RESEARCH ARTICLE

Single-Organelle Quantification Reveals Stoichiometric and Structural Variability of Carboxysomes Dependent on the Environment

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17 **Short title**: Carboxysome protein stoichiometry and variability

One Sentence Summary: Determination of absolute protein stoichiometry reveals the organizational
 variability of carboxysomes in response to microenvironmental changes

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26 ABSTRACT

27 The carboxysome is a complex, proteinaceous organelle that plays essential roles in carbon assimilation in cvanobacteria and chemoautotrophs. It comprises hundreds of protein homologs that 28 29 self-assemble in space to form an icosahedral structure. Despite its significance in enhancing CO₂ 30 fixation and potentials in bioengineering applications, the formation of carboxysomes and their 31 structural composition, stoichiometry and adaptation to cope with environmental changes remain 32 unclear. Here we use live-cell single-molecule fluorescence microscopy, coupled with confocal and 33 electron microscopy, to decipher the absolute protein stoichiometry and organizational variability of 34 single β-carboxysomes in the model cyanobacterium Synechococcus elongatus PCC7942. We 35 determine the physiological abundance of individual building blocks within the icosahedral 36 carboxysome. We further find that the protein stoichiometry, diameter, localization and mobility 37 patterns of carboxysomes in cells depend sensitively on the microenvironmental levels of CO₂ and 38 light intensity during cell growth, revealing cellular strategies of dynamic regulation. These findings, 39 also applicable to other bacterial microcompartments and macromolecular self-assembling systems, 40 advance our knowledge of the principles that mediate carboxysome formation and structural 41 modulation. It will empower rational design and construction of entire functional metabolic factories in heterologous organisms, for example crop plants, to boost photosynthesis and agricultural productivity. 42 43

44 Keywords

45 Bacterial microcompartment, carboxysome, protein stoichiometry, self-assembly, single-molecule 46 fluorescence imaging, structural flexibility

48 INTRODUCTION

Organelle formation and compartmentalization within eukaryotic and prokaryotic cells provide 49 50 the structural foundation for segmentation and modulation of metabolic reactions in space and time. Bacterial microcompartments (BMCs) are self-assembling organelles widespread 51 among bacterial phyla (Axen et al., 2014). By physically sequestering specific enzymes key 52 for metabolic processes from the cytosol, these organelles play important roles in CO₂ fixation, 53 pathogenesis, and microbial ecology (Yeates et al., 2010; Bobik et al., 2015). According to 54 their physiological roles, three types of BMCs have been characterized: the carboxysomes for 55 CO₂ fixation, the PDU microcompartments for 1,2-propanediol utilization, and the EUT 56 microcompartments for ethanolamine utilization. 57

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The common features of various BMCs are that they are ensembles composed of purely 59 60 protein constituents and comprise an icosahedral single-layer shell that encases the catalytic enzyme core. This proteinaceous shell, structurally resembling virus capsids, is self-61 assembled from several thousand polypeptides of multiple protein paralogs that form 62 hexagons, pentagons and trimers (Kerfeld and Erbilgin, 2015; Sutter et al., 2016; Faulkner et 63 al., 2017). The highly-ordered shell architecture functions as a physical barrier that 64 concentrates and protects enzymes, as well as selectively gating the passage of substrates 65 and products of enzymatic reactions (Yeates et al., 2010; Bobik et al., 2015). 66

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Carboxysomes serve as the key CO₂-fixing machinery in all cyanobacteria and some 68 chemoautotrophs. The primary carboxylating enzymes, ribulose-1,5-bisphosphate 69 carboxylase oxygenase (Rubisco) (Rae et al., 2013), are encapsulated by the carboxysome 70 shell that facilitates the diffusion of HCO₃⁻ and probably reduces CO₂ leakage into the cytosol 71 (Dou et al., 2008). Based on the form of enclosed Rubisco, carboxysomes can be categorized 72 into two different classes, α -carboxysomes and β -carboxysomes (Rae et al., 2013; Kerfeld 73 and Melnicki, 2016). The β -carboxysomes in the cyanobacterium Synechococcus elongatus 74 75 PCC7942 (Syn7942) have been extensively characterized as the model carboxysomes. The shell of β-carboxysomes from Syn7942 is composed of the hexameric proteins CcmK2, 76 CcmK3 and CcmK4 that form predominately the shell facets (Kerfeld et al., 2005), the 77 pentameric protein CcmL that caps the vertices of the polyhedron (Tanaka et al., 2008), as 78 well as the trimeric proteins CcmO and CcmP (Cai et al., 2013; Larsson et al., 2017). The 79 core enzymes of β-carboxysomes consist of a paracrystalline arrangement of plant-type 80

Rubisco (comprising the large and small subunits RbcL and RbcS) and β-carbonic anhydrase 81 (β -CA, encoded by the *ccaA* gene). The colocalized β -CA dehydrates HCO₃⁻ to CO₂ and 82 creates a CO₂-rich environment in the carboxysome lumen to favor the carboxylation of 83 Rubisco. In addition, CcmM and CcmN function as "linker" proteins to promote Rubisco 84 packing and shell-interior association (Kinney et al., 2012). CcmM in the β -carboxysome 85 appears as two isoforms, a 35-kDa truncated CcmM35 and a full-length 58-kDa CcmM58 86 (Long et al., 2007; Long et al., 2010; Long et al., 2011). CcmM35 contains three Rubisco 87 small subunit-like (SSU) domains that interact with Rubisco (Hagen et al., 2018b; Wang et al., 88 2019), whereas CcmM58 has an N-terminal γ -CA-like domain in addition to the SSU domains 89 and recruits CcaA to the shell. RbcX is recognized as a chaperonin-like protein for Rubisco 90 assembly (Emlyn-Jones et al., 2006; Saschenbrecker et al., 2007; Occhialini et al., 2016); it 91 has been recently revealed to serve as one component of the carboxysome and play roles in 92 93 mediating carboxysome assembly and subcellular distribution (Huang et al., 2019).

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Understanding the physiological composition and assembly principles of carboxysome 95 building blocks is of key importance not solely to unravel the underlying molecular 96 mechanisms of carboxysome formation and biological functions, but also for heterologously 97 engineering and modulating functional CO₂-fixing organelles to supercharge photosynthetic 98 carbon fixation in synthetic biology applications. Previous estimations of the carboxysome 99 protein stoichiometry from either the whole cell lysates or the isolated forms using immunoblot 100 and mass spectrometry illustrated the relative abundance of carboxysome proteins (Long et 101 al., 2005; Long et al., 2011; Rae et al., 2012; Faulkner et al., 2017). Moreover, it was revealed 102 that carboxysome biosynthesis in Syn7942 is highly dependent upon environmental 103 conditions during cell growth, such as light intensity (Sun et al., 2016) and CO₂ availability 104 (McKay et al., 1993; Harano et al., 2003; Woodger et al., 2003; Whitehead et al., 2014). The 105 exact stoichiometry of all building components in the functional carboxysome and how 106 carboxysomes manipulate their compositions, organizations and functions to cope with 107 environmental changes have remained elusive. 108

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Here, we construct a series of Syn7942 mutants with individual components of carboxysomes functionally tagged with the bright and fast-maturing enhanced yellow fluorescent protein (YFP) and report the *in vivo* characterization of protein stoichiometry of carboxysomes at the single-organelle level, using real-time single-molecule fluorescence microscopy, confocal and

114 electron microscopy combined with a suite of biochemical and genetic assays. Quantification of the protein stoichiometry of β -carboxysomes in Syn7942 grown under different conditions 115 116 demonstrates the organizational flexibility of β -carboxysomes, and their ability to modulate 117 functions towards local alterations of CO₂ levels and light intensity during cell growth, as well as the regulation of the spatial localization and mobility of β -carboxysomes in the cell. This 118 study provides fundamental insight into the formation and structural plasticity of 119 carboxysomes and their dynamic organization towards environmental changes, which could 120 be extended to other BMCs and macromolecular systems. A deeper understanding of the 121 protein composition and structure of carboxysomes will inform strategies for rational design 122 and engineering of functional and adjustable metabolic modules towards biotechnological 123 applications. 124

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127 **RESULTS**

128 Protein stoichiometry of functional carboxysomes at the single-organelle level

129 We constructed ten Syn7942 strains expressing individual β -carboxysome proteins (CcmK3. 130 CcmK4, CcmK2, CcmL, CcmM, CcmN, RbcL, RbcS, CcaA, RbcX) fused with YFP at their Ctermini individually (Supplemental Figure 1). Fluorescence tagging at the native chromosomal 131 locus under the control of their native promoters ensures expression of the fluorescently-132 tagged proteins in context and at physiological levels (Sun et al., 2016). Eight of these strains, 133 in which YFP was fused to CcmK3, CcmK4, CcmL, CcmM, CcmN, RbcS, CcaA, and RbcX 134 respectively, are fully segregated (Supplemental Figures 1C and 2) and exhibit wild-type 135 levels of cell size, growth and carbon fixation within experimental error (Supplemental Table 136 1), consistent with previous observations (Savage et al., 2010; Cameron et al., 2013; Sun et 137 al., 2016; Faulkner et al., 2017; Huang et al., 2019). 138

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By contrast, RbcL-YFP and CcmK2-YFP strains are only partially segregated, in agreement 140 with previous studies (Savage et al., 2010; Cameron et al., 2013; Sun et al., 2016). Through 141 immunoblot analysis using anti-fluorescence protein, anti-RbcL and anti-CcmK2 antibodies 142 (Supplemental Figure 2B), we estimate that 29.2 \pm 7.1 % (mean \pm standard deviation (SD), n 143 = 4) of total RbcL and 6.0 \pm 0.7 % (*n* = 3) of total CcmK2 were tagged with YFP in RbcL-YFP 144 and CcmK2-YFP strains. Nevertheless, we excluded the stoichiometric quantification of RbcL 145 and CcmK2 in this study, in view of the partial segregation which could result in quantification 146 inaccuracy. 147

