

Characterization of a Planctomycetal Organelle: a Novel Bacterial Microcompartment for the Aerobic Degradation of Plant Saccharides

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Bacterial microcompartments (BMCs) are organelles that encapsulate functionally linked enzymes within a proteinaceous shell. The prototypical example is the carboxysome, which functions in carbon fixation in cyanobacteria and some chemoautotrophs. It is increasingly apparent that diverse heterotrophic bacteria contain BMCs that are involved in catabolic reactions, and many of the BMCs are predicted to have novel functions. However, most of these putative organelles have not been experimentally characterized. In this study, we sought to discover the function of a conserved BMC gene cluster encoded in the majority of the sequenced planctomycete genomes. This BMC is especially notable for its relatively simple genetic composition, its remote phylogenetic position relative to characterized BMCs, and its apparent exclusivity to the enigmatic *Verrucomicrobia* and *Planctomycetes*. Members of the phylum *Planctomycetes* are known for their morphological dissimilarity to the rest of the bacterial domain: internal membranes, reproduction by budding, and lack of peptidoglycan. As a result, they are ripe for many discoveries, but currently the tools for genetic studies are very limited. We expanded the genetic toolbox for the planctomycetes and generated directed gene knockouts of BMC-related genes in *Planctomyces limnophilus*. A metabolic activity screen revealed that BMC gene products are involved in the degradation of a number of plant and algal cell wall sugars. Among these sugars, we confirmed that BMCs are formed and required for growth on L-fucose and L-rhamnose. Our results shed light on the functional diversity of BMCs as well as their ecological role in the planctomycetes, which are commonly associated with algae.

Bacterial microcompartments (BMCs) are organelles bounded by a proteinaceous shell. They were first observed in electron micrographs of cyanobacteria (1) and later in some chemoautotrophs (2). Subsequent characterization showed they contain ribulose biphosphate carboxylase/oxygenase (RuBisCO) and were required for carbon fixation at atmospheric concentrations of CO₂, and they were dubbed carboxysomes (3). Apparently icosahedral, carboxysome shells are formed by three types of proteins. BMC-H proteins contain a single copy of the Pfam00936 domain and form hexamers (4), while BMC-T proteins are a fusion of two Pfam00936 domains and form trimers (pseudo-hexamers) (5, 6). BMC-H and BMC-T proteins are thought to compose the facets of the icosahedron, while BMC-P proteins, composed of a single Pfam03319 domain, form pentamers that cap the vertices (7, 8). Several years after the identification of the proteins that make up the carboxysome shell, operons involved in 1,2-propanediol (PDU) and ethanolamine (EUT) degradation in *Salmonella enterica* were sequenced, and it was discovered that these loci contain homologs to genes encoding carboxysomal BMC-H, BMC-T, and BMC-P proteins (9–12). The formation of polyhedral bodies in *S. enterica* was subsequently confirmed by electron microscopy and shown to be regulated by the availability of 1,2-propanediol or ethanolamine (12, 13). These catabolic BMCs are referred to as metabolosomes, in contrast to the carbon-fixing carboxysomes.

Several lines of evidence suggest that at least one common function of carboxysomes and metabolosomes is to concentrate a volatile or toxic metabolite (14–17). The carboxysome shell acts as a CO₂ diffusion barrier, increasing the local concentration of RuBisCO substrate and thus improving carbon fixation (17). The PDU metabolosome houses enzymes that form and degrade propanaldehyde, a toxic metabolite (15). Similarly, the EUT metabolosome encapsulates enzymes that generate and degrade acetalde-

hyde, a volatile and reactive metabolite (14). It has been proposed that substrate channeling due to the close packing of these functionally sequential enzymes prevents the buildup of toxic and/or volatile metabolites (13).

The PDU and EUT metabolosomes likewise appear to share common core biochemistry. In both, an aldehyde is generated that is then oxidized by an aldehyde dehydrogenase to form an acyl coenzyme A (CoA) species, reducing NAD⁺ to NADH in the process (18, 19). Because this reaction is isolated from the cytosolic substrate pool, the NAD⁺ and CoA need to be regenerated within the metabolosome. In order to regenerate NAD⁺, an alcohol dehydrogenase reduces a second aldehyde to an alcohol, which exits the BMC (20). To regenerate CoA, a phosphotransacylase replaces the CoA moiety with a phosphate (21). The acyl phosphate then presumably participates in substrate level phosphorylation via an acyl kinase also encoded within the BMC locus (22).

Bioinformatic surveys of sequenced bacterial genomes have revealed that roughly 20% contain gene clusters with the potential to form BMCs, for which only speculative functions have been assigned, based on gene annotations (23–25). One of these loci has recently been investigated further in *Clostridium phytofermentans*

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TABLE 1 Strains used in this study

Strain name	Genotype	Term used in text	Reference or source
Mü 290 ^T	Wild type	Wild type	31
OE3	$\Delta plim_{1104}::npt$	$\Delta plim_{1104}$, control strain	This work
OE4	$\Delta plim_{1751}::IN(npt)$	$\Delta pvmJ::IN(npt)$	This work
OE22	$\Delta lim_{1751}::npt$	$\Delta pvmJ$	This work
OE23	$\Delta(plim_{1756-plim_{1755}})::npt$	$\Delta pvmDE$, shell-less mutant	This work
OE24	$\Delta plim_{1747}::npt$	$\Delta pvmN$	This work

and has been shown to utilize a glycol radical enzyme (GRE) to degrade 1,2-propanediol (26); this is in contrast to the PDU metabolosome of *S. enterica*, which uses a vitamin B₁₂-dependent enzyme. The recent *C. phytofermentans* study marked the first reverse-genetics approach to confirm the presence of a bioinformatically predicted BMC, and surprisingly, it revealed an alternative mechanism with which to degrade 1,2-propanediol. However, the majority of bioinformatically identified BMC loci remain with tentative or no functional prediction whatsoever.

We identified one such locus conserved in the majority of sequenced planctomycete genomes and in one clade of the phylum *Verrucomicrobia*. Depending on the species, the locus contains between 11 and 13 genes, which we refer to as *Planctomycetes* and *Verrucomicrobia* metabolosome genes A through O (*pvmA* through *pvmO*) (see Table S1 in the supplemental material). The phyla *Planctomycetes* and *Verrucomicrobia* are closely related; they are united by their morphological similarity, in particular intracellular membrane organization (27, 28), and were among the first bacteria to challenge the view that bacteria are simple organisms lacking organizational complexity. The planctomycetes form three distinct clades: autotrophic species that contain the anammoxosome (reviewed by van Niftrik and Jetten [29]); the class *Phycisphaerae*, which are facultative aerobes and divide by binary fission (30); and the class *Planctomycetia*, members of which are predominantly aerobic or facultatively aerobic, lack anammoxosomes, and divide by budding. Nine members from the last clade representing seven genera isolated from very different habitats have sequenced genomes; eight contain the BMC gene cluster. This locus is apparently restricted to the *Planctomycetes* and *Verrucomicrobia*, and its function provides clues to the ecological role of the ubiquitous planctomycetes, as well as the diversity of BMCs.

MATERIALS AND METHODS

Strains, media, and culture growth conditions. The strains used in this study are described in Table 1. *P. limnophilus* Mü 290^T (31) was grown in YVT, a modified PYGV medium (DSMZ medium 621 [http://www.dsmz.de]): 0.1% yeast extract, 10 mM Tris-HCl (pH 7.5), vitamin solution, and salt solution. YVT medium was supplemented with a 10 mM concentration of an additional carbon source as needed. Agar (1.5%) was added to make solid medium. Liquid cultures for all experiments were grown aerobically in a volume of medium that was one-fifth the volume of the flask: generally 50 ml of medium in a 250-ml Erlenmeyer flask. Flasks were grown in the dark at 30°C with shaking at 150 rpm. L-Fucose and L-rhamnose were purchased from Sigma-Aldrich, and *Laminaria* fucoidan was purchased from Glycomix.

