Concerted Metabolic Shifts Give New Insights Into the Syntrophic Mechanism Between Propionate-Fermenting *Pelotomaculum thermopropionicum* and Hydrogenotrophic *Methanocella conradii*

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Microbial syntrophy is a thermodynamically-based cooperation between microbial partners that share the small amounts of free energy for anaerobic growth. To gain insights into the mechanism by which syntrophic microorganisms coordinate their metabolism, we constructed cocultures of propionate-oxidizing *Pelotomaculum thermopropionicum* and hydrogenotrophic *Methanocella conradii* and compared them to monocultures. Transcriptome analysis was performed on these cultures using strand-specific mRNA sequencing (RNA-Seq). The results showed that in coculture both *P. thermopropionicum* and *M. conradii* significantly upregulated the expression of genes involved in catabolism but downregulated those for anabolic biosynthesis. Specifically, genes coding for the methylmalonyl-CoA pathway in *P. thermopropionicum* and key genes for methanogenesis in *M. conradii* were substantially upregulated in coculture compared to monocultures. The putative flavin-based electron bifurcation/confurcation systems in both organisms were also upregulated in coculture. Formate dehydrogenase encoding genes in both organisms were markedly upregulated, indicating that formate was produced and utilized by *P. thermopropionicum* and *M. conradii*, respectively. The inhibition of syntrophic activity by formate and 2-bromoethanesulphonate (2-BES) but not H\textsubscript{2}/CO\textsubscript{2} also suggested that formate production was used by *P. thermopropionicum* for the recycling of intracellular redox mediators. Finally, flagellum-induced signal transduction and amino acids exchange was upregulated for syntrophic interactions. Together, our study suggests that syntrophic organisms employ multiple strategies including global metabolic shift, utilization of electron bifurcation/confurcation and employing formate as an alternate electron carrier to optimize their metabolisms for syntrophic growth.

**Keywords:** *P. thermopropionicum*, *M. conradii*, syntrophy, propionate, formate, interspecies electron transfer, RNA-Seq
INTRODUCTION

In anaerobic habitats, microbial consortia, of which each member has a specific metabolic function, perform the degradation of complex organic matter to carbon dioxide (CO₂) and methane (CH₄) (Schink, 1997; Conrad, 1999). Syntrophic oxidation of key intermediates (e.g., propionate and butyrate) requires tightly coupled, mutualistic interactions between fermentative bacteria and methanogenic archaea (Schink and Stams, 2006; McInerney et al., 2008; McInerney et al., 2009). How syntrophic consortia conserve energy from reactions that operate close to the thermodynamic equilibrium and which molecular mechanisms are involved in the establishment and maintenance of syntrophic interactions still are open questions (Sieber et al., 2012).

Propionate is next to acetate as an important intermediate during methanogenic degradation of organic matter in anaerobic habitats (Rothfuss and Conrad, 1993; Glissmann and Conrad, 2000; McInerney et al., 2009). Two types of syntrophic bacteria are found to oxidize propionate in association with methanogens. The first type utilizes the methylmalonyl-CoA (MMC) pathway in which three pairs of electrons from propionate oxidation are released in the form of hydrogen (H₂) and/or formate. Organisms of this type include Syntrophobacter spp. and Pelotomaculum spp. (McInerney et al., 2008; Müller et al., 2010). The second type oxidizes propionate through a dismutating pathway in which two propionate molecules are first jointed to form a C₆ compound which is then dismutated to acetic and butyrate before being degraded via beta oxidation. The main organisms of this type include Smithella spp. (de Bok et al., 2001).

Pelotomaculum spp. are widespread in bioreactors (Imachi et al., 2002; Nobu et al., 2015) and in natural environments like rice field soils (Lueders et al., 2004; Gan et al., 2012) where they cooperate with methanogens to oxidize propionate and other organic compounds. Pelotomaculum thermopropionicum has been used as a model organism for studies of the physiology and ecology of propionate-oxidizing syntrophs (Ishi et al., 2005, 2006; Kato et al., 2009). The genome of P. thermopropionicum reveals that several peripheral pathways are associated with the central MMC pathway, indicating that these organisms can have a broad substrate specificity (Imachi et al., 2002; Kosaka et al., 2006, 2008). Genes coding for most of the MMC pathway form a large gene cluster (the mmc cluster), which may facilitate its coordinate expression and hence save transcription costs (Kosaka et al., 2006, 2008). The genome encodes four hydrogenases and two formate dehydrogenases, some of which are considered to be capable of mediating electron communication (Kosaka et al., 2008; Kato et al., 2009; Sieber et al., 2010). In addition, the genome encodes a large number of signal-transduction mechanisms (Kosaka et al., 2008). Transcriptomic analysis revealed that P. thermopropionicum has complex regulatory mechanisms for gene expression and for metabolism in response to substrate addition and coculture conditions (Kato et al., 2009). There is also evidence that P. thermopropionicum may be capable of flagellum-mediated communication with the methanogen partners (Shimoyama et al., 2009).

Different methanogens have been used for studies on syntrophic metabolism (Kato et al., 2009; Enoki et al., 2011; Worm et al., 2011b; Walker et al., 2012; Sieber et al., 2015). Methanocella spp. (order Methanocellales), of which so far three strains were isolated, are all obtained from paddy field soil (Sakai et al., 2008, 2010; Lü and Lu, 2012). However, environmental surveys indicate that Methanocellales are widespread in nature including not only paddy field soil but also peat bogs, river and lake sediment, and even upland and desert soils (Conrad et al., 2006; Angel et al., 2011, 2012; Aschenbach et al., 2013). Methanocella spp. are hydrogenotrophic and well-adapted to low H₂ concentrations (Lu et al., 2005; Sakai et al., 2009). Consequently, they are often detected in association with secondary fermenters for syntrophic oxidation of fatty acids (e.g., propionate) occurring as intermediates of the anaerobic degradation of organic matter in paddy field soil (Lueders et al., 2004; Liu and Conrad, 2010; Liu P. et al., 2011; Rui et al., 2011; Gan et al., 2012). The genome analysis of Methanocella spp. has reconstructed their core metabolism and predicted certain novel features, including a set of genes potentially involved in utilizing low concentrations of H₂ (Lyu and Lu, 2015). A novel hydrogenase, tentatively named as disulfide-reducing hydrogenase (Drh), was discovered that is phylogenetically more closely related to Coo hydrogenase (carbon monoxide-induced hydrogenase) of bacterial sulfate reducers than to any canonical hydrogenase of archaeal methanogens (Lyu and Lu, 2015). We have previously constructed syntrophic cocultures using Methanocella conradii in combination with either Syntrophomonas wolfei or P. thermopropionicum (Liu et al., 2014), which revealed that M. conradii in coculture tended to upregulate the expression of genes coding for methanogenesis pathway but downregulate the genes coding for anabolic biosynthesis. However, only a few functional genes were examined and moreover the response of the bacterial partners was not evaluated. Therefore, a holistic understanding on how syntrophic organisms coordinate their metabolism under energy-limiting conditions is still lacking.