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We used single-molecule Slimfield microscopy (Plank et al., 2009) to visualize individual 149 carboxysomes that were fused with YFP (Figure 1, Supplemental Figure 3). This technique 150 allows detection of fluorescently-labelled proteins with millisecond sampling, enabling real-151 time tracking of rapid protein dynamics inside living cells, exploited previously to study 152 functional proteins involved in bacterial DNA replication and remodeling (Reves-Lamothe et 153 154 al., 2010; Badrinarayanan et al., 2012), gene regulation in budding yeast cells (Wollman et al., 2017; Leake, 2018), bacterial cell division (Lund et al., 2018), and chemokine signaling in 155 lymph nodes (Miller et al., 2018). Our prior measurements using relatively fast-maturing 156 fluorescent proteins such as YFP suggest that less than 15% of fluorescent proteins are likely 157 to be in a non-fluorescent immature state (Leake et al., 2008; Shashkova et al., 2018). 158

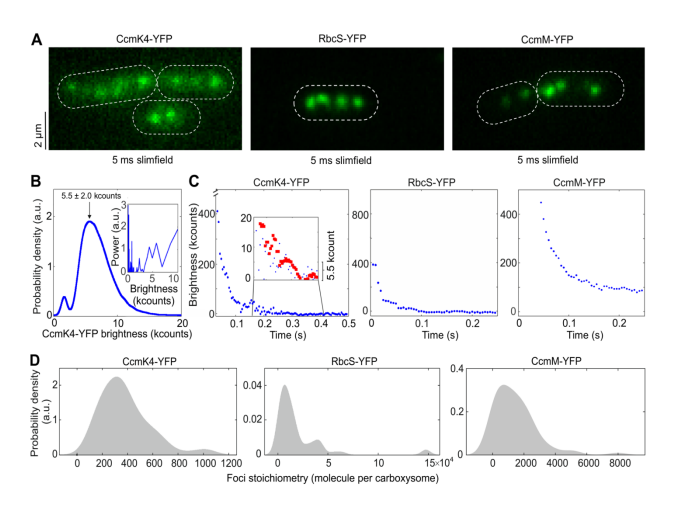


Figure 1. Slimfield quantification of cells grown under ambient air/moderate light Air/ML condition.

(A) Averaged Slimfield images of YFP fluorescence (green) over 5 frames of strains expressing shell component CcmK4-YFP, the interior enzyme RbcS-YFP and the shell-interior linker protein CcmM-YFP. White dash lines indicate cell body outlines.

(B) Distribution of automatically detected foci intensity from the end of the photobleaching, corresponding to the characteristic intensity of *in vivo* YFP. Inset shows the Fourier spectrum of 'overtracked' foci, tracked beyond photobleaching, showing a peak at the characteristic intensity.

(C) Representative fluorescence photobleaching tracked at ultra-fast speed. The CcmK4 plot shows an inset 'zoomed in' on lower intensity range with step-preserving Chung-Kennedy filtered data in red, with individual photobleaching steps clearly visible at the characteristic intensity. Brightness (kcounts), counts measured per camera pixel multiplied by 1,000.

(D) Distribution of YFP copy number detected for individual carboxysomes in corresponding mutants, rendered as kernel density estimates using standard kernel width. Heterogeneity of contents was observed, also a "preferable" copy number, represented by kernel density peak values could be determined. Statistics of copy numbers (Peak value ± HWHM) are listed in Table 1 for ML conditions. The corresponding Slimfield images and histogram for complete strain sets are shown in Supplemental Figure 3.

160 Figure 1A shows the Slimfield images of three representative Syn7942 strains RbcS-YFP,

161 CcmK4-YFP and CcmM-YFP that grow under ambient air and moderate light (hereafter

denoted Air/ML), to determine the protein stoichiometry from different carboxysome structural 162 domains. Single carboxysomes are detected as distinct fluorescent foci in cells of the YFP-163 164 fused strains (Figure 1A, Supplemental Figure 3), whose sigma width is approximately 250 nm (n = 100), comparable to the diffraction-limited point spread function width of our imaging 165 system. We use the number of YFP molecules per fluorescent focus as an indicator of the 166 stoichiometry of the fluorescently-labelled protein subunits in each individual carboxysomes. 167 which we determined by quantifying step-wise photobleaching of the fluorescent tag during 168 the Slimfield laser excitation process (Figures 1B to 1C, Table 1) using a combination of 169 Fourier spectral analysis and edge-preserving filtration of the raw data (Leake et al., 2003; 170 Leake et al., 2004; Leake et al., 2006) (see details in Materials and Methods). The resulting 171 broad distributions of protein stoichiometry, rendered as kernel density estimates, suggest a 172 variable content of individual components per carboxysome (Figure 1D), indicative of the 173 174 structural heterogeneity of β -carboxysomes. The modal average stoichiometry of each protein subunit per carboxysome was defined by the measured peak from each distribution of the raw 175 stoichiometric data (Figure 1D, Supplemental Figure 3), after subtracting the background 176 fluorescence distribution, primarily from chlorophylls, which was determined from the WT cells 177 (Supplemental Figure 4). 178

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In the β -carboxysome synthesized in cells grown under Air/ML, Rubisco enzymes are the 180 predominant components, as indicated by the RbcS content (Table 1). CcmM is the second 181 most abundant element; there are over 700 copies of CcmM molecules per β-carboxysome. 182 In addition, the CcmK4 content is greater than that of CcmK3 by a factor of 3.8. CcmL, CcmN, 183 CcaA and RbcX are the minor components in the β -carboxysome. Our results reveal that 184 there are 37 CcmL subunits per carboxysome, with the raw stoichiometry distribution showing 185 some indications of peaks at multiples of ~5 molecules indicative of multiples of CcmL 186 pentamers (Supplemental Figure 4C), consistent with the atomic structure of CcmL (Tanaka 187 et al., 2008). A modal average of 37 CcmL molecules thus suggests that a single 188 189 carboxysome contains an average of 7.4 CcmL pentamers, less than the 12 CcmL pentamers that were postulated to occupy all the vertices of the icosahedral shell (Bobik et al., 2015; 190 Kerfeld et al., 2018). It is feasible that not all vertices of the carboxysome structure are 191 capped by CcmL pentamers, as BMC shells deficient in pentamers could still be formed 192 without notable structural variations (Cai et al., 2009; Lassila et al., 2014; Hagen et al., 2018a). 193

194 Our study represents a direct characterization of protein stoichiometry at the level of single 195 functional carboxysomes in their native cellular environment.

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As a control, we fused RbcL with mYPet, a monomeric-optimized variant of YFP. The RbcL-YFP and RbcL-mYPet cells show no significant difference in the subcellular distribution of carboxysomes as well as cell doubling times and carbon fixation (Supplemental Figure 5), demonstrating that there are no measurable artefacts due to putative effects of dimerization of the YFP tag.

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We also examined the relative abundance of individual carboxysome proteins in the YFP-203 fusion Syn7942 strains in cell lysates, using immunoblot probing with an anti-fluorescent 204 protein antibody (Supplemental Figure 2A, Supplemental Table 2). To compare with the 205 206 stoichiometry obtained from Slimfield, we normalized the abundance of carboxysome proteins estimated from immunoblot analysis, using the RbcS content per carboxysome determined by 207 Slimfield. It appears that the content of β -carboxysome proteins determined by 208 immunoblotting is generally greater than that within the carboxysome characterized by 209 Slimfield. Despite the potential effects caused by YFP fusion, this could suggest the presence 210 of a "storage pool" of carboxysome proteins located in the cytoplasm that are involved in the 211 biogenesis, maturation and turnover of carboxysomes. The ratio of RbcL/S detected from cell 212 lysates fraction is about 8:5.8 (n = 4) (Supplemental Table 2), in line with previous results 213 (Long et al., 2011) but distinct from the *in vitro* reconstitution observations (Ryan et al., 2018; 214 Wang et al., 2019). 215

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217 Stoichiometry of carboxysome proteins exhibit a dependence on the 218 microenvironment conditions of live cells

Our previous study showed that the content and spatial positioning of β -carboxysomes in 219 Syn7942 are dependent upon light intensity during cell growth, revealing the physiological 220 regulation of carboxysome biosynthesis (Sun et al., 2016). Whether the stoichiometry of 221 different components in the carboxysome structure changes in response to fluctuations in 222 environmental conditions is unknown. Here we addressed this question by taking advantage 223 of the far greater throughput of confocal microscopy compared to Slimfield, whilst still using 224 the single-molecule precise Slimfield data as a calibration to convert the intensity of detected 225 foci from confocal images into estimates for absolute numbers of stoichiometry. We achieved 226

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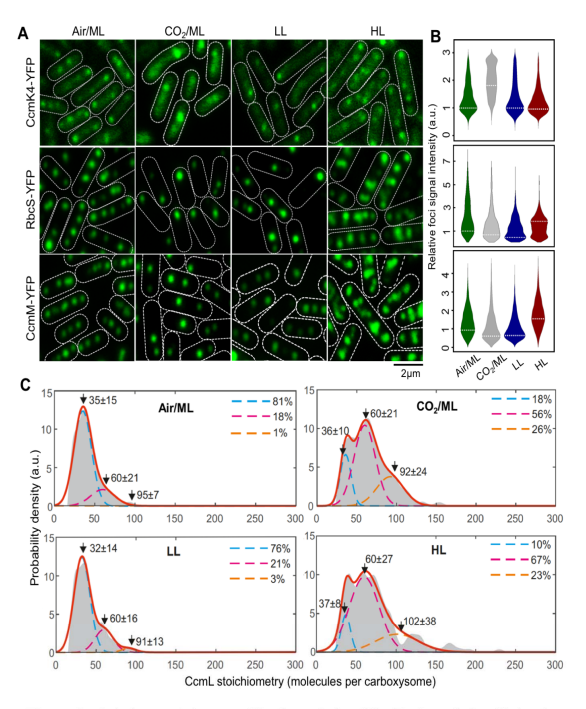


Figure 2. Relative protein quantification of CcmK4, RbcS and CcmM in the carboxysome under different CO_2 levels and light intensities using confocal microscopy.

