Biochemical and Structural Insights into Bacterial Organelle Form and Biogenesis*

Many heterotrophic bacteria have the ability to make polyhedral structures containing metabolic enzymes that are bounded by a unilamellar protein shell (metabolosomes or enterosomes). These bacterial organelles contain enzymes associated with a specific metabolic process (e.g. 1,2-propanediol or ethanolamine utilization). We show that the 21 gene regulon specifying the pdu organelle and propanediol utilization enzymes from Citrobacter freundii is fully functional when cloned in Escherichia coli, both producing metabolosomes and allowing propanediol utilization. Genetic manipulation of the level of specific shell proteins resulted in the formation of aberrantly shaped metabolosomes, providing evidence for their involvement as delimiting entities in the organelle. This is the first demonstration of complete recombinant metabolosome activity transferred in a single step and supports phylogenetic evidence that the pdu genes are readily horizontally transmissible. One of the predicted shell proteins (PduT) was found to have a novel Fe-S center formed between four protein subunits. The recombinant model will facilitate future experiments establishing the structure and assembly of these multiprotein assemblages and their fate when the specific metabolic function is no longer required.

It has been recognized for more than 30 years that all cyanobacteria (1) and some other chemoheterotrophic bacteria (2) contain carboxysomes. These polyhedral inclusions consist of a proteinaceous shell enclosing an active enzyme, ribulose bisphosphate carboxylase/oxygenase (RuBisCO). Their function is to enhance the fixation of carbon dioxide (3), a reaction of planetary significance in that marine cyanobacteria are responsible for the majority of global carbon fixation (4, 5). More recently, sequence similarity was noticed between carboxysome shell genes and metabolic operon genes associated with propanediol utilization (pdu) and ethanolamine utilization (eut) in a variety of heterotrophic bacteria found in the mammalian gut (3) and the environment. In growth conditions that induce these metabolic operons, polyhedral organelles resembling carboxysomes were observed on electron microscopy of Salmonella enterica serovar Typhimurium (6), Klebsiella oxytoca, Citrobacter freundii, and Escherichia coli (7). Bioinformatics analysis also locates genes resembling carboxysome shell genes in metabolic operons in Clostridium perfringens (8), Clostridium tetani (9), Listeria monocytogenes and Listeria innocua (10), Enterococcus faecalis (11), Lactobacillus collinoides (12), Citrobacter rodentium, Yersinia enterocolitica (13) among other organisms. The non-carboxysome polyhedral structures have been referred to as enterosomes (3) or metabolosomes (14), emphasizing their role in cellular metabolism.

There is some considerable interest in how these proteinaceous organelles form and the arrangement of protein subunits that give rise to these remarkable macromolecular assemblies. In carboxysomes, there are thought to be a number of shell proteins that encase the RuBisCO and carbonic anhydrase. The situation is more complex in metabolosomes, where there are at least five shell proteins that encase ancillary factors, metabolic enzymes, and activating factors. There are thus between 17 and 21 genes associated with the ethanolamine and propanediol metabolosomes, respectively, and although some functional studies have been undertaken, little is known about the topological arrangement of the encoded protein components within the organelle. However, sequence analysis reveals that the shell proteins found in carboxysomes and metabolosomes are similar, indicating that they have evolved from a common ancestor.

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** The on-line version of this article (available at http://www.jbc.org) contains supplemental Figs. S1–S3 and Tables S1 and S2.

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1 These authors contributed equally to this work.

2 To whom correspondence may be addressed. Tel.: 44-1227-824690; E-mail: m.j.warren@kent.ac.uk.

3 To whom correspondence may be addressed. Tel.: 353-21-4901420; E-mail: m.prentice@ucc.ie.
Structural studies on some of the individual shell proteins have given an insight into how they may function. The main carboxysome shell protein, CCMK1, has been shown to have a hexameric crystal structure with a charged pore (15), suggesting a selective permeability mechanism making the structure a prokaryotic functional equivalent to a eukaryotic organelle. Structures of the CsoS1A carboxysome protein from Halothioabacillus neapolitanus (16) and the EutN shell protein of the ethanolamine utilization enterosomes from E. coli (17) also reveal similar hexameric arrangements with charged residues surrounding a central pore. Cryoelectron microscopy of extracted carboxysomes shows they have an icosahedral symmetry with triangular faces, but a complete image reconstruction was not possible because of a variability in individual carboxysome size, lack of symmetry in carboxysome contents, and absence of obvious capsomers on the surface (18). A similar imaging project has provided even further detail, suggesting that carboxysomes isolated from a Synechococcus strain contain ~250 RuBisCOs, which are organized into three or four concentric layers (19).

