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Novel energy conservation strategies and behaviour of *Pelotomaculum schinkii* driving syntrophic propionate catabolism

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Summary

Under methanogenic conditions, short-chain fatty acids are common byproducts from degradation of organic compounds and conversion of these acids is an important component of the global carbon cycle. Due to the thermodynamic difficulty of propionate degradation, this process requires syntrophic interaction between a bacterium and partner methanogen; however, the metabolic strategies and behaviour involved are not fully understood. In this study, the first genome analysis of obligately syntrophic propionate degraders (*Pelotomaculum schinkii* HH and *P. propionicicum* MGP) and comparison with other syntrophic propionate degrader genomes elucidated novel components of energy metabolism

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behind Pelotomaculum propionate oxidation. Combined with transcriptomic examination of P. schinkii behaviour in co-culture with Methanospirillum hungatei, we found that formate may be the preferred electron carrier for P. schinkii syntrophy. Propionatederived menaquinol may be primarily re-oxidized to formate, and energy was conserved during formate generation through newly proposed proton-pumping formate extrusion. P. schinkii did not overexpress conventional energy metabolism associated with a model syntrophic propionate degrader Syntrophobacter fumaroxidans MPOB (i.e., CoA transferase, Fix and Rnf). We also found that P. schinkii and the partner methanogen may also interact through flagellar contact and amino acid and fructose exchange. These findings provide new understanding of syntrophic energy acquisition and interactions.

Originality-Significance Statement

Syntrophic interaction between fatty acid-degrading *Bacteria* and methanogenic *Archaea* is a critical component of methanogenic decomposition of organic compounds. This study reports the first genome and transcriptome analyses of obligately syntrophic propionate oxidizers (e.g., *Pelotomaculum schinkii*). The investigation reveals novel insights into the energy metabolism and interspecies interactions of *P. schinkii* and differences between two well-known syntrophic propionate-oxidizing genera *Syntrophobacter* and *Pelotomaculum*.

Introduction

Syntrophy is a mutualistic microbial interaction, where for energetic reasons the degradation of a substrate can only occur when the reaction products, such as H_2 , acetate and formate are consumed by a partner microorganism. Methanogens and sulfate–reducing bacteria serve often as interacting partners of syntrophic bacteria by efficiently consuming their products to low concentrations (McInerney *et al.*, 2009). Syntrophy is necessary for the

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degradation of alcohols, organic acids and aromatic compounds in methanogenic ecosystems and is thus important in the global carbon cycle (McInerney et al., 2009; Schink and Stams, 2013). Degradation of propionate is thermodynamically challenging а conversion $[\Delta G = +73.8 \text{ kJ mol}_{\text{propionate}}, \text{ for H}_2 \text{ generation at } 37^{\circ}\text{C}$ and pH 7 (Thauer et al., 1977; Hanselmann, 1991; Amend and Shock, 2001)]. Although any bacterium requires a reaction to have at least -15 kJ mol⁻¹ to harness energy, the above propionate degradation can only reach this when H₂ (or formate) is kept at a low concentration (2.85 Pa H₂ or 1.58 µM formate) at 37°C, pH 7, 1 mM propionate, 1 mM acetate and 50 mM HCO3-. Thus, syntrophic propionate degraders must cope with highly energy-constrained conditions (Scholten et al., 2000: Adams et al., 2006).

Only four bacterial genera are known to syntrophically degrade propionate: Syntrophobacter, Pelotomaculum, Smithella and Desulfotomaculum (Harmsen et al., 1993; Wallrabenstein et al., 1995; Harmsen et al., 1998; Chen et al., 2005; de Bok et al., 2005). Biochemical and genomic analyses of Syntrophobacter fumaroxidans MPOB (Harmsen et al., 1998) and Pelotomaculum thermopropionicum SI (Imachi et al., 2002) have provided insight into their thermodynamically optimized way of life. This includes the discovery of use of energy from exergonic reactions and proton motive force to drive parallel endergonic reactions (i.e., energy conservation and reverse electron transport respectively; Houwen et al., 1990; Kosaka et al., 2008; Plugge et al., 2012; Sieber et al., 2012) and close physical interaction with the partner methanogen (Shimoyama et al., 2009; Stams and Plugge, 2009; Felchner-Zwirello et al., 2013). These studies have been instrumental to the development of our understanding of syntrophic propionate oxidation. However, S. fumaroxidans MPOB and P. thermopropionicum SI are not obligately syntrophic and can live as fermenters and/or sulfate reducers, so the biochemical and thermodynamic strategies and behaviour of species dedicated to syntrophy remain to be explored [e.g., P. schinkii HH and P. propionicicum MGP (de Bok et al., 2005; Imachi et al., 2007)].

