



Large centric diatoms allocate more cellular nitrogen to photosynthesis to counter slower RUBISCO turnover rates

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Diatoms contribute ~40% of primary production in the modern ocean and encompass the largest cell size range of any phytoplankton group. Diatom cell size influences their nutrient uptake, photosynthetic light capture, carbon export efficiency, and growth responses to increasing $p\text{CO}_2$. We therefore examined nitrogen resource allocations to the key protein complexes mediating photosynthesis across six marine centric diatoms, spanning 5 orders of magnitude in cell volume, under past, current and predicted future $p\text{CO}_2$ levels, in balanced growth under nitrogen repletion. Membrane bound photosynthetic protein concentrations declined with cell volume in parallel with cellular concentrations of total protein, total nitrogen and chlorophyll. Larger diatom species, however, allocated a greater fraction (by 3.5-fold) of their total cellular nitrogen to the soluble Ribulose-1,5-bisphosphate Carboxylase Oxygenase (RUBISCO) carbon fixation complex than did smaller species. Carbon assimilation per unit of RUBISCO large subunit ($\text{C RbcL}^{-1} \text{ s}^{-1}$) decreased with cell volume, from ~8 to ~2 $\text{C RbcL}^{-1} \text{ s}^{-1}$ from the smallest to the largest cells. Whilst a higher allocation of cellular nitrogen to RUBISCO in larger cells increases the burden upon their nitrogen metabolism, the higher RUBISCO allocation buffers their lower achieved RUBISCO turnover rate to enable larger diatoms to maintain carbon assimilation rates per total protein comparable to small diatoms. Individual species responded to increased $p\text{CO}_2$, but cell size effects outweigh $p\text{CO}_2$ responses across the diatom species size range examined. In large diatoms a higher nitrogen cost for RUBISCO exacerbates the higher nitrogen requirements associated with light absorption, so the metabolic cost to maintain photosynthesis is a cell size-dependent trait.

Keywords: *Coscinodiscus*, diatom, ocean acidification, photosynthesis, resource allocation, RUBISCO, size scaling, *Thalassiosira*

INTRODUCTION

Nitrogen is essential to life as a major constituent of proteins, nucleic acids, and other macromolecules. In photosynthetic organisms, nitrogen is particularly required for building the abundant complexes mediating photosynthetic carbon assimilation (Dortch et al., 1984; Campbell et al., 2003; Falkowski et al., 2008; Shi and Falkowski, 2008). These nitrogen-rich protein complexes are required for harvesting light as well as driving photosynthetic electron flow that generates the energy and reducing power for inorganic nutrient assimilation, including CO_2 fixation by Ribulose-1,5-bisphosphate Carboxylase Oxygenase (RUBISCO). Nitrogen availability in turn operates as the proximal limiting factor for phytoplankton carbon assimilation (Gruber and Galloway, 2008; Moore et al., 2013) in much of the modern ocean. Phytoplankton span a wide size range (Finkel et al., 2010). Under steady state growth, smaller cells have higher rates of nutrient diffusional delivery and lower cellular nutrient requirements for growth (Pasciak and Gavis, 1974; Gavis, 1976; Finkel et al., 2010; Marañón et al., 2013). Consequently, smaller cells dominate nitrogen-limited regions whereas larger cells are typically constrained to well mixed

waters where nitrogen and other nutrients are replete (Irwin et al., 2006), although some large species exploit nitrogen limited waters through symbiotic N_2 fixation (Foster et al., 2011). Larger phytoplankton cells also incur significant optical packaging effects (Falkowski, 1981; Morel and Bricaud, 1981; Kirk, 1994; Finkel, 2001), which lower the photons absorbed per unit of pigment-protein complex. These changes in optical performance decrease the return of captured photons per unit of metabolic nitrogen invested into pigment protein complexes (Raven, 1984). After the captured photons initiate charge separation in a photosystem, the electrons flow along an electron transport chain comprised of abundant protein complexes, to deliver reductant for the Calvin Cycle to generate the Ribulose-1,5-bisphosphate acceptor molecules for carbon fixation by RUBISCO (Ribulose-1,5-bisphosphate Carboxylase Oxygenase). Some fraction of the reductant is then ultimately retained in assimilated organic material (Kroon and Thoms, 2006; Kromkamp et al., 2008; Suggett et al., 2009; Halsey et al., 2010, 2013).

Seawater $p\text{CO}_2$ concentrations are currently too low to saturate RUBISCO carboxylation activity via passive CO_2 diffusion

into the cell (Badger et al., 1998; Tortell et al., 2000) and phytoplankton thus employ carbon concentrating mechanisms (CCMs) to increase the CO₂ concentration around their RUBISCO complexes (Raven et al., 2011, 2014). This need for CCMs is particularly pronounced in larger cells where CO₂ diffusion is limited by a smaller surface area to volume ratio (Giordano et al., 2005; Wu et al., 2014). RUBISCO represents a moderate fraction (~2–5%) of total protein in marine phytoplankton (Brown et al., 2008; Losh et al., 2013; Li et al., 2014; Young et al., 2015) which, combined with C assimilation rates, mean that achieved Rubisco catalytic carboxylation turnover rates (C RbcL⁻¹ s⁻¹) in phytoplankton are generally faster than achieved rates in terrestrial plants. Such high achieved rates of carboxylation, at least in the smallest (nano–pico) phytoplankton, are in part attributable to taxonomic differences in RUBISCO catalytic properties (Badger et al., 1998; Young et al., 2012) as well as the structure and operation of CCMs (Losh et al., 2013; Raven, 2013) that likely impose substantial energy and macromolecular costs for phytoplankton to meet their CO₂ requirements in the current ocean. As such, increased pCO₂ availability via ocean acidification could potentially afford cells a means to lower these costs (Rost et al., 2008; Hopkinson et al., 2011; Wu et al., 2014). However, experiments simulating elevated pCO₂ predicted for future oceans demonstrate variable growth responses across phytoplankton (Tortell and Morel, 2002; Schippers et al., 2004; Boelen et al., 2011; Gao et al., 2012; McCarthy et al., 2012; Li and Campbell, 2013) since interacting factors including light (Gao et al., 2012), nutrient limitation (Beardall et al., 2009), taxa (Langer et al., 2006), and cell size (Wu et al., 2014) can strongly influence the net growth outcomes under ocean acidification.

