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Effect of bio-engineering on size, shape, composition and rigidity of bacterial microcompartments

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Bacterial microcompartments (BMCs) are proteinaceous organelles that are found in a broad range of bacteria and are composed of an outer shell that encases an enzyme cargo representing a specific metabolic process. The outer shell is made from a number of different proteins that form hexameric and pentameric tiles, which interact to allow the formation of a polyhedral edifice. We have previously shown that the *Citrobacter freundii* BMC associated with 1,2-propanediol utilization can be transferred into *Escherichia coli* to generate a recombinant BMC and that empty BMCs can be formed from just the shell proteins alone. Herein, a detailed structural and proteomic characterization of the wild type BMC is compared to the recombinant BMC and a number of empty BMC variants by 2D-gel electrophoresis, mass spectrometry, transmission electron microscopy (TEM) and atomic force microscopy (AFM). Specifically, it is shown that the wild type BMC and the recombinant BMC are similar in terms of composition, size, shape and mechanical properties, whereas the empty BMC variants are shown to be smaller, hollow and less malleable.

Some bacteria have an inherent ability to compartmentalize specific metabolic processes by encasing them within a proteinaceous organelle called a bacterial microcompartment (BMC)^{1–3}. In so doing they generate cytoplasmic supramolecular complexes that look similar to phage, with a diameter of between 100–200 nm, when thin sections of BMC-containing bacteria are viewed by transmission electron microscopy^{4–6}. BMCs consist of an outer semi-permeable protein shell that envelopes and secludes enzymes and metabolites from the main cellular milieu^{7–11}. They are associated with either anabolic processes such as carbon fixation (carboxysomes)⁵ or with catabolic processes connected with the utilization of either carbon or nitrogen (metabolosomes)¹². Carboxysomes house ribulose-1,5-bisphosphate carboxylase/oxygenase (RuBisCO) and carbonic anhydrase and act to promote a higher local concentration of carbon dioxide in order to increase the efficiency of the carboxylation reaction¹³. Metabolosomes are associated with fermentative processes with substrates such as ethanolamine, choline, fucose, rhamnose or 1,2-propanediol¹. A recent bioinformatics analysis has indicated that there are at least 23 distinct BMCs found across a range of bacteria phyla¹⁴.

Of the metabolosomes, the best characterized is the 1,2-propanediol (1,2-PD) utilizing organelle (Fig. 1) from organisms such as *Salmonella enterica*^{4,15,16} and *Citrobacter freundii*¹⁷. The genetic information for the organelle is found within the propanediol utilization (*pdu*) operon, which contains 23 genes that encode for not only the shell proteins of the BMC but also the enzymes associated with the transformation of 1,2-PD into propionaldehyde and its subsequent disproportionation into propionic acid and propanol⁴. This includes the adenosylcobalamin-dependent diol dehydratase^{4,16} as well as enzymes linked to the reactivation of the coenzyme^{18–20}. In this respect the metabolosomes house a more complex repertoire of enzymes than the carboxysomes and have a broader requirement for coenzymes and cofactors. The protein shell of the metabolosomes is proposed to protect the cytoplasm from toxic aldehyde intermediates²¹.

Structural data from X-ray studies of individual shell proteins from both carboxysomes and metabolosomes have revealed that they form hexameric-shaped tiles with a central pore that piece together to form the faces of the organelle whereas pentameric tiles act as the vertices of the structure^{9,10,22–30}. The Pdu BMC contains 7 shell proteins, PduA, PduB, PduB' PduJ, PduK, PduT and PduU, which generate the hexagonal tiles that represent the

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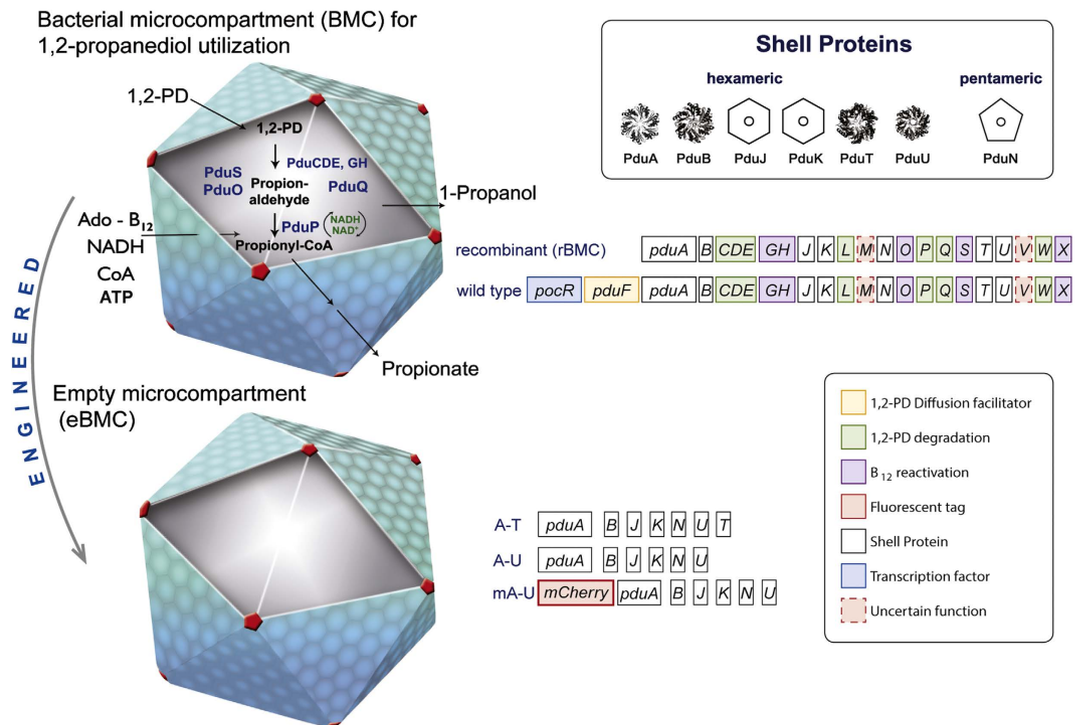


