# CO<sub>2</sub>-fixing one-carbon metabolism in a cellulosedegrading bacterium *Clostridium thermocellum*

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Clostridium thermocellum can ferment cellulosic biomass to formate and other end products, including CO2. This organism lacks formate dehydrogenase (Fdh), which catalyzes the reduction of CO<sub>2</sub> to formate. However, feeding the bacterium <sup>13</sup>C-bicarbonate and cellobiose followed by NMR analysis showed the production of <sup>13</sup>C-formate in *C. thermocellum* culture, indicating the presence of an uncharacterized pathway capable of converting CO<sub>2</sub> to formate. Combining genomic and experimental data, we demonstrated that the conversion of CO<sub>2</sub> to formate serves as a CO<sub>2</sub> entry point into the reductive one-carbon (C1) metabolism, and internalizes CO<sub>2</sub> via two biochemical reactions: the reversed pyruvate:ferredoxin oxidoreductase (rPFOR), which incorporates CO2 using acetyl-CoA as a substrate and generates pyruvate, and pyruvate-formate lyase (PFL) converting pyruvate to formate and acetyl-CoA. We analyzed the labeling patterns of proteinogenic amino acids in individual deletions of all five putative PFOR mutants and in a PFL deletion mutant. We identified two enzymes acting as rPFOR, confirmed the dual activities of rPFOR and PFL crucial for CO<sub>2</sub> uptake, and provided physical evidence of a distinct in vivo "rPFOR-PFL shunt" to reduce CO2 to formate while circumventing the lack of Fdh. Such a pathway precedes CO<sub>2</sub> fixation via the reductive C1 metabolic pathway in C. thermocellum. These findings demonstrated the metabolic versatility of C. thermocellum, which is thought of as primarily a cellulosic heterotroph but is shown here to be endowed with the ability to fix CO<sub>2</sub> as well.

*Clostridium thermocellum* | pyruvate:ferredoxin oxidoreducase | formate | <sup>13</sup>C-isotopic tracing | one-carbon metabolism

he gram-positive *Clostridium thermocellum* is a thermophilic and strict anaerobic bacterium. It has gained a great amount of interest due to its cellulolytic abilities. By taking advantage of an extracellular cellulase system called the cellulosome (1), C. thermocellum can depolymerize cellulose into soluble oligosaccharides. The latter are further transported into the cells and fermented through its glycolytic pathway to pyruvate, the precursor to an array of fermentation products (e.g., H<sub>2</sub>, formate, lactate, acetate, ethanol, secreted amino acids) (2, 3). This capability makes C. thermocellum an attractive candidate for consolidated bioprocessing of lignocellulosic biomass, a process configuration that directly converts plant biomass into biofuels and chemicals without separate additions of enzymes (4). To date, there have been numerous works describing the molecular and genetic details of the cellulolytic system and fermentation pathways in C. thermocellum (5-9), whereas other metabolic characteristics of this species have not been adequately understood.

Currently, few details regarding inorganic carbon utilization are known for *C. thermocellum*. However, several investigators routinely add bicarbonate, a dissolved form of  $CO_2$ , into the medium to promote *C. thermocellum* growth (6, 10–12), implying its ability to use  $CO_2$ . This observation raises the questions of how  $CO_2$  plays a role in promoting *C. thermocellum* growth (13), and whether  $CO_2$  is incorporated into the metabolism. There is limited information on pathways relating to  $CO_2$  uptake thus far. One proposed pathway involving  $CO_2$  utilization in *C. thermocellum* is the malate shunt and its variant pathway (7, 14), in which  $CO_2$  is first incorporated by phosphoenolpyruvate carboxykinase (PEPCK) to form oxaloacetate and then is released either by malic enzyme or via oxaloacetate decarboxylase complex, yielding pyruvate. However, other pathways capable of carbon fixation may also exist but remain uncharacterized.

In recent years, isotopic tracer experiments followed by mass spectrometry (MS) (15, 16) or NMR measurements (17) have emerged as powerful tools for metabolic analysis. This methodology traces the transit of isotopic atoms throughout the biochemical network and sheds light on active reactions as well as molecular fluxes in vivo. Coupled with classic genetic manipulations, this technology has been successfully applied to provide insights into novel metabolic pathways in bacteria (18), yet it has not been used to study *C. thermocellum*.

