Diversity and Expression of RubisCO Genes in a Perennially Ice-Covered Antarctic Lake during the Polar Night Transition

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The autotrophic communities in the lakes of the McMurdo Dry Valleys, Antarctica, have generated interest since the early 1960s owing to low light transmission through the permanent ice covers, a strongly bimodal seasonal light cycle, constant cold water temperatures, and geographical isolation. Previous work has shown that autotrophic carbon fixation in these lakes provides an important source of organic matter to this polar desert. Lake Bonney has two lobes separated by a shallow sill and is one of several chemically stratified lakes in the dry valleys that support year-round biological activity. As part of an International Polar Year initiative, we monitored the diversity and abundance of major isoforms of RubisCO in Lake Bonney by using a combined sequencing and quantitative PCR approach during the transition from summer to polar winter. Form ID RubisCO genes related to a stramenopile, a haptophyte, and a cryptophyte were identified, while primers specific for form IA/B RubisCO detected a diverse autotrophic community of chlorophytes, cyanobacteria, and chemolithoautotrophic proteobacteria. Form ID RubisCO dominated phytoplankton communities in both lobes of the lake and closely matched depth profiles for photosynthesis and chlorophyll. Our results indicate a coupling between light availability, photosynthesis, and chlorophyll. The distinct water chemistries of the east and west lobes have resulted in depth- and lobe-dependent variability in RubisCO diversity, which plays a role in transcriptional activity of the key gene responsible for carbon fixation.

The McMurdo Dry Valleys form the largest ice-free region (∼4,000 km²) on the Antarctic continent (5). Owing to both the unique ecosystems they harbor and their sensitivity to environmental perturbation, the dry valleys are protected as an Antarctic Special Managed Area under environmental protection protocols of the Antarctic Treaty. Despite classification as a hyper-arid cold desert (with an average temperature of −17°C and precipitation equivalent to <10 mm water), the dry valleys contain a mosaic of hydrologically closed lake basins that act as microbial oases, providing a year-round liquid water habitat for phototrophic, chemoautotrophic, and heterotrophic microorganisms that are isolated from direct contact with the atmosphere by permanent 3- to 6-m-thick ice covers (38). The perennial ice cover prevents wind-driven turbulence, allowing strong vertical stratification of both chemical and biological characteristics to exist in the water column. Although the ice attenuates more than 95% of incident light and narrows the solar spectrum to blue-green wavelengths, the liquid water columns of these lakes support primary producers, dominated by low-light-adapted phytoplankton, which provide fixed carbon for the strictly microbial food web (30, 40). Microscopic and pigment analyses of samples collected during the spring and summer showed that the phytoplankton communities are strongly vertically stratified within the water columns in response to gradients in light and nutrients. Maximum productivity occurs close to the chemocline (39).

Lake Bonney is one of several lakes in the Taylor Valley, Antarctica, which have been studied since 1993 as part of the McMurdo Long Term Ecological Research Program (http://www.mcmater.org/). Dry valley lake aquatic systems support distinct microbial communities containing bacteria, algae, and flagellated and ciliated protozoans that interact to form truncated food webs dominated exclusively by microorganisms. Stratified phototrophic protist populations, including cryptophytes, chlorophytes, and chrysophytes, play key roles in primary productivity and carbon cycling in the lake food web (21, 27, 40). In one of the first phylogenetic studies conducted in Lake Bonney, Bielewicz et al. (2) characterized protist diversity through the stratified water columns of both lobes of Lake Bonney. The 18S rRNA libraries revealed that more than 85% of the sequences aligned with those of known phototrophic microbial eukaryotes. Shallow depths in the lake were dominated by a cryptophyte population related to Geminigera cryphila, mid-depths (13 to 15 m) harbored haptophytes and stramenopiles related to Isochrysis and Nannochloropsis, respectively, and the deepest photic zone was dominated by a variety of chlorophytes (2).

