Comparative Genomics of Syntrophic Branched-Chain Fatty Acid Degrading Bacteria

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The syntrophic degradation of branched-chain fatty acids (BCFAs) such as 2-methylbutyrate and isobutyrate is an essential step in the production of methane from proteins/amino acids in anaerobic ecosystems. While a few syntrophic BCFA-degrading bacteria have been isolated, their metabolic pathways in BCFA and short-chain fatty acid (SCFA) degradation as well as energy conservation systems remain unclear. In an attempt to identify these pathways, we herein performed comparative genomics of three syntrophic bacteria: 2-methylbutyrate-degrading “Syntrophomonas wolfei subsp. methylbutyrica” strain JCM 14075T (=4J5T), isobutyrate-degrading Syntrophothermus lipocalidus strain TGB-C1T, and non-BCFA-metabolizing S. wolfei subsp. wolfei strain Göttingen1. We demonstrated that 4J5 and TGB-C1 both encode multiple genes/gene clusters involved in β-oxidation, as observed in the Göttingen genome, which has multiple copies of genes associated with butyrate degradation. The 4J5 genome possesses phylogenetically distinct β-oxidation genes, which may be involved in 2-methylbutyrate degradation. In addition, these Syntrophomonadaceae strains harbor various hydrogen/formate generation systems (i.e., electron-bifurcating hydrogenase, formate dehydrogenase, and membrane-bound hydrogenase) and energy-conserving electron transport systems, including electron transfer flavoprotein (ETF)-linked acyl-CoA dehydrogenase, ETF-linked iron-sulfur binding reductase, ETF dehydrogenase (FixABCX), and flavin oxidoreductase-heterodisulfide reductase (Flox-Hdr). Unexpectedly, the TGB-C1 genome encodes a nitrogencase complex, which may function as an alternative H2 generation mechanism. These results suggest that the BCFA-degrading syntrophic strains 4J5 and TGB-C1 possess specific β-oxidation-related enzymes for BCFA oxidation as well as appropriate energy conservation systems to perform thermodynamically unfavorable syntrophic metabolism.

Key words: syntroph, branched-chain fatty acid, genomics, energy conservation

Under methanogenic conditions, the degradation of amino acids and proteinaceous materials inevitably generates fatty acids as byproducts (26). Fatty acid-oxidizing bacteria and methanogens are known to form syntrophic interactions in order to accomplish the endergonic oxidation of these fatty acids (9, 19, 26). Although the biochemical pathways and genes involved in the syntrophic degradation of short-chain fatty acids (SCFA; e.g., propionate and butyrate) have already been described, they have not yet been elucidated for branched-chain fatty acids (BCFAs; e.g., isobutyrate, isovalerate, and 2-methylbutyrate) derived from branched-chain amino acids (13, 26). Syntrophic BCFA degradation to acetate and propionate has been observed in isolates and mixed cultures (14, 26, 34, 35, 41). Only three strains of the family Syntrophomonadaceae are currently known to syntrophically degrade 2-methylbutyrate (“Syntrophomonas wolfei subsp. methylbutyrica” strain JCM 14075T (=4J5T) and S. bryantii strain CuCal1) and isobutyrate (Syntrophothermus lipocalidus strain TGB-C1T) (29, 33, 40). These Syntrophomonadaceae species are considered to be important for fatty acid degradation in anaerobic ecosystems, including the sludge digestion process (19), rice paddy fields (12), and the termite gut (42). Furthermore, an uncultivated Syntrophaceae member has been proposed to degrade BCFA syntrophically in a methanogenic bioreactor through metagenomic and metatranscriptomic approaches (21). However, the key catabolic enzymes and energy conservation systems necessary to drive thermodynamically unfavorable BCFA and SCFA degradation remain unclear.

In the present study, the genomes of strains 4J5 (20) and TGB-C1 (4) were investigated in order to identify the metabolic pathways for 2-methylbutyrate and isobutyrate catabolism and energy conservation systems for syntrophic metabolism. A comparative genomic analysis between BCFA- and non-BCFA-degrading syntrophs within the family Syntrophomonadaceae (i.e., S. wolfei subsp. wolfei strain Göttingen1 [30]) provides genomic insights into the degradation of BCFA in methanogenic ecosystems.

