¹) and 82% in WH0402 (from 300 to 25 μmol quanta m⁻² s⁻¹) and were negatively correlated with mean specific growth rates (WH0401, r = −0.91, n = 4; WH0402, r = −0.92, n = 5) and mean cell volumes (WH0401, r = −0.89, n = 4; WH0402, r = −0.71, n = 5; Fig. 1f).

CO2 experiments (experiments 3 and 4)

For unknown reasons, the growth rates of WH0401 were lower in the CO2 experiment than in the CO2–light experiment with the same CO2 concentrations at relatively equivalent light intensities. Measured pH values in bubbled cultures of the CO2 experiments were comparable to bubbled cultures in the CO2–light experiments (Table 2). The partial pressure of CO2 did not have a significant effect on growth rates in the isolate with large cells (WH0402; P > 0.05; Fig. 2). In contrast, growth rates in the small-celled strain (WH0401) were significantly lower at 190 ppm pCO2 than those at higher pCO2 concentrations (P < 0.002; Fig. 2) but were not significantly different between the present-day and elevated pCO2 treatments (P > 0.05). Mean gross : net N2-fixation rate ratios decreased with increasing pCO2 by 42% for WH0401 (r = 0.82, r = 0.9, n = 5) and 27% for WH0402 (r = 0.58, r = 0.89, n = 4; WH0402, r = −0.71, n = 5; Fig. 2).

CO2–light experiments (experiments 5 and 6)

Measured TCO2 concentrations and pH values in our cultures were within the expected range for the respective pCO2 treatments (Table 2). The specific growth rates of WH0401 were significantly lower in the low pCO2 treatment than in higher pCO2 treatments (F₁,₁₈ > 55, P < 0.001; Fig. 3a) but were not significantly different between the present-day and elevated pCO2 treatments (F₁,₁₈ = 0.92, P = 0.66). The growth rates of WH0402 were not significantly different between the present-day and elevated pCO2 treatments at all light intensities that we investigated (F₂,₁₈ = 3.2, P = 0.09; Fig. 3b). Cell-normalized CO2-fixation rates were positively affected by pCO2 in WH0401 (F₂,₁₈ = 4.7, P = 0.02), but the interactive
effect between light and pCO2 was not significant ($F_{4,18} = 0.13, P = 0.97$; Fig. 3c). Light and pCO2, however, did have a significant positive interactive effect on cellular CO2-fixation rates in cultures of WH0402 ($F_{1,20} = 13, P = 0.002$; Fig. 3d), indicating that the effect of elevated pCO2 significantly increased with increasing light.

Gross cellular N2-fixation rates of WH0401 were not affected by light between 100–300 μE m$^{-2}$ s$^{-1}$ treatments ($F_{2,18} = 3.0, P = 0.1$), or by pCO2 between the present-day and elevated pCO2 treatments ($F_{1,18} = 0.22, P = 0.65$), but were significantly lower in the 190 ppm treatment compared to higher pCO2 treatments ($F_{1,18} \geq 7.8, P \leq 0.01$; Fig. 4a). Similarly,
for WH0402, gross cellular N₂-fixation rates were not significantly different between the present-day and elevated pCO₂ treatments ($F_{1,20} = 3.1, P = 0.09$; Fig. 4b), but significantly increased as a function of increasing light between all light treatments ($F_{1,20} > 7.2, P < 0.02$).

Trends in cell-normalized net $^{15}$N₂-fixation rates by WH0401 were similar to those observed for growth rates: low pCO₂ had a significant negative effect on net $^{15}$N₂-fixation rates in comparison with higher pCO₂ levels ($F_{1,18} ≥ 8.2, P ≤ 0.01$) and rates did not differ between the air and elevated pCO₂ concentrations ($F_{1,18} = 0.2, P = 0.67$; Fig. 4c). In WH0402, cell-normalized net $^{15}$N₂-fixation rates were not significantly different between the air and elevated pCO₂ treatments ($F_{1,20} = 0.08, P = 0.77$; Fig. 4d), but were strongly affected by light ($F_{4,20} = 64, P < 0.0001$). PN-accumulation rates by WH0401 were lower than gross N₂-fixation rates but considerably higher than net $^{15}$N₂-fixation rates (Fig. 4e). In WH0402, PN-accumulation rates were similar to gross N₂-fixation rates and higher than net $^{15}$N₂-fixation rates (Fig. 4f; see the Materials and methods section for methodological differences in the acetylene assay between experiments). Both light and pCO₂ had significant positive effects on gross N-specific N₂-fixation rates by WH0401 ($F_{2,18} = 26, P < 0.001$ $F_{2,18} = 8.0, P = 0.003$) and differences in gross N-specific N₂ fixation between the 190 ppm pCO₂ treatment and higher pCO₂ treatments were more pronounced compared to gross cell-normalized N₂-fixation rates (Fig. 4g). In WH0402, gross N-specific N₂-fixation rates were not significantly different between pCO₂ treatments and were light saturated near 100 μmol quanta m⁻² s⁻¹ ($P > 0.05$, Fig. 4h). In both strains, trends in...
N-specific $^{15}$N$_2$-fixation rates (Fig. 4i, j) were very similar to trends in growth rates (Fig. 3a, b).