(A) Confocal images of CcmK4-YFP, RbcS-YFP and CcmM-YFP strains under Air/ML, CO_2/ML , LL and HL. Fluorescence foci (green) indicate carboxysomes and cell borders were outlined by white dashed lines. Scale bar indicates 2 μ m.

(B) Violin plot of carboxysome intensities under Air/ML, CO₂/ML, LL and HL, normalized to kernel density ML peak values (peaks marked by white dash lines).

(C) Kernel density estimates of CcmL carboxysome copy number grown under Air/ML, CO₂, LL and HL detected by Slimfield and corrected for chlorophyll. Triple Gaussian fits are indicated as colored dashed lines with summed fit in red. The percentage in each Gaussian is indicated aside.

this by identifying the peak value of the foci intensity distribution from each given cell strain

obtained from confocal imaging with the peak value of the measured Slimfield foci stoichiometry distribution for the equivalent cell strain under Air/ML. This approach allows us to generate a conversion factor which we then applied to subsequent confocal data acquired under lower light (LL), higher light (HL) and ML with the air supplemented by 3% CO₂, and to estimate relative changes in the stoichiometry of carboxysome building components using large numbers of cells, without the need to obtain separate Slimfield datasets for each condition (Figure 2, Supplemental Figures 6 to 8).

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Figure 2A shows confocal fluorescence images of RbcS-YFP, CcmK4-YFP, and CcmM-YFP 236 strains grown under Air/ML, 3% CO₂ (CO₂/ML), LL and HL. The confocal images reveal 237 classic patterns of cellular localization of carboxysomes similar to those observed with 238 Slimfield microscopy (Supplemental Figure 6). We analyzed the confocal images to detect 239 240 carboxysome fluorescent foci within the cells and quantify their fluorescence intensities (Figure 2B, Supplemental Figures 7 and 8). We find that the number of carboxysomes per cell 241 is dependent on growth conditions; it is reduced under CO₂/ML in contrast to Air/ML, whereas 242 HL increases the abundance of β -carboxysomes (Supplemental Table 3), consistent with 243 previous findings (Whitehead et al., 2014; Sun et al., 2016). The slightly different 244 carboxysome contents estimated in individual YFP-fused strains might suggest potential 245 mechanisms of the cells that tune carboxysome organization. As a common feature, the 246 abundance of all the proteins in the β -carboxysome is apparently modulated under distinct 247 growth conditions. For instance, both RbcS and CcmM have a higher content per 248 carboxysome under HL compared with that under other conditions, whereas the CcmK4 249 content per β -carboxysome increase under 3% CO₂ (Figure 2B). The dependence of 250 carboxysome protein stoichiometry inferred from the peak values of the stoichiometry 251 distributions under different cellular microenvironmental conditions is summarized in Table 1. 252

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Interestingly, we find that the variation of CcmL abundance per carboxysome rises with increasing light illumination and CO₂ availability (Figure 2C). The measured stoichiometry distribution of CcmL pentamers suggests the presence of three populations: (I) carboxysomes with < 60 CcmL subunits (in the range of 32-37); (II) carboxysomes with 60 CcmL subunits, consistent with the expectation that 12 vertices of the icosahedral carboxysome are fully occupied by CcmL pentamers (Tanaka et al., 2008; Rae et al., 2013; Kerfeld et al., 2018); (III) carboxysomes with > 60 CcmL subunits (in the range of 91-102). Using a nearest-neighbor

model to estimate the probability for the diffraction-limited optical images of individual 261 carboxysomes in a cell, we find that the Population III carboxysomes represent random 262 overlap of two or more carboxysomes from the Population I and II (Figure 2C). Population I 263 represents a "non-complete capped" state in which not all vertices in the icosahedron are 264 occupied by CcmL pentamers. We find the characteristic stoichiometry of the Population I 265 carboxysomes increases with the enhancement of light intensity during cell growth, from 32 266 CcmL molecules (LL) to 35 (ML) and 37 (HL), with HL having a significantly smaller proportion 267 (23%) of "non-complete capped" carboxysomes compared to ~80% under LL and ML 268 conditions. Supplementing the air with 3% CO₂ under ML similarly results in a substantial 269 decrease in the proportion of "non-complete capped" carboxysomes in the population (18%) 270 comparable to the HL condition in the absence of any supplemental CO₂. These findings 271 suggest a dependence of carboxysome assembly which may allow adaptation towards 272 273 microenvironmental changes, i.e. the increase in the population of capped carboxysomes in situations which are favorable towards photosynthesis (HL conditions and locally-raised levels 274 of CO_2). 275

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277 This finding is also validated by the changes in protein abundance of other carboxysome components under environmental regulation (Table 1, Supplemental Figures 7 and 8). Cells 278 were maintained under different growth conditions prior to microscopy imaging, to ensure 279 their full acclimation. Variations of protein content in carboxysomes under CO₂/ML vs. Air/ML, 280 and HL vs. LL conditions indicate distinct fashions of stoichiometric regulation of 281 carboxysome building blocks (Figure 3, Supplemental Table 4). The abundance of CcmK3 282 and CcmK4, whose encoding genes are distant from the ccmKLMNO operon (Sommer et al., 283 2017), increases under 3% CO₂ and remains relatively constant under HL/LL, contrary to the 284 changes in the abundance of CcmN and CcmM that are located in the ccm operon. In 285 addition, the ratio of CcmK4:CcmK3 per carboxysome appear to be relatively constant, in the 286 range of 3.6-4.1 (Supplemental Table 5), indicating the organizational correlation between 287 CcmK3 and CcmK4 within the β -carboxysome structure. We find the rise of CcaA content and 288 reduction of RbcS content under CO₂/ML vs. Air/ML, whereas both increase under HL, 289 suggesting distinct regulation of the two components. It has been recently demonstrated that 290 the putative Rubisco chaperone RbcX is part of the carboxysome and plays roles in mediating 291 carboxysome formation (Huang et al., 2019). The fold changes of RbcX content in each 292 carboxysome under different conditions are close to 1 (Figure 3), probably ascribed to the fact 293

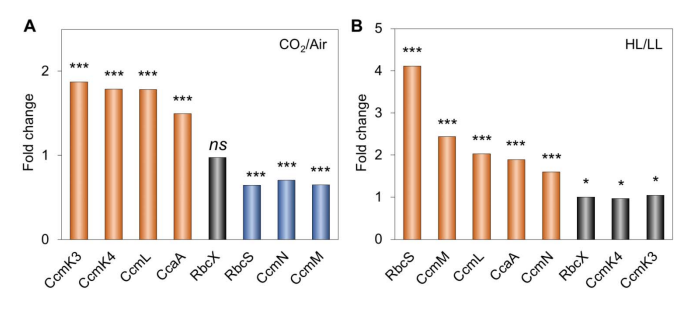


Figure 3. Changes in carboxysome protein stoichiometry by increase in CO_2 levels and light intensity.

(A) Comparison of carboxysome protein stoichiometry under CO_2 treatment. Increase in the CO_2 concentration resulted in the rise of CcmK3, CcmK4, CcaA and CcmL contents and the decline of RbcS, CcmN and CcmM contents.

(B) Comparison of carboxysome protein stoichiometry under light intensity treatment. Increased light intensity led to the elevation of RbcS, CcmM, CcmL, CcaA and CcmN contents, whereas the abundance of RbcX, CcmK3 and CcmK4 contents per carboxysome does not change dramatically.

Man-Whitney U-tests were performed to compare the numbers of functional units of individual carboxysome proteins changed from CO_2/ML to Air/ML (A) and from HL to LL (B). *, p < 0.05; ***, p < 0.005; ns, p > 0.05.

- that its encoding gene is distant from the *rubisco* and *ccm* operons in Syn7942. Collectively,
 these results highlight the highly flexible stoichiometry of individual components within the
 natural carboxysomes in response to environmental changes.
- 297

298 Variation of carboxysome diameter represents a strategy for manipulating 299 carboxysome activity to adapt to environmental conditions

The change in the protein content per carboxysome signifies the variation of β -carboxysome size and organization among different cell growth conditions. Indeed, electron microscopy (EM) of Syn7942 WT cells substantiates the variable structures of β -carboxysomes in response to the changing environment (Figures 4A and 4B). The average diameter of β carboxysomes is 192 ± 41 nm (n = 33) in Air/ML, 144 ± 24 nm (n = 25) in 3% CO₂, 151 ± 22 nm (n = 27) in LL, and 208 ± 28 nm (n = 51) in HL (Figure 4B, Supplemental Table 5, Supplemental Figure 9). These results reveal that both the CO₂ level and light intensity can

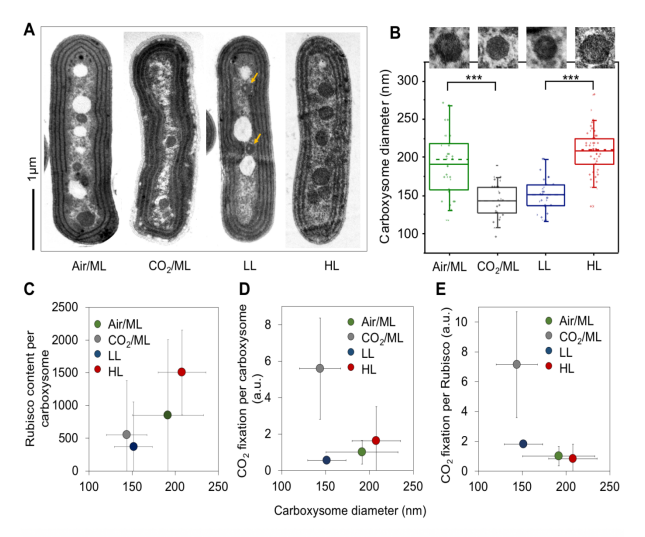


Figure 4. Variations of the carboxysome size and carbon fixation under Air/ML, CO₂, LL and HL.

(A) Thin-section electron microscopy (EM) images showing individual carboxysomes in the Syn7942 WT cells under Air/ML, CO_2 , LL and HL treatments Yellow arrows indicate the carboxysomes with spaces of low protein density under LL. More EM images are shown in Supplemental Figure 9. Scale bar indicates 1 μ m.