Both hydrogen and formate transfer are important interspecies electron transfer mechanisms (de Bok et al., 2004; Sieber et al., 2012; Schink et al., 2017; Sedano-Núñez et al., 2018). It has been reported that formate is temporarily produced by P. thermopropionicum when it grows on ethanol under syntrophic association with a hydrogenotrophic methanogen (Kosaka et al., 2008). However, it is still unclear whether P. thermopropionicum also produces formate when growing on propionate. Although this possibility has been suggested to be unlikely (Imachi et al., 2002), a significant upregulation of formate dehydrogenases was observed in P. thermopropionicum when grown in propionate degrading coculture (Kato et al., 2009). M. conradii cannot use formate as sole substrate for growth and methane production (Lü and Lu, 2012). However, we previously showed that M. conradii can use formate for methane production if H₂ is supplied in addition. Furthermore, formate dehydrogenase was upregulated in syntrophic cocultures with propionate- or butyrate- degrading syntrophs, indicating that M. conradii is able to perform formate-dependent methanogenesis (Liu et al., 2014). However, since such upregulation may be merely a response to H₂ limitation (Nölling and Reeve, 1997; Wood et al., 2003; Hendrickson et al.,...
2007), the importance of formate as electron shuttle between *P. thermopropionicum* and *M. conradii* is still not clear.

In the present study, high throughput sequencing of transcriptomes (RNA-Seq) was used to study the global gene expression in a coculture of *P. thermopropionicum* and *M. conradii*. RNA-Seq is believed to offer a greater dynamic range with less background noise and reveal sequence identity directly (Giannoukos et al., 2012; McGettigan, 2013; Hrdlickova et al., 2017). The strand-specific RNA-Seq in particular resolves the correct expression levels of coding or non-coding overlapping transcripts and provides accurate quantification of gene expression (Levin et al., 2010). We hypothesized that interactions during syntrophic growth will trigger global transcriptional responses in both *P. thermopropionicum* and *M. conradii*. A further aspect was the role of formate in interspecies electron transfer.

**MATERIALS AND METHODS**

**Cultivation and Cell Collection**

*P. thermopropionicum* strain SI^T^ (DSM 13744) monoculture (abbreviated as PM) was grown in basal medium with 20 mM pyruvate as substrate under N\(_2\)/CO\(_2\) (80/20, v/v) at 170 kPa (Imachi et al., 2002; Liu et al., 2014). *M. conradii* strain HZ254^T^ (DSM 24964) was grown in monoculture (abbreviated as MM) under H\(_2\)/CO\(_2\) (80/20, v/v) at 170 kPa or in coculture (abbreviated as PMC) with *P. thermopropionicum* by using 20 mM propionate as substrate under N\(_2\)/CO\(_2\) (80/20, v/v) at 170 kPa, as previously described (Lü and Lu, 2012; Liu et al., 2014). All the components of the basal medium were the same for all cultures except for the substrates. To obtain adequate amounts of RNA for sequencing, the culture volume was 100 ml in 550-ml bottle for PM, 500 ml in 2.3-L bottle for MM, and 500 ml in 1.1-L bottle for PMC. Cultivation was carried out at 55°C in the dark without agitation. The growth of PM was monitored by measuring optical density at 578 nm (OD578) and those of MM and PMC were monitored by analysis of CH\(_4\) production in the headspace as previously described (Liu et al., 2014). All cultures were harvested for transcriptomic analysis at the mid- to late exponential growth phases (Figure S1). Cell suspensions were centrifuged at 23,000 g at 4°C for 20 min (Avanti J-26XP, Beckman Coulter, USA), supernatants were discarded and 1.8 ml RNAlater (Ambion, USA) was added to the pellets immediately, which were then incubated at 4°C overnight. Cells were recollected by centrifuging at 20,000 g at 4°C for 60 min. Supernatants were discarded carefully and the pellets were stored immediately at −80°C until RNA extraction.

In order to test the effects of 2-BES, H\(_2\)/CO\(_2\) and formate on PMC, 100 ml cultures in 250-ml bottle were used. 2-BES served as a specific methanogenic inhibitor (Liu H. et al., 2011). Other culture conditions were the same as described above. Consumption of propionate, H\(_2\), and formate, and the production of CH\(_4\) and acetate were monitored by GC or HPLC as previously described (Liu et al., 2014). Five mM 2-BES, 20 mM formate and/or H\(_2)/CO\(_2\) (80/20, v/v, 170 kPa) were added after the initiation of propionate degradation. The headspace of the bottles treated with 20 mM formate were re-flushed with N\(_2)/CO\(_2\) (80/20, v/v) at 170 kPa.

**Total RNA Extraction and mRNA Enrichment**

Total RNA was extracted using AllPrep DNA/RNA Mini Kit (Qiagen, Germany) according to the manufacturer’s instruction. Lysis buffer RLT plus was supplemented with 2% (v/v) 2 M Dithiothreitol (DTT) (Sangon Biotech, China). Lysis buffer RLT plus (600 µl) was added to each cell pellet. Cell pellets of PM and PMC were homogenized for 120 s (5.0 m/s, 3 × 40 s with 2 min intervals on ice) in a FastPrep-24 (MP Biomedicals, USA) with 0.5 g RNase free beating beads (0.1 mm, Sigma, USA). Cell pellets of MM were homogenized at 5.0 m/s for 20 s without beating beads. Residue DNA digestion, verification of absence of DNA, and RNA purification were carried out as previously described (Liu et al., 2014). RNA was finally dissolved in 75 µl RNase free H\(_2\)O. RNA integrity values (RIN) were determined by an Agilent 2100 Bioanalyzer (Agilent Technologies, USA) according to the manufacturer’s instruction (Figure S2). Aliquots of total RNA were kept for quantitative real-time PCR (qRT-PCR), and the remainder (≥10 µg) was sent to Beijing Genomics Institute (BGI) (http://www.genomics.cn, China) on dry ice for mRNA enrichment by using the Ribo-Zero™ Magnetic Kit for Bacteria (Epicenter, USA).

**Strand-Specific Library Construction, Illumina Sequencing, and Data Analysis**

RNA fragmentation, strand-specific cDNA synthesis (Parkhomchuk et al., 2009), library preparation, sequencing (2 × 90 paired-end reads on an Illumina HiSeq 2000 sequencer) and base calling were performed by BGI according to their in-house procedures (Liu L. et al., 2011). Four biological replicates were sequenced for PM, MM and PMC, respectively, yielding ≥25 million clean paired-end reads per replicate (Table S1). Reads were mapped to reference genomes using the function BWA-MEM of bwa program version 0.7.5a-r405 (Li and Durbin, 2009) with parameters: -t 4. Uniquely mapped reads were filtered by using samtools-0.1.19 (Li et al., 2009), with parameters: -f 0x2, -q 30, -F 0x100, NM ≤ 5. On average, 0.10, 6.74, and 4.98% of the total reads were mapped to rRNAs for PM, MM and PMC, respectively. These rRNA reads were excluded for further analysis (Table S1). To exclude reads that may map ambiguously (that is, to either genome), we did a cross reference mapping by using the parameter described above (Figure S3). Mapping of reads from PM and MM to the opposite genome showed only 0.62–1.42% and ~0.1% of the total reads from PM and MM, respectively (Table S1). These reads were filtered out (masked from downstream analysis). Unmapped reads from PM and MM were mapped to the genome of *P. thermopropionicum* (dataset PMp) and *M. conradii* (dataset MMm), respectively. For PMC, reads were mapped to the genome of either *M. conradii* or *P. thermopropionicum*. Reads unmapped to *M. conradii* were mapped to the genome *P. thermopropionicum* (dataset PMCp) and reads unmapped...
to the genome of *P. thermopropionicum* were mapped to the genome of *M. conradii* (dataset PMCm), respectively. Library strand specificity was checked by using RSeQC (Wang et al., 2012) (Table S2). Unique mapped reads were analyzed and assigned to individual CDS (coding DNA sequence) by HTSeq (Anders et al., 2015) according to the genome annotation provided by GenBank (*P. thermopropionicum*: NC_009454 and *M. conradii*: NC_017034). Following parameters set for HTSeq were: -m union, -nonunique = none, -t gene, -i locus_tag, -s reverse (strand specificity libraries, according to the results of RSeQC).