Diatoms contribute ~40% of productivity in the modern ocean, and span the widest size range of any phytoplankton group, from several μm to several 100 μm in cell diameter (Finkel et al., 2010). Within the diatoms, fewer than 50 species contribute >90% of all diatom biomass globally, with centric species dominating (LeBlanc et al., 2012). With increasing cell volume across species, diatoms exhibit a decrease in many key traits, notably growth rate, light capture per chlorophyll, photosynthesis rates per unit chlorophyll-*a* and susceptibility to photoinactivation of Photosystem II (Mullin et al., 1966; Burkhardt et al., 1999; Finkel, 2001; Marañón et al., 2007, 2013; Key et al., 2010). It is, however, unknown how these size-scaling patterns interact with (re) distributions of nitrogen amongst the catalytic complexes governing CO₂ assimilation and growth, and whether size-scaling in resource allocations alter in response to elevated pCO₂. We recently found (Wu et al., 2014) that increased pCO₂ differentially stimulates the growth of larger vs. smaller centric diatoms, apparently through a release of diffusional limitations upon CO₂ delivery to the surfaces of larger cells. We therefore grew 6 strains of marine centric diatoms, spanning five orders of magnitude in cell volume, under 3 pCO₂ levels, with growth-saturating light and nutrient repletion, to investigate interactive effects of cell size and pCO₂ on photosynthetic performance and resource allocations of cellular nitrogen quotas. We determined nitrogen investment into core photosynthetic constituents across the species and pCO₂ range, and analyzed in parallel how changing macromolecular investments offset changes in the achieved photosynthetic

turnover rates of the abundant catalytic complexes mediating photosynthesis.

MATERIALS AND METHODS

SPECIES, CULTURE CONDITIONS, SAMPLING AND GROWTH RATES

Six marine centric diatom strains, *Thalassiosira pseudonana* (CCMP 1335); *Thalassiosira oceanica* (CCMP 1003); *Thalassiosira guillardii* (CCMP 988); *Thalassiosira weissflogii* (CCMP 1336); *Thalassiosira punctigera* (CCAP 1085/19) and *Coscinodiscus wailesii* (CCMP 2513) were maintained in mid-exponential growth through semi-continuous culturing at 20°C (as per Wu et al., 2014) in 1 L flasks. Cultures were illuminated with cool fluorescent tubes at a photon flux density of 350 μmol m⁻² s⁻¹, measured with Li-Cor spherical PAR sensor in the center of the incubation bottle (8 × 8 × 16 cm), under a 12:12 light:dark cycle, and always sampled 6 h after lights on. After the initial measurement of light levels in the center of the flask, we did not perform further light measures as the cultures grew, but the cultures were optically thin and the light level was saturating for growth of these species under these conditions (data not shown). The analyses in this paper draw upon the same culture experiments and some of the underlying data sets supporting (Wu et al., 2014), but include data from an additional species *Thalassiosira oceanica* (CCMP 1003).

Nutrient replete *f/2* media was made using artificial seawater prepared with a total alkalinity (TA) of ~2300 μmol kg⁻¹, at three different pH (8.42, 8.16, and 7.90) corresponding to pCO₂ levels of ca. 190, 380, and 750 ppmv (Riebesell et al., 2010; Wu et al., 2014) and filtered to remove debris. The dissolved inorganic carbon (DIC) concentration in the media was measured with an infrared gas analyzer (QuBit, Canada) (McCarthy et al., 2012) and absolute pCO₂ subsequently computed using CO₂SYS software (Pierrot et al., 2006) from the known values of DIC, pH, salinity, and alkalinity of the media. Culture vessels were sealed to avoid gas exchange with the atmosphere, and maintained at low biomass with chlorophyll *a* < 10 μg L⁻¹ by regular dilution with pre-equilibrated medium. Using this approach, culture pH variability remained within 0.05 pH units of the target levels throughout the growth experiments. Flasks were manually agitated 4–5 times daily and distributed arbitrarily within the growth chamber.

Daily samples from the different pCO₂ cultures were fixed with Lugol's acid solution for determination of growth rate and cell size, with analyses within a week of fixation. Species with diameters <40 μm were counted with a Coulter Counter (Multisizer 3, Beckman) using the filtered media as a blank. Larger species were counted under the microscope using a Sedgwick Rafter chamber. The cell specific growth rate (μ, d⁻¹) for each of the strains was determined by ordinary least squares linear regression of ln cell density, corrected for culture dilutions with media, over time. Cell volume for all species was estimated from linear dimensions from images taken with a camera attached to a light microscope calibrated with a micrometer, using Image-J software and assuming the cells were cylinders. For all samples a minimum of 100 cells were measured.

After at least eight rounds of cell division with steady growth rate (Wu et al., 2014) to ensure acclimation to the pCO₂ regime,

cell samples for further analyses were harvested by mild vacuum filtration onto glass fiber filters (0.22 μm pore size, Whatman; vacuum pump at 0.05–0.06 MPa). Filters were pulled dry of bulk media but were not further rinsed for most analyses. During harvest about half of the culture volume was taken, across multiple sub-samples for different analyses. Cells on filters were flash frozen in liquid nitrogen and stored at -80°C until later analyses of protein composition (25 mm filters) or CN elemental analyses (13 mm pre-combusted filters). Each combination of taxa and $p\text{CO}_2$ was cultured independently in at least 3 separate flasks. All subsequent analyses and data transformations were conducted independently using matched samples and measures taken at the same sampling time from each flask separately. Final plotted points present the average of the determinations from each flask. In this way we avoided propagation of errors in the multi-step data transformations.

PROTEIN QUANTITATION AND NITROGEN RESOURCE ALLOCATION ESTIMATES

Total protein was extracted from the cells on the frozen glass fiber filters using an FastPrep-24 with the 24×2 rotor (MPBio). Each frozen filter was transferred to a 2 mL tube containing bead lysing matrix D (MPBio) and 700 μL of extraction buffer (140 mM TRIS base, 100 mM TrisHCL, 75 mM LDS, 0.5 mM EDTA, 0.4 mM protease inhibitor (AEBSE, BioShop), 10% w/v glycerol). The FastPrep was set to run at 6.5 m/s for three 1-min cycles. Samples were held on ice for 1 min between each cycle. Following cell lysis, each sample was centrifuged for 3 min at $10,000 \times g$ and the supernatant assayed for total protein concentration using the BioRad DC protein assay kit with the mixed protein preparation Bovine Gamma Globulin (Pierce) standards of known concentration.

RUBISCO, Photosystem II (PSII), Cytb₆f, and Photosystem I (PSI) protein complex contents were estimated using quantitative immunoblotting of representative subunits RbcL, PsbA, PetC, and PsaC respectively (Brown et al., 2008), by comparison to four point calibrated protein quantitation standard curves for each specific antibody (AgriSera, AS03-037, and AS01-017S; AS05-084 and AS01-016S; AS08 330 and AS08 330S, AS10 939 and AS04 042S). The secondary antibody goat anti-rabbit IgG was used for all blots in a dilution of 1: 50,000 (ImmunoReagents Inc, lot 14-122-042810).