Both light ($F_{2,18} > 20.4, P < 0.0001$) and pCO$_2$ ($F_{2,18} = 5.4, P = 0.01$) had a significant negative effect on the ratio of gross : net $N_2$ fixation in WH0401 but the interactive effect of light and pCO$_2$ on the ratio was not significant ($P > 0.05$; Fig. 5a). In WH0402, light and pCO$_2$ did have a significant interactive effect on the ratio of gross : net $N_2$ fixation; the ratio decreased with increasing light by 53% in the air treatment but by only 37% in the 750 ppm CO$_2$ treatment (from 300 to 18 μmol quanta m$^{-2}$ s$^{-1}$). Thus, the effect of elevated pCO$_2$ on gross : net $N_2$ fixation significantly increased with decreasing light ($F_{4,20} = 3.9, P = 0.02$; Fig. 5b), suggesting that the effect of elevated pCO$_2$ on cellular N retention was strongest under low light. Growth rates of WH0402 were strongly negatively correlated with the gross : net $N_2$-fixation rate ratio ($r = -0.95$). Light was the most important factor controlling the gross : PN accumulation.

Fig. 4. Cellular gross $N_2$-fixation rates (a, b), cellular net $N_2$-fixation rates (c, d), calculated cellular particulate nitrogen (PN) accumulation rates (e, f), N-specific gross $N_2$-fixation rates (g, h), and N-specific net $N_2$-fixation rates (i, j) in semi-continuous cultures of *Crocosphaera watsonii*, isolates WH0401 and WH0402, as a function of pCO$_2$ and light. WH0401 and WH0402 were grown under present-day and elevated pCO$_2$ levels and WH0401 was also grown under 190 ppm pCO$_2$. Data correspond to experiments 5 and 6 in Table 1. Open symbols are 750 or 761 ppm pCO$_2$ treatments; grey symbols are air treatments; closed symbols are 190 ppm pCO$_2$ treatments. Error bars are the standard errors of the means of three experimental replicates.
accumulation ratio in WH0401 \( (F_{2,18} = 3.5, P = 0.05; \text{Fig. } 5c) \), which declined with increasing light. While pCO2 had no effect on the gross : PN accumulation ratio in WH0402 \( (F_{1,20} = 0.17, P = 0.69) \), the two-way ANOVA test suggested that light had a significant negative effect on this ratio \( (F_{4,20} = 5.6, P = 0.003; \text{Fig. } 5d) \), although this was driven mostly by the large increase in the ratio at 50 \( \mu \text{mol quanta m}^{-2} \text{s}^{-1} \). In both strains, the range of the gross : PN accumulation ratio was substantially lower than the range of the gross : net N2-fixation rate ratio (Fig. 5).

**Discussion**

Our results identified both similarities and differences in the physiological responses to changing pCO2 and light between large-celled and small-celled strains of *Crocosphaera watsonii* isolated from the western equatorial region of the Atlantic Ocean. In our light experiments, maximum growth responses \( (\mu_{\text{max}}) \) differed between strains but other Monod functional growth parameters \( (K_\mu \text{ and } E_c) \) were similar. Overall, our data indicate that the strain with large cells (WH0402) had higher growth, N2-fixation, and CO2-fixation rates at near-saturating light, compared to the strain with small cells (WH0401), despite having similar photosynthetic efficiencies at high light (data not shown). These high growth and fixation rates may give WH0402 an ecological advantage in regions of the ocean where nutrient concentrations are relatively high, whereas the smaller-celled WH0401 strain, with a higher cell surface area : volume ratio, may be better able to survive in lower-nutrient oceanic waters, because it may be able to acquire nutrients more readily when concentrations are low. In both strains, however, the diameters of cells acclimated to high light were ~20% greater than those acclimated to low light, suggesting that light controls a range of nutrient acquisition rates based on highly plastic cell surface area : volume ratios, as well as cellular quotas of elements.