In several species of *Clostridium*, the one-carbon (C1) metabolism is initiated from CO<sub>2</sub> reduction to become formate via formate dehydrogenase (Fdh) (19). This pathway provides C1 units for many crucial reactions, including amino acids and purine biosynthesis. Formate serves as a precursor to the folate biosynthesis pathway that forms the "methyl" branch of the Wood–Ljungdahl pathway, yielding acetyl-CoA from CO<sub>2</sub> (19). However, *C. thermocellum* lacks the first key enzyme, Fdh, based on genome information. In the present study, we investigate CO<sub>2</sub>-fixing pathway(s) in *C. thermocellum* involving C1 metabolism by using <sup>13</sup>C-tracer experiments. We

## Significance

High carbon yield in the bioengineering of heterotrophic bacteria is hindered by carbon loss to  $CO_2$  production. We provide evidence showing *Clostridium thermocellum*, a cellulosedegrading bacterium and a model consolidated bioprocessing organism, can fix  $CO_2$  while growing predominantly on cellobiose, a cellulose-derived disaccharide. By adding <sup>13</sup>C-bicarbonate to the bacterial culture and tracking <sup>13</sup>C-labeled metabolites, we discovered active reductive one-carbon (C1) metabolism in this bacterium. We further identified critical enzymes responsible for fixing  $CO_2$  and channeling the fixed carbon to the C1 metabolic pathway. Our findings pave the way to future engineering of this bacterium to use cellulose and  $CO_2$  simultaneously as a means to improve microbial carbon efficiency that is constrained by theoretical limitation and to reduce  $CO_2$  in the environment.



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measure the isotopomers of fermentation products and proteinogenic amino acids using NMR and GC/MS, and uncover a route *C. thermocellum* employs to fix  $CO_2$  when grown in a primarily heterotrophic mode supplemented with sodium bicarbonate.

#### Results

Effects of Bicarbonate on C. thermocellum Growth and Metabolism. To examine C. thermocellum's ability to use inorganic carbon, we compared cell growth and metabolic product formation during cellobiose fermentation with and without bicarbonate supplementation (20 mM). To eliminate any potential buffering effect exerted by bicarbonate, fermentation was performed in batch mode with pH controlled at 7.0 throughout. Upon consumption of all cellobiose, cultures with added bicarbonate yielded  $\sim 40\%$ more cell biomass than in its absence (Table 1). Without accounting for the excreted amino acids that have been observed in overflow metabolism (3), this result increased the possibility that extra total carbon output (i.e., cell biomass, lactate) could emerge from incorporation of the inorganic carbon (bicarbonate) as suggested by the calculated apparent carbon efficiency (65.7% without bicarbonate and 75.5% with bicarbonate; Table 1). We also observed higher apparent carbon efficiency in bicarbonate-fed cultures in non-pH-controlled fermentation (Table S1) buffered with 50 mM 3-(N-morpholino)propanesulfonic acid (MOPS).

Reduction of Bicarbonate to Formate as Revealed by NMR Analysis Using <sup>13</sup>C-Tracer. To track how bicarbonate could be metabolized by the bacterium, we fed C. thermocellum <sup>13</sup>C-bicarbonate (20 mM) and unlabeled cellobiose (14.6 mM) and analyzed the labeled fermentation products that were excreted into the medium with <sup>13</sup>C-NMR. Data from Fig. 1A show 1D <sup>13</sup>C spectra of the culturespent medium after cells were grown to late log phase. In the medium, we detected small peaks at 160.3 and 128.0 ppm (Fig. 1A, double asterisks), which correspond to residual bicarbonate and CO<sub>2</sub>, respectively (Fig. 1A, Top and Middle), and a signature formate carbonyl peak with a chemical shift at 171.1 ppm (Fig. 1A, *Middle* and *Bottom*). To confirm the formation of  $^{13}$ C-formate further, common fermentation products were also assessed for the assignment of the new peak at 171.1 ppm. Organic acids, including succinate, malate, butyrate, citrate, and isocitrate, have carbonyl carbons with chemical shifts that do not match the observed peak. One class of doubtable compounds is alpha-keto acids (i.e., pyruvate, 2-oxoglutarate), which have a carboxylic carbon linked to a keto group, appearing at around 170 ppm. To validate further, the <sup>1</sup>H decoupling was inactivated during signal acquisition. A one-bond carbon-hydrogen scalar coupling,  $J_{\rm HC}$ , of 195-Hz splits the peak at 171.1 ppm (Fig. 1B) due to a hydrogen directly bound to the carbonyl carbon. Pyruvate and 2-oxoglutarate, however, do not have directly bound <sup>1</sup>H on their respective carbonyl carbons, leading to the conclusion that the new peak at 171.1 ppm is indeed derived from formate. In addition to <sup>13</sup>C-formate, two small peaks at 182.5 and 174.4 ppm (Fig. 1A, marked with asterisks) correspond to other fermentation products that may also be derived from <sup>13</sup>C-bicarbonate labeling but in very small amounts. Other peaks from 5 to 65 ppm likely arose from natural abundance components in the defined media and did not change in relative intensity from sample to sample.