Cells were harvested onto 13 mm GF/F filter for measurement of total particulate organic nitrogen, which was assumed to be equivalent to cellular nitrogen, measured using an elemental analyzer (VarioIII EL, Elementar), and normalized to cell counts (Wu et al., 2014). The fraction of total N allocated to RUBISCO, to Photosystem II and to the summed protein complexes of the photosynthetic system was estimated using molar quantitations of Chl a and Chl c by absorbance spectrophotometry (Jeffrey and Humphrey, 1975), and the representative protein subunits PsbA, PsbD, PetC, PsaC, and RbcL by quantitative immunoblotting. We multiplied the molar content of each indicator molecule by its molecular nitrogen content. We estimated nitrogen contents for each protein complex as the sum of all amino acid nitrogen atoms in the protein subunit amino acid sequences predicted from the genome of *Thalassiosira*

pseudonana (Armbrust et al., 2004), multiplied by the stoichiometry for conserved subunits within the complex as determined from crystal structures of PSII (Umena et al., 2011) (for PsbD or PsbA), the Cytochrome b₆f complex (Baniulis et al., 2008) (for PetC), the PSI complex (Amunts et al., 2010) (for PsaC) or the RUBISCO complex (Mizohata et al., 2002) (for RbcL). To estimate the nitrogen allocation to the fucoxanthin-chlorophyll protein antenna complex (FCP) we used a content of two Chl c per FCP monomer (Premvardhan et al., 2010) to estimate the molar content of FCP monomers, and multiplied by the nitrogen content of a representative FCP protein monomer (Nagao et al., 2013) based upon the protein sequence (FCP8 *Thalassiosira pseudonana* NCBI XP_002290137.1). It should be noted that this approach of summed nitrogen allocation to the photosynthetic system is conservative since we did not include the ATP synthase complex, nor protein complexes mediating carbon concentration nor the Calvin Cycle aside from RUBISCO.

To compare our elemental analyses of cellular C and N to photosynthetic parameters, the electron equivalent content of the accumulated C and N per biovolume was estimated as:

$$\begin{aligned} & \text{assimilated } e^- \text{ cell volume}^{-1} \\ & = \{(C/\text{cell} \times 4 e^-/C) + (N/\text{cell} \times 10 e^-/N)\}/V(\mu\text{m}^3 \text{cell}^{-1}) \end{aligned} \quad (1)$$

C/cell and N/cell were estimated from elemental analyses of particulate organic carbon or particulate organic N and cell counts. To the extent that our cultures contained non-living particulate debris (Agusti and Sanchez, 2002) our estimates of C & N assimilation rates will be exaggerated. The source for assimilated C in our cultures is dissolved CO_2 which requires a minimum of $4 e^-/C$ for reductive assimilation into biomass, although this value is likely an underestimate since diatoms accumulate C in the form of lipids (Halsey et al., 2010, 2013). The source for assimilated N was dissolved NO_3^- which requires a minimum of $10 e^-/N$ for reductive assimilation into biomass.

PHOTOSYNTHETIC PARAMETERS

The PSII electron transport rate ($e^- \bullet \text{PSII}^{-1} \bullet \text{s}^{-1}$) during the 12 h illuminated growth period was estimated (Suggett et al., 2004, 2009) based on photosynthetic parameters measured via FRe fluorometry (Satlantic) as:

$$\begin{aligned} & \text{Electron transport rate } (e^- \bullet \text{PSII}^{-1} \bullet \text{s}^{-1}) \\ & = \text{PPFD} \bullet \sigma_{\text{PSII}'} \bullet \Phi_{\text{PSII}'} / \Phi_{\text{PSII}} \bullet A \end{aligned} \quad (2)$$

Where PPFD is the growth photon flux ($350 \mu\text{mol photon m}^{-2} \text{s}^{-1}$) measured with a Walz microspherical quantum sensor (Walz, Germany). σ'_{PSII} ($\text{m}^2 \text{quanta}^{-1}$) is the PSII effective absorption cross section (Kolber et al., 1998) measured with a background actinic light approximately equivalent to the growth light level. σ'_{PSII} is estimated by fast repetition rate fluorometry induction (Laney, 2003) with a blue light emitting diode source from the FRe fluorometer and thus was corrected to the white growth light using a spectral correction factor for diatom absorbance of blue vs. white light (Suggett et al., 2009). $\Phi_{\text{PSII}'}$

is the PSII photochemical efficiency under growth light, while Φ_{PSII} is the maximum PSII efficiency in a dark-acclimated state. The factor A converts 1.0 mol photon absorbed by PSII to 1.0 mol electron generated by PSII (Suggett et al., 2009).

Apparent carbon assimilation per RUBISCO ($C \text{ RbcL}^{-1} \text{ s}^{-1}$) was estimated as:

$$\begin{aligned} \text{RUBISCO turnover rate } (C \text{ RbcL}^{-1} \text{ s}^{-1}) \\ = (C \text{ cell}^{-1}) \bullet (e^{\mu} - 1) \bullet (\text{cell RbcL}^{-1}) \bullet 2 \quad (3) \end{aligned}$$

$C \text{ cell}^{-1}$ was estimated from elemental analyses of particulate organic carbon normalized to cell counts, μ was the cell specific exponential growth rate (s^{-1}), and RbcL was from immunoquantitation data normalized to cell counts. This estimator is based upon net assimilated carbon per cell, and upon cellular growth rate integrated over a full 12:12 diel illumination cycle. Carbon assimilation is actually restricted to the illuminated fraction of the day, and thus we also include the factor of 2 to account for carbon assimilation $\text{s}^{-1} \text{ RbcL}^{-1}$ during the illuminated fraction of the 12:12 diel cycle. This likely underestimates actual carbon fixation per RUBISCO during the illuminated period, since we neglect C lost from the cell through respiration or release of extracellular products (López-Sandoval et al., 2013), but as noted above, any contribution of non-living particulate organic carbon will exaggerate the apparent C living cell^{-1} . The protein-specific electron generation rate ($e^{-} \mu\text{g protein}^{-1} \text{ s}^{-1}$) and the protein specific carbon assimilation rate ($C \mu\text{g protein}^{-1} \text{ s}^{-1}$) were then derived from the PSII electron transport rate and the apparent carbon assimilation per RUBISCO, both normalized to cellular protein content.