Materials and Methods

Genome sequencing and annotation

This study analyzed the “S. wolfei subsp. methylbutyrica” strain JCM 14075T (=4J5T) draft genome (DDBJ/GenBank/EMBL accession: BBQT01000001–BBQT01000092) (20), S. lipocalidus strain TGB-C1T complete genome (CP002048) (4), and S. wolfei strain Göttingen complete genome (CP000448) (30). As reported previously (20), the genomic DNA of strain 4J5 was provided by the RIKEN BRC through the National Bio-Resource Project of MEXT, Japan, and sequenced using the Illumina MiSeq platform (Illumina, San Diego, CA, USA) at FASMAC (Atsugi, Japan). Briefly, we constructed and sequenced a 300-bp paired-end library totaling ca. 2.2 Gb of MiSeq data. Assemblies were performed using SPAdes
version 3.1.1 (2). The strain 4J5 draft genome comprises 89 scaffolds and has an estimated genome size of 3.2 Mbp with an average G+C content of 45.55%. The quality of the genome sequence was evaluated using the Check M version 1.0.5 program with a marker gene set of the class *Clostridia* (23). A total of 2,964 protein coding genes were annotated with Prokka version 1.11 (see Supplemental Information) (28). Basic local alignment search tool (BLAST) (ver. 2.2.30) with a non-redundant protein sequence database (nr) and the protein sequence database of Göttingen and TGB-C1 (11) and BLASTKoala of Kyoto Encyclopedia of Genes and Genomes (KEGG) (8) were used to search for functional domains and characterize potential protein functions. Proteins associated with energy conservation systems were identified by criteria based on the genomic and physiological information of previously reported energy conservation pathways (21). Transport systems were identified using TransportDB (25).

Results and Discussion

**BCFA degradation**

The *Syntrophomonadaceae* strains 4J5 (40), TGB-C1 (29), and Göttingen (1, 15, 16) have very similar genetic compositions (see Supplemental information, Table S1, S2, and S3), reflecting previously observed physiological similarities, such as central metabolic pathways (e.g., glycolysis and the TCA cycle) and pilus and flagellum formation. Syntrophic BCFA degradation is considered to proceed through β-oxidation (13, 21), as observed for butyrate degradation by the strain Göttingen (16, 27, 30, 38). These *Syntrophomonadaceae* genomes all encode multiple genes for β-oxidation (Fig. 1, Table S4), potentially with varying specificities to particular fatty acid structures (i.e., alkyl branching, chain length, and saturation), affinities to specific concentration ranges, or adaptation to other environmental conditions (30). Strain 4J5 possesses several β-oxidation genes with high similarities (>90%) to those in Göttingen, presumably involved in butyrate metabolism (Table S4, Fig. S1) (27, 32). However, we also identified several strain 4J5 β-oxidation-related genes with relatively low similarities to those of Göttingen (42.3%–62.6% amino acid identity), which may specifically be involved in 2-methylbutyrate degradation—acyl-CoA dehydrogenase (Swmb 1942), enoyl-CoA hydratases (Swmb_01703 and Swmb_03023), 3-hydroxybutyryl-CoA dehydrogenase (Swmb_01947), and acetyl-CoA C-acetyltransferase (Swmb_02509). The TGB-C1 genome encodes several β-oxidation genes not only with relatively high similarities (30–85%), but also with no significant similarity (<30%) to those in the mesophilic strains Göttingen and 4J5 (Table S5), implying that TGB-C1 drives the β-oxidation pathway by using thermostable/thermophilic enzymes. Further biochemical, transcriptomic, and/or proteomic analyses are needed in order to clarify the activities and specificities of the β-oxidation-related enzymes of these strains.

