Modulation of the Major Paths of Carbon in Photorespiratory Mutants of \textit{Synechocystis}

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Abstract

\textbf{Background:} Recent studies using transcript and metabolite profiles of wild-type and gene deletion mutants revealed that photorespiratory pathways are essential for the growth of \textit{Synechocystis} sp. PCC 6803 under atmospheric conditions. Pool size changes of primary metabolites, such as glycine and glycolate, indicated a link to photorespiration.

\textbf{Methodology/Principal Findings:} The $^{13}$C labelling kinetics of primary metabolites were analysed in photoautotrophically grown cultures of \textit{Synechocystis} sp. PCC 6803 by gas chromatography-mass spectrometry (GC-MS) to demonstrate the link with photorespiration. Cells pre-acclimated to high CO$_2$ (5%, HC) or limited CO$_2$ (0.035%, LC) conditions were pulse-labelled under very high (2% w/w) $^{13}$C-NaHCO$_3$ (VHC) conditions followed by treatment with ambient $^{13}$C at HC and LC conditions, respectively. The $^{13}$C enrichment, relative changes in pool size, and $^{13}$C flux of selected metabolites were evaluated. We demonstrate two major paths of CO$_2$ assimilation via Rubisco in \textit{Synechocystis}, i.e., from 3PGA via PEP to aspartate, malate and citrate or, to a lesser extent, from 3PGA via glucose-6-phosphate to sucrose. The results reveal evidence of carbon channelling from 3PGA to the PEP pool. Furthermore, $^{13}$C labelling of glycolate was observed under conditions thought to suppress photorespiration. Using the glycolate-accumulating \textit{AglcD1} mutant, we demonstrate enhanced $^{13}$C partitioning into the glycolate pool under conditions favouring photorespiration and enhanced $^{13}$C partitioning into the glycine pool of the glycine-accumulating \textit{AgcvT} mutant. Under LC conditions, the photorespiratory mutants \textit{AglcD1} and \textit{AgcvT} showed enhanced activity of the additional carbon-fixing PEP carboxylase pathway.

\textbf{Conclusions/Significance:} With our approach of non-steady-state $^{13}$C labelling and analysis of metabolite pool sizes with respective $^{13}$C enrichments, we identify the use and modulation of major pathways of carbon assimilation in \textit{Synechocystis} in the presence of high and low inorganic carbon supplies.

Introduction

Cyanobacteria are considered the first organisms to have evolved the capacity for oxygenic photosynthesis around three billion years ago [1]. The endosymbiotic uptake of an ancient cyanobacterial ancestor by a eukaryotic cell initiated the evolution of phototrophic algae and plants. Many of the initial cyanobacterial proteins are still detectable within the chloroplasts and nuclear genomes of current higher plants [2,3]. In both cyanobacteria and C3 plants, CO$_2$ fixation is primarily catalysed by the enzyme ribulose-1,5-bisphosphate carboxylase/oxygenase (Rubisco). The carboxylation reaction generates two molecules of 3-phosphoglycerate (3PGA) from ribulose-1,5-bisphosphate and CO$_2$, whereas O$_2$ competition at the reaction centre leads to the oxygenase products 3PGA and 2-phosphoglycerate (2PG) [4]. The product 2PG is a cellular toxin that needs to be detoxified, as it inhibits Calvin-Cycle enzymes [5–7]. In plants, 2PG is scavenged by a sequence of reactions called the photorespiratory C2 pathway [8–10], which regenerates one molecule of 3PGA for every two molecules of 2PG, at the cost of CO$_2$ and NH$_4^+$ release.

In contrast to higher plants, 2PG metabolism was thought to exert negligible effects in cyanobacteria. Early studies only indicated the formation of glycolate from 2PG [11]. Furthermore, the discovery of a sophisticated inorganic carbon (Ci) concentrating mechanism (CCM) demonstrated the potential of cyanobacteria to increase the internal concentration of CO$_2$ in the vicinity of Rubisco and thus to compensate for the low CO$_2$ affinity of the cyanobacterial enzyme [12]. As a consequence, the CCM was thought to be sufficient to suppress the oxygenase reaction and to make photorespiratory detoxification irrelevant for cyanobacterial metabolism.

Recent studies, however, demonstrated not only that the CCM is insufficient to prevent ribulose-1,5-bisphosphate oxygenation in an O$_2$-containing atmosphere but also that there is active 2PG
metabolism. The photorespiratory pathways were found to be essential for growth under atmospheric conditions [13,14]. Photorespiratory 2PG metabolism in the Synechocystis sp. strain PCC 6803 (hereafter Synechocystis) comprises three alternative paths: a plant-like C2 cycle, a bacterial glycerate pathway, and the complete decarboxylation of glyoxylate via oxalate. Mutants defective in specific enzymes within these pathways including glycolate dehydrogenase (DglcD1, sll0404) and glycine decarboxylase (DgcV, sll0171) displayed growth retardation. In addition, intermediates of photorespiratory 2PG metabolism accumulated in agreement with the enzymatic defect; these intermediates included glycolate in DglcD1 and glycine in DgcV. These phenomena were already apparent under elevated CO2 conditions (5%, high CO2, HC), and were enhanced under the low-CO2-availability conditions of ambient air (~0.035%, low CO2, LC) [14,15]. The Ci affinity and the maximum rate of photosynthesis were found to be higher in cells grown under LC conditions than in cells grown under HC conditions [14].

Transcript and metabolite profiles of the wild-type (WT) strain compared to the above-mentioned mutants indicated that the photorespiratory mutants, when grown under HC conditions, display gene expression and metabolite patterns that are characteristic for WT cells when shifted from HC to LC conditions [15,16]. Characterisation of changes of metabolite pools opened the path towards currently unanswered questions, including: (1) Is the previously observed accumulation of intermediates, such as glycolate and glycine, indeed linked to photorespiration? (2) How is the ratio of Rubisco oxygenation to carboxylation influenced by altered Ci availability? (3) Which routes of carbon fixation are used under HC or LC conditions? To approach these questions, we applied a non-steady state 13C pulse and chase experimental design similar to work previously published by our lab [17]. The appearance and dilution of 13C labelled intermediates within the primary metabolism of photoautotrophically grown WT and selected mutant cells was monitored following the suggestions made for dynamic flux analyses [18]. With this approach, we reveal the use and modulation of the major paths of carbon assimilation in Synechocystis.