DESeq2 package (version 1.6.2) in R (R Core Team, 2016) was then used to assess differential expression across the transcriptome (Love et al., 2014). The function DESeq was called with design formula "~condition," in which monocyte conditions of either *M. conradii* or *P. thermopropionicum* were used as control groups and the coculture were used as treatment groups. DESeq2 performs a hypothesis test on each gene to see whether evidence is sufficient to decide against the null hypothesis that there is no effects of the coculture on the gene expression and that the observed difference between coculture and monoculture was merely caused by experimental variability. Only absolute log2-fold change values (L2fc) ≥ 1.0 with false discovery rate (FDR) < 0.05 in transcription were considered significant. Principal components analysis (PCA) was performed using the function "prcomp" in R using the variance stabilizing transformed data from DESeq2. “Fragments Per Kilobase of CDS (or exon) per Million reads” (FPKM) was calculated according to the below formula (Trapnell et al., 2010):

$$\text{FPKM} = \frac{\text{Raw counts} \times 10^9}{\text{(length of that CDS} \times \text{total number of mapped reads)}}$$

The Pearson correlation coefficients based on log2 FPKM value of each CDS were calculated, which showed that biological replicates within each group were reproducible (Table S3). Furthermore, we calculated the log2 value of mean FPKM (L2mFPKM) for each dataset, with the median of 6.9, 6.8, 6.7, and 6.3 for PMp, PMCp, MMm, and PMCm, respectively. The median was used as an indication of expression activity (Shrestha et al., 2013).

### qRT-PCR Verification of Transcriptomic Data

To verify the differential expression level detected by sequencing, qRT-PCR was performed for 6 differential expressed genes of *P. thermopropionicum* and *M. conradii*, respectively. Primers targeting the selected genes were designed by using Primer Premier 6 (Premier, Canada) and synthesized by Life Technologies (Shanghai, China) (Table S4). cDNA synthesis was carried out as previously described (Liu et al., 2014). qRT-PCR and data analysis was performed with the 7500 real-time PCR system (Applied Biosystems, USA) (Liu et al., 2014). The expression levels of 16S rRNA gene and DNA-directed RNA polymerase beta subunit encoding gene (*rpoC*) of *P. thermopropionicum* and 16S rRNA gene and DNA-directed RNA polymerase A subunit encoding gene (*rpoA1*) of *M. conradii* were chosen as the references, respectively.

### Retrieval of Microbial Signal Transduction System (STS) Related Genes

Genes related to signal transduction in *P. thermopropionicum* and *M. conradii* were retrieved from the microbial signal transduction database (MiST 2.2, [http://mistdb.com/](http://mistdb.com/)) (Ulrich and Zhulin, 2010).

### Nucleotide Sequence Accession Number

The sequence reads determined in this study have been submitted to the GEO databases under accession number GSE103596.

### RESULTS

#### Overview of RNA-Seq Transcriptomes

In monoculture, *P. thermopropionicum* on pyruvate and *M. conradii* on H2/CO2 grew with generation times of 6–7 and 7–8 h, respectively. In syntrophic coculture on propionate, generation times were in a range of 36–40 h (Figure S1). These generation times were consistent with those obtained in previous studies (Imachi et al., 2002; Lü and Lu, 2012; Liu et al., 2014).

Of all the sequenced RNA-Seq reads, 75.6 and 78.0% were mapped to coding DNA sequences (CDSs) of the corresponding genomes of all reads from PM and MM, respectively. Of the reads from PMC, 9.3 and 68.4% were mapped to *P. thermopropionicum* and *M. conradii* CDSs, respectively (Table S1). The remaining reads were mapped to genome regions without annotation or with reverse complemented directions (antisense) to CDSs (data not shown).

Of all the 2920 CDSs of *P. thermopropionicum* and 2455 CDSs of *M. conradii*, 676 (23%) and 363 (15%) were significantly upregulated and 374 (13%) and 361 (15%) were significantly downregulated in the coculture compared to the respective monocultures ([Figures 1A,B](#fig1){refig} and Supplementary Datasets Pt1, Mc1). Principal component analysis (PCA) analysis showed that the gene expression profiles were distinct between monoculture and coculture conditions ([Figures 1C,D](#fig1){refig}). Differential expression of selected genes determined by qRT-PCR correlated well with that by RNA-Seq (Pearson *r* > 0.99) and had a slope that approached one (Figure S4).

In order to evaluate global transcriptional responses of *P. thermopropionicum* and *M. conradii* to syntrophic growth, the numbers of differentially expressed genes were summarized as functional categories according to the clusters of orthologous groups of proteins (COGs) database (Tatusov et al., 2003) (Table I). For both *P. thermopropionicum* and *M. conradii*, the upregulated genes in coculture were mainly distributed in “Energy production and conversion,” “Cell wall/membrane/envelope biogenesis,” “Carbohydrate transport and metabolism,” and “Signal transduction mechanisms.” The genes downregulated in coculture were mainly categorized into “Amino acid transport and metabolism,” “Translation, ribosomal structure, and biogenesis,” “Replication, recombination, and repair,” “Lipid transport and metabolism,” and “Cell motility” were upregulated in coculture.
Metabolic Shifts in *P. thermopropionicum*

The large *mmc* gene cluster (PTH_1356-1369) in *P. thermopropionicum* consists of coding genes for propionyl-CoA: oxaloacetate transcarboxylase (Pot), methylmalonyl-CoA epimerase (Mce), methylmalonyl-CoA mutase (Mcm), succinyl-CoA synthase (Scs), fumarase (Fht), malate dehydrogenase (Mdh), and pyruvate: ferredoxin oxidoreductase (Por). The *mmc* cluster was upregulated more than 4-fold in coculture (Figure 2 and Supplementary Dataset Pt2). The other three key enzymes of the MMC pathway include acyl-CoA: acetate/3-ketoacid CoA transferase (Pct), succinate dehydrogenase (Sdh), and acetyl-CoA synthetase (Acs) (Supplementary Dataset Pt2). The genes coding for one Sdh (Sdh I) were upregulated similarly as the *mmc* cluster, while the expression of the second Sdh (Sdh II) was one order of magnitude lower and did not show much response. The coding genes for the two Pct (PTH_2042-2043 and PTH_2044-2045) were upregulated more than 16-fold (L2fc > 4) and the Acs coding gene (PTH_2131) was upregulated to a greater extent (L2fc > 7). For the conversion of pyruvate to acetyl-CoA, the gene coding for pyruvate formate lyase (Pfl)
TABLE 1 | Distribution of differential expressed genes in clusters of orthologous groups (COGs) categories.

