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Daniel G. Olson
Dartmouth College

Richard J. Giannone
Oak Ridge National Laboratory

Robert L. Hettich
Oak Ridge National Laboratory

Lee R. Lynd
Dartmouth College

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Role of the CipA Scaffoldin Protein in Cellulose Solubilization, as Determined by Targeted Gene Deletion and Complementation in *Clostridium thermocellum*

Daniel G. Olson,^{a,d} Richard J. Giannone,^{b,d} Robert L. Hettich,^{b,d} Lee R. Lynd^{a,b,c,d}

Dartmouth College, Hanover, New Hampshire, USA^a; Oak Ridge National Laboratory, Oak Ridge, Tennessee, USA^b; Mascoma Corporation, Lebanon, New Hampshire, USA^c; BioEnergy Science Center, Oak Ridge, Tennessee, USA^d

The CipA scaffoldin protein plays a key role in the *Clostridium thermocellum* cellulosome. Previous studies have revealed that mutants deficient in binding or solubilizing cellulose also exhibit reduced expression of CipA. To confirm that CipA is, in fact, necessary for rapid solubilization of crystalline cellulose, the gene was deleted from the chromosome using targeted gene deletion technologies. The CipA deletion mutant exhibited a 100-fold reduction in cellulose solubilization rate, although it was eventually able to solubilize 80% of the 5 g/liter cellulose initially present. The deletion mutant was complemented by a copy of *cipA* expressed from a replicating plasmid. In this strain, Avicelase activity was restored, although the rate was 2-fold lower than that in the wild type and the duration of the lag phase was increased. The *cipA* coding sequence is located at the beginning of a gene cluster containing several other genes thought to be responsible for the structural organization of the cellulosome, including *olpB*, *orf2p*, and *olpA*. Tandem mass spectrometry revealed a 10-fold reduction in the expression of *olpB*, which may explain the lower growth rate. This deletion experiment adds further evidence that CipA plays a key role in cellulose solubilization by *C. thermocellum*, and it raises interesting questions about the differential roles of the anchor scaffoldin proteins *OlpB*, *Orf2p*, and *SdbA*.

Clostridium thermocellum is an anaerobic thermophilic bacterium noted for its ability to rapidly solubilize crystalline cellulose, a process mediated by the cellulosome (1). The cellulosome is composed of tightly bound enzymatic and structural components. At the heart of the cellulosome is the scaffoldin protein, CipA (also known as S_L and S_1) (2). This protein has been shown to be capable of crystalline cellulose solubilization in conjunction with cellulosomal cellulase Cel48S (3). Analysis of the DNA sequence of *cipA* has revealed a set of nine repeated elements known as type I cohesins (4). These cohesins bind to the type I dockerins found on cellulosomal enzymes (5). Subsequent analysis of the CipA protein has revealed three additional modules, the type II dockerin, the carbohydrate binding module (CBM), and the x domain.

The type II dockerin comprises a duplicated set of 22 amino acid residues located near the C terminus of CipA (4). The type II dockerin binds to type II cohesins located on the anchor scaffoldin proteins, *OlpB*, *Orf2p*, and *SdbA*. *OlpB* has seven type II cohesins, while *Orf2p* has two and *SdbA* has one. The anchor scaffoldins have a C-terminal sequence called the S-layer homology (SLH) domain that mediates binding to the cell surface (6). In CipA the CBM is located between the second and third type I cohesins and binds to crystalline cellulose with a K_D (equilibrium dissociation constant) of 0.4 μ M (1). Thus, the current understanding of the adhesion of *C. thermocellum* to cellulose involves the following 3 interactions: (i) binding of glycoside hydrolase enzymes in proximity to each other to promote enzyme-enzyme synergy; (ii) binding of enzymes to the cellulosic substrate via the CBM; and (iii) anchoring of the cellulosome to the cell surface, where CipA binds to the anchor scaffoldin (*OlpB*, *Orf2p*, or *SdbA*) via its type II dockerin and the anchor scaffoldins are attached to the cell by their SLH domains.

Finally, CipA has one additional module, located between the

ninth type I cohesin and the type II dockerin, called the x module. Its function in *C. thermocellum* remains unknown, although it has been shown to improve the solubility of recombinantly expressed type II dockerins and seems to enhance the affinity of the type II cohesin-dockerin interaction (7).

