Abstract

Cyanobacterial CO₂-fixation is supported by a CO₂-concentrating mechanism which improves photosynthesis by saturating the primary carboxylating enzyme, ribulose 1, 5-bisphosphate carboxylase/oxygenase (RuBisCO), with its preferred substrate CO₂. The site of CO₂-concentration is a protein bound micro-compartment called the carboxysome which contains most, if not all, of the cellular RuBisCO. The shell of β-type carboxysomes is thought to be composed of two functional layers, with the inner layer involved in RuBisCO scaffolding and bicarbonate dehydration, and the outer layer in selective permeability to dissolved solutes. Here, four genes (ccmK2-4, ccmO), whose products were predicted to function in the outer shell layer of β-carboxysomes from Synechococcus elongatus PCC 7942, were investigated by analysis of defined genetic mutants. Deletion of the ccmK2 and ccmO genes resulted in severe high-CO₂-requiring mutants with aberrant carboxysomes, whilst deletion of ccmK3 or ccmK4 resulted in cells with wild-type physiology and normal ultrastructure. However, a tandem deletion of ccmK3-4 resulted in cells with wild-type carboxysome structure, but physiologically deficient at low CO₂ conditions. These results revealed the minimum structural determinants of the outer shell of β-carboxysomes from this strain: CcmK2, CcmO and CcmL. An accessory set of proteins was required to refine the function of the pre-existing shell: CcmK3 and CcmK4. These data suggested a model for the facet structure of β-carboxysomes with CcmL forming the vertices, CcmK2 forming the bulk facet, and CcmO, a “zipper protein,” interfacing the edges of carboxysome facets.

Introduction

Cyanobacteria (blue-green algae) have a CO₂-concentrating mechanism (CCM) which provides a growth advantage under conditions of limiting inorganic carbon (Ci; principally CO₂ and HCO₃⁻ in aquatic environments). The CCM has two functional features of note: the presence of a set of transcriptionally and post-translationally regulated Ci uptake systems which accumulate Ci as a cytoplasmic bicarbonate pool [1]; and the carboxysome, an icosahedral, protein-bound micro-compartment in which the action of carboxysomal carbonic anhydrase (CA) enzymes provides a CO₂-rich micro-environment that supports the carboxylation reaction of d-ribulose 1,5-bisphosphate carboxylase/oxygenase (RuBisCO: EC: 4.1.1.39), thereby reducing the wasteful oxygenation reaction [2,3]. Thus the concerted action of the cyanobacterial CCM improves photosynthetic rate under Ci limitation and reduces the energetic impact of the photosynthetic pathway.

Two evolutionarily distinct forms of carboxysome exist for which two monophyletic groups of cyanobacteria are named [4–6]. The α-type β-carboxysome is found in α-cyanobacteria and probably first arose in chemo-autotrophic bacteria [4,7], whereas the ccm-type β-carboxysome is found only in β-cyanobacteria [4,6]. α- and β-carboxysomes appear to have evolved in parallel [5]. Whilst some of their constituent small proteins bear marked similarity to each other, other larger structural proteins have no observable similarity [2–4]. Specifically, the RuBisCO enzyme and the proteins predicted to form a semi-permeable shell structure are each distinct lineages of the same evolutionarily-related proteins [4,6,8,9].

Whilst details of the interior structure of α-carboxysomes are scarce, these bodies are known to contain form-1A RuBisCO as well as a number of shell-associated proteins [10] including that presumed to organise the interior, CsoS2 [11], and the α-carboxysomal CA enzyme, CsoSCA [12,13]. In contrast, the interior structure of β-carboxysomes is better understood [2,3,14]. In Synechococcus elongatus PCC 7942 and Synechocystis PCC 6803, an inner-shell bicarbonate dehydration complex and an interior para-crystalline matrix of form-1B RuBisCO molecules are linked and organised by at least two isoforms of the CcmM protein [15–17]. Current models of β-carboxysome organisation place a semi-permeable shell structure on the cytoplasmic face of a second, sub-shell, layer which contains the carboxysomal carbonic anhydrase enzymes [3,14–19], thus two functionally distinct shell layers are present in β-carboxysomes: the semi-permeable shell layer and the bicarbonate dehydration complex/RuBisCO organising layer.

The outer shell layer of bacterial micro-compartments is thought to consist of oligomeric proteins containing the Bacterial
Micro-Compartment (BMC) domain (pfam: PF00936) with the first known sequence for this family originating from *S. elongatus* PCC 7942 [20]. In crystal structures, BMC protomers form flattened hexagonal oligomers (hexamers or trimers of proteins containing one or two BMC domains respectively) which themselves tessellate into sheet or strip-like higher-level oligomers [18,19,21–27]. Current structural models suggest that these sheets and/or strips of BMC oligomers form the facets of the icosahedral carboxysome, the vertices being closed by pentameric proteins that contain a distinct type of protein domain (pfam: PF03319) [9,19,28,29]. Charged pores at the six-fold axis of symmetry of BMC proteins are thought to underlie the selective permeability of the outer carboxysome shell [9,30]. In addition to their potential as pores for charged solute transit, some BMC proteins have a complex pore conformation in crystal structures, evoking the potential for gated transit of larger metabolites [26]. On the other hand, some BMC proteins from related types of micro-compartment involved in ethanolamine and propanediol detoxification have absent or obfuscated pores [21–23,25].

In *S. elongatus* PCC 7942, multiple low-mass bands corresponding to BMC proteins are sometimes observed in SDS-PAGE and western immunoblots, though there is a single band under normal conditions [15]. In *PCC 7942*. In *PCC 7942*, however, multiple shell proteins are evident in SDS-PAGE and western immunoblots [10,12,31], thus the complexity of the putative outer-shell complex appears to differ between carboxysome type and experimental conditions.