Results

Experimental design of metabolic flux analysis

Pulse labelling of photoautotrophic Synechocystis cultures was performed by adding aliquots of a saturated solution of 13C labelled NaHCO3 to a final concentration of 2% (w/w). This procedure resulted in a very high carbon (VHC) pulse and was chosen to ensure a step change with the highest 13C enrichment possible. Moreover, the VHC conditions should suppress the oxygenase activity of Rubisco. We combined the 13Ci pulse of stably labelled bicarbonate with a chase using unlabelled CO2. Our experimental procedure generated an optimal rectangular step change during the 13Ci pulse and sufficient enrichment for short isotope dilution times of 10–60 min (Figure 1). In the following study, we focused on the metabolite pools that reached high 13C enrichment and thus allowed optimal GC-MS-based analysis. The VHC pulse was applied to cells that pre-acclimated to 5% CO2 (HC) or to 0.035% CO2 (LC) conditions, as previously reported [15]. For the chase, a quick medium exchange was
performed. The cells were subsequently incubated under continuous aeration with either HC or LC identical to the initial pre-acclimation. Our previous work on metabolic profiling revealed highly reproducible metabolic patterns after HC or LC acclimation and indicated that near-steady-state conditions could be achieved using standardized batch cultivation [15]. To analyse changes in carbon partitioning within the primary metabolism, we combined metabolic flux and pool size assessments as previously suggested [17]. The metabolic data were extended by parameters derived from mass isotopomer distribution analyses, i.e., the initial rate of $^{13}$C accumulation during the first 10 min and the maximum $^{13}$C enrichment at 20–60 min. Both parameters were calculated from the transient labelling kinetics of each of the monitored metabolite pools. Based on the analysis of 3PGA as the main entry point of the $^{13}$C label in photoautotrophic cultures of Synechocystis, we estimated the general features of the chosen experimental design. The 3PGA levels did not change after pre-acclimation to HC and LC or throughout the VHC pulse but increased clearly upon initiation of both the HC and LC chase (Figure 1A). As both chase conditions represented a shift to lower $C_i$ availability, these observations were in agreement with the previously observed increase of the 3PGA pool 3 h after a shift from HC to LC [15]. In parallel, the transient $^{13}$C labelling of the 3PGA pool was assessed during pulse and chase. Under our labelling conditions, the 3PGA pool reached its maximum saturated $^{13}$C enrichment 10 min after the pulse in both the HC and LC acclimated cells. LC cells showed a more rapid $^{13}$C accumulation. During the chase period with HC conditions, labelling returned below 10% within 30 min. Under LC conditions, a delayed chase response was observed. As expected, the empirical mass distribution vectors of 3PGA and PEP indicated homogenous $^{13}$C labelling. The kinetic behaviour of the mass distribution vectors (Figure S1) was generally in agreement with the modelled predictions reported earlier [19].

The presence of unlabelled ambient CO$_2$ resulted in a perceptible isotope dilution compared to the 98% enrichment of the applied NaHCO$_3$. For example, the final $^{13}$C enrichment in the 3PGA pool was on average $>82\%$ and $>89\%$ for the HC and LC conditions, respectively (Table S1). Moreover, the label exchange for the chase phase indicated a step change only for the HC condition. Under LC conditions, the label exchange was clearly delayed (Figure 1). Dilution and carryover effects at chase initiation were unavoidable in our hands, especially when implementing the LC chase conditions. Therefore, the slow return of $^{13}$C label in 3PGA could be caused by both an insufficient physical dilution of $^{13}$C label and by a physiological effect of the CCM, which is activated under HC conditions but suppressed under HC conditions [12]. To account for these effects, we used the enrichment data of the first assimilation products, for example 3PGA, to correct the influence of ambient CO$_2$ and any potential label carryover.

In this study, we analysed the pool size and labelling kinetics as $^{13}$C enrichment parameters of the observed metabolite pools. Each of these parameters will first be reported in context of the HC versus LC pre-acclimation for the WT strain. Subsequently, the respective phenotypes of the mutant strains will be presented. We focused on the mutants $AglD1$ ($Dl0404$), which is deficient for one of the two isoforms of glycolate dehydrogenase, and $AgeT$ ($Dl0171$), deficient for the T-protein subunit of the glycolate decarboxylase complex.

The entry points of the $^{13}$CO$_2$ label: 3PGA and PEP

The two most rapidly and completely labelled metabolite pools were the 3PGA pool, as expected, and, unexpectedly, the PEP pool (Figure 1). The $^{13}$C enrichment was higher and the rate of $^{13}$C labelling was faster in the PEP pool compared to the 3PGA pool under all conditions and in all analysed mutants. The Student’s t-test of the differences in $^{13}$C enrichment at 20–60 min after the pulse and in the initial $^{13}$C accumulation rate at 0.5–10.0 min showed that these differences were highly significant in both cases, $p < 0.00009$ and $p = 0.050$, respectively. These results are documented in more detail by the PEP/3PGA ratios of both measurements, which were consistently >1.0 (Table 1). The kinetic analysis of the PEP/3PGA ratio of $^{13}$C enrichment (Figure 2) demonstrated that the PEP pool was labelled faster in the pulse phase and was also de-labelled faster in the chase phases. This observation coincided with a general increase in both pool sizes upon shift from VHC during the pulse phase to HC or LC during the chase phase (Figure 1). The 3PGA pool increased by more than the PEP pool, as demonstrated by a reduction of the PEP/3PGA pool size ratio from above 1 to about 0.15 (Figure 2).