| COG categories                        | P. thermopropionicum | M. conradii |
|---------------------------------------|----------------------|-------------|
|                                       | Total | Up\(^{a}\) | Down\(^{b}\) | Total | Up\(^{a}\) | Down\(^{b}\) |
| Energy production and conversion (C)  | 181   | 53\(^{c}\) | 33         | 173   | 29        | 15          |
| Translation, ribosomal structure and biogenesis (J) | 138   | 3        | 37\(^{c}\) | 163   | 5         | 86          |
| Amino acid transport and metabolism (E) | 183   | 30       | 47         | 147   | 14        | 29          |
| Coenzyme transport and metabolism (H) | 122   | 15       | 19         | 133   | 19        | 20          |
| Inorganic ion transport and metabolism (P) | 86    | 21       | 24         | 113   | 11        | 10          |
| Transcription (K)                     | 139   | 29       | 20         | 100   | 19        | 25          |
| Replication, recombination and repair (L) | 181   | 38       | 17         | 97    | 9         | 7           |
| Cell wall/membrane/envelope biogenesis (M) | 145   | 44       | 16         | 90    | 17        | 2           |
| Carbohydrate transport and metabolism (G) | 82    | 18       | 9          | 83    | 17        | 9           |
| Posttranslational modification, protein turnover, chaperones (O) | 78    | 17       | 11         | 83    | 8         | 12          |
| Signal transduction mechanisms (T)    | 135   | 32       | 9          | 66    | 22        | 7           |
| Nucleotide transport and metabolism (F) | 66    | 6        | 10         | 54    | 9         | 7           |
| Intracellular trafficking, secretion, and vesicular transport (U) | 63    | 11       | 6          | 42    | 9         | 9           |
| Cell cycle control, cell division, chromosome partitioning (Q) | 35    | 2        | 2          | 31    | 6         | 1           |
| Defense mechanisms (V)                | 39    | 3        | 9          | 31    | 5         | 2           |
| Lipid transport and metabolism (I)    | 71    | 22       | 5          | 22    | 3         | 9           |
| Cell motility (N)                     | 65    | 13       | 2          | 18    | 5         | 8           |
| Cytoskeleton (Z)                      | n.a.  | n.a.     | n.a.       | 18    | 4         | 0           |
| Secondary metabolites biosynthesis, transport and catabolism (Q) | 26    | 6        | 3          | 5     | 0         | 2           |
| Chromatin structure and dynamics (B)  | 3     | 1        | 0          | 3     | 0         | 2           |
| RNA processing and modification (A)   | n.a.  | n.a.     | n.a.       | 2     | 2         | 0           |
| General function prediction only (R)  | 250   | 46       | 29         | 308   | 47        | 30          |
| Function unknown (S)                  | 191   | 38       | 29         | 249   | 35        | 34          |

\(^{a}\)Log₂ fold change value (L2fc) ≥ 1 and false discovery rate (FDR) < 0.05.

\(^{b}\)L2fc ≤ −1 and FDR < 0.05.

\(^{c}\)COG categories mentioned in the main text are shown in bold and red and blue colors depict that corresponding categories are featured by certain number of up or down regulated genes.

was also upregulated, but the total transcript abundance was much lower than Por coding genes within the mmc cluster. Other enzymes related to propionate oxidation, like oxaloacetate decarboxylase (Odc) showed low abundances and the gene coding for malic enzyme (Sfc) was downregulated (Figure 2 and Supplementary Dataset Pt2).

P. thermopropionicum contains four hydrogenases, i.e., one membrane bound NiFe hydrogenase (NiFe-H₂ase), one membrane-bound Fe-hydrogenase (Fe-H₂ase I) and two cytoplasmic Fe-only hydrogenases (Fe-H₂ases II and III); and two formate dehydrogenases, i.e., a membrane bound formate dehydrogenases (Fdh I) and a cytoplasmic one (Fdh II) (Kosaka et al., 2008). The genes coding for Fe-H₂ase III (PTH_2010-2012) and Fdh II (PTH_2645-2649) were significantly upregulated (L2fc > 1.2 for Fe-H₂ase III and L2fc > 4.2 for Fdh II, respectively) (Figure 2, Table 2, and Supplementary Dataset Pt2). The membrane bound Fdh I did not show significant upregulation, while the other three hydrogenases showed low transcript abundances and were moderately downregulated (Figure 2 and Supplementary Dataset Pt2). Multiple sets of putative gene clusters are present in P. thermopropionicum coding for electron transfer flavoproteins (ETF) that are homologous to the Fix complex in nitrogen-fixing microorganisms. This complex comprises two subunits of the electron transfer flavoprotein (FixA and FixB), one ferredoxin-like protein (FixX), and one membrane-bound ETF-menadione oxidoreductase (FixC). Most of the genes of the Fix system were significantly upregulated in coculture (Figure 2, Table 2, and Supplementary Dataset Pt2).

P. thermopropionicum uses intermediates of the MMC pathway, like 2-oxaloacetate, succinyl-CoA and pyruvate as building bricks for biosynthesis. The transcription of the related genes was generally downregulated in coculture (Figure 2). The transcription of genes for enzymes catalyzing early steps of the glycolysis pathway were also downregulated (Figure S5). P. thermopropionicum relies on the assimilation of ammonium and amino acids as nitrogen sources. Notably, the ammonium permease encoding gene (amtB) showed very low transcription (L2mFPKM~ = 2). While several ABC-type branched-chain amino acid transport systems were actively transcribed and upregulated (e.g., PTH_0073-0077 and PTH_0458-0462) in coculture (Figure 2 and Supplementary Dataset Pt3). Glutamate synthase coding genes were downregulated, though the expression of the glutamate dehydrogenase gene (gdhA) with
FIGURE 2 | Conceptual metabolism scheme of *P. thermopropionicum* and *M. conradii* syntrophic coculture, highlighting the central energy conservation and early steps of biosynthesis pathways for both organisms. Enzymes catalyzing each step are shown in bold. Prefix omitted locus tags are shown for steps without a specific enzyme abbreviation. Significant changes (See Supplementary Dataset for details) in transcripts abundance during syntrophic growth are shown by red (up) and blue (down) coloration. Black coloration indicates statistically insignificant change. Monoculture of *P. thermopropionicum* and *M. conradii* served as control growth conditions. Enzymes encoded by genes with transcripts abundance lower than the median (as specified in the Materials and Methods section) of both monoculture and coculture conditions are underscored. *Mvh2* and *Hdr2* of *M. conradii* are also shown in red since coding genes for them and *Fwd* are in the same gene cluster (See Figure S9). Green arrows indicate intermediates for biosynthesis. Black and gray dash arrows indicate electron transfer and missing steps in the genome, respectively. Putative bifurcation and confurcation mediating complexes are shown in blue cycles. Abbreviation of enzyme names of *P. thermopropionicum* shown in the scheme: Pot, propionyl-CoA: oxaloacetate transcarboxylase; Mce, methylmalonyl-CoA epimerase; Mcm, methylmalonyl-CoA mutase; Scs, succinyl-CoA synthase; Fht, fumarase; Mdh, malate dehydrogenase; Por, pyruvate: ferredoxin oxidoreductase. Coding genes for above enzymes are assembled in the *mmc* gene cluster (PTH_1356-1369). Pct, acyl CoA: acetate/3-ketoacid CoA transferase; Sdh, succinate dehydrogenase (I & II); Acs, acetyl-CoA synthetases/AMP-(fatty) acid ligases; Odc, oxaloacetate decarboxylase; Plt, pyrurate formate lyase; Fe-H$_2$ase, Fe-hydrogenase (I, II & III); NiFe-H$_2$ase, NiFe-hydrogenase; Fdh, formate dehydrogenase (I & II); Fix complex, containing two subunits of the electron transfer flavoprotein (FixA and FixB), one ferrodox-like protein (FixO), and one membrane-bound ETF-mercaptoquinone oxidoreductase (FeC); AmtB, ammonia permease; GhaA, glutamine synthetase; AA, amino acid; MQ, Menaquinone; here was used to simplify the Menaquinone cycle.
### TABLE 2 | Transcripts abundance and differential expression of genes coding for putative flavin-based confurcation and bifurcation complexes in *P. thermopropionicum* and *M. conradii*.