Ribulose-1,5-bisphosphate carboxylase/oxygenase (RubisCO) catalyzes the first step in CO₂ fixation. Four holoenzyme forms, I, II, III, and IV, are recognized in both phototrophic and chemotrophic organisms (47). Form I is the major CO₂ fixation enzyme found in plants, eukaryotic algae, cyanobacteria, and most phototrophic and chemolithoautotrophic proteobacteria (46). Most phytoplankton populations harbor one of four subclasses of form I RubisCO, including marine α-cyanobacteria (form IA), chlorophytes and β-cyanobacteria (form IB), and chromophytic algae such as diatoms and prymnesiophytes (form ID) (47). The
large subunit of Rubisco form I, encoded by the \(rbcL\) gene, is highly conserved and has been used as a phylogenetic tool in a variety of aquatic environments (e.g., see references 6, 11, and 15).

Logistical constraints have restricted most research in the McMurdo Dry Valleys to the austral spring, summer, and autumn months (10, 39), leaving us with little information on how the stratified microbial communities residing in the water columns of the lakes in this region respond to extended dark periods. We present data from the first study to examine responses of the dry valley lake autotrophic communities residing in Lake Bonney to the transition from full sunlight to the polar night. We focused specifically on the influence of the polar night transition on trends in primary production as well as on Rubisco dynamics through the photic zone.

**MATERIALS AND METHODS**

**Site description.** Lake Bonney lies immediately east of the Taylor Glacier, a major outlet glacier of the East Antarctic Ice Sheet. An ice-covered bedrock sill separates the lake into two basins, an east (ELB; 3.5 km²) and a west (WLB; 1.3 km²) lobe. The sill isolates saline, nutrient-rich deep waters while allowing the surface oxygenated waters (above 13 m in depth) to flow from WLB to ELB during the summer. The lake is fed at the west lobe by intermittent glacial meltwater at the terminus of the Taylor Glacier. An iron-rich subglacial brine pool called Blood Falls also flows seasonally into WLB (29). The lack of turbulent mixing leads to density-driven stratification of the water columns, producing stable gradients in temperature and conductivity. Approximately 1 to 5% of incident irradiance penetrates the ice cover, and the photic zones of both lobes extend from the bottom of the ice cover to the permanent nutricline (25). Both lobes exhibit unbalanced ratios of nitrogen to phosphorus: shallow phytoplankton populations are phosphorus limited (24, 39), while the phytoplankton residing near the chemocline depends upon upward nutrient diffusion from nutrient-rich deep waters (39). Lastly, long separation and differential evaporative histories between the two lobes have led to distinctive water chemistry differences in the isolated bottom waters. In the west lobe, oxygenated surface waters overlay anoxic layers with low nitrate and high ammonium levels, while the east lobe exhibits subsurface waters below the chemocline, with extremely high nitrate and nitrous oxide levels (50, 52).

**Field sampling.** Our study was conducted over one field season, from 17 February to 15 April 2008. A total of 28 water samples from ELB and 20 water samples from WLB were collected from late February to early April (sampling dates were 24 February, 2 March, 9 March, 16 March, 24 March, 29 March, and 10 April), a period representing the transition from 24 h of sunlight to complete darkness beneath the ice covers. Holes in the ice covers of the east and west lobes were drilled with a 10-cm-diameter ice auger and enlarged by melting with circulating heated ethylene glycol through a copper pipe. Sampling depths (6 m, 13 m, 18 m, and 20 m for ELB and 10 m, 13 m, 15 m, and 20 m for WLB) were measured from the piezometric water level in the ice hole (approximately 30 cm below the ice surface), using a depth-calibrated hand winch. Depths chosen coincided with chlorophyll \(a\) (Chl \(a\)) maxima reported in earlier studies (24) and spanned the photic zone (25). In addition, the locations of the Chl \(a\) maxima were confirmed on the day of sampling by performing a depth profile of Chl \(a\) in situ, using a bbe Moldaenke profiling spectrophotometer (1). At each time point, lake water was collected from both lobes at each sampling depth, using a 5-liter Niskin bottle sampler (General Oceancics, FL), and 1 liter was concentrated onto sterile 47-mm, 0.45-µm pore-size Durapore polycryliden fluoride membrane filters (Millipore, MA), using a vacuum of 0.3 atm. The filters were then frozen immediately in liquid nitrogen before being transported to McMurdo Station, where they were shipped on dry ice to U.S. laboratories and stored at −80°C until nucleic acid extraction.