Cells pre-acclimated to the HC or LC conditions exhibited several characteristic differences in 3PGA and PEP labelling, but the initial rate of $^{13}$C accumulation and the maximum $^{13}$C enrichment of both metabolites remained strictly correlated during the pulse phase. Firstly, under LC conditions, the initial rates of $^{13}$C accumulation in both metabolite pools were faster by a factor of $\sim 1.3$ as compared to HC conditions. Also the final $^{13}$C enrichment was $\sim 1.1$-fold higher (cf. Table 1) with LC conditions. Secondly, the chase phase revealed a substantial difference. While metabolites in cells from both conditions behaved according to the assumption of a rectangular step change after the pulse, only metabolites under the HC chase conditions appeared to follow this assumption after the chase. In contrast, the LC cultures retained high $^{13}$C enrichment during the first 20 min of the chase until the onset of a delayed and slow de-labelling. This irregular behaviour probably resulted either from the physical carryover effect of $^{13}$C label from the preceding VHC pulse, as reported above, or from the intracellular CCM activity of Synechocystis, which is highly active under LC conditions [12] and should favour the re-assimilation of the $^{13}$CO$_2$ generated from internal sources.

The 3PGA and PEP labelling of the glycine decarboxylase-deficient $AgeT$ mutant strain was similar to that of the WT strain. In contrast, the glycolate dehydrogenase mutant, $AglD1$, showed large differences in 3PGA labelling but only slight changes in PEP $^{13}$C enrichment kinetics. Specifically, the rate of $^{13}$C accumulation and the final $^{13}$C enrichment in the 3PGA pool were lowest in the $AglD1$ mutant (Table 1). This observation could be explained by the toxic effects of mutant-specific glycolate accumulation on Calvin-Benson cycle activities. These findings were in agreement with previous observations of reduced photosynthetic and growth rates for glycolate dehydrogenase-deficient mutants [14].

Monitoring the flux into the glucose-6-phosphate and sucrose pools

To analyse the downstream fate of the initial labelling products, $^{13}$C partitioning into the glucose, glyceraldehyde-6-phosphate (G6P) and sucrose pools was monitored. The maximum $^{13}$C enrichment of the glucose pool was unchanged, i.e. 16% for LC ($n = 2$) and 13% for HC ($n = 2$). In contrast, the pools of both G6P and sucrose were rapidly labelled during the pulse and de-labelled during the chase. The maximum $^{13}$C enrichment of G6P was $82\%$ for LC ($n = 3$) and $76\%$ for HC ($n = 3$). The maximum $^{13}$C enrichment of sucrose was slightly lower at $68\%$ for LC ($n = 3$) and 65% for HC ($n = 3$). Sucrose labelling was slightly delayed compared to G6P after the pulse, and sucrose de-labelling was consistently more rapid under the LC chase conditions (Figure S2). The differences between G6P and sucrose
were less obvious under the HC chase conditions. Taken together these observations indicated a carbon flux from 3PGA via G6P towards sucrose synthesis when *Synechocystis* was exposed to excess Ci during the VHC pulse conditions. Furthermore, sucrose was mobilised when the cells were shifted back to lower carbon conditions. The sucrose flux appeared to represent one of the primary paths of photosynthetic CO₂ assimilation under our conditions. The sucrose flux results (Table 2) were in agreement with earlier findings [14], that the acclimation of *Synechocystis* to the LC condition leads to a higher rate of photosynthesis and sucrose flux than acclimation to HC.

Monitoring photorespiratory flux: Glycolate, glycine and serine

In our study, glycolate was the first photorespiratory intermediate in the pathway that was amenable to ¹³C tracing analysis. The 2PG pool was typically at or below detection limits. As was expected of WT cells, the initial rate of ¹³C accumulation and the maximum ¹³C enrichment of the glycolate pool were low under VHC pulse conditions. Nevertheless, even under these conditions, we formally demonstrated ¹³C labelling of the glycolate pool with ~0.6 atom% min⁻¹ for HC conditions and up to ~1.4 atom% min⁻¹ for LC conditions. The glycolate/3PGA ratios of these measurements were ~0.09-fold and ~0.18-fold, respectively (Table S1A). Under the HC and LC chase conditions, however, the glycolate pool dropped below the detection limits in WT cells. The *DgicD1* mutant had previously been shown to exhibit a large increase in the glycolate pool, as shown by Eisenhut et al. [15], and therefore allowed detailed analysis of glycolate labelling. The glycolate pool size in the *DgicD1* mutant responded to the reduced CO₂ availability under HC and LC chase conditions (Figure 3). Initially, ¹³C rapidly partitioned into the constantly increasing glycolate pool. To compare the HC and LC chase responses, the kinetics of the glycolate/3PGA ¹³C enrichment ratio were analysed in the *DgicD1* mutant. If glycolate were predominantly or exclusively generated by the oxygenase activity of Rubisco, we would expect nearly identical glycolate/3PGA ratios under both HC and LC conditions, even though the apparent chase kinetics of 3PGA ¹³C enrichment were different. This expectation is shown to be true in Figure 3 (bottom).

Furthermore, we attempted to quantify the flux of carbon into the glycolate and 3PGA pools under LC and HC chase conditions in the *AglD1* mutant. The glycolate turnover time was shorter than the 3PGA turnover time under LC conditions; specifically, it was 2.9 min (glycolate; k = 0.3337 min⁻¹) compared to 90.9 min (3PGA; k = 0.0062 min⁻¹). In contrast, the turnover times were in

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**Table 1.** Rate of ¹³C accumulation during a 0.5–10.0 min ¹³Ci-VHC pulse using 2% (w/w) ¹³C-NaHCO₃ and ¹³C enrichment at maximum labelling (20–60 min) in the pools of 3-phosphoglycerate (3PGA) and phosphoenolpyruvate (PEP).