In this study, we sequenced the first genomes of obligately syntrophic organisms *P. schinkii* HH and *P. propionicicum* MGP, sequenced an additional *Pelotomaculum* sp. for comparison (*P.* sp. FP), analysed their propionate oxidation and electron transfer strategies, and compared these genomes with other syntrophic propionatedegraders (SPDs). To further investigate the behaviour of an obligate syntrophic organism, we examined the gene expression of *P. schinkii* HH in co-culture with *Methanospirillum hungatei* JF-1 through transcriptomics. In total, we describe novel energy conservation strategies and interactions for *Pelotomaculum* SPDs and differences between *Pelotomaculum* and *Syntrophobacter*.

Results and Discussion

General genome information and the propionate oxidation pathway

The sequenced genomes of *Pelotomaculum* spp. (P. schinkii HH, P. propionicum MGP and P. sp. FP) were 3.95-4.47 Mb in size (sequencing depth of 69-122× coverage), had a GC content of 49.8%-50.8%, harboured 3847-4302 coding sequences and (Supporting Information Table S1). All genomes are available under NCBI BioProject PRJNA454509. All genomes encoded the conventional propionate oxidation pathway: propionate uptake, propionate activation to propionyl-CoA, carboxylation to methylmalonyl-CoA, subsequent isomerization to succinyl-CoA, oxidative decarboxylation to acetyl-CoA via oxaloacetate and pyruvate, acetyl-CoA dethiolation to acetate, and acetate export (Fig. 1; Supporting Information Table S3). While the first two steps (propionate activation and propionyl-CoA carboxylation) required energy input, these Pelotomaculum spp. all encoded CoA transferases and carboxyltransferases to couple the energy-requiring (i.e., endergonic) steps with downstream energy-yielding (i.e., exergonic) steps: acetyl-CoA dethiolation and oxaloacetate decarboxylation

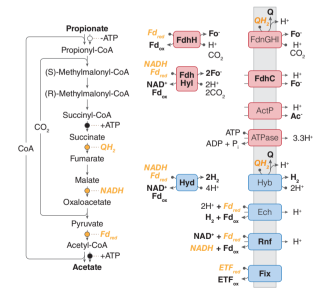


Fig. 1. Propionate degradation pathway of *Pelotomaculum schinkii* HH. Enzyme complexes involved in energy conservation, H₂ metabolism, or formate metabolism are coloured based on whether they were among the top 100 genes (red) or not (blue) in terms of gene expression (RPKM). The reduced electron carriers responsible for transducing reducing power from propionate oxidation to H₂ and formate generation are highlighted (orange). Enzymes related to H₂ and formate metabolism found across all sequenced syntrophic propionate degraders genomes are bolded.

respectively. All species also encoded AMP-dependent acyl-CoA synthetases to activate propionate and dethiolate acetyl-CoA independently. Based on these features, propionate oxidation to acetate by these *Pelotomaculum* spp. might yield 1 mol ATP and $6e^-$ (1 mol each of NADH, reduced ferredoxin (Fd_{red}) and menaquinol) per mol propionate.

Electron transfer proteins

Under methanogenic conditions, favourable electron acceptors are unavailable for electron disposal. Syntrophic organisms like Pelotomaculum spp. must resort to using H^+ and CO_2 as electron sinks for the reducing power generated from substrate oxidation (e.g., NADH, Fd_{red} and menaquinol). This is quite challenging as H⁺ reduction to H₂ and CO₂ reduction to formate both have very low reduction potentials (E' = -420 mV). For example, NADH (E' = -230 mV) is a highly unfavourable donor for H⁺/CO₂ reduction. To overcome this thermodynamic obstacle, all Pelotomaculum spp. encoded confurcating hydrogenases and formate dehydrogenases capable of driving endergonic H₂/formate-generating NADH oxidation using exergonic H₂/formate-generating oxidation of Fd_{red} (E' = -430 mV; Fig. 1 and Supporting Information Table S3). Electron confurcation systems have also been described for other syntrophic organisms (de Bok et al., 2003; McInerney et al., 2007; Kosaka et al., 2008; Sieber et al., 2010). Strains HH and FP also encoded membrane-bound hydrogenases (Ech) that can directly oxidize Fd_{red}, taking advantage of the positive energy margin between Fd_{red} and H₂ and extruding protons to conserve energy in the form of proton motive force.