**Putative Reductive C1 Metabolism Based on Genome Annotation.** The above labeling experiments provided evidence of a significant CO<sub>2</sub>-to-formate reduction pathway that remains to be characterized. To identify genes that are putatively responsible for this conversion, we reconstructed the C1 metabolic pathway de novo, based on the genomic information of *C. thermocellum* DSM1313 strain (20). Its genome sequence suggests that *C. thermocellum* is capable of converting formate to methionine, serine, and pyruvate via the methyl branch of the Wood–Ljungdahl pathway (Fig. 2). However, it lacks Fdh, the enzyme microbes typically use to convert CO<sub>2</sub> to formate (Fig. 2).

Lacking the *fdh* gene, the only other gene encoding formate production found in C. thermocellum genome is pyruvate-formate lyase (PFL), catalyzing the conversion of pyruvate and CoA to acetyl-CoA and formate (21). We thereby hypothesize that a bypass consisting of two reactions catalyzed by pyruvate:ferredoxin oxidoreductase (PFOR) in the reversed direction and PFL may realize the CO<sub>2</sub>-to-formate conversion. Presumably, the reversed PFOR activity fixes CO<sub>2</sub> by catalyzing the formation of pyruvate using CO<sub>2</sub> and acetyl-CoA as cosubstrates. The fixed carbon atom from  $CO_2$  forms the carboxylic group of pyruvate, which was then split to make formate by PFL (Fig. 2), with the remaining two carbons in pyruvate making up the acetyl group in acetyl-CoA. Both *pfl* and *pfor* genes are annotated in the *C. thermocellum* genome. Genes for pyruvate formate lyase (pflB, clo1313\_1717) and its activating enzyme (pflA, clo1313\_1716) are adjacent to each other on the genome. Several putative genes encoding for PFOR (clo1313 0673, clo1313 0020-0023, clo1313 0382-0385, and clo1313\_1353-1356) were also annotated, hence making our hypothesis genetically feasible.

<sup>13</sup>C-Tracer Data Demonstrated the Reversed PFOR Activity. To investigate this hypothesized pathway, we first test if a reversed PFOR (rPFOR) activity exists in vivo. We set up a steady-state tracer experiment by feeding <sup>13</sup>C-bicarbonate (20 mM) along with unlabeled cellobiose (14.6 mM) to the cells and detecting the labeling pattern of proteinogenic amino acids by GC/MS. The isotopomer fingerprint of pyruvate can be traced by the isotopomer fingerprint of alanine, which is a direct transaminated product of pyruvate. As shown in Fig. 3A, M ala 085 is a GC/MS fragment of alanine and reveals the labeling pattern of a pyruvate fragment with carbon atoms at positions 2 and 3 (Pyr2-3). For Pyr2-3, a dominant peak at m0 accounted for 95% (n = 3, SE  $\leq 1\%$ ) of total isotopes, suggesting that the carbon atoms at positions 2 and 3 remained almost entirely unlabeled. M ala 057 is another MS fragment of alanine, representing the isotopomer pattern of pyruvate with all carbon atoms (Pyr1-3). The detected m0 and m1 signals accounted for 55% and 43% of total isotopes, respectively. indicating that a significant portion of this MS fragment either remained fully unlabeled or had only one <sup>13</sup>C-carbon (~43%). These results led us to conclude that the labeled carbon is predominantly at position 1 (carboxylic group) of pyruvate, which is consistent with the hypothesized rPFOR reaction to fix CO<sub>2</sub>

Table 1. Carbon input and output in pH-controlled bioreactors (pH 7.0) with and without bicarbonate

HCO <sub>3</sub> , 20 mM	Cellobiose consumed, mM	Lactate, mM	Formate, mM	Acetate, mM	Ethanol, mM	Final OD <sub>600</sub>	C <sub>biomass</sub> ,* mM	Apparent carbon efficiency <sup>†</sup>
With bicarbonate	17.2 ± 0.6	10.3 ± 1.1	3.1 ± 0.6	9.1 ± 0.4	19.9 ± 0.5	$\begin{array}{c} 1.4  \pm  0.2 \\ 1.0  \pm  0.3 \end{array}$	63.8 ± 8.8	75.5 ± 3.0%
Without bicarbonate	17.3 ± 0.3	8.3 ± 0.1	3.8 ± 1.2	11.1 ± 0.6	21.3 ± 0.4		42.9 ± 12.1	65.7 ± 5.5%

\*Carbon molarity in cell biomass ( $C_{biomass}$ ) is calculated based on the *C. thermocellum* formula  $C_{s}H_{8}O_{2}N$  and the measured  $OD_{600}$  values ( $OD_{600} = 1.04$  g/L cell dry weight).