Genetic transformation of *P. limnophilus*. The kanamycin resistance gene, including 137 bp upstream of the ATG start codon, was amplified from plasmid pUH-37 via PCR using primers OE53 (TATAGAATTCAG ATCTCGGAATTGCCAGCT) and OE54 (TATAGGATCCTCAGAAGA

ACTCGTC). EcoRI and BglII restriction sites were added to the 5' end, a BamHI restriction site was added to the 3' end, and the gene was cloned into the p2741 cloning vector. This vector was chosen for its small size, paucity of restriction enzyme sites, and high copy number; these features make it amenable to cloning of multipart constructs.

For homologous recombination, genomic DNA sequences 500 to 1,000 bp upstream and downstream of the gene(s) to be replaced with the kanamycin resistance cassette were amplified via PCR with EcoRI and BglII restriction sites on the 5' end and a BamHI restriction site on the 3' end (primer information is in Table S2 in the supplemental material). Knockout cassettes were constructed in the p2741 vector using the Bgl-Bricks cloning strategy (32). Circular plasmids were used for electroporation. Linearized plasmids and PCR fragments were also tested and could also transform *P. limnophilus*. Aliquots (100 μ l) of *P. limnophilus* competent cells were prepared and electroporated as described previously (33). Electroporated cells were recovered in 1 ml of liquid YVT supplemented with D-glucose for 1.5 h with shaking at 30°C, and all cells were plated onto YVT plates supplemented with D-glucose and 30 μ g/ml of kanamycin for the selection. Plates were incubated at 30°C in the dark until colonies formed, after 5 to 7 days. Colonies were struck onto fresh selection plates, grown for an additional 3 to 5 days, and then genotyped by colony PCR or PCR using purified genomic DNA as the template.

Phylogenetic analysis. For the 16S rRNA species tree, full-length 16S rRNA sequences were obtained from IMG (http://img.jgi.doe.gov) and NCBI (http://www.ncbi.nlm.nih.gov), aligned using MUSCLE (34, 35), and curated with GBLOCKS (36), and phylogeny was constructed using PhyML 3.0 (37) with 500 bootstraps using the Phylogeny.fr web interface (38, 39). For the gene concatenation tree, orthologs of *pduL* (see gene tree in Fig. S1A in the supplemental material) in *S. enterica* were concatenated to orthologs of *pduP* (gene tree in Fig. S1B) present in a list of 67 genomes that contain a BMC gene cluster (see Table S3 in the supplemental material), sequences were aligned using MUSCLE as described above, and gapped positions were manually trimmed using Jalview (40). The phylogeny was constructed using PhyML as described above, with 100 bootstraps. The gene clusters were selected with the goal of maximizing the number of phyla represented. Metabolosome type was assigned based on the identity of the defining enzyme of the gene cluster: propanediol dehydratase (PDU) or ethanolamine ammonia lyase (EUT). Both the PDU2 and glycol radical enzyme (GRE) clades contain a GRE homolog; the gene concatenations that branched close to the PDU clade were defined as PDU2, and those that branched close to the EUT clade were designated GRE. The presence of an aldolase and the absence of any of the aforementioned signature enzymes are the defining features of the *Planctomycetes* and *Verrucomicrobia* (PV) and PV-like BMC gene clusters. The PV-like gene clusters have the same features (gene annotations) as the PV type, but the gene order is different.

Gene comparisons. Pairwise BLAST comparisons were performed using the IMG genome BLASTP utility with a minimum E value of 1e-2 and minimum identity of 30% as criteria for a positive hit, and orthologs were defined as those genes that returned bidirectional BLAST hits by the IMG server (see above).

Biolog assay. Cells were grown for 5 days on solid YVT medium supplemented with glucose, then scraped off using a sterile cotton swab, and resuspended in IF0a medium, obtained from Biolog, to an optical density at 600 nm (OD₆₀₀) of 0.24 as measured in by a NanoDrop 2000c in the cuvette holder. Although the cells were not washed in fresh medium to control for nutrient transfer when resuspended, the A1 well of each Biolog plate contains no carbon source and was used as a blank when measuring dye reduction, to serve as a control. Cell suspension (1.76 ml) was added to 22.24 ml of inoculating fluid (YVT with 0.005% yeast extract instead of 0.1% and 240 μ l of dye G, obtained from Biolog). Phenotypic microarray plates 1 and 2 were inoculated with 100 μ l of final cell suspension, wrapped in aluminum foil to retard evaporation, and incubated at 30°C for 5 days. The absorbance at 590 nm was recorded on a BioTek EON microplate spectrophotometer.

Growth curves. *P. limnophilus* strains were grown on plates for 5 days and then resuspended into YVT medium using a sterile cotton swab to scrape cells from the plate. Cultures were resuspended in liquid YVT to an OD₆₀₀ of about 0.5, as measured by a BioTek EON microplate spectrophotometer, and inoculated into liquid medium for growth curves to a starting OD₆₀₀ of 0.01 ± 0.02. Cultures were grown as described above.

Electron microscopy. Cultures were grown as described above (YVT medium supplemented with a 10 mM concentration of an additional carbon source) for 9 days prior to sample preparation for electron microscopy. Cells were concentrated by centrifugation and loaded into 50-μm-deep specimen carriers (catalog no. 390; Wohlwend Engineering, Sennwald, Switzerland) and ultrarapidly frozen in a Bal-Tec HPM 010 high-pressure freezer (Bal-Tec Ag, Liechtenstein). Frozen samples were freeze substituted over a period of 2 h using the superquick-freeze substitution (SQFS) method of McDonald and Webb (41). Cells were rinsed in pure acetone, infiltrated with Epon resin (25, 50, 75, and 100% [the last value for three times]) for 15 min at each concentration, and then polymerized for 2 days in a 60°C oven. Sections 60 nm thick were cut and poststained for 4 min in 2% aqueous uranyl acetate and 2 min in lead citrate (42). Images were taken with a Gatan Ultrascan 1000 camera (Gatan, Inc., Pleasanton, CA) bottom mounted on a Tecnai Spirit transmission electron microscope with an accelerating voltage of 120 kV (FEI, Hillsboro, OR). All original electron micrographs were adjusted in the same manner using Adobe Photoshop CS6 as follows: brightness was decreased 20%, and contrast was increased 90%.

RESULTS

A conserved BMC gene cluster is present in members of the phyla Planctomycetes and Verrucomicrobia. The *Planctomycetes* and *Verrucomicrobia*-type (PV) BMC gene cluster is defined by the presence of two (or three in *Singulisphaera acidiphila* and *Isosphaera pallida*) genes encoding BMC-H proteins and three genes encoding BMC-P proteins (Fig. 1A). In addition to the shell proteins, five genes code for putative enzymes, and the cluster is preceded by a transcriptional regulator, suggestive of operon-like regulation (Fig. 1A). Interestingly, the gene clusters present in the *Verrucomicrobia* either do not have this transcriptional regulator or have it in the opposite orientation relative to the rest of the gene cluster; this may reflect a different type of regulation between the two phyla.

Based on the species tree, it appears as though the PV gene cluster was present in the last common ancestor of the classes *Phycisphaerae* and *Planctomycetia*, with *Gemmata obscuriglobus* having lost the gene cluster (Fig. 1A). A similar situation is apparent in the *Verrucomicrobia*, where *Coralimargarita akajimensis* apparently lost the gene cluster. The bootstrap values for the nodes at which *G. obscuriglobus* bifurcates within the class *Planctomycetia* are very low, and this could be because the “true” position of *G. obscuriglobus* is most likely basally located within the class, as has been shown in other, more phylogenetically focused studies (43–45). As a result, we cannot definitively determine whether the *G. obscuriglobus* genome lost the BMC locus or never contained it. However, in either case the species tree is consistent with vertical inheritance of the gene cluster within the respective phyla, as opposed to horizontal gene transfer, highlighting the retention of the loci over evolutionary timescales. Conservation of genes in a specific order across species typically suggests that they function together in a given biological phenomenon (46); accordingly, we hypothesized that the PV gene cluster encodes a bona fide BMC.