As for menaguinol oxidation, all studied Pelotomacuencoded cytochrome b-linked quinonelum spp. dependent hydrogenases (HybABCO) and formate dehydrogenases (FdnGHI and FdnG-HybAB). We found that, in all Pelotomaculum spp., the cytochrome b subunits (i.e., HybB and FdnI) had guinone-binding sites on the periplasmic side [quinone-binding domain (Fisher and Rich, 2000) on periplasmic end of transmembrane helix (Kall et al., 2004)], indicating that Hyb- and Fdn-mediated menaquinol re-oxidation could potentially release protons on the periplasmic side and contribute to proton motive force formation. This is contrasting to the wellcharacterized Escherichia coli FdnGHI that has a cytoplasm-oriented quinone-binding site and couples formate oxidation with proton extrusion (Jormakka et al., Further, the *Pelotomaculum* coupling 2002). of menaquinol-oxidizing H₂/formate generation with proton extrusion is guite surprising as menaguinone has a much less negative reduction potential (E' = -74 mV) than H⁺ and CO₂. Energy acquisition from menaquinol oxidation would be a highly valuable biochemical strategy for syntrophic organisms, but simple thermodynamic calculations cannot justify this reaction, which is endergonic ($\Delta G = +38.20 \text{ kJ mol}^{-1}$) even if the H₂ concentration were as low as 1 Pa (or 2.96 μ M formate based on equilibrium at 37°C with 0.3 atm CO₂). Perhaps these *Pelotomaculum* spp. Hyb and Fdn employ an unbeknownst anaerobic energy conservation strategy.

While both H₂ and formate are known to play a critical role in syntrophic electron transfer, several studies have reported that SPDs may prefer formate transfer (Schmidt and Ahring, 1995; de Bok et al., 2002). For example, P. schinkii could not grow in co-culture with a methanogen incapable of utilizing formate (de Bok et al., 2005). Unlike H₂, formate is an anion, so it cannot freely pass the cell membrane and will inevitably accumulate in the cytosol. To address this, all studied Pelotomaculum spp. encoded potential formate transporters (FdhC) related (> 40% amino acid similarity) to Escherichia coli and Methanobacterium formicicum formate transporters FocA and FdhC (White and Ferry, 1992; Suppmann and Sawers, 1994; Supporting Information Table S3). We suspect that FdhC can take advantage of an accumulating formate gradient and couple formate extrusion to the symport of protons with generation of proton motive force. In agreement, the FdhC were adjacent to the electronconfurcating formate dehydrogenases FdhAB-HyIABC in the genomes of *P. schnkii* and *P.* sp. FP. This is a novel energy conservation mechanism for syntrophic propionate degradation.

The studied Pelotomaculum spp. also encoded energyconserving enzymes that can potentially mediate electron transfer from propionate oxidation to H₂/formate generation. P. schinkii and P. sp. FP encoded Fd:NADH oxidoreductases (RnfABCDEG) that can couple exergonic electron transfer from Fd_{red} to NAD⁺ with proton extrusion. In addition, all studied Pelotomaculum spp. harboured electron transfer flavoprotein dehydrogenases (FixABCX) that can couple quinol-reducing electron transfer flavoprotein oxidation with proton pumping. Both complexes have been proposed to play key roles in syntrophic metabolism (Sieber et al., 2012; Nobu et al., 2015). In total, each of these Pelotomaculum spp. possessed at least five potential approaches to pairing electron transfer with proton motive force generation, which suggests metabolic optimization towards energy conservation.

Comparison with other syntrophic propionate degraders

Besides the three mesophilic *Pelotomaculum* strains discussed so far, two other SPDs have been genomically characterized by previous studies: thermophilic *Pelotomaculum thermopropionicum* and mesophilic *Syntrophobacter fumaroxidans* (Kosaka *et al.*, 2008; Plugge *et al.*, 2012). All these SPDs had the general propionate