<sup>†</sup>Apparent carbon efficiency is defined as the ratio of produced organic carbons to consumed organic carbons following the calculation: ( $C_{lactate} + C_{formate} + C_{acetate} + C_{ethanol} + C_{biomass}$ )/ $C_{cellobiose}$ . Data are representative biological replica (mean  $\pm$  SD, n = 2).



**Fig. 1.** <sup>13</sup>C-NMR analysis of supernatant from *C. thermocellum* (Ct) culture fed with 20 mM <sup>13</sup>C-bicarbonate and 14.6 mM unlabeled cellobiose. (A) <sup>13</sup>C-NMR signal at 171 ppm is consistent with a formate carbonyl carbon that is observed in the spent medium of *C. thermocellum* (red spectrum). Defined medium supplemented with 20 mM <sup>13</sup>C-bicarbonate (green spectrum) and 20 mM <sup>13</sup>C-formate (black spectrum) was analyzed as the references. The single asterisk indicates other products of bicarbonate metabolism. The double asterisks indicate a contaminant in the <sup>13</sup>C-bicarbonate. The signals within the braces indicate natural abundance of aliphatic and methyl carbons from the medium or metabolites. (*B*) *J*<sub>HC</sub> dipolar coupling of 195 Hz further confirms the identification of the peak at 171 ppm as formate.

(Fig. 3*B*) and strongly supports the existence of rPFOR activity in vivo under the tested physiological condition. We also detected <sup>13</sup>C-lactate with a labeling pattern consistent with the labeling pattern of pyruvate (Figs. 2 and 3), presumably propagated from pyruvate.

To confirm rPFOR activity as the main mechanism contributing to the labeled pyruvate (position 1), and not to the label exchange reaction, we fed *C. thermocellum* [2-<sup>13</sup>C]acetate (40 mM) supplemented with unlabeled bicarbonate (20 mM) when cells were grown primarily in cellobiose (5 g/L). Results showed labeled carbons at the expected position in pyruvate (position 3) and acetyl-CoA (position 2) in both wild type ( $\Delta hpt$ ; Fig. 3*C*) and PFL-inactivated mutant ( $\Delta hpt\Delta pflA$ ; Fig. S1). For confirmation and characterization of the  $\Delta hpt\Delta pflA$  mutant, see Fig. 5 and Figs. S2 and S5*A*. These results also ruled out the possibility of labeled pyruvate transiting from acetyl-CoA via reversed PFL activity. Results are described in greater detail in *SI Results*. The observed labeled carbon at position 3 of pyruvate is independent of the label exchange carbon specific to the carboxyl group, unlike previously reported in *Clostridium acetobutylicum* (22).

To determine which pathway(s) may also lead to  $^{13}$ C-labeled pyruvate at the carboxylic carbon, we examined the fate of carbon atoms among all possible candidate pathways, including the malate shunt (7, 14) which is known to assimilate and release CO<sub>2</sub> in *C. thermocellum.* As shown in Fig. 3*B*, CO<sub>2</sub> fixation is catalyzed by PEPCK, resulting in  $^{13}$ C-labeling at position 4 of oxaloacetate. This labeled carbon is propagated to position 4 of malate via malate dehydrogenase and is subsequently lost as

 $^{13}$ CO<sub>2</sub> by malic enzyme. Therefore, malate shunt, per se, does not contribute to the  $^{13}$ C-labeling at the carboxylic group of pyruvate. Another possible carboxylase enzyme that could presumably result in the labeling at the carboxylic group of pyruvate is ribulose bisphosphate carboxylase/oxygenase (RuBisCO). However, genes encoding for the highly conserved RuBisCO enzyme are not present in the *C. thermocellum* genome or in other sequenced *Clostridium* species. Thus, RuBisCO was ruled out.

We also considered the possibility of a reversible conversion between malate and fumarate catalyzed by malate hydrolyase (Clo1313\_0640). The symmetry of the fumarate molecule can interchange labeling on carbon atoms in positions 1 and 4; hence, it could contribute to the resulting <sup>13</sup>C-carboyxlic group in pyruvate. Although the alanine labeling pattern (Fig. 3*A*) cannot rule out the malate hydrolyase reaction, data from genetic KOs (Fig. 4) presented below are consistent with rPFOR activity and not malate hydrolyase activity.