| Enzyme | Gene ID | Gene | L2fc<sup>a</sup> | L2mFPKM | Function |
|--------|---------|------|-----------------|----------|-----------|
|        |         |      | Monoculture | Coculture |           |
|        |         |      | *P. thermopropionicum* | | |
| Fe-hydrogenase (Fe-H₂ase III) | PTH_2010 | | 1.49 | 12.05 | 13.51 | Hypothetical hydrogenase subunit |
| | PTH_2011 | nuoF | 1.19 | 11.62 | 12.79 | NADH: ubiquinone oxidoreductase, NADH-binding 51 kD subunit |
| | PTH_2012 | nuoE | 1.68 | 11.23 | 12.88 | NADH: ubiquinone oxidoreductase, 24 kD subunit |
| Formate dehydrogenase (Fdh II) | PTH_2645 | | 4.26 | 7.20 | 11.53 | Hypothetical formate dehydrogenase |
| | PTH_2646 | | 4.42 | 7.06 | 11.57 | Hypothetical formate dehydrogenase |
| | PTH_2647 | | 4.45 | 7.00 | 11.53 | Hypothetical membrane protein |
| | PTH_2648 | nuoF | 4.59 | 6.90 | 11.56 | NADH: ubiquinone oxidoreductase, NADH-binding 51 kD subunit |
| Putative Fix complex<sup>b</sup> | PTH_0016 | fixA | 1.57 | 1.61 | 3.10 | Electron transfer flavoprotein, beta subunit |
| | PTH_0017 | fixB | 1.76 | 3.97 | 5.71 | Electron transfer flavoprotein, alpha subunit |
| | PTH_0018 | fixC | 4.55 | 4.61 | 9.21 | Dehydrogenases |
| | PTH_0019 | fixX | 2.06 | 4.40 | 6.45 | Ferredoxin-like protein |
| Putative Fix complex II<sup>b</sup> | PTH_0597 | fixB | 3.04 | 1.03 | 4.42 | Electron transfer flavoprotein, alpha subunit |
| | PTH_0598 | fixC | 2.33 | 1.68 | 4.15 | Dehydrogenases |
| | PTH_0599 | fixX | 2.47 | 2.15 | 4.79 | Ferredoxin-like protein |
| Putative Fix complex III<sup>b</sup> | PTH_1765 | fixX | n.a. | n.a. | n.a. | Ferredoxin-like protein |
| | PTH_1766 | fixC | 2.26 | 1.26 | 3.52 | Dehydrogenases |
| | PTH_1767 | fixB | 1.70 | 1.39 | 3.10 | Electron transfer flavoprotein, alpha subunit |
| | PTH_1768 | fixA | 2.35 | 0.69 | 3.09 | Electron transfer flavoprotein, beta subunit |
| **M. conradii** | | | | | |
| Fwd-Mvh-Hdr complex<sup>d</sup> | Mtc_2468 | fwdC | 0.81 | 11.70 | 12.08 | Tungsten-containing formylmethanofuran dehydrogenase, subunit C |
| | Mtc_2469 | fwdA | 1.31 | 11.91 | 12.79 | Tungsten-containing formylmethanofuran dehydrogenase, subunit A |
| | Mtc_2470 | fwdB | 1.02 | 11.25 | 11.84 | Tungsten-containing formylmethanofuran dehydrogenase subunit B |
| | Mtc_2471 | fwdD | 0.89 | 11.29 | 11.74 | Tungsten-containing formylmethanofuran dehydrogenase subunit D |
| | Mtc_2476 | fwdF | 0.59 | 10.17 | 10.32 | Tungsten-containing formylmethanofuran dehydrogenase, subunit F |
| | Mtc_2477 | fwdG | 0.08 | 9.36 | 8.99 | Tungsten-containing formylmethanofuran dehydrogenase subunit G |
| | Mtc_2472 | mvhD-2 | 0.95 | 11.58 | 12.09 | F₄₂₀-non-reducing hydrogenase subunit D |
| | Mtc_2473 | hdaA-2 | 0.85 | 11.16 | 11.56 | CoB-S-S-CoM heterodisulfide reductase subunit A |
| | Mtc_2474 | hdb-2 | 0.55 | 10.87 | 10.98 | CoB-S-S-CoM heterodisulfide reductase, subunit B |
| | Mtc_2475 | hdc-2 | 0.75 | 11.05 | 11.35 | CoB-S-S-CoM heterodisulfide reductase, subunit C |
| Formate dehydrogenase (Fdh)<sup>e</sup> | Mtc_2124 | fdhB | 2.81 | 10.23 | 12.80 | F₄₂₀-reducing formate dehydrogenase, beta subunit |
| | Mtc_2125 | fdhA | 2.76 | 10.24 | 12.55 | F₄₂₀-reducing formate dehydrogenase, alpha subunit |

L2fc, log₂-fold change value; L2mFPKM, log₂ value of mean FPKM (with 4 replicates); n.a., not applicable.

<sup>a</sup>Differential expression tested by DESeq2 based on raw read counts. Only genes with absolute L2fc ≥ 1.0 and false discovery rate (FDR) < 0.05 were considered to be significantly regulated. Since genes listed here all with FDR < 0.05 except PTH_1765 (See below), FDR are not shown (See Supplementary Datasets Pt2, Mc2 for details).

<sup>b</sup>For the Fix complex encoding genes, only putative gene clusters with all four fix genes (fixA, fixB, fixC, and fixX) are shown (See Supplementary Dataset Pt2 for more details).

<sup>c</sup>The transcripts of PTH_1765 is not detected under current sequencing depth.

<sup>d</sup>Shown is the large transcript unit which codes for Fwd, Hdr2, and the subunit D of Mvh2. Proteins may eventually form Fwd-Hdr-Mvh complex for electron bifurcation (See also Figure 2, Figure S7, and Supplementary Dataset Mc2).

<sup>e</sup>Genes coding for Fdh in M. conradii do not form transcript cluster with Hdr coding genes. But Fdh-Hdr complex is likely to form for electron bifurcation in methanogens according to Costa et al. (2010).
low abundance apparently increased in coculture (Figure 2 and Supplementary Dataset Pt3). Four of glutamine synthetase encoding genes (glnA) were downregulated. The transcription of other amino acids synthetic pathways were downregulated in most cases (Figure S6).

Metabolic Shifts in M. conradii
The methanogenesis pathway dominated the transcriptomes of M. conradii (Supplementary Dataset Mc2). The methyl-CoM reductase (Mcr) coding genes showed the highest transcription followed by methylene-tetrahydromethanopterin (H₂MPT) dehydrogenase (Mtd), methylene-H₂MPT reductase (Mer) and methyl-H₂MPT-CoM methyltranferase (Mtr). The Mcr and Mer coding genes were significantly upregulated. The genes coding for formylmethanofuran dehydrogenase (Fwd), heterodisulde reductase (Hdr) and F₄₂₀-non-reducing hydrogenase (Mvh) form a large gene cluster (Mtc_2468-2479) (Liu et al., 2014; Lyu and Lu, 2015). This cluster was moderately to significantly upregulated (Figure 2, Table 2, and Supplementary Dataset Mc2). The genes coding for formylmethanofuran-H₂MPT formyltransferase (Ftr) and methenyl-H₂MPT cyclohydrolase (Mch) were not affected.