Figure 1. Schematic illustration of the Pdu shell and shell proteins associated with the wtBMC, rBMC and eBMC variants (A-U, mA-U and A-T). Shell Proteins: The protein structures for PduA, PduB, PduT and PduU are shown, revealing the central pores within the hexameric or pseudohexameric symmetry. Hexameric shell proteins form the facets of the structure. The pentameric vertex protein (PduN) is thought to take up the vertex position of the structure. Bacterial microcompartment (BMC) for 1,2-propanediol utilization: wtBMCs and rBMCs incorporate many of the enzymes necessary for 1,2-PD degradation as well as enzymes for adenosylcobalamin reactivation. The rBMC plasmid lacks the genes encoding for the diffusion facilitator protein (yellow, *pduF*) and the transcription factor (cyano, *pocR*). Substrate (1,2-PD) and cofactors (ATP, CoA, NAD⁺ and B₁₂) are thought to diffuse through the gated pores in the shell protein. Empty microcompartment (eBMC): Three different shell protein constructs consisting of shell protein genes (A-T, A-U and mA-U) can be produced in *E. coli* (Supplementary). A-T is the basic eBMC that houses all the shell proteins (PduA, B, B', J, K, N, U, T). A-U is composed of six Pdu shell protein, PduA, B, B', J, K, N, U. The mA-U construct has a C-terminal fusion of mCherry (red, fluorescent tag) to PduA and thus consists of mCherry-PduA, B, B', J, K, N, U.

facets of the polyhedral structure (Fig. 1). Most of the shell proteins contain a single BMC domain that consists of about 90 amino acids, which fold into a β - α - β motif allowing them to aggregate as hexamers⁹. However, PduB, its N-terminally truncated derivative from an alternative translation start site (PduB') and PduT contain tandem BMC domains that generate a pseudohexamer formed by a trimeric structure^{23,27,28}. The vertices of the BMC are thought to be formed from the pentameric PduN²⁹⁻³¹.

All shell proteins contain central pore(s) (Fig. 1), some of which appear to have alternative conformations^{22,23,26,27}. Passage of substrates, products and cofactors across the shell is thought to be mediated by diffusion via the pores within the hexameric components of the shell face, movement that may be facilitated by a differential charge distribution of amino acid residues surrounding the pore. It is envisaged that engineering of these pores may change the ability of the BMC to allow passage of substrates, products or cofactors into or out of the macromolecular structure^{32,33} as has recently been evidenced through the manipulation of PduA pores³⁴. Overall, the structure of the BMC has to be flexible enough to allow shell assembly and closure but also stable enough to maintain structural integrity. However, to date there is little information on the structural rigidity of these complexes.

BMCs represent genetically tractable systems that can be successfully cloned into organisms in which they are not normally found. In this respect, the Pdu system from *C. freundii* has been engineered into *E. coli* to allow for the formation of functional recombinant organelles^{17,35,36}. Similarly, reports demonstrate that the ethanolamine utilization (Eut) system can be manipulated to allow for recombinant BMC production and protein targeting^{37,38}. Recombinant carboxysomes, too, have been produced in *E. coli*, generating structures that are capable of carbon dioxide fixation³⁹. Empty *C. freundii* Pdu microcompartments (eBMCs) have been formed from the coordinated recombinant production of PduA, B, B', J, K, N and U³⁶. However, the composition, structure and physical properties of purified wild-type BMCs (wtBMCs) have not been characterized in detail and compared to the full recombinant Pdu BMC (rBMC) or its empty derivatives. Herein, we compare the shell protein constituents of wtBMCs, rBMCs and eBMCs, and investigate their composition, form and nano-mechanical properties.

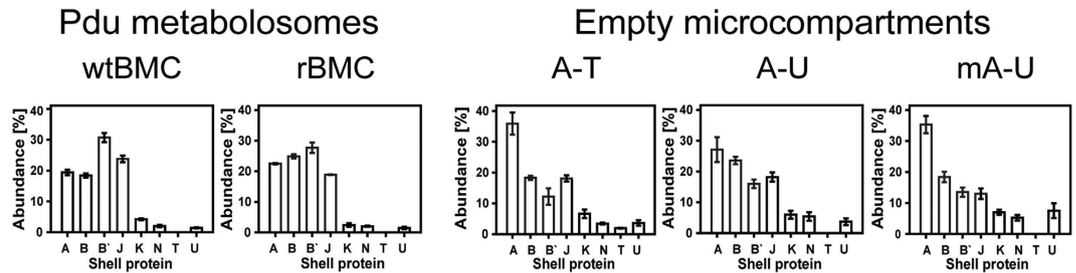


Figure 2. BMC shell protein composition. The shell protein composition is shown as a percentage of the total protein of wtBMC, rBMC and the eBMCs A-T, A-U, mA-U. The data represents the mean values of three independent repeats. The error-bars indicate one standard error of mean.

Results

***In vivo* characterization of recombinant and wild type Pdu BMCs.** In order to compare wild type BMCs with recombinant and empty BMCs, *in vivo*, the sizes of the various bacterial organelles were first compared *in situ* after thin sectioning and visualization of the host bacteria by transmission electron microscopy (TEM) (Supplementary Figure 1). In this respect wild type Pdu BMCs (wtBMCs) from *C. freundii* were compared initially to recombinant *C. freundii* Pdu BMCs (rBMCs) produced in *E. coli*. The rBMC construct includes all shell proteins, metabolic enzymes for 1,2-PD degradation and reactivation factors for adenosylcobalamin. Subsequently, the wtBMC and rBMC were compared to a range of engineered empty recombinant BMCs (eBMCs) that were produced as a result of (i) the expression of the genes encoding all the shell proteins PduA-B-J-K-N-U-T (A-T), (ii) a reduced set of shell proteins PduA-B-J-K-N-U (A-U), and (iii) an empty BMC containing a fluorescent protein fused to PduA, mCherry-PduA-B-J-K-N-U (mA-U).

Growth of *C. freundii* in the presence of 1,2-propanediol resulted in the production of Pdu wtBMCs, which were easily visualized after embedding and thin sectioning of the bacteria. As expected, the wtBMCs appeared as heterogeneous polyhedral bodies that were evenly dispersed around the cell, as had been previously observed with the Pdu BMCs that are found in *S. enterica*⁴ and *L. reuterii*⁴⁰. In contrast, rBMCs were observed densely packed in thin-sectioned *E. coli* cells harboring a plasmid containing the *pdu* operon. In this case the rBMCs take up the majority of available space within the cytoplasm of the cell. However, despite the difference in the quantity of organelles within the cell, both the wtBMCs and rBMCs were found to be of a similar size distribution.