RESULTS

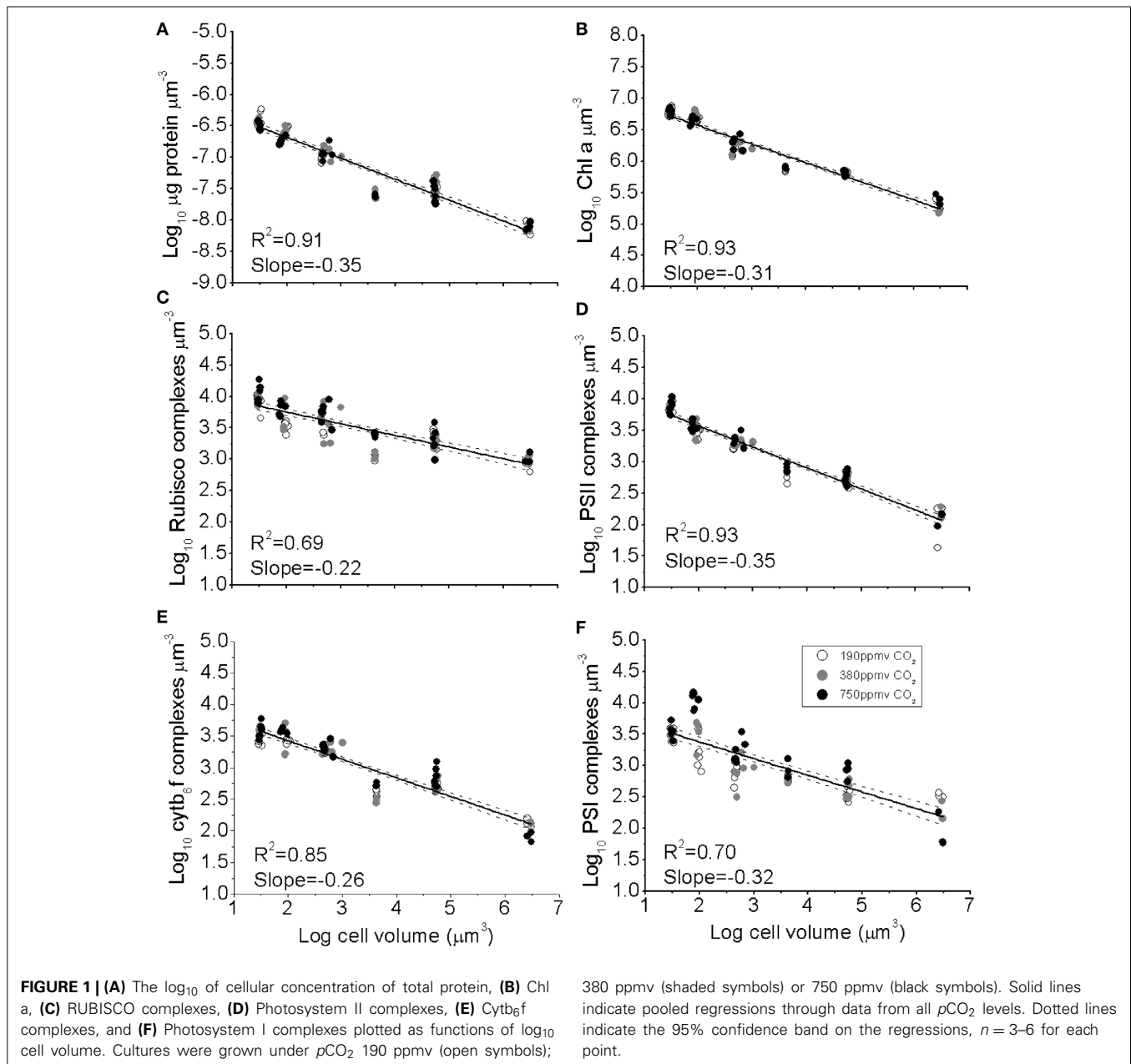
Total protein concentration per unit biovolume decreased with increasing cell volume, from $\sim 0.37 \times 10^{-6} \mu\text{g protein } \mu\text{m}^{-3}$ for the smallest species, *T. pseudonana*, to $\sim 8 \times 10^{-9} \mu\text{g protein } \mu\text{m}^{-3}$ for the largest species, *C. wailesii*, (Figure 1A). The extracted protein quantified through the dye binding assay represented $88 \pm 7\%$ ($n = 89$, $\pm 95\%$ confidence interval) of the particulate organic nitrogen from the same cultures, determined using elemental analyses. The ratio of extracted protein to particulate nitrogen varied somewhat among the species. In particular, in the largest species *Coscinodiscus wailesii* detected protein represented $50 \pm 9\%$ ($n = 9$, $\pm 95\%$ confidence interval) of total (particulate) nitrogen. We are currently pursuing this issue to determine whether the variation is biological, technical, or both. For subsequent analyses, levels of protein complexes are expressed as number of complexes per cell, or as the fraction of cellular (particulate) nitrogen allocated to the complex, and issues of protein quantitation cancel out in the data transformations.

The concentrations of photosynthetic complexes per unit biovolume also declined with increasing cell volume (Figures 1B–E), from $\sim 7500 \text{ PSII } \mu\text{m}^{-3}$, $\sim 3500 \text{ PSI } \mu\text{m}^{-3}$, $\sim 3500 \text{ Cytb}_{6f} \mu\text{m}^{-3}$, and $\sim 9500 \text{ RUBISCO (L8S8 complex) } \mu\text{m}^{-3}$ for the smallest species, down to $\sim 130 \text{ PSII } \mu\text{m}^{-3}$, $\sim 170 \text{ PSI } \mu\text{m}^{-3}$, $110 \text{ Cytb}_{6f} \mu\text{m}^{-3}$, and $\sim 960 \text{ RUBISCO } \mu\text{m}^{-3}$ for the largest species. We did not find significant effects of $p\text{CO}_2$ on the model 2 regression

slopes (Figure 1), which parameterise the size-scaling for each complex; ($\log_{10}(Y) = b * \log_{10}(\text{cell volume}) + Y \text{ intercept}$). Thus, a single size-scaling model was used to describe the cell size dependency for each protein pool across all $p\text{CO}_2$ treatments pooled; specifically $b = -0.349 \pm 0.010$ ($\pm 95\%$ confidence intervals) for total protein (Figure 1A); -0.306 ± 0.007 for Chl a (Figure 1B); -0.347 ± 0.013 for PSII (Figure 1D), -0.256 ± 0.009 for Cytb_{6f} (Figure 1E) and -0.317 ± 0.021 for PSI (Figure 1F). RUBISCO, in contrast, showed a significantly smaller size scaling exponent ($b = -0.223 \pm 0.012$, Figure 1C), showing that relatively more of the total protein pool was allocated to RUBISCO as cell size increases.

The fraction of total cellular nitrogen allocated to the summed photosynthetic complexes increased with cell volume, from the smaller species to larger species (Figure 2A; Table 1). In spite of this increased relative allocation to photosynthesis, the cell specific growth rates across the range of species decreased significantly with cell size (Wu et al., 2014) with scaling exponents of -0.071 for cells growing under 190 ppmv $p\text{CO}_2$, -0.064 under 380 ppmv $p\text{CO}_2$ and -0.052 under 750 $p\text{CO}_2$. We detected $p\text{CO}_2$ effects upon protein allocations in individual species; notably, the photosynthetic N-fraction in *T. oceanica* and *T. pseudonana* increased with elevated $p\text{CO}_2$ treatments (Figure 2A) but showed no consistent trend for larger species (Figure 2A). The total nitrogen-fraction allocated to PSII ranged from 0.003 to 0.016 and increased significantly under elevated $p\text{CO}_2$ conditions in the smallest species, *T. oceanica* and *T. pseudonana* ($p < 0.05$) (Figure 2B). The RUBISCO N-fraction ranged widely from 0.01 to 0.08, and increased with cell volume (Table 1) consistent with the size scaling of cellular RUBISCO content per μm^{-3} relative to total protein concentration (above, Figures 1A,C). Interestingly, under our nitrogen replete growth conditions, there was no correlation between growth rate and the fraction of cellular nitrogen allocated to photosynthesis nor to RUBISCO (data not shown). We are currently conducting analogous experiments under nitrogen limitation to pursue this issue.