Our findings do not support previous studies that documented increased growth rates of *C. watsonii* in response to elevated CO2 concentrations when compared to present-day CO2 concentrations (in South Atlantic strain WH8501 from 28°S, 48°W: Fu et al., 2008); in both of our strains, mean specific growth rates did not differ significantly between the present-day and elevated pCO2 treatments under any of the light levels that we tested. The growth rates of WH0401, however, were significantly lower under low pCO2 (190 ppm) at all light levels (100–300 \( \mu \text{mol quanta m}^{-2} \text{s}^{-1} \)) than in treatments with higher pCO2, whereas the growth rates of WH0402 were only slightly lower at low pCO2 and near-saturating light (155 \( \mu \text{mol quanta m}^{-2} \text{s}^{-1} \)) when compared to higher pCO2 treatments. These data suggest that WH0402 has a low \( K_\mu \) for growth with respect to pCO2 compared to WH0401, and also that the present-day concentration of pCO2 is near growth-saturating levels for both strains. We do not know why growth rates of WH0401 were lower in the CO2 experiment (experiment 3) than in the CO2–light experiment (experiment 5) but the CO2 experiments with WH0401 and WH0402 were done in parallel and

![Fig. 5. Gross : net N2-fixation rate ratios (a, b) and gross : net PN accumulation ratios (c, d) in semi-continuous cultures of *Crocosphaera watsonii*, isolates WH0401 and WH0402, as a function of pCO2 and light. WH0401 and WH0402 were grown under present-day and elevated pCO2 levels and WH0401 was also grown under 190 ppm pCO2. Data correspond to experiments 5 and 6 in Table 1. Open symbols are 750 or 761 ppm pCO2 treatments; grey symbols are air treatments; closed symbols are 190 ppm pCO2 treatments. Error bars are the standard errors of the means of three experimental replicates.](image-url)
provide comparative data between these strains. Collectively, these data support the notion that there may be strong differences between strains of *Crocosphaera* in terms of their ability to sequester inorganic carbon.

**Mechanistic effect of elevated CO\textsubscript{2} on N\textsubscript{2}-fixation rates**

Several authors have suggested that N\textsubscript{2}-fixation rates in *Trichodesmium* benefit from elevated pCO\textsubscript{2} by higher rates of diffusion of CO\textsubscript{2} across the cell membrane (Barcelos e Ramos *et al.*, 2007; Hutchins *et al.*, 2007; Levitan *et al.*, 2007, Kranz *et al.*, 2009, 2010; Garcia *et al.*, 2011). These higher rates then decrease the energy demand associated with active transport of bicarbonate (HCO\textsubscript{3}\textsuperscript{–}), the main source of inorganic carbon that fuels CO\textsubscript{2} fixation in both *Trichodesmium* and *Crocosphaera* (Giordano *et al.*, 2005; Badger *et al.*, 2006; Price *et al.*, 2008). This ‘saved’ energy might then be used to support high CO\textsubscript{2}-fixation and N\textsubscript{2}-fixation rates. This model must be modified for dark N\textsubscript{2} fixers like *Crocosphaera*, however, because cellular inorganic carbon uptake is driven by light (Badger *et al.*, 2006). Thus, the indirect effect of elevated pCO\textsubscript{2} on N\textsubscript{2} fixation by *Crocosphaera* seems to be to allow the cell to accumulate larger photosynthetic reserves during the preceding light period and the energy acquired from respiration of those reserves can then be used to drive N\textsubscript{2} fixation during the dark hours.

Studies of eukaryotic phytoplankton have supported this model for cellular energetic benefits from elevated pCO\textsubscript{2}, suggesting that a doubled pCO\textsubscript{2} could lead to a saving of ~20% of the energy consumed by the CCM, or up to 6% of the total cellular energy budget (Hopkinson *et al.*, 2011). Further studies are needed to confirm this in *Trichodesmium* and *Crocosphaera*, however, because some data suggest that inorganic C cycling in other cyanobacteria is related to light energy dissipation rather than CO\textsubscript{2} saturation of Rubisco and that cells may constitute a source of CO\textsubscript{2} rather than a sink (Tchernov *et al.*, 1998, 2003).

**Potential effects of cell size**

Differing abilities to acquire inorganic carbon could be related to cell size. Larger cells have a lower surface area : volume ratio than smaller cells, and so have lower volume-normalized CO\textsubscript{2} diffusion rates into the cell. However, in our study, the strain with larger cells (WH0402) seemed to have a lower K\textsubscript{m} with respect to CO\textsubscript{2}. Assuming that differences in K\textsubscript{m} are proportional to K\textsubscript{15} for CO\textsubscript{2} between strains, it is apparent that simple diffusion-based surface area : volume relationships cannot explain our results. New physicochemical modelling (Flynn *et al.*, 2012) suggests that as phytoplankton cell size increases, pH changes in the bulk medium have less physiological effect because the chemistry of the cell’s diffusive boundary layer is progressively more influenced by cellular metabolic processes. Differences in K\textsubscript{m} with respect to CO\textsubscript{2} might also be caused by differences in the efficiency of transmembrane HCO\textsubscript{3}–-transport systems, but such attribution of cause and effect must await further studies with multiple *C. watsonii* isolates. Although Price *et al.* (2008) identified different mechanisms by which strain WH8501 acquires carbon, there is no literature describing differences in these mechanisms between this and other isolates of *C. watsonii*.