We also investigated the isotopomer pattern for serine, which could be derived from phosphoglycerate, a triose phosphate interconverted with pyruvate in the glycolytic pathway (Fig. 2). Assuming that serine is exclusively derived from glycolysis and shares an identical synthetic route with alanine, the labeling pattern of serine and alanine should be very similar. However, in contrast to mostly unlabeled Pyr2–3, we observed a nonnegligible m1 peak of Ser2–3 with 12% enrichment (Fig. 3*A*). One carbon labeled at Ser2–3 implies a distinct serine pathway from pyruvate, consistent with the metabolic pathway in which a <sup>13</sup>C-methylene-THF from reductive C1 metabolism is combined with glycine by glycine hydroxymethyltransferase (*glyA*, *clo1313\_1155*) and forms the hydroxylmethylene group of serine at position 3



**Fig. 2.** Schematic of reductive C1 metabolism initiated from rPFOR and reversed PFL. The question mark indicates a missing reaction in *C. thermocellum* DSM1313 genome annotations. Pink-shaded pathways illustrate the reduction of CO<sub>2</sub> to formate and CO<sub>2</sub> to pyruvate. Yellow-shaded pathways include steps of serine and methionine biosynthesis. Fdh is not annotated in *C. thermocellum* genome. Fhs, formyl-tetrahydrofolate synthase (*clo1313\_0030*); FoID, methylene-tetrahydrofolate dehydrogenase/cyclohydrolase (*clo1313\_0030*); GlyA, glycine hydroxymethyltransferase (*clo1313\_1155*); MetF, methylene-tetrahydrofolate reductase (*clo1313\_1531\_155*); MS, methionine synthase (*clo1313\_1587-1588*, *clo1313\_1581-1582*, *clo1313\_0372*). Dotted lines indicate multistep catalysis. Carbons in red indicate <sup>13</sup>C-isotopes propagated from <sup>13</sup>C-bicarbonate or <sup>13</sup>C-formate in the tracer experiments.



**Fig. 3.** Isotopomer analysis of proteinogenic amino acids and lactate in cultures grown on 20 mM <sup>13</sup>C-sodium bicarbonate and 14.6 mM unlabeled cellobiose. (*A*) Relative abundance of mass isotopomers for pyruvate, serine, and lactate (n = 3, SD < 1%) in wild type ( $\Delta hpt$ ). (*B*) Schematics of carbon atom transits via the reversed PFOR (pink box) and the malate shunt (gray box). OAADC, oxaloacetate decarboxylase complex; MDH, malate dehydrogenase; ME, malic enzyme. The labeled (<sup>13</sup>C) and unlabeled (<sup>12</sup>C) carbon atoms are represented by closed and open circles, respectively. (*C*) Isotopomer analysis of proteinogenic alanine and leucine in  $\Delta hpt$  in cultures fed with [2-<sup>13</sup>C]acetate and unlabeled bicarbonate (20 mM) when the bacteria were grown primarily in 14.6 mM (5 g/L) cellobiose. Carbons at positions 1 and 2 of *N*-tert-butyldimethylsilyl-*N*-methyltrifluoroacetamide–derivatized leucine can be fragmented by GC/MS and are directly derived from acetyl-CoA.

(Fig. 2). These results implied that  $^{13}$ C isotope propagated to pyruvate was able to flow into the C1 pathway via formate, contributing to the biosynthesis of serine in *C. thermocellum*.