Bioinformatic comparison of PV BMC genes to other metabolosome genes. We focused on the gene complement in *P. limnophilus* as the representative BMC-associated gene cluster be-

cause this species is rapidly becoming a model organism with which to study planctomycete biology (33, 47). We compared its composition to that of the EUT and vitamin B₁₂-dependent PDU BMC loci found in *S. enterica* and to the glycol radical enzyme utilizing the PDU (hereafter referred to as PDU2) BMC locus found in *C. phytofermentans* (Table 2). While the PDU, PDU2, and EUT loci encode at least one BMC-T protein, *P. limnophilus* does not have any, nor does any planctomycete or verrucomicrobium. Furthermore, the PV locus contains three genes encoding BMC-P proteins, while the EUT and PDU BMC loci each have only one. In addition to the differences in shell protein composition, it also appears that all four BMCs use distinct enzymes to generate an aldehyde (Table 2). Downstream of aldehyde generation, all four types of BMC loci are similar, encoding enzymes required to participate in the core BMC biochemistry (Table 2). Furthermore, all of the aldehyde-generating enzymes as well as the aldehyde dehydrogenases encoded in all of the loci contain a BMC encapsulation peptide (23) not present in non-BMC-associated homologs. Thus, it appears that an aldehyde can be generated and partially catabolized in the putative PV BMC, as in the PDU, EUT, and PDU2 BMCs.

The aldolase encoded in the PDU2 locus has been reported to generate lactaldehyde from L-fucose and L-rhamnose (26) and is homologous to the aldolase encoded in the PV locus, suggesting a similar function in these BMCs. While pairwise sequence comparisons (see Fig. S2 in the supplemental material) supported a possible common function in the BMCs, a few key differences are apparent. First, the propanol dehydrogenase encoded in the PDU2 BMC gene cluster (*Cphy_1179*) does not have any bidirectional hits to genes in the *P. limnophilus* genome, indicating that no ortholog is present (48) (see Fig. S2). Second, the gene order within the PV BMC gene cluster is markedly different from that in other BMC gene clusters; such dissimilarity has been proposed to be indicative of operon evolution associated with diverging function (49). Finally and most critically, both the PV BMC gene cluster and the *P. limnophilus* genome lack any homologs to the propanediol dehydratase (forming propionaldehyde from 1,2-propanediol) genes found in the PDU and PDU2 loci (see Fig. S2), so it is unlikely that propanediol is processed by this BMC. Given these observations, we posited that the PV BMC is involved in the degradation of L-fucose, L-rhamnose, and/or related sugars via a pathway different from that of *C. phytofermentans*.

Phylogenetic analysis of PV-type BMC genes. Only two protein domain families are conserved across all BMC types found in microbial genomes: Pfam00936 (in BMC-H and BMC-T) and Pfam03319 (BMC-P). Because these domains are small (approximately 100 amino acids) and are present in different copy numbers in different BMC gene clusters, sequence alignments result in low phylogenetic signal. However, for catabolic BMCs, the three enzymes involved in the core biochemistry are common to all (Table 2). Accordingly, we searched for homologs of these enzymes in sequenced microbial genomes and constructed a phylogeny using a concatenation of the aldehyde dehydrogenase and phosphotransacylase genes from 67 diverse genomes (see Table S3 in the supplemental material). Because some gene clusters encoded multiple putative alcohol dehydrogenases, that core enzyme was not included in the concatenation.

In the resulting gene tree, a number of distinct clades were apparent (Fig. 1B). BMCs containing glycol radical enzymes (GREs) either grouped near those involved in metabolism of pro-