PSII electron transfer rates estimated from chlorophyll fluorescence parameters in the light Equation (2), ranged from $\sim 330 e^{-} \text{ PSII}^{-1} \text{ s}^{-1}$ in the smallest species, *T. pseudonana*, down to $\sim 150 e^{-} \text{ PSII}^{-1} \text{ s}^{-1}$ in the largest species examined, *C. wailesii* (Figure 3A), as a consequence of packaging effects on excitation delivery to PSII (Figure 4). The apparent C assimilated per RUBISCO Equation (3) decreased from $\sim 8 C \text{ RbcL}^{-1} \text{ s}^{-1}$ in the smallest species down to $\sim 2 C \text{ RbcL}^{-1} \text{ s}^{-1}$ in the largest species (Figure 3B). Thus, the decrease in achieved turnover with increasing cell size was greater for RUBISCO than for PSII. The apparent RUBISCO turnover rate was significantly lower under elevated $p\text{CO}_2$ for *T. oceanica* and *T. weissflogii* ($p < 0.05$) (Figure 3B, Table 1). PSII electron generation normalized to total protein did not exhibit any general trend with cell volume but did exhibit $p\text{CO}_2$ effects for individual species (Figure 3C, Table 1). Across the size spectrum, carbon assimilation rates normalized to total cellular protein did not show a trend from small to large diatoms (Figure 3D, Table 1), since large species generally allocated more of their cellular nitrogen into RUBISCO, which countered the lower apparent turnover rate of their RUBISCO complexes (Figure 3B).



A measure of PSII light capture capacity, the effective absorption cross section serving PSII (σ_{PSII}), decreased from $320 \text{ A}^2 \text{ PSII}^{-1}$ in the smallest species down to $130 \text{ A}^2 \text{ PSII}^{-1}$ in the largest species (Figure 4A). In contrast, the molar ratio of Chl a (Figure 4B) and Chl c (Figure 4C) to PSII reaction center content varied amongst species, with an average of $\sim 1000 \text{ mol Chla: mol PSII}$, and $\sim 220 \text{ mol Chlc: mol PSII}$ (Figure 4C), but with no significant trend with cell size. We found no consistent $p\text{CO}_2$ effects for these light capture capacity parameters.

Carbon accumulated per generated electron (carbon e^{-1}), varied from 0.11 to 0.28 across species (Figure 5A), with no consistent trends observed with cell size or with $p\text{CO}_2$ (Table 1). A regression of accumulated electron equivalents invested to assimilated N and C per cell volume Equation (1) vs. electrons generated

by PSII per cell volume ($e^{-} \mu\text{m}^{-3} \text{ s}^{-1}$) shows a ratio of assimilated organic matter conserving $\sim 95\%$ of the predicted electrons generated, across the range of diatom sizes (Figure 5B).

DISCUSSION

We grew a size spectrum of marine centric diatoms under nutrient replete and light-saturated growth at three levels of $p\text{CO}_2$. As expected (Mullin et al., 1966; Finkel, 2001; Marañón et al., 2007, 2013) the cellular concentrations of total protein, chlorophyll and the membrane bound photosynthetic complexes Photosystem II, Photosystem I and Cytochrome b_6f all declined with increasing cell volume, with similar negative scaling exponents. RUBISCO, however, showed a less negative size scaling, so that larger diatoms allocated a larger fraction of their total protein toward RUBISCO,

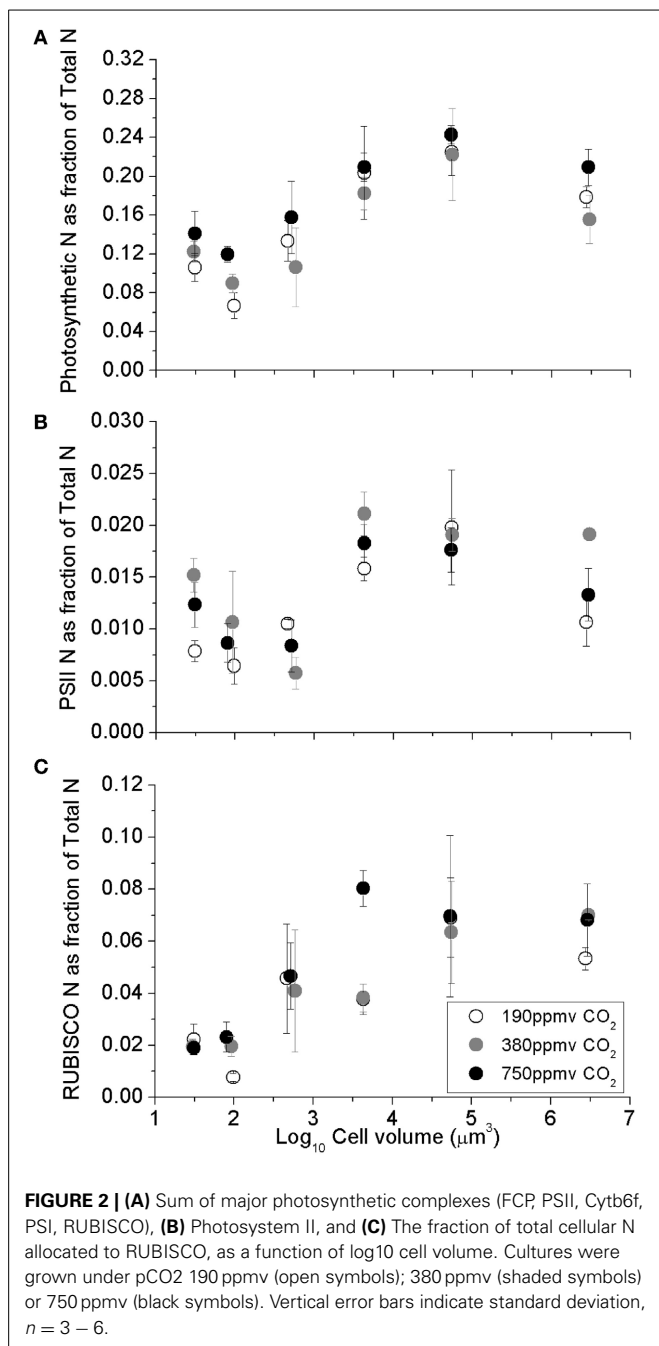


FIGURE 2 | (A) Sum of major photosynthetic complexes (FCP, PSII, Cytb6f, PSI, RUBISCO), **(B)** Photosystem II, and **(C)** The fraction of total cellular N allocated to RUBISCO, as a function of log_{10} cell volume. Cultures were grown under $p\text{CO}_2$ 190 ppmv (open symbols); 380 ppmv (shaded symbols) or 750 ppmv (black symbols). Vertical error bars indicate standard deviation, $n = 3 - 6$.

which in turn drove up the fraction of their total nitrogen allocated to their summed photosynthetic system. As with previous studies (Losh et al., 2013; Li et al., 2014; Young et al., 2015), RUBISCO represented only $\sim 2\%$ of total cellular nitrogen in the smaller diatoms but increased to $\sim 8\%$ in the larger diatoms (Figure 2C; Table 1). Thus, bigger diatoms make a greater investment of cellular nitrogen into carbon-assimilation capacity relative to photosynthetic electron generation, compared to smaller diatoms.