The distribution and relatedness of both the structural genes for the protein shell (15) and the associated metabolic operons (20, 21) strongly suggest repeated horizontal transmission in evolution. In keeping with this, we show that plasmid-based expression in E. coli of a pdu operon from C. freundii containing metabolosome structural and enzyme genes results in synthesis of recombinant metabolosomes with fully functional enzyme activity. In this way we have compartmentalized the bacterial cytoplasm, showing that it is possible to create novel compartments by horizontal genetic transfer in bacteria and to vary the shape and topology of the organelle by overproducing individual shell proteins. This process has the potential to be used for metabolic engineering whereby bacteria can be shielded from toxic metabolic intermediates. Thus, this project will contribute to the emerging discipline of synthetic biology.

**EXPERIMENTAL PROCEDURES**

A more detailed account of the techniques, including the cloning and sequencing of the pdu operon is given in the supplemental material.

*Citrobacter sp. Library*—The genomic DNA of Citrobacter sp. was partially digested with Sau3AI, and 20–30-kb fragments were purified after separation by agarose gel electrophoresis, cloned into the BglII site of the cosmid pLA 2917, and encapsidated in vitro following the Packagene® Lambda DNA packaging system instructions (Promega). The library was used to infect E. coli LE392. One cosmid, which had the ability to degrade 1,2-propanediol, was isolated and named pAR3114 (supplemental Table S1).

*Bacteria and Growth Conditions*—All strains and plasmids are described in supplemental Table S1. E. coli JM109 wild-type and recombinant strains harboring pLA2917 (22) and pAR3114 were grown either in LB medium or trypticase soy agar (10 g of tryptone, 5 g of NaCl, and 15 g of agar in 1 liter of distilled water) without or with tetracycline (20 μg/ml), respectively. pAR3114 is a pLA2917 cosmid containing an ~30-kb (pdu operon, cbiA and pocR genes) insert derived from Sau3AI digestion of the C. freundii chromosome.

**Sequencing of Insert, Gene Annotation, and Recombinant Manipulation**—The pdu operon sequence was obtained from pAR3114 after it was sent to be sequenced by Qiagen Ltd. The insert was found to contain 28,618 bp. The sequencing coverage was higher than 12, and the data quality has an error rate of <1/10000 bp and a base accuracy of >99.99%. The cloning and overproduction of individual Pdu components was achieved as described in the supplemental material.

**Propanediol Utilization Assay**—The utilization of 1,2-propanediol was tested by a MacConkey triple-plate method (20). Bacteria were streaked on to MacConkey agar (20 g Soy peptone, 5 g of ox bile, 5 g of NaCl, 0.075 g of neutral red, 1% 1,2-propanediol, and 12 g of agar in 1 liter of distilled water) plates containing 1,2-propanediol, 20 μM cobalt chloride, or 100 mM cobinamide. Plates were incubated at 37 °C for 24 h. Formation of red colonies indicated the utilization of 1,2-propanediol by the reduction in pH due to the formation of propionate, and a more rapid appearance of red color with the addition of cobinamide indicated cobalamin dependence of propanediol utilization.

**Isolation of Metabolosomes**—Metabolosomes were isolated by the organelle purification procedure as described previously (7) and outlined further in the supplemental material.

**Electron Microscopy**—Late logarithmic phase cells of E. coli pLA2917 and pAR3114 were prefixed in 2.5% (v/v) glutaraldehyde, 10 mM Hapes, pH 7.0, for 72 h. After immobilization in 2% (w/v) water-agar they were postfixed in 1% (w/v) osmium tetroxide in 75 mM cacodylate, pH 7.2, for 60 min at 4 °C followed by dehydration in an ethanol series. The 70% dehydration step, supplemented with 1% (w/v) uranylacetate, was done overnight at ambient temperature following the method of Bobik et al. (6). Ultrathin sections (90 nm) were post-stained with 4% (w/v) aqueous uranylacetate and analyzed at zero-loss bright field mode in an energy-filtered transmission electron microscope (Zeiss CEM 902, Oberkochen, Germany).