**Limnological variables.** Conductivity, temperature, photosynthetically available radiation (PAR), light-mediated primary productivity (PPR), and Chl \(a\) were collected and measured using previously described methods (37). Briefly, temperature and conductivity were measured with a Seabird model 25 profiler as described by Spigel and Priscu (43). Depth profiles for in situ PAR were measured with a Li-Cor LI-193 spherical quantum sensor (Li-Cor Biosciences, NE). PAR levels were recorded manually at 0.5-m intervals from the piezometric water level to approximately 5 m below the photic zone (depth of 25 m). Levels of all inorganic nitrogen species were determined with a Lachat autoanalyzer, using methods described by Priscu (37). Owing to the low levels of soluble reactive phosphorus (SRP) in Lake Bonney, SRP was analyzed manually using the antimony-molybdate method (45) with a 10-cm-path-length cuvette. In situ Chl \(a\) was determined with a bbe Moldaenke profiling spectrophotometer (1). This instrument was lowered at a rate that produced −10 measurements m⁻¹, which was adequate to define the highly layered phytoplankton species. Vertical profiles of primary productivity were made by measuring \(^{14}C\)bicarbonate incorporation into particulate matter over a 24-h in situ incubation of 2 light bottles and 1 dark bottle at selected depths. The radioactivity in the dark bottles was subtracted from that in the light bottles to correct for nonphotosynthetic incorporation. Following incubation, samples were filtered onto Whatman GF/F filters, acidified with 0.5 ml 3 N HCl, air dried at 50°C, and counted using standard scintillation spectrometry. The concentration of dissolved inorganic carbon required for productivity rate calculations was determined by passing the gas from acid-sparged samples through a calibrated infrared gas analyzer. Depth-integrated estimates for PPR and Chl \(a\) in shallow (4 to 8 m), middle (10 to 15 m), and deep (18 to 20 m) populations residing within the photic zone were calculated as the sums of the measurements at sampling depths of 4 to 8 m, 10 to 15 m, and 18 to 20 m, respectively, according to the method of Lizotte et al. (27).

**Nucleic acid isolation.** Frozen filters were cut in half to allow for extraction of environmental RNA and DNA from the same filter. For environmental DNA, half of a filter was cut into small pieces and DNA was isolated using a FastDNA spin kit for soil (MP Biomedicals, OH) following the manufacturer’s protocol. Total RNA was isolated from the other half of the filter by use of a combination of RNeasy miniprep (Qiagen, CA) and FastRNA Pro Soil-Direct (MP Biomedicals, OH) kits (see the supplemental material for more details), according to the method of Kong and Nakatsu (17). For reverse transcription, total RNA ranging from 5 to 30 ng was reverse transcribed to single-stranded cDNA by use of an iScript cDNA synthesis kit (Bio-Rad, CA) as specified by the manufacturer. Negative controls (replacing RNA with water in the reaction mixture) were included in all experiments.