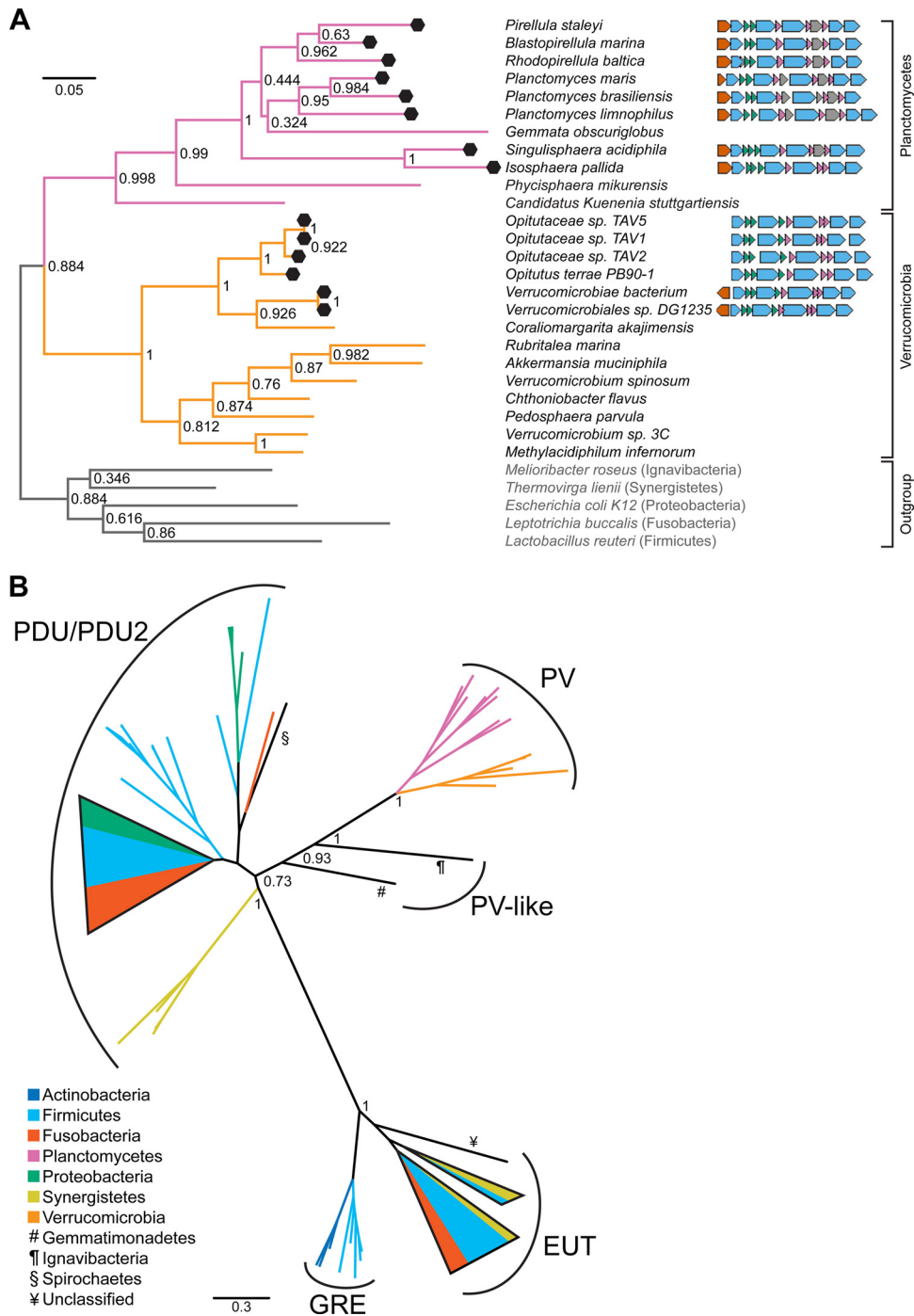


FIG 1 Phylogenetic trees. (A) 16S rRNA tree of representative species of the phyla *Planctomycetes* and *Verrucomicrobia* with sequenced genomes (as of June 2013) and five outgroup species. Bootstrap values are shown at nodes, and black hexagons indicate species with a BMC gene cluster. Each species' BMC gene cluster is shown schematically to the right of the species name, color-coded by gene annotation as follows: orange, transcriptional regulator; blue, enzyme; green, BMC-H shell protein; pink, BMC-P shell protein; gray, hypothetical protein. (B) Gene tree of the concatenation of aldehyde dehydrogenase (*pduP*) and phosphotransferase (*pduL*) homologs (Table 2; see also Table S3 in the supplemental material). Branches with less than 50% bootstrap support were collapsed and drawn in cartoon form. Branch color or symbol corresponds to phylum. Collapsed branches were similarly colored to represent phyla present within. Scale bar represents number of substitutions per site. Clades are annotated by the key enzymes present in the gene clusters (see Materials and Methods). PDU, vitamin B₁₂-dependent propanediol utilization; PDU2, vitamin B₁₂-independent (glycyl radical enzyme-utilizing) propanediol utilization; PV, *Planctomycetes* and *Verrucomicrobia* type; EUT, vitamin B₁₂-dependent ethanolamine utilization; GRE, locus of unknown function containing a glycyl radical enzyme. Bootstrap values of branches separating major clades are shown.

panediol (PDU2) or formed a distinct clade close to representative sequences of the EUT metabolosome (Fig. 1B, GRE). In addition to the PDU/PDU2, GRE, and EUT groups, PV enzymes form a well-defined clade (Fig. 1B). The branch length separating the PV-type clade from the PDU/PDU2 clade is comparable to that separating the EUT and PDU/PDU2 clades, suggesting that the enzymes within the PV-type BMC are functionally divergent from propanediol metabolism. Within the PV clade, the tree topology is essentially congruent with the species tree (data not shown), supporting our hypothesis that the BMC-associated genes have descended vertically through the *Planctomycetes* and *Verrucomicrobia*. In contrast, the other major clades are paraphyletic, indicating horizontal gene transfer (Fig. 1B). Due to the clear distinction of the PV lineage, we inferred that the PV BMC has a novel function distinct from any other characterized to date.

The PV BMC appears to be involved in several saccharide degradation pathways. In order to experimentally identify the substrate of the PV BMC and the related biochemical reactions that take place, we conducted an aerobic metabolic screen using Biolog phenotype microarrays (Biolog, Hayward, CA). In this assay, 96-well plates have prospective substrates dried in each well and are inoculated with a cell suspension containing a redox-sensitive dye. Reducing equivalents are generated when a sample has metabolic activity in the well; these reduce the dye, turning it purple. Given that all BMCs characterized to date are involved in carbon metabolism, we selected phenotype microarrays containing a library of carbon sources. One of the presumed core PV BMC enzymes, encoded by *pvmJ*, is a NAD⁺-dependent dehydrogenase, which should produce NADH when the BMC substrate is degraded. We predicted that a mutant of *P. limnophilus* deficient in this gene would display little to no metabolism-associated signal, and comparison with wild-type growth would be an effective method to rapidly generate a list of candidate functions for the BMC. *pvmJ* was deleted using homologous recombination by replacing the gene with the *npt* kanamycin resistance gene under the control of its own promoter, in an orientation opposite to that of the rest of the gene cluster (Fig. 2A). This marks the first instance a directed gene knockout has been made in the phylum *Planctomycetes*; all other studies have employed random transposon insertion (33, 47). We confirmed that the selectable marker replaced *pvmJ* by PCR genotyping (Fig. 2B). All tested clones were double recombinants, not requiring a secondary selection step, highlighting the potential of reverse genetics to study planctomycetal biology. In order to control for any adverse metabolic effects that expressing the *npt* gene may have, we also generated a control strain, with the *npt* gene replacing *plim_1104*, a pseudogene in a relatively barren region of the chromosome (see Fig. S3A and B in the supplemental material). Thus, we have also identified a neutral site in the *P. limnophilus* genome in which to introduce genetic constructs for heterologous expression for other types of genetic studies.

Surprisingly, the aldehyde dehydrogenase $\Delta pvmJ$ mutant displayed a metabolic deficiency under several conditions, as opposed to one or two (see Fig. S4 in the supplemental material). There were also some compounds with which the mutant appeared to produce a stronger signal than the control strain, but we did not expect these compounds to readily inform about the function of the BMC and so they were not further investigated. The substrates that displayed the strongest metabolic deficiency were pectin, mannose-containing compounds (D-mannose, mannan,

TABLE 2 Protein compositions of functionally distinct metabolosomes

BMC type	No. and name of shell protein gene(s)			Gene(s) for aldehyde-generating enzymes				Gene for core metabolosome enzymes			
	BMC-H	BMC-T	BMC-P	1,2-Propanediol oxidoreductase	Aldolase	Propionaldehyde generating diol dehydratase	Acetaldehyde generating	Aldehyde dehydrogenase	Phosphotransferase	Acyl kinase	Alcohol dehydrogenase
PDU (<i>S. enterica</i>)	3; <i>pdhA</i> , - <i>J</i> , - <i>K</i> , - <i>U</i>	2; <i>pdhB</i> , - <i>T</i>	1; <i>pdhN</i>	None	None	<i>pdhCD²</i> (B ₁₂ -dependent diol dehydratase)	None	<i>pdhP²</i>	<i>pdhU²</i>	<i>pdhW²</i>	<i>pdhQ²</i>
EUT (<i>S. enterica</i>)	3; <i>eutK</i> , - <i>M</i> , - <i>S</i>	1; <i>eutL</i>	1; <i>eutN</i>	None	None	None	<i>eutBC²</i> (ethanolamine ammonia lyase)	<i>eutE²</i>	<i>eutD²</i>	<i>ack²</i>	<i>eutG²</i>
PDU2 (<i>C. phytofermentans</i>)	4; <i>Cphy_1176</i> , <i>Cphy_1180</i> , <i>Cphy_1181</i> , <i>Cphy_1182</i>	1; <i>Cphy_1186</i>	1; <i>Cphy_1184</i>	<i>Cphy_1185</i>	<i>Cphy_1177</i>	<i>Cphy_1174</i> (B ₁₂ -independent diol dehydratase)	None	<i>Cphy_1178</i>	<i>Cphy_1183</i>	<i>Cphy_1327</i>	<i>Cphy_1179</i>
PV (<i>P. limnophilus</i>)	2; <i>pvmD</i> , - <i>E</i>	0	3; <i>pvmH</i> , - <i>K</i> , - <i>M</i>	None	<i>pvmN</i>	None	None	<i>pvmJ</i>	<i>pvmB</i>	<i>pvmG</i>	<i>pvmO</i>

^a Experimentally verified function. Accession numbers for genes are provided in Tables S1 and S4 in the supplemental material.

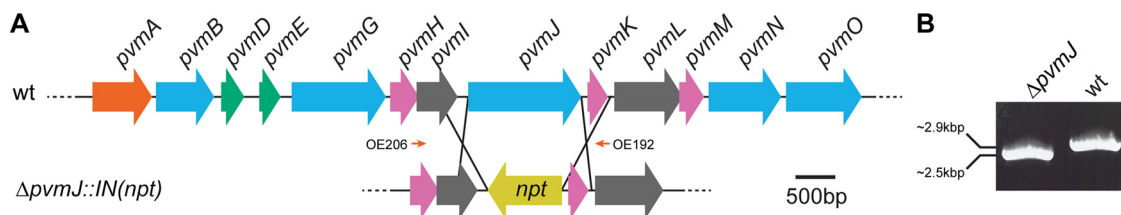


FIG 2 Homologous recombination to generate knockout mutants in *P. limnophilus*. (A) Schematic of the BMC gene cluster in *P. limnophilus* and the homologous recombination used to create the $\Delta pvmJ::IN(npt)$ mutant. Orange, transcriptional regulator; blue, enzyme; green, BMC-H shell protein; pink, BMC-P shell protein; gray, hypothetical gene; yellow, kanamycin resistance gene (*npt*). Orange arrows represent primers OE206 and OE192 (see Table S2 in the supplemental material) used for PCR genotyping. OE206 is outside the homology region, showing that recombination happened at the desired locus. (B) PCR amplification using primers OE206 and OE192 showing that amplicons are different sizes for the wild-type (wt) and knockout strains.