(B) Changes in the carboxysome diameter under Air/ML, CO₂, LL and HL measured from EM (n = 33, 25, 27 and 51, respectively), with representative carboxysome images depicted above. Dashed lines indicate medians and solid lines indicate means. Differences in the carboxysome diameter are significant between CO₂ and air ($p = 1.92 \times 10^{-14}$) and between LL and HL ($p = 8.29 \times 10^{-7}$), indicated as ***.

(C) Correlation between the carboxysome size and the Rubisco content per carboxysome under Air/ML, CO₂, LL and HL.

(D) Correlation between the carboxysome size and CO_2 fixation per carboxysome.

(E) Correlation between the carboxysome size and CO_2 fixation per Rubisco of the carboxysomes. Carboxysome diameters and CO_2 fixation were present as average ± SD, whereas the carboxysome total protein content and Rubisco content were shown as Peak value ± HWHM.

307 result in alternations of carboxysome size (Figure 4B). Larger β -carboxysomes can

accommodate more Rubisco enzymes (estimated on the basis of RbcS content) (Figure 4C). 308 An exception is the carboxysomes under LL, which are around 5% larger than the 309 310 carboxysomes under 3% CO₂ but comprises only 67% of Rubisco per carboxysome under CO_2 (Figure 4C, Supplemental Table 5). EM images reveal that the lumen of β -carboxysomes 311 synthesized under LL often contain regions with low protein density (Figure 4A, arrows; 312 Supplemental Figure 9), 59% for LL (16 out of 27 carboxysomes) compared with 9% for 313 Air/ML (3 out of 33), 12% for CO₂/ML (3 out of 25) and 8% for HL (4 out of 51), which likely 314 accounts for the reduced and uneven Rubisco loading within the β -carboxysome. 315

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We also find that CO₂-fixing activity per carboxysome increases as the β -carboxysome 317 structure enlarges, which is correlated to strong light intensity during cell growth (Figure 4D), 318 demonstrating the correlation between β -carboxysome structure and function *in vivo*. 319 320 Moreover, under HL the CO₂-fixation activity per Rubisco of the β -carboxysome declines as the carboxysome size and Rubisco density in the carboxysome lumen increase (Figure 4E, 321 Supplemental Table 5). This may suggest that Rubisco density and local Rubisco packing are 322 important for determining CO₂-fixation activity of individual Rubisco (Supplemental Table 5). 323 324 Interestingly, the relatively small β -carboxysomes under 3% CO₂ exhibit high CO₂-fixing activities per Rubisco and per carboxysome, compared with β-carboxysomes under other 325 conditions. The enhanced carbon fixation capacity under 3% CO₂ might be correlated with the 326 increase in CcmK3 and CcmK4 content (Figure 3A, Table 1), as it has been shown that 327 depletion of CcmK3/CcmK4 impedes carbon fixation of carboxysomes (Rae et al., 2012). 328

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Patterns of spatial localization and diffusion of β-carboxysomes in live cells change dynamically depending upon light intensity during growth

The patterns of β -carboxysome localization within the cyanobacterial cells appears to be 332 crucial for carboxysome biogenesis and metabolic function (Savage et al., 2010; Sun et al., 333 2016). We measured the organizational dynamics of β -carboxysomes with distinct diameters 334 in Syn7942 under different light intensities, using time-lapse confocal fluorescence imaging on 335 the RbcL-YFP Syn7942 strain. Previous studies have shown that tagging of RbcL with 336 fluorescent proteins does not obstruct β -carboxysome assembly and function in Syn7942 337 (Savage et al., 2010; Cameron et al., 2013; Chen et al., 2013; Sun et al., 2016). During time-338 lapse confocal imaging, we applied illumination on the cell samples, similar to that used for 339 cell growth, in order to maintain cell physiology. We find that the overall mobility of individual 340

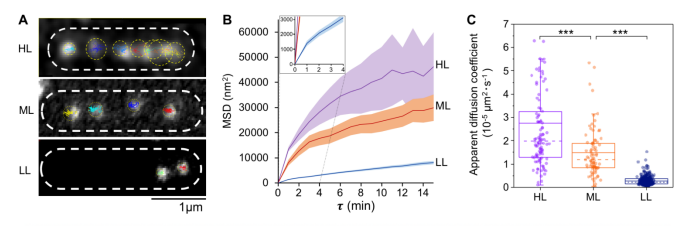


Figure 5. Spatial localization and diffusion dynamics of carboxysomes in Syn7942 cells are dependent on light intensity.

(A) Tracking of carboxysome diffusion in cells grown under HL, ML and LL. Colored lines indicate the diffusion trajectories of each carboxysomes and circles represent the diffusion areas of each carboxysomes over 60 mins. Scale bar indicates $1 \,\mu$ m.

(B) Non-linear MSD (Mean Square Displacement) *vs.* the time interval (τ) profiles suggest the mobility of carboxysomes in Syn7942 cells grown under HL, ML and LL. Inset, zoom-in view of the MSD profile under LL.

(C) Diffusion coefficient of carboxysomes *in vivo* decreases significantly when the light intensity reduces: $2.76 \pm 2.83 \times 10^{-5} \mu m^2 \cdot s^{-1}$ for HL (mean \pm SD, n = 105), $1.48 \pm 1.03 \times 10^{-5} \mu m^2 \cdot s^{-1}$ for ML (n = 84), and $0.28 \pm 0.19 \times 10^{-5} \mu m^2 \cdot s^{-1}$ for LL (n = 336). $p = 3.05 \times 10^{-5}$ between HL and ML; $p = 2.77 \times 10^{-5}$ between ML and LL, two-tailed Student's t-test).

β-carboxysomes within cyanobacterial cells is non-Brownian (Figure 5A, Supplemental Movie
1). Carboxysomes under HL display larger diffusive regions than those under LL. The mean

343 square displacement (MSD) of tracked carboxysomes increased with the rise of light intensity

344 (Figure 5B), as did the mean microscopic diffusion coefficient of individual carboxysomes

(Figure 5C): an average diffusion coefficient of 2.76 ± 2.83 x $10^{-5} \mu m^2 \cdot s^{-1}$ for HL (mean ± SD,

346
$$n = 105$$
), 1.48 ± 1.03 x 10⁻⁵ µm² · s⁻¹ for ML ($n = 84$), and 0.28 ± 0.19 x 10⁻⁵ µm² · s⁻¹ for LL ($n = 84$)

336). It is interesting that the mobility of carboxysomes does not exhibit typical constrained 347 diffusion – asymptotic MSD values at higher values of τ (Robson et al., 2013) – but rather 348 exhibits anomalous diffusion at higher values of τ characterized by a non-linear relation. 349 which can be observed in the intracellular protein mobility traces of other cellular systems 350 (Lenn et al., 2008; Wollman et al., 2017). These results indicate the intracellular restrictions, 351 for example the proposed interactions with the cytoskeletal system (Savage et al., 2010), 352 McdA and McdB (MacCready et al., 2018) and ParA-mediated chromosome segregation (Jain 353 et al., 2012), may mediate carboxysome positioning, but do not completely confine the 354 mobility of carboxysomes. Notably, carboxysomes with a larger diameter (Figure 4) generated 355

under HL present a higher diffusion coefficient compares with carboxysomes with relatively smaller size under ML and LL. However, there is no apparent correlation between the diffusion coefficient of carboxysomes and their size in the same light conditions (Supplemental Figure 10).

360

362 Discussion

Precise quantification of the protein stoichiometry and organizational regulation of 363 carboxysomes provides insight into their assembly principles, structure and function. In this 364 work, we functionally fused fluorescent protein tags to the building blocks in β -carboxysomes 365 and exploited advanced "Physics of Life" technologies, in particular using bespoke single-366 molecule fluorescence microscopy to count the actual protein stoichiometry of β-367 carboxysomes in Syn7942 cells, at the single-organelle level. This approach minimizes the 368 ensemble averaging encountered in bulk estimations from proteomic and immunoblot 369 analysis. We characterized the stoichiometric flexibility of carboxysome proteins within 370 individual polyhedral structures towards environmental variations. Variability of the protein 371 stoichiometry and size of carboxysomes likely provides the structural foundation for the 372 physiological regulation of carboxysome formation and carbon fixation activity. Given the 373 374 shared structural features of carboxysomes and other BMCs, we believe that this work opens up new opportunities to quantitatively evaluate protein abundance and decipher the formation 375 of all BMC organelles, in both native forms and synthetic variants. 376

377

Despite prior efforts on understanding carboxysome structure and function, the relative 378 stoichiometry of functional carboxysome components in their native cell environment - key 379 information required for reconstituting entire active carboxysome structures in synthetic 380 biology (Fang et al., 2018), was still unclear. The major challenges have been the poor 381 specificity of immunoblots and mass spectrometry, given the homology of carboxysome 382 proteins and the lack of effective purification of intact carboxysomes from host cells, as well 383 as the heterogeneity of carboxysome structures (Long et al., 2005). The previous model of 384 carboxysome protein stoichiometry was based on the total amount of proteins in cell lysates 385 (Long et al., 2011) and does not directly reflect the stoichiometry of carboxysome proteins in 386 the organelle, given the possible free-standing carboxysome components in the cytosol (Dai 387 et al., 2018). We have recently reported the isolation of β -carboxysomes from Syn7942 and 388 the structural and mechanical exploration of the organelles (Faulkner et al., 2017). 389 Interestingly, some components, i.e. CcmO, CcmN, CcmP and RbcX, were not detectable by 390 mass spectrometry in the isolated carboxysomes, likely due to their low content or potential 391 loss of carboxysome components during isolation. Here, as demonstrated, fluorescence 392 tagging and Slimfield and confocal imaging enable single-organelle analysis of the protein 393 stoichiometry of eight β-carboxysome proteins (including RbcX) and their regulation in their 394