The M. conradii genome encodes four types of canonical hydrogenases (Frh, Ech, Mvh1, Mvh2), plus one novel hydrogenase (Drh) that is not found in other methanogens and one F₄₂₀-reducing formate dehydrogenase (Fdh). Apart from Mvh mentioned above the putative membrane-associated novel hydrogenase (Drh) was upregulated while the transcription of F₄₂₀-reducing hydrogenase (Frh) and energy-converting hydrogenase (Ech) did not show an explicit response. Conversely, the transcription of Fdh and formate transporter coding genes were upregulated over 6-fold (L2fc > 2.7) in coculture (Figure 2, Table 2, and Supplementary Dataset Mc2). For ATP synthesis, two of four ATP synthases (Atp1 and Atp11) showed high transcript abundances but no response to coculture conditions, while the other two (Atp2 and Atp12) with low abundances were significantly upregulated (Figure 2 and Supplementary Dataset Mc2).

M. conradii employs the AMP-forming acetyl-CoA synthetase (Acs) and pyruvate: ferredoxin oxidoreductase (Por) for acetate assimilation and pyruvate conversion. Acs1 and Acs2 were downregulated 3- to 16-fold in coculture (Figure 2 and Supplementary Dataset Mc3). The transcription of inorganic pyrophosphatase (Ppa) coding gene (ppa) and Por coding genes also declined but not significantly. One ADP-forming acetyl-CoA synthetase (Acd) coding gene was significantly downregulated. The transcript abundances of pyruvate dehydrogenase (Pdh) coding genes were far lower compared with other carbon assimilation genes. Manual curation revealed that Methanocella strains contained citrate synthase and aconitate, indicating the presence of a partial oxidative TCA cycle (Lyu and Lu, 2015). Some of aconitase (Mtc_1230-1231) and isocitrate dehydrogenase (Icd) coding genes were downregulated (Figure 2 and Supplementary Dataset Mc3). A novel citrate synthase encoding gene (Mtc_1389) was actively transcribed but remained unaffected (Figure 2 and Supplementary Dataset Mc3).

Most of the genes involved in nitrogen assimilation system in M. conradii were repressed in coculture (Figure 2 and Supplementary Dataset Mc4). The transcription of genes coding for ammonium transporter (Amt) and ABC type amino acid transporter (GluQ) was downregulated 3-fold in coculture. M. conradii relies on glutamate and glutamine system for ammonium assimilation. Genes coding for glutamate synthase (Glt) and glutamine synthetase (Gln) were downregulated up to 100-fold in coculture (Figure 2 and Supplementary Dataset Mc4). Although ammonium was present in the culture medium, the transcripts of nitrogenase coding genes were detected. Their transcription however was either unaffected (nifH2) or downregulated (nifH2DKENO) in coculture (Figure 2 and Supplementary Dataset Mc4).

Genes for Signal Transduction System and Flagellum
P. thermopropionicum and M. conradii contain 218 and 101 genes, respectively, coding for proteins involved in signal transduction systems (MiST 2.2, http://mistdb.com/). In P. thermopropionicum, the transcription of 50 one-component system genes (29 up and 21 down) and 28 two-component system genes (23 up and 5 down) showed explicit shifts in coculture (Supplementary Dataset Pt4). Some of these genes were among those which were most differentially expressed. For example, a gene coding for the response regulator OmpR (PTH_1670) was upregulated 88-fold (L2fc = 6.47) in coculture compared to monoculture (Supplementary Dataset Pt4). In addition, the transcription of two genes (flIC and flID) coding for flagella were upregulated significantly in coculture (Figure 2 and Supplementary Dataset Pt5). In M. conradii, the expression of 24 one-component system genes (13 up and 11 down) and 16 two-component system genes (11 up and 5 down) changed significantly in coculture (Supplementary Dataset Mc5).

The Effects of Sodium 2-bromoethanesulphonate (2-BES), H₂, and Formate on Coculture
The contribution of H₂ vs. formate for interspecies electron transfer remains elusive (de Bok et al., 2004; Schink et al., 2017). Examination of the inhibitory effects of H₂ and formate has been shown to be an effective way in discerning their contribution (Ahring and Westermann, 1988; Dong et al., 1994). Therefore, 2-BES (5 mM), H₂/CO₂ (80/20, v/v, 170 kPa) and formate (20 mM) were added separately or in combination after the initiation of propionate oxidation and CH₄ production (Figure 3). Methane production and propionate oxidation were completely inhibited after the addition of 2-BES. Hydrogen accumulated to about 55 Pa in the end (Figure 3A), but formate was below the detection limit (∼50 μM). The addition of formate halted propionate oxidation and CH₄ production immediately (Figure 3B). But the added formate was consumed (to CH₄) within 4 days and the oxidation of propionate resumed 10 days after formate consumption. Hydrogen accumulated to 45 Pa at day 25 and then leveled off. Notably, the addition of H₂/CO₂ (80/20, v/v) did not significantly affect propionate oxidation.
Methane production, propionate oxidation, acetate accumulation and H₂ utilization occurred concomitantly. Addition of both formate and H₂/CO₂ (80/20, v/v) inhibited the oxidation of propionate for about 4 days (Figure 3D). In this experiment, formate was converted to CH₄ within 1 day after addition of formate and the resumption of propionate oxidation was faster than in the treatment with formate addition alone.

DISCUSSION
Catabolism vs. Anabolism Shifts in P. thermopropionicum and M. conradii
A large proportion of genes in P. thermopropionicum and M. conradii were differentially transcribed in coculture compared to monoculture. Distinct transcriptome profiles revealed by PCA ordination showed that global metabolism responses occurred in both P. thermopropionicum and M. conradii under syntrophic coculture condition (Figure 1). It is worth noting that conditions in monoculture are not identical to those in coculture, which can hardly be avoided, since the different microbial communities affect the conditions. The factors (e.g., H₂ partial pressure, interspecies interactions, the type and concentration of substrate) that may be important for gene expression are to some extent the result of the different organismic compositions, and should be carefully evaluated. However, comparison of gene expression between monoculture and coculture is certainly warranted.

Genes coding for the entire MMC pathway were significantly upregulated in coculture. Although bypaths are present for the conversion of malate to pyruvate (Figure 2), the transcript abundances indicated that P. thermopropionicum used malate dehydrogenase and propionyl-CoA: oxaloacetate transcarboxylase instead of malic enzyme and oxaloacetate decarboxylase, consistent with a previous report (Kato et al., 2009). Succinate dehydrogenase (Sdh) catalizes the thermodynamically most endergonic step in propionate oxidation (Schink, 1997). Due to the low redox potential of the electron acceptor, the oxidation of succinate to fumarate must involve reverse electron transport that is probably driven by proton motive force (PMF) (Müller et al., 2010; Schink et al., 2017). PMF is likely formed by ATPase. Unlike the...
MMC pathway, transcription of the ATPase encoding gene in *P. thermopropionicum* did not significantly respond to coculture conditions. Our results are consistent with previous proteomic analyses, in which subunits of ATPase were major spots in the gel but showed less response to syntrophic propionate coculture conditions than the protein spots of the MMC pathway catalyzing enzymes (Kosaka et al., 2006). We assume that the insignificant response of ATPase implies a strategy for avoiding the waste of PMF or the slowdown of growth under coculture condition. The enzyme Pct is proposed to catalyze the formation of propionyl-CoA from propionate by coupling the conversion of acetyl-CoA to acetate, i.e., the first and the last step of the MMC pathway, respectively (Kosaka et al., 2006, 2008) (Figure 2). The enzyme Acs has been proposed to also catalyze the hydrolysis of acetyl-CoA, coupled to ATP formation (the last step of MMC pathway) (Kosaka et al., 2006) (Figure 2). Notably, the expression of Pct and Acs genes was upregulated to greater extents than that of the mmc cluster, especially the Acs gene showing the largest upregulation. High pyrophosphate levels and a high AMP-to-ATP ratio in syntrophic bacterium *Syntrophus aciditrophicus* cells support the operation of Acs in the direction of acetate and ATP forming (James et al., 2016). If similar high AMP-to-ATP ratio was maintained by *P. thermopropionicum*, then the activity of Acs together with succinyl-CoA synthase (Scs) probably supported CoA recycling and ATP generation via substrate-level phosphorylation. Otherwise, Acs would operate in the direction of acetate-CoA formation, securing the pool of acetyl-CoA. The upregulation of the first and last steps could help securing the activity of the MMC pathway and the conservation of energy under coculture condition. In contrast, most of the genes involved in biosynthetic processes were downregulated (Figure 2 and Figures S5, S6). This downregulation coincided with a reduced growth rate of *P. thermopropionicum* in coculture (Figure S1).