Electron microscopy has revealed hemispherical protuberances on the outside of *C. thermocellum* cells, which are known as polycellulosomes (8). In their resting state they are about 200 nm in diameter, but they form a protracted conformation in the presence of cellulose (9). Immunolabeling has identified the presence of CipA (10) and *OlpB* (6) in the polycellulosomes, though the protuberances may contain other cellulosomal components as well.

There have been two previous reports of mutants of *C. thermocellum* deficient in cellulase activity. Both were isolated by screening for cells unable to adhere to cellulose. *C. thermocellum* AD2 was isolated by mixing cells with cellulose and allowing the cellulose to settle. Adherent cells were pulled out of solution upon binding to cellulose, thus enriching the supernatant for nonadherent cells. After five rounds of this sedimentation enrichment, strain AD2 was isolated by single-colony purification (11). The AD2 strain was analyzed by SDS-PAGE and found to be missing a band associated with CipA when grown on cellobiose, although

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Address correspondence to Lee R. Lynd, lee.lynd@dartmouth.edu.

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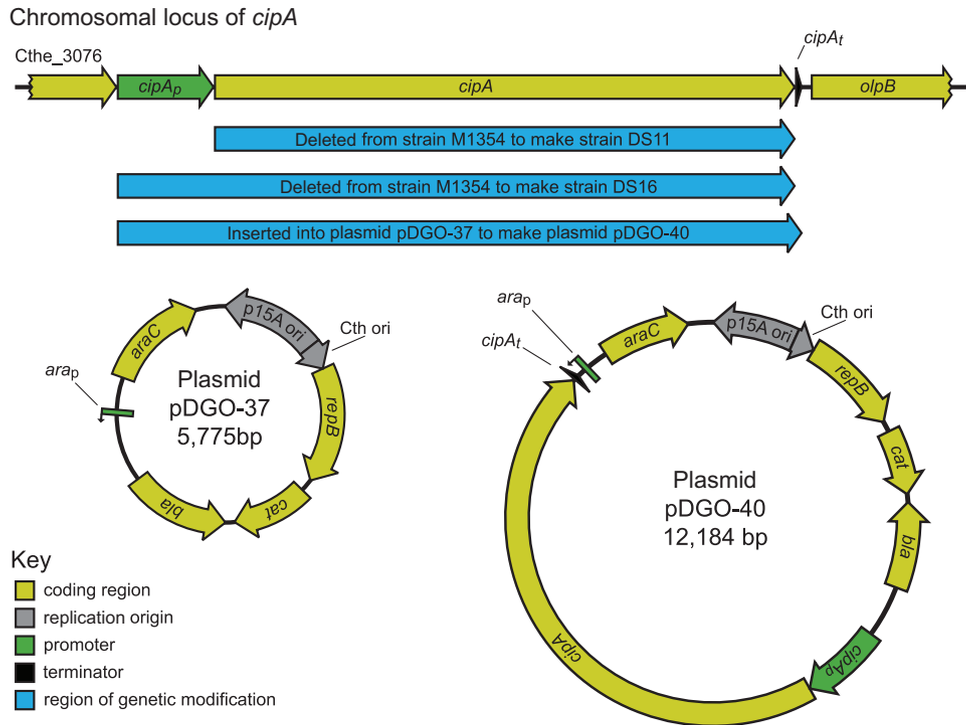


FIG 1 Diagram of genetic elements used in this work.

the band reappeared when the strain was grown on cellulose (8). Further analysis of AD2 by scanning electron microscopy revealed the complete absence of polycellulosomes when the strain was grown on cellobiose (12).

Strains SM1, SM4, SM5, and SM6, which also are deficient in cellulase activity, were isolated using a procedure similar to that used for strain AD2, though augmented by an initial chemical mutagenesis step followed by a screen on cellobiose plates with an Avicel overlay (13). This final screen was designed to identify cells that were deficient in cellulose solubilization. These mutants were analyzed by SDS-PAGE, and all were found to be missing a 210-kDa band associated with CipA. DNA sequence analysis revealed the presence of an IS1447 insertion element disrupting the *cipA* coding sequence in each mutant. Strain SM1 had an insertion in the first type I cohesin and appeared to be completely lacking functional type I cohesins. This strain was unable to grow on MN300 cellulose and exhibited a 15-fold reduction in enzymatic activity compared to the wild-type (WT) strain (13).