Isolated polyhedral bodies were fixed in 1% (v/v) glutaraldehyde, and after adsorption to Formvar-carbon-coated grids they were negatively stained with 2% (w/v) uranylacetate, pH 4.5. Samples were analyzed by energy-filtered transmission electron microscope, and images were recorded, in general, with a charge-coupled device camera (CCD; Proscan Electronic Systems, Scheuring, Germany). Fast Fourier transformations (FFT) of individual particles was done with CRISP software (Calidris, Sollentuna, Sweden).

SuperSTEM microscopy was carried out on a VG HB501 dedicated STEM fitted with a Nion second generation spherical aberration corrector, high angle annular dark field detector (angular range 70–210 mrad), and a Gatan Enfina electron microscope, and images were recorded, in general, with an Oxford helium flow cryostat as described previously. Redox
titrations were performed in a Belle Technology glove-box under a nitrogen atmosphere, essentially as described previously (24, 25).

RESULTS

Utilization of 1,2-Propanediol—Transformation of E. coli JM109 cells with a cosmider library made from a partial digest of the C. freundii genome resulted in the identification of a single red colony when the library was plated on propanediol-MacConkey medium (PM medium). The appearance of this red colony was likely due to the ability of the recombinant strain to metabolize 1,2-propanediol, a phenotype absent in wild-type E. coli. The cosmider, termed pAR3114, was found to have a \\( \sim 30 \) kb insert.

The insert was sequenced, which revealed that it encoded 30 contiguous genes involved in cobalamin biosynthesis and propanediol utilization and five other genes localized at one end of the insert. These genes and the putative functions of the encoded protein products are described in Table 1. The resulting plasmid, when transformed into JM109, bacterial colonies turned red when grown on PM indicator plates, demonstrating that the transformed bacteria had gained the ability to metabolize propanediol. A further deletion was made whereby \( \text{pduA} \) and \( \text{pduB} \) were removed from the insert in pED460 to give pED461 (supplementary Table S1). The resulting plasmid, when transformed into E. coli JM109, had a greatly reduced ability to metabolize propanediol on PM medium in that the colonies turned only slightly pink. Reconstitution of propanediol metabolism was achieved by adding \( \text{pduA-B} \) back in trans on a separate, compatible plasmid (pLysS). However, the individual addition of either \( \text{pduA} \) or \( \text{pduB} \) did not restore propanediol metabolism, demonstrating that both proteins are required for this activity.

Electron Microscopy—Cobalamin-dependent propanediol utilization is associated with the formation of intracellular inclusions called carboxysome-like bodies, enterosomes, or metabolosomes (6). We therefore examined our transformed cells for evidence of metabolosomes. Ultrathin sections of E. coli pAR3114 grown on minimal medium in the presence of 1,2-propanediol showed intracellular metabolosomes (Fig. 2b). E. coli pLA2917 (empty cosmider) control strain grown on mini-