It is also possible that a higher CO\textsubscript{2} diffusion rate for larger cells might be facilitated by the existence of acidic zones around the cell, as suggested by Raven *et al.* (2008). Future studies should investigate this possibility for large-celled strains of *C. watsonii*, given the much larger amounts of potentially acidic extracellular polysaccharide exudates associated with large-celled strains, compared to small-celled strains (Webb *et al.*, 2009; Sohn *et al.*, 2011). In addition, strain-specific differences might be caused by biogeochemical differences between their sites of origin; for instance WH0401 was collected near the Amazon River plume, and WH0402 is likely not adapted to this type of terrestrial influence environment. Further studies should address the global diversity of N\textsubscript{2}-fixing cyanobacteria such as *Crocosphaera* in relation to changing pCO\textsubscript{2}, as our work makes it clear that even closely related strains of a diazotrophic species may have very different responses to the environmental changes that will occur in a future acidifying and warming ocean.

**Gross : net N\textsubscript{2} fixation ratios**

Mean gross : net N\textsubscript{2}-fixation rate ratios were negatively correlated with mean growth rates, suggesting that high light and high pCO\textsubscript{2} enhanced the incorporation of fixed N\textsubscript{2} into biomass when growth rates were maximal. Thus, based on previous studies of cellular N retention (Mulholland *et al.*, 2004; Mulholland, 2007), we infer that the loss of fixed N might be minimized in a high light, high pCO\textsubscript{2} environment. Assuming that *C. watsonii* will be grazed upon to a significant degree in the future, we might expect nitrogen to flow more efficiently through food webs within the next 100 years, thereby fuelling higher secondary and tertiary production rates. For instance, a high rate of N loss would tend to favour production within the microbial loop, thereby decreasing the efficiency of N transfer to higher trophic levels. These higher secondary and tertiary production rates may, in turn, accelerate carbon drawdown from surface layers of the oceans.

Recently, Mohr *et al.* (2010b) have addressed potential problems associated with the execution of
the $^{15}$N$_2$ isotope uptake method, suggesting that gas solubility issues can potentially lead to large underestimates of actual net N$_2$-fixation rates if the technique is not applied properly. The PN accumulation rate is another method that estimates net N$_2$-fixation rates (Kranz et al., 2009). In our experiments, the gross : PN accumulation ratio was close to 1 at light levels that were non-limiting to growth. A ratio of 1 seems more reasonable than the very high estimates of the gross : net N$_2$-fixation rate ratios (up to 15) that we documented using the isotope uptake method. But in support of the $^{15}$N$_2$ isotope uptake method, $^{15}$N$_2$-fixation rates and growth rates were strongly correlated in all of our experiments. In addition, $^{15}$N$_2$ injections probably equilibrated with non-isotope N$_2$ gas during our 12-h incubations. We note that gross N$_2$-fixation rates in the CO$_2$–light experiment with WH0401 were amplified, in comparison with other experiments, because of the modification of the acetylene assay (see N$_2$ fixation in the Materials and methods section). These higher gross N$_2$-fixation rates amplified gross : net ratios in the CO$_2$–light experiment with WH0401. Because the method for the acetylene assay technique was the same in the light and CO$_2$ experiments, our best comparisons of the gross : net N$_2$-fixation rate ratios between strains are those shown in Figs 1 and 2.

In summary, the growth rates of the large-celled strain (WH0402) were higher than those observed for the smaller-celled strain (WH0401). Our data also imply that WH0402 might have a stronger ability to sequester inorganic carbon than WH0401 at 155 μE m$^{-2}$ s$^{-1}$. This conclusion is based on the difference in growth rate reduction between isolates in response to low pCO$_2$ compared to air treatments in the CO$_2$ experiments (experiments 3 and 4; ≥ 40% for WH0401; 10–15% for WH0402). These data indicate that $K_p$ for WH0401 with respect to CO$_2$ might be close to or higher than 190 ppm pCO$_2$, whereas that for WH0402 is lower. A strong ability to sequester inorganic carbon may be the reason that WH0402 has higher growth, N$_2$-fixation and CO$_2$-fixation rates than WH0401.

Our study suggests that unicellular diazotrophic cyanobacteria may have strain-specific responses to interacting variables such as CO$_2$ and light. Similar differences may exist in the responses of different strains to changes in temperature, or the availability of essential nutrients such as phosphorus and iron, as well as to the interactions between all of these factors. Because global change in the ocean involves simultaneous shifts in each of these variables, our work emphasizes the need to understand multivariate effects in the context of the diversity within the genus *Crocosphaera*, rather than making broad generalizations based on studies using only single isolates.

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