Similar to *P. thermopropionicum*, *M. conradii* tended to upregulate catabolism in coculture compared to monoculture. Genes coding for Mer and Mcr were not only upregulated but exhibited the highest transcript abundances (Figure 2 and Supplementary Dataset Mc2). Notably, FdhII-reducing formate dehydrogenase (Fdh) but not FdhI-reducing hydrogenase (Fhr) was significantly upregulated (Figure 2, Table 2, and Supplementary Dataset Mc2). This result possibly indicates that formate plays an important role (see more discussion below). The concomitant upregulation of Fdh and Mer activity indicates that larger proportions of electron flow were channeled into methanogenesis pathway in coculture compared to monoculture. Another route for electron flow is the putative Fwd-Mvh-Hdr complex, which was proposed to connect the first and last steps of the methanogenesis pathway and mediates electron bifurcation (Liu et al., 2014). The genes coding for this route were significantly upregulated (Figure 2, Table 2, and Supplementary Dataset Mc2). In addition, genes coding for two ATPases (Atp12 and Atp2) were upregulated. ATPases are proposed to produce ATP using the Na⁺/H⁺ motive force generated from Mtr activity (Lyu and Lu, 2015). Thus, under coculture condition *M. conradii* enhanced several key steps for methanogenic energy formation, i.e., the methyl CoM reductase (Mcr), two routes of electron flow and ATPase. By contrast, most of the genes associated with carbon assimilation and nitrogen metabolisms were significantly downregulated in coculture (Figure 2, Figures S7, S8).

Taken together, both *P. thermopropionicum* and *M. conradii* enhanced catabolism but decreased anabolism under syntrophic conditions. It is believed that growth rate affects gene expression in both bacteria and archaea. Therefore, the responses observed may include the effect of growth rate in addition to syntrophy. Because the growth rate of both organisms was lower under coculture than monoculture conditions, this kind of metabolic shift probably reflects an universal strategy for microbes to survive environmental stress or harsh conditions (Goodchild et al., 2005; Kato et al., 2008, 2009; Enoki et al., 2011).

### Flavin-Based Electron Bifurcation/Confurcation

Under conditions of thermodynamic limitation the ability to maximize energy conservation should be an advantage for the growth of syntrophic partners (Sieber et al., 2012). Flavin-based electron bifurcation/confurcation that couples exergonic with endergonic redox reactions has been proposed as the third form of energy conservation in microbial life (Herrmann et al., 2008; Buckel and Thauer, 2018). This process seems to be widespread in anaerobic microorganisms and offers a solution for their energetic dilemma (Buckel and Thauer, 2013; Peters et al., 2016). Three putative electron bifurcation/confurcation systems are found in *P. thermopropionicum*. First, the cytoplasmic Fe-H₂ase III contains NADH dehydrogenase and shows sequence homology with the electron confurcating Hyd complex of *Thermotoga maritima*. Therefore, it could be a putative site for electron confurcation (Sieber et al., 2010, 2012). Second, the genes coding for catalytic subunits of Fdh II are associated with genes coding for NADH-quinone oxidoreductase, thus suggesting that this enzyme could catalyze electron confurcation. Thus, H₂ and formate could be produced by coupling the reoxidation of NADH with the reduction of ferredoxin (Fdred) by Fe-H₂ase III and Fdh II, respectively. Notably, both Fe-H₂ase III and Fdh II were actively transcribed and significantly upregulated (Figure 2 and Table 2). Third, the Fix complex has been shown to perform electron bifurcation in the N₂-fixing *Azotobacter vinelandii* for the production of Fdred from NADH (Ledbetter et al., 2017). Most of Fix encoding genes were markedly upregulated in coculture (Figure 2 and Table 2). The Fix system was also found in the genome of butyrate-oxidizing *S. wolfei* and was proposed to produce Fdred, which is needed for the biosynthesis of hydrogen and/or formate from NADH (Sieber et al., 2010). However, the necessity of electron bifurcation/confurcation in *S. wolfei* has been questioned recently, since the beta oxidation pathway in butyrate degradation does not involve Fdred production, and since the H₂ partial pressure under syntrophic conditions was sufficiently low for direct reoxidation of NADH (Losey et al., 2017). However, for propionate oxidation in *P. thermopropionicum* where pyruvate: ferredoxin oxidoreductase coding genes were actively transcribed and upregulated under coculture condition (Figure 2), production of Fdred from pyruvate oxidation...
and coupling of NADH oxidation with Fd_red reduction is quite likely. The activity of the Fix enzyme may play an anaplerotic role in maintaining NADH and Fd_red equilibrium during propionate oxidation and/or supporting the needs of biosynthesis. Further investigations are necessary to elucidate why genes for all putative electron bifurcation/confurcation systems (Fe-H2ase III, Fdh II, and Fix) are upregulated in \textit{P. thermopropionicum} during syntrophic growth.

In hydrogenotrophic methanogens, Mvh and Hdr proteins form a complex that performs flavin-based electron bifurcation to simultaneously reduce CoM-S-S-CoB heterodisulfide and ferredoxin, which is required for the reduction of CO2 to formylmethanofuran (Kaster et al., 2011; Wagner et al., 2017). This coupling mechanism together with the anaplerotic reaction catalyzed by energy converting hydrogenase (Lie et al., 2012) completes the reduction of CO2 to methane in a full cycle (i.e., the Wolfe cycle), with ferredoxin coupling the first and last steps (Kaster et al., 2011; Thauer, 2012). The presence of a multifunctional supercomplex in \textit{Methanococcus maripaludis}, comprising not only Hdr and Vhu (equivalent to Mvh in \textit{M. conradii}) but also Fwd and Fdh, indicates that ferredoxin pools are probably compartmentalized and formate dehydrogenase is directly involved in this process (Costa et al., 2010; Thauer, 2012). We found that the transcription of Fdh coding genes was upregulated under coculture condition to a significantly greater extent than that of Mvh genes. Therefore, it is probable that formate donates electrons to the electron bifurcation Mvh-Hdr complex via Fdh in \textit{M. conradii}, similarly as in \textit{M. maripaludis}. The \textit{M. conradii} genome predicts the presence of a large transcript unit comprising genes coding for Hdr, Fwd and a subunit of Mvh (Lyu and Lu, 2015). The transcriptome analysis disclosed that this cluster was transcribed indeed as a single unit in \textit{M. conradii} (Figure S9 and Table S5) and was significantly upregulated in coculture (Figure 2 and Table 2). Comparative genomics indicates that the large transcript unit comprising Hdr and Fwd is only present in \textit{Methanocella} strains among all methanogens known to date (Lyu and Lu, 2015). This feature may make \textit{Methanocella} unique for constituting the Wolfe cycle and assembling its bifurcating multi-enzyme complex, and thus optimize energy conservation. Optimized energy conservation in \textit{Methanocella} may be the reason for the extraordinary adaption to H2-limiting conditions (Lu et al., 2005) and the ability to outcompete other methanogens for forming a syntrophic relationship with fatty acid oxidizing bacteria (Sakai et al., 2007; Liu P. et al., 2011; Rui et al., 2011; Gan et al., 2012).