α -methyl-D-mannoside), turanose-containing compounds (turanose and melezitose), D-raffinose, and L-fucose (see Fig. S4). For all of these sugars, the mutant still displayed a basal level of metabolic signal except L-fucose, which was completely devoid of metabolic signal (see Fig. S5 in the supplemental material). Furthermore, we observed only minimal metabolic signal in the well containing propanediol (PM1 well D5) (see Fig. S5), as well as no significant difference in signal between the mutant and control strain with this substrate. Thus, the PV BMC aldehyde dehydrogenase appears to be involved in the catabolism of a number of carbon sources and is absolutely necessary for the degradation of L-fucose and L-rhamnose, which appears to proceed via a pathway independent of propanediol.

PV-type BMC genes are required to degrade L-fucose and L-rhamnose. In order to focus on the physiological roles of the putative aldehyde dehydrogenase (encoded by *pvmJ*) and the putative aldolase (encoded by *pvmN*), we constructed deletion mutants designed specifically to avoid any polarity effects (see Fig. S3C to F in the supplemental material). We confirmed that these mutations produced no transcript of the deleted gene and did not significantly disrupt the expression of downstream genes when cells were grown on L-fucose (data not shown).

We investigated the growth of the $\Delta pvmJ$ and $\Delta pvmN$ strains in the presence or absence of L-fucose and the structurally similar deoxy sugar L-rhamnose. Due to the slow doubling time of *P. limnophilus* (about 20 h on D-glucose; data not shown), it was not feasible to conduct the growth experiments with either L-fucose or L-rhamnose as the sole carbon source. Instead, we continued to supplement the medium with 0.1% yeast extract to reach suitable growth rates. The growth profiles of the wild type and the $\Delta plim_{1104}$ control strain were comparable, allowing us to control for any metabolic stress associated with the expression of the *npt* gene product (see Fig. S6 in the supplemental material).

The $\Delta pvmJ$ mutant had a detectable growth defect in the presence of L-fucose and a slight defect for L-rhamnose, but it had a growth rate similar to that of the control strain on yeast extract alone, indicating that this aldehyde dehydrogenase mutant could not utilize those sugars for energy but could grow on nutrients supplied by the yeast extract (Fig. 3A and B). Interestingly, the $\Delta pvmN$ (aldolase mutant) strain did not grow at all in the presence of either L-fucose or L-rhamnose (Fig. 3A and B). This is suggestive of a toxic effect, since nutrition from the yeast extract is present. We verified this by spiking D-glucose-grown $\Delta pvmN$ cultures with L-fucose when they reached early exponential growth. The cultures that were challenged with L-fucose displayed a dramatically reduced growth rate, and the cell density after 6 days of growth was

substantially lower than that of the control (Fig. 3C). These observations are consistent with an accumulation of an inhibitory factor when the $\Delta pvmN$ strain is grown on L-fucose or L-rhamnose. We further suggest that the *pvmN* gene product serves to detoxify this inhibitory factor. In addition, it appears that this gene cluster is not catabolically repressed by the presence of D-glucose. Collectively, the data indicate that the putative aldolase and aldehyde dehydrogenase of the PV BMC gene cluster are involved in the catabolic pathway of L-fucose and L-rhamnose and that this pathway produces toxic intermediates.

Planctomycetes assemble a metabolosome to degrade L-fucose and L-rhamnose. Since metabolosomes have been proposed to sequester reactions that produce toxic intermediates (15), we examined whether the formation of a BMC (composed of a shell and encapsulated enzymes) was involved in the metabolism of L-fucose and L-rhamnose. In order to confirm a role for the BMC shell proteins, we deleted *pvmD* and *pvmE*, the only two BMC-H shell protein genes present in the gene cluster (Table 2; see also Fig. S3G and H in the supplemental material); this deletion was expected to abolish metabolosome shell formation or integrity. In the presence of L-rhamnose, the $\Delta pvmDE$ mutant grew slower than the control, mimicking cells grown on yeast extract alone. However, when grown in the presence of L-fucose, the mutant exhibited growth inhibition similar to that of the $\Delta pvmN$ mutant (the putative aldolase mutant) and grew roughly 1 order of magnitude slower than the control strain (Fig. 4A). This is in contrast to previous studies on the EUT and PDU metabolosome in which individual shell proteins were deleted and relatively minor growth defects were observed (14, 50). These data suggest that the shell proteins are required for the optimal metabolism of both L-fucose and L-rhamnose.

To confirm that a BMC is formed, we visualized wild-type *P. limnophilus* cells grown on L-fucose, L-rhamnose, or D-glucose by electron microscopy. It has been reported that BMCs are most apparent in stationary-phase cultures of *S. enterica* (12, 13) and *C. phytofermentans* (26), so we grew *P. limnophilus* to stationary phase before preparing thin sections. BMC-like structures were present in L-fucose- and L-rhamnose-grown cell populations but not in D-glucose-grown cultures (Fig. 4B, C, and D). These structures are 55 ± 9 nm ($n = 47$) in diameter and resemble in size and morphology the BMCs observed in *S. enterica* (51) and *C. phytofermentans* (26). Although amorphous bodies are present in D-glucose-grown cells, they do not have the distinct edges seen for the BMCs and do not otherwise resemble the structures of metabolosomes (Fig. 4D). Collectively, these data strongly suggest