395 native context, and extends analyses of the assembly and action of carboxysomes. Microscopy imaging of fluorescently-tagged β -carboxysomes has been used to reveal their 396 397 patterns of cellular localization, biogenesis pathways and light-dependent regulation in Syn7942 (Savage et al., 2010; Cameron et al., 2013; Chen et al., 2013; Sun et al., 2016; 398 Niederhuber et al., 2017; MacCready et al., 2018). Although we cannot completely exclude 399 the potential effects of YFP tags on carboxysome structure, we validate that YFP tagging to 400 most of the structural components does not impede formation of functional carboxysome 401 structures, suggesting the physiological relevance of the determined protein stoichiometry in 402 the carboxysome in the presence of fluorescence tags. This flexibility emphasizes the 403 extraordinary capacity of the carboxysome structure in adjusting their protein stoichiometry 404 and accommodating foreign proteins while maintaining functionality, indicating the possibility 405 of manipulating carboxysome organization in bioengineering for diverse purposes. 406 407 Exceptionally, fluorescence tagging on CcmP and CcmO does not show normal carboxysome assembly and localization compared to other YFP-tagged strains (Supplemental Figure 11). 408 In this work, therefore, we did not include estimation of the protein abundance of CcmP and 409 CcmO, as well as RbcL and CcmK2 that cannot be fully tagged with YFP. 410

411

Numerous studies have described the regulation of carboxysome protein expression at the 412 transcriptional level (McGinn et al., 2003; Woodger et al., 2003; Schwarz et al., 2011). 413 Counting protein abundance of β -carboxysomes at different cell growth conditions enables 414 direct characterization of the stoichiometric plasticity of carboxysome building components in 415 the cells grown under not only the same environmental condition but also a range of various 416 conditions (Figure 6A). Our observations elucidate the size variation of β -carboxysomes in 417 Syn7942 cells grown under distinct environmental conditions (Figure 6B) and adjustable 418 carbon fixation capacities of carboxysomes that may be closely linked to the protein 419 organization and size of carboxysomes. Variations in the diameter of intact carboxysomes, 420 ranging from 90 to 600 nm, have been also shown in previous studies not only in single 421 species but also among distinct species (Shively et al., 1973; Price and Badger, 1991; Iancu 422 et al., 2007; Liberton et al., 2011), suggesting the adaptation strategies exploited by 423 cyanobacteria for regulating their CO₂-fixing machines to survive in diverse niches. It may be 424 related to the environment-sensitive protein-protein interactions that drive protein self-425 assembly and BMC formation (Faulkner et al., 2019). Moreover, the spatial positioning and 426 mobility of β-carboxysomes in live cells appear to be independent of carboxysome diameter 427

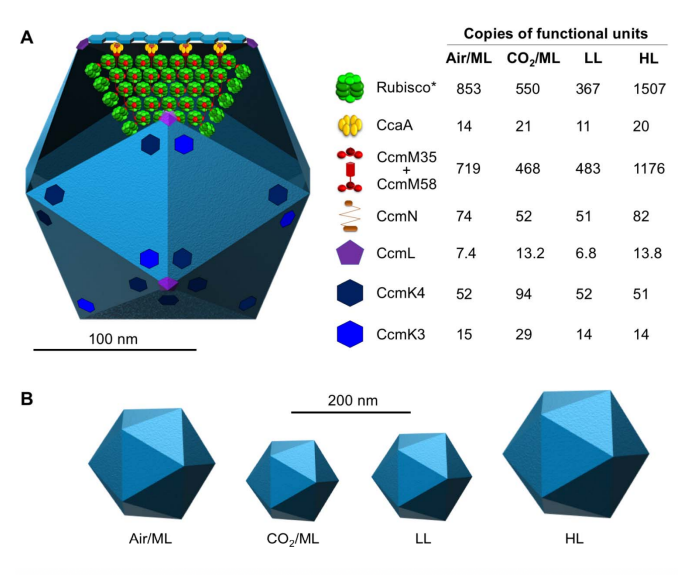


Figure 6. Model of the β -carboxysome structure and protein stoichiometry.

(A) Diagram of an icosahedral carboxysome structure and organization of building components. The stoichiometry of each building component within the carboxysome and its variations in response to changes in CO_2 and light intensity are shown on the right (See also Table 1). *Rubisco content was estimated from RbcS stoichiometry based on the RbcL₈S₈ Rubisco structure. The majority of shell facets shown in light blue is tiled by the major shell protein CcmK2. The total abundance of CcmM58 and CcmM35 was estimated. The components RbcL, CcmK2, CcmO and CcmP were not directly determined in this work and thus are not shown in this model.

(B) The carboxysome diameter is variable in response to changes in the CO_2 level and light intensity.

- 428 but show a strong dependence to light intensity, suggesting that light-dependent mechanisms
- 429 might mediate carboxysome location and diffusion. Carboxysome spacing and partitioning
- 430 have been suggested to be driven by different possible mechanisms, such as the cytoskeletal
- 431 proteins ParA and MreB (Savage et al., 2010), ParA-mediated chromosome segregation (Jain

et al., 2012) via filament-pull model (Ringgaard et al., 2009) or a diffusion-ratchet model (Vecchiarelli et al., 2013) as well as very recently the McdA and McdB that utilize a Brownianratchet mechanism to position carboxysomes (MacCready et al., 2018). Altogether, the organizational flexibility of β -carboxysomes, including modulatable protein stoichiometry, diameter and mobility, may represent the natural strategies for modifying shell permeability and enzyme encapsulation and ensuring structural and functional adaptations dependent on the local cellular environment.

439

The estimated number of CcmL pentamers per carboxysome could be less than 12, 440 demonstrating explicitly that it is not a prerequisite for CcmL pentamers to occupy all 12 441 vertices of the icosahedral shell to ensure complete formation of functional carboxysomes. 442 This hypothesis has been validated by previous observations that BMC shells in the absence 443 444 of pentamers have no significant morphological changes (Cai et al., 2009; Lassila et al., 2014; Hagen et al., 2018a). These "non-complete capped" forms appear to be prevalent among the 445 resultant carboxysomes under Air/ML and LL (Figure 2C), unlike the procarboxysomes 446 (Cameron et al., 2013) or "immature" carboxysomes which are incapable of establishing an 447 oxidative microenvironment for cargo enzymes (Chen et al., 2013). Whether the loss of 448 capping CcmL will create large space within the shell, as a possible mechanism of modulating 449 shell permeability, or will be compensated for by incorporation of other shell proteins, for 450 example the additional CcmP trimers that are speculated to be responsible for permeability, 451 remains to be further investigated. Our results also suggest that carboxysomes could possess 452 a flexible molecular architecture, resonating with the observation of structural "breathing" of 453 virus capsids which has been reported to be key to cope with temperature change (Roivainen 454 et al., 1993; Li et al., 1994). Carboxysomes, though structurally resembling virus capsids, 455 have been shown to be mechanically softer than the P22 virus capsid by a factor of ~10, 456 suggesting greater flexibility of protein-protein interactions within the carboxysome structure 457 (Faulkner et al., 2017). The capping flexibility of pentamers may represent the dynamic nature 458 of shell assembly probably in the second timescale and tunable protein-protein interactions in 459 the shell, as characterized recently (Sutter et al., 2016; Faulkner et al., 2019). 460

461

It was proposed that CcmM58 proteins are confined to a subshell layer for linking Rubisco, CcaA and CcmN to the shell, whereas CcmM35 molecules are predominantly located in the core to stimulate Rubisco aggregation (Rae et al., 2013). A recent study revealed that

CcmM35 and CcmM58 display similar distribution profiles in carboxysomes and are both 465 integrated within the core of the carboxysome (Niederhuber et al., 2017). Fluorescence 466 467 tagging at the protein C-terminus exploited in this work allowed us to only estimate the total amounts of CcmM but not distinguish CcmM35 and CcmM58, which can be addressed by N-468 terminal labeling of CcmM58 in our future study. Compared with the previous model that was 469 based on protein stoichiometry of cell lysates (Long et al., 2011), our relative quantifications 470 determined under the Air/ML condition show the 4.9-fold and 2.2-fold increases in the ratios of 471 Rubisco/CcmM and Rubisco/CcaA, respectively (Figure 6A, Supplemental Table 5). The 472 discrepancy may be caused by different sampling methods and cultivation conditions. 473

474

Based on immunoblot analysis of cell lysates, the previous model has proposed an 475 imbalanced ratio of RbcL to RbcS (~8:5), likely due to the binding of CcmM to Rubisco 476 477 replacing 3 RbcS subunits (Long et al., 2011). This result was similar to our immunoblot 478 quantification from cell lysates (Supplemental Table 2). Recent studies indicate that CcmM interacts with Rubisco (RbcL $_8S_8$) at distinct sites, without displacing RbcS (Ryan et al., 2018; 479 Wang et al., 2019). Based on the L_8S_8 ratio and RbcS abundance per carboxysome 480 determined, we estimate that there are approximately 853, 550, 367, and 1507 copies of 481 482 Rubisco per β -carboxysome under Air/ML, CO₂/ML, LL, and HL, respectively (Figure 6A, Table 1). Even the lowest Rubisco abundance per β -carboxysome (an average diameter of 483 484 151 nm) under LL is still greater than the Rubisco abundance per α -carboxysome (an average diameter of 123 nm) (lancu et al., 2007) by a factor of 1.6. This finding confirms the different 485 interior organization of the two classes of carboxysomes: densely packed with Rubisco 486 forming paracrystalline arrays inside the β -carboxysome (Faulkner et al., 2017) and random 487 packing of Rubisco in the α -carboxysome (lancu et al., 2007; lancu et al., 2010). The different 488 interior structures may be ascribed to their distinct biogenesis pathways: biogenesis of β-489 carboxysomes is initiated from the nucleation of Rubisco and CcmM35 and then the shell 490 encapsulation (Cameron et al., 2013); whereas α -carboxysome assembly appears to start 491 from shell formation (Menon et al., 2008) or a simultaneous shell-interior assembly (lancu et 492 al., 2010). 493

494

While the abundance of most of the structural components varies, the ratio of CcmK4 and CcmK3 is relatively unaffected (ranging from 3.6 to 4.1, Supplemental Table 5) under the tested growth conditions, implying their spatial colocalization within the carboxysome shell (Figure 6A). This is reminiscent of the recent observation that CcmK3 and CcmK4 can form a heterohexameric complex with a 1:2 stoichiometry and further form dodecamers in a pHdependent manner (Sommer et al., 2019). The *ccmK3* and *ccmK4* genes are located in the same operon that is distant from the *ccm* operon and they may have different expression regulation compared with other carboxysome components (Rae et al., 2012; Sommer et al., 2017). The balanced expression and structural cooperation of CcmK3 and CcmK4 may be crucial for the fine-tuning of carboxysome permeability towards environmental stress.