4506 Hidalgo C. A. et al.

oxidation pathway, electron-confurcating hydrogenase, electron-confurcating formate dehydrogenase, monomeric formate dehydrogenase (FdhH), and formate transporter in common (Supporting Information Table S3). This suggests that electron confurcation and the newly proposed energy acquisition from formate extrusion are core features of syntrophic propionate degradation. *P. thermopropionicum* also shares many other features (e.g., guinone-dependent hydrogenases and formate dehydrogenases) with the three mesophilic Pelotomaculum strains, but lacks Ech and Rnf (Supporting Information Table S3). As S. fumaroxidans is from a different genus, we found several major differences. While the four Pelotomaculum spp. only encoded propionyl-CoA:acetate CoA transferases, S. fumaroxidans also harboured a propionvl-CoA:succinate CoA transferase. Thus, unlike Pelotomaculum, Syntrophobacter may also activate propionate using succinyl-CoA hydrolysis as an energy input. Conversely, Syntrophobacter lacked membranebound hydrogenases (Ech and HybABCO) and formate dehydrogenases (FdnGHI) associated with proton extrusion, and rather encoded a cytosolic NiFe hydrogenase (HoxEFUHY) and periplasmic heterodimeric formate dehydrogenase that were not directly involved in proton pumping. The energy conservation strategies of Pelotomaculum and Syntrophobacter clearly differ, where Syntrophobacter seemed to have a more intricate energy conservation mechanism. A recent study has also reported contrasting energy conservation mechanisms among syntrophic acetate and aromatic compound

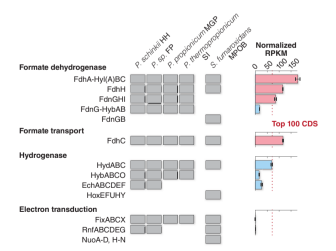


Fig. 2. Energy conservation strategies of *Pelotomaculum* and *Syntrophobacter* species (left) and gene expression of these enzymes by *P. schinkii* HH under co-culture with *Methanospirillum hungatei* JF-1 (right). The gene expression of the subunit with the highest expression level is shown as RPKM (reads per kilobase of transcript per million mapped reads) normalized to the *P. schinkii*'s median RPKM. Bars are coloured based on whether the subunits have expression levels (RPKM) among the top 100 genes (red) or not (blue). Error bars are also shown.

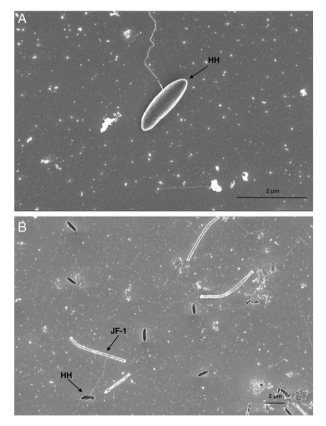
oxidizers (Nobu *et al.*, 2017; Manzoor *et al.*, 2018), suggesting diversity in energy metabolism across syntrophic organisms.

Pelotomaculum schinkii gene expression during syntrophic propionate degradation

To further investigate the energy conservation strategies of syntrophic propionate degradation, we studied the gene expression profile of a co-culture of an obligate SPD (P. schinkii) and H₂/formate-using methanogen M. hungatei. Transcriptomes were sequenced for triplicate co-cultures (4.9-5.2 million reads and 736-778 Gb) and, as a reference, triplicate cultures of *M. hungatei* as well (3.8-4.7 million reads and 577-704 Gb; Supporting Information Table S2). For the co-cultures, 22.2%-26.9% and 7.6%-11.2% of the transcriptomic reads mapped to P. schinkii and M. hungatei CDS respectively. For the M. hungatei pure culture, 35.1%-51.3% mapped to M. hungatei CDS. While most P. schinkii genes involved in propionate uptake, propionate oxidation, and acetate export were highly expressed (3rd-81st on the expression ranking, refer to Supporting Information Table S3 for specific gene expression levels and standard deviation), CoA transferases ranked very low (2367th and below; Supporting Information Table S3). Instead, P. schinkii expressed an AMP-dependent acyl-CoA synthetase (Acs1; 181st). Thus, P. schinkii likely decouples propionate activation and acetyl-CoA hydrolysis and does not conserve energy here. This is quite dissimilar from other characterized SPDs who are thought to depend on energy conservation via CoA transferases: Syntrophobacter fumaroxidans and Pelotomaculum thermopropionicum (Kato et al., 2009; Plugge et al., 2012). Clearly, even strategies for propionate oxidation can vary, suggesting overlooked diversity among SPDs in thermodynamic optimization. The use of AMP-dependent pyrophosphate-forming acyl-CoA synthetase for substrate activation has also been observed for Syntrophus aciditrophicus during syntrophic benzoate degradation, indicating that P. schinkii shares similarities in biochemical strategies with other syntrophic organisms. In addition, like S. aciditrophicus and Clostridium ultunense (James et al., 2016; Manzoor et al., 2018), P. schinkii may conserve energy from substrate activation by coupling proton extrusion with hydrolysis of the byproduct pyrophosphate via a membrane-bound pyrophosphatase (rank 342nd; Supporting Information Table S4).