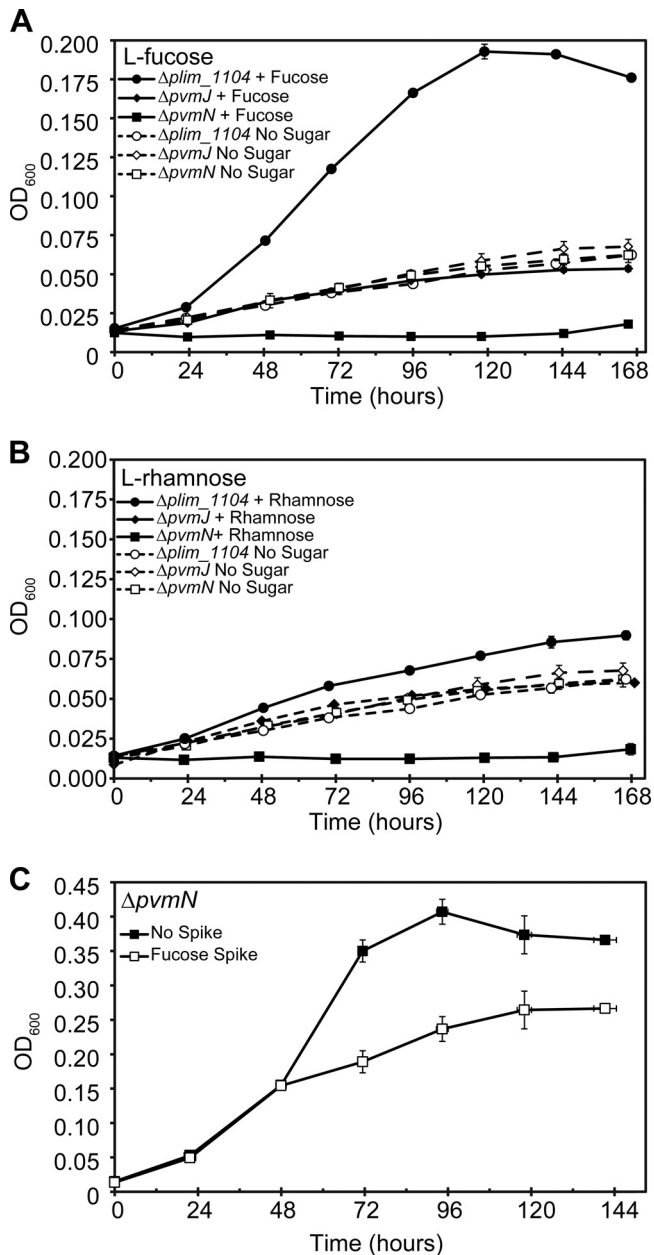


FIG 3 Growth curves of $\Delta plim_{1104}$, $\Delta pvmJ$, and $\Delta pvmN$ strains grown on L-fucose (A) and L-rhamnose (B). "No sugar" data in panel B are reproduced from panel A. (C) Spiking experiment in which the $\Delta pvmN$ strain was grown on 5 mM D-glucose for 2 days, at which point 5 mM L-fucose was added to half of the cultures. Solid squares represent nonspiked cultures, and open squares represent cultures with L-fucose added. In all graphs, each data point is an average of three independently grown cultures ($n = 3$); each error bar represents 1 standard deviation.

that a bona fide BMC is synthesized in *P. limnophilus* in order to degrade fucose and rhamnose.

BMC genes are required for growth on fucoidan, a sulfated polysaccharide from macroalgae. L-Fucose and L-rhamnose rarely occur in nature as monosaccharides; they are usually components of polysaccharides. We next considered possible sources of these saccharides that may be encountered by planctomycetes. Planctomycete genomes are known to encode more sulfatase

genes than those of most organisms; moreover, some planctomycetes have been shown to grow on sulfated polysaccharides (52). The sulfated polysaccharide fucoidan is composed of 95% L-fucose (53), and we hypothesized that *P. limnophilus* could utilize this compound as a source for L-fucose. We conducted a growth assay using the $\Delta plim_{1104}$ control strain, the $\Delta pvmDE$ shell mutant, and the $\Delta pvmN$ aldolase mutant on YVT medium supplemented with fucoidan extract from the *Laminaria* genus of brown algae. While the $\Delta plim_{1104}$ strain exhibited appreciable growth, the shell protein $\Delta pvmDE$ mutant appeared to grow about the same as on yeast extract alone, indicating that the fucoidan could not be metabolized and that no inhibitory factor was accumulating (Fig. 5). The $\Delta pvmN$ aldolase mutant exhibited the same growth suppression seen on L-fucose and L-rhamnose (Fig. 5), suggesting that fucoidan is desulfated and broken down into fucose monomers and then further metabolized to an inhibitory metabolite.

DISCUSSION

We have shown that bacterial microcompartments are formed in *P. limnophilus* and that they are required for aerobic growth on L-fucose, L-rhamnose, and the sulfated polysaccharide fucoidan. In order to study the BMC, we generated the first reported series of directed gene knockouts in a planctomycete, advancing the genetic toolset available for the phylum. Analysis of these mutants suggests that the organelle serves to detoxify products generated in the aerobic degradation pathway(s) of those sugars. In addition, we have observed a BMC mutant to be deficient in the degradation of several other carbon sources present in a metabolic screen.

Based on our bioinformatic and experimental data, and previous genetic characterization of homologs in the EUT, PDU, and clostridial PDU2 metabolosomes, we propose a biochemical model for the function of the PV BMC, with two alternative upstream pathways to generate lactaldehyde from either L-rhamnose or L-fucose: phosphorylative (54) and nonphosphorylative (55–57) (Fig. 6). It is not clear which of these strategies *P. limnophilus* uses, as we could not identify homologs for the full enzyme complement for either pathway in its genome (Fig. 6; see also Fig. S2 in the supplemental material); these activities may be provided via nonorthologous gene displacement (58). The BMC-encoded aldolase is capable of acting on intermediates of both pathways to produce lactaldehyde as well as a compound that can readily enter central metabolism: either dihydroxyacetone phosphate or pyruvate (55, 59–61). Downstream of the aldolase reaction, the remaining four enzymes in the PV BMC gene cluster participate in reactions that degrade lactaldehyde to lactate while allowing for cofactor recycling within the compartment; this is required because the BMC shell prevents the diffusion of NAD^+/H and coenzyme A to the cytosol (20, 21) (Fig. 6). The lactate can then enter glycolysis via pyruvate and be respired to generate several reducing equivalents for cellular energy. The propanediol is produced as a result of cofactor recycling and may be excreted; there is no obvious evidence of *P. limnophilus* being able to metabolize this compound (see Fig. S5 in the supplemental material, PM1 well D4), an observation corroborated by other investigators (62). Due to the PV BMC metabolism being aerobic, the end products of the BMC metabolism (DHAP and/or pyruvate) likely enter the tricarboxylic acid (TCA) cycle and generate ATP through oxidative phosphorylation. We posit that the competitive advantage provided by the PV BMC outweighs the

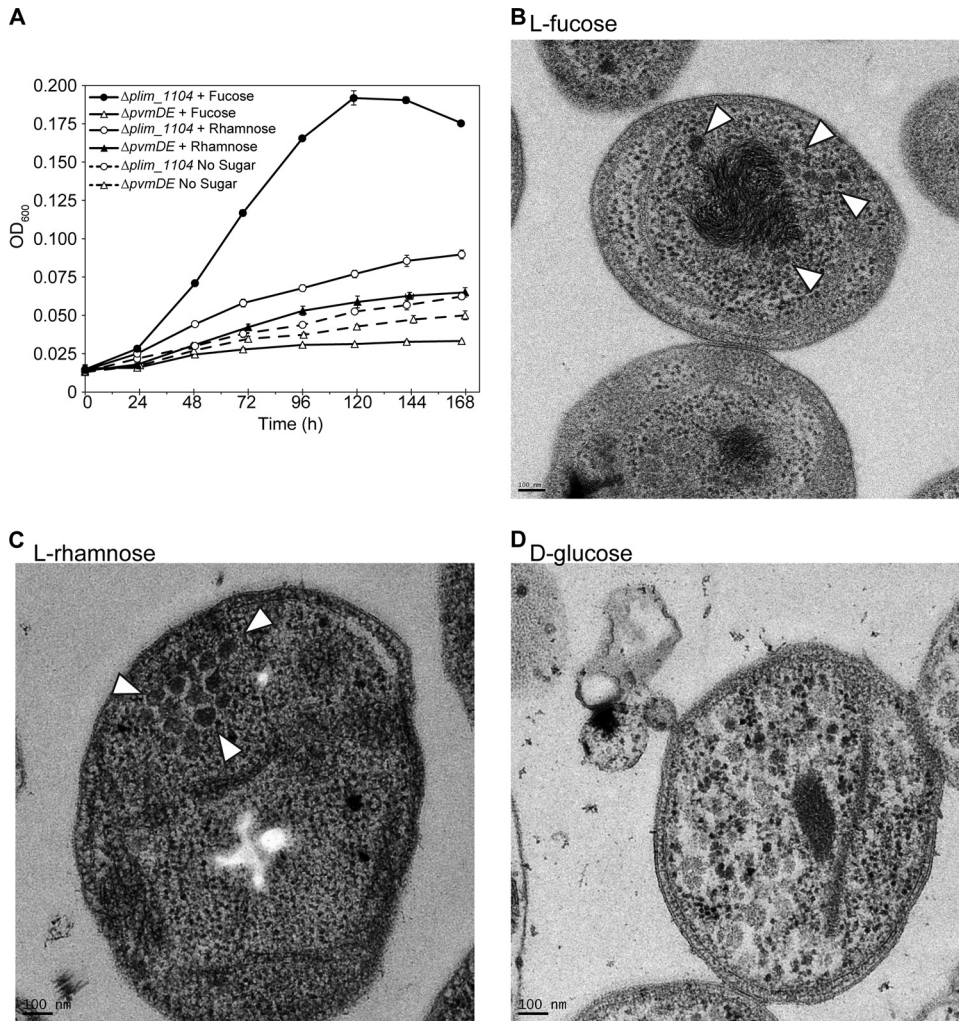


FIG 4 BMCs are expressed during growth on fucose and rhamnose. (A) Growth curves using L-fucose or L-rhamnose of the $\Delta pvmDE$ and $\Delta plim_{1104}$ strains grown with or without sugars. Data for the $\Delta plim_{1104}$ strain are reproduced from Fig. 3A and B. Each data point is an average of three independently grown cultures ($n = 3$); each error bar represents 1 standard deviation. (B to D) Electron microscopy of wild-type *P. limnophilus* showing BMCs (indicated by white arrowheads) when grown for 9 days on L-fucose (B) and L-rhamnose (C) but not D-glucose (D).