### TABLE 1

| Gene  | Start | End | Function                                      | Length | Direction of transcription | Predicted mass |
|-------|-------|-----|-----------------------------------------------|--------|---------------------------|----------------|
| cbkB  | 835   | 1   | Cobalamin biosynthesis protein                | 834    | —                         | 17,817         |
| cbIA  | 2,211 | 832 | Cobyricin acid a,c-diamide synthase            | 1379   | —                         | 49,669         |
| pocr  | 3,718 | 2,807| Transcriptional regulator                     | 911    | —                         | 49,636         |
| pduE  | 4,729 | 3,920| Propanediol diffusion facilitator             | 809    | —                         | 28,125         |
| pduA  | 5,255 | 5,532| Shell protein                                 | 277    | +                         | 9,376          |
| pduB  | 5,536 | 6,348| Shell protein                                 | 812    | +                         | 28,021         |
| pduC  | 6,367 | 8,031| Diol dehydratase large subunit                | 1664   | +                         | 60,280         |
| pduD  | 8,042 | 8,716| Diol dehydratase medium subunit               | 674    | +                         | 24,280         |
| pduE  | 8,731 | 9,249| Diol dehydratase small subunit                | 518    | +                         | 19,233         |
| pduG  | 9,265 | 11,097| Diol dehydratase reactivation protein         | 1832   | +                         | 64,163         |
| pduH  | 11,087| 11,437| Shell protein                                 | 350    | +                         | 12,632         |
| pduL  | 11,457| 11,732| Shell protein                                 | 275    | +                         | 9,053          |
| pduK  | 11,823| 12,227| Shell protein                                 | 404    | +                         | 13,921         |
| pduL  | 12,227| 12,859| Phosphotransacetylase                         | 632    | +                         | 23,048         |
| pduM  | 12,856| 13,347| Unknown                                       | 491    | +                         | 18,175         |
| pduN  | 13,351| 13,626| Shell protein                                 | 275    | +                         | 9,205          |
| pduO  | 13,635| 14,642| Cobalamin adenosyltransferase                 | 1007   | +                         | 45,886         |
| pduP  | 14,639| 16,024| CoA-dependent propionylcarboxylic dehydrogenase | 1385   | +                         | 48,671         |
| pduQ  | 16,035| 17,147| Propanol dehydrogenase                        | 1112   | +                         | 39,702         |
| pduS  | 17,144| 18,499| Cobalamin reductase                           | 1355   | +                         | 48,568         |
| pduT  | 18,502| 19,056| Shell protein                                 | 554    | +                         | 19,038         |
| pduU  | 19,056| 19,406| Shell protein                                 | 350    | +                         | 12,483         |
| pduV  | 19,411| 19,863| Shell protein                                 | 452    | +                         | 16,331         |
| pduW  | 19,848| 21,062| Shell protein                                 | 1214   | +                         | 43,634         |
| pduX  | 21,062| 22,021| Unknown                                       | 929    | +                         | 33,908         |
| pduY  | 22,021| 22,636| Shell protein                                 | 1058   | +                         | 39,618         |
| DacD  | 23,694| 24,372| DD-carboxysomepeptidase                        | 1172   | —                         | 43,085         |
| plsc  | 26,450| 26,683| Thiosulfate reductase cytochrome B subunit     | 767    | —                         | 28,545         |
| plsb  | 27,035| 27,456| Thiosulfate reductase protein                  | 579    | —                         | 21,622         |
| plsu  | 28,618| 27,043| Thiosulfate reductase precursor                | 1575   | —                         | 53,778         |
mal medium in the presence of 1,2-propanediol did not show metabolosomes inside the cells (Fig. 2a). When grown on rich medium (LB), *E. coli* pAR3114 contained metabolosomes in the presence or absence of added propanediol. In cells grown on rich medium, the metabolosomes were closely packed and formed up to 90% of the observed section surface. Again, control strains containing the empty cosmid did not reveal any such structures when grown under similar conditions.

In ultrathin cell sections, the metabolosomes measured on average 101 nm across their widest diameter (n/1000 = 63; S.D./1000 = 25 nm). There were two maxima in the frequency distribution of measured diameters, compatible with two different size bodies or different sections of a non-spherical polyhedral structure with shorter (77–92 nm) and longer (108–123 nm) axial dimensions (Fig. 2d). These dimensions are similar to those reported for wild-type metabolosomes formed in *S. enterica* (7), and carboxysomes from *H. neapolitanus* also show variable sizes with two distinct maximal diameters (18). When looked at in detail (Fig. 2c, white arrows), apparent sections of polyhedral bodies showed a weak linearly ordered substructure, indicating a certain degree of interior package order. Additionally, polar deposits of electron-dense amorphous granules could be observed in many recombinant cells (Fig. 2b, asterisks).

When the *E. coli* strain containing the *pdu* operon, but missing *pduA-B*, was analyzed by electron microscopy, no metabolosomes were observed (data not shown). Metabolosomes were observed again when this strain was complemented with a compatible plasmid containing *pduA-B* (data not shown). However, no organelles or other structures were seen in strains in which only *pduA* or *pduB* was added back in trans (data not shown). This result indicates that both PduA and PduB are required for organelle biogenesis. Previous research on the *S. enterica pdu* operon had shown that PduA is essential for organelle formation (26).

**Diol Dehydratase Activity**—The activity of diol dehydratase was measured from the protein extracted from *E. coli* pAR3114 grown to stationary phase in the presence and absence of 1,2-propanediol. Diol dehydratase activity was detected at 2.23 units/mg whole-cell protein for *E. coli* pAR3114 grown in the presence of 1,2-propanediol. Activity was reduced to 0.72 unit/mg when *E. coli* pAR3114 was grown in the absence of 1,2-propanediol. Recombinant metabolosomes were extracted from cells grown in minimal medium with propanediol as per published methods for the extraction of wild-type metabolosomes (7). Purified metabolosomes derived from *E. coli* pAR3114 grown in the presence of 1,2-propanediol showed diol dehydratase activity of 32.03 units/mg protein, ~14 times higher than the enzyme activity found in the crude protein extract.