Increased $p\text{CO}_2$ differentially stimulated growth rates by up to 15% in the largest of these centric diatoms (Wu et al., 2014). In our relatively short-term acclimatory $p\text{CO}_2$ manipulation

experiment we observed significant nitrogen allocation responses to $p\text{CO}_2$ in some species. For example the fraction of total N allocated to the photosynthetic complexes increased with increased $p\text{CO}_2$ in the small species *T. oceanica* and *T. weissflogii* (Figure 2A), although at our level of replication these changes did not generate significant correlations with growth rate. Such species-specific responses are consistent with the growing evidence of variable responses to $p\text{CO}_2$ availability across taxa, even within closely related phytoplankton taxonomic groups (Trimborn et al., 2009, 2013; Brading et al., 2013; Gao and Campbell, 2014). Nevertheless, increased CO_2 did not significantly change the size scaling of nitrogen allocations to the photosynthetic system complexes across our size range of centric diatoms. This generalized size scaling pattern across $p\text{CO}_2$ levels suggest that the size-dependent increase in nitrogen costs to maintain C-assimilation outweighs any changes in costs associated with changing $p\text{CO}_2$ (Raven et al., 2014).

To assess comparative physiological returns upon nitrogen investments across the diatom size spectrum and $p\text{CO}_2$, we estimated PSII electron transport per PSII complex, following Suggett et al. (2009), but using PsbA protein content as a proxy for the content of functional PSII. As expected given the optical packaging effects that decrease effective absorption cross sections with increasing diatom cell size (Figure 4) (Kirk, 1994; Finkel, 2001; Key et al., 2010; Wu et al., 2014), achieved electron transport per PSII decreased significantly with increasing cell size. In parallel, we used the cellular growth rate, carbon per cell, and cellular RbcL protein content to estimate the achieved carbon assimilation per RUBISCO active site. As with (Losh et al., 2013; Young et al., 2015), the smaller diatoms showed apparent carbon assimilation rates per RUBISCO close to the catalytic turnover maxima expected for RUBISCO (Badger et al., 1998; Badger and Bek, 2007) but these apparent rates declined with increasing cell size.

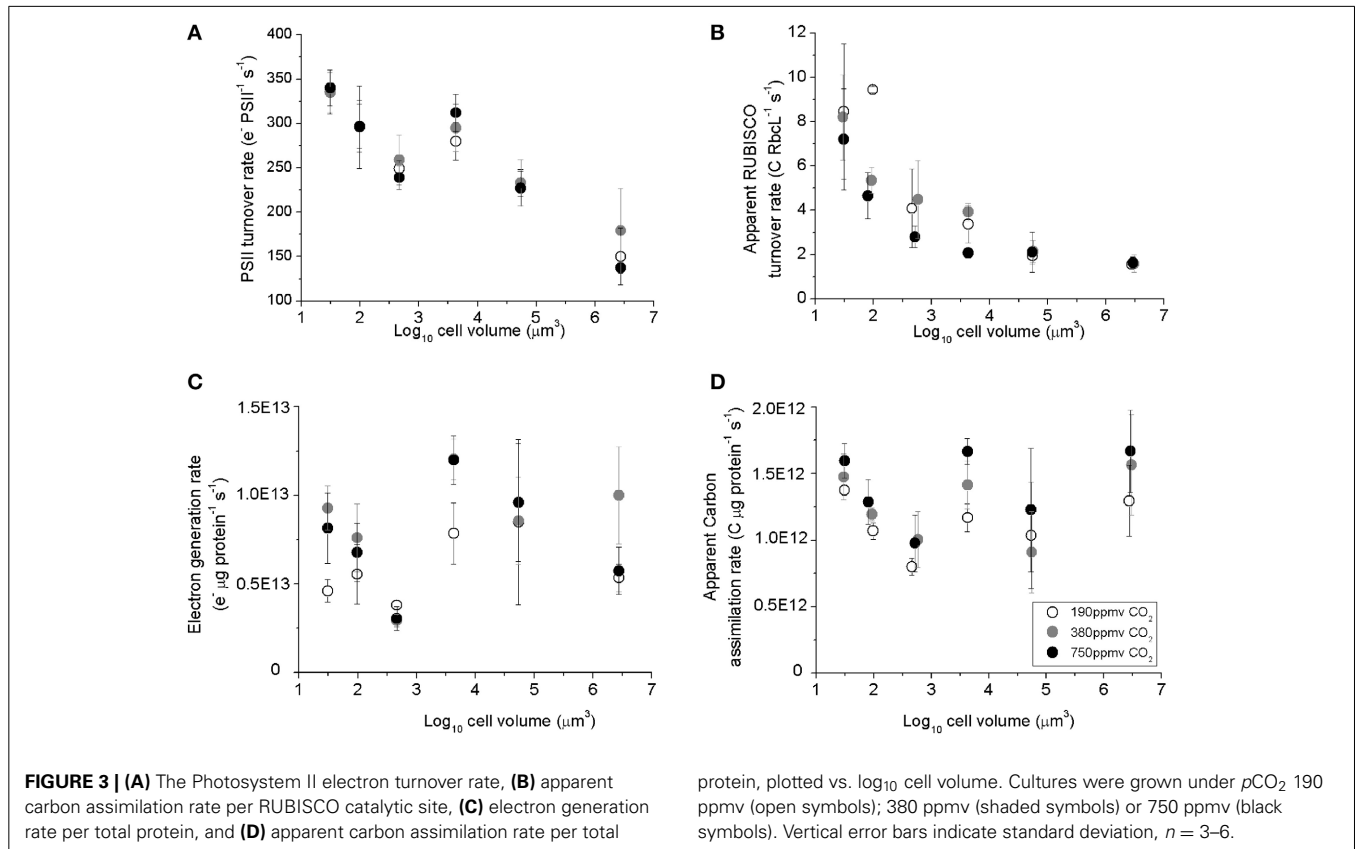
We propose initial, non-exclusive hypotheses to explain the cell size-dependent decline in apparent C assimilation per RUBISCO.