**Real-time qPCR.** Quantification of the Rubisco gene copy number and transcript level was performed using real-time quantitative PCR (qPCR) with primer sets that target different forms of Rubisco genes (see the supplemental material for more details). The primer sets targeting form IA/B and form ID Rubisco genes were developed as previously described (33). Form ID and IA/B Rubisco clades represent the dominant isoforms in marine and freshwater systems (33, 35, 54). The primer set GemF2 plus GemR2 was designed for this study to target the cryptocyste Gemmiger a cryophila, possessing the form ID Rubisco gene. This organism was previously found in 185 rRNA clone libraries (2) but could not be detected in this study by use of the form ID primer set. The species-specific primer set GemF2 plus GemR2 (GemF2, GCGTTTCTATCTGGTATGGA; and GemR2, GGGCCACGTAGTAATCCACCT) was designed using the program Primer3, based on the rbcL sequence of *G. cryophila* (GenBank accession no. AB164411).

qPCR was performed according to a previously described method (2), using a 25-µl reaction mixture containing 1 µl of cDNA or DNA (10 ng), 1.5 µl of each primer (10 pmol µl⁻¹), and 12.5 µl iQ SYBR Green Supermix (Bio-Rad, CA). Amplification conditions were 5 min at 95°C followed by 35 cycles of 1 min at 94°C, 20 s at 52°C, and 30 s at 72°C, and fluorescence intensity was acquired at 83°C (above the melting point of primer dimers). To determine the melting temperature and PCR product specificity, a melting curve was acquired by heating from 50°C to 95°C.
Phytoplankton residing in the shallow and mid-depths responded to the decline in PAR during the summer-winter transition by a reduction in PPR (Fig. 2C to F). Photosynthetic rates in the deepest layer were very low in both lobes throughout the sampling period. Deep populations in ELB exhibited a rapid decline in PPR in early January, while WLB deep populations exhibited low photosynthetic rates throughout the polar night transition (Fig. 2G and H). In contrast with phytoplankton photosynthesis, Chl a levels generally increased during the summer-winter transition (Fig. 2C to F). However, while shallow populations exhibited a steady rise in Chl a levels throughout the sampling period, Chl a stabilized at mid-depths by mid-March (Fig. 2E and F) and exhibited a decline in deep populations beginning in mid-February (Fig. 2G and H). To determine whether PPR or Chl a covaried with declining PAR, Pearson correlation coefficients were determined at four sampling depths throughout the photic zone between sampling dates 24 February and 10 April (Table 1). Depths at or above the chemocline (i.e., 6 m and 13 m in ELB and 10 m and 13 m in WLB) exhibited a positive correlation between PPR and PAR in both lobes ($r = 0.82$ to 0.99; $P < 0.05$) (Table 1) during the polar night transition. In contrast to the case for photosynthetic rates, shallow populations from both lobes exhibited a negative correlation between PAR and Chl a ($r = -0.792; P < 0.1$) (Table 1).

**RubisCO diversity.** To gain a better understanding of autotrophic gene diversity in Lake Bonney, clone libraries were generated for the RubisCO large-subunit gene, using DNA and mRNA (reverse transcribed to cDNA) as templates. A total of 4 phylotypes were detected in the form ID DNA and cDNA clone libraries (Fig. 3). The vast majority of form ID $rbcL$ sequences were

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**RESULTS**

**General lake characteristics.** Physical and chemical profiles in Lake Bonney revealed a highly stratified vertical structure of the water column within each lobe (Fig. 1). The water temperature reached a maximum in the middle of the water column and ranged from $-2.1$ to $4.9^\circ C$ in ELB and from $-4.3$ to $2.2^\circ C$ in WLB (Fig. 1A). ELB and WLB both have steep mid-depth conductivity gradients, with values ranging from 0.36 to 109.13 mS cm$^{-1}$ and from 0.74 to 79.60 mS cm$^{-1}$, respectively (Fig. 1B). Both lobes showed distinct light-dependent PPR and Chl a maxima at 13.5 m, or just above the chemocline (Fig. 1C and D). Maximum PPR was 4- to 5-fold higher in ELB (Fig. 1C), while the Chl a maximum was 2-fold higher in WLB than in ELB (Fig. 1D).