Knocking Out PFOR Genes Impairs Pyruvate Synthase Activity for CO<sub>2</sub> Fixation. The CO<sub>2</sub>-fixing C1 pathway initiated by PFOR in the reductive direction was further examined through the deletion of putative PFOR genes and the analysis of altered labeling patterns in the KO mutants. Multiple genes or gene clusters are annotated as PFOR (clo1313 0020-0023, 1353-1356, 0673, and 0382-0385) or indole pyruvate ferredoxin oxidoreductase (clo1313 1615-1616) in the C. thermocellum genome. These genes were respectively deleted via homologous recombination following the schemes shown in Fig. S2A, and validation of the genotype via colony PCR is shown in Fig. S3 and Table S2. All PCR products were further confirmed by commercial sequencing. The resulting mutants, ΔhptΔclo1313 0020-0023, ΔhptΔclo1313 1353-1356, ΔhptΔclo1313 1615-1616, ΔhptΔclo1313 0673, and  $\Delta hpt \Delta clo1313 \ 0382-0385$ , were fed with 20 mM <sup>13</sup>C-bicarbonate and 14.6 mM unlabeled cellobiose for isotope tracer experiments. The relative abundance of m0 and m1 shown for Pyr1-3 (M ala 057; Fig. 4) and for Pyr2-3 (M ala 085; Fig. S4) indicates that the carbon is labeled predominantly on the carboxylic group in pyruvate in both wild type ( $\Delta hpt$ ) and mutants, but the relative abundance of m1 is significantly lower in mutants ΔhptΔclo1313 0020-0023 and ΔhptΔclo1313 1353-1356 (Fig. 4). By taking the ratio of pyruvate labeled at the carboxylic group (m1) to unlabeled ones (m0), data showed that m1/m0 ratios are not significantly different in individual PFOR deletion mutants  $\Delta hpt \Delta clo1313$  1615-1616 and  $\Delta hpt \Delta clo1313$  0673, or exhibited little difference in mutant  $\Delta hpt \Delta clo1313$  0382-0385. However, the m1/m0 ratios are significantly lower in  $\Delta hpt \Delta clo1313$  0020-0023 and  $\Delta hpt \Delta clo1313$  1353-1356, with reduced labeling at the carboxylic group. The results identified Clo1313 0020-0023 and Clo1313 1353-1356 functioning as a pyruvate synthase, catalyzing the incorporation of CO<sub>2</sub> originating from <sup>13</sup>C-bicarbonate to form pyruvate. Based on the relative ratios of m1 to m0, Clo1313 0020-0023 and Clo1313 1353-1356 account for ~50% and 30% (Fig. 4) of pyruvate synthase activity, respectively, for carbon fixation. The differential m1/m0 ratios in these mutants also suggest that rPFOR contributes to the <sup>13</sup>C-carboxylic carbon labeling in pyruvate, which cannot originate from the interconversion between malate and fumarate alone. However, potential interplay between the putative PFORs and the interconversion between malate and fumarate is beyond the scope of this study, and hence is not addressed here.

involvement of PFL in reductive C1 metabolism, we inactivated PFL by deleting the *pflA* gene as illustrated in Fig. S2B, adapting a scheme developed previously (23). Deletion of the *pflA* gene was confirmed by PCR analysis (Fig. S5A) and commercial sequencing of the PCR products. Mutant  $\Delta hpt\Delta pflA$  grew slower than the wild type in defined medium, consistent with previous observations (12). We also measured the batch fermentation products when all substrates were consumed (Fig. S5B). As expected, although the wild type produces 2.3 mM formate, formate is no longer detected in the  $\Delta hpt\Delta pflA$  culture-spent medium (Fig. S5B), confirming complete inactivation of PFL and the fact that PFL is the only enzyme responsible for formate production from pyruvate.

PFL KO Mutant Is Deficient in Reductive C1 Metabolism. To assess the

To investigate how PFL deficiency affects carbon flux toward C1 metabolism, we compared the labeling pattern of isotopomers in both  $\Delta hpt\Delta pflA$  and wild type by feeding <sup>13</sup>C-formate (20 mM) and <sup>13</sup>C-bicarbonate (20 mM), respectively (Fig. 5). In either case, unlabeled cellobiose (14.6 mM) was added to the cell culture. We collected isotopomer data for proteinogenic amino acids that may be synthesized from this pathway, including alanine, serine, and methionine. With the addition of <sup>13</sup>C-formate, an m1 peak with 16% enrichment in Pyr1–3 was observed for wild type, whereas the corresponding m1 ratio was merely 4% in  $\Delta hpt\Delta pflA$ . In the isotopomer of Pyr2–3, we observed no



**Fig. 4.** Isotopomer analysis of proteinogenic alanine in  $\Delta hpt$  and in individual deletions of all five putative PFORs. All strains were grown on 20 mM <sup>13</sup>C-sodium bicarbonate and 14.6 mM unlabeled cellobiose. (*Inset*) Ratio of m1/m0 in individual putative PFOR deletion mutants is plotted. wt, wild type.



significant difference in labeling pattern between the strains. Taken together, the results supported that the <sup>13</sup>C-carbon atom in formate was internalized by the cells and became the carboxylic carbon of pyruvate, which was catalyzed by the reversed PFL activity. We next analyzed the Ser and Met isotopomers that could be synthesized from the C1 pathway downstream of <sup>13</sup>C-formate. Based on the labeling patterns of Ser1–3, Ser2–3, and Ser1–2 (Fig. 5*A*), we deduced that the <sup>13</sup>C-carbon propagated from <sup>13</sup>C-formate is mainly at position 3 of Ser (hydroxylmethylene group), consistent with our constructed de novo C1 metabolic pathway toward serine biosynthesis (Fig. 2). As for the isotopomer of Met shown in Fig. 5*A*, both  $\Delta hpt\Delta pflA$  and wild type indeed demonstrated a dominant peak at m1 upon <sup>13</sup>C-formate addition, validating the incorporation of the methyl group propagated from <sup>13</sup>C-formate into Met. Detailed descriptions of Ser and Met isotopomer labeling analyses are provided in *SI Results*.