**Extracted Polyhedral Metabolosomes**—Metabolosomes were purified from *E. coli* pAR3114 as described under “Experimental Procedures” and in the supplemental material and were viewed both by an energy-filtered transmission electron microscope in the elastic bright field mode and a novel aberration-corrected non-contrast electron microscopy technique using a SuperSTEM instrument. No such structures were obtained from *E. coli* control strains, indicating that the metabolosomes were generated as a result of the presence of the *pdu* operon. The interior of negatively stained polyhedral bodies of *E. coli* pAR3114, fixed with glutaraldehyde (Fig. 3a), showed a finely
branching substructure. These particles had a mean diameter of 92 nm (n = 143; S.D. = 32 nm) with a frequency distribution showing 1 maxima (presumed short axis) from 62 to 77 nm and a presumed long axis from 108 to 123 nm (Fig. 2e). Higher magnifications of those particles showed regularly arranged substructures with a layer spacing of about 7 nm (Fig. 3b, white arrows). Intrinsic order on or below this scale within isolated polyhedral bodies was detected by FFT (Fig. 3c, FFTs 2 and 3) relative to the amorphous foil substratum (Fig. 3c, FFT 1). Distinct frequency intensities within a 128 × 128 matrix were obvious at 0.30 and 0.41 per nm, corresponding to a particulate resolution of 3.3 and 2.4 nm (Fig. 3c, FFT 2). That is, the electron micrographs shown resolve structures as small as 2.4 nm, confirming their ability to detect 7-nm layers.

Further evidence for regular metabolosome substructure was obtained using SuperSTEM microscopy of the recombinant metabolosomes both in cell sections and protein extracts. A number of particles displayed apparent multicomponent substructure, with smaller particles containing electron density typical of proteins (Fig. 4, a and b). This suggests that the interior of the metabolosome is not an amorphous mixture of component molecules but constitutes a regular assembly of structural proteins. Such a regular assembly has recently been reported for carboxysomes from Synechocystis, where the RuBisCOs are organized into three or four concentric layers (19). Currently crystal structures of carboxysome and metabolosome shell proteins show them forming flat hexagonal sheets, although it has not been determined how this planar structure can bend or fold up to form a closed shell (15, 16). Our SuperSTEM images showing discrete electron-dense foci spaced around the recombinant metabolosomes suggest the presence of non-planar protein complexes acting as structural linkage points joining sheets of shell protein to form the faceted surface of the metabolosome (Fig. 4, c and d) in the same way that vertex proteins link planar proteins to form the viral capsid structure (27, 28). As in viral capsids, these complexes are presumably pentameric, connecting planes of hexameric shell proteins (27, 28). The likely candidate for this metabolosome com-
ponent is one (or more) of the three predicted shell proteins, just as minor variants of viral capsid proteins normally forming hexameric sheets form pentameric vertex proteins (28).

**Liquid Chromatography-Tandem Mass Spectrometry SDS-PAGE Analysis**—To identify proteins present within the metabolosome, purified metabolosome extracts were subjected to SDS-PAGE, and individual protein bands were extracted and analyzed (Fig. 5 and supplemental Table S2). The proteome pattern of metabolosomes isolated from recombinant *E. coli* pAR3114 grown in the presence of 1,2-propanediol is similar to that reported for wild-type *S. enterica* (7). By using MALDI-TOF on trypsin digests of the extracted proteins run out on SDS-PAGE, it was possible to identify PduA, C, D, E, J, O, P as the major components of the recombinant metabolosome. The presence of these proteins as well as PduB’ was further confirmed by two-dimensional gel electrophoresis of the isolated organelle (see supplemental Fig. S1 and Table S2). The 11 major bands seen on the SDS-gel of the purified organelle (Fig. 5) resolved into around 100 spots on the two-dimensional gel, indicating a further layer of complexity. A stoichiometric analysis of the *pdu* organelle proteome from *C. freundii* has yet to be determined. Overall, however, the protein composition of the isolated organelle appears to be similar to the proteomic analysis of the *S. enterica pdu* metabolosome (7).

**Two Forms of PduB: B and B’**—The *S. enterica* PduB appears to be produced in two forms, one of which is about 5 kDa smaller than the other (7). It was suggested that this was caused by the presence of two translational start sites giving rise to two forms of the protein, referred to as PduB and PduB’ (7). To investigate whether this is also the case in the *C. freundii* Pdu metabolosome, the recombinant production of PduB and B’ from pduB was investigated.