When the three eBMCs variants (A-T, A-U and mA-U) were compared to one another, no phenotypical differences were observed in the TEM images of the thin-sectioned *E. coli* strains containing the eBMC constructs. All contained heterogeneous polyhedral bodies that were crammed into the cytoplasm of the cell. However, overall, all three eBMCs variants appeared smaller in diameter in comparison to the complete BMCs (wtBMCs and rBMCs, for images see Supplementary Figure 1).

Purification of intact wtBMCs, rBMCs and eBMCs. In order to characterize and compare the structural and biochemical properties of wtBMCs, rBMCs and eBMCs, a number of purification methods to isolate intact BMCs were investigated involving the use of mild detergent lysis procedures, employing either a bacterial specific lysis reagent (B-PERTM) or a yeast specific lysis reagent (Y-PERTM). The bacterial protein extraction (B-PERTM) method is based on lysing cells with B-PERTM and entails consecutive centrifugation steps to separate cell debris from Pdu metabolosomes⁴¹. This procedure was found to work best for the isolation of complete BMCs (wtBMCs and rBMCs). The yeast protein extraction reagent (Y-PERTM) method is based on lysis of bacterial cells and subsequent salt precipitation and was found to be effective for the purification of eBMCs³⁵. The purification and isolation of the various BMCs was followed by SDS-PAGE (Supplementary Figure 2). As the eBMCs variants were found to purify under different conditions to the rBMC and wtBMCs it was concluded that they must have different physical and chemical properties.

Determination of the shell protein composition of wtBMCs, rBMCs and eBMCs. The purification of wtBMCs, rBMCs and eBMCs allowed their shell protein composition to be compared through a proteomic approach. Such holistic analysis gives information on the composition of the building blocks of these supramolecular structures, which is essential in order to understand their assembly and mechanical properties, as well as their ability to acquire substrates and cofactors. Differences in the shell protein composition of purified wtBMCs, rBMCs and eBMCs were examined by 2D-PAGE (Supplementary Figure 3) and MALDI MS-MS tandem mass spectrometry in order to identify the individual building blocks and to quantify their relative abundance.

When the purified wtBMCs and rBMCs were analyzed by 2D-PAGE all the shell proteins could be detected by MALDI TOF-TOF. The abundance of the various shell proteins, in terms of per cent of total protein, was estimated by densitometry (Fig. 2). In wtBMCs, PduB' (31%) was the most abundant shell protein followed by PduJ (24%), PduA (19%) and PduB (18%). In contrast, PduK, PduN, PduT and PduU were found to be minor components of the complex. PduT was not quantified due to its low spot intensity. The minor shell components were estimated to account for less than 8% of the capsid in total. In rBMCs the order of shell protein abundance is slightly different, with the order of PduB' (28%) -PduB (24%) -PduA (22%) -PduJ (18%). The remaining shell proteins representing minor components of the structure. Thus, overall, the profile of the rBMC to the wtBMC

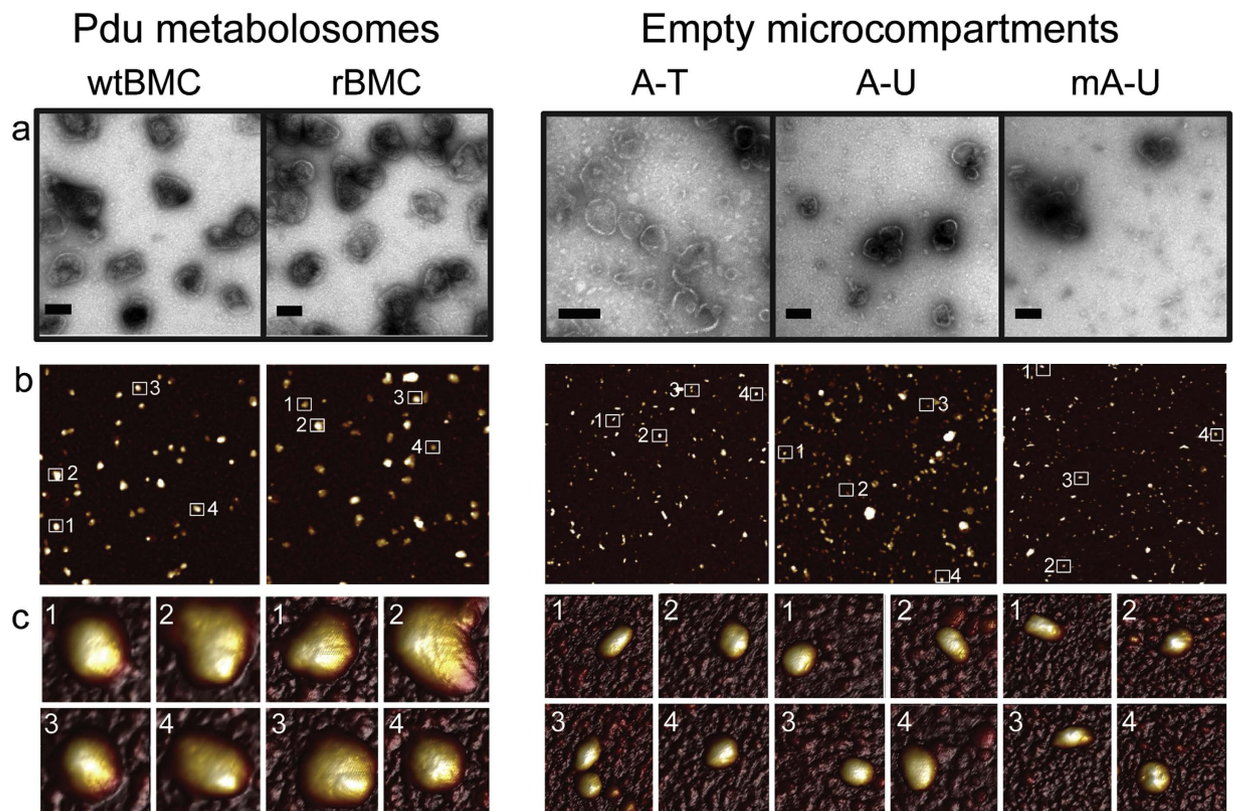


Figure 3. Comparison of TEM and AFM images of isolated BMCs. (a) TEM images of purified wtBMCs, rBMCs and eBMCs (A-U, mA-U and A-T). The scale bars represent 100 nm. (b) Overview of zoomed-in $4 \times 4 \mu\text{m}$ height AFM scans. White squares represent microcompartment particles shown in Fig. 3c. (c) 3D topology images of single zoomed-in particles ($240 \times 240 \text{ nm}$ each) from (b).

is remarkably similar indicating that they have a very analogous composition, as expected. This analysis is also similar to that reported to the purified *S. enterica* Pdu BMC¹⁶.