In syntrophic isobutyrate degradation, strain TGB-C1 has been reported to perform alkyl isomerization in order to form butyrate for subsequent β-oxidation (29). Strain TGB-C1 encodes an isobutyryl-CoA mutase gene cluster including the

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**Fig. 1.** Branched-chain fatty acid metabolism in (A) *“Syntrophomonas wolfei* subsp. methylbutyratica” strain 4J5 and (B) *Syntrophothermus lipocalidus* strain TGB-C1. Each reaction is labeled with the protein name (italicized) of the gene encoding the function and locus tag(s). Figures in brackets indicate the number of multiple gene copies. Four (Swmb_00646–00651, Swmb_1699–1704, Swmb_1496–1500, and Swmb_2505–2510) and three (Slip_0499–0503, Slip_0577–0581, and Slip_1236–1241) β-oxidation gene clusters were identified in strains 4J5 and TGB-C1, respectively (shown in bold face).
C- and N-terminal domain proteins, cob(I)alamin adenosyl-transferase and MeaB-like protein (Slip_0757–0760) to facilitate the rearrangement step (Fig. 1B, Table S5). Homologous gene clusters have been found in “Ca. Caldisyntrophus multiacidovorans”, a syntrophic BCFA degrader in a scale anaerobic bioreactor (21) and other prokaryotes (Fig. S2). The isobutyryl-CoA mutase genes of TGB-C1 and “Ca. Caldisyntrophus” are phylogenetically related to a group associated with sulfate-reducing bacteria (Fig. S2), suggesting potential evolutionary relationships between isobutyrate-isomerizing syntrophs and sulfate reducers.

As final products of the predicted β-oxidation pathways, the syntrophic degradation of isobutyrate generates two acetyl-CoA and 2-methylbutyrate produces acetyl-CoA and propionyl-CoA (Fig. 1A). Acetyl-CoA yields ATP through dethiolation to acetate by phosphate acetyltransferase (Swmb_02801 and Slip_0902) and acetate kinase (Swmb_02802 and Slip_0903). Regarding strain 4J5, these enzymes may perform the dethiolation of 2-methylbutyrate-derived propionyl-CoA because the active site structure of previously known propionate kinase resembles those of acetate kinase and butyrate kinase (7). AMP-dependent acyl-CoA synthetases found in the 4J5 and TGB-C1 genomes (Swmb_02363 and Swmb_02710; Slip_0475, Slip_0583, and Slip_1686) potentially serve as an alternative acyl-CoA degradation pathway, as suggested by McInerney et al. (17). However, Swmb_02710 and Slip_1686 have high identities (>64% by amino acid sequences) to that of strain Göttingen (Swol_1180), which has been predicted to function in biosynthesis (30). The other homolog found in 4J5 (Swmb_02363) may be involved in poly-β-hydroxybutyrate metabolism due to an association with the poly-β-hydroxybutyrate polymerase gene, as observed in strain Göttingen (Swol_1144). The remaining TGB-C1 homologs (Slip_0475 and Slip_0583) have low amino acid sequence identities (<32%) with the biosynthesis-associated acyl-CoA synthetase (Table S4), implying that these acyl-CoA synthetase genes are responsible for the production of acetate from acetyl-CoA through the degradation of isobutyrate.

**Energy conservation and electron flow**

A syntrophic substrate metabolizer uses energy conservation systems, such as reverse electron transfer and electron bifurcation, to conduct thermodynamically unfavorable proton (H⁺) reduction to hydrogen (H₂) (26, 31). Since the 4J5, Göttingen, and TGB-C1 genomes lack the Rhodobacter nitrogen fixation (Rnf) complex, these Syntrophomonadaceae syntrophs may not perform reverse electron transport from NADH to oxidized ferredoxin (Fdox) through the Rnf found in several syntrophs (e.g., Syntrophus and Syntrophobacter) (17, 39). Instead, we identified other types of reverse electron transport systems employing electron transfer flavoprotein (ETF) dehydrogenases described in the Göttingen genome: ETF-linked iron-sulfur binding (Fe-S) reductase and ETF-oxidizing hydrogenase complex (FixABCX) systems (6, 10, 22, 30) (Fig. 2A, Table S6). These results support a previous finding on the expression of Fe-S reductase with strain Göttingen during syntrophic butyrate degradation (18, 27). Although these Syntrophomonadaceae genomes also encode an electron-bifurcating ETF-associated butyryl-CoA dehydrogenase (Bcd/EtfAB), this system may support the capacities of these organisms to perform crotonate reduction (10, 18). In addition, the Syntrophomonadaceae syntrophs encode a syntroph-associated flavin oxidoreductase-heterodisulfide reductase (Flox-Hdr) system (22, 24) expressed by the BCFA-
degrading syntroph “Ca. Caldisyntrophus multiacidivorans” and aromatic substrate-degrading syntrophs (Pelotomaculum spp. and Syntrophorhabdus spp.) along with FixABCX (21), suggesting the involvement of these energy conservation systems in the syntrophic degradation of various organic compounds including BCFA.