We also analyzed the isotopomers of proteinogenic amino acids for <sup>13</sup>C-bicarbonate labeling experiments in  $\Delta hpt \Delta pflA$ . Based on our hypothesized "rPFOR-PFL shunt," incorporation of the <sup>13</sup>C-carbon from bicarbonate into the C1 metabolism is mediated by PFL, which could be validated by comparing <sup>13</sup>Cinflux into the C1 metabolism in the presence and absence of an active PFL. Similar pyruvate labeling patterns were observed for wild-type and  $\Delta hpt \Delta pflA$  strains, both of which showed m0 and m1 dominant peaks for Pyr1-3 and m0 signal for Pyr2-3 fragment. Results suggest that pflA deletion does not affect CO<sub>2</sub> incorporation catalyzed by the pyruvate synthase activity. As for serine labeling, both Ser1–3 and Ser2–3 fragments containing the hydroxyl methylene group had a lower labeling ratio in  $\Delta hpt \Delta pflA$ mutant than in wild type. For Ser2–3, mutant  $\Delta hpt \Delta pflA$  had a much lower m1 peak (2.6%) than the control (12.7%). Correspondingly, the unlabeled m0 peak of Ser2–3 for  $\Delta hpt \Delta pflA$  strain is 10.1% higher than in the parental strain (97.4% vs. 87.3%). Collectively, reduced isotope enrichment in the mutant strongly supports that C1 metabolism originating from <sup>13</sup>C-bicarbonate is disrupted by knocking out pflA. It is also evident that PFL activity is essential in internalizing the <sup>13</sup>C-carbon in bicarbonate in C. thermocellum, and our hypothesis that PFL coupled with the rPFOR activity initiates the reductive C1 pathway in C. thermocellum was corroborated. We observed a lower extent of fractional labeling of methionine in the  $\Delta hpt\Delta pflA$  mutant than in wild type, which is described in greater detail in SI Results.

## Discussion

This study reveals that during cellobiose metabolism, *C. thermocellum* also assimilates  $CO_2$  via a reductive C1 metabolic pathway that is initiated by the dual actions of reversed PFOR and PFL. PFOR has been greatly considered in the forward direction in *C. thermocellum*, catalyzing the oxidative decarboxylation

Fig. 5. Isotopomer analysis of GC/MS fragments for proteinogenic amino acids in  $\Delta hpt\Delta pflA$  mutant (white columns) and wild type ( $\Delta hpt$ , black columns) grown on 14.6 mM unlabeled cellobiose with 20 mM <sup>13</sup>C-formate (A) or 20 mM <sup>13</sup>C-sodium bicarbonate (B). Panels for each GC/MS fragment represent the labeling pattern of the amino acid or its precursor with indicated C atoms. The technical variance is less than 1% in tracer experiments.

of pyruvate to acetyl-CoA and CO2 and generating reduced ferredoxin (Fd<sub>red</sub>). The PFOR activity in C. thermocellum has been associated with the oxidative decarboxylation of pyruvate to acetyl-CoA. Uyeda and Rabinowitz (24, 25) and Raeburn and Rabinowitz (26) demonstrated in vitro that low potential electron donors, such as Fd<sub>red</sub>, can drive PFOR to catalyze the reductive carboxylation of acetyl-CoA as a pyruvate synthase. The rPFOR serves as a key enzyme for autotrophic CO<sub>2</sub> fixation via the Wood-Ljungdahl pathway in acetogens (27). In this work, the in vivo isotopomer data measured postfeeding *C. thermocellum* <sup>13</sup>C-biocarbonate and [2-<sup>13</sup>C] acetate double-confirmed rPFOR activity and revealed that rPFOR activity accounted for over 40% of the labeled carbon in pyruvate. Measurements in individual PFOR mutants further identified rPFOR activity predominantly catalyzed by Clo1313 0020-0023 and Clo1313\_1353-1356. Consistent with previous global expression studies, Clo1313 0020-0023 is the most highly expressed PFOR based on proteomics (28) and transcriptomics (29) analyses. Clo1313 1353-1356 is the third most highly expressed PFOR based on proteomics data, but it has a low transcript level based on microarray data, and the difference could be due to different growth conditions in these studies.

