Genome-Scale Fluxome of *Synechococcus elongatus* UTEX 2973 Using Transient $^{13}$C-Labeling Data

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*Synechococcus elongatus* UTEX 2973 (*Synechococcus 2973*) has the shortest reported doubling time (2.1 h) among cyanobacteria, making it a promising platform for the solar-based production of biochemicals. In this meta-analysis, its intracellular flux distribution was recomputed using genome-scale isotopic nonstationary $^{13}$C-metabolic flux analysis given the labeling dynamics of 13 metabolites reported in an earlier study. To achieve this, a genome-scale mapping model, namely iSyu593, was constructed using the iSyu593 mapping model for *Synechocystis* sp. PCC 6803 (*Synechocystis 6803*) as the starting point encompassing 593 reactions. The flux elucidation revealed nearly complete conversion (greater than 96%) of the assimilated carbon into biomass in *Synechococcus 2973*. In contrast, *Synechocystis* 6803 achieves complete conversion of only 86% of the assimilated carbon. This high biomass yield was enabled by the reincorporation of the fixed carbons lost in anabolic and photorespiratory pathways in conjunction with flux rerouting through a noncarbonic reductase reaction such as phosphoketolase. This reincorporation of lost CO$_2$ sustains a higher carbon efficiency drive, acetyl-coenzyme A was entirely produced using the carbon-efficient phosphoketolase pathway. Comparison of the *Synechococcus 2973* flux map with that of *Synechocystis* 6803 revealed differences in the use of Calvin cycle and photorespiratory pathway reactions. The two species used different reactions for the synthesis of metabolites such as fructose-6-phosphate, glycine, sedoheptulose-7-phosphate, and Ser. These findings allude to a highly carbon-efficient metabolism alongside the fast carbon uptake rate in *Synechococcus 2973*, which explains its faster growth rate.

Photosynthesis produces a wide range of metabolites from only water, sunlight, and CO$_2$. Cyanobacteria are oxytrophic phototrophs that are more amenable to genetic transformations than microalgae and plants, making them ideal photosynthetic production platforms for industrially relevant chemicals and biofuels (Ducat et al., 2011; Quintana et al., 2011). While the industrial relevance of cyanobacteria has been restricted by their slow growth rates under photoautotrophic conditions, the fast-growing cyanobacterium *Synechococcus elongatus* UTEX 2973 (here referred to as *Synechococcus 2973*), which has a reported doubling time of 2.1 h under high-light and high-CO$_2$ conditions, has emerged as a promising platform (Yu et al., 2015). The puzzling DNA sequence similarity with the well-studied and comparatively slower-growing cyanobacterium *Synechococcus elongatus* PCC 7942 (*Synechococcus 7942*) has made *Synechococcus 2973* a model organism that could enable us to uncover the factors responsible for a faster photoautotrophic growth (Yu et al., 2015; Abernathy et al., 2017; Mueller et al., 2017). This has led to the rapid development of genetic tools, including a CRISPR/Cas9 system (Wendt et al., 2016). The faster growth rate and availability of genetic tools in *Synechococcus 2973* make it a promising solar-based production platform for biochemicals. Photosynthesis-based Suc production already has been demonstrated in this organism (Song et al., 2016). Ongoing efforts to understand and engineer *Synechococcus 2973* into a successful production host will benefit from an understanding of its metabolic flows supporting fast growth. This can be accomplished using $^{13}$C-metabolic flux analysis ($^{13}$C-MFA; Long and Antoniewicz, 2014; Young, 2014; Niedenführ et al., 2015; Allen, 2016).