505

Rational design, construction and modulation of bioinspired materials with structural and 506 functional integrity are the major challenges in synthetic biology and protein engineering. 507 Given their self-assembly, modularity and high efficiency in enhancing carbon fixation, 508 carboxysomes have attracted tremendous interest to engineering this CO₂-fixing organelle 509 510 into other organisms, for example C_3 plants, with the intent of increasing photosynthetic efficiency and crop production (Lin et al., 2014b; Lin et al., 2014a; Occhialini et al., 2016; 511 Long et al., 2018). Recently, we have reported the engineering of functional β -carboxysome 512 structures in *E. coli* – a step towards constructing functional β -carboxysomes in eukaryotic 513 organisms (Fang et al., 2018). Our present study, by evaluating the actual protein 514 stoichiometry and structural variability of native β-carboxysomes, sheds light on the molecular 515 basis underlying the assembly, formation and regulation of functional carboxysomes. It will 516 empower bioengineering to construct BMC-based nano-bioreactors and scaffolds, with 517 functional and tunable compositions and architectures, for metabolic reprogramming and 518 targeted synthetic molecular delivery. A deeper understanding of carboxysome structure and 519 the developed imaging techniques will be broadly extended to other BMCs and 520 521 macromolecular systems.

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- 523

524 Materials and Methods

525 Bacterial strains, growth conditions, light and CO₂ treatment, and generation of 526 mutants

527 Wild-type (WT) and mutant *Synechococcus elongatus* PCC7942 (Syn7942) strains were 528 grown in BG-11 medium in culture flasks with constant shaking or on BG-11 plates containing 529 1.5% (w/v) agar at 30°C. Syn7942 WT and mutants were maintained and grown under 530 different intensities of constant white LED light illumination: 80 μ E·m⁻²·s⁻¹ as HL (higher light

in ambient air), 50 μ E·m⁻²·s⁻¹ as Air/ML (moderate light in ambient air), 10 μ E·m⁻²·s⁻¹ as LL (lower light in ambient air) to ensure full acclimation, respectively. Cultures were grown in air without an additional CO₂ source, except for the CO₂ treatment experiment in which Syn7942 cultures in the growth incubators were aerated with 3% CO₂ under moderate light (CO₂/ML).

536 Cultures were constantly diluted with fresh medium to maintain exponential growth phase for 537 the following imaging and biochemical analysis. *Escherichia coli* strains used in this work, 538 DH5a and BW25113, were grown aerobically at 30 or 37°C in Luria-Broth medium. Medium 539 supplements were used, where appropriate, at the following final concentrations: ampicillin 540 100 mg·mL⁻¹, chloramphenicol 10 mg·mL⁻¹, apramycin 50 mg·mL⁻¹, and arabinose 100 mM.

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535

All YFP-fusion mutants were generated following the REDIRECT protocol (Supplemental Figure 1) (Gust et al., 2002), by inserting the *eyfp:apramycin* DNA fragment to the C-terminus of individual carboxysome genes based on homologous recombination (Supplemental Table 6). Primers used in this work were listed in Supplemental Table 7. The same strategy was also applied for the mYPet mutant. For these mutant strains, BG-11 medium was supplemented with apramycin at 50 μ g·mL⁻¹.

548

549 Cell doubling time and growth curve measurement

550 Cultures were inoculated at OD_{750} of 0.05-0.1 with fresh BG-11. Growth of cells was 551 monitored at OD_{750} using a spectrophotometer (Jenway 6300 spectrophotometer, Jenway, 552 UK) every 24 hours. Doubling times were calculated using exponential phase of growth from 553 day 1 to day 4. Four biological replicates from different culture flasks were recorded. Data are 554 presented as mean ± standard deviation (SD). For each experiment, at least three biological 555 replicates from different culture flasks were analyzed.

556

557 Slimfield microscopy and data analysis

Live cells were applied at the small volume onto the BG-11 agarose pad at 0.25 mm thickness to maintain physiological growth, air dried to remove excessive medium and then assembled with plasma cleaned (Harrick-Plasma) glass cover slips. A dual-color bespoke laser excitation single-molecule fluorescence microscope was used utilizing narrow epifluorescence excitation of 10 µm full width at half maximum (FWHM) in the sample plane

to generate Slimfield illumination using narrowfield epifluorescence (Wollman and Leake, 563 2016; Wollman et al., 2016b; Wollman et al., 2017). This was incident on a sample mounted 564 565 on a Mad City Labs nanostage built on an inverted Zeiss microscope body consisting of a 20 mW 514 nm wavelength laser. A Chroma GFP/mCherry dichroic was mounted under the 566 Olympus 100x NA = 1.49 TIRF (total internal reflection fluorescence) objective, which delivers 567 10 mW excitation power. The image was split into YFP and chlorophyll channels using a 568 bespoke color splitter utilizing a Chroma dichroic split at 560 nm with 542 nm and 600 nm, 25 569 nm bandwidth filters. Imaging was done with an Andor iXon 128 x 128 pixel EMCCD camera 570 (iXon DV860-BI, Andor Technology, UK), at a pixel magnification of 80 nm/pixel using 5 ms 571 camera exposure time. Excitation intensity was initially reduced by 100x using and ND = 2 or 572 1 attenuation filter for high copy number strains (all except CcmL and RbcX) to avoid pixel 573 saturation on the EMCCD camera detector before a full-power photobleaching. Sample sizes 574 575 for individual strains are 60 (RbcS), 219 (CcmK3), 77 (CcmK4), 316 (CcmL), 71 (CcmM), 86 (CcmN), 95 (CcaA) and 211 (RbcX), respectively. Each population of carboxysomes comes 576 from 20-30 fields of view, with 1-7 cells per field of view. 577

578

579 The analysis was performed using bespoke MATLAB (Mathworks) software (Miller et al., 2015) with previously outlined methods (Llorente-Garcia et al., 2014; Wollman et al., 2016a; 580 Beattie et al., 2017; Lund et al., 2018; Stracy et al., 2018). In brief, candidate bright 581 fluorescent foci were identified in images using morphological transformation and thresholding. 582 The sub-pixel centroids of these foci were determined using iterative Gaussian masking and 583 their intensity quantified as the summed intensity inside a 5-pixel radius region of interest 584 (ROI) corrected for the mean background intensity inside a surrounding 17 x 17 pixel ROI 585 (Delalez et al., 2010; Leake, 2014). Foci were accepted and tracked through time if they had 586 a signal-to-noise ratio, defined as the mean intensity in the circular ROI divided by the 587 standard deviation in the outer ROI, over 0.4. The characteristic intensity of single 588 YFP/mYPet was measured from the distribution of detected foci intensity towards the end of 589 590 the photobleaching (Figure 1), confirmed by comparing the obtained value to individual photobleaching steps obtained using edge-preserving filtration (Figure 1) (Leake et al., 2003; 591 Leake et al., 2004). The stoichiometry of foci was then determined through cell-by-cell based 592 Slimfield imaging using numerical integration of pixel intensities (Wollman and Leake, 2015) 593 594 in each carboxysome divided by the intensity of a single YFP (Figure 1B).

595

596 For high-copy-number strains, intensity of carboxysomes was very high compared to the chlorophyll but for CcmL (typically ~2x, compare Supplemental Figure 3 with Supplemental 597 598 Figure 4A) the fluorescence intensity per carboxysome was comparable (although generally 599 brighter) to small regions of bright chlorophyll, detected as foci by our software, as confirmed by looking at the parental strain with no YFP present. To correct for this chlorophyll content, 600 we tracked parental WT Syn7942 cells as YFP-labelled cells to calculate the apparent 601 602 chlorophyll stoichiometry distribution (Supplemental Figure 4A). The CcmL distribution was then corrected by subtracting the apparent chlorophyll distribution. To investigate putative 603 periodic features in the stoichiometry distribution, we used the raw uncorrected values to 604 minimize dephasing artefacts (Figure 4C) using a kernel width of 0.5 molecules (equivalent to 605 the error in determining the characteristic intensity). The peak values in other strains were far 606 from the chlorophyll peak and so unaffected by this correction. 607

608

609 Confocal microscopy imaging and data analysis

Preparation of Syn7942 cells for confocal microscopy was performed as described earlier (Liu 610 et al., 2012; Casella et al., 2017). Cells were maintained under different growth conditions 611 prior to microscopy imaging, to ensure full acclimation. Confocal fluorescence images (12-bit, 612 512 x 512 pixels) were recorded using a Zeiss LSM780 with an alpha Plan-Fluor 100x oil 613 immersion objective (NA 1.45) and excitation at 514 nm from an Argon laser. YFP and 614 chlorophyll fluorescence were captured at 520-550 nm and 660-700 nm, respectively. The 615 image pixel size was 41.5 nm. The pixel dwell time was 0.64 µs and the frame averaging was 616 8, resulting in an effective frame time of ~ 1.5 s. The pinhole was set to give z axis resolution 617 of 1 µm. Live-cell confocal fluorescence images were recorded from at least five different 618 619 cultures. The sample stage was pre-incubated and thermo-controlled at 30°C before and during imaging. Zoom settings were set to have each carboxysome visualized with a 620 minimum of 8 x 8 pixels array to allow sufficient profiling of carboxysome signals by peak 621 intensity recognition and measurement. All images were captured with all pixels below 622 saturation. 623