<table>
<thead>
<tr>
<th>Genotype</th>
<th>Condition</th>
<th>Rate of ¹³C Accumulation ¹³C Enrichment</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>atom% min⁻¹</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>3PGA</td>
<td></td>
<td></td>
</tr>
<tr>
<td>WT</td>
<td>HC</td>
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<td></td>
<td>LC</td>
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</tr>
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</tr>
<tr>
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<tr>
<td></td>
<td>LC</td>
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<td>JgicD1</td>
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<tr>
<td></td>
<td>LC</td>
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</tr>
<tr>
<td>JgcvT</td>
<td>HC</td>
<td>12.6</td>
</tr>
<tr>
<td></td>
<td>LC</td>
<td>18.3</td>
</tr>
<tr>
<td>PEP/3PGA ratio</td>
<td>Rate of ¹³C Accumulation ¹³C Enrichment</td>
<td></td>
</tr>
<tr>
<td></td>
<td>fold</td>
<td>RSD (%)</td>
</tr>
<tr>
<td>WT</td>
<td>HC</td>
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<td>1.28</td>
</tr>
<tr>
<td></td>
<td>LC</td>
<td>1.23</td>
</tr>
</tbody>
</table>

The PEP/3PGA ratios were calculated from paired observations within each sample. Cultures were pre-acclimated and subjected to a chase of either 5% CO₂ (HC) or 0.035% CO₂ (LC) of ambient isotope composition. Note that compared to the HC condition, the LC condition resulted in faster and higher labelling of both the 3PGA and PEP pools. Also note that the PEP pool exhibited more rapid and higher labelling than the 3PGA pool. Each replicate experiment (WT n = 3, mutants n = 2) was an independent time series of 7–8 time points sampled ~0.5–60.0 min after the pulse.

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the same order of magnitude under HC conditions, i.e., 39.8 min (glycolate; \( k = 0.0251 \text{ min}^{-1} \)) and 25.1 min (3PGA; \( k = 0.0399 \text{ min}^{-1} \)). Taking the pool sizes of both metabolites into consideration, we calculated the molar rates of appearance under chase conditions, which may serve as estimates of the rates of oxygenation (glycolate) and carboxylation (3PGA). Under the LC condition, the molar rates of appearance were \( 4.80 \times 10^{-6} \mu\text{mol min}^{-1} \mu\text{g}^{-1} \text{Chl} \) and \( 1.21 \times 10^{-6} \mu\text{mol min}^{-1} \mu\text{g}^{-1} \text{Chl} \) for glycolate and 3PGA, respectively. The ratio of the molar rates of appearance was \( \approx 4.0 \) (glycolate/3PGA). *Synechocystis* cells in the HC condition showed a 100-fold lower ratio (\( \approx 0.04 \) glycolate/3PGA). The molar rates of appearance were \( 4.27 \times 10^{-7} \mu\text{mol min}^{-1} \mu\text{g}^{-1} \text{Chl} \) and \( 1.18 \times 10^{-7} \mu\text{mol min}^{-1} \mu\text{g}^{-1} \text{Chl} \) for glycolate and 3PGA, respectively.

Moreover, the photorespiratory pathway was monitored with respect to the \(^{13}\text{C} \) enrichment in the glycine, serine, and glycerate pools. Again, ratios of \(^{13}\text{C} \) enrichment to 3PGA were used because this approach corrects for the differences in 3PGA labelling among the different experiments (Figure 4). The serine \(^{13}\text{C} \) enrichment ratios of WT cells were higher than the glycine \(^{13}\text{C} \) enrichment ratios throughout the VHC pulse (Figure 4, left panel). The \(^{13}\text{C} \) enrichment ratios of glycine were equal to or less than the \(^{13}\text{C} \) enrichment ratios of serine (data not shown). These observations were consistent under all conditions and in all genotypes (Figure 4). Our observations thus indicated that the synthesis of serine from glycine was minimal under VHC conditions, while the conversion of glycerate to serine or the opposite reaction seemed to occur under HC and LC conditions.

During the chase periods, the glycolate-accumulating *D*\textit{glcD1} mutant showed a transient increase in glycine and serine \(^{13}\text{C} \) enrichment ratios that was similar to the effect observed in the WT (Figure 4, right panel). The *D*\textit{gcvT} mutant, which is deficient in glycine decarboxylase and is known to accumulate glycine under LC conditions [15], showed the expected enhanced accumulation of the \(^{13}\text{C} \) label in the glycine pool under LC chase conditions (Figure 4, central panel).

### Monitoring PEP utilization: Aspartate, malate and citrate

Oxaloacetate, the product of PEP carboxylase activity, was not measured in our study. Instead PEP utilisation was monitored by assessing the kinetics of \(^{13}\text{C} \) enrichment in three metabolic oxaloacetate products: aspartate, malate, and citrate (Figure 5). To normalise the \(^{13}\text{C} \) labelling of these pools, we calculated ratios of the initial rate of \(^{13}\text{C} \) accumulation and maximum \(^{13}\text{C} \) enrichment with respect to PEP instead of the previously chosen 3PGA (Table S1B).

During the VHC pulse, \(^{13}\text{C} \) accumulation was highest in the citrate pool, followed by the malate and aspartate pools. Unexpectedly, the order changed over the monitored time (Figure 5). At the onset of the pulse phase, \(^{13}\text{C} \) enrichment was highest in the aspartate pool, followed by the malate and citrate pools. The reversal of \(^{13}\text{C} \) enrichment was noticeable in cells pre-acclimated to HC and became obvious under LC conditions. The LC condition also demonstrated the mobilisation of unlabelled carbon during the early phase of the VHC pulse. This internal

### Table 2. Sucrose flux estimated from the initial kinetics of a 0–5 min \(^{13}\text{C} \)-VHC pulse.

<table>
<thead>
<tr>
<th>Genotype</th>
<th>Condition</th>
<th>Sucrose Flux</th>
<th>( \mu\text{mol min}^{-1} \mu\text{g} \text{chlorophyll a} )</th>
<th>Avg</th>
<th>RSD (%)</th>
</tr>
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<tbody>
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<td>WT</td>
<td>HC</td>
<td>4.72E-05</td>
<td>104.3</td>
<td></td>
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</tr>
<tr>
<td></td>
<td>LC</td>
<td>1.38E-04</td>
<td>104.3</td>
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<tr>
<td><em>D</em>\textit{glcD1} (sll0404)</td>
<td>HC</td>
<td>1.80E-04</td>
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<td>LC</td>
<td>2.25E-04</td>
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<tr>
<td><em>D</em>\textit{gcvT} (sll0171)</td>
<td>HC</td>
<td>6.63E-05</td>
<td>141.0</td>
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<tr>
<td></td>
<td>LC</td>
<td>1.29E-04</td>
<td>141.0</td>
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</tbody>
</table>

Each replicate experiment (WT \( n = 3 \), mutants \( n = 2 \)) was an independent time series of 7–8 time points sampled \( \approx 0.5–60 \) min after the pulse. The sucrose flux was calculated based on the influx of carbon from the G6P pool according to previously described methods [39]. Cultures were pre-acclimated to 5% CO\(_2\) (HC) or 0.035% CO\(_2\) (LC) of ambient isotope composition.