Overproduction of the *C. freundii* PduB in *E. coli* was achieved by cloning the gene into pET14b to allow the protein to be produced with an N-terminal His tag. Analysis of the protein profile of the recombinant strain overproducing the *C. freundii* PduB revealed the presence of two major protein bands by SDS-PAGE. When the crude cell extract containing the overproduced PduB was passed through a Ni²⁺ column, both the higher (37 kDa) and lower (32 kDa) molecular mass proteins copurified after elution in imidazole (supplemental Fig. S2a). The bands were confirmed as PduB-type proteins by MALDI-TOF analysis after their excision from the SDS-gel and trypsin digestion. Significantly, only the higher molecular mass protein cross-reacted with anti-His tag antibodies when analyzed by Western blotting (supplemental Fig. S2b). These results are consistent with pduB having two translation start sites. The two observed protein bands represent one with an N-terminal His tag and the second one starting at Met-38. The two proteins must form a tight complex, as they were able to copurify on the nickel column. Confirmation of the two start sites was obtained by mutating the codon for Met-38 within *pduB* to Ala (see sup-
The resulting protein (PduB only) was produced and purified as a single product (data not shown).

Overproduction of the Shell Proteins, PduA, -B, -J, -K, -T, and -U, in Situ with the Metabolosome Results in Aberrant Structures—Although the absence of pduA or pduB from the pdu operon leads to the loss of metabolosome formation, it was found that overproduction of any one of PduA, -B, -B’/H11032, -B/H11032, -J, -K, -T, and -U, in a background in which recombinant metabolosomes formed, resulted in significant effects upon size, shape, and aggregation of the organelle (Fig. 6). For overproduction of the individual shell proteins, the relevant gene was cloned into pLys, as described in the supplemental material, a plasmid that is compatible with pED460 (which contains all the pdu genes from pduA to pduX). The pLysS derivative was then transformed into an E. coli strain containing pED460 and the resulting derivative analyzed for the presence of cytoplasmic structures. To determine whether the gene product encoded within pLysS was being overproduced in situ, cell lysates of the corresponding transformed strains (containing the pLys derivative plus pED460) were analysed. In all but one case, a clear band corresponding to the expected size of the protein could be observed on the SDS-gel at an intensity well above the control strain (supplemental Fig. S3). The one exception was PduU, where no overproduced protein band was seen (supplemental Fig. S3).

Overproduction of PduA within a strain housing pED460 resulted in the appearance of long thin structures in the recombinant E. coli, indicating that excess PduA prevents the formation of rounded organelles (Fig. 6B). A similar observation had been made previously on PduA overproduction in S. enterica (26). Overproduction of PduB and B’ gave large single homogeneous structures with no delimiting feature (Fig. 6C). The structure also appeared more granular indicating that too much PduB and B’ prevents normal metabolosome biogenesis. Striking changes in organelle morphology were obtained by the separate overproduction of PduB and PduB’. In the case of the former, large circular bodies were observed containing concentric rings (Fig. 6D), whereas with the latter regular straight-lined structures were generated (Fig. 6E). From this it would appear that the presence of PduB is important for generating curved structures and that the appropriate level of PduB and PduB’ is important for the shape of the metabolosome. The curvature effect must be a property of the first 38 amino acids present in the N terminus of PduB.

No aberrant structures were observed when pduA and pduB were expressed together to complement the pduA,B knock-out strain. This is likely because the levels of PduA and PduB are low when the two genes are expressed together, in comparison to the much higher levels of protein production that would be expected with the single gene expression studies.

With the overproduction of PduJ, the organelle took on a rosette appearance (Fig. 6F). In this case, several layers of (presumably) shell protein produced a whorled architecture. Overproduction of PduK led to the appearance of a single large aggregate complex that still maintained the delimiting substructure feature of individual metabolosomes (Fig. 6G). A noticeable change in organelle appearance was observed with attempts to overproduce PduT. Here, cells were either found