- (i) We observed a significant $p\text{CO}_2$ enhancement of growth rate of $\sim 15\%$ under 750 ppmv $p\text{CO}_2$ in the largest centric diatoms (Wu et al., 2014), which was explicable via relief from passive CO_2 diffusion limitation to the cell surface of larger diatoms. An increase in RUBISCO protein allocation in larger cells could be driven to compensate for a size-dependent limitation on CO_2 diffusion. As the immediate sink for CO_2 , higher cellular RUBISCO content could drive faster CO_2 uptake per cell. Although we have not yet tested pennate diatoms, their higher surface to volume ratios could alter these responses to $p\text{CO}_2$.
- (ii) The decline in achieved RUBISCO performance could reflect negative scaling of metabolic rates with cell size (Burkhardt et al., 1999; Finkel, 2001; Marañón et al., 2013), which could limit the rate of metabolic regeneration of the Ribulose-1,5-bisphosphate acceptor molecule for RUBISCO carbon fixation (Mott et al., 1984; Gotoh et al., 2010).
- (iii) Since our C-assimilation estimates are based upon retained particulate organic carbon, an increase in extracellular carbon release from larger diatoms, or a size-dependent increase in respiratory losses to support metabolic needs,

Table 1 | Results of Two-Way ANOVA, testing if $p\text{CO}_2$ or cell size had individual or interactive effects on key parameters.

Factor	Photosyn N/ Total N	PSII N/ Total N	Rubisco N/ Total N	PSII ETR	C Rubisco ⁻¹ s ⁻¹	e- μg protein ⁻¹ s ⁻¹	C μg protein ⁻¹ s ⁻¹	C/e ⁻
$p\text{CO}_2$	NS	NS	NS	NS	S	NS	NS	NS
cell size	HS	S	HS	HS	HS	NS	NS	NS
Interaction	NS	NS	NS	NS	NS	NS	NS	NS

$n = 3-6$ for each factor combination. NS, refers to p -values that were not significant ($p > 0.05$), S, significant $0.05 > p > 0.001$, or HS, highly significant ($p < 1 \times 10^{-6}$).



could lower the apparent C assimilation per RUBISCO, although our consistent ratio of accumulated electrons to generated electrons across the cell size spectrum argues against this scenario. If larger diatoms experienced a disproportionate increase in respiratory losses, we would expect a decrease in the ratio of accumulated to generated electrons, but we did not see such a decrease, consistent with previous discussions (Marañón et al., 2013).

- (iv) Our cultures were growing under nutrient repletion, the C:N was around 6.7:1 with no consistent $p\text{CO}_2$ effects (Wu et al., 2014), so the increased allocation to RUBISCO in larger cells could reflect a size-dependent relative increase in luxury accumulation of RUBISCO protein, that in turn lowers achieved performance per unit RUBISCO. Such luxury protein accumulation could relate to increased vacuolization in larger species (Stolte and Riegman, 1995). A comparable experiment with small and large centric diatoms under

nitrogen-limited chemostats to limit luxury accumulation is now underway to test this hypothesis.

- (v) Micro-evolutionary changes in RUBISCO structure/function (Young et al., 2012) could separate small from large diatoms. We compared the RbcL (large subunit of RUBISCO) sequences across our panel of diatoms (NCBI gi118411104, gi315320529, gi98990579, gi98990561, gi98990581, gi98990595, gi329343318) as well as RbcS (small subunit of RUBISCO) sequences available for the studied species or con-generics (NCBI gi327258980, gi327258982, gi315320528, gi283569001, gi468111925). As expected, the sequences are highly conserved across the species with no amino acid differences apparent within the active site nor within the RbcL:RbcL homodimer interface surfaces. There are, however, suggestive amino acid point changes within regions annotated as the heterodimer RbcL:RbcS dimer interfaces, including a change in RbcS

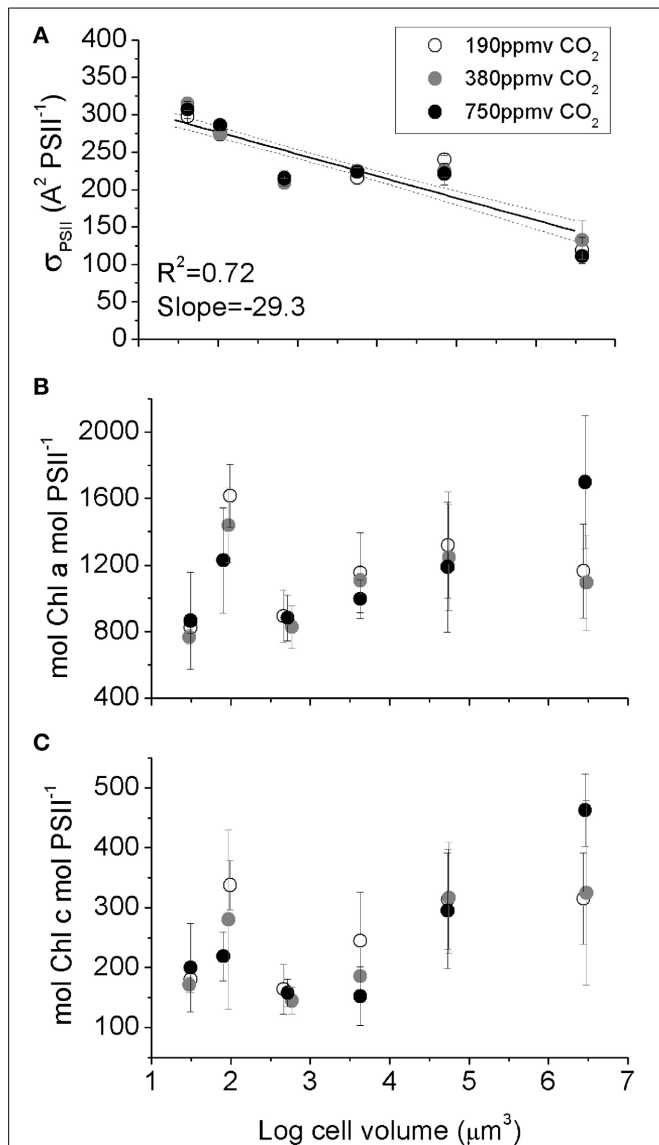


FIGURE 4 | (A) Effective blue light absorbance cross section for Photosystem II, **(B)** Chl a PSII^{-1} , **(C)** Chl c PSII^{-1} plotted vs. \log_{10} cell volume. The significant decrease in σ_{PSII} with increasing cell size **(A)** reflects optical packaging effects because Chl per PSII **(B,C)** does not decline with cell size. Solid line indicates pooled linear regression for samples from the three $p\text{CO}_2$ levels. Dotted lines indicate the 95% confidence band on the regression. Cultures were grown under $p\text{CO}_2$ 190 ppmv (open symbols); 380 ppmv (shaded symbols) or 750 ppmv (black symbols). $n = 3-6$.

at amino acid 46 from NYWEL (amino acid sequence) conserved across the smaller *Thalassiosira* lineages (NCBI gi327258980, gi327258982, gi315320528, gi283569001) compared to SYWEL in an RbcS sequence from a larger *Coscinodiscus* (NCBI gi468111925).