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**Figure 2. Ratios of PEP/3PGA \(^{13}\text{C} \) enrichment (A) and respective pool sizes (B).** The pulse and chase conditions used in HC and LC acclimated *Synechocystis* sp. PCC 6803 wild-type were identical to the conditions reported in the legend of Figure 1. The initiation of the pulse (0 min) and chase (60 min) are indicated by arrows.

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carbon source first entered the aspartate pool (Figure 5, arrow) and appeared to persist for the longest time in the citrate pool. Under the same conditions, the ornithine analyte showed essentially the same internal carbon mobilisation effect as aspartate but with lower $^{13}$C enrichment (data not shown). Note that due to chemical conversions during the derivatisation of metabolites for GC-MS analysis, the measured ornithine analyte represents the sum of the ornithine, arginine, and citrulline pools.

Indications of additional $^{13}$C fixation through PEP carboxylase activity were found by analysing the ratios of the initial rates of $^{13}$C accumulation for the various compounds with respect to PEP (Table S1B). In WT cells, the 3PGA/PEP, citrate/PEP, and malate/PEP ratios of the initial rates of $^{13}$C accumulation were $\sim 0.96$, $\sim 0.78$, and $\sim 0.62$, respectively. These ratios were not influenced by pre-acclimation of the WT cells to the HC or LC conditions. The aspartate/PEP ratio was, however, increased by...

**Figure 3. Glycolate pool size, $^{13}$C enrichment, and ratio of glycolate/3PGA $^{13}$C enrichment.** The data were acquired during the pulse and chase phases under HC and LC conditions with the glycolate-accumulating $\Delta$gldC1 ($\Delta$ssl0404) mutant. Note the strong increase in the glycolate/3PGA $^{13}$C enrichment ratio under both the HC and LC conditions (cf. Table S1). The initiation of the pulse (0 min) and chase (60 min) are indicated by arrows.

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1.34 fold under the LC condition (0.65) compared to the HC condition (0.48). This LC acclimation effect was even more apparent in the two photorespiratory mutants, \( \text{DglcD1} \) and \( \text{DgcvT} \), with aspartate/PEP ratios approximating 1 under LC conditions in both mutants. A similar but less extreme effect in the mutants was observed for the malate/PEP ratios. These ratios increased by 1.3-fold in cells pre-acclimated to LC compared to those receiving HC. The citrate/PEP ratios showed the same tendency under the LC condition and were only marginally increased in the mutant strains compared to WT. Under HC, the citrate/PEP ratios were, however, reduced in both mutants compared to WT.

**Discussion**

The major paths of carbon assimilation in *Synechocystis*

Our data were essentially consistent with the established fact that 3PGA is the first assimilation product of the Calvin-Benson cycle. Rapid labelling of 3PGA with \(^{13}\text{C}\) suggests that C3-photosynthesis is the dominant carbon fixation pathway in *Synechocystis*. Fixation of \(^{13}\text{C}\) via PEP carboxylase seems to be less important, but occurred in LC acclimated cells (see below). Following the dilution of the initial \(^{13}\text{C}\) label of 3PGA into the metabolite pools that were measured by our GC-MS-based technology, we observed two major branches of carbon flow (Figure 6). The first branch leads from 3PGA to sucrose via G6P with minimal leakage of carbon into the free glucose pool. The flux of carbon from 3PGA through this branch into sucrose appears to reflect the photosynthesis rate of the respective pre-acclimated cells [15] because the enrichment of the sucrose pool is higher in LC cells than in HC cells (cf. Table 2). The observed conversion of 3-PGA into sucrose and probably also into glycogen shows that surplus carbon is converted into storage compounds. The second major path of carbon appears to lead to aspartate, malate, and citrate (Figure 6) via PEP and probably further into the amino acid pools via 2-oxo-glutarate. This second branch of 3PGA utilisation directly fuels the biosynthesis of primary metabolites, specifically amino acids that are generated via the incomplete tricarboxylic acid cycle of *Synechocystis*. In agreement

![Figure 4. Ratios of glycine and serine \(^{13}\text{C}\) enrichment to 3PGA \(^{13}\text{C}\) enrichment. Pulse and chase phases under HC conditions (top) and LC conditions (bottom) were used with the wild-type (left), the \( \text{AgcvT (Dts0171)} \) mutant (middle) and the \( \text{DglcD1 (Dts0404)} \) mutant (right) strains. Glycolate data from the \( \text{DglcD1} \) mutant are included in the right panel. The initiation of the pulse (0 min) and chase (60 min) are indicated by arrows. doi:10.1371/journal.pone.0016278.g004](Image)
with the interruption of the canonical tricarboxylic acid cycle in Synechocystis, fumarate and succinate were labelled more slowly than malate (data not shown).

We further conclude that C4 carboxylation via PEP carboxylase also occurs under our experimental conditions, specifically in LC acclimated cells. The joint action of C3 and C4 carboxylation via Rubisco and PEP carboxylase, respectively, has been revealed by 13C labelling experiments using different cyanobacterial strains, as shown by Dohler [20]. However, the relative ratio is still a matter of discussion. The oxaloacetate product, which cannot directly be measured by our current technique, appears to be rapidly converted into aspartate and malate via the aspartate aminotransferase and malate dehydrogenase reactions. Of the potential oxaloacetate products, citrate is the most rapidly labelled, possibly because it receives part of its carbon backbone via acetyl-CoA. Acetyl-CoA should be highly labelled because it is generated from the rapidly labelled PEP pool via the final steps of the glycolytic pathway. However, as judged by the rates of 13C accumulation, the C3 carboxylating pathway from 3PGA via PEP appears to be the main route to transfer newly fixed organic carbon into the central metabolism of Synechocystis. The 13C enrichment of oxaloacetate products shows that primary 13C fixation via Rubisco is dominant under our experimental conditions.