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Isotope tracing using a one-carbon substrate results in uniform labeling of metabolites under an isotopic steady state (Shastri and Morgan, 2007). As a consequence of this, isotopic nonstationary ¹³C-metabolic flux analysis (INST-MFA) is needed to elucidate photoautotrophic flux distribution. This methodology has been used extensively to study the metabolism of cyanobacteria (Adebiyi et al., 2015), microalgae (Wu et al., 2015), and plants (Ma et al., 2014). In contrast to steady-state MFA, INST-MFA elucidates fluxes so as to recapitulate experimentally measured metabolite-labeling dynamics as well as steady-state labeling distributions (Nöh et al., 2007). The complexity of the underlying nonlinear least squares problem and the number of balance equations required to predict the labeling distribution of the metabolites has so far placed significant constraints on the scope of the metabolic reactions considered at one time. Generally, INST-MFA has been restricted to core models spanning central metabolism only (Nöh and Wiechert, 2011). Such simplifications introduce biases regarding pathway utilization (Gopalakrishnan and Maranas, 2015), which can result in inaccurate uncertainty calculations and limit meaningful inferences about energy metabolism in an organism. Furthermore, reaction lumping restricted the metabolite-labeling dynamics afforded by a core model, leading to significant disagreements between experimentally measured labeling dynamics and those predicted using simplified models (Nöh and Wiechert, 2011). In order to mitigate the effect of these factors on the accuracy of flux estimation, the scope of the INST-MFA technique has been expanded to genome-scale models with the implementation of an efficient algorithm for the simulation of metabolite-labeling dynamics (Gopalakrishnan et al., 2018).

Meta-analysis of the labeling dynamics observed under photoautotrophic conditions in *Synechocystis* sp. PCC 6803 (*Synechocystis* 6803) using genome-scale INST-MFA showed that the genome-scale mapping model (GSMM), namely *msyn617*, fitted the experimental data better than the core model (Gopalakrishnan et al., 2018). Interestingly, the flux distribution obtained using *msyn617* was qualitatively different from the one obtained using the core model, as indicated by the reliance on transaldolase (TAL) for the synthesis of fructose-6-phosphate (*f6p*), a bifurcated photosynthetic C2 cycle, and off-gassing of fixed CO₂ by anabolic reactions outside core metabolism. These prediction departures from a core mapping model demonstrate the importance of using comprehensive GSMMs while performing INST-MFA to glean an accurate description of internal metabolic fluxes.

 Genome-scale INST-MFA requires a high-quality genome-scale model (GSM; Gopalakrishnan and Maranas, 2015) spanning all the reactions encoded by the genome annotation without any presumption about their activity under a given condition. The GSM must contain a detailed biomass equation, and the synthesis pathways of macromolecular precursors such as amino acids, lipids, and nucleotides should be traced completely. The lumping of reactions should be kept to a minimum, and care should be taken to account for all the reactions in the amino acid and peripheral metabolism that produce central carbon intermediates such as acetate, fumarate, pyruvate, and CO₂ as by-products. Once a GSM is available, the next step is to construct a GSMM describing the carbon mapping from the reactants to products for each reaction. Carbon-mapping information for reactions can be obtained from pathway databases such as MetaCyc (Caspi et al., 2018), existing GSMMs, and reaction mechanisms available from the literature. Atom-mapping algorithms such as Determination of Reaction Mechanisms (First et al., 2012), Minimum Weighted Edit-Distance (Latendresse et al., 2012), InfoChem-Map (Kraut et al., 2013), Canonical Labeling for Clique Approximation (Kumar and Maranas, 2014), and Reaction Decoder Tool (Rahman et al., 2016) also can be used to generate plausible mappings when this information is unavailable.

In this study, the photoautotrophic metabolism of *Synechococcus* 2973 under optimal growth conditions is elucidated using genome-scale INST-MFA. For this purpose, we constructed the GSMM *msyu593* using the GSM *sSyu683* (Mueller et al., 2017). Flux variability analysis (FVA; Mahadevan and Schilling, 2003) was used to identify the reactions that are active under photoautotrophic conditions by constraining the model with experimentally measured bicarbonate uptake and growth rate. Flux elucidation using INST-MFA reveals a high Rubisco activity capable of reincorporating the fixed carbon lost in photorespiration and peripheral metabolism. The analysis also predicted that this organism relies entirely on the carbon-efficient phosphoketolase (Pketolase) pathway instead of the conventional pyruvate dehydrogenase (PDH) reaction for the synthesis of acetyl-CoA. The oxidative pentose phosphate pathway carried a negligible flux. These factors facilitate a nearly perfect routing of assimilated carbon toward biomass. Prediction departures from the core model (Abernathy et al., 2017) include the usage of TAL for sedoheptulose-7-phosphate (*s7p*) synthesis and Pketolase for acetyl-CoA synthesis. Comparison with the genome-scale flux distribution of *Synechocystis* 6803 (Gopalakrishnan et al., 2018) grown under its optimal growth conditions (Young et al., 2011) revealed interesting differences in pathway recruitment that were not apparent through the core model-based studies or their respective GSMMs. *Synechococcus* 2973 has a higher overall carbon conversion (96%) compared with *Synechocystis* 6803 (86%). As a result, the biomass yield (per fixed carbon) of *Synechococcus* 2973 was 2% higher than that of *Synechocystis* 6803. This increase in the biomass yield accounted for 2.9% of the difference in the growth rate between these two species, with the remaining 97.1% due to differences in carbon uptake kinetics. The two species also differed in the choice of Calvin cycle reactions for the synthesis of *f6p* and *s7p*. In *Synechococcus* 2973, Ser and Gly were synthesized using the photosynthetic C2 cycle, whereas
*Synechocystis* 6803 relied instead on the phosphoSer (PSER) pathway and C2 cycle. The new flux map obtained in this study provides valuable insight into a cyanobacterial strain of great industrial importance, directly supporting any rational redesigning of its metabolism to generate a superior production host.