As with the complete BMCs, all the encoded shell proteins of the purified eBMC variants could also be detected by MALDI TOF-TOF analysis of spots excised from 2D-PAGE. Surprisingly, in all three eBMC variants (A-T, A-U, mA-U) that were analyzed, as a percentage of the total protein, PduA was found to be the major Pdu shell protein component (27–35%), followed by PduB (18–24%), PduB' (12–16%) and PduJ (13–18%) (Fig. 2). PduK, PduN, PduT (only in A-T) and PduU were all found to be minor shell components comprising less than 20% of the shell. Although some differences in the shell protein composition were noted for the eBMCs variants, the overall trend is similar (Fig. 2).

The major conclusion from this comparative proteomic analysis is that eBMCs appear to incorporate considerably more PduA relative to all other protein components, in comparison to their wtBMC and rBMC counterparts. Whether this reflects a different expression pattern or results from a lack of internal cargo has yet to be determined. However, it is likely that differences in shell protein composition will affect the nano-mechanical and functional properties of these bioengineered structures.

Structural characterization by single particle analysis. The structure/function relationship of BMCs is currently poorly understood but yet this knowledge is vital in order to understand and develop principles for engineering encapsulated processes³³. To learn more about how variation in BMC composition alters the physical properties of the structures a combination of AFM and TEM (Fig. 3) were employed to image and characterize purified intact wtBMCs, rBMCs and eBMCs (A-T, A-U and mA-U).

When viewed by negative stain TEM, the isolated Pdu microcompartment samples (wtBMC, rBMC, A-T, A-U and mA-U) all appeared to be pure (i.e. not contaminated with other bodies) (Fig. 3). The TEM images therefore confirm the efficiency of the purification method used and are consistent with the proteomic analysis described above, which did not detect any major contaminating proteins. The isolated wtBMCs were observed to adopt a wide variety of different polyhedral shapes and sizes (Fig. 3) and appeared to be irregular in overall topology. No major differences in size and shape were visualized between the wtBMC and rBMC structures. This suggests that the heterologous formation of Pdu BMCs in *E. coli* does not affect BMC size. No detectable structural differences were visualized between the different eBMC variants. This suggests that the presence of the minor BMC shell building block PduT does not affect the BMC structure and that the fusion of mCherry to the N-terminus of the major BMC shell protein building block PduA does not disrupt BMC assembly, size or shape. However, as

		Diameter/nm	Area/ μm^2	Height/nm
wtBMC	TEM	127 \pm 13.2	0.0128 \pm 0.0026	
	AFM	131 \pm 15.4	0.0137 \pm 0.0032	49.99 \pm 8.10
rBMC	TEM	122 \pm 26.4	0.0122 \pm 0.0053	
	AFM	133 \pm 23.6	0.0143 \pm 0.0053	50.15 \pm 8.36
A-T	TEM	77 \pm 18.5	0.0049 \pm 0.0024	
	AFM	59 \pm 7.9	0.0028 \pm 0.0007	18.74 \pm 3.31
A-U	TEM	65 \pm 7.4	0.0034 \pm 0.0008	
	AFM	66 \pm 8.4	0.0034 \pm 0.0009	18.42 \pm 2.41
mA-U	TEM	74 \pm 23	0.0047 \pm 0.0033	
	AFM	56 \pm 8.0	0.0025 \pm 0.0007	18.46 \pm 3.16

Table 1. Image analyses of microcompartment size distributions. Values denote mean \pm SD.

observed with the *in vivo* thin-section data, a clear size difference was observed between complete BMCs (wtBMCs and rBMCs) and the eBMC variants.

To complement the TEM data of the purified BMCs, AFM was also employed to provide both information on the 3D shapes of the structures and their nano-mechanical properties. The various microcompartments (wtBMC, rBMC, A-T, A-U, mA-U) were scanned in air using AFM. The general observations made by AFM are consistent with those made by TEM in that all of the isolated microcompartments appeared to be pure (i.e. not contaminated with other bodies). Both wtBMCs and rBMCs appeared heterogeneous in shape and size, but no size distribution differences were observed between the wtBMC and rBMC structures. The eBMC variants (A-T, A-U, mA-U) also appeared irregular in shape and heterogeneous in size. Despite the variation in size and shape of the eBMCs, the AFM analysis shows that they are indistinguishable in terms of their physical dimensions. As with the TEM imaging, both wtBMCs and rBMCs appear significantly larger than the eBMC variants when viewed by AFM (Fig. 3).

From the TEM and AFM data sets that were collected, the diameter, apparent cross-sectional area and peak height of the wtBMCs and rBMCs were quantified and compared (Table 1). The diameter of the *C. freundii* wtBMCs was 127 \pm 13.2 nm and 131 \pm 15.4 nm as measured by TEM and AFM, respectively (Fig. 4, Table 1, Supplementary Table 1). The isolated rBMC structures showed a slightly wider variation (TEM: 122 \pm 26.4 nm, AFM: 133 \pm 23.6 nm). However, the average diameter size of the wtBMCs and rBMCs were alike (122–133 nm). Indeed, these average values of the *C. freundii* wtBMCs and rBMC are consistent with the reported average diameter of the Pdu BMCs that had been purified from *S. enterica* (123 \pm 30 nm¹⁰).

To take the heterogeneous and irregular shape of Pdu BMCs into account, the apparent cross-sectional area occupied by wtBMC and rBMC particles were also quantified and compared (Table 1, Supplementary Table 2). The area occupied by individual wtBMCs was 0.0128 \pm 0.0026 μm^2 and 0.0137 \pm 0.0032 μm^2 as measured by TEM and AFM, respectively, whereas the single particle area of rBMCs was observed to be 0.0122 \pm 0.0053 μm^2 by TEM and 0.0143 \pm 0.0053 μm^2 by AFM. Therefore, as with the average diameter, rBMC particles displayed a higher variation in area size, although the average area size of wild-type and rBMCs showed no difference.