By feeding <sup>13</sup>C-bicarbonate to the PFL-inactivated mutant and the wild type, we confirmed the incorporation of these tracers into formate (Fig. 1), lactate (Fig. 3A), and the downstream amino acids linked to C1 metabolism (Figs. 3 and 5). We provided multiple lines of evidence showing that the fixed  $CO_2$  was channeled to the C1 metabolism through PFL, which could operate reversibly (21, 30) and contribute to the biosynthesis of essential amino acids, including serine and methionine. In addition, the reductive C1 pathway mediated by PFL may play a critical role in nucleotide biosynthesis (reconstructed purine biosynthesis pathway in C. thermocellum shown in Fig. S6). The impaired growth observed in PFL-inactivated mutant is consistent with a previous finding (12) and underscores PFL's importance. A transcriptome study reported that folate-dependent enzymes are in the most highly expressed category, which also suggests an active C1 metabolism in vivo (31).

Carbon loss in the form of  $CO_2$  from the conversion of pyruvate to acetyl-CoA across a wide range of bacterial heterotrophs reduces the theoretical maximum yield of a carbon product of interest per carbon consumed (carbon efficiency) downstream of acetyl-CoA. A synthetic nonoxidative glycolysis has been shown to bypass the loss of  $CO_2$  in the conversion of pyruvate to acetyl-CoA (32). The rPFOR activity shown here allows some of the lost  $CO_2$ to be recaptured into pyruvate, a critical branch point and precursor to numerous natural and nonnatural products. Based on our data,  $CO_2$  utilization has increased the titers of organic carbon compounds (Table 1) by incorporating  $CO_2$  into the biomass formula, as detected in proteinogenic serine and methionine, and, as such, supplementing cell growth.

It is interesting that C. thermocellum uses the rPFOR-PFL shunt for CO<sub>2</sub> uptake in the absence of Fdh. We speculate that although pyruvate is a key branching point in central carbon metabolism and a key node connecting glycolysis and the C1 metabolic pathway, the rPFOR-PFL route for CO<sub>2</sub> uptake may provide added metabolic flexibility and versatility to cells in response to environmental changes affecting their carbon and energy (redox) needs. To gain insights into how widespread the rPFOR-PFL shunt could be for CO<sub>2</sub> uptake, we surveyed the presence of PFOR, PFL, and Fdh in 67 sequenced Clostridium species. We discovered that only 20 species have Fdh, whereas 51 species have both PFOR and PFL (Fig. S7). Although the in vivo reversibility of PFOR can be organism- and/or growth condition-specific, and information as to whether these bacteria can use inorganic carbon during organic substrate feeding may not be readily available, the presence of these genes renders it genetically feasible.

It should be noted that carbon fixation via the rPFOR-PFL shunt requires  $Fd_{red}$  as an electron donor. On the other hand, PFOR functioning in the oxidative direction serves as a major pathway for generating  $Fd_{red}$  to yield reduced biofuels and chemicals. Thus, the metabolic flux toward biosynthesis may be a trade-off of the flux toward electron generation. We speculate that CO<sub>2</sub> enhances growth by regenerating oxidized Fd, which, in turn, provides a driving force for faster cellobiose consumption. Feeding CO<sub>2</sub> may also enhance the "transient" CO<sub>2</sub> fixation via the malate shunt, with GTP produced via PEPCK promoting *C. thermocellum* growth. The biosynthesis vs. energy requirement at the pyruvate node needs to be evaluated comprehensively with

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other major redox pathways (i.e.,  $H_2$  production) to enhance the carbon economy without causing redox imbalance. To satisfy our curiosity, we tested *C. thermocellum* autotrophic growth by feeding either bicarbonate plus 100%  $H_2$  or an 80%  $H_2/20\%$ (vol/vol) CO<sub>2</sub> gas mixture, but detected no growth (Fig. S8). Nevertheless, the rPFOR-PFL shunt uncovered in this study may have important ramifications in the metabolic plasticity of a cellulose degrader and in other heterotrophs, and hence warrants more in-depth investigations.

#### **Materials and Methods**

A more complete discussion is provided elsewhere (15, 16, 23, 33–35), as well as in *SI Methods*.

Unless otherwise noted, *C. thermocellum* DSM1313-derived strains,  $\Delta hpt$  (*hpt* encoding for hypoxanthine phosphoribosyl transferase is used for counterselection), and other mutants were routinely grown anaerobically at 55 °C on 5 g/L cellobiose. The *hpt* gene deleted strain is used as the wild type for the studies herein. To test the inorganic carbon utilization, 20 mM so-dium bicarbonate was added to cell cultures also fed with cellobiose. We used the DSM122-defined medium referred to as CTFUD medium (36). Full details on strain construction, fermentation conditions, and <sup>13</sup>C-tracer experiments are provided in *SI Methods*.

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