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Confocal microscopic images were processed using FIJI Trackmate plugins (Tinevez et al., 2017) to retrieve peak intensities of carboxysomes based on the Find Maxima detection algorithm. Noise tolerance was determined by background intensities in empty regions. Imaging for different treatments in the same strain was performed under the same imaging

settings. For strains with visible cytosolic signals, the cytosolic background intensity was 629 determined by the average peak intensities in non-carboxysome regions over the central line 630 631 of the cell and was subtracted to obtain peak intensities. Raw data were processed by Origin Lab and MATLAB (Mathworks) for profile extraction and statistical analysis and the goodness-632 of-fit parameter for Violin plot visualization. Violin plots were generated by R to illustrate the 633 fluorescence intensity distribution of individual building proteins per carboxysome fitted by 634 kernel smooth fitting. The representative values and deviations of signal intensities were 635 represented by Peak value ± half width at half maximum (HWHM) measured from kernel 636 density fitted profiles, respectively. The significance of differences between treatments was 637 evaluated by Mann-Whitney U-tests pair-wisely (Supplemental Table 4). Standard errors of 638 sampling were determined through randomized grouping of intensity entries, with each group 639 containing a minimum of 70-100 entries. Errors were controlled below 5% to have accurate 640 641 estimation from the distributions. The relative protein abundance of carboxysomes was estimated by confocal imaging under Air/ML, CO₂/ML, LL, and HL was normalized by the 642 definite copy number of each strain under Air/ML determined by Slimfield imaging. 643

644

645 Live-cell time-lapse confocal imaging and data analysis

A 2 mm-thick BG-11 agar mat was prepared in stacked sandwiches to accommodate drops of 646 diluted Syn7942 cells. Cells were incubated on the BG-11 agar mat on the microscope for 1-2 647 hours before imaging. The continuous light illumination was provided at the intensity relatively 648 equal to HL, ML, or LL that were used for cell growth, in order to maintain cell physiology. The 649 650 same illumination was applied to the cells during time-lapse imaging with a hand-made module that switched off the light during laser scanning (less than 5 s per minute intervals). 651 The interval time was set to 60 s to guarantee sufficient light illumination between imaging. 652 The laser power was set to the minimum (1%) to reduce the bleaching for signals during long-653 term tracking. Images were initially corrected for horizontal drifting by Descriptor-based series 654 registration (2d/3d+T) plugin, and then were processed by the Trackmate plugin in FIJI for 655 particle tracking. Retrieved track data was analyzed using bespoke MATLAB (Mathworks) 656 scripts for MSD. Diffusion coefficient calculations and data visualization were modified as 657 previously described (Ewers et al., 2005; Sbalzarini and Koumoutsakos, 2005). Diffusion 658 coefficients were calculated by fitting the first 6 points of the MSD vs. τ curves. As the MSD 659 vs. τ curves indicated potentially non-Brownian diffusion at higher τ values, we described the 660 diffusion coefficients as "apparent diffusion coefficients". Tracking and diffusion coefficient 661

determination were tested by computational simulations (Supplemental Movie 2). Bespoke 662 Matlab code was written to generate simulated image stacks of carboxysomes diffusing inside 663 664 cells. Images were simulated by integrating a model 3D point spread function over a 3D model for the cell structure (Wollman and Leake 2015). This model comprises an inner 665 cytosol surrounded by thylakoid membranes (indicated by chlorophyll fluorescence) and 3 666 carboxysomes with a diameter of 200 nm. Each component's intensity was adjusted to match 667 real images before representative Poisson noise was applied. Carboxysomes were simulated 668 undergoing Brownian motion with a diffusion coefficient of $1.3 \times 10^{-5} \mu m^2 \cdot s^{-1}$ over 40 image 669 frames. Trackmate tracking and diffusion coefficient calculation yielded a mean diffusion 670 coefficient of $1.32 \pm 0.02 \times 10^{-5} \mu m^2 \cdot s^{-1}$, giving a 1.5% error. 671

672

673 Immunoblot analysis

Immunoblot examination was carried out following the procedure described previously (Sun et 674 al., 2016). 150 µg of cell lysate, measured by Pierce Coomassie (Bradford) Protein Assay Kit 675 (Thermo Fisher Scientific), was loaded on 10% (v/v) denaturing SDS-PAGE gels. Immunoblot 676 analysis was performed using the primary mouse monoclonal anti-GFP (Invitrogen, 33-2600), 677 capable of recognizing series of GFP variants including YFP, the rabbit polyclonal anti-RbcL 678 (Agrisera, AS03 037), the horseradish peroxidase-conjugated goat anti-mouse IgG secondary 679 antibody (Promega, W4021) and a Goat anti-Rabbit IgG (H&L), HRP conjugated (Agrisera 680 AS10 1461). Anti-CcmK2 antibody was kindly provided by the Kerfeld lab (Michigan State 681 University, US) (Cai et al., 2016). Protein quantification from immunoblot data was carried out 682 using FIJI. Our nominal assumption that the ratios of YFP-tagged to total RbcL or CcmK2 in 683 carboxysomes are similar to those in cell lysates. 684

685

686 *In vivo* carbon fixation assay

In vivo carbon fixation assay was carried out to determine carbon fixation of Syn7942 WT and mutant cells, as described in the previous work (Sun et al., 2016). For each WT and mutant, at least three biological replicates from different culture flasks were assayed. Significance was assessed by two-tailed Student's t-tests.

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692 Electron microscopy and carboxysome size measurement

Electron microscopy was carried out as described previously (Liu et al., 2008; Sun et al., 2016). Carboxysome diameter was measured as described previously (Faulkner et al., 2017) and was analyzed using Origin.

696

697 Accession Numbers

698 Accession numbers of genes in this article are provided in Supplemental Table 6.

699

700 Supplemental Data

- Supplemental Figure 1. Construction and verification of Syn7942 strains with YFP fusion to
 individual carboxysome proteins.
- **Supplemental Figure 2.** Immunoblot analysis of the YFP-tagged Syn7942 strains using the anti-GFP, anti-RbcL and anti-CcmK antibodies of soluble fractions in this study based on SDS-PAGE.
- Supplemental Figure 3. Slimfield images of YFP-fusion cells under Air/ML and stoichiometric
 histogram of copies of YFP molecules per carboxysome.
- Supplemental Figure 4. Normalization of chlorophyll during Slimfield imaging for Syn7942
 strains.
- **Supplemental Figure 5.** Comparison of YFP and mYPet tagging to RbcL reveals no differences in carboxysome localization, cell growth and carbon fixation, suggesting that there
- are no measurable artefacts due to putative effects of dimerization of the YFP tag.
- 713 **Supplemental Figure 6.** Confocal images of YFP-tagged cells.
- 714 Supplemental Figure 7. Confocal images of RbcS-YFP, CcmM-YFP, CcmK4-YFP and
- 715 CcmK3-YFP cells under Air/ML, CO₂, LL, and HL and distribution profiles of carboxysome 716 protein signal intensity.
- 717 Supplemental Figure 8. Confocal images of CcmL-YFP, CcmN-YFP, CcaA-YFP and RbcX-
- 718 YFP cells under Air/ML, CO₂, LL, and HL and distribution profiles of carboxysome protein 719 signal intensity (continuing Supplemental Figure 7).
- Supplemental Figure 9. Thin-section EM images of WT Syn7942 cells under Air/ML, CO₂/ML,
 LL and HL.
- 722 **Supplemental Figure 10.** Changes in the diffusion coefficient of carboxysomes in Syn7942
- cells under HL, ML and LL are not dependent on the carboxysome size.
- 724 **Supplemental Figure 11.** CcmP-YFP and CcmO-YFP Syn7942 cells.

- 725 Supplemental Table 1. Cell growth, carbon fixation and cell dimensions of Syn7942 WT and
- 726 YFP-fusion mutants under Air/ML.
- Supplemental Table 2. Immunoblotting estimation of the stoichiometry of carboxysomal
 proteins in cell lysates.
- 729 Supplemental Table 3. Carboxysome content per cell under Air/ML, CO₂/ML, LL and HL
- 730 determined by confocal imaging.
- 731 **Supplemental Table 4.** Evaluation and quality control of quantitative microscopy.
- 732 Supplemental Table 5. Carboxysome properties in Syn7942 vary under Air/ML, CO₂/ML, LL
- and HL, determined by Slimfield, confocal and EM imaging.
- 734 **Supplemental Table 6.** Accession numbers for genes/proteins in this work.
- 735 **Supplemental Table 7.** PCR primers used in this study for gene cloning and sequencing.
- **Supplemental Movie 1.** Time-lapse confocal imaging reveals different diffusion dynamics of carboxysomes in the RbcL-YFP Syn7942 cells grown under HL, ML and LL conditions.
- Supplemental Movie 2. Simulations of diffusing carboxysomes *in cellulo* validate tracking
 and diffusion coefficient determination.
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741 Acknowledgements

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754 Author contributions

L.-N.L and M.C.L designed research; Y.Q. A.J.M.W and F.H. performed research and analyzed data; L.-N.L, Y.Q., M.C.L., and A.J.M.W. wrote the paper.

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759 Competing interests

The authors declare no conflict of interest.

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763 Figure legends

Figure 1. Slimfield quantification of cells grown under ambient air/moderate light Air/ML conditions.

- (A) Averaged Slimfield images of YFP fluorescence (green) over 5 frames of strains
 expressing shell component CcmK4-YFP, the interior enzyme RbcS-YFP, and the shell interior linker protein CcmM-YFP. White dashed lines indicate cell body outlines.
- **(B)** Distribution of the intensities of automatically detected foci from the end of photobleaching, corresponding to the characteristic intensity of *in vivo* YFP. Inset shows the Fourier spectrum of 'overtracked' foci, tracked beyond photobleaching, showing a peak at the characteristic intensity.
- (C) Representative fluorescence photobleaching tracked at ultra-fast speed. The CcmK4 plot
 shows an inset 'zoomed in' on lower intensity range with step-preserving Chung-Kennedy
 filtered data in red, showing individual photobleaching steps clearly visible at the
 characteristic intensity. Brightness (kcounts), counts measured per camera pixel multiplied by
 1,000.
- (D) Distribution of YFP copy number detected for individual carboxysomes in corresponding
 mutants, rendered as kernel density estimates using standard kernel width. Heterogeneity of
 contents was observed, and a "preferable" copy number, represented by kernel density peak
 values could be determined. Statistics of copy numbers (Peak value ± HWHM) are listed in
 Table 1 for ML conditions. The corresponding Slimfield images and histogram for complete
 strain sets are shown in Supplemental Figure 3.
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Figure 2. Relative protein quantification of CcmK4, RbcS and CcmM in the carboxysome under different CO₂ levels and light intensities using confocal microscopy.