The gene expression profile of electron transfer enzymes also revealed unexpected behaviour. *P. schinkii* highly expressed formate generation through three formate dehydrogenases: FdhAB-HyIABC (27th), FdhH (59th) and FdnGHI (86th; Figs. 1 and 2; Supporting Information Table S3). Conversely, the hydrogenases all

had lower expression levels (all ranked less than 100th). To compare, the electron-confurcating and guinonedependent hydrogenase catalytic subunits only had $48.5 \pm 1.8\%$ (p = 0.011) and 17.8 $\pm 1.1\%$ (p = 0.002) of the expression of the respective formate dehydrogenase catalytic subunits. This suggests that P. schinkii may preferentially funnel reducing power into CO2-reducing formate generation rather than H₂ production, like S. fumaroxidans and P. thermopropionicum (Worm et al., 2011; Liu and Lu, 2018). In particular, the markedly different expression levels of Hvb and Fdn indicates that P. schinkii primarily re-oxidizes menaguinol through formate generation. This is the first evidence for a syntrophic organism segregating different electron carriers towards different electron acceptors. To complement the highly active formate metabolism. P. schinkii also highly expressed the formate transporter (FdhC; 58th) to acquire energy through the newly proposed formate export. This further evidences the importance of formatemediated proton pumping for syntrophic propionate degradation (Fig. 2). Similarly, the methanogenic partner M. hungatei in the co-culture expressed formate uptake (FdhC, Mhun 1811; 13th) and oxidation (two FdhAB, Mhun 1813-14 and Mhun 3238-37; 19th and 23rd)



higher than H₂ oxidation (e.g., FrhABDG; 40th). The *M. hungatei* formate dehydrogenase expression level was significantly higher when grown in co-culture than in an H₂-fed axenic culture (14.54×, p = 0.05; Supporting Information Tables S5 and S6), indicating that *M. hungatei* was oxidizing *P. schinkii*-derived formate.

Interestingly, the highly expressed formate dehydrogenases can directly transfer electrons from propionate oxidation to formate generation. Propionate oxidation to acetate generates NADH, Fdred and menaguinol as reducing equivalents at a 1:1:1 ratio (Fig. 1). The expressed formate dehydrogenases can stoichiometrically re-oxidize these electron carriers: FdhAB-HyIABC putatively oxidizes NADH and Fd_{red} at a 1:1 ratio and FdnGHI oxidizes menaguinol, using CO₂ as an electron acceptor. This is quite unique as syntrophic organisms typically depend on electron transduction mechanisms to couple substrate degradation with electron disposal (e.g., NADH:Fd oxidoreductase Rnf and electron transfer flavoprotein dehydrogenase Fix; Sieber et al., 2012). The genome of P. schinkii encoded such syntrophy-associated enzymes (Rnf and Fix) but did not express them highly (Rnf 1863rd and Fix 3095th; Supporting Information Table S3). Another syntrophic organism, Syntrophaceticus schinkii, has also been reported to possess but not express Rnf (Manzoor et al., 2016). Thus, P. schinkii, and perhaps other syntrophic organisms, employ a minimalist energy conservation approach distinct from those that are well-characterized (e.g., Syntrophobacter fumaroxidans and Syntrophus aciditrophicus; McInerney et al., 2007; Plugge et al., 2012).