**Effect of polar night transition on photosynthesis and phytoplankton biomass.** We monitored responses of the phototrophic community through the photic zone during the summer-winter transition. Beginning in mid-February, daily under-ice irradiance declined rapidly in both lobes of Lake Bonney (Fig. 2A and B). Depth-integrated Chl a concentrations and photosynthetic rates were calculated for the three major phytoplankton assemblages, i.e., shallow, mid-depth, and deep populations.
related to a stramenopile, *Nannochloropsis* (GenBank accession no. DQ977734; 97% similarity), and a haptophyte, *Isochrysis* sp. (GenBank accession no. AB043693 and AY119783; 91% similarity). Using cryptophyte-specific *rbcL* primers, we also detected a single cryptophyte, which was related to *Geminigera cryophila* (GenBank accession no. AB164411; 99% similarity). The dominant form ID phylotypes were detected in both lobes of Lake Bonney and were present in both DNA and cDNA libraries; however, haptophyte sequences dominated cDNA libraries in both lobes (Fig. 3; also see Fig. S1 in the supplemental material).

A total of 15 phylotypes were detected in the form IA/B DNA and cDNA clone libraries, related to cyanobacteria, chlorophytes, and chemoautotrophic proteobacteria (see Fig. S2 to S4 in the supplemental material). More than 95% of cyanobacterial sequences were found in the DNA clone libraries only and were related to *Nostocales* and *Oscillatoriales* (see Fig. S2 in the supplemental material). *Nostocales* sequences were restricted to WLB, while *Oscillatoriales* sequences were detected throughout the water columns of both lobes. While cyanobacterial *rbcL* genes were rare in the cDNA clone libraries, chlorophyte *rbcL* genes related to those in *Micractinium* and *Chlorella* spp. (GenBank accession no. EF113451 and AB260909; 94% similarity), as well as the Antarctic chlorophytes *Chloromonas* and *Chlamydomonas* spp. (GenBank accession no. U80809 and AY731086; 90 and 99% similarity, respectively), dominated form IA/B RubisCO cDNA clone libraries (see Fig. S3 and S5 in the supplemental material). Lastly, form IA/B sequences related to chemoautotrophic proteobacteria (an endosymbiont of *Oligobrachia haakonmosbi* and *Thiobacillus* [GenBank accession no. AB057772 and AY914807; 83 and 87% similarity, respectively]) were recovered from DNA libraries from WLB samples located below the chemocline. No chemoautotroph sequences were recovered from ELB clone libraries (see Fig. S4 and S5 in the supplemental material).

**Depth- and lobe-specific RubisCO trends.** Given the current lack of data on abundance or expression levels of autotrophic genes in dry valley environments, we used qPCR to measure ver-
TABLE 1 Polar night transition-dependent Pearson correlations for PPR, Chl a, rbcL gene copy number (DNA), and rbcL transcript level (mRNA) versus PAR

<table>
<thead>
<tr>
<th>Sampling depth (m)</th>
<th>PPR</th>
<th>Chl a</th>
<th>rbcL DNA</th>
<th>Form ID</th>
<th>Form IA/B</th>
<th>rbcL mRNA</th>
<th>Form ID</th>
<th>Form IA/B</th>
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<tr>
<td>6</td>
<td>0.934**</td>
<td>−0.792*</td>
<td>−0.219</td>
<td>−0.797*</td>
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<tr>
<td>13</td>
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<td>−0.276</td>
<td>0.808*</td>
<td>−0.375</td>
<td>0.754*</td>
<td>0.844**</td>
<td></td>
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<tr>
<td>18</td>
<td>0.994**</td>
<td>0.386</td>
<td>0.815*</td>
<td>0.843**</td>
<td>0.554</td>
<td>0.737*</td>
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<tr>
<td>20</td>
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<td>0.939*</td>
<td>0.604</td>
<td>0.586</td>
<td>0.731</td>
<td>−0.700</td>
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<tr>
<td>WLB sampling depths</td>
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<tr>
<td>10</td>
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<td>−0.792*</td>
<td>0.468</td>
<td>0.517</td>
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<tr>
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<td>−0.313</td>
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<tr>
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<td>0.916**</td>
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<td>−0.809</td>
<td>−0.030</td>
<td>−0.038</td>
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</table>

*Samples were collected between 24 February and 10 April 2008. All data were log transformed prior to analysis. Sampling depths were measured from the piezometric water level within the sampling hole. *, significant correlation (P < 0.1); **, significant correlation (P < 0.05).