One of the unexpected results was the observation that first aspartate and then malate and citrate receive 13C from internal stores during the 13C-VHC pulse (Figure 5). We interpret this observation as a carbon mobilisation from internal sources into the aspartate and ornithine/arginine/citrulline pools. The source could be fast cyanophycin degradation. Cyanophycin is a storage polymer in cyanobacteria that has a poly-aspartate backbone and arginine side chains. Cyanophycin accumulates mainly as a response to excess nitrogen in Synechocystis [21]. The VHC pulse results in an environment characterised by high C₂ availability and relatively low nitrogen. This should apply especially to LC cells, which were pre-acclimated to rather high C/N conditions. Cyanophycin degradation mobilises stored nitrogen and may ultimately lead to rebalancing of the intercellular C/N ratio. Such a mobilisation process should be considered in future studies that aim at dissecting the relative contribution of C₃ and C₄ metabolism to the biosynthesis of organic acid intermediates such as citrate.

Indications for carbon channelling from 3PGA into the PEP pool

Had Calvin and Benson [22,23] used Synechocystis to unravel the photosynthetic carbon cycle, PEP would have likely been misinterpreted as the first assimilation product of the carboxylation reactions. We found clear evidence that the PEP pool of Synechocystis is not only more rapidly labelled but also has a slightly but significantly higher final 13C enrichment than the 3PGA pool under all of our experimental conditions (Table 1, Figure 2). Even today, this phenomenon could be interpreted as evidence for the presence of alternative CO₂ fixation mechanisms in cyanobacteria. Such pathways for CO₂ fixation, e.g., the reductive tricarboxylic acid cycle or the carbamoylphosphate synthetase pathway, have previously been suggested to be operative in cyanobacteria [24]. Both pathways have been implicated for ancillary CO₂ fixation. In view of the incomplete tricarboxylic acid cycle in Synechocystis and considering our data, i.e., the lower rates of 13C accumulation in the citrate and malate pools compared to the PEP pool, we consider it unlikely that under our growth conditions Synechocystis utilizes any of the above ancillary routes of CO₂ fixation or the hydroxypropionate pathway for CO₂ fixation, which was recently discovered in Chloroflexus aurantiacus [25]. This view is supported by the absence of genes for such alternative C fixation pathways from the complete genome sequence of Synechocystis (CyanoBase at http://genome.kazusa.or.jp/cyanobase/Synechocystis).

Instead, we favour the interpretation that the majority of newly assimilated carbon is channelled from 3PGA to PEP, and the much smaller PEP pool is more quickly saturated with 13C. With this interpretation, we need to assume that two sub-pools of 3PGA exist in Synechocystis. The carboxysome, a cyanobacterial micro-compartment that is a component of the CCM and harbours most of the cellular Rubisco, may well be a rapidly labelled 3PGA pool from which PEP is produced. The second 3PGA pool may be cytosolic. The cytosolic 3PGA pool could originate from a second pool of Rubisco. Rubisco was recently indicated to exist outside of the carboxysome but attached to thylakoids and in close association with other Calvin-Benson cycle enzymes [26]. Carbon should be channelled into PEP either within the carboxysomes or along with the export of 3PGA from the carboxysome into the cytosol. The cytosolic 3PGA pool may well be diluted with unlabelled carbon as a result of the turnover or mobilisation of previously assimilated soluble sugars or glycogen. Channelling into PEP may also indicate that the PEP carboxylase reaction for Synechocystis is more important than previously described [27]. This conclusion is well supported by the rapid accumulation of assimilated carbon in pools downstream of oxaloacetate (cf. discussion above) and may explain enigmatic early reports of high accumulations of label in unexpected metabolite pools, such as the aspartate pool [20,24].
Formal demonstration of photorespiratory flux

We detected a significant partitioning of the $^{13}$C label into the glycolate pool, primarily in the $AgcD1$ mutant, but also in WT cells under our VHC pulse conditions. Thus we show that even when *Synechocystis* cells are exposed to very high $C_i$ concentrations, the oxygenation reaction of Rubisco is not completely suppressed. This finding is in agreement with the reduced growth and photosynthesis caused by glycolate accumulation in cells of the $AgcD1$ mutant grown under HC conditions [13].

The glycolate-accumulating $AgcD1$ mutant and the glycine-accumulating $AgcT$ mutant proved to be valuable tools to generate a detailed demonstration of photorespiratory flux (Figure 6). The $AgcD1$ mutant allowed the demonstration of enhanced carbon flux into the glycolate pool after a shift to lowered $C_i$ during the chase phase (Figure 3). In addition, this mutant enabled an estimation of the rates of oxygenation and carboxylation under HC and LC chase conditions by quantification of the molar rates of $^{13}$C appearance in the glycolate pool compared to the 3PGA pool. An enhanced photorespiratory flux was found under LC conditions compared to HC conditions. In contrast, WT cells displayed a very low photorespiratory flux under ambient CO$_2$ conditions, which was consistent with previous reports on the quantitatively low importance of photorespiratory 2PG metabolism [11].

The $AgcT$ mutant demonstrated enhanced $^{13}$C flux into the glycolate pool under chase conditions. This evidence may explain the coinciding increase in glycine levels (Figure 4), which is possibly linked to the increased production of glycine and reduced glycine utilisation of this slow-growing mutant [13].