There may be multiple hydrogenase and formate dehydrogenase genes in the 4J5 and TGB-C1 genomes (Fig. 2, Table S6). Both strains harbor systems for hydrogen generation through electron-bifurcating Fe-only hydrogenase (NADH/Fd-oxidizing), Hyd hydrogenase (quinoi), and cytoplasmic hydrogenase (Fd) with relatively high similarities (≥68%) to those of strain Göttingen (30). In addition, the genomes of strains 4J5 and TGB-C1 encode the Hya hydrogenase gene cluster (Swmb_01823–01933 and Slip_0383–0392), which are lacking in the Göttingen genome. Strain TGB-C1 also has an energy-conserving hydrogenase (Ech; Fd-oxidizing) gene cluster (Slip_0665–0660). Regarding formate generation, we identified putative electron-bifurcating formate dehydrogenases (37) in strain 4J5, cytoplasmic formate dehydrogenase (NADH-oxidizing) in strain TGB-C1, and Fdo formate dehydrogenase (quinol-oxidizing) in both. These results suggest that BCFA-degrading syntrophs flexibly employ multiple energy conservation systems to maintain a syntrophic lifestyle in thermodynamically limited environments.

A nitrogenase gene (NifBDEHIK) was detected in the genome of strain TGB-C1 (Slip_2124–2130) along with a molybdate transporter (ModABC, Slip_2121–2123) and ammonia transporter (AmtB, Slip_2119) (Fig. 2C, Table S7). A previous microbial genome survey revealed that nitrogen fixation-related proteins (Nif) are distributed in phylogenetically diverse microbes including some fermentative bacteria and syntrophic substrate metabolizers (5). The nitrogenase activity of the fermentative H2-producing organism Clostridium butyricum strain CWB11009 has been proposed to generate H2 and enhance tolerance to acidification through the consumption of protons in the reaction, which produces ammonia as a base (3). We observed high amino acid sequence similarities (up to 98%) between the nitrogenase, molybdate transporter, and ammonia transporter genes of TGB-C1 and CWB11009, and also moderate similarities (up to 68%) with two other Nif-encoding syntrophs (Thermacetogenium phaeum strain PB and Syntrophobacter fumaroxidans strain MPOB) (Fig. 3, Table S7). Among syntrophs, nitrogen fixation may serve as a mechanism to tolerate acidification and provide hydrogen and ammonia for partner hydrogenotrophic methanogens to survive under hydrogen/ammonia-limited conditions.

In summary, a comparative genomic analysis of the Syntrophomonadaceae strains revealed multiple genes for the β-oxidation of BCFA/SCFA and energy conservation systems employing various types of electron carriers for electron disposal. In theory, the syntrophic degradation of 2-methylbutyrate and isobutyrate have Gibbs free energy values identical to butyrate under standard conditions (calculations based on Thauer et al. [36]):

$$\text{Butyrate} + 2\text{H}_2\text{O} \rightarrow 2\text{Acetate}^- + 2\text{H}_4 + \text{H}^+$$

$$\text{2-methylbutyrate} + 2\text{H}_2\text{O} \rightarrow \text{Acetate}^- + \text{Propionate}^+ + 2\text{H}_4 + \text{H}^+$$

$$\text{Isobutyrate}^- + 2\text{H}_2\text{O} \rightarrow 2\text{Acetate}^- + 2\text{H}_4 + \text{H}^+$$

$$\Delta G_0' = +48.3 \text{kJ mol}^{-1}$$

Thus, while Syntrophomonadaceae strains degrading the above fatty acids have unique substrate-specific genes (e.g., β-oxidation), they share similar energy conservation systems reflecting the thermodynamic similarity shown above. TGB-C1 has several distinct energy conservation pathways from Göttingen and 4J5 potentially due to differences in their temperature preferences. These genome-based results further provide insights into how syntrophic BCFA-degraders interact with H2/formate-utilizing methanogen in an anaerobic ecosystem.

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