Rubisco gene dynamics during the polar night. Seasonal dynamics of Rubisco during the polar night transition were determined throughout the photic zones of both lobes of Lake Bonney (Fig. 5). Form ID Rubisco dominated the autotrophic communities of both lobes throughout the summer-winter transition, at the levels of both transcript abundance and copy number. In ELB, the abundances of form ID mRNA and DNA in samples from 13 m, as well as form ID DNA in samples from 18 m, declined during the polar night (Fig. 5A and C) and were positively correlated with PAR (r = 0.81 to 8.85; P < 0.1) (Table 1). In contrast, samples from both 13 m and 15 m from WLB exhibited an initial 7-fold decline in form ID mRNA abundance, followed by a 3- to 5-fold increase in form ID mRNA levels at a later (8 March) sampling time (Fig. 5E and G). No positive correlation (P < 0.1) between form ID mRNA and PAR was observed in samples collected from WLB (Table 1). Haptophyte sequences dominated the cDNA clone libraries at mid-depth (13 m) for both lobes at early (24 February) and late (29 March) time points (see Fig. S1 in the supplemental material). In contrast to the case for samples from 13 m, haptophytes replaced stramenopiles during the summer-winter transition at a shallow (6 m) depth in ELB, while the opposite trend was observed for a shallow (10 m) depth in WLB (see Fig. S1 in the supplemental material).

Form IA/B mRNA and DNA levels exhibited trends that were both depth and lobe dependent. In ELB, form IA/B exhibited a positive correlation between PAR levels and mRNA abundance at 13 m and both mRNA and DNA abundances in 18-m populations (r = 0.74 to 0.85; P < 0.1) (Fig. 5B and D; Table 1). However, samples from 6 m exhibited a 7-fold rise in form IA/B DNA during the polar night transition, as well as a negative correlation with PAR (r = −0.80; P < 0.1) (Table 1). Form IA/B mRNA and DNA levels were generally lower in the west lobe (Fig. 5F and H). Furthermore, no significant trends between form IA/B abundance and PAR levels were observed in WLB, and form IA/B mRNA and DNA levels increased in most sampling depths by 1.6- to 3.9-fold during the polar night transition (Table 1; Fig. 5F and H). Form IA/B DNA clone libraries (generated from samples taken on 2 March) revealed a mixed population of cyanobacteria, chlorophytes, and chemotrophs in WLB. However, ELB form IA/B clone libraries harbored a less diverse population of cyanobacteria and chlorophytes, and form IA/B cDNA libraries from ELB were dominated almost exclusively by chlorophyte mRNAs (see Fig. S5 in the supplemental material).

**DISCUSSION**

Permanently ice-covered Antarctic lakes harbor few metazoans, do not mix seasonally, have low allochthonous input, and are exposed to minimal direct human impact. As such, the vast majority of the organic carbon that supports this ecosystem is provided by carbon fixation by autotrophic microorganisms. Past studies have shown that phytoplankton productivity in the short austral summer is limited by under-ice PAR levels (10, 12, 23) as well as by nutrient availability (39). Our study extends this earlier work and shows that Lake Bonney phytoplankton populations exhibit distinct seasonal and spatial trends in photosynthesis and Rubisco gene dynamics.
In high-latitude aquatic habitats such as dry valley lakes, seasonal transitions during winter-spring and autumn-winter are accompanied by extreme changes in day length and incident irradiance. Phytoplankton residing in either lobe of Lake Bonney responded to the seasonal loss in PAR during the summer-winter transition by an overall reduction in photosynthesis throughout the photic zone. This trend was most pronounced in middle and deep populations, which experienced lower PAR levels earlier in the summer-winter transition than those of phytoplankton residing in the shallow layers (Fig. 2). Conversely, Chl $a$ in the shallow populations increased throughout our sampling period, while Chl $a$ in the deeper populations remained unchanged (mid-depths) or