FIGURE 6. Effect of shell protein overproduction on metabolosome formation. A, a thin section of E. coli pAR3114 (housing pduA-X) with recombinant metabolosomes. The effect of overproducing certain individual shell proteins within E. coli pAR3114 by expressing the respective gene on a separate compatible plasmid is shown in B–I. B shows the effect of overproducing PduA; C, the effect of overproducing PduB and PduB’; D, the effect of overproducing PduB; E, the effect of overproducing PduB’; F, the effect of overproducing PduJ; G, the effect of overproducing PduK; H, the effect of overproducing PduT; I, the effect of overproducing PduU; and J, a negative control (E. coli containing the empty cosmids vector pLA2914 and pLysS).
with deposits or having lamina-like structures (Fig. 6f) not dissimilar to that observed with PduA overproduction, where a layered appearance was observed. When PduU was overproduced, multiple smaller structures were observed, which appeared to clump together (Fig. 6f, j). In contrast, overproduction of the only remaining putative shell protein, PduN, had no effect upon the appearance of the organelle (data not shown). From these studies it can be concluded that PduA, -B, -B', -J, -K, -T, and -U play important roles in organelle biogenesis and help to shape the structure. Moreover, the data indicate that organelles require a specific ratio of these shell proteins to orchestrate the correct assembly of the metabolosome.

No structures were observed in strains that contained only the pLysS derivatives. The aberrant organelles were observed only in strains that contained the pLysS derivatives. Normal organelles were observed when pLysS (with no insert) was transformed into E. coli cells containing pED460. Therefore, the changes in organelle morphology must be due to the excess protein encoded within the pLysS derivative.

PduT Contains an Fe-S Center—Overproduction of PduT resulted in the isolation of a brown-colored protein in which the UV-visible spectrum had a broad absorption maximum at 420 nm (Fig. 7a), suggesting the presence of an Fe-S center. This was confirmed by EPR analysis, where X-band EPR spectra of PduT (Fig. 7c) indicate the presence of a 4Fe-4S iron-sulfur cluster in this protein. The protein as isolated gives rise to a rhombic EPR spectrum at 15 K with $g_{||} = 1.91$ and $g_{\perp} = 2.04$. This spectrum was not visible at 70 K (data not shown). Both of these properties are indicative of the presence of a $[4Fe-4S]^{1+}$ cluster in the protein as isolated, suggesting that a proportion of the cluster is reduced in the protein under these conditions. Reduction with dithionite increases the amount of $[4Fe-4S]^{1+}$ detected and confirms the assignment of the signal to a reduced iron-sulfur cluster. The properties of the iron-sulfur cluster suggest that it has a relatively high midpoint redox potential ($E_m$) and that it is accessible to external redox agents. This was confirmed by redox potentiometry, which revealed that the midpoint potential of the cluster was +99 mV (Fig. 7b). Gel filtration analysis (data not shown) suggests that the protein exists as a homotetramer, and as there are less than four conserved cysteines in the protein, the possibility exists that this center is formed from the coordination of several separate subunits.

To investigate further this possibility, we also examined Cys → Ala mutants of the three cysteine residues found in this protein PduT using EPR spectroscopy. Dithionite-reduced proteins (Fig. 8) exhibit the same characteristic $[4Fe-4S]^{1+}$ EPR spectrum in the C136A and C108A mutants, but this spectrum cannot be obtained for the C38A mutant. This suggests that only one cysteine residue from each PduT protein ligates a 4Fe-4S cluster, a highly unusual arrangement but consistent with the proposed homotetrameric structure. This arrangement may also explain the unusually low $g^\perp$ value. Cys-38 is the only cysteine residue conserved in known PduT sequences.

**DISCUSSION**

Phylogenetic evidence strongly suggests that horizontal transfer of the propanediol utilization operon has occurred several times in the evolution of the related Enterobacteriaceae,
Recombinant Organelle Biogenesis

S. enterica and E. coli (20, 21), despite the number of genes involved (>20 kb of DNA). We attempted to reproduce this event in vitro between C. freundii and E. coli to show that the complex systems involved were indeed transferable in one step. Electron microscopy of E. coli cells carrying plasmids or cosmids harboring the pdu genes that had been grown on propanediol medium and not on succinate alone showed polyhedral bodies closely resembling in size and shape those previously reported from other Enterobacteriaceae in which metabolosomes have been induced (3, 6, 7, 29). Assays and protein analysis suggested that these bodies were functioning metabolosomes. In E. coli strains containing the cosmids pAR3114, their presence was induced by propanediol and the recombinant host acquired cobalamin-dependent propanediol utilization capacity. In strains carrying pED460, where the regulatory gene had been trimmed, the metabolosomes were constitutively produced.