potential loss of energy in the form of excreted propanediol. Furthermore, although we grew *P. limnophilus* aerobically only, our model also supports fermentative growth on L-fucose and L-rhamnose, as the pathway would net one NADH or ATP,

depending on if the phosphorylative or nonphosphorylative pathway was being utilized, respectively.

Both *pvmN*, which generates an aldehyde, and *pvmJ*, which degrades the aldehyde, are predicted to encode a BMC encapsulation peptide (23). Similar peptides are found in a subset of enzymes associated with PDU (63, 64), PDU2, and EUT BMCs that are shown or predicted to generate or degrade an aldehyde intermediate (23). These peptides interact with shell proteins (23, 65), although it is not clear whether they associate with the inner or the outer surface of the shell; for example, it has recently been proposed that ethanolamine ammonia lyase associates with the outside the shell and “injects” acetaldehyde into the lumen (21). It is possible that *pvmN* is similarly associated with the outside of the shell, as it presumably would be difficult for its six-carbon substrate to efficiently diffuse across a shell designed to sequester metabolites (25). Although some crystal structures of shell proteins contain large gated pores that could allow larger metabolites across (5, 6, 66), these are thus far confined to BMC-T proteins, which PV BMC loci lack (Table 2). As ADP and ATP are slightly larger than L-fucose, we predict that the final product of the core

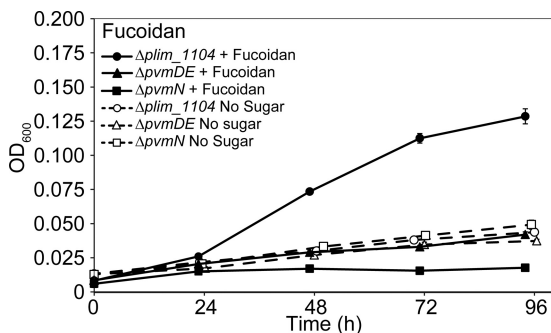


FIG 5 Growth curves of $\Delta plim_{1104}$, $\Delta pvmDE$, and $\Delta pvmN$ strains grown on fucoidan from *Laminaria* spp. “No sugar” data are reproduced from Fig. 3A and 4A. Each data point is an average of three independently grown cultures ($n = 3$); each error bar represents 1 standard deviation.

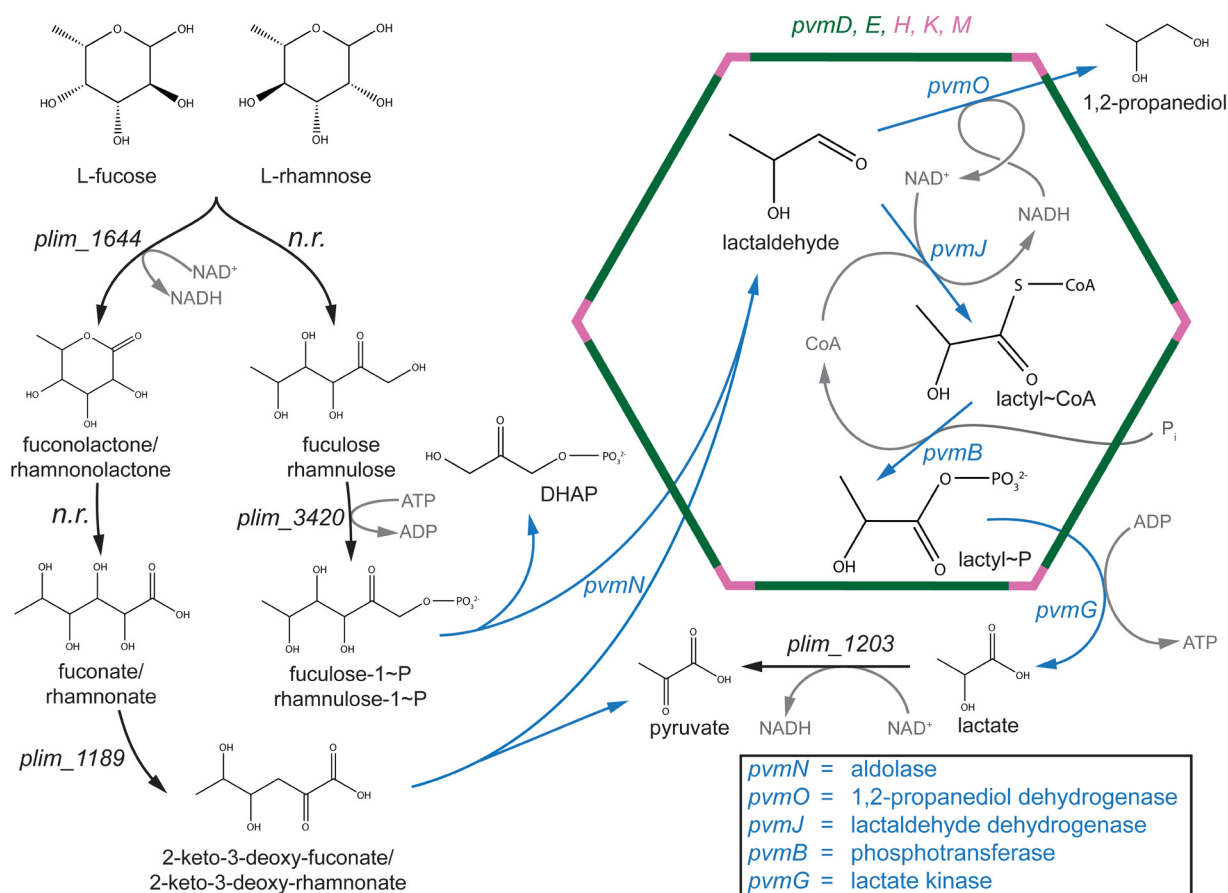


FIG 6 Proposed model of L-fucose metabolism in *P. limnophilus*. Two potential pathways to reach lactaldehyde are shown. Enzymes encoded by the *pvm* gene cluster are shown in blue, and shell proteins are shown in green (BMC-H) or pink (BMC-P). Cofactors are depicted in gray. Enzymes that potentially catalyze reactions not directly involved in the BMC chemistry are shown in black. "n.r." stands for no result, meaning that there was no significant BLAST hit using characterized enzymes as a query. DHAP, dihydroxyacetone phosphate.

BMC chemistry, lactyl phosphate, diffuses out of the compartment and is converted to lactate by the cytosolic product of *pvmG*. It must be noted, however, that the PV BMC shell gene complement is unique and could allow for hetero-oligomeric shell units that permit the larger substrates and cofactors to diffuse across the shell, functionally replacing BMC-T shell units (discussed below).