Further analyses of a broader range of diatom RbcS sequences for which we had cell size estimates ($n = 24$) (Finkel, 2007) demonstrated that the 18 taxa with the NYWEL sequence at amino

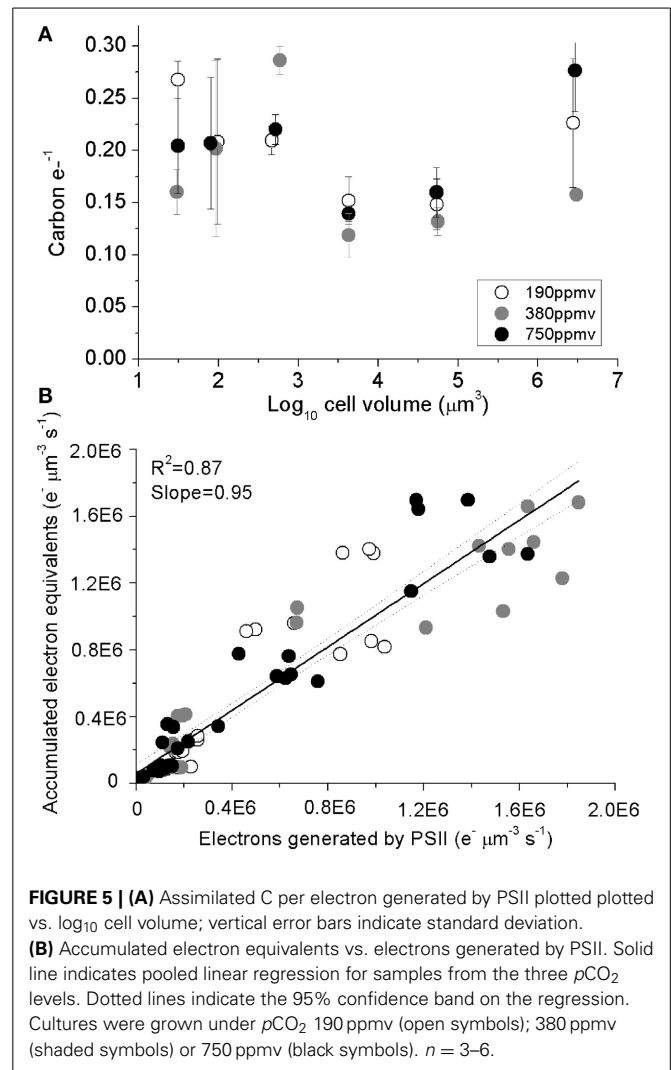


FIGURE 5 | (A) Assimilated C per electron generated by PSII plotted vs. \log_{10} cell volume; vertical error bars indicate standard deviation. **(B)** Accumulated electron equivalents vs. electrons generated by PSII. Solid line indicates pooled linear regression for samples from the three $p\text{CO}_2$ levels. Dotted lines indicate the 95% confidence band on the regression. Cultures were grown under $p\text{CO}_2$ 190 ppmv (open symbols); 380 ppmv (shaded symbols) or 750 ppmv (black symbols). $n = 3-6$.

acid 46 had a median cell biovolume of $\sim 88 \mu\text{m}^3$ compared to the 6 taxa with SYWEL with a median cell biovolume of $\sim 3043 \mu\text{m}^3$ ($p = 0.08$). In parallel RbcL position 171, also annotated as falling at the RbcL:RbcS heterodimer interface, shifts from NKYGTP conserved across the smaller *Thalassiosira* lineages (NCBI gi118411104, gi315320529, gi98990579, gi98990561, gi98990595) to NKYGIP in *Thalassiosira weissflogii* (NCBI gi98990581) and to NKYGAP in the large *Coscinodiscus wailesii* (NCBI gi329343318). Similarly, an even wider alignment of diatom RbcL sequences for which we had cell size estimates ($N = 390$) (Finkel, 2007) showed that the 152 taxa with NKYGTP had a median cell biovolume of $\sim 132 \mu\text{m}^3$, compared to a median cell biovolume of $\sim 452 \mu\text{m}^3$ for the 238 taxa with a hydrophobic amino acid such as I at position 171.

Even though we cannot yet discern the cause(s) for the decreasing C-assimilation per RUBISCO with increasing cell size, larger centric diatoms allocate an increased fraction of their total cellular nitrogen to RUBISCO, and this increase largely offsets the slower achieved C-assimilation rates per unit RUBISCO enzyme.

The reductant equivalents accumulated in the form of cellular C and N represented about 95% of the estimated electrons

generated by PSII, across our diatom size spectrum. This ratio is high compared to other studies based upon ^{14}C uptake or oxygen evolution (Suggett et al., 2009; Halsey et al., 2010, 2013; Lawrenz et al., 2013). Our estimate of cellular C and N was based upon total Particulate Organic Carbon and Particulate Organic Nitrogen, and so likely exaggerates the actual content of C and N from those living cells that contribute to the estimate of photosynthetic electron transport, since our cultures contained some dead cells (Agusti and Sanchez, 2002) and other debris. Even so, the ratio of accumulated to generated reductant did not vary with cell size nor with $p\text{CO}_2$ treatment, indicating a conserved allocation of photosynthetically derived reductant to total C on a per cell volume basis, at least for diatoms under nutrient and light depletion. Furthermore, Antarctic diatoms (Young et al., 2015) during a bloom approached our estimates for the ratio of accumulated reductant per generated electrons, estimated over a diel cycle (Lawrenz et al., 2013) supporting the validity of our approach.

Nitrogen allocation to the macromolecular complexes of the photosynthetic apparatus now represents a predictably size dependent biosynthetic cost for diatoms. We show that large diatoms incur a higher demand upon their nitrogen metabolism capacity to accumulate larger pools of RUBISCO, apparently to counter their slower achieved carbon uptake per RUBISCO. Thus, in addition to the cost of decreased light harvesting per unit chlorophyll imposed by optical packaging, slower C-uptake per RUBISCO places a further size-dependent burden upon phytoplankton nitrogen metabolism, thereby reinforcing the competitive disadvantage upon larger cells relative to smaller cells in acquiring the key resources required for photosynthesis. Our experimentally derived outcome for centric diatoms is consistent with observations of their distribution in the open ocean; specifically, larger centric diatom species appear restricted to deeper depths closer to the nutricline whereas smaller species are more abundant in the high light-low nutrient waters (Venrick, 2002). The size-dependent increase in the burden upon nitrogen metabolism could also explain why some larger diatoms form associations with N_2 -fixing bacteria in open ocean surface waters (Zehr et al., 2000; Foster et al., 2011). We therefore suggest that the combined burdens of not only light harvesting but also CO_2 acquisition upon nitrogen metabolism may place an effective upper limit on the cell size that can sustain growth of diatoms.

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