In addition to diameter and area quantification, the maximum height of particles can be measured by AFM. However, from the AFM scans of wtBMC and rBMC particles the maximum height of individual microcompartments was found to be smaller than the radius. This suggests that particle deformation may have occurred during sample deposition. Nonetheless, the maximum height information still provided new information concerning the size distribution of Pdu microcompartments. The peak height of wtBMCs and rBMC particles were both 50 \pm 8 nm. As with both diameter and area, no significant differences were observed between the wtBMC and rBMC structures (Fig. 4, Table 1, Supplementary Table 3).

The application of AFM to the eBMC variants again highlighted a wide variety of sizes in terms of diameter (Table 1, Supplementary Table 1). Importantly, no substantial differences were observed for the average diameters of A-T, A-U and mA-U, ranging between 56 \pm 8 nm and 77 \pm 19 nm. The cross-sectional areas of all three tested eBMCs were also compared (Fig. 4, Table 1, Supplementary Table 2). Interestingly, no significant differences in area size between A-T, A-U and mA-U were identified. The vast majority of all tested eBMC variants had average values between 0.0025 \pm 0.0007 μm^2 and 0.0049 \pm 0.0024 μm^2 . Finally, and as seen with the wtBMC and rBMC particles, the peak heights associated with the eBMC variants appeared distorted, as their peak heights were smaller than their diameter (Fig. 4, Table 1, Supplementary Table 3). The average peak height of the eBMC particles ranged from 18.4 \pm 2.4 nm to 18.7 \pm 3.3 nm. The average values of all three eBMC were similar with peak heights of 18–19 nm.

Nano-mechanical mapping. To further investigate the difference between complete BMCs (wtBMCs and rBMCs) and the eBMC variants (A-T, A-U and mA-U), the nano-mechanical properties of the microcompartments were quantified using AFM quantitative nano-mechanical mapping (QNM) (Bruker). Application of a force titration between 200 pN and 2 nN showed no deformation for any of the purified microcompartments and no change in the height of the compartments was observed, confirming that imaging of the microcompartments did not cause an observable distortion. However, in contrast to the low force experiments, QNM experiments at higher than 2 nN applied force showed significant nano-mechanical differences between complete BMCs and eBMCs (Fig. 5). At a force set point of 250 nN, both wtBMCs and rBMC showed an average maximum deformation of 2.9–3.0 nm, whereas eBMCs showed an average maximum deformation of 1.7–1.9 nm. The same QNM

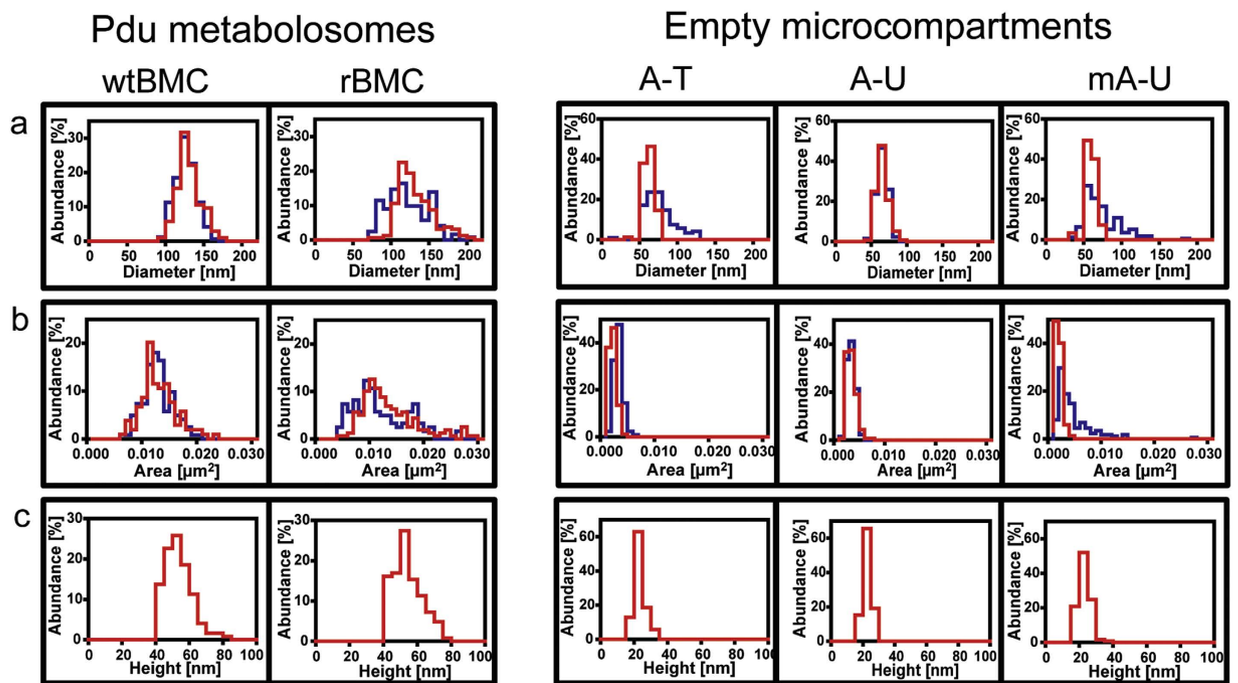


Figure 4. Comparative representation of diameter, area and height distributions of wtBMC, rBMC and eBMCs (A-T, A-U, mA-U). (a) Diameter distribution of normalized data of TEM (blue) and AFM (red), bin size is 10 nm. (b) Area distribution of normalized TEM (blue) and AFM (red) data, bin size is $0.001 \mu\text{m}^2$. (c) Height distribution of normalized AFM data, bin size is 5 nm.