Previous work has shown that the abilities to bind and to solubilize cellulose are linked (11, 13). Thus, mutants deficient in cellulose binding are also deficient in cellulose solubilization. The functional link between these abilities is consistent with our understanding of the component modules of CipA. In this study, we further evaluated the extent to which *cipA* is responsible for this dysfunctional phenotype and explored the cellulase activity of *C. thermocellum* in the absence of a complexed cellulase system.

MATERIALS AND METHODS

Strains and media. All *C. thermocellum* strains described here are derived from *C. thermocellum* strain DSM 1313 and were grown in modified DSM 122 broth as described previously (14). Cellobiose or Avicel-PH105 microcrystalline cellulose (Sigma-Aldrich) was used as the primary carbon

source at a concentration of either 5 or 10 g/liter. Cells were grown at 55°C. Strain M1354 was a generous gift from the Mascoma Corporation (Lebanon, NH) (15). This strain is derived from *C. thermocellum* strain DSM 1313 and has a deletion of the *hpt* gene (Clo1313_2927) to allow for use of the *hpt* gene as a counterselectable marker with the antimetabolite 8-aza-hypoxanthine.

Molecular biological methods. Plasmids were constructed using yeast-mediated ligation (16), In-Fusion PCR cloning (TaKaRa Bio Inc.), or standard cloning techniques (17). Plasmids were maintained in *Escherichia coli* TOP10 cells (Invitrogen Corporation) and prepared using the Qiagen plasmid minikit (Qiagen Inc.). Sequences of chromosomal DNA were obtained by PCR using genomic DNA from *C. thermocellum* strain DSM 1313. Primers were designed using genome sequences provided by the Joint Genome Institute (<http://www.jgi.doe.gov/>). The *repB* and *cat* genes are derived from plasmid pMU102 (18). The yeast origin of replication is derived from plasmid pMQ87 (16). The pMB1 *E. coli* origin of replication is derived from plasmid pUC19 (Invitrogen Corp.). The p15A *E. coli* origin of replication and arabinose-inducible promoter are derived from plasmid pBAD30 (19). The *hpt* and *tdk* genes (Tsac_0936 and Tsac_0324, respectively) are derived from *Thermoanaerobacterium saccharolyticum* JW/SL-YS485. The glyceraldehyde 3-phosphate dehydrogenase promoter consists of the 525-bp region upstream of the *C. thermocellum* glyceraldehyde 3-phosphate dehydrogenase gene (Clo1313_2095). The *cbp* promoter consists of the 621-bp region upstream of the cellobiose phosphorylase gene (Clo1313_1954).

Plasmid pDGO-37 (GenBank accession number JX966413) was created by combining the p15A *E. coli* origin of replication and *Pbad* promoter with the thermophilic Gram-positive origin of replication from plasmid pMU102. Plasmid pDGO-40 (GenBank accession number JX966414) was created by inserting the *cipA* coding sequence, including 819 bp upstream of the start codon (putative promoter region) and 67 bp downstream of the stop codon (putative terminator region), into plasmid pDGO-37 (Fig. 1; Table 1).

PCR was performed using either *Taq* or Phusion DNA polymerase

TABLE 1 Description of *C. thermocellum* strains

Strain	Chromosomal genetic elements			Plasmid genetic elements		Genotype	Source or reference
	<i>cipAp</i>	<i>cipA</i>	<i>hpt</i>	<i>cat</i>	<i>cipAp-cipAt</i>		
DSM1313 (WT)	+	+	+			Wild type	DSMZ ^a
M1354	+	+				Δhpt	15
DS11	+					M1354 $\Delta cipA$	20
DS16						M1354 $\Delta(cipAp-cipA)$	20
DS18				+		DS16/pDGO-37	This study
DS19	+	+		+		M1354/pDGO-37	This study
DS20				+	+	DS16/pDGO-40	This study
DS22	+	+		+	+	M1354/pDGO-40	This study

^a DSMZ, Deutsche Sammlung von Mikroorganismen und Zellkulturen GmbH, Germany.