**Figure 6. Scheme of central carbon and nitrogen metabolism in *Synechocystis* sp. strain PCC 6803.** Ellipses highlight the $^{13}$C labelled metabolites detected and evaluated in this study. The enzymatic steps affected in the $AgcD$ and $AgcT$ mutants are marked by boxes.
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Mutants and WT compared under HC and LC conditions

HC and LC acclimated *Synechocystis* cells were labelled with $^{13}$C to assess the differential acclimation to high and limiting $C_i$ availability. Labelling of 3PGA and sucrose confirmed previous observations on the respective photosynthesis rates, i.e., that HC acclimation resulted in lower maximal assimilation rates than LC acclimation, and that $AgcD$ mutants appear to be partially deficient in photosynthesis. Moreover, HC acclimation was linked to greater channelling of assimilate into the PEP pool (Figure 2). The apparent differences between the LC and HC acclimations were most clearly assessed when comparing relative $^{13}$C enrichment after normalisation to either 3PGA or PEP in the different experiments (Figures 4 and 6). The ratios demonstrated only minor differences of $^{13}$C partitioning into the glycolate pool after a HC or LC chase in the $AgcD1$ mutant. In addition, flux through PEP carboxylase appeared to be slightly affected by the different conditions in WT cells. Only an increased aspartate/PEP-labelling ratio was observed in LC acclimated WT cells (Table S1B), which indicates PEP carboxylase activation under LC conditions. In contrast, in addition to the aspartate/PEP ratio, the mutants exhibited increased malate/PEP and citrate/PEP ratios.
for the initial rates of $^{13}$C accumulation, which is consistent with an enhanced flux through the PEP carboxylation reaction under LC conditions. The enhanced $^{13}$C fixation through PEP carboxylase in both photosynthetic mutants, and to a lesser extent in WT cells, could indicate that this alternative carboxylating enzyme is activated in LC cells to compensate for the inhibitory effects of glycolate or glycine on Rubisco and other Calvin-Benson cycle enzymes.

**Materials and Methods**

**Strains and culture conditions**

The strain *Synechocystis* sp. PCC 6803 was obtained from Prof. Murata (National Institute for Basic Biology, Okazaki, Japan) and served as the WT for this study. The generation and characterisation of the mutants in the *Synechocystis* sp. PCC 6803 photorespiratory pathway, i.e., *AgeD1*, which bears a defect in the glycolate dehydrogenase-coding gene *sll0404*, and *AgeT*, which bears a defect in the gene *sll0171* that encodes the T-protein subunit of the glycine decarboxylase complex, have been described elsewhere [13,28]. Cultivation was performed in BG11 medium [29] at pH 8.0 with an initial optical density (OD$_{680}$) of ~0.8, which is equal to $\sim 10^5$ cells mL$^{-1}$. Mutants were grown in the presence of 50 µg mL$^{-1}$ kanamycin (Km) or 20 µg mL$^{-1}$ spectinomycin (Sp). Potential contamination by heterotrophic bacteria was ruled out by spreading 0.2 mL of culture on LB plates.

Axenic cultures were grown phototrophically at 29°C in batch cultures using 3-cm glass vessels with 5-mm glass tubes for aeration with 5% CO$_2$-enriched air (HC) with a bubbling flow rate set to 5 mL min$^{-1}$. Cultures were continuously illuminated with warm light using an Osram L58 W32/3 at 130 µmol photons s$^{-1}$ m$^{-2}$. Pre-cultures were split into equal parts. One part continued to be grown under HC conditions while the second subculture was grown under Ci limiting conditions (LC conditions) by bubbling with ambient air containing 0.035% CO$_2$. Acclimation to low Ci by pre-cultivation with ambient air was performed 24 h prior to the isotope-labelling experiments. The pH of the growth medium was stable under the chosen HC and LC cultivation conditions, which were essentially as described earlier [15].

**Transient $^{13}$C$_i$ isotope-labelling experiments**

Pulse experiments with $^{13}$C$_i$ were performed by transferring 40 mL of HC or LC pre-culture into a new cultivation vessel that was kept without bubbling under otherwise identical conditions. The transient isotope pulse was started under very-HC conditions (VHC), i.e. 2% (w/w) sodium hydrogen carbonate, by adding 10 mL of a $^{13}$C-bicarbonate stock solution prepared by dissolving 1 g sodium hydrogen carbonate 98 atom% $^{13}$C (Sigma-Aldrich) in 10 mL BG11. The pH of the $^{13}$C-bicarbonate stock solution was adjusted to 8.0. Subsequent chase experiments were performed after a 60 min pulse according to the pre-acclimation, either under 5% CO$_2$ (HC) or 0.035% CO$_2$ (LC) conditions. The BG11 medium was exchanged by centrifugation (3000g, 2 min, 22°C), careful removal of the supernatant and re-suspension in the original volume. The fresh medium was pre-adjusted to the ambient $^{13}$C/$^{12}$C isotope ratio and CO$_2$ concentration by bubbling with ambient air. Care was taken to minimise the carryover of $^{13}$C upon medium exchange. To enable fast sampling of the chase kinetics, a wash of the cell pellet had to be avoided. Samples were taken immediately after pulse or chase and at 1, 3, 5, 10, 15, 20 (or 30), and 60 min. Observations of pulse and chase kinetics were repeated with 2–3 independent cultivations.

In our hands a clear step change of $^{13}$C label was only possible using high bicarbonate concentrations at the cost of a change of osmolarity within the growth medium. The use of low enrichments for pulse and chase experiments, which are possible using the highly sensitive radioactive $^{14}$C detection methods, are not applicable to the less-sensitive $^{13}$C detection methods. To control for unavoidable imperfect step changes, we normalised the initial rates of $^{13}$C accumulation and the $^{13}$C enrichment in downstream metabolite pools of a pathway to the respective parameters of the first $^{13}$C assimilation products, i.e., 3PGA and PEP (cf. supplementary Methods S1).

**GC-EI-TOF-MS analysis of metabolite pool sizes and mass isotopomer distributions**

The previously described GC-EI-TOF-MS metabolite profiling technology for methoxyaminated and trimethylsilylated methanol/water-soluble metabolites from *Synechocystis* [15,30] was applied to assess a combination of both the metabolite pool sizes and the respective $^{13}$C mass isotopomer distributions from $^{13}$C labelled samples. Culture samples of 2–7 mL, equivalent to about $10^5$ cells mL$^{-1}$, were harvested and separated from the media by fast filtration in the light using a glass vacuum filtration device with controlled temperature and illumination. To minimize the time between sample collection and analysis, no wash was performed. This sampling method, which uses a filter disc to remove secreted metabolites and components of the growth medium, has been previously described [31]. In addition, we performed rapid metabolic inactivation by immediately shock-freezing the cells on the filter disk in liquid N$_2$ to obtain the quickest possible samples for tracing studies [22,23]. Recovery checks by internal standardisation using chemically synthesised stable isotope-labelled reference compounds had to be omitted so as not to interfere with the $^{13}$C tracing experiments. Instead, metabolite pools were normalised to the chlorophyll a content [14]. The relative changes of normalised metabolite pool sizes were based on the peak intensities. In addition to the conventional metabolite profiling procedure, the sum of all observed mass isotopomers of characteristic fragments was calculated to enable pool size quantification in the presence of shifting mass isotopomer distributions. The metabolites glycolate, glycine, serine, glucose-6-phosphate (G6P), sucrose, glyceraldehyde, phosphoenolpyruvate (PEP) and 3-phosphoglycerate (3PGA) were externally calibrated using a dilution series of nine concentration points, ranging from 0.04 ng µL$^{-1}$ to 166.67 ng µL$^{-1}$ of a chemically defined mixture of authenticated reference compounds in equal amounts at 1.0 mg mL$^{-1}$ for each compound [32].