Phylogeny and evolution of metabolosomes. An accurate phylogeny of the various BMC types has not been previously reported. In this study, we have used a core metabolic gene concatenation strategy to begin to explore the evolutionary history of functionally distinct BMCs (Fig. 1B). Although useful for our purposes, our analysis has limitations, mainly that the sample size is small ($n = 67$; see Table S3 in the supplemental material) and that carboxysome and EUT operons were excluded due to the lack of *pdul* homologs in those loci. Nevertheless, we observed that metabolosomes distinctly group by function, potentially allowing predicted catabolic BMCs to be preliminarily assigned a function based on the phylogenetic position of their core enzymes. Supporting these assignments, two classes of BMCs that use glycol radical enzymes (GREs) emerged, and they are consistent with two different functional assignments: one presumably involved in 1,2-propanediol utilization (26, 67) (PDU2, grouping with PDU loci) and another clade, GRE, closely related to the EUT BMC,

proposed to metabolize choline, a compound structurally similar to ethanolamine (68).

Notably, the PV BMC appears to be restricted to the phyla *Planctomycetes* and *Verrucomicrobia* (Fig. 1). This is a curiosity, as BMC gene clusters appear to be prone to horizontal gene transfer, explaining their widespread presence in bacterial genomes (25, 69, 70). However, there are multiple biochemical strategies used by bacteria to degrade L-fucose and L-rhamnose (55, 57, 61, 71), and the PV BMC would not be expected to provide any further fitness advantage to organisms that could already degrade those compounds; this perhaps explains their restricted distribution. A corollary to these assumptions is that ancient planctomycetes could not fully metabolize L-fucose and L-rhamnose until acquiring a BMC locus via horizontal gene transfer.

Environmental sources of PV BMC substrates. Optimal L-fucose and L-rhamnose metabolism could be very important for planctomycete niche specialization. These sugars are commonly found in the extrapolsaccharides of bacteria and fungi (72), lipids (73), and plant and animal saccharides (53, 74, 75). Planctomycetes have been shown to attach to detritus (marine snow), which is composed of decaying plant and animal materials that most likely contain L-fucose and L-rhamnose, and are thought to be involved in carbon cycling in oceans (76). Furthermore, a recent study has

shown that two planctomycetes, one marine and one freshwater, are capable of degrading several other plant-derived sugars (62). Seaweeds and other marine algae contain an abundance of sulfated polysaccharides (77), two of which are fucoidan and ulvan, which are composed of 95% L-fucose (53) and 55% L-rhamnose (75), respectively. Members of the phylum *Planctomycetes* dominate seaweed-associated biofilms (78), frequently co-occur with algal blooms (79), and contain an abnormal abundance of sulfatase genes that are used to degrade sulfated polysaccharides (52); all of these observations are consistent with the prediction that many *Planctomycetes* species use a BMC in degrading algal polysaccharides.

Sulfated polysaccharides have been associated with a plant response to salinity (80), and almost all identified sulfated polysaccharides are derived from marine algae, so it is logical that marine planctomycetes would encode the PV BMC for use in polysaccharide degradation. In addition to planctomycetes, a marine verrucomicrobium of relatively close relation to *Opitutus terrae*, a species that contains a *pvm* locus, was shown to degrade fucoidan from several algal sources (81). Intriguingly, we have shown that *P. limnophilus*, a freshwater planctomycete, can grow on fucoidan and that BMC components are required for its degradation. This strongly suggests that the organism is exposed to sulfated polysaccharides in its typical niche. Furthermore, several other freshwater planctomycetes also contain the *pvm* locus. It could be that the abundance of sulfated polysaccharides in freshwater systems has been underestimated; at least one freshwater phytoplankton species has been observed to possess a sulfated fucan (82).

Recently, a comprehensive metabolic screen of *P. limnophilus* suggested that the organism is specialized to degrade plant-derived sugars and polysaccharides (62). Our results confirmed these observations and further implicated the involvement of the BMC in degrading some of these compounds. Notably, the $\Delta pvmJ::IN(npt)$ strain displayed reduced metabolism on pectin, mannan, and raffinose. These sugars do not yield lactaldehyde or propionaldehyde in known degradation pathways, but they do produce some compounds, such as aldols and glyceraldehyde, that could be substrates of *pvm* gene products. Thus, assuming that shell formation could be regulated, enzymes within the gene cluster would still be beneficial to planctomycetes as soluble proteins, even when L-fucose- and L-rhamnose-rich compounds are not present. Whether or not the organism uses just a subset of enzymes from the gene cluster or an entire BMC is synthesized to degrade those sugars remains to be investigated.

Diversity of bacterial microcompartments. Compared to other BMC loci, the *pvm* locus appears to be relatively simple, encoding only a transcriptional regulator, enzymes, and BMC-H and BMC-P shell proteins. Other BMC loci encode BMC-T shell proteins and several accessory proteins that are required to form functional organelles (23, 50, 83–86). Although some *pvm* loci contain hypothetical genes that could encode accessory proteins (Fig. 1A), their presence is not conserved across the various *pvm* loci. It will be interesting to determine if these hypothetical genes are required for BMC function and, if so, why some planctomycetes need them while others do not. Also perplexing is why the *pvm* locus contains three genes for BMC-P. BMC-P proteins are minor components of BMC shells: only 60 BMC-P polypeptide chains are required per icosahedral organelle to form the vertices (7, 87), and PDU, PDU2, and EUT operons each contain only one

BMC-P gene. It may be the case that the BMC-P proteins encoded in the *pvm* locus assemble into heteropentamers to cap the vertices or incorporate into the facets of the shell with the BMC-H subunits.

The PV BMC marks the sixth experimentally characterized function for a BMC, joining the carboxysome, PDU, EUT, PDU2, and ethanol-utilizing ETU (88, 89) BMCs. Metabolosome specialization can be correlated with the environmental niche of the organism. For example, *S. enterica* and *E. coli* are enteric bacteria that are generally exposed to anaerobic conditions in the gut. When they are provided with L-fucose as a carbon source, the sugar is anaerobically degraded to 1,2-propanediol, which is then excreted (71). When the *pdu* operon was acquired by an ancestral strain, it most likely provided a substantial fitness advantage; the organism could now utilize the propanediol previously wasted (69). Similarly, *C. phytofermentans* is an anaerobe associated with plant cell wall degradation and would also be expected to ferment L-fucose and L-rhamnose, which are common components of plant cell wall polysaccharides, such as the pectin rhamnogalacturonan I. On the other hand, *Planctomycetes* and *Verrucomicrobia* that contain *pvm* loci are usually associated with aerobic habitats, and some are even strict aerobes and would not need to ferment L-fucose and L-rhamnose but instead degrade them (and potentially other polysaccharides) aerobically through the PV BMC.

With continued sequencing of microbial genomes, the phylogenetic and functional diversity of metabolosomes is increasingly apparent, while the core biochemistry is strikingly conserved. Based on our comparison of the PV metabolosome to other characterized examples, the key difference among metabolosome types is the identity of the enzyme that generates the aldehyde; the biochemical transformations downstream of aldehyde generation are remarkably similar. Coupled with the observation that BMCs tend to be organized in readily transmissible units (i.e., tightly clustered loci), it is possible to envision that new functional types of BMCs could have evolved simply by horizontally transferring a section of a metabolosome locus that encodes core enzymes and shell proteins. All the core enzymes would need to be transferred, as cofactor recycling is necessary for the macromolecular biochemical machine to proceed. Assuming a certain degree of enzymatic promiscuity, the transplanted BMC would degrade short-chain aldehydes (similar to propionaldehyde, lactaldehyde, and acetaldehyde), provide a new metabolic function for the host microbe, and over time be specialized by selective pressures into a new functional type of BMC. This could account for both the apparent biochemical diversity and the pervasiveness of metabolosomes among the bacteria and could have implications for bacterial speciation. Carboxysomes, in contrast, are essentially anabolic BMCs; they do not share the core biochemistry of the catabolic metabolosomes. Likewise, understanding their position in the evolutionary history of BMCs is required to develop a comprehensive theory of BMC evolution and phylogenesis.

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