### RESULTS

#### The Genome-Scale Carbon-Mapping Model of *Synechococcus* 2973

The elementary metabolite unit (EMU) network, obtained from imSyu593, capturing the carbon paths that can contribute to the labeling of the 13 fitted metabolites, spanned 232 metabolites and 352 reactions and consisted of 1,271 EMUs. Scale-up to imSyu593 introduces 41 novel carbon paths that were not included during flux elucidation using a core model (Abernathy et al., 2017). This includes acetyl-CoA production via the P Ketolase pathway (Fig. 1), Ser production via the PSER pathway, and carbon recycling from peripheral metabolic pathways such as fatty acid biosynthesis, pantothenic and CoA biosynthesis, riboflavin metabolism, and porphyrin and chlorophyll biosynthesis. The glyoxylate shunt was excluded in imSyu593 based on the lack of genomic evidence (Fig. 1; Yu et al., 2015; Abernathy et al., 2017; Mueller et al., 2017). These differences in the carbon paths captured by the core model and imSyu593 cause significant differences in the predicted labeling dynamics. Compared

![Figure 1. Schematic of the central carbon metabolism of *Synechococcus* 2973.](image-url)
with imSyu617, imSyu593 contains two novel carbon paths producing acetyl-CoA and erythrose-4-phosphate (e4p). The PKetolase pathway consisting of the enzymes PKetolase, ATP:acetyl phosphotransferase, and acetate:CoA ligase allows acetyl-CoA production from the sugar phosphates (f6p and xylulose-5-phosphate) in the Calvin cycle while bypassing the lower glycolysis. The second novel carbon path was due to the presence of 5,6-dimethylbenzimidazole synthase (EC 1.13.11.79), which degraded FMN into dilaurate, dimethyl benzimidazole, and the central carbon intermediate e4p. This resulted in an alternative carbon path from ribose-5-phosphate (r5p) to e4p through riboflavin biosynthesis.

The Flux Distribution of Synechococcus 2973

The estimation of intracellular fluxes using imSyu593 led to a better fit (sum of squared residuals [SSR] = 363, degrees of freedom = 402) of the labeling data when compared with that with the core model (SSR = 610, degrees of freedom = 535; Fig. 2). The F test was used to analyze the statistical significance of this improvement in the fit. The obtained F of 2.079 (P = 1.39 × 10⁻⁸) indicates with 95% confidence that the improvement is due to additional active carbon paths accounted for in imSyu593 compared with that in the core model and not due to the mere increase in the number of parameters. A large reduction in the SSR was observed for metabolites Glc-6-P, r5p, s7p, 6-phosphogluconate, 3-phosphoglycerate, 2-phosphoglycerate, phosphoenolpyruvate, citrate, succinate, and malate. The estimated 95% confidence intervals showed that 609 reactions were resolved with flux ranges narrower than 10 mmol g⁻¹ dry weight h⁻¹ for a bicarbonate uptake rate of 100 mmol g⁻¹ dry weight h⁻¹. Of this, only 334 reactions were resolved due to growth coupling, whereas the rest of the reactions were resolved due to MFA. Of the 78 unresolved reactions, 45 reactions are outside the purview of the EMU balances and are involved in pathways such as photosynthesis, oxidative phosphorylation, and the exchange and transport of oxygen, photons, protons, and water. The remaining 33 reactions, involved in pathways such as Ala and Asp metabolism, citrate cycle, Glu metabolism, glycolysis/gluconeogenesis, pentose phosphate pathway, purine metabolism, and starch and Suc metabolism, participate in unresolved futile cycles expending ATP. This is because the photosynthesis reactions responsible for generating ATP are not captured by the EMU balances and, therefore, are not constrained by the labeling data.

Comparison of flux ranges generated using MFA and FVA (Supplemental Table S1) showed an average of 56% bound contraction (reduction in the MFA flux range relative to the FVA flux range). Reactions with the most impact flux range reductions (greater than 99% bound contraction) were from central metabolism and included inactivated metabolic pathways such as the oxidative pentose phosphate pathway, lower tri-carboxylic acid cycle, the PSER pathway, and glycolate degradation. This aids in the resolution of most reactions from central metabolism and decreases the uncertainty in biomass yield prediction using MFA by 74% relative to that using FVA. This 74% reduction in biomass flux range caused 334 growth-coupled reactions to reach a 74% bound contraction using MFA. The highly efficient carbon routing to biomass limited the possibility of the production of by-products such as sulfoquinovosyl-diacylglycerol, digalactosyl-diacylglycerol, and phosphatidylglycerol, contributing to the contraction of flux ranges in 43 peripheral metabolic reactions using MFA relative to FVA. Nevertheless, 78 reactions, including those involved in energy metabolism (photosynthetic light reactions and oxidative phosphorylation), remained poorly resolved by MFA as with FVA.

The flux distributions through the carbon fixation pathways obtained using imSyu593 and the core model were qualitatively different (Fig. 3). s7p was produced via the TAL reaction in imSyu593 as opposed to via the sedoheptulose-1,7-bisphosphate aldolase (SBA) and

![Figure 2](https://example.com/image2.png)

Figure 2. Comparison of variance-weighted SSR obtained, in the case of Synechococcus 2973, using the core model and imSyu593 for the various metabolite fragments. Green bars correspond to the core model, and brown bars correspond to imSyu593. Lower SSR indicates a better recapitulation of the labeling dynamics for the corresponding fragment. *Fragments for which the difference in SSR is greater than 25.
sedoheptulose-1,7-bisphosphatase (SBPase) reactions in the core model. To test the robustness of this inference, we removed TAL from imSyu593 and repeated the flux elucidation. Upon removing TAL from imSyu593, the minimum SSR increased to 368 and carbon flux was redirected through the SBA and SBPase reactions. The SSR increase was attributed to an increased disagreement between predicted and measured labeling dynamics of s7p and sedoheptulose-1,7-bisphosphate (sbp) fragments. The F test ($F = 5.61$, $P = 0.018$) indicated that the reduction in the SSR caused by including TAL was statistically significant and formed the basis for rejecting the model lacking TAL. The oxidative pentose phosphate pathway carried a negligible flux, which is consistent with flux predictions using the core model. The 95% confidence interval for this flux identified a lower bound of zero, indicating the dispensability of the oxidative pentose phosphate pathway under photoautotrophic conditions in *Synechococcus* 2973, which is consistent with the experimentally observed nonessentiality of the zwf gene in *Synechococcus* 2973 (Abernathy et al., 2017).

Flux elucidation using imSyu593 revealed that acetyl-CoA is synthesized predominantly from f6p via the PKetolase pathway instead of PDH as reported by the core model (Fig. 3). The PKetolase reaction splits f6p into e4p and acetyl phosphate, which is converted to acetyl-CoA via the ATP:acetate phosphotransferase and acetate-CoA ligase reactions (Fig. 1). No flux was seen through PDH, indicating that the PKetolase pathway is the sole source of acetyl-CoA in *Synechococcus* 2973. Since acetyl-CoA biosynthesis via the PKetolase pathway bypasses lower glycolysis, flux through lower glycolysis was reduced by 50% compared with that in the core model. PKetolase also catalyzes the conversion of xylulose-5-phosphate to glyceraldehyde-3-phosphate. However, this reaction was found to carry no flux. Removal of the f6p-utilizing PKetolase reaction resulted in a statistically acceptable fit with an increased SSR of 379. An F test ($F = 18.15$, $P = 0.000025$) indicated that the increase in the SSR was statistically significant enough to reject the model that lacks the PKetolase, demonstrating that the labeling data support that the PKetolase pathway was active under the experimental conditions. The pathway is usually not included in the core models of cyanobacterial metabolism; therefore, this is the first report that supports the operation of the native PKetolase pathway under photoautotrophic conditions. Flux through this pathway was corroborated by proteomic analysis reporting that the PKetolase enzyme level is comparable with that of phosphoenolpyruvate carboxylase, citrate synthase, and subunits of the PDH complex under photoautotrophic conditions in *Synechococcus* 2973 (Yu et al., 2015).

Rubisco oxygenase carried a 3-fold higher flux in imSyu593 compared with that in the core model. The ratio of Rubisco carboxylase to Rubisco oxygenase flux was found to be 83:17. High photorespiration was expected, since the nonstationary labeling experiment was performed under high-light conditions (Abernathy et al., 2017). Under similar conditions, previous studies have reported a 2.4-fold higher rate of photorespiratory oxygen uptake in *Synechococcus* 2973 compared with that in *Synechococcus* 7942 (Ungerer et al., 2018a). The high rate of photorespiration caused a large flux through the C2 cycle for phosphoglycolate salvage, which generates all of the Gly and Ser needed for biomass synthesis. The PSER pathway for Ser synthesis was found to be dispensable, as it was found to carry no flux.

In agreement with the core model predictions, malic enzyme (ME) was found to be dispensable and pyruvate was produced through pyruvate kinase. This departure from other cyanobacterial species, where the major route for pyruvate synthesis is via ME (Young et al., 2011; Jazmin et al., 2017). Consistent with the core model predictions, the tricarboxylic acid cycle was incomplete and exhibited a bifurcated topology, since there was no flux through 2-oxoglutarate decarboxylase and succinic semialdehyde dehydrogenase. This organism lacks the γ-aminobutyric acid shunt (Mueller et al., 2017). The flux and directionality of malate dehydrogenase (MDH) remain unresolved in this study.

Figure 3. Comparison of the flux ranges (95% confidence interval) for relevant reactions obtained using the core model (Abernathy et al., 2017) and imSyu593 under photoautotrophic conditions. Green bars correspond to the core model, and brown bars correspond to imSyu593. The left end of each bar indicates the lower bound, and the right end of each bar indicates the upper bound. The fluxes are normalized to a carbon uptake of 100 mmol g$^{-1}$ dry weight (DW) h$^{-1}$. See Figure 1 for abbreviations and details of individual reactions.
The genome annotation did not identify any gene that codes for MDH; therefore, this organism may not contain this enzyme. However, earlier labeling experiments with this species have shown the incorporation of carbons from malate into the amino acid Asp derived from oxaloacetate, suggesting the existence of either a previously uncharacterized MDH (Abernathy et al., 2017) or a pathway facilitating the conversion of malate to oxaloacetate.

The secretion rates of organic acids were negligible, and carbon loss in the form of CO2 evolution was insignificant. This implies that CO2 released in photosynthesis and peripheral metabolism was incorporated into the biomass through the Rubisco carboxylase reaction. Of the 100 mmol of bicarbonate taken up by *Synechococcus* 2793, 8 mmol was fixed by phosphoenolpyruvate carboxylase, indicating that at least 92 mmol should be fixed by Rubisco. However, Rubisco was found to fix 111 mmol of CO2, which exceeded the expected flux by 19 mmol. The excess CO2 fixed accounted for the 11 mmol of CO2 generated in the photosynthetic pathway and the 8 mmol of CO2 generated in growth-coupled anabolic pathways. The net carbon balance indicated that there is a nearly complete conversion (greater than 96%) of assimilated carbon into biomass, significantly higher than what was obtained for *Synechocystis* 6803 (86%). The biomass yield was predicted to be 2.48 g dry weight dmo1-1 bicarbonate uptake. This is 2% higher than the predicted biomass yield in *Synechocystis* 6803 (Gopalakrishnan et al., 2018).

**DISCUSSION**

In this study, intracellular fluxes of the fast-growing cyanobacterium *Synechococcus* 2973 were recomputed using the recently developed genome-scale INST-MFA procedure (Gopalakrishnan et al., 2018), the dynamic labeling distribution of 13 intracellular metabolites (Abernathy et al., 2017), and the constructed GSMM imSyu593. Comparison with imSyn617 showed that imSyu593 is more exhaustive in terms of its coverage of cyanobacterial metabolism by accounting for carbon paths formed by pathways not included in imSyn617, such as the PKetolase pathway. The statistically significant improvement in the fit, with an F statistic of 2.079 (*P* = 1.39 × 10-8), indicates that the extra carbon paths included in imSyu593 are necessary to explain the labeling data.

The complete reliance on the TAL reaction for the synthesis of s7p instead of the SBA/SBPase route is a striking deviation from the core model prediction. The use of TAL enzyme explains the observed faster labeling of s7p over that of sbp (Supplemental Fig. S1) in the labeling data (Abernathy et al., 2017). In the SBA/SBPase route, sbp is formed before s7p and sbp would label faster than s7p. Although a larger pool size for sbp would slow down its labeling, the resultant slow turnover of sbp would, in turn, slow down the labeling of all central carbon metabolites, since sbp is present in the Calvin-Benson-Bassham pathway (Nöh et al., 2007; McAtee et al., 2015). Thus, sbp is unlikely to have a larger pool size in this case, as we do not see this effect in the labeling data for downstream metabolites. In contrast, s7p is synthesized before sbp via TAL. However, under photoautotrophic conditions, the enzyme level of TAL is much smaller than that of SBA and SBPase (Yu et al., 2015). A faster labeling of s7p over sbp along with a lower TAL level can be explained with a metabolic channel resulting from the formation of a multi-enzyme complex consisting of SBA and SBPase. In such a multi-enzyme complex, there will be direct labeling of the s7p pool, bypassing sbp, since the product of the first enzyme, sbp, would be transferred directly to the second enzyme without labeling the sbp pool in the cytoplasm. A relatively higher flux through such a metabolic channel also might result in faster labeling of s7p compared with sbp with low levels of TAL. This explains both the labeling dynamics as well as the existing proteomics data (Yu et al., 2015); however, there is no direct experimental evidence for such a channel. In contrast, *Synechocystis* 6803 used the SBA/SBPase route to produce s7p and TAL to produce f6p from s7p and glyceraldehyde-3-phosphate (Fig. 4). This indicates that, even though the set of reactions in the Calvin cycle is conserved among cyanobacteria, the configuration in which they operate differs between species.

PKetolase was reported to be active under mixotrophic conditions in cyanobacteria (Alagesan et al., 2013; Xiong et al., 2015). However, there are no reports on the activity of the native enzyme under the photoautotrophic conditions. Nevertheless, heterologous PKetolase enzymes were shown to increase the intracellular acetyl-CoA levels under photoautotrophic conditions in *Synechocystis* 6803 (Anfelt et al., 2015). Flux balance analysis predicted this reaction to be active under photoautotrophic conditions (Yu et al., 2012; Knoop et al., 2013). Interestingly, this pathway was not included in the mapping models of INST-MFA studies targeting the photoautotrophic metabolism of cyanobacteria. In this study, the f6p-utilizing variant of this pathway was found to be the major route for acetyl-CoA synthesis. The PKetolase route generates acetyl-CoA from two molecules of CO2, whereas the glycolysis route via PDH requires three CO2 molecules. This higher conservation of the fixed carbon reduces the number of CO2 molecules that need to be fixed by Rubisco to produce 1 mol of acetyl-CoA and confers both a kinetic and an energetic advantage for the production of acetyl-CoA-based products (Anfelt et al., 2015). Indeed, the overexpression of heterologous PKetolase enzyme has been shown to enhance the production of acetyl-CoA-based products such as n-butanol (Anfelt et al., 2015) in *Synechocystis* 6803 and acetone (Chwa et al., 2016) in *Synechococcus* 7942. The reliance of the wild-type *Synechococcus* 2973 on this pathway for the synthesis of acetyl-CoA makes it inherently suitable for the production of the above biochemicals.
and PDH, and the usage of nondecarboxylating reactions such as PKetolase. A direct comparison with the biomass yield and growth rate of *Synechocystis* 6803 under its optimal growth conditions would enable us to quantify the relative contribution of the observed carbon-efficient metabolism to the growth rate difference between the two species. Such a comparative analysis between fast- and slow-growing species would shed light on the metabolic factors that might contribute to the faster growth rates. The carbon-efficient metabolism of *Synechococcus* 2973 resulted in a 2% higher biomass yield over that of *Synechocystis* 6803 (Gopalakrishnan et al., 2018). This difference in the biomass yield accounted for only 2.9% of the difference in the growth rate between *Synechococcus* 2973 (0.33 ± 0.05 h⁻¹; Abernathy et al., 2017) and *Synechocystis* 6803 (0.09 h⁻¹; Young et al., 2011). Thus, most of the difference in the growth rate between these two species is due to the higher CO₂ uptake rate in *Syneccoccus* 2973 (12.2 mmol g⁻¹ dry weight h⁻¹; Abernathy et al., 2017) compared with that of *Synechocystis* 6803 (3.7 mmol g⁻¹ dry weight h⁻¹; Young et al., 2011). This higher rate of CO₂ fixation in *Syneccoccus* 2973 requires a higher rate of ATP and NADPH generation. Studies guided by comparative genomics of *Synechococcus* 2973 and *Synechococcus* 7942 have shown that the single-nucleotide polymorphisms in ATP synthase and NAD⁺ kinase genes lead to high rates of ATP and NADPH generation in *Syneccoccus* 2973 and that this contributes significantly to its faster growth rate (Ungerer et al., 2018b). This high rate of energy currency generation requires very high light intensities, and the optimal light intensity for *Syneccoccus* 2973 (500 μE m⁻² s⁻¹) is higher than that for *Synechocystis* 6803 (300 μE m⁻² s⁻¹; Yu et al., 2015).

This meta-analysis using genome-scale INST-MFA showed that the fast-growing cyanobacterium *Synechococcus* 2973 possesses a more carbon-efficient metabolism when compared with that of *Synechocystis* 6803 and pointed out certain key features that contribute to this characteristic. Repeating the analysis on mutants lacking some of these features (e.g., PKetolase) will further improve our understanding about their role in increasing the carbon efficiency of photosynthetic metabolism. Also, it would be worthwhile to compare the genome-scale flux distribution of *Syneccoccus* 2973 with that of the slow-growing close relative *Synechococcus* 7942 to elucidate the metabolic impact of the 55 single-nucleotide polymorphisms that exist between the genomes of these two strains (Yu et al., 2015). A comparison with the core model flux distribution of *Synechococcus* 7942 (Abernathy et al., 2017) revealed some interesting differences in central carbon metabolism. *Synechococcus* 7942 had zero flux through the photosynthetic pathway, whereas *Synechococcus* 2973 displayed a significant amount of photosynthesis. There was a higher flux through the Calvin cycle in *Synechococcus* 2973 compared with *Synechococcus* 7942 in order to replenish the ribulose-1,5-bisphosphate lost in photorespiration. *Synechococcus* 2973 used TAL for...
the synthesis of s7p, whereas *Synechococcus* 7942 made use of the SBA/SBPass route. Since the PKetolase pathway was not included in the core model of *Synechococcus* 7942, there was a higher flux through glycolysis compared with that in *Synechococcus* 2973. Interestingly, both *Synechococcus* 2973 and *Synechococcus* 7942 used the pyruvate kinase enzyme instead of the ME for pyruvate production. Unfortunately, further comparison cannot be carried out, since the available *Synechococcus* 7942 fluxes are generated using the core model. Comparison of genome-scale flux distribution for these two species will reveal several other interesting differences. Since our analysis used the transition labeling data for a limited number of metabolites, the flux through certain reactions such as MDH could not be resolved. Usage of an expanded repertoire of metabolite-labeling data would resolve these uncertainties. The study used FVA to identify the active set of reactions in the GSM, and these reactions formed the metabolic network used for the genome-scale INST-MFA. However, a more precise way of identifying the active set of reactions would be to use the transcriptome and proteome data generated under similar conditions. The data also could act as additional constraints resulting in the enhanced resolution of the fluxes.

**MATERIALS AND METHODS**

**Construction of imSyu593**

The GSM used for flux elucidation was adapted from Syu683 (Mueller et al., 2017) upon eliminating reactions incapable of carrying metabolic flux and reactions involved in loops by performing FVA (Mahadevan and Schilling, 2003) constrained by the measured growth rate and bicarbonate uptake rate (Supplemental Table S2; Abernathy et al., 2017). The genes involved in the PSER pathway were identified recently in *Synechocystis* sp. PCC 6803 (Klenke et al., 2015). BLAST analysis showed that the corresponding homologs are in *Synechococcus elongatus* UTEX 2973; therefore, this pathway was included in the model. Although Syu683 does not include the MDH reaction, metabolite-labeling data suggest the presence of flux through this reaction (Abernathy et al., 2017), which led to the inclusion of MDH in the GSM and GSM. The final GSM used for the analysis contained 700 reactions (Supplemental Table S3) and 667 metabolites (Supplemental Table S4). Apart from the metabolic reactions, reactions modeling the mixing of metabolically active and inactive pools were included for all the fitted metabolites. The GSMM for *Synechococcus* 2973, namely imSyu593, traces the carbon transitions through 593 reactions (Supplemental Table S5) in the GSM. The remaining reactions were outside the purview of the EMU balances and were not mapped, since they do not contribute to the labeling of the fitted metabolites. Carbon mapping for 519 shared reactions with *Synechocystis* 6803 were imported from the GSMM imSyu593 (Gopalakrishnan et al., 2018). Twenty-one reactions, mostly from fatty acid biosynthesis pathways, were mapped manually based on the reaction mechanism and the mapping information available in MetaCyc (Caspi et al., 2018). Mapping information for the remaining 50 reactions is unavailable in the literature and, therefore, was generated using the Canonical Labeling for Cliques Approximation algorithm and verified manually (Kumar and Maranas, 2014).

**Flux Estimation and Statistical Analysis of the Fit**

The labeling data for flux elucidation were obtained from a previous study (Abernathy et al., 2017) where *Synechococcus* 2973 was grown under photoautotrophic conditions supporting optimal growth and 100% [13C]bicarbonate was used for the dynamic carbon-labeling experiment. The transition labeling data of the following 13 metabolites (Supplemental Table S6) were used for flux estimation: 2-phosphoglycerate, 6-phosphogluconate, r5p, G6c-6-P, phosphoenolpyruvate, s7p, succinate, malate, 3-phosphoglycerate, f6p, citrate, sbp, and Fru bisphosphate. The dynamic labeling data included the mass isotope distributions (MIDs) at time points 20, 30, 40, 60, 90, 150, 300, 600, 1,200, and 3,600 s. Scanned bicarbonate uptake rate (10 mmol g⁻¹ dry weight h⁻¹) and growth rate (0.27 h⁻¹) also were fitted along with the labeling dynamics. Flux elucidation was performed as described earlier (Gopalakrishnan et al., 2018). This methodology uses the EMU-based decomposition of the carbon-mapping model to reduce the number of balance equations (ordinary differential equations [ODEs]) required to simulate the fragment MIDs (Antoniewicz et al., 2007). There were 1,271 EMUs after decomposing the imSyu593 to track the labeling of 13 metabolites that were fitted. This resulted in a total of 3,813 EMU mass fractions that need to be simulated to capture the labeling dynamics of the fitted metabolites. The ODEs capturing the labeling dynamics are then solved using Richardson’s extrapolation integrators after discretizing them in the time domain. A modified Levenberg-Marquardt algorithm was used to solve the nonlinear least squares problem that finds the genome-scale flux distribution that closely predicts the experimentally measured labeling dynamics. The numbers of free fluxes and pool sizes in the model were 202 and 217, respectively. Thus, for every step taken in the parameter space, a total 15.6 × 10⁵ ODEs were solved to simulate 3,813 EMU fractions and calculate their sensitivity to 419 parameters. The statistical significance of the final fit was assessed by subjecting the SSR to a χ² test (Antoniewicz et al., 2006). The χ² test was used to check the statistical significance of the improvement in the fit obtained using the GSMM over the core model. The 95% confidence interval for the estimated fluxes was calculated by determining the sensitivity of the minimum SSR as a function of individual fluxes (Antoniewicz et al., 2006). The estimated fluxes and confidence intervals are available in Supplemental Table S7.

**Supplemental Data**

The following supplemental materials are available.

- Supplemental Figure S1. Comparison of the relative fraction of M+0 isotopomer in sbp and s7p at various time points.
- Supplemental Table S1. Comparison of flux ranges generated using MFA and FVA.
- Supplemental Table S2. Experimental values of parameters such as growth rate and CO₂ uptake.
- Supplemental Table S3. The list of reactions part of the imSyu593.
- Supplemental Table S4. The list of metabolites part of the imSyu593.
- Supplemental Table S5. The atom-mapping information for imSyu593.
- Supplemental Table S6. The MIDs of metabolites used for flux estimation.
- Supplemental Table S7. Fluxes and confidence intervals estimated using imSyu593 in this study.

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**LITERATURE CITED**


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