(New England BioLabs Inc.) according to the directions provided by the manufacturer. When using whole cells as the PCR template, a 10-min heating step was included at the beginning of the thermocycling protocol to lyse the cells. When using *Taq* DNA polymerase, the lysing temperature was 95°C. When using Phusion DNA polymerase, the lysing temperature was 98°C. DNA sequencing was performed using standard techniques with an ABI model 3100 genetic analyzer (Applied Biosystems).

Strain construction. Previously, two different deletions of *cipA* were made. In strain DS11, the *cipA* coding sequence was deleted from the start codon to the stop codon using plasmid pDGO-03 (GenBank accession number JX489218.1). In strain DS16, the *cipA* promoter sequence was deleted in addition to the *cipA* coding region using plasmid pDGO-34 (GenBank accession number JX489219.1) (20). Plasmids were transformed into *C. thermocellum* using previously described techniques (21). To avoid the potential for homologous recombination between the plasmid and chromosomal copy of the *cipA* promoter region, plasmid pDGO-40 was transformed only into strain DS16.

Fermentation conditions. Strains were grown in modified DSM 122 broth (18) at 55°C with cellobiose or Avicel microcrystalline cellulose as the primary carbon source. When fermentations were performed in a 125-ml glass bottle sealed with a butyl rubber stopper (22), the fermentation volume was 50 ml, 5 g/liter substrate (Avicel or cellobiose) was used, the headspace was purged with nitrogen, and the bottles were shaken at 200 rpm. When fermentations were performed in a computer-controlled fermentor (Sartorius GmbH), the fermentation volume was 2 liters, 10 g/liter substrate (Avicel or cellobiose) was used, the headspace was purged with a mixture of 20% CO₂ and 80% N₂, the vessel was stirred at 200 rpm, and the pH was controlled to 7.0 with 4 N potassium hydroxide. For some fermentations, an automated sampling device was used to take 6-ml samples at regular intervals (23).

Analytical techniques. Concentrations of cellobiose, glucose, lactate, acetate, ethanol, and formate were measured by high-performance liquid chromatography (HPLC) as previously described (24). Total carbon and total nitrogen concentrations (TOCN) were measured with a Shimadzu TOC-V CPH elemental analyzer with TNM-1 and ASI-V modules (Shimadzu Corp.) on 0.5- to 1.0-ml aliquots washed twice with water. The Avicel concentration was determined from these measurements by assuming that Avicel contained no nitrogen and that cells contained carbon and nitrogen in a 4.67:1 molar ratio. A detailed description of the theory and calculations is being prepared for publication (E. K. Holwerda and L. R. Lynd, unpublished data).

Samples for protein identification were prepared as previously described (14). Briefly, cell pellets were separated from supernatant by centrifugation (2,000 × g) and washed twice with Tris-buffered saline (100 mM Tris HCl, 150 mM NaCl, pH 8.0) to remove residual supernatant proteins. Cells were lysed by the addition of SDS lysis buffer (SDS LB) (4% SDS [wt/vol] in 100 mM Tris HCl, pH 8.0), boiled (5 min), sonically disrupted (Branson Sonifier), and boiled again. Supernatants were concentrated 50-fold (50 ml to 1 ml) via spin filtration using a 3,000-molec-

ular-weight (MW)-cutoff membrane (Vivaspin 20, 3 kDa, PES [GE Healthcare]), adjusted to 2% SDS with 1 ml of SDS LB, and boiled (5 min). Both fractions, whole cell (WC) and supernatant (SN), were pre-cleared by centrifugation (21,000 × g) and protein concentrations determined by bicinchoninic acid (BCA) assay (Pierce). Sample fractions were then combined in a 2:1 (wt/wt) WC-to-SN ratio, reduced with 25 mM dithiothreitol (DTT), and trichloroacetic acid (TCA) precipitated (3 mg of combined crude lysate adjusted to 20% TCA on ice for 1 h). Precipitated proteins were then washed, resolubilized in denaturation buffer (8 M urea, 100 mM Tris HCl, 5 mM DTT, pH 8.0), digested with trypsin and prepared for MudPIT liquid chromatography-tandem mass spectrometry (LC-MS/MS) as previously described (14). In total, 50 µg of peptides was analyzed per sample via a 24-h MudPIT analysis using an LTQ XL mass spectrometer (Thermo Scientific). The resulting peptide fragmentation data were then searched with the MyriMatch database search algorithm (25) against the *C. thermocellum* DSM 1313 proteome (with decoy sequences) as previously described. Identified peptides were then score filtered (false-discovery rate [FDR], <2% peptide spectrum match) and assembled into protein identifications (minimum of 2 distinct peptides per protein call) by IDPicker 3 (26). Proteins were then spectrally balanced to deal with nonunique peptides and normalized by normalized spectral abundance factors (NSAF), and abundance values were adjusted to normalized spectral counts (nSpC) as previously described (27). Protein-to-protein abundance was then assessed across all samples to identify those that were differentially expressed (Student's *t* test).