Flagellum and amino acid-mediated syntrophy. We further explored genomes and the transcriptome for noncatabolic behaviour relevant to syntrophy. Previous studies demonstrate that P. thermopropionicum can interact with its partner methanogen through flagella (Ishii et al., 2005; Shimoyama et al., 2009; Liu and Lu, 2018). We also found that P. schinkii expressed flagella-related genes despite being immotile (average rank 484th; Supporting Information Table S3; Fig. 3; de Bok et al., 2005). Given that all Pelotomaculum SPDs encoded complete flagellum biosynthesis machinery (Supporting Information Table S3), 'flagellummediated symbiosis' may be a core mechanism for Pelotomaculum-initiated syntrophy. While most flagellar components were quite homologous between Pelotomacu*lum* SPDs (average amino acid sequence similarity > 60%), the specific subunit that interacts with methanogens [FliD (Shimoyama et al., 2009)] can be guite different between strains (HH-SI 29.7%, HH-MGP 45.4% and HH-FP 97.0%). Though highly speculative, perhaps FliD (and thus Pelotomaculum) may have some level of specificity or affinity to methanogenic partner. In future work, it would be interesting to delve into investigation of the stimuli behind such interspecies interactions and the importance in H_2 and formate transfer.

The transcriptomics also revealed potential nutrient exchange between *P. schinkii* and *M. hungatei*. Although

4508 Hidalgo C. A. et al.

P. schinkii had complete pathways for synthesizing all standard amino acids (AAs), it had depressed expression of biosynthesis for five AAs (i.e., lower than median gene expression; Fig. 4). Conversely, we found that *M. hungatei* expressed synthesis for three of these AAs (arginine, tyrosine and methionine; greater than median gene expression). Similarly, P. schinkii highly expressed biosynthesis of four other AAs that M. hungatei did not (proline, serine, phenylalanine and tryptophan). M. hungatei indeed increased expression of arginine and methionine synthesis and decreased expression of tryptophan synthesis in co-culture compared to axenic, providing evidence for changes in AA biosynthesis behaviour in the presence of P. schinkii. This suggests division of labour for AA biosynthesis and AA exchange between the two syntrophically interacting strains. However, these results must be interpreted with caution as the co-culture contains yeast extract, which contains amino acids. Some recent studies also identify coordinated crossfeeding between organisms (Hubalek et al., 2017; Liu et al., 2018; Zengler and Zaramela, 2018), suggesting nutrient exchange may be a prevalent and critical interaction for syntrophy.

Other anaerobic metabolism. P. schinkii is reported to syntrophically degrade propionate and also co-utilize fumarate in the presence of propionate, but not metabolize any other tested compounds (de Bok et al., 2005). However, P. schinkii genome encoded genes for the degradation of many amino acids, fructose and ethanol and fermentative production of butyrate, lactate, ethanol and propanediol (Supporting Information Table S7). Interestingly, catabolismspecific pathways of fructose (using phosphofructokinase rather than anabolic fructose-1,6-bisphosphatase) and branched-chain AAs were highly expressed by P. schinkii (all genes have average RPKM greater than the median RPKM for Pelotomaculum; Supporting Information Table S7) while biosynthesis of these compounds were expressed by M. hungatei (average RPKM normalized to the median: 1.01-15.07 for fructose and 2.56-24.97 for branched-chain AAs). This suggests that exchange of these compounds may also play a role in syntrophy between the two partners and, unlike AA transfer discussed in the previous section. fructose and branched-chain AAs may serve as a supplementary energy sources co-utilized with propionate. While aspartate and alanine metabolism were also highly expressed, these pathways are reversible and may be used for anabolism. Most other catabolism- (i.e., threonine, methionine and lysine degradation) or fermentation-specific (i.e., ethanol, lactate, butyrate and propanediol metabolism) pathways were not highly expressed (Supporting Information Table S7). Given that *P. schinkii* was reported to not be able to utilize or produce (in excess) these compounds (de Bok *et al.*, 2005), *P. schinkii* may employ these genes for other untested environmental conditions and/or alternative lifestyles that we have yet to uncover.

As syntrophic propionate catabolism only has a small energy margin (-15 kJ mol⁻¹) that can be shared between the syntroph and its partner (de Bok et al., 2004; Schink and Stams, 2013), it is critical for SPD genera to optimize the energetics. Comparative genomics and transcriptomics revealed that Pelotomaculum syntrophy entails formatedominated interspecies electron transfer, energy acquisition through formate extrusion, potential flagellum-mediated interaction, exchange of AA for anabolism, exchange of AA and fructose for potentially supplementary catabolism and unexpected independence from conventional energy conservation pathways typically associated with other well-studied syntrophs (e.g., Syntrophobacter fumaroxidans, Syntrophomonas wolfei and Syntrophus aciditrophicus; McInerney et al., 2007; Sieber et al., 2010; Plugge et al., 2012; Sieber et al., 2012). Most importantly, these results reinforce a previous observation that formate may play a larger role in syntrophy than H₂ (de Bok et al., 2002) and provide further evidence for cross-feeding between syntrophs and their partners in parallel with H₂ and formate transfer.