**Formate as a Mediator for Interspecies Electron Transfer**

It has been assumed that \textit{P. thermopropionicum} is not likely to use formate as electron mediator for syntrophic growth with methanogens (Imachi et al., 2002). Indeed, \textit{M. conradii} in pure culture cannot grow on formate alone (Lü and Lu, 2012). In the present experiment, however, we found that the cytoplasmic Fdh II in \textit{P. thermopropionicum} was upregulated to a significantly greater extent than any of its cytoplasmic Fe-H2ases. Similarly, the membrane bound Fdh I was also upregulated (albeit not significantly) while its H2ase counterparts appeared to be downregulated under syntrophic condition. Coincidently, Fdh in \textit{M. conradii} was upregulated in coculture to a greater extent than both F420-reducing (Frh) and F420-non-reducing hydrogenases (Mvh). Our experiments showed that the addition of formate to an active coculture suppressed the syntrophic activity immediately (Figure 3B). It took a substantial time lag before the coculture resumed its activity even though formate was already consumed. On the contrary, the addition of H2 instead of formate resulted in very little effect (Figure 3C). Notably, the addition of H2 together with formate caused inhibition but not as significant as formate alone. H2 did not result in an additional negative effect but instead accelerated the consumption of formate by \textit{M. conradii} and pushed the recovery of syntrophic activity (Figure 3D). Taken together, our study suggests that formate played an important role in interspecies electron transfer between \textit{P. thermopropionicum} and \textit{M. conradii}. This conclusion is supported by our previous observation that the expression of a gene coding for the alpha subunit of Fdh in \textit{M. conradii} is upregulated under syntrophic conditions and \textit{M. conradii}, albeit not using formate alone, used formate in the presence of H2 (Liu et al., 2014). A previous study showed that H2 was essential for \textit{M. maripaludis} to grow on formate where H2 was either produced internally (from formate) or supplied externally (Lie et al., 2012). This essentiality was considered to be associated with the need for the activity of a membrane bound energy-converting hydrogenase (Eha). Whether a similar mechanism works in \textit{M. conradii} deserves further investigations.

Despite the importance of formate, a role of H2 for interspecies electron transfer cannot be ruled out. Genes coding for hydrogenases in both \textit{P. thermopropionicum} (Fe-H2ase III in particular) and \textit{M. conradii} (Mvh and Frh) were actively transcribed. Production and consumption of H2 occurred simultaneously during syntrophic growth (Figure 3). Syntrophic growth was not inhibited by high partial pressure of H2 might be due to diffusion limits under static culture conditions. We assume that optimal syntrophic growth requires both formate and H2 for interspecies electron transfer in coculture. The necessity of formate together with H2 was also demonstrated in \textit{Syntrophobacter fumaroxidans} (Dong et al., 1994; Dong and Stams, 1995; Worm et al., 2011a,b; Sedano-Núñez et al., 2018).

**Other Possible Mechanisms**

Additional mechanisms, such as signal transduction by flagella or amino acid transport, may be important for syntrophic interactions between \textit{P. thermopropionicum} and \textit{M. conradii}. Flagella are considered allowing syntrophic bacteria to effectively interact with their methanogen partners (Ishii et al., 2006; Krumholz et al., 2015). The flagellar cap protein (Flid) apparently induces signal transduction between \textit{P. thermopropionicum} and \textit{M. thermautotrophicus} (Shimoyama et al., 2009). Our transcriptome analysis showed that transcription of flagellum biosynthesis genes (especially Flid coding gene) was markedly enhanced in coculture suggesting its implication for syntrophic growth.

Transcriptome analysis also showed that the abundance of ammonium transporters was exceptionally low while amino
acid transporter coding genes were upregulated, indicating that *P. thermopropionicum* may rely on amino acid uptake. This is consistent with proteomic analyses in which a subunit of the branched-chain amino acid ABC transporter was the major soluble protein under coculture conditions (Kosaka et al., 2006). The importance of amino acid transport for syntrophic conditions is also in line with the previous observation that *P. thermopropionicum* can grow syntrophically with *M. thermautotrophicus* in the absence of yeast extract but requires yeast extract in monoculture (Kato et al., 2009). Transfer of amino acids has also been proposed for a syntrophic coculture of *Desulfovibrio vulgaris* and *M. maripaludis* (Walker et al., 2012). Exchange of amino acids may be an evolutionarily optimizing strategy to reduce the biosynthetic burden while promoting the cooperative interactions between different bacteria in the microbiome (Mee et al., 2014). We hypothesize that amino acid transfer from *M. conradii* to *P. thermopropionicum* compensates the low free energy gain during propionate degradation.

CONCLUSIONS

Our coculture experiments and transcriptome analyses disclosed several aspects of coordinated metabolic shifts in both *P. thermopropionicum* and *M. conradii* to cope with syntrophic growth. Firstly, both organisms tended to decrease energy-consuming biosynthesis pathways and enhance energy-generating catabolic pathways. This strategy is in accordance with the reduced growth rate of both organisms in coculture compared to monoculture. Secondly, genes coding for flavin-based electron bifurcation/confurcation systems were substantially upregulated in *P. thermopropionicum* and *M. conradii* under coculture condition. Fdh-based multienzyme complex might be formed in addition to the Mvh-Hdr complex in *M. conradii* for electron bifurcation. This study shows the importance of electron bifurcation/confurcation for syntrophic partners to maximize their energy conservation. Thirdly, formate was found to act besides H₂ as an important mediator for interspecies electron transfer between the two syntrophic partners and possibly renders syntrophic propionate oxidation more efficient. Previously studies only revealed the utilization of formate by *M. conradii* in monoculture in the presence of H₂ and the capacity of *P. thermopropionicum* to produce formate for the recycling of redox mediators (e.g., NADH) during syntrophic propionate oxidation was unclear. The transcriptome analysis also indicated that additional mechanisms like the flagellum mediated signal transduction and amino acids exchange was important for syntrophic growth. Apparently, *P. thermopropionicum* and *M. conradii*, when growing syntrophically, are employing multiple mechanisms for optimizing their cooperation under energy limitation condition. Syntrophic interactions under in situ conditions, however, may be even more complicated. Redundant copies of functional genes like multiple cytoplasmic and membrane-bound hydrogenases are present in both *P. thermopropionicum* and *M. conradii* genomes. How these different gene products coordinate and work together under varying environmental conditions shall deserve further investigations.

AUTHOR CONTRIBUTIONS

PL and YL designed the experiment. PL performed the experiment and analyzed data. PL and YL wrote the main manuscript. PL and YL revised and approved the final version of the manuscript.

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SUPPLEMENTARY MATERIAL

The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fmicb.2018.01551/full#supplementary-material

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Conflict of Interest Statement: The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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