**FIG 3** Neighbor-joining phylogenetic tree of representative form ID *rbcL* sequences (615 bp) retrieved from environmental DNA (blue) and mRNA (red) from Lake Bonney. Bar, 0.02 substitution per nucleotide position. The GenBank accession number is listed after each sequence name. Clones were named as follows: time point with lake name and depth-primer name-template type-index number.
declined (deep layer). These data show that sustained light-dependent CO₂ fixation in shallow and middle populations supported phytoplankton biomass accumulation later into the summer-winter transition than that of the deep chlorophyll layer in both lobes of Lake Bonney. Similarly, during the early spring, phytoplankton biomass develops sequentially from the shallow to the deep layers of the photic zone in response to the seasonal increase in PAR (27).

While a recent study provided molecular evidence for vertical stratification and seasonal variability in the Lake Bonney protist community (2), little is known of the relative contributions of specific phytoplankton groups to primary productivity in the dry valley lake food web. In agreement with previous reports (2, 19, 26, 41), our functional analyses of a key photosynthetic gene confirmed that a stratified autotrophic community dominated by photosynthetically active protists occupies the photic zone of Lake Bonney. First, a population of stramenopiles related to *Nannochloropsis* was abundant throughout the water columns of both lobes. Results from cDNA clone libraries from this study indicate that *Nannochloropsis* is an active member of the primary producer population (see Fig. S6 in the supplemental material). *Nannochloropsis*-dominated algal blooms have been detected in other aquatic systems during cold-water periods (7, 8, 18), implying that they are adapted to low temperatures. Second, a cryptophyte species identified as *Geminigera cryophila* was identified as an active member in the shallow layers (see Fig. S6 in the supplemental material). Finally, a haptophyte related to an *Isochrysis* sp. made up 95 to 100% of sequences recovered from cDNA libraries generated from 13-m samples for both lobes (see Fig. S1 in the supplemental material). *Isochrysis* populations also occur at the same depth as high concentrations of dimethylsulfide in Lake Bonney, and they may produce biogenic sulfur compounds as cryoprotectants (20).
While regulation of RubisCO enzyme activity is complex, most phototrophic organisms actively transcribe the gene encoding the large subunit, and the \textit{rbcL} gene copy number, transcript abundance, and phylogenetic diversity have been applied as molecular indicators of carbon fixation potential in aquatic environments \cite{13-15, 28, 32, 53, 54}. Phototrophs harboring form ID RubisCO dominate the photic zone of Lake Bonney, with maximum transcript levels and gene copy numbers occurring at mid-depth (15 m) in both lobes. Vertical trends in form ID abundance matched those of PPR and phytoplankton biomass (this study), as well as 18S rRNA transcript abundance \cite{2}, suggesting that phytoplankton populations harboring form ID RubisCO are key primary producers in the dry valley lake food web. Phytoplankton populations residing in Lake Bonney balance light availability with a sufficient supply of nutrients by residing at depths directly above the chemocline \cite{22, 27, 39}. Furthermore, we observed a strong correlation between declining PAR and both form ID gene copy number and mRNA level in this phytoplankton population (Table 1). Taken together, these data imply that at the depth of maximum productivity, there is a coupling between PAR, photosynthetic rates, and transcriptional activity of the major isofrom of RubisCO.