Methods

Cultivation experiment. M. hungatei $JF-1^{T}$ (=DSM 864^T) was obtained from the Deutsche Sammlung von Mikroorganismen and Zellkulturen (Braunschweig, Germany). The co-culture of *P. schinkii* and *M. hungatei* was obtained from internal strain collection. A bicarbonate-buffered mineral medium was used, with the following composition: 3 mM Na₂HPO₄, 3 mM KH₂PO₄, 5.6 mM NH₄Cl, 0.75 mM CaCl₂, 0.5 mM MgCl₂, 5 mM NaCl, 50 mM NaHCO₃, 1 mM Na₂S, 7.5 mM FeCl₂, 1 mM H₃BO₃, 0.5 mM ZnCl₂, 0.1 mM CuCl₂, 0.5 mM MnCl₂, 0.5 mM CoCl₂, 0.1 mM NiCl₂, 0.1 mM Na₂SeO₃, 0.1 mM Na₂WO₄, 0.1 mM Na₂MOO₄, 0.5 mg

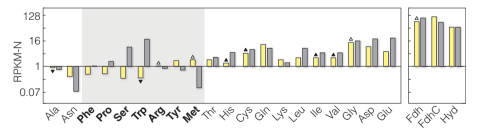


Fig. 4. Expression of amino acid biosynthesis, formate metabolism (Fdh), formate transport (FdhC) and H₂ metabolism (Hyd), by *M. hungatei* JF-1 (yellow) and *P. schinkii* HH (grey). The expression level of the gene with the lowest expression level (RPKM normalized to the median RPKM of the corresponding genome) of each pathway is shown. Biosynthesis pathways with significantly different expression levels ($p \le 0.05$) between co-culture and axenic culture for *M. hungatei* JF-1 are marked (black triangle). Pathways are marked with a white triangle if the expression levels are significantly different only when axenic culture replicate 2 is excluded. The direction of the triangle indicates whether *M. hungatei* JF-1's expression increases (up) or decreases (down) in the co-culture compared to the axenic culture.

EDTA I^{-1} and the following vitamins (mg I^{-1}): 0.02 biotin, 0.2 nicotinic acid, 0.5 pyridoxine, 0.1 riboflavin, 0.2 thiamin, 0.1 cvanocobalamin, 0.1 p-aminobenzoic acid, 0.1 pantothenic acid, 0.1 lipoic acid and 0.1 folic acid. To obtain the biomass for genomic sequencing, M. hungatei was cultured at 37°C in 1 I serum bottles with 500 ml medium and a gas phase of 1.7 atm 80:20 v/v of H₂/CO₂. The P. schinkii co-culture was inoculated into a pregrown M. hungatei culture (1:10). Prior to inoculation, CH₄ and H₂ was removed by exchanging with 1.7 atm of N₂/CO₂ gas (80:20 v/v) and 0.1 g/l of yeast extract and 20 mM of propionate was added. Co-culture bottles were done in triplicates and incubated at 37°C during approximately 30 days. For transcriptome sequencing, 10% of co-culture inoculum was transferred directly to 500 ml bottles with 250 ml medium and incubated at 37°C during approximately 15 days. Pure cultures of M. hungatei were grown at 37°C during 4 days in 250 ml bottles with 100 ml medium and 1.7 atm 80:20 v/v of H_2/CO_2 in the headspace. All culturing and sequencing for transcriptomics was done in biological triplicates.

Nucleic acids extraction. Biomass was harvested by centrifugation at 15.000 g at 4°C under sterile conditions. DNA was extracted using MasterPure™ Gram Positive DNA Purification Kit following the manufacturer specifications. Quality and quantity of total DNA was measured using a NanoDropTM 2000 spectrophotometer and QubitTM dsDNA BR Assay Kit. Total RNA extraction was performed from cells harvested in the middle of exponential growth phase (~6 mM CH₄ produced and 10 mM propionate consumed). Lysis and protein precipitation was performed using the solutions and enzymes from MasterPure™ Gram Positive DNA Purification Kit as follows: Lysozyme incubation at room temperature for 10 min, after lysis 3 uL of β -mercaptoethanol were added, cells were sonicated using Bendelin SONO-PULS HD 3200 ultrasonic homogenizer (6 cycles of 20 s. pulse 30 s. pause) and proteinase K incubation at 60°C for 10 min. Protein precipitation was performed according to the kit specifications. Automated RNA purification was performed using Maxwell[®] 16 MDx instrument and LEV simplyRNA Purification Kit (DNAse treatment and 16S rRNA depletion was included on the kit).