Significantly, cyanobacterial genomes can contain up to nine genes whose products contain a recognisable BMC domain (Table S1), however the specific roles that each of these proteins play, if any, is unclear [9]. Recent evidence suggests that simple micro-compartmental interactions can be derived from the expression of a single BMC gene [32], so the shell structures of carboxysomes are not necessarily complex. Why then, do all cyanobacteria have between three and nine BMC homologues? It is well known that genes in the ‘core’ *ccm* operon *ccmK2* and *ccmK3* are essential for biogenesis of *β*-carboxysomes in *S. elongatus* PCC 7942 [20,33–42]. Of these, *ccmK2* and *ccmO* have obvious BMC domains, indeed the *β*-cyanobacterial model *S. elongatus* PCC 7942 contains four genes with products containing BMC domains, *ccmK2* (named for sll1028 from *Synechocystis* PCC 6803, previously referred to as *ccmK* or *ccmK1*), *ccmK3*, *ccmK4* and *ccmO*, with the latter containing two BMC domains in tandem. Of this group, only CcmK2 has been observed in *β*-carboxysome-enriched fractions [15,16,43], thus other potential BMC proteins have no predicted functional role. Whilst fluorescently tagged CcmK4 proteins were shown to possess carboxysomes whose appearance and dimensions were compatible with the selectivity permeability of the outer carboxysome shell [9,30]. In addition to their potential as pores for charged solute transit, some BMC proteins have a complex pore conformation in crystal structures, evoking the potential for gated transit of larger metabolites [26]. On the other hand, some BMC proteins from related types of micro-compartment involved in ethanolamine and propanediol detoxification have absent or obfuscated pores [21–23,25].

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### Results

Following initial growth testing, the carboxysomal ultrastructure of genetic mutants in BMC genes was investigated by transmission electron microscopy, and the protein components of wild-type and mutant *β*-carboxysomes identified there were further studied by carboxysome purification and western immunoblot analysis. To examine the physiological consequence of *β*-carboxysome shell perturbation the same strains were also assessed for their photosynthetic affinity for CO₂ and growth rates under CO₂-replete and limited conditions. To maintain consistency with prior literature, the partially characterised HIND insertion mutant of *ccmK2* [20] was subjected to the same analyses but does not form a major part of the analyses presented here. In light of the similarity of the Δ*ccmK3* and Δ*ccmK4* mutants to wild type and their lack of a high-CO₂ requiring phenotype, these genes were subsequently deleted in tandem. The resulting double-deletion mutant, Δ*ccmK3*-4, was subjected to the same analyses and, as described below, was deficient in some aspects of its CCM.

### Ultrastructural phenotypes of carboxysome shell mutants

Transmission electron microscopy revealed two classes of ultrastructural phenotype in the mutants generated (Figure 1, Table 1), namely normal wild-type appearance or aberrant polar bodies. The wild-type strain had carboxysomes whose cross-sectional diameter (175±23 nm, Figure 1-A) compared well to previously published studies (172±26 nm [16]).

Like the wild-type strain, the Δ*ccmK3* and Δ*ccmK4* mutants possessed carboxysomes whose appearance and dimensions compared favourably to wild-type (167±26 and 169±26 nm respectively, Figure 1-E, F). The Δ*ccmK3*-4 mutant had carboxysomes with a mean diameter of 183±24 nm. However, the spatial arrangement of carboxysomes within the cell was altered in this strain with carboxysomes aggregated (Figure 1-G) rather than well-separated as observed in wild-type cells (Figure 1-A).

In contrast to the wild-type-like mutants, the HIND, Δ*ccmK2* and Δ*ccmO* mutants possessed large, polar, electron-dense structures which lacked the characteristic faceted geometry of carboxysomes (Figure 1-C,D, H, Table 1). The maximum cross-sectional diameter of these bodies was much greater than wild-type, though still smaller than wild-type *β*-carboxysomes from some *β*-cyanobacterial species [46].

Occasional aberrant carboxysomes were observed in a complemented mutant strain generated in this study, Δ*ccmO* + pH6-Ub-*ccmO* (Figure S1-C), however, the majority of carboxysomes in this...
The Outer Shell of β-Carboxysomes

strain, whilst being slightly larger than wild-type (208±49 nm), appeared normal and the strain was not physiologically impaired (Figure S2). Nonetheless, occasional aberrant carboxysomes observed in this strain included rod-like carboxysomes reminiscent of those described for the PVU mutant \(\text{ccmL}::\text{CmR}\) \(\text{PCC} 7942\). A, Wild-type \(\text{PCC} 7942\). B, \(\Delta\text{ccmM}\) mutant \(\text{ccmK2}::\text{CmR}\ \text{PCC} 7942\). C, \(\Delta\text{ccmK2}\) mutant \(\text{ccmK2}\) \(\text{PCC} 7942\). D, \(\Delta\text{ccmK3}\) mutant \(\text{ccmK3}::\text{CmR}\ \text{PCC} 7942\). E, \(\Delta\text{ccmK3}::\text{CmR}\) mutant \(\text{ccmK3}::\text{CmR}\ \text{PCC} 7942\). F, \(\Delta\text{ccmK4}\) mutant \(\text{ccmK4}::\text{CmR}\ \text{PCC} 7942\). G, \(\Delta\text{ccmK3}::\text{CmR}\) mutant \(\text{ccmK3}::\text{CmR}\ \text{PCC} 7942\). H, \(\Delta\text{ccmO}\) \(\text{PCC} 7942\). Scale bars are 500 nm and HIND and \(\Delta\text{ccmK4}\) were embedded in Eponaralite rather than LR-white resin.

doi:10.1371/journal.pone.0043871.g001

Physiological phenotypes of BMC mutants

Although two types of carboxysome ultrastructure were observed, three classes of physiological phenotype were apparent within the mutants generated in this study (Table 1, Figure 2). Mutants were assessed for differences in photosynthetic affinity for \(\text{Ci}\) \(K_{1/2} \text{Ci}\) derived from oxygen evolution curves plotted against \(\text{Ci}\) concentrations \(\text{47}\). The \(\Delta\text{ccmK3}\) \(K_{1/2} \text{Ci}\) \(0.35±0.05 \text{mm}\) and \(\Delta\text{ccmK4}\) \(0.34±0.04 \text{mm}\) mutants had \(\text{Ci}\)-uptake physiology that closely matched the wild-type strain \(0.27±0.05 \text{mm}\). In contrast, the polar body mutants HIND \(17.8±1.5 \text{mm}\), \(\Delta\text{ccmK2}\) \(16.5±0.6 \text{mm}\) and \(\Delta\text{ccmO}\) \(17.3±0.6 \text{mm}\) had very highly attenuated CCM activities (Figure 2), which agreed with the physiological findings of previous studies of \(\text{ccmK2}\) \(\text{20}\) and \(\text{ccmO}\) mutants \(\text{41,42}\). Indeed, the photosynthetic physiology of these mutants was similar to \(\Delta\text{ccmM}\) which completely lacks carboxysomes \(\text{16}\). A third, unusual phenotype was observed for the \(\Delta\text{ccmK3}::\text{CmR}\) mutant which displayed a photosynthetic physiology that was intermediate to the wild-type and carboxysome mutants \(10.0±2.3 \text{mm}\;\text{Figure 2}\). Hence, although essentially a high-\(\text{CO}_2\) requiring mutant, \(\Delta\text{ccmK3}::\text{CmR}\) could photosynthesise and grow in air.

These physiological findings were supported by the comparative growth rate analysis of these mutant strains under conditions of high and low \(\text{CO}_2\) (Table 1). All strains grew at a similar rate under 4% \(\text{CO}_2\) where the doubling times ranged from 7.7±0.7 to 10.8±0.5 h. The wild-type, \(\Delta\text{ccmK3}\) and \(\Delta\text{ccmK4}\) strains also grew at a similar rate in air \(9.6±0.4 – 10.7±0.6 \text{h}\). However, the carboxysome mutants HIND, \(\Delta\text{ccmK2}\) and \(\Delta\text{ccmO}\) were not capable of growth without \(\text{Ci}\) supplementation. Like its \(\text{Ci}\)-uptake phenotype, the \(\Delta\text{ccmK3}::\text{CmR}\) mutant grew in air, albeit very slowly (doubling time 29.9±0.9 h).

Protein composition of isolated β-carboxysome preparations

The protein components of carboxysomes and polar bodies were assessed by western immunoblots against protein extracts enriched in carboxysomes using the \(\text{Mg}^{2+}\) precipitation and Triton X-100-Percoll (TP pellet) methods \(\text{15,36,43,48,49}\). Since approximately 73% of CcmK2 remains in the insoluble pellet fraction after cell lysis \(\text{14}\), the Triton X-100-Percoll purification was appropriate for purification of carboxysome-like polar bodies because the effectiveness of this method is independent of the presence of an intact native outer shell structure. As described below, the outcomes of both Triton X100-Percoll purification closely matched those from Mg2+ precipitation which is a structure independent method for visualising carboxysomal proteins, thus validating these assumptions.

The pattern of carboxysome protein presence and absence from TP pellets (Figure 3), and \(\text{Mg}^{2+}\) fractions (Figure S3), was similar for all strains. In terms of the inner shell bicarbonate dehydration layer and carboxysome lumen, all detectable components of the RuBisCO organising complex (CcaA, CcmM-58, CcmM-35, and RuBisCO) were present in carboxysomes and polar bodies (Figure 3, Figure S3), however the presence of CcmN could not be reliably confirmed using our low-titre antibody. The 35 kDa form of CcmM, which predominantly links RuBisCO holoenzymes into an interior matrix, was present in the carboxysome enriched fractions of all of the mutants, even those possessing polar bodies.

CcmK2 and CcmO were not detected in TP pellets of \(\text{ccmK2}\) and \(\text{ccmO}\) mutants respectively, and a relationship was shown between CcmK2 and CcmO such that the absence of one of this pair resulted in the absence of the other (Figure 3). This pattern was in-part repeated in Mg2+-precipitated fractions, where the presence of CcmO was dependent on the presence of CcmK2 without the reverse situation being true (Figure S3). Intriguingly, for these mutants the CcmO or CcmK2 proteins were not found in the Mg2+ supernatant. Thus CcmO, like CcmK2 \(\text{14}\) may largely be lost from the shell and aggregated after cell breakage, especially in the absence of other shell proteins. This is the first evidence of a
valid structural interaction between CcmK2 and CcmO in the β-carboxysome and is not that surprising given that these proteins share the BMC protein domain.

**IMAC purification of interacting BMC shell proteins**

To investigate the observed relationship between CcmK2 and CcmO in the shell of β-carboxysomes (Figure 3), immobilised metal affinity chromatography (IMAC) was used to IMAC-purify hexahistidine tagged CcmO proteins along with any protein-interaction partners from ΔccmO + pSE2-H6-Ub-CcmO (Figure 4). To enhance resolution of complexed proteins, the analysis was pre-enriched for carboxysome proteins by using TP pellets prepared from ΔccmO + pSE2-H6-Ub-CcmO as a substrate for IMAC. Tagged and complexed proteins were identified by western immunoblots. As observed by Long et al. [16], partial denaturation with 1.0 M urea was required for IMAC co-purification of β-carboxysome protein complexes. This confirms previous work showing that BMC proteins like CcmK2 and CcmO form extremely stable oligomers which often cannot be dissociated, even with high urea concentrations [14–16]. Thus a small amount of urea is necessary to sufficiently solubilise or weaken the BMC protein oligomers, allowing effective purification of these proteins and their binding partners from the outer shell of β-carboxysomes.

**Table 1. Ultrastructural, physiological and growth parameters for BMC mutants.**

| Genotype     | Carboxysome type and diameter | $K_{1/2}$ (Cl) (mm) | Doubling time (h) | Physiology$^a$ |
|--------------|-------------------------------|---------------------|-------------------|----------------|
| PCC 7942     | CBX 175±37 (35)               | 0.27±0.05           | 8.6±0.4           | 10.7±0.6       | WT |
| HIND$^c$     | PB 318±56 (4)                | 17.8±1.5            | 8.4±0.1           | n.g.           | HCR |
| ΔccmK2       | PB 361±78 (56)               | 16.5±0.6            | 10.8±0.5          | n.g.           | HCR |
| ΔccmK3       | CBX 167±26 (50)              | 0.34±0.04           | 9.0±0.3           | 10.0±0.2       | WT |
| ΔccmK4       | CBX 169±36 (33)              | 0.35±0.05           | 7.7±0.7           | 9.6±0.4        | WT |
| ΔccmK3-4     | CBX 183±24 (50)              | 10.0±2.3            | 28.9±0.9          | HCR$^b$       |
| ΔccmO        | PB 302±54 (56)               | 17.3±0.6            | 9.3±0.7           | n.g.           | HCR |
| ΔccmM$^b$    | None n.a.                    | 19.2±2.38           | 8.1±0.4           | n.g.           | HCR |

Carboxysome diameters are the mean and standard deviation of a sample size shown in brackets and photosynthetic half-saturation constant $K_{1/2}$ (Cl) and growth rates are expressed as the mean and standard deviation of at least three replicate cultures.

$^a$, Wild-type like carboxysomes (CBX) and carboxysome-like polar bodies (PB).

$^b$, Wild-type like (WT) or high CO2 requiring (HCR).

$^c$, The partially characterised HIND mutant (ccmK2::CmR) [20] and the carboxysomeless ΔccmM mutant [37] are included for reference.

$^d$, ΔccmK3-4 has an intermediate physiology which is essentially HCR.

n.g., no growth.

n.a., ΔccmM has no carboxysome ultrastructure.

doi:10.1371/journal.pone.0043871.t001

**Figure 2. Photosynthetic oxygen evolution in response to external Cl by BMC shell mutants.** Shown are a representative set of mass-spectrometric measurements of Cl-dependent O2 evolution by wild type and mutant strains of S. elongatus PCC 7942 over a range of Cl concentrations. Cl is the sum of CO2 and HCO3 in solution (pH 7.9).

doi:10.1371/journal.pone.0043871.g002

**Figure 3. Carboxysomal proteins in wild-type S. elongatus PCC 7942 and BMC mutants.** Western immunoblots show the presence or absence of carboxysomal proteins in carboxysomes purified to the Triton X-100-Percoll stage of the Epps-EDTA method for β-carboxysome purification [43,48]. Polyclonal antibodies against CcaA, RbcL5, RbcLS, CcmM-35, CcmK2, and CcmO were used [15]. The CcmM-35 antisera detects CcmM-58 and CcmM-35, as well as a non-specific band (ns) corresponding to RbcL that is routinely visible in CcmM western immunoblots [15]. The CcmK2 antibody detects all three CcmK homologues, but not CcmO, whereas the CcmO antibody detects CcmO specifically.

doi:10.1371/journal.pone.0043871.g003
CcmK2 was confirmed as an in vivo protein binding partner of CcmO (Figure 4) as suggested by the western immunoblot analysis above (Figure 3). It was not clear whether this was due to the presence of hetero-oligomers formed from CcmK2 and CcmO or interactions between adjacent protein oligomers in the β-carboxysome shell. The untagged form of CcmO was the predominant form co-purified from TP pellets of ΔccmO + pH6-Ub-ccm0 (Figure 4). Complicated expression of both tagged and untagged forms of H6-Ub chimeric constructs are well known [50,51] explaining the presence of both forms of recombinant CcmO in the TP pellet. Significantly, that untagged CcmO was co-purified by H6-Ub-CcmO confirms speculation that CcmO forms oligomeric proteins like other BMC proteins.

The small amount of CcmM-58 detected in the IMAC bound fraction (Figure 4) was probably not due to an interaction between CcmO and CcmM-58. A similar amount of CcmM58 is present under both urea treatments (Figure 4). This is hardly surprising given that CcmM-58 is an abundant cellular protein [52], and is certainly one of the most abundant proteins in TP pellets [16,43] suggesting non-specific carryover of CcmM-58 into the IMAC eluate. Nonetheless, the presence of CcmM-58 in the eluate would not be unexpected given its proven interaction with CcmK2 [17].

Discussion

CcmK2 and CcmO are the predominant proteins of the outer β-carboxysome shell in S. elongatus PCC 7942

The outer shell of β-carboxysomes is predicted to be formed from a number of subtly different CcmK homologues [9]. In Synechocystis PCC 6803, CcmK1 and CcmK2 have been postulated as major shell proteins whereas CcmK3 and CcmK4 are probably minor [29]. Of these, S. elongatus PCC 7942 has the ccmk2, k3, k4 and ccmO genes. We argue that CcmK2 and CcmO are the major proteins, and CcmK3 and CcmK4 are minor proteins forming the outer shell of β-carboxysomes from this strain.

In this report, we showed that some BMC mutants formed aberrant carboxysome ultrastructure. The polar body mutants H1ND, ΔccmK2 and ΔccmO were phenotypically high-CO2 requiring and thus incapable of growth in air (Table 1). Previously, some partially characterised ccmN and RuBisCO small subunit mutants have exhibited the polar body carboxysome phenotype [33,39,40], and these polar bodies were shown to contain most, if not all of the RuBisCO of the cell [39]. Interestingly, the ultrastructure of ΔccmO carboxysomes presented here (Figure 1-H) contradicts previous analyses of ccmO mutants which revealed either no carboxysomes [41], or a large proportion of aberrant carboxysomes [42]. We suggest that our specific genetic deletion provides a better experimental model than the insertional ccmO mutant investigated previously, hence the phenotypic difference.

The structural basis for polar bodies was clear. It was apparent that a structurally relevant interaction between CcmK2 and CcmO represents a key interaction for the integrity of the β-carboxysome outer shell (Figure 3, Figure 4). Previous work showed no interaction between CcmO and CcmK2 [17] but we showed that genetic deletion of either of the genes abolished not just that protein, but also the other from carboxysome-enriched TP pellets (Figure 3), and these were subsequently shown to interact in vivo (Figure 4). In contrast to the proteins identified from TP pellets (Figure 3), very small amounts of CcmK2 were present in the Mg²⁺ pellet from the ΔccmO mutant (Figure S3). This may indicate the presence of a partial, or incomplete shell structure on polar bodies in these mutants, or the spontaneous and extraneous association of outer shell sub-complexes to the polar body due to the well-established interaction between CcmK2 and CcmM [15-17]. Indeed, polar-body mutants had high CO₂-requiring (HCR) physiological phenotypes which were similar to that of the carboxysomelss ΔccmO mutant (Figure 2), indicating the lack of an effective shell structure.

Polar bodies are ordered structures containing active RuBisCO

In terms of the inner structure of polar bodies, western immunoblot against known β-carboxysome components showed that polar bodies contained all of the protein components expected of β-carboxysomes except the major outer shell proteins (Figure 5). Indeed, the maximum cross-sectional diameters of polar bodies (as well as carboxysomes) increased consistently as expected if these bodies were to contain regularly increasing RuBisCO layers bound together by CcmM-35 [14]. It is well known that cyanobacteria require high RuBisCO activity to survive, and because polar bodies were previously shown to contain all of the RuBisCO of the cell [39]. Thus in contrast to the supposition of Kinney et al. [33], we argue that polar bodies are physiologically relevant, ordered structures containing active RuBisCO. During β-carboxysome biogenesis there must therefore be a mechanism by which nascent carboxysome-like complexes are constrained into smaller carboxysomes rather than polar bodies. Kinney et al. [33] proposed that the CcmN protein is important for this process, and that this protein would bridge the inner and outer shell layers. Based on the shared ultrastructure of ccmk2, ccmO and ccmN mutants we agree that this is a reasonable proposal.

So far, no β-carboxysome mutant has been reported that results in polar bodies, perhaps stemming from their relative lack of internal structure and the less complete set of genetic knock-outs so far completed. However, within the propanediol utilisation micro-compartment, which has eight putative shell proteins, the AphAαBBB mutant results in polar bodies. These bodies are much smaller than those reported here, and they probably represent aggregated active protein components from the PDU compartment [53]. The shell proteins identified in PDU micro-compartment are varied, and two protein products of AphAαBBB make up approximately 25% of the PDU micro-compartment [54]. Thus polar body formation after disruption of the major shell proteins could be a common
feature of bacterial micro-compartments, and not necessarily due to general protein aggregation.

CcmK3 and CcmK4 are minor outer-shell components

In contrast to the HCR mutants containing polar bodies, the remaining mutants, ΔccmK3, ΔccmK4 and ΔccmK3-4, had wild-type like carboxysomes (Figure 1) but those from ΔccmK3-4 were physiologically impaired (Figure 2). The phenomenon of physiologically impaired carboxysomes with normal ultrastructure is not novel, as spherical carboxysomes lacking the vertex proteins CsoS4A and CsoS4B were shown to have increased CO2 leakage from ostensibly normal carboxysomes [20]. In terms of their protein composition, comparison of the CcmK western immunoblot detected the absence of detectable quantities of CcmK3 and CcmK4 from TP pellets (Figure 3). This suggests that multiple CcmK bands, observed here and previously [15], may be breakdown or alternative products of CcmK2 as they are only abolished by the ccmK2 and ccmO deletions (Figure 3). Therefore the protein evidence is consistent with CcmK3 and CcmK4 being low-abundance carboxysome proteins. The physiology and ultrastructure of β-carboxysomes from ΔccmK3-4 support this claim. This is not unexpected given their BMC domain and their Ci-responsive co-expression with other carboxysome components [55,56]. Indeed, Savage et al. [44] showed that the Ci-responsive co-expression with other carboxysome components [55,56]. Indeed, Savage et al. [44] showed that the Ci-responsive co-expression with other carboxysome components [55,56]. Indeed, Savage et al. [44] showed that the Ci-responsive co-expression with other carboxysome components [55,56]. Indeed, Savage et al. [44] showed that the Ci-responsive co-expression with other carboxysome components [55,56]. Indeed, Savage et al. [44] showed that the Ci-responsive co-expression with other carboxysome components [55,56]. Indeed, Savage et al. [44] showed that the Ci-responsive co-expression with other carboxysome components.

CcmK3 and CcmK4 are required for correct subcellular localisation of β-carboxysomes

Recently, the spatial arrangement of β-carboxysomes was shown to depend on the bacterial cytoskeleton [44]. The precise interactions underlying β-carboxysome localisation and partitioning are unclear, however the cytoskeleton proteins ParA and MreB are essential in S. elongatus PCC 7942 [44] and we present evidence in this report that CcmK3 and CcmK4 are individually required for correct subcellular localisation of β-carboxysomes (Figure 1-G). The phenotype of seemingly normal, aggregated, micro-compartment was also observed in the paxA mutant of Salmonella enterica, whose PDU micro-compartments are involved in degradation of 1,2-propanediol [53]. This suggests a common mechanism for arrangement and partitioning of different types of bacterial micro-compartment across vast phylogenetic distances.

Aberrant carboxysome partitioning is insufficient to explain the physiological phenotype of the ΔccmK3-4 strain. Savage et al. [44] showed that the paxA mutant was unable to correctly apportion β-carboxysomes within the cell, and to daughter cells at mitosis. However this strain had only a very slight growth rate disadvantage whereas the ΔccmK3-4 mutant had a severely reduced growth-rate in air. We perceive similar implications for carboxysome partitioning during mitosis in the polar body mutants HIND, ΔccmK2 and ΔccmO. Supporting this, numerous cells without obvious PBs were observed in TEM sections, indicating that the PBs, and thus the cellular RuBisCO, were not partitioned to daughter cells in the same even manner as in β-carboxysomes in wild-type cells. Thus we reason that for the mutants with polar bodies, half of the daughter cells at mitosis will inherit all of the RuBisCO of the parental cell. Hence some part of the HCR phenotype of HIND, ΔccmK2 and ΔccmO is probably caused by the lack of recruitment of CcmK3 and CcmK4 to the carboxysome, and subsequent deficiencies in localisation and inheritance of carboxysomes during mitosis. This is perhaps an extreme example of the type of β-carboxysome partitioning deficit described by Savage et al. [44].

Variable numbers of ccmK genes

We have shown that in S. elongatus PCC 7942 there are two major β-carboxysome shell components, CcmK2 and CcmO, and two minor, CcmK3 and CcmK4. Thus the minimum BMC-gene requirement for a structurally relevant carboxysome is ccmK2 and ccmO, whereas accessory BMC genes such as ccmK3-4 refine the functionality of the shell. Intriguingly, other β-cyanobacteria have as many as nine identifiable BMC genes and most of the variability in β-cyanobacterial BMC genes is due to accessory minor shell proteins (Table S1). The potential for CcmK3 and CcmK4 proteins to act on carboxysome localisation and partitioning suggests that the variable BMC gene complement reflects the varied morphology of cyanobacterial cells. Perhaps the wide variation in cyanobacterial morphology and subcellular structure is matched by a varied cytoskeletal structure and dynamic, thus the carboxysome must be able to flexibly alter its shell structure in order to maintain cognate interaction with a similarly varied or flexible cytoskeleton. However there is no obvious correlation between number of accessory ccmK genes and lifestyle or environment niche of β-cyanobacterial strains.

In Synechocystis PCC 6803, ccmK4 was previously shown to be essential for phototrophic growth [45]. Thus there are questions over the roles that accessory CcmK proteins play in different β-carboxysome systems. One wonders what aspect of Synechocystis carboxysomes makes CcmK4 an essential protein. It is possible that the multiple transposon mutants of ccmK4 identified by Zhang et al. [45] were also polar mutants of ccmK3. This seems implausible, leading to the conclusion that different ccmK homologues have different functions, or are under different selective pressures in different β-cyanobacteria.

Does the CcmO protein recruit the vertex pentamer CcmL?

CcmL is predicted to occupy the vertices of β-carboxysomes [19] and is required for effective shell function. Its absence leads to rod-like carboxysomes in β-cyanobacteria [20,34,36]. Rod-like and potentially tetrahedral carboxysome-like inclusions were sometimes observed in the δccmO + pH6-Ub-ccmO complemented strain (Figure S1), hinting at the possibility of a structural interaction between CcmO and CcmL. As evidenced by protein analysis, the predominant CcmO form in carboxysome-enriched TP pellets was the untagged form (Figure 4) which probably arises from some proteolytic activity in vivo. Hence two possibilities may underlie the occasional occurrence of structural aberration: Structural perturbation caused by the 11 kDa hexahistidine-
ubiquitin tag itself, or an improper stoichiometry between shell proteins due to the non-native promotor from which the tagged ccmO gene was expressed.

The rod-like carboxysome phenotype is usually attributed to a syndrome of CcmL insufficiency where the vertices of the carboxysome cannot close. Thus, if CcmL is recruited to the β-carboxysome by CcmO, and the quantity of CcmO is sub-stoichiometric due to expression from the non-native promotor, the result may be a low rate of mutant carboxysomes that resemble the authentic ccmL mutant. Alternatively, the CcmO protein may tolerate some forms of modification; indeed it has variable N- and C-terminal domains which make it the least conserved BMC protein in β-cyanobacteria. Based on these data, we would argue that CcmO interacts with CcmL at the vertices of the carboxysomal icosahedron. Indeed, it is possible that CcmO forms a possible bridging protein, or “zipper”, that helps fix the interface between neighbouring outer-layer triangular facets.

CcmP: potential RuBP pore for the β-carboxysome outer shell?

The α-carboxysomal BMC gene csoS1D was recently shown to have a β-cyanobacterial homologue, conP [56]. This gene was not obviously related to the β-carboxysome at the outset of this study, thus it is absent from the analysis presented here. Nonetheless it is an obvious focus for future work, the α-carboxysomal homologue CsoS1D was shown to be exceedingly rare in the isolated α-carboxysomes of Prochlorococcus sp. str. MED1, existing at less than one functional unit per α-carboxysome facet [31]. This protein was also shown to be important, but not essential for correct α-carboxysome ultrastructure in an ectopic α-carboxysome expression study [57]. Early assertions that the CsoS1D protein may form a gated pore for RuBP [26] seem not to be borne out by experimental studies where its role appeared to be primarily structural [57]. Thus it remains to be seen whether the structure of β-carboxysomes is dependent on the protein product of conP.

A tentative model for the interaction of outer shell proteins in S. elongatus PCC 7942

Given the carboxysome ultrastructure and physiology of ∆ccmK3-4, it appears that as few as three proteins, CcmK2, CcmO and CcmL, form the minimum set required to construct an outer shell that is capable of supporting photo-autotrophic growth in air levels of CO2—bearing in mind that an outer shell cannot form without the 35 and 58 kDa isoforms of the RuBisCO-organising CcmM protein [15,16,17]. Based on observed structures in some ∆ccmO + pH6-Ub-ccmO cells the vertex protein CcmL could interact with CcmO, perhaps suggesting a structural role for the CcmO protein at the vertices of the carboxysome. The CcmK2 protein is known to interact
with the inner shell proteins CcmM-58 and CcmN [15,17], and is likely to form the bulk of the facets. CcmK3, CcmK4, and potentially CcmP, probably have niche roles, and CcmK3 and CcmK4 are individually required to close the carboxysome shell, potentially at very specific locations.

The data presented in this study suggest potential models describing the outer-shell structure of facets in idealised β-carboxysomes from *S. elongatus* PCC 7942 (Figure 5). The first facet model for the outer-shell of idealised β-carboxysomes (Figure 5-A) has CcmK2 as the predominant protein of the β-carboxysome facet, covering ~70% of the facet surface (Table 2). CcmO is shown at the vertex-facet interface as well as the facet-facet interface and covers ~30% of the surface. The second model (Figure 5-B) has CcmO trimers at defined locations along the edges of the β-carboxysome, as a ‘zipper’ protein that forms ~10% of the β-carboxysome surface (Table 2).

The location of CcmO in both models (Figure 5-A,B) is supported by the observation that CcmO was not detected in the Mg²⁺ pellet of ΔccmK2 and HIND mutants, but that CcmK2 was detected at low levels in the Mg²⁺ pellet of ΔccmO (Figure S3): CcmK2 can associate to the polar body through its dual interactions with CcmM and CcmN, but CcmO does not (Figure 4), probably because the opportunity for interaction with internal proteins is minimised at the facet-facet interface. Similarly, the stoichiometry of CcmK2 hexamers to CcmO trimers in model A (Table 2) is consistent with the observed amount of CcmK2 in β-carboxysomes, which by current estimates covers only 63–75% of the carboxysome surface in *S. elongatus* PCC 7942 [14]. Thus we postulate that the amount of CcmO protein in model A is consistent with reported shortfall in β-carboxysome surface coverage.

Also, CcmO requires interaction with the CcmK2 proteins of the carboxysome facet for incorporation into the carboxysome. Thus we postulate that CcmO could form the entire edge structure and interfaces adjacent facets (Figure 5-A). This speculative model accounts for the minimum set of components encoding a functional outer shell structure CcmK2, CcmO and CcmL, and accounts for structural requirements at the facet-facet interface by assuming that CcmO is able to fill two structural roles: flat hexagonal trimers at the interface with CcmK2, and bent trimers at the interface between the facets. This assumption is supported by the behaviour of EutS which is able to adopt flat and bent conformations in the ethanalamine detoxification micro-compartment [23,32].

Our alternative model where CcmO only fulfils the EutS-like bent-oligomer role (Figure 5-B, Table 2) is more economical with regard to CcmO content per carboxysome, and may be unsatisfactory with respect to observed amounts of CcmK2 proteins reported by Long et al. [14]. Nonetheless, the model B demands only a single oligomeric conformation for the CcmO protein – the ‘bent’ EutS-like conformation. To date, the exact nature of CcmO trimers has not been revealed by protein crystallography, thus the exact role played by CcmO in the outer β-carboxysome shell remains speculative.

### Materials and Methods

**Bacterial strains and culture conditions**

*S. elongatus* PCC 7942 and mutant derivatives were maintained on modified BG-11 medium [58–60] solidified with 1.2% agar. The same medium was used for growth analyses and liquid culturss but for physiological analysis by membrane-inlet mass spectrometry NaNO₃ was replaced with 20 mm NaCl and the medium was buffered with 50 mm bis-tris propane (pH 7.9).

Gene inactivation plasmids were generated in pUC18 which does not replicate in *S. elongatus* PCC 7942. Primers used for PCR are listed in Table S2. Gene inactivation plasmids were constructed in the pUC18 plasmid backbone as described previously [37] using a selectable chloramphenicol resistance marker (Cm₄⁰) [61]. DNA sequences of all constructs were confirmed by DNA sequencing and the final plasmids were transformed into *S. elongatus* PCC 7942 as described previously [34,37]. Segregated transformants were confirmed by diagnostic PCR and restriction digestion, these strains are listed in Table S3. The DNA oligonucleotides used for PCR are listed in Table S4.

**Complementation of mutant strains**

Phenotypically mutant strains were subsequently complemented by reintroduction of their respective genes from an E. coli/S. elongatus shuttle vector, ccmK2 was reintroduced into HIND and ΔccmK2 under control of its native promotor in the vector pSE41 (derived from pSE4 [62,63] in this work), ccmO was reintroduced into ΔccmO under control of the lac operator in the pSE2 vector [64], and ccmK3-4 were reintroduced into ΔccmK3-4 under control of their native promotor in the pSE41 vector. The pSE41 E. coli/S. elongatus expression vector was constructed by the in-frame insertion of the promotorless ampicillin resistance gene from pUC18 as well as additional synthetic restriction sites (NheI, BmiI, BglIII, Scal and XhoI) into the NcoI/XbaI restriction sites within the polylinker sequence of pSE4 [63]. The pSE41 plasmid is ampicillin and spectinomycin resistant in E. coli. The pSE41 plasmid map is shown in Figure S4, and the complemented strains are listed in Table S3. The DNA oligonucleotides used for PCR are listed in Table S4.

Previous studies have shown that effective IMAC purification of β-carboxysomal proteins was enhanced by insertion of a spacer domain between the protein and the hexahistidine tag (chloramphenicol acetyltransferase, CAT in [16]). In the current study we used the pHUE hexahistidine-ubiquitin tagging system to produce a single plasmid construct for both complementation of ΔccmO and IMAC-purification of CcmO-interacting proteins [50,51]. The ubiquitin tag has been shown to be a non-interacting tag that has been effective in the improved solubility of otherwise problematic proteins [50,51].

The mutant strains that had detectable ultrastructural or physiological phenotypes were complemented to wild type carboxysome structure, arrangement and function, showing that the mutant phenotypes were entirely attributable to the specific gene deletions (Figure S1, Figure S2, Table S2).

**Physiological analyses**

Physiological measurements by membrane-inlet mass spectrometry (MIMS) were performed as described previously [65,66], and maximum growth rates were measured as described by Long et al. [16].

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**Table 2.** Protein components of the outer shell of idealised models of the β-carboxysomes presented in Figure 5.

|                | CcmK2 | CcmO | CcmL |
|----------------|-------|------|------|
| **Model A**    |       |      |      |
| Multimers per carboxysome | 1700  | 720  | 12   |
| Surface area (%)  | 69.9  | 29.6 | 0.5  |
| **Model B**    |       |      |      |
| Multimers per carboxysome | 2180  | 240  | 12   |
| Surface area (%)  | 89.6  | 9.9  | 0.5  |

doi:10.1371/journal.pone.0043871.t002
Transmission electron microscopy

Cells were prepared for transmission electron microscopy (TEM) essentially as described previously [34] the exception being that the cells were embedded in LR-White resin or epon-arlaidite rather than Spurrs resin. Sections were stained with 2% uranyl acetate and Reynolds lead citrate. The sections were viewed in a Hitachi H7000 transmission electron microscope (Hitachi Ltd, Tokyo, Japan) at 75 kV. Measurement of carboxysome diameter was performed using ImageJ 1.45 [67]. Carboxysome measurements were made at the widest cross-sectional width in longitudinal median sections.

Protein purification and western immunoblots

Carboxysomes were enriched to the Triton-Percoll pellet stage of the Epps-EDTA method for β-carboxysome enrichment as described previously [43,48]. Crude β-carboxysome preparations were also made using the Mg2+ precipitation method [49]. IMAC purification of hexahistidine-tagged protein complexes was carried out as described previously [16] and protein samples were concentrated by precipitation with one volume of 100% trichloroacetic acid on ice for 30 minutes. After incubation the sample was collected by centrifugation at 4°C and washed twice with ice cold 80% acetone. The pellet was finally resuspended in 1x NuPAGE sample buffer and 50 mm dithiothreitol.

Western immunoblots were performed as described previously, using the same CcaA, CcmM-35, CcmK2 (called CcmK1 in our previous work), and RuBisCO antibodies [15]. We have also performed western immunoblots using a polyclonal rabbit CcmO antibody raised against recombinant S. elongatus PCC 7942 CcmO. We have previously shown that the RuBisCO antibody reacts to the large and small RuBisCO subunits [16], and that the CcmM-35 antibody reacts with the 35 and 58 kDa isoforms of CcmM [15]. The CcmK2 antibody detects CcmK3 and CcmK1 proteins produced in E. coli but not CcmO produced using the same expression system. In contrast, our CcmO antibody specifically produced in E. coli but not CcmO produced using the same expression system.

Supporting Information

Figure S1 Carboxysome ultrastructures in complemented mutant strains. A, the HIND ccmK2 insertional mutant complemented with pSE41-ccmK2. B, ∆ccmK3-4 complemented with pSE41-ccmK3-4. C, ∆ccmO complemented with pH6-Ub-ccmO. Scale bars are 500 nm. (TIFF)

Figure S2 Photosynthetic O2 evolution in response to external Ci by complemented BMC mutants. Shown are a representative set of mass-spectrometric measurements of Ci-dependent O2 evolution by wild type and mutant strains of S. elongatus PCC 7942 over a range of Ci concentrations. Ci is the sum of CO2 and HCO3− in solution (pH 7.9). (TIFF)

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