As for wild-type metabolosomes (7), enzymatic assay of B13-dependent diol dehydratase showed enriched activity in extracted metabolosomes compared with the whole bacterial cell, suggesting concentration of the metabolic enzymes within the organelle. Moreover, the protein components of the purified organelle were found to correspond to the encoded Pdu proteins, thereby confirming the efficacy of the recombinant metabolosome.

The host E. coli JM109 contains a functional ethanolamine utilization (eut) operon as shown by a standard bioassay (20), implying possession of a set of shell protein genes associated with the E. coli ethanolamine utilization operon closely resembling that in S. enterica (30). However, no metabolosomes were seen in the control E. coli strain that were capable of ethanolamine utilization but lacking pdu operon genes when grown on propanediol medium, indicating that these genes were not significantly expressed in propanediol medium. This is the first demonstration that the full functionality of a metabolosome can be horizontally transferred by standard recombinant DNA procedures in vitro.

Electron microscopy of extracted recombinant metabolosomes revealed that a high percentage of extracted metabolosomes had ragged surfaces, unlike the previous report in which the surface protein layer was mainly intact (7), even though the metabolosome extracts were fixed prior to negative staining to minimize rupture. This may be because of slight differences in the way the samples were prepared. The dimensions of the extracted organelles were still close to those found for intracellular ones, and the disruption was therefore presumably confined to the surface layer. Apart from this, electron microscopy appearances were similar overall to those reported previously (2, 3, 6, 7), although we noted two features not previously commented on for metabolosomes but recently established in carboxysomes. First, the frequency distribution of metabolosome diameter showed two distinct maxima, suggesting a common elongated nonspherical structural plan. The apparently variable polyhedral appearances seen on electron microscopy may be a combination of fixation artifacts and random cross-sections. Second, a higher intrinsic order within the organelle was detected by FFT, which is compatible with assembly of the organelles from protein subunits of particulate resolution 3.3 and 2.4 nm. A regular substructure is compatible with cryoelectron microscopic studies of H. neapolitanus (18) and Synchococcus (19) carboxysomes.

It was also observed that an E. coli strain carrying a plasmid with the main C. freundii pdu genes but deleted in pduA and pduB had greatly reduced propanediol metabolizing ability. Analysis by electron microscopy of this strain revealed that it had no metabolosomes. However, complementation with a compatible plasmid containing both pduA and pduB resulted in restoration of full propanediol-utilizing activity and the formation of intracellular organelles. Thus, the presence of the metabolosome appears to enhance the metabolic activity of the organelle-associated enzymes. Furthermore, as the plasmid harboring pduA and pduB, but not plasmids containing pduA or pduB, was able to form metabolosomes in this particular E. coli strain, it suggests that PduA and PduB are both independently required for metabolosome formation. Moreover, the presence of both PduB and B’ on two-dimensional gels indicates that both proteins are components of the metabolosome.

Organelle formation is dependent upon the appropriate concentration of certain components, as too little or too much of some of the individual proteins can prevent organelle formation or lead to the formation of aberrant structures. Signifi-
cantly, overproduction of PduA, -B, -B’, -B and -B’, -J, -K, -T, and -U, which are all predicted to be shell proteins, in an E. coli strain that harbors the pdu operon gives rise to abnormal bodies. The differences in shape and form of the organelle when these proteins are overproduced certainly support a role for these proteins in the outer shell of the organelle. The various aberrant organelles were not studied further, and no attempts were made to compare their protein composition with that of wild-type structures. Similarly, no protein localization studies have yet been undertaken to demonstrate whether the putative shell proteins really do form the shell of the structure.

Unexpectedly, isolation of one of these proteins, PduT, revealed that it contained a 4Fe-4S center, with an unusual Cys ligation pattern within a PduT homotetramer, indicating that the organelle has some form of molecular wiring. The presence of such a redox group represents a major engineering challenge, on the same scale as the carboxysome. It is interesting to note that the cobalamin reductase, PduS, also contains a 4Fe-4S center, with an unusual Cys ligation pattern within a PduT homotetramer, indicating that single electron transfer or redox sensing can be carried out through intact organelle shells without using the structurally established pores or other gaps in the shell proteins. This is an important insight into the metabolic circuitry of the metabolosome, which may also be shared by the carboxysomes, such as how the components are pieced together, how the enzymes are internalized, and how substrates and products are moved in and out of the organelle (33). These structures offer the potential for future exploitation in engineered reactions using the organelle architecture to house novel active metabolic processes. Our studies represent the first step toward engineering such systems and toward the rational design of specific prokaryotic cytosolic compartments.

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