Microscopy. For scanning electron microscopy analysis, the culture was adhered to poly L-lysin 12 mm coated coverslips (Corning, BioCoat) and incubated for 1 h at room temperature. The cells were then fixed in 2.5% glutaraldehyde in 0.1 M phosphate buffer (pH 7.4) for 1 h at room temperature, rinsed 3 times with 0.1 M phosphate buffer (pH 7.4) and postfixed with 1% osmium tetroxide for 60 min. Hereafter, the cells were dehydrated in a graded alcohol series (10%, 30%, 50%, 70%, 80%, 96% and 100%), dried to critic point in 100% ethanol with CO_2 in the Leica EM CPD300 system and mounted onto aluminium stubs and coated with tungsten. Cells were subsequently studied with a FEI Magellan 400 scanning electron microscope.

Sequencing and bioinformatics. To sequence the genomes of *P. schinkii*, *P. propionicum* and *P.* sp. FP, we performed DNA extraction, Illumina (San Diego, CA, USA) MiSeq sequencing, quality control with Trimmomatic v0.36

Pelotomaculum schinkii syntrophic catabolism 4509

(SLIDINGWINDOW:6:30 MINLEN:78 LEADING:3 TRAILING:3; Bolger et al., 2014), and genome assembly with SPAdes v3.11 as previously described (Narihiro et al., 2016a,b). For the P. schinkii and M. hungatei co-culture, genomic DNA was also sequenced using PacBio (PacBio RS II, > 6000 bp reads, 500 Mb raw data), and the resulting sequences were coassembled using SPAdes v3.10 with default settings. The M. hungatei sequences were removed by comparison with the publically available *M. hungatei* genome. The *Pelotomaculum* genomes were annotated using Prokka 1.12 (Seemann, 2014: Nobu et al., 2015). As for the transcriptomics. RNA extracted from the P. schinkii and M. hungatei co-culture was sequenced using RNA-seg strand specific library constructed using a Kapa Hyper Prep Kit (Kapa Biosystems) including 16S depletion and subjected to 250-300 bp insert paired-end sequencing on an Illumina HiSeq 4000 platform (sequenced by Beijing Novogene Bioinformatics Technology), trimmed using Trimmomatic v0.36 (SLIDINGWINDOW:6:30 MINLEN:50 LEADING:3 TRAILING:3), and mapped to the P. schinkii and M. hungatei genomes using the Burrows Wheeler Aligner with default settings (Li and Durbin, 2009) to calculate gene expression levels as previously described (Nobu et al., 2017).

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4510 Hidalgo C. A. et al.

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Supporting Information

Additional Supporting Information may be found in the online version of this article at the publisher's web-site:

- Table S1. Genome statistics
- Table S2. Transcriptome statistics

Table S3. Gene annotation of propionate catabolis, hydrogen/formate generation and electron transduction pathways.

Table S4. Gene expression levels for *P. schinkii* HH in coculture with JF-1 in terms of reads per kilobase transcript per million mapped reads normalized to the median.

Table S5. Gene expression levels for *M. hungatei* JF-1 in co-culture with *P. schinkii* HH (left) and in axenic culture (right) in terms of reads per kilobase transcript per million mapped reads normalized to the median. The ratio of the expression and *p*-value of each gene in co-culture against axenic culture is shown (p < 0.05 coloured yellow). **p*-value calculated from replicate 1 and 3 of the co-culture and all replicates of the axenic culture (p < 0.05 coloured blue). This may be justifiable as the standard deviation of the replicate 1 and 3 is less than that of all three replicates for 82% of the JF-1 genes with detected expression. In addition, the standard deviation of replicates 1 and 3 is less than any other combination (i.e., rep. 1 + 2, 2 + 3 and 1 + 2 + 3) for 70% of the JF-1 genes with detected expression.

Table S6. Gene expression levels for *M. hungatei* JF-1 in an axenic culture in terms of reads per kilobase transcript per million mapped reads normalized to the median.

Table S7. Other catabolic/fermentation pathways found in the *P. schinkii* HH genome and the corresponding genes' average expression level (RPKM) during syntrophic propionate degradation.