**Data processing and compound identification**

GC-TOF-MS chromatograms were processed using TagFinder-Software [33]. Analytes were manually identified using the TargetFinder plug-in of the TagFinder-Software and a reference library of ambient and $^{13}$C labelled mass spectra and retention indices (RI) from the Golm Metabolome Database (GMD, http://gmd.mpimp-golm.mpg.de/) [34,35]. A peak intensity matrix containing all available mass isotopomers of characteristic mass fragments that represented the primary metabolites under investigation was generated by TagFinder. This matrix was processed using the CORRECTOR software tool (http://www-en.mpimp-golm.mpg.de/~03-research/researchGroups/01-dept1/Root_Metabolism/smp/CORRECTOR/index.html). Using this batch processing tool, we calculated the sum of mass isotopomer intensities and the $^{13}$C enrichments of mass fragments that had been annotated previously [17] using previously described methods [36,37].
Calculations and statistical data mining
Data management, data transformation, calculations, and statistical analyses were performed using Microsoft Office Excel 2003 software, the R 2.9.1 statistical programming package, and SigmaPlot 11.0 software (Systat Software Inc., San Jose, CA, USA). Calculations of the molar rate of appearance (Ra) for the Rubisco reactions were performed according to Wolfe and Chinkes [38] using the equation Ra = k*Q, where Q represents the pool size of the respective metabolite. To obtain k, the turnover rate, the experimental data were fitted to the equations Et = E0(1 - e^(-kt)) or Et = E0 - E0(1 - e^(-kt)). The turnover rate k^(-1) of a pool is given in min^(-1), E0 represents the isotopic enrichment at time t = 0, Et the plateau enrichment and E0 the enrichment at t = 0. The sucrose flux was calculated based on the influx of the precursor G6P, as described by Roessner-Tunali and co-authors [39].

Error estimates and error minimisation are relevant when judging labelling results. The technical precision when determining the 13C enrichment of a metabolite pool is generally below 2% relative standard deviation (RSD) when using the GC-TOF-MS metabolite profiling method [17]. In contrast, the technical error of a metabolite pool size determination using the same method is about one order of magnitude higher, in the range of 5–20% RSD [40]. In addition to the technical error, the biological variability, i.e., the culture-to-culture differences of replicated Synechocystis experiments, was considered using the 3PGA measurement as a test case. The precision of 3PGA pool size determinations in replicate WT and mutant cultures had previously been determined as 25.6% RSD (n = 9), similar to the average 22.9% RSD (n = 9) of all metabolites observable by GC-TOF-MS profiling (cf. Table S1 of [15]). The culture-to-culture variation of the parameters was smaller. The maximum 13C enrichment, determined as atom% at 20–60 min, of the 3PGA or PEP pools had, on average, 6.3% RSD, while the initial rate of 13C accumulation, determined as the atom% min^(-1) by linear regression (r^2 = 0.85–0.99), in either of the pools exhibited, on average, 17.1% RSD, as assessed by triplicate WT and duplicate mutant experiments (Table 1).

Supporting Information
Methods S1 Motivation, detailed description and error assessment of the experimental design chosen for the comparative metabolic flux phenotyping of Synechocystis sp. PCC 6803 wild-type and photorespiratory mutants. (DOCX)

Figure S1 Head-to-tail view of ambient (red) and maximally 13C labeled (blue) mass spectra from the 3-phosphoglycerate (4TMS) and phosphoenolpyruvate (3TMS) analytes of Synechocystis. Mass fragments suitable for mass isotopomer distribution analysis are indicated by an asterisk [17]. Insets indicate the identifiers of the Golm Metabolome Database [35] and the expected retention index (RI) and experimental deviation (ΔRI%) within Synechocystis extracts. The empirical mass distribution vectors of the M-15 fragments, i.e., C14H23O2PSi4 (M0-3 = 459-462) from the 3-phosphoglycerate (4TMS) analyte and C14H21O2PSi4 (M0-3 = 369-372) from the phosphoenolpyruvate (3TMS) analyte, are shown on the right, as determined from an experiment performed under the HC condition. (TIF)

Figure S2 13C enrichment of the glucose, glucose-6-phosphate and sucrose pools in the wild-type strain Synechocystis sp. PCC 6803. Data from LC acclimation followed by a VHC pulse at t = 0 min and LC chase at t = 60 min or 120 min are shown. Note that glucose was detectable only under VHC pulse conditions in 2 of 3 replicate experiments. (TIF)

Table S1 Initial rates of 13C accumulation. Data were acquired during a 0.5-10.0 min 13C pulse and 13C enrichment at maximum labelling (20-60 min) in (A) the glycolate pool compared to the 3PGA or PEP pools and (B) aspartate, malate, and citrate pools compared to the PEP pool. The ratios of the initial rates of 13C accumulation and the 13C enrichment at maximum labelling were calculated from paired observations within each sample. Cultures were pre-acclimated and subjected to a chase with ambient isotope composition of either 5% CO2 (HC) or 0.035% CO2 (LC).

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Author Contributions
Conceived and designed the experiments: JH JG DS HB MH JK. Performed the experiments: JH JG DS. Analyzed the data: JH JG DS HB MH JK. Contributed reagents/materials/analysis tools: JH JG DS HB MH JK. Wrote the paper: JH JG DS HB MH JK.

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