Categorization of cellulase and cellulosomal proteins. A list of all proteins that could participate in cellulose solubilization was generated, based on membership in the Carbohydrate Active Enzyme (CAZY) database (28) or presence of a cohesin or dockerin domain as determined by the Pfam database (29) (see Data Set S1 in the supplemental material). Proteins with Pfam domain PF00963 were designated “cohesin containing.” Proteins with Pfam domain PF00404 (and no cohesin domain) were designated “dockerin containing.” Three additional proteins, Clo1313_1300, Clo1313_2479, and Clo1313_2861, were added to this list, based on analysis done by Ed Bayer and colleagues (Ed Bayer, personal communication). Proteins in the CAZY database that did not have a cohesin or dockerin domain were designated “CAZY, no cohesin, no dockerin.”

Mathematical analysis of fermentation data. To determine the rate of substrate consumption, the substrate consumption data points were fitted with the 5-parameter sigmoidal Richards equation (30) per the work of Holwerda and Lynd (unpublished data):

$$s(t) = A_0 + \frac{A_t - A_0}{1 + e^{\left(\frac{t_0 - t}{\text{slope}}\right)_{\text{asymm}}}} \quad (1)$$

where A_0 is the lower horizontal asymptote, A_t is the higher horizontal asymptote, t is time, t_0 is the inflection point, slope is the slope at the inflection point, and asymm is the asymmetry parameter. The time (t)

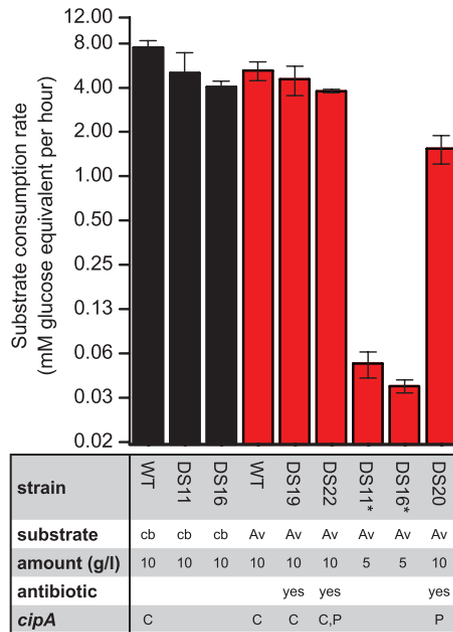


FIG 2 Substrate consumption rates for strains of *C. thermocellum* grown on either cellobiose (cb) or Avicel (Av) at an initial concentration of 5 or 10 g/liter. Antibiotic selection was used to maintain the plasmid in plasmid-containing strains. The presence of the *cipA* coding sequence is indicated as either chromosomal (C), plasmid based (P), or both. Error bars represent one standard deviation and were determined based on biological replicates, where $n \geq 2$. *, due to the difficulties of growing strains DS11 and DS16 on Avicel in fermentors, they were grown in sealed glass bottles instead.

when the slope of the fitted curve was greatest was determined by taking the second derivative with respect to time, setting it equal to zero, and solving for t , which yielded the following equation:

$$t_{\text{max slope}} = x_0 \ln(\text{asymm}) * \text{slope} \quad (2)$$

The first derivative of equation 1 with respect to time was then evaluated at the time determined by equation 2 to determine the maximum rate of substrate consumption. