Carboxysome Mispositioning Alters Growth, Morphology, and Rubisco Level of the Cyanobacterium *Synechococcus elongatus* PCC 7942

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**ABSTRACT** Cyanobacteria are the prokaryotic group of phytoplankton responsible for a significant fraction of global CO$_2$ fixation. Like plants, cyanobacteria use the enzyme ribulose 1,5-bisphosphate carboxylase/oxidase (Rubisco) to fix CO$_2$ into organic carbon molecules via the Calvin-Benson-Bassham cycle. Unlike plants, cyanobacteria evolved a carbon-concentrating organelle called the carboxysome—a proteinaceous compartment that encapsulates and concentrates Rubisco along with its CO$_2$ substrate. In the rod-shaped cyanobacterium *Synechococcus elongatus* PCC 7942, we recently identified the McdAB system responsible for uniformly distributing carboxysomes along the cell length. It remains unknown what role carboxysome positioning plays with respect to cellular physiology. Here, we show that a failure to distribute carboxysomes leads to slower cell growth, cell elongation, asymmetric cell division, and elevated levels of cellular Rubisco. Unexpectedly, we also report that even wild-type *S. elongatus* undergoes cell elongation and asymmetric cell division when grown at the cool, but environmentally relevant, growth temperature of 20°C or when switched from a high- to ambient-CO$_2$ environment. The findings suggest that carboxysome positioning by the McdAB system functions to maintain the carbon fixation efficiency of Rubisco by preventing carboxysome aggregation, which is particularly important under growth conditions where rod-shaped cyanobacteria adopt a filamentous morphology.

**IMPORTANCE** Photosynthetic cyanobacteria are responsible for almost half of global CO$_2$ fixation. Due to eutrophication, rising temperatures, and increasing atmospheric CO$_2$ concentrations, cyanobacteria have gained notoriety for their ability to form massive blooms in both freshwater and marine ecosystems across the globe. Like plants, cyanobacteria use the most abundant enzyme on Earth, Rubisco, to provide the sole source of organic carbon required for its photosynthetic growth. Unlike plants, cyanobacteria have evolved a carbon-concentrating organelle called the carboxysome that encapsulates and concentrates Rubisco with its CO$_2$ substrate to significantly increase carbon fixation efficiency and cell growth. We recently identified the positioning system that distributes carboxysomes in cyanobacteria. However, the physiological consequence of carboxysome mispositioning in the absence of this distribution system remains unknown. Here, we find that carboxysome mispositioning triggers changes in cell growth and morphology as well as elevated levels of cellular Rubisco.

**KEYWORDS** Rubisco, carbon dioxide assimilation, carbon dioxide concentration mechanism, carbon dioxide fixation, carboxysome, cell division, cyanobacteria, filamentation

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As the evolutionary ancestor of algae and plant chloroplasts, all cyanobacteria perform oxygenic photosynthesis and fix carbon dioxide through the Calvin-Benson-Bassham cycle. Unlike chloroplasts, cyanobacteria encapsulate their ribulose 1,5-bisphosphate carboxylase/oxidase (Rubisco) and carbonic anhydrase within large selectively permeable protein-based organelles called carboxysomes. This mechanism generates an environment around Rubisco that is significantly enriched in CO2, which increases the carboxylation activity of Rubisco, while simultaneously reducing photorespiration (1).

In the model rod-shaped cyanobacterium *Synechococcus elongatus* PCC 7942 (hereafter *S. elongatus*), carboxysomes were found to be uniformly distributed down the length of individual cells (2). This equidistant positioning supports equal inheritance of carboxysomes following cell division and may assist in maximizing the diffusion of substrates and products across the carboxysome shell. Carboxysomes are essential for the growth and survival of all cyanobacteria and are responsible for ~35% of global carbon fixation through atmospheric CO2 assimilation (3, 4). However, it remains unknown how the subcellular organization of carboxysomes influences cyanobacterial physiology, a question of considerable ecological, evolutionary, and biotechnological importance.

Savage et al. were first to report that a ParA-type ATPase (hereafter McdA [for maintenance of carboxysome distribution A]) is required for positioning carboxysomes in *S. elongatus* (2). ParA family members have well-established roles in the segregation of bacterial chromosomes and plasmids (5, 6). Less studied are ParA family members shown to be required in the positioning of diverse protein complexes, such as those involved in secretion (7, 8), chemotaxis (9–11), conjugation (12), cell division (13, 14), and cell motility (15, 16), as well as bacterial microcompartments (BMCs), such as the carboxysome (2, 17). We recently identified a small novel protein, McdB, responsible for generating dynamic McdA gradients on the nucleoid (17). McdB colocalizes and directly interacts with carboxysomes and removes McdA from the nucleoid in their vicinity. We proposed that carboxysomes use a Brownian-ratchet mechanism whereby McdB-bound carboxysome motion occurs in a directed and persistent manner toward increased concentrations of McdA on the nucleoid. We also recently found that the McdAB system is widespread among β-cyanobacteria (18) and carbon-fixing proteobacteria (19). Together, the findings suggest that the equidistant positioning of carboxysomes in carbon-fixing bacteria is important, but the physiological consequences of carboxysome mispositioning in the absence of the McdAB system remain unclear.

When Savage et al. first identified the McdA requirement for carboxysome positioning, a minor decrease in CO2 fixation was found in a ΔmcdA strain, but growth at 30°C under ambient CO2 was the only condition studied (2). Rising atmospheric CO2 levels and sea surface temperatures are pervasive effects of climate change (20), and both have direct influences on Rubisco activity and cyanobacterial growth (21, 22). Here, we show that a failure to distribute carboxysomes leads to a temperature-dependent decrease in growth rate, cell elongation, asymmetric cell division, and elevated levels of cellular Rubisco. We also report that, unexpectedly, wild-type *S. elongatus* undergoes filamentous growth when switched from a high- to ambient-CO2 environment or when grown at 20°C, an environmentally relevant growth temperature for *S. elongatus* not commonly used in the lab. We propose that carboxysome positioning by the McdAB system functions as part of an autotrophic growth strategy that maintains the carbon fixation efficiency of Rubisco by preventing carboxysome aggregation. In the absence of carboxysome positioning, we propose the changes in cell growth, morphology, and Rubisco levels are all responses to organic carbon limitation.

**RESULTS**

We performed *in vivo* microscopy to determine how carboxysome organization was altered in *mcdA, mcdB*, and *mcdAB* deletion strains compared to wild-type *S. elongatus*. Immunoblot analysis against McdA and McdB proteins verified the loss of expression from the deletion strains and verified that the remaining *mcdA* or *mcdB* gene was still
expressed at levels similar to that of the wild type (see Fig. S1A in the supplemental material). In S. elongatus, each Rubisco enzyme is composed of eight large (RbcL) and eight small (RbcS) subunits to form RbcL₈S₈ (23). Therefore, to image carboxysomes, the fluorescent protein mTQ (monomeric Turquoise2) was fused to the C terminus of RbcS to make RbcS-mTQ (see Materials and Methods for strain construction details). 

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In the lab, S. elongatus is typically grown at 32°C with a doubling time on the order of several hours depending on growth conditions (temperature, light, and CO₂ availability). This was the temperature used in our previous study that identified the McdAB carboxysome positioning system (17). However, S. elongatus was recently shown to grow faster at 40°C (25, 26). Therefore, in anticipation of a growth defect in the absence of carboxysome positioning in our McdAB system mutants, we began our studies by growing cultures at 40°C in constant light under ambient (0.04%) or high (2%) CO₂ concentrations.

Carboxysome mispositioning correlates with minor defects in cellular physiology at 40°C. We first verified that carboxysomes are organized by the McdAB system at 40°C, as we found previously at 32°C (17). We examined carboxysome positioning under high CO₂ conditions (2% CO₂) to maximize growth. We found that RbcS-mTQ-labeled carboxysomes were equally spaced down the long axis of wild-type S. elongatus cells (Fig. 1A). The ΔmcdA, ΔmcdB, and ΔmcdAB mutants lost this uniform positioning of carboxysomes (Fig. 1B to D), as observed previously at 32°C (17).

We compared the nearest neighbor spacing of carboxysome foci as a function of cell length (Fig. 1E). Wild-type cells showed uniform carboxysome spacing (0.55 ± 0.20 μm) regardless of cell length. All three mutants, on the other hand, displayed an increase in the nearest neighbor spacing of carboxysome foci as cell length increased. Both the median and variability in spacing were greater in the mutants compared to the wild type (Fig. 1F). The increased spacing resulted in fewer carboxysome foci per unit cell length (Fig. 1G), as well as fewer carboxysome foci per cell (Fig. 1H). When quantifying the carboxysome fluorescence intensity (Fig. 1I), it became apparent that the increased spacing in all three mutant populations was not likely a result of fewer carboxysomes being assembled, but rather, carboxysomes were coalescing into massive aggregates. Indeed, when comparing carboxysome focus number across cell populations, we find that ~95% of wild-type cells (n = 440) had three or more foci, whereas ~45% of all mutant populations had three or more foci (Fig. 1J). Roughly 20% of cells from all three mutant populations had a single carboxysome aggregate or no carboxysomes at all, whereas wild-type cells never had less than two foci. The 40°C data show that the McdAB system equally spaces carboxysomes down the cell length and also serves as an antiaggregator, preventing carboxysomes from coalescing.

We then asked whether carboxysome mispositioning affected cell physiology at 40°C. Wild-type S. elongatus cells were 3.1 ± 0.5 μm in length (Fig. 2A) and 1.25 ± 0.05 μm in width (Fig. 2B). We found that ΔmcdA cells were of a similar length but thinner compared to the wild-type cells (Fig. 2A and B). The ΔmcdB and ΔmcdAB cells had a slightly longer median cell length and wider distribution (Fig. 2A). These mutants were also thinner compared to the wild type (Fig. 2B). The presence of both longer and shorter cells suggested asymmetric cell division events in the ΔmcdB and ΔmcdAB cell populations. We quantified the frequency of symmetric (mid-cell) versus asymmetric (non-mid-cell) division events and found that asymmetric division was exclusive to the
$D_{mcdB}$ and $D_{mcdAB}$ populations (Fig. 2C). The data show that while both McdA and McdB are required for positioning carboxysomes, the loss of McdB elicits an asymmetric division phenotype at 40°C.

The changes in cell morphology suggested that, even with high CO$_2$ where carboxysomes are not essential for viability, carboxysome mispositioning may still alter cell growth. Interestingly, a reduction in growth rate was observed for all three mutants, but only when grown at high CO$_2$ (Fig. 2D and F and Fig. S3A). When the cells were grown slowly in ambient CO$_2$ (0.04%), we did not observe significant differences in growth rate compared to the wild type (Fig. 2E and F). Overall, at 40°C, carboxysomes are mispositioned in all three McdAB system mutants, with moderate increases in

**FIG 1** McdAB system mutants display fewer and mispositioned carboxysome aggregates at 40°C. (A to D) Microscopy images of the specified cell strains (wild type [WT] and $D_{mcdA}$, $D_{mcdB}$, and $D_{mcdAB}$ mutants) grown at 40°C in 2% CO$_2$. Phase contrast micrographs are shown in black and white, and carboxysomes are shown in green. The phase contrast channel is blue in the merge. (E) Spacing between carboxysome foci as a function of cell length. (F) Spacing between carboxysome foci in the same cell. (G) Number of carboxysome foci per unit cell length. (H) Carboxysome foci number per cell. (I) Carboxysome peak focus intensity (in arbitrary units [AU]). (J) Population percentages of cells with the specified number of carboxysome foci. $n \geq 1,000$ carboxysomes from 440 cells of each strain.
Carboxysome spacing due to aggregation, which ultimately results in fewer carboxysome foci per cell. We also found an asymmetric cell division phenotype exclusive to strains lacking McdB. However, in all deletion strains, the growth rate was significantly lower than that of the wild type only when high CO2 was provided.

**Carboxysome mispositioning causes cell elongation and asymmetric cell division at 30°C.** We continued our study at 30°C, a growth temperature closer to what we used to first identify the McdAB carboxysome positioning system (17). As we showed previously, and similar to our 40°C data, wild-type *S. elongatus* cells have equally spaced carboxysomes (Fig. 3A), while in all three mutants, carboxysomes were mispositioned (Fig. 3B to D). Wild-type cells displayed the same carboxysome spacing distance (0.50 ± 0.20 μm) regardless of cell length, while all three mutants had increased carboxysome spacing and variability in spacing as cell length increased (Fig. 3E and F). Intriguingly, we found that mutant cell lengths were significantly longer compared to the wild-type length. Cell elongation was more extreme in the ΔmcdB and ΔmcdAB mutants, resulting in more distantly spaced carboxysome foci (Fig. 3F). The data once again show that McdB plays a currently unknown role in carboxysome function, separate from its role in carboxysome positioning with McdA.

The increased spacing resulted in fewer carboxysome foci per unit cell length (Fig. 3G) and fewer carboxysomes per cell (Fig. 3H). Carboxysome foci in all three mutants were significantly higher in intensity than that of wild type, consistent with aggregation (Fig. 3I). While ~90% of wild-type cells (*n* = 486) had three or more foci, ~50% of all three mutant populations had three or more foci (Fig. 3J). Again, ~20% of cells in all three mutant populations contained a single carboxysome aggregate or no
Carboxysomes at all. This is a striking reduction in carboxysome foci when considering the cell elongation phenotype exclusive to the mutant cell lines.

As the carboxysome spacing data suggested, all three mutants had significantly longer cell lengths compared to the wild type when grown at 30°C (Fig. 4A). Median cell width was similar across all strains; however, \( \Delta mcdB \) and \( \Delta mcdAB \) cell populations displayed significantly wider distributions (Fig. 4B). Once again, the \( \Delta mcdB \) and \( \Delta mcdAB \) mutant populations displayed a significant number of asymmetrical division events (~70% of dividing cells) compared to the wild-type or \( \Delta mcdA \) strain (Fig. 4C). We propose that carboxysome mispositioning and aggregation at 30°C elicits cell division arrest, cell elongation, and when McdB is absent, asymmetric cell division; phenotypes that were less apparent when cells were grown at 40°C.

**FIG 3** McdAB system mutants house few and mispositioned carboxysome aggregates at 30°C. (A to D) Microscopy images of the specified cell strains grown at 30°C in 2% CO₂. Phase contrast micrographs are shown in black and white, and carboxysomes are shown in green. The phase contrast channel is blue in the merge. (E) Spacing between carboxysome foci as a function of cell length. (F) Spacing between carboxysome foci in the same cell. (G) Number of carboxysome foci per unit cell length. (H) Carboxysome foci number per cell. (I) Carboxysome peak foci intensity (in arbitrary units [AU]). (F to I) Solid bars represent the median and the 95% confidence interval. Statistical significance was based on a nonparametric Mann-Whitney test and indicated as follows: ***, \( P < 0.001 \). (J) Population percentages of cells with the specified number of carboxysome foci (\( n = 1,000 \) carboxysomes from 440 cells of each strain).
The cell elongation phenotype suggested that the mutants may also display lower growth rates, even when grown at high CO₂. Indeed, with high CO₂, a significant reduction in growth rate was observed for all three mutants compared to the wild type (Fig. 4D and F and Fig. S3B). However, once again, when cells were grown in ambient CO₂, there was little to no reduction in growth rate compared to the wild type (Fig. 4E and F).

Overall, we find that at 30°C, carboxysomes are mispositioned in the ΔmcdA, ΔmcdB, and ΔmcdAB mutants with drastic increases in carboxysome spacing due to aggregation, resulting in fewer carboxysome foci per cell. We unveiled a cell elongation phenotype for all three mutant populations, and strikingly, we also found an asymmetric cell division phenotype that was exclusive to cells lacking McdB, a phenotype also found at 40°C but exacerbated at 30°C (compare Fig. 2C and 4C). Despite these carboxysome aggregation and cell morphology phenotypes, only minor decreases in growth rate were observed, and only under high CO₂ conditions. We propose that carboxysome aggregation decreases the carbon-fixing activity of encapsulated Rubisco, which triggers a carbon limitation response that results in cell elongation and asymmetric cell division.

Wild-type *S. elongatus* elongates at 20°C and growth is slower in McdAB system mutants. A 10 degree drop in growth temperature unveiled the physiological consequences of carboxysome mispositioning at 30°C, which were largely absent at 40°C. The catalytic rate of carboxysome-encapsulated Rubisco is often considered the bottle neck of photosynthesis because the enzyme is inefficient and temperature dependent, whereas the light reactions of photosynthesis are temperature independent (27–29). *S. elongatus* PCC 7942 was originally isolated from a freshwater source in the San Francisco Bay area, with an annual temperature range of 8°C to 25°C (30).
therefore studied the effects of carboxysome mispositioning when cells were grown at 20°C, which is within the environmentally relevant temperature range.

Unexpectedly, we found that even wild-type *S. elongatus* undergoes extreme cell elongation when grown at 20°C (Fig. 5A). *S. elongatus* has been previously studied at this temperature (31), but to our knowledge, this is the first time its temperature-dependent cell elongation phenotype has been directly observed and reported. Despite the remarkably long cell lengths, carboxysomes were still well aligned down the entire longitudinal axis of the cell (Fig. 5A). All three mutants were also elongated, but to a lesser extent, and carboxysomes were clearly mispositioned (Fig. 5B to D).

**FIG 5** At 20°C, wild-type and McdAB system mutants elongate, but only the mutants display few and mispositioned carboxysome aggregates. (A to D) Microscopy images of the specified cell strains grown at 20°C in 2% CO₂. Phase contrast micrographs are shown in black and white, and carboxysomes are shown in green. The phase contrast channel is blue in the merge. (E) Spacing between carboxysome foci as a function of cell length. (F) Spacing between carboxysome foci in the same cell. (G) Number of carboxysome foci per unit cell length. (H) Carboxysome foci number per cell. (I) Carboxysome peak foci intensity (in arbitrary units [AU]). (F to I) Solid bars represent the median and the 95% confidence interval. Statistical significance was based on a nonparametric Mann-Whitney test and indicated as follows: ***, P < 0.001. (J) Population percentages of cells with the specified number of carboxysome foci (*n* = 1,000 carboxysomes from 440 cells of each strain).
The wild type showed the same uniform carboxysome spacing distance (0.50 ± 0.30 µm) even in cells as long as 20 µm (Fig. 5E). All three mutants, on the other hand, displayed dramatically increased carboxysome spacing and variability in spacing as cell length increased (Fig. 5F). Since all cell types dramatically elongated at 20°C, the loss of carboxysome positioning resulted in a massive reduction in the number of carboxysome foci per unit cell length (Fig. 5G) and per cell (Fig. 5H). Carboxysome foci in all three mutants were significantly larger than that of the wild type, once again suggesting aggregation (Fig. 5I). While 90% of wild-type cells (n = 388) had six or more foci, only 20% of all three mutant populations had six or more foci (Fig. 5J). The data emphasize the importance of the McdAB system in positioning carboxysomes in cells that are dramatically elongated when grown at cool, but environmentally relevant temperatures.

We quantified cell length at 20°C and found that the median length of wild-type cells was significantly longer than those for all three mutant populations (Fig. 6A). The ΔmcdA cells were similar in width to that of the wild type, while ΔmcdB and ΔmcdAB cell populations were notably thinner (Fig. 6B). Consistent with the wide distributions in cell length (Fig. 6A), a significant fraction of division events at this growth temperature were asymmetric across all cell populations, including the wild type (Fig. 6C). However, the frequency of asymmetric division events was still highest in mutants lacking McdB. When growth rates were assayed at high CO2 (Fig. 6D and F) or ambient CO2 (Fig. 6E and F), we found statistically significant reductions in growth rate under both conditions for all three mutants compared to the wild type.

**FIG 6** At 20°C, wildtype and McdAB mutants elongate and undergo asymmetric cell division, but only McdAB system mutants show lowered growth rates. (A and B) Cell lengths (A) and widths (B) of the WT population compared to McdAB system mutants. Cells were grown at 20°C in 2% CO2. Statistical significance was based on a nonparametric Mann-Whitney test and indicated as follows: ***, P < 0.001; **, P < 0.01. (C) Percentage of cells undergoing asymmetric (non-mid-cell) division (n ≥ 440 dividing cells of each strain). (D and E) Quantification of mean exponential growth rate in 2% CO2 (D) or no CO2 added (E). (F) Quantification of dry cell biomass for cells grown in 2% CO2 or no CO2 added. In panels C to F, error bars represent the standard deviations from at least three independent biological replicates. Statistical significance was based on an unpaired t test and indicated as follows: **, P < 0.01; *, P < 0.05; n.s., not significant.
Overall, we identified a cell elongation phenotype in wild-type *S. elongatus* that occurs when grown at colder, but environmentally relevant temperatures. All three mutants elongated to a lesser extent and had aggregated carboxysomes, which resulted in drastically fewer carboxysome foci per cell compared to the wild type. Only at this cooler temperature, we found significant decreases in growth rate with or without CO\(_2\) added. We speculate that colder growth temperatures decrease the carbon fixation activity of Rubisco to a point where cell division arrest, elongation, and asymmetric cell division are triggered in response to carbon limitation, even in wild-type cells. At 30°C, this response was found only in the McdAB system mutants likely due to reduced carbon fixation efficiency resulting from carboxysome aggregation. At 40°C, higher Rubisco activity compensates for carboxysome aggregation, which would explain why McdAB system mutants displayed cell morphologies and growth rates closer to those of the wild type.

**McdAB mutants have elevated levels of cellular Rubisco.** The Rubisco encapsulated in carboxysomes is the sole enzyme providing organic carbon for cellular biomass production and the phototrophic growth of *S. elongatus*. We set out to determine whether carboxysome mispositioning and the changes in cell morphology of McdAB system mutants correlated with altered cellular levels of Rubisco using immunoblot analysis against the large subunit of Rubisco (RbcL) (Fig. S4). RbcL content from cell cultures was normalized based on Atp\(_\beta\) quantity from immunoblot analysis as shown previously (32, 33). We used high CO\(_2\) when assaying the McdAB system mutants, as this was the growth condition that showed the strongest physiological defects across all temperatures tested. At 40°C, Rubisco abundance in the mutant strains was not significantly different than that of wild type (Fig. 7A). However, at 30°C, Rubisco levels were three- to sixfold higher in all three mutants compared to the wild type (Fig. 7B). This trend was also observed at 20°C (Fig. 7C). Overall, our findings show that carboxysome aggregation correlates with a temperature-dependent increase in Rubisco abundance in the cell.

**Wild-type *S. elongatus* cells elongate, asymmetrically divide, and have elevated Rubisco levels in response to CO\(_2\) down-shift.** Finally, we set out to determine whether cell elongation and asymmetric cell division were consequences of reduced carbon fixation by performing CO\(_2\) down-shift assays on wild-type *S. elongatus*. Similar to our McdAB deletion mutants, we found that wild-type cells had increased levels of Rubisco at higher growth temperatures and when grown under ambient CO\(_2\) compared to when grown at high CO\(_2\) (Fig. 7D). The CO\(_2\) down-shift caused wild-type cells to elongate (Fig. 7E) and undergo asymmetric cell division, compared to cells maintained at high CO\(_2\) (Fig. 7F). Together, the data are consistent with our proposal that carboxysome aggregation in McdAB system mutants results in reduced carbon fixation activity by encapsulated Rubisco and a carbon limitation response—cell elongation, asymmetric cell division, and elevated levels of Rubisco.

**DISCUSSION**

We recently identified the McdAB system that is responsible for the equidistant positioning of carboxysomes in the rod-shaped cyanobacterium *S. elongatus* PCC 7942 (17). We also recently found that McdAB systems are widespread among β-cyanobacteria (18) and carboxysome-containing proteobacteria (19). The findings suggest important and widespread, but currently unknown, physiological roles for carboxysome positioning in carbon-fixing bacteria. Here, we show that a failure to distribute carboxysomes in *S. elongatus* leads to several changes in cell physiology: slower growth, cell elongation, asymmetric cell division, and a significant increase in cellular levels of Rubisco.

At the three growth temperatures tested (20, 30, and 40°C), all three mutants housed few and irregularly spaced carboxysome aggregates, whereas wild-type cells have uniformly spaced and sized carboxysome foci. The data provide quantitative support for our previous finding that the McdAB system equally distributes carboxysomes to opposite sides of the cell to ensure inheritance following cell division (2, 17), akin to ParA-based plasmid partition systems in bacteria (5). However, in addition, and
particularly important for protein-based cargoes, the McdAB system serves to prevent carboxysome aggregation. This “antiaggregation” activity serves as a homeostasis mechanism that regulates carboxysome size, number, composition, positioning, and ultimately, its carbon-fixing function in the cell. It has recently been found that the pyrenoid, the functional analog of the carboxysome in chloroplasts of the model alga *Chlamydomonas*, is also spatially regulated and this activity affects carbon fixation (34).

Physiological defects associated with carboxysome mispositioning are temperature dependent. Despite the drastic mispositioning and aggregation of carboxysomes in the mcdA, mcdB, and mcdAB deletion strains of *S. elongatus*, only minor changes in cell morphology and growth rate were found at the highest temperature used in this study (40°C) (Fig. 2). At 30°C, however, all three mutants displayed a cell elongation phenotype, and a significant fraction of cell division events in the mutant populations lacking McdB were asymmetric (Fig. 4). Along with these changes in cell morphology, the growth rates of all three mutants were lower than that of wild type, but only at high CO₂. At our coldest but environmentally relevant growth temperature of 20°C, we found that all cell populations, even the wild type, underwent filamentous growth and asymmetric cell division (Fig. 6). Wild-type cells at 20°C were as much as
10 times longer than the median length when grown at 30°C or 40°C. Even in these extremely elongated cells, the McdAB system robustly distributed carboxysomes down the entire cell length (Fig. 5A). This mode of filamentous growth was slower in all McdAB system mutants, resulting in significantly shorter filaments compared to the wild type (Fig. 6A). Overall, we found temperature-dependent physiological defects associated with carboxysome mispositioning and aggregation, phenotypes that were masked at higher temperatures typically used in the lab, and unveiled here at lower but environmentally relevant temperatures.

**Cell elongation and asymmetric cell division—a stress response to carbon limitation?** Many bacteria can change shape in response to growth conditions (35). *Escherichia coli* and *Bacillus subtilis* produce larger cells under nutrient-rich conditions and smaller cells under nutrient-limited conditions (36, 37), *Pseudomonas aeruginosa* elongates to enhance nutrient uptake during carbon and nitrogen starvation (38), *Caulobacter crescentus* cells increase cell area in response to phosphate starvation (39), and abundant human gut species such as *Bacteroides thetaiotaomicron* elongate under sugar-limited conditions (40). We propose that *S. elongatus* filamentation and asymmetric cell division are responses triggered by carbon limitation, resulting from reduced carbon fixation activity of Rubisco in this obligate photoautotroph. Cell filamentation (i) suppresses the birth of cells devoid of carboxysomes, (ii) gives the cell an opportunity to increase carbon fixation by producing more Rubisco, and (iii) increases cell surface area, which would increase the light-harvesting capability for photochemistry.

A carbon limitation response parsimoniously explains the three triggers identified here: (i) carboxysome aggregation, (ii) colder growth temperatures, and (iii) CO₂ downshift. For the carboxysome aggregation trigger, carbon limitation may result from a decrease in Rubisco efficiency due to losses in carboxysome integrity, substrate/product permeability, and/or enzyme stability. Our data suggest that all McdAB system mutants respond by significantly increasing the total cellular Rubisco content (Fig. 7B and C). We propose that this carbon limitation response can also be triggered in wild-type *S. elongatus* by low-temperature growth. At high temperature (i.e., 40°C), Rubisco activity may be sufficient to compensate for carboxysome aggregation in the McdAB system mutants. Therefore, the carbon limitation response is not triggered. At 30°C, Rubisco activity is still high enough to prevent a carbon limitation response in wild-type cells, but not in the McdAB system mutants with aggregated carboxysomes. As a result, cell filamentation is triggered only in the mutants. At 20°C, we speculate that Rubisco activity has reduced to a level that, even in wild-type cells, triggers a carbon limitation response. At this temperature, McdAB system mutants are starved for carbon to a point where growth is slowed, resulting in shorter filaments compared to the wild type (Fig. 6A). We were not able to perform *in vivo* carbon fixation assays to show a direct link between carbon fixation activity and the mutant phenotypes observed. However, we did find that wild-type *S. elongatus* cells elongate, asymmetrically divide, and increase Rubisco levels when shifting cultures from a high- to low-CO₂ environment. The identical phenotypes implicate carbon limitation as the trigger for these physiological responses.

**Spatial regulation of carboxysomes influences Rubisco activity and abundance.** Previous studies have shown positive relationships between growth rate and Rubisco abundance (41–46). Elevated carbon fixation rates during blooms, for example, have been shown to be associated with severalfold increases in Rubisco. The 20°C temperature is well below the thermal optimum for Rubisco activity (47) and well below the optimal growth temperature of *S. elongatus*. We propose that the increased level of cellular Rubisco in the McdAB system mutants and the associated cell elongation phenotype both represent acclimation responses that compensate for the decreased activity of aggregated Rubisco at low-temperature growth. It is important to emphasize that cell elongation, lowered growth rate, and increased Rubisco levels were found when the cells were grown at high CO₂—a condition whereby *S. elongatus* does not require the carbon-concentrating activity of carboxysomes for growth (48). This strengthens the argument that it is not CO₂ substrate limitation causing these
phenotypes. Instead, it is Rubisco activity that is significantly compromised when complexed in carboxysome aggregates. These findings highlight the physiological importance of carboxysome antiaggregation by the McdAB system in maximizing the carbon fixation activity of Rubisco.

Overall, we show that carboxysome distribution by the McdAB system in cyanobacteria can appropriately respond to variability in Rubisco abundance and activity along a wide temperature gradient. We speculate that this mechanism of carboxysome homeostasis provided by the McdAB system is part of an autotrophic growth strategy particularly in elongated cells—Rubisco is uniformly distributed into homogeneously sized carboxysomes at low temperature to overcome the lower activity of this temperature-dependent enzyme (46).

McdB plays a role in carboxysome function outside of its positioning with McdA. Regardless of growth temperature, \( \Delta mcdB \) and \( \Delta mcdAB \) mutants displayed a stronger asymmetric-division phenotype compared to the \( \Delta mcdA \) mutant. The finding suggests that McdB plays a currently unknown but critical role in carboxysome function, outside of its role in positioning carboxysomes with McdA. Consistently, our recent bioinformatics analysis of McdAB systems across cyanobacteria (18) and proteobacteria (19) identified numerous species with orphan McdB proteins, once again suggesting a functional role independent of McdA. We have also shown previously via bacterial two-hybrid analysis that McdA does not physically associate with any carboxysome component, while McdB directly interacts with a number of shell proteins (17). We have also recently found that purified McdB undergoes liquid-liquid phase separation (LLPS) \textit{in vitro} (18). This activity is intriguing given recent studies showing that both \( \alpha \)- and \( \beta \)-carboxysomes, as well as the algal pyrenoid, have intrinsically disordered proteins that form liquid-like condensates with Rubisco (49–51). Collectively, these studies suggest that LLPS is a common feature underlying carboxysome biogenesis. The LLPS activity of McdB and other carboxysome components may be related, and potentially influence carboxysome function.

Cold growth temperature triggers cell elongation and asymmetric cell division. Across the bacterial world, complex systems maintain cell size homeostasis. We find here that wild-type \( S. \) elongatus maintains cell size homeostasis when grown at 30°C or 40°C but undergoes filamentation at the environmentally relevant temperature of 20°C. A growing number of bacterial species are known to elongate amid environmental changes to promote survival. For example, \( E. \) coli cells become filamentous during infection (52, 53), and during DNA damage, the SOS response blocks cell division until damage has been repaired (54). Several other forms of stress induce elongation, including host environment, antibiotics, nutrient access, pH, heat shock, and osmotic fluctuations (55–58). Bacteria can clearly exist in diverse morphological states, in part dictated by their environment (35, 59). To our knowledge, we find here the first example of bacterial filamentation caused by cold growth temperature. We propose colder temperatures trigger filamentation indirectly as a carbon limitation response due to the reduced carbon-fixing activity of Rubisco.

Alternatively, transition into a filamentous morphology has been proposed as a way to protect against predation in aquatic environments (56, 60–62). The growth temperature of 20°C coincides with the seasonal temperature during which predation occurs (61). Therefore, it is attractive to speculate that \( S. \) elongatus may use temperature as a cue to elongate so as to avoid planktivorous protists. Consistent with this possibility, previous studies have demonstrated that several freshwater bacteria, including \( C. \) crescentus (56), exhibit high phenotypic plasticity and can transition to a filamentous morphology—a transition that may be specifically triggered in the presence of a size-selective protistan predator (61, 63, 64). Therefore, some aquatic bacteria may undergo filamentation at temperatures that coincide with grazing season.

How bacterial cells enter and exit filamentous states to ensure survival during changes in environment remains poorly characterized. We find here that \( S. \) elongatus elongates when grown at low temperature or when carboxysomes are mispositioned. When these filaments divide, it is asymmetric, with one daughter cell of “normal” length. \( S. \) elongatus also elongates under dim-light stress, and then divides asymmetrically to form
daughter cells of the correct size when brought back into well-lit conditions (65). The Min system has been shown to play a role in the asymmetric cell division and daughter cell sizing of filamentous E. coli (58), Vibrio parahaemolyticus (66, 67), and S. elongatus (65). Division restoration at the poles of these filaments has recently been shown to be regulated by a combination of Min oscillations, FtsZ levels and terminus segregation, resulting in daughter cells of the right length (68, 69). It should be noted that in addition to sensing light and inorganic carbon substrates, S. elongatus responds to lowering temperatures by slowing and even turning off circadian rhythms, thereby downregulating cell growth, division, and metabolic flux (70, 71). The mechanism by which filamentation and asymmetric division occurs in S. elongatus is an area of future research. Taken together, the conserved ability for various bacterial species to undergo filamentation and morphological recovery, some of which showed direct selective benefits, strongly suggests that this differentiation plays an important role in survival and proliferation.

MATERIALS AND METHODS

Construct designs. All constructs in this study were generated using Gibson Assembly (72) from synthesized double-stranded DNA (dsDNA) and verified by sequencing. Constructs contained flanking DNA that ranged from 500 to 1,500 bp in length upstream and downstream of the targeted insertion site to promote homologous recombination into target genomic loci (73).

Generation of bacterial strains. All Synechococcus elongatus PCC 7942 transformations were performed as previously described (73). Strains used in this study are shown in Table 1. Plasmid constructs of mcdA, mcdB, and mcdAB deletions were created by replacing the respective coding sequences with a kanamycin resistance cassette. All fluorescent strains were transformed using plasmid pAH40, which contains a chloramphenicol resistance cassette and a second copy of the kanamycin resistance cassette. All strains were maintained and grown under constant light-emitting diode (LED) illumination of 60 μE m⁻² s⁻¹ at either 20°C, 30°C, or 40°C in 2% or 0.04% CO₂ as specified. Cultures grown in 0.04% CO₂ (ambient) were grown in air with no additional CO₂. Cultures were regularly diluted with fresh medium to maintain exponential growth phase for subsequent imaging and immunoblot analyses. For cloning, One Shot TOP10 Chemically Competent E. coli (ThermoFisher) were grown aerobically at 37°C in Luria broth medium.

Fluorescence microscopy. Microscopy was performed using exponentially growing cells at an optical density (OD) of 0.4. Two milliliters of culture was spun down at 15,000 × g for 60 s and resuspended in 100 μl of BG-11. Five microliters was transferred to a square 1.5% agarose plus BG-11 pad, which was then flipped onto a 35-mm cell culture dish with a #1.5 glass coverslip bottom (ManTek). All images were captured using a Nikon Eclipse Ti2 inverted microscope with a PlanApo Objective lens (100×, 1.45 numerical aperture [NA], oil immersion), with phase contrast trans-illumination, and with a SOLA LED light source for imaging chlorophyll fluorescence (excitation, 560/40 nm [540 to 580 nm]; emission, 630/70 nm [593 to 668 nm], dichroic mirror, 585 nm) and RbcS-mTQ fluorescence (excitation, 436/20 nm [426 to 446 nm]; emission, 480/40 nm [460 to 500 nm]; dichroic mirror, 455 nm). Images were acquired using a Photometrics Prime 95B back-illuminated scientific complementary metal oxide semiconductor (sCMOS) camera. Image analysis was performed using Fiji (74) and the MicrobeJ plugin (75).

Growth curve measurements. Cultures were inoculated at a starting optical density at 750 nm (OD750) of 0.1 to 0.2 with fresh BG-11. Cell growth was monitored at OD750 using a DS-11 spectrophotometer.

| Strain | Description | Reference |
|--------|-------------|-----------|
| Wild type | | 17 |
| ΔmcdA | | 17 |
| ΔmcdB | | 17 |
| ΔmcdAB | | 17 |
| rbcS-mTQ | Wild type transformed with pAH40 | |
| ΔmcdA + rbcS-mTQ | ΔmcdA mutant transformed with pAH40 | |
| ΔmcdB + rbcS-mTQ | ΔmcdB mutant transformed with pAH40 | |
| ΔmcdAB + rbcS-mTQ | ΔmcdAB mutant transformed with pAH40 | |
Biomass measurements. All strains were inoculated at a starting OD₅₇₀ of 0.1 for 24 h (at 30 and 40°C), 48 h (at 20°C with 2% CO₂), or 10 days (at 20°C with ambient CO₂). Cultures were filtered using MF-Millipore membrane filter, 0.22-μm pore size (catalog no. JGWP04700; Sigma-Aldrich). Membrane filters with cells were dried at 80°C for 20 h. Biomass was calculated by subtracting the weight of a membrane filter without cells (60 mg) from the weight of dried membrane filters with cells.

Microbe quantification. Multiple fields of view were taken for each cell strain using three channels: phase contrast microscopy provided cell length, width, and perimeter, while fluorescence microscopy provided chlorophyll autofluorescence and RbcS-mtQ intensities. Chlorophyll autofluorescence was used to determine the site of cell division in dividing cells. These data were analyzed using MicrobeJ (75). In each cell line, cell length detection was performed using the rod-shapeddescriptor and thresholding set to 0.4 μm < area < maximum, 0.71 μm < width range < 2 μm, 0 μm < width variation < 0.2 μm, and 0 μm < angularity amplitude < 0.35 μm. Carboxysome detection was performed using the point function with a tolerance of 20 and an intensity minimum of 500. Associations, shape descriptors, profiles, and distances were recorded for each strain. All MicrobeJ quantification was also verified manually. Graphs and statistical analyses were generated with Graph Pad Prism.

Immunoblot analysis. Cultures were concentrated to an OD₅₇₀ of 3 when harvesting. An equal volume of 2× Laemmli sample buffer was added prior to boiling for 20 min. Samples (55 μl) were loaded on a 4% to 12% Bis-Tris NuPAGE gel with wedge wells (Invitrogen). Gels were transferred onto a mini-size polyvinylidene difluoride membrane (Bio-Rad) using a Trans-Blot Turbo system (Bio-Rad). The membrane was immunoprobed using rabbit polyclonal antisera against RbcL and the beta subunit of ATP synthase, Atpb (Agrisera) and then horseradish peroxidase (HRP)-conjugated anti-rabbit IgG secondary antibody (Millipore Sigma). Membrane signals were developed with femto maximum sensitivity sub- strate (Thermo Scientific) and visualized and quantified using Li-Cor Image Studio (three biological replicates). We normalized RbcL signal using Atpb, a method previously used in Zhang et al. (33) and specifically with S. elongatus PCC 7942 in Sun et al. (32).

CO₂ down-shift assay. Wild-type S. elongatus cultures were grown at 30°C with 2% CO₂ until reaching OD₅₇₀ of 0.6. Cultures were spun down, and the spent medium was discarded. Cell pellets were resuspended in fresh BG-11 medium and split into equal volumes to continue growth at 30°C with 2% CO₂ or ambient CO₂. After 72 h, phase contrast images of ≥300 cells for each condition from 15 fields of view were collected to quantify cell length and asymmetric cell division.

SUPPLEMENTAL MATERIAL
Supplemental material is available online only.

FIG S1, TIF file, 2.6 MB.
FIG S2, TIF file, 2.6 MB.
FIG S3, TIF file, 2.4 MB.
FIG S4, TIF file, 2.7 MB.

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