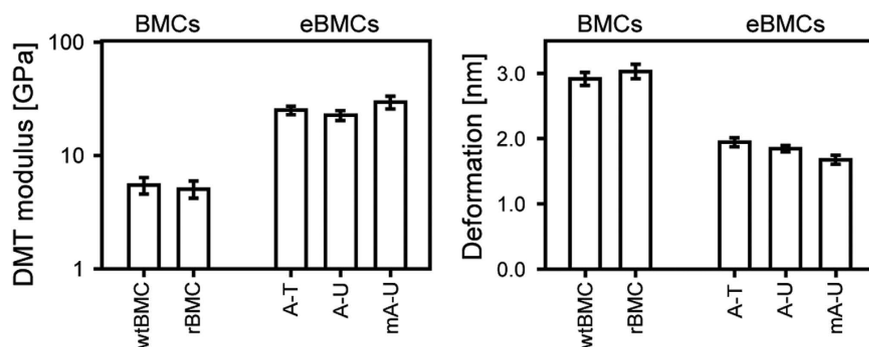


Figure 5. Quantitative nano-mechanical mapping of BMCs. The QMN mapping at 250 nN force set point for wtBMCs, rBMCs and eBMCs (A-U, mA-U, A-T). Average values and error-bars representing standard error of mean (of 20 measured particles) of reduced DMT modulus [GPa] and deformation [nm] are shown.

experiments also revealed that the reduced elastic modulus (E^*) is on average 5.0–5.4 GPa for both wtBMCs and rBMCs, whilst eBMCs displayed values between 22.8–29.8 GPa. In summary, single particle AFM measurements show striking differences in size distribution and mechanical properties for wtBMCs and rBMCs in comparison to eBMCs.

Internal structural organization. To investigate and compare the internal lumen of the various BMC structures for the presence of internalized protein cargo, isolated complete BMCs (wtBMCs and rBMCs) and eBMCs (A-T, A-U, mA-U) were embedded in resin, thin sectioned and imaged by TEM (Fig. 6). When visualized, both wtBMCs and rBMCs structures showed an intact outline with a pentameric or hexameric shape. Particles appeared heterogeneous in size and shape, as described previously, and the sampled diameter and area values corresponded to the analysis shown in Fig. 4. Strikingly, the lumen of wtBMCs and rBMC appeared highly electron dense with granular spots (Fig. 6). Such density is consistent with the physiological role of Pdu microcompartments to sequester enzymes for 1,2-PD utilization. However, no higher order arrangement of internal density was observed for the wtBMC and rBMC particles as has been observed in the carboxysome^{7,11}. In contrast, the eBMC variants (A-T, A-U, mA-U) showed intact outlines that appeared heterogeneous in size and shape as observed previously, but all had low electron density in the lumen (Fig. 6). This suggests that eBMCs form empty or hollow

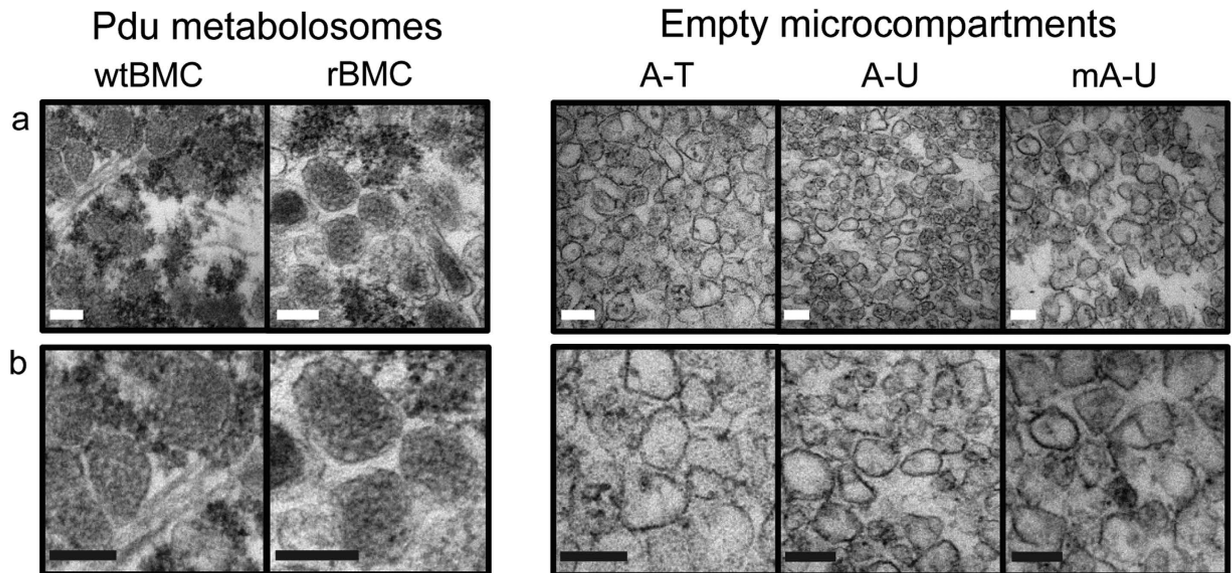


Figure 6. TEM of thin-sectioned BMCs. (a) Internal structural organization of wtBMCs, rBMCs and eBMCs (A-U, mA-U and A-T) after 70 nm sectioning. Scale bars correspond to 100 nm each. (b) Magnified images of samples in (a) showing structural organization of wtBMCs, rBMCs and eBMCs (A-U, mA-U and A-T).

structures. The formation of intact and hollow structures is not affected by the presence of PduT or mCherry fused to PduA. These observations support the hypothesis that eBMCs form intact, but empty, structures.

Discussion

In this study on the wtBMCs and their redesigned variants we have quantified structural differences between complete and empty BMCs and shown that the observed differences could be attributable to the variations in the shell protein composition of the BMC supramolecular structures. We also reveal that the presence of a protein cargo, while not required for the assembly of the BMCs, could mediate changes in the incorporation of the various shell proteins. To enable quantitative structural characterization of the BMCs, we have developed and refined purification methods in order to obtain these nano-structures in a high purity and yield. Interestingly, eBMCs purified more effectively with Y-PER™ whereas rBMCs and wild-type BMCs purified more readily with B-PER™. The differences in the purification likely reflect differences in biochemical and biophysical properties. To investigate this further, the protein composition of wtBMCs, rBMCs and the eBMCs variants (A-T, A-U and mA-U) were analyzed by a combination of 2D-PAGE and MALDI mass spectrometry. This resulted in a series of well-resolved protein spots that allowed a detailed proteomic analysis of the organelles to be undertaken. In all cases the shell proteins were identifiable, including the vertex protein PduN, whose presence had hitherto evaded identification due to its low abundance. The *C. freundii* Pdu BMC and the recombinant version produced in *E. coli* produced a similar proteomic profile to that observed with the *S. enterica* Pdu BMC¹⁶.