While phytoplankton populations residing in Lake Bonney are always light limited \cite{24}, productivity in the surface waters is under extreme nutrient deprivation for most of the austral summer \cite{39}. Nutrient deficiency develops under the ice in early spring and persists throughout the summer, until either nutrient pools are recharged in the winter or a transient influx of glacial meltwater introduces glacial flour particles to the water column \cite{25}. In support of earlier reports that observed lower photosynthetic productivity and photosynthetic efficiency within surface layers of Lake Bonney \cite{22, 27, 39}, \textit{rbcL} DNA and mRNA levels measured in shallow samples from either lobe were >100-fold lower than those of mid-depth populations (Fig. 4). Despite a strong relationship between light availability and photosynthetic rates in surface populations, no positive correlation was observed between PAR and RubisCO abundance in the shallow layers of Lake Bonney (Table 1). This lack of a relationship between PAR and RubisCO suggests that additional factors are likely to influence \textit{rbcL} expression in the phytoplankton residing in the shallow layers of Lake Bonney. A recent study on diel responses in natural phytoplankton populations from the Mississippi and Orinoco River plumes found that daily patterns of \textit{rbcL} transcript abundance were not only influenced by the diurnal light cycle but also impacted by nutrient deficiency \cite{13}. We suggest that nutrient deficiency may have played a role in the differential seasonal patterns in \textit{rbcL} expression observed between the shallow and deeper phytoplankton populations. Nutrient stress may also influence coupling between RubisCO transcription and carbon fixation capacity in the shallow populations, which has been observed in other nutrient-deficient environments \cite{13}.

It should be noted that RubisCO is regulated not only transcriptionally but also at the level of enzyme activity \cite{31}. Studies have suggested that dissipation of excess energy products via photorespiration is an environmental stress response to nutrient deficiency in some phytoplankton communities \cite{13, 16}. Thus, regulation of enzyme activity could also influence carbon fixation rates during the polar night transition, particularly in nutrient-stressed shallow populations. Furthermore, while the application of qPCR to environmental samples has been adopted widely as a proxy for estimating the distribution and activity of microbial populations (e.g., see references \textit{4, 9, 32, 44}, and \textit{49}), there are caveats that should be considered in applying qPCR approaches to environmental samples. These considerations include variability in gene copy and genome numbers between organisms, as well as differences in nucleic acid extraction efficiencies and mRNA stability (reviewed in reference \textit{42}).

Both lobes of Lake Bonney support distinct subsets of metabolically active autotrophic communities that carry either form ID or IA/B RubisCO. The strong relationship between PAR, rates of primary production, and transcript levels of form ID RubisCO indicates that transcriptional regulation of this RubisCO isofrom could be important for controlling rates of autotrophic carbon fixation. However, variability in vertical and temporal trends for both isoforms within and between the lobes suggests that light availability is not the sole factor influencing RubisCO dynamics in Lake Bonney autotrophic populations. Studies on other aquatic systems have shown that phytoplankton community composition contributes to (i) light-dependent patterns in \textit{rbcL} transcript levels \cite{13, 15, 34}, (ii) the coupling between RubisCO gene expression and rates of photosynthesis \cite{13}, and (iii) the coupling between RubisCO activity and carbon fixation potential \cite{16}. Differences in autotrophic community structure and activity in the two lobes presumably reflect the climate evolution of the two lobes of the lake. Using helium isotope data, Poreda et al. \cite{36} showed that these lake basins were isolated and possibly ice-free for thousands of years, until a major climate shift led to an overflow of water from the west lobe to the east lobe \cite{43}. This climate shift produced an important tipping point in the lake ecosystem, in terms of the development of a permanent ice cover and mixing of near-surface waters between the lobes. As a result of this evolution in physical and chemical character, biodiversity would have been altered significantly, culminating in the vertical structure that we observed in our present study. As such, climate must have played a key role in the differences in RubisCO diversity and regulation measured in our study. As the climate of Antarctica warms \cite{51}, we can expect to see new tipping points that may lead to further changes in dry valley lake food web structure. Thus, while the transition to polar night represents one of the most extreme seasonal changes experienced by the autotrophic organisms in Lake Bonney, the legacy of evolution of lake chemistry over the past 6,000 years may exert long-term selection pressures on the autotrophic biology of Lake Bonney, and presumably other lakes within the McMurdo Dry Valleys.

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