- (A) Confocal images of CcmK4-YFP, RbcS-YFP and CcmM-YFP strains under Air/ML,
 CO₂/ML, LL and HL. Fluorescence foci (green) indicate carboxysomes, and cell borders were
 outlined by white dashed lines. Scale bar indicates 2 µm.
- **(B)** Violin plot of carboxysome intensities under Air/ML, CO₂/ML, LL and HL, normalized to kernel density ML peak values (peaks marked by white dashed lines).
- (C) Kernel density estimates of CcmL carboxysome copy number grown under Air/ML, CO₂,
 LL and HL detected by Slimfield and corrected for chlorophyll. Triple Gaussian fits are
 indicated as colored dashed lines with the summed fit in red. The percentage in each
 Gaussian is indicated aside.
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Figure 3. Changes in carboxysome protein stoichiometry upon increases in CO₂ levels and light intensity.

(A) Comparison of carboxysome protein stoichiometry under CO_2 treatment. Increase in the CO_2 concentration resulted in the rise of CcmK3, CcmK4, CcaA and CcmL contents and the decline of RbcS, CcmN and CcmM contents.

(B) Comparison of carboxysome protein stoichiometry under light intensity treatment. Increased light intensity led to the elevation of RbcS, CcmM, CcmL, CcaA and CcmN contents, whereas the abundance of RbcX, CcmK3 and CcmK4 contents per carboxysome did not change dramatically.

Mann-Whitney U-tests were performed to compare the numbers of functional units of individual carboxysome proteins changed from CO₂/ML to Air/ML (A) and from HL to LL (B). *, p < 0.05; ***, p < 0.005; ns, p > 0.05.

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Figure 4. Variations of the carboxysome size and carbon fixation under Air/ML, CO₂, LL and HL.

- (A) Thin-section electron microscopy (EM) images showing individual carboxysomes in the
 Syn7942 WT cells under Air/ML, CO₂, LL and HL treatments Yellow arrows indicate the
 carboxysomes with spaces of low protein density under LL. More EM images are shown in
 Supplemental Figure 9. Scale bar indicates 1 μm.
- (B) Changes in the carboxysome diameter under Air/ML, CO₂, LL and HL measured from EM (n = 33, 25, 27 and 51, respectively), with representative carboxysome images depicted above. Dashed lines indicate medians and solid lines indicate means. Differences in the carboxysome diameter are significant between CO₂ and air ($p = 1.92 \times 10^{-14}$) and between LL and HL ($p = 8.29 \times 10^{-7}$), indicated as ***.
- 822 **(C)** Correlation between the carboxysome size and the Rubisco content per carboxysome 823 under Air/ML, CO₂, LL and HL.
- (**D**) Correlation between the carboxysome size and CO₂ fixation per carboxysome.
- (E) Correlation between the carboxysome size and CO_2 fixation per Rubisco of the carboxysomes. Carboxysome diameters and CO_2 fixation are presented as average \pm SD, whereas the carboxysome total protein content and Rubisco content are shown as Peak value \pm HWHM.
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Figure 5. Spatial localization and diffusion dynamics of carboxysomes in Syn7942 cells are dependent on light intensity.

- (A) Tracking of carboxysome diffusion in cells grown under HL, ML and LL. Colored lines
 indicate the diffusion trajectories of each carboxysomes and circles represent the diffusion
 areas of each carboxysomes over 60 mins. Scale bar indicates 1 µm.
- (B) Non-linear MSD (Mean Square Displacement) *vs.* the time interval (τ) profiles suggest the mobility of carboxysomes in Syn7942 cells grown under HL, ML and LL. Inset, zoom-in view of the MSD profile under LL.
- (C) Diffusion coefficient of carboxysomes *in vivo* decreases significantly when the light intensity reduces: $2.76 \pm 2.83 \times 10^{-5} \mu m^2 \cdot s^{-1}$ for HL (mean ± SD, *n* = 105), $1.48 \pm 1.03 \times 10^{-5}$
- 840 $\mu m^2 \cdot s^{-1}$ for ML (*n* = 84), and 0.28 ± 0.19 x 10⁻⁵ $\mu m^2 \cdot s^{-1}$ for LL (*n* = 336). *p* = 3.05 x 10⁻⁵ 841 between HL and ML; *p* = 2.77 x 10⁻⁵ between ML and LL, two-tailed Student's t-test).
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Figure 6. Model of the β -carboxysome structure and protein stoichiometry.

(A) Diagram of an icosahedral carboxysome structure and organization of building components. The stoichiometry of each building component within the carboxysome and its variations in response to changes in CO_2 and light intensity are shown on the right (See also Table 1). *Rubisco content was estimated from RbcS stoichiometry based on the RbcL₈S₈ Rubisco structure. The majority of shell facets shown in light blue is tiled by the major shell

protein CcmK2. The total abundance of CcmM58 and CcmM35 was estimated. The components RbcL, CcmK2, CcmO and CcmP were not directly determined in this work and thus are not shown in this model.

(B) The carboxysome diameter is variable in response to changes in the CO_2 level and light intensity.

Table 1. Protein stoichiometry of the Syn7942 β-carboxysome and its variability in cells grown under Air/ML, CO₂/ML, LL and HL 855 conditions determined from Slimfield and confocal microscopy. Stoichiometry is presented as Peak value ± HWHM and the sample sizes 856 are indicated as n. Peak values were determined from Slimfield stoichiometry profiles of each carboxysome proteins (Figure 1, 857 Supplemental Figure 3). Quantification of CcmL under the four conditions was acquired from Slimfield for accurate measurement of 858 copies of shell pentamers for capping the carboxysome structure. Copies of other carboxysome proteins were calculated using Slimfield 859 results (bold) with definitive counts of protein copies under Air/ML (See also Supplemental Figure 3) in combination with relative 860 quantification of each protein under the four conditions from confocal imaging (See also Supplemental Figure 7 and 8). Protein structures 861 were derived from previous studies (Kerfeld et al., 2005; Long et al., 2007; Tanaka et al., 2007; Tanaka et al., 2008; Long et al., 2011; 862 Kinney et al., 2012; McGurn et al., 2016). *Monomeric unit of CcmM was designated to CcmM35 that is the majority of CcmM; CcmM58 863 is postulated as a trimer. 864

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| | Structure | Protein | Air/ML | | CO ₂ /ML | | LL | | HL | |
|----------------------|-----------|---------|---------------------------------|----------------------------------|----------------------------------|----------------------------------|----------------------------------|----------------------------------|------------------------------------|----------------------------------|
| Category | | | Peak value ± HWHM | Number of functional units | Peak value ± HWHM | Number of functional units | Peak value ± HWHM | Number of functional units | Peak value ± HWHM | Number of functional units |
| Shell proteins | Hexamer | CcmK3 | 92 ± 148 (<i>n</i> = 219) | 15 ± 25 | 172 ± 83 (<i>n</i> = 2048) | 29 ± 14 | 83 ± 31 (<i>n</i> = 1516) | 14 ± 5 | 87 ± 52 (<i>n</i> = 2155) | 14 ± 9 |
| | | CcmK4 | 314 ± 194 (<i>n</i> = 77) | 52 ± 32 | 562 ± 263 (<i>n</i> = 1918) | 94 ± 44 | 313 ± 121 (<i>n</i> = 1766) | 52 ± 20 | 304 ± 95 (<i>n</i> = 3215) | 51 ± 16 |
| | Pentamer | CcmL | 37 ± 17 (<i>n</i> = 316) | 7.4 ± 3.4 | 66 ± 24 (<i>n</i> = 311) | 13.2 ± 4.8 | 34 ± 15 (<i>n</i> = 394) | 6.8 ± 3.0 | 69 ± 24 (<i>n</i> = 220) | 13.8 ± 4.8 |
| Structural proteins | Monomer* | CcmM | 719 ± 1433 (<i>n</i> = 71) | 719 ± 1433 | 468 ± 425 (<i>n</i> = 2313) | 468 ± 425 | 483 ± 366 (<i>n</i> = 3655) | 483 ± 366 | 1176 ± 691 (<i>n</i> = 2318) | 1176 ± 691 |
| | Monomer | CcmN | 74 ± 51 (<i>n</i> = 86) | 74 ± 51 | 52 ± 28 (<i>n</i> = 3143) | 52 ± 28 | 51 ± 20 (<i>n</i> = 4022) | 51 ± 20 | 82 ± 34 (<i>n</i> = 5074) | 82 ± 34 |
| CA | Hexamer | CcaA | 86 ± 81 (<i>n</i> = 95) | 14 ± 14 | 129 ± 86 (<i>n</i> = 1354) | 21 ± 14 | 65 ± 21 (<i>n</i> = 217) | 11 ± 4 | 122 ± 59 (<i>n</i> = 2837) | 20 ± 10 |
| Rubisco enzyme | L_8S_8 | RbcS | 6822 ± 9200 (<i>n</i> = 60) | 853 ± 1150 | 4401 ± 6655 (<i>n</i> = 894) | 550 ± 832 | 2934 ± 5492 (<i>n</i> = 752) | 367 ± 687 | 12057 ± 5186 (<i>n</i> = 1974) | 1507 ± 648 |
| Rubisco chaperone | Dimer | RbcX | 39 ± 32 (<i>n</i> = 211) | 20 ± 16 | 38 ± 10 (<i>n</i> = 1370) | 19 ± 5 | 40 ± 9 (<i>n</i> = 1402) | 20 ± 5 | 40 ± 9 (<i>n</i> = 1861) | 20 ± 5 |

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