The quantification of the various shell proteins associated with BMC formation (Fig. 2 and Supplementary Table 4) provided some interesting observations. With wtBMCs and rBMCs, PduB is the most abundant shell protein. In contrast, with all eBMC variants PduA is the major shell protein. In both the complete (wtBMC and rBMC) and empty (A-T, A-U, mA-U) BMCs the major shell proteins included PduA, PduB, PduB' and PduJ, with the remaining shell proteins (PduK, PduN, PduT, PduU) making only a minor contribution to the overall composition of the shell of the organelle. Significantly, the presence of the mCherry-PduA fusion, the most abundant BMC shell building block, did not affect the formation of the eBMC. This helped to confirm the idea that the Pdu BMC assembly is a flexible process that is able to accommodate some variation in composition but at the same time giving rise to highly stable macro-molecular assemblies.

The size, nano-mechanical properties and internal structural organization of the wtBMCs, rBMCs and eBMCs were investigated by a complementary combination of TEM and AFM. AFM has recently been reported to provide detail on a uniformly orientated shell protein sheet⁴², and has also been used to characterize a variety of virus assemblies⁴³. These together with the present study suggest that AFM has the potential to provide detailed structural and mechanical information on native and complete BMCs *in situ*. Here, significant size distribution differences were observed between complete BMCs (wtBMCs and rBMCs) and empty eBMCs (A-T, A-U and mA-U) by single particle measurements. The average diameter of all tested eBMC variants was approximately half of that observed for wtBMC and rBMC particles. The average apparent cross-sectional area of eBMC variants was approximately 1/3 of the average apparent cross-sectional area of Pdu BMCs (wtBMC and rBMC). The average peak height of all eBMC variants imaged by AFM was around 40% of the average peak height of the wtBMCs and rBMCs. The nano-mechanical stability of wtBMC and rBMC structures measured by AFM-QNM were compared to the eBMC variants. This revealed that the complete BMCs (wtBMCs and rBMCs) are less stiff and more deformable than the smaller eBMCs. While these nano-mechanics data do not solely reflect the intrinsic elasticity

of the shell protein material since they are also influenced by the size and shape of the BMCs, they highlight the difference in mechanical behavior of the different BMC nano-structures, and corroborates the smaller size distribution and the absence of protein cargo in eBMCs compared with the complete BMCs. Moreover, detailed imaging of thin-sectioned isolated compartments confirmed that the complete BMCs (wtBMCs and rBMCs) contain a protein cargo.

What the combined biochemical and the image data tell us is that some of the structural differences between complete and empty BMCs could be attributable to variations in the shell protein composition of the supramolecular structures. Similarly, the presence of a protein cargo could also mediate changes in the incorporation of the various shell proteins. Larger BMCs, containing a protein cargo, would also be expected to be more malleable to deformation under pressure. It may be that these two latter points are connected in that the presence of a protein cargo helps in the formation of complete, larger BMCs. For instance, the current model of BMC self-assembly, which has been deduced from a study of carboxysome biogenesis, involves a nucleation event that involves the assembly of cargo proteins prior to encapsulation of preformed shell protein faces^{44,45}. However, in the absence of an aggregating protein cargo the shell proteins may self-assemble into smaller structures that are represented by the eBMCs.

Significantly, TEM imaging (Fig. 6) and 2D-PAGE data (Supplementary Figure 3) all suggest that the cellular cytoplasmic content appears to be excluded from the lumen of wtBMCs, rBMCs and eBMCs, during their formation^{36,46,47}. Similarly, it is important to note that the absence of PduT or the inclusion of a fused fluorescent protein to PduA has no influence on eBMC formation. This supports our hypothesis that BMC formation is a stable and flexible process that can be utilized in synthetic biology for the enhanced design of nano-compartments. Indeed, engineering of chimeric BMC shells in carboxysomes has been shown to be successful³². Both the incorporation of protein cargo into the BMC and shell protein composition are important parameters in defining the size and shape of the BMC assemblies. Altering the shell protein composition may represent an interesting approach for tailoring of the compartment shell size to its redesigned functions. BMCs represent a model system for which novel design principles can be determined^{33,35}. Such insights will allow the development of bioengineered internal chassis that will have application especially with metabolic pathways that involve the biosynthesis of toxic intermediates. Moreover, the ability to isolate eBMCs easily, and to unfold and refold them with a new cargo of protein or small molecule, could also allow their use as alternative delivery systems.

Material and Methods

Chemicals and reagents. All chemicals and reagents were purchased from Sigma unless stated otherwise.

Bacterial Strains and Media. A complete list of all strains and plasmids used in this study is presented in Supplementary Table 5. *E. coli* BL21(DE3) and BL21(DE3)-pLysS (Promega, Southampton, UK) transformants were grown in 2YT media in presence of appropriate antibiotics (50 µg mL⁻¹ ampicillin and/or 34 µg mL⁻¹ chloramphenicol). All strains were grown at 37 °C to an OD₆₀₀ of 0.4. Protein expression was induced with the addition of 100 µM IPTG and cultures were grown overnight at 19 °C prior to microcompartment purification and electron microscopy. Wild type *C. freundii* ballerup 7851 were grown in NCE media with succinate as the carbon source as described previously⁴¹. At an OD₆₀₀ of 1.0, 1,2-PD (1% v/v) was added to the NCE media and the cultures were grown overnight at 19 °C before microcompartment purification. For electron microscopy 50 mL of overnight culture were resuspended in fresh NCE media containing 2% (v/v) 1,2-propanediol without succinate and grown for 1 h at 37 °C/160 rpm.

Purification of Pdu Microcompartments. eBMCs (A-U, mA-U and A-T) were purified using yeast protein extraction reagent (Y-PERTM), which included an essential salt precipitation step³⁵. BMCs (rBMCs and wild type) were purified using bacterial protein extraction reagent (B-PERTM). Pdu metabolosomes were separated from cell debris at 12,000 × g. BMCs were pelleted at 20,000 × g⁴¹.

Thin sections of cells and purified microcompartments. Samples were fixed in 2 mL 2.5% (v/v) glutaraldehyde in 100 mM sodium cacodylate pH 7.2 for 2 hours, pelleted by centrifugation at 5,000 × g for 2 min and washed twice in 100 mM sodium cacodylate pH 7.2 for 15 min. Samples were post-fixed with 1% (w/v) osmium tetroxide in 100 mM sodium cacodylate pH 7.2 for 2 h followed by two washes in dH₂O. Samples were first dehydrated by incubation in an ethanol gradient: 50% (v/v) EtOH for 10 min, 70% (v/v) EtOH overnight, and twice in 100% (v/v) anhydrous EtOH for 10 min. This was followed by two final dehydration steps in propylene oxide for 15 min. Samples were equilibrated in 1 mL of a 1:1 propylene oxide and Agar LV Resin for 30 min. Samples were embedded in two steps of 100% Agar LV resin for 1.5 h each and resuspended in 0.5 mL Agar LV resin, centrifuged for 10 min at 3,000 × g in Beem capsules and polymerized at 60 °C for 16 h. Samples were thin sectioned on a RMC MT-XL ultramicrotome. Sections were stained with 4.5% (w/v) uranyl acetate in 1% (v/v) acetic acid for 45 min and 0.1% (w/v) Reynolds lead citrate for 8 min¹⁷.

TEM and AFM analysis of purified microcompartments. For EM analysis 10 µL of purified microcompartments were mounted onto Formvar, carbon coated 400 mesh, copper grids for 2 min, followed by the addition of 20 µL of 2.5% (v/v) glutaraldehyde in PBS for 1 min. The grids were washed three times in 20 µL drops of 2.5% (v/v) glutaraldehyde in PBS and then three times in dH₂O. Finally, the grids were dried and stained with 2% (w/v) aqueous uranyl acetate. Grids were analyzed at zero-loss bright field mode in an energy-filtered transmission electron microscope (Jeol 1230 TEM operated at an accelerating voltage of 80kV).

For AFM, purified microcompartment samples (wtBMC, rBMC, A-U, mA-U, A-T) were deposited onto freshly cleaved mica surfaces for five min. Microcompartments were fixed using 2.5% (v/v) glutaraldehyde in PBS for 5 min. The surface was washed three times with 1 mL sterile filtered deionized H₂O and dried under a gentle stream of N₂ gas. Images were collected in air using Bruker Multimode 8 scanning probe microscope (Bruker).

ScanAsyst Air cantilever probes (Bruker) with a nominal spring constant of 0.4 N/m were used. Height images of 2048 × 2048 to 4096 × 4096 pixels in size, covering surface areas of 10 × 10 μm to 20 × 20 μm were acquired at 20 °C in air. Sample tilt and scanner bow were removed using NanoScope 1.40 (Bruker) as described previously³⁵. Images were further analyzed using Image J 1.48 v⁴⁸ or NanoScope 1.40. Size distribution measurements of TEM and AFM images were conducted using Image J 1.48 v. Diameters were calculated assuming a circular geometry model for the microcompartments. Maximum height information of 120 single particles for each construct were obtained using NanoScope 1.40. For AFM Peak-Force QNM experiments, a TAP525 probe was used with a peak force setpoint of 250 nN. The spring constant (116 N/m) was calculated by using Sader's method⁴⁹, and the deflection sensitivity (45.23 nm/V) was calculated by ramping onto a sapphire surface. The tip geometry (8.6 nm tip radius) was estimated using an absolute method with a titanium roughness surface following the standard procedure in NanoScope 1.40. The nano-mechanical properties of twenty particles for each sample were analyzed using the DMT model, the DMT modulus and deformation was quantified (Supplementary Table 6) using NanoScope 1.40.

2D-PAGE. Purified microcompartments (160 μg of wtBMC, rBMC, A-U, mA-U or A-T) were used for isoelectric focusing. Strips (pH 3.0–10) were sealed on top of 15% (v/v) Tris-glycine gels, and run at 150 V for 150 min⁵⁰. Gels were stained using Coomassie Brilliant Blue and spots were picked for MALDI ToF-ToF. The protein content was quantified relative to PduA using Image J 1.48 v of 3 independent cultures (N = 3). Standard errors of mean are given.

In-gel digestion from MS and MS-MS analysis. Excised, diced protein gel spots were washed, reduced and S-alkylated as described previously⁵¹. A sufficient volume of 2 ng μL⁻¹ of trypsin (modified sequencing grade, Promega, Southampton, UK) in 25 mM ammonium bicarbonate was added to cover the gel pieces and digestion was performed overnight at 20 °C. The digests were then acidified by the addition of half the volume of 50% (v/v) acetonitrile with 5% (v/v) formic acid prior to MS and MS-MS analysis⁵².

MS and MS-MS analysis. The sample (1 μL of the above peptide digest) was placed on the sample target (AnchorChip standard, 800 μm, Bruker) and dried. 0.5 μL of matrix was added and dried. The matrix was α-cyano-4-hydroxy-cinnamic acid (α-CHCA, 0.7 mg mL⁻¹ in 85% (v/v) acetonitrile, 0.1% (v/v) TFA and 1 mM NH₄H₂PO₄, Sigma-Aldrich, Gillingham, UK). For external calibration in the protein mass range, Peptide Calibration Standard I (Bruker) was used. MALDI-TOF MS and MALDI TOF-TOF MS-MS analysis was performed in the positive ion mode using a Bruker UltrafleXtreme. The spectra were obtained in reflector mode with an acceleration voltage of 25 kV and a pulse ion extraction time of 80 ns. The mass range for MS was generally between 700 and 3500 m/z. The number of laser shots summed in MS was 3500. The number of laser shots summed in MS-MS was 3000. The software FlexAnalysis (Bruker, Bremen, Germany) was used for peak picking prior to using the standard Mascot search engine (Matrix Science Ltd, London, UK). The peptide mass fingerprint was searched against the NCBI proteobacteria protein database⁵².

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Author Contributions

M.M. undertook most of the experimental work and helped write the manuscript. R.J. and I.R.B. developed systems for the thin-sectioning of BMCs and undertook the majority of the TEM imaging. E.D. helped with the construction of the various plasmids used in this study, whilst D.M.B. and D.J.P. assisted with the AFM. W.F.X., S.F. and M.J.W. conceived the experiments, were involved with data interpretation and helped write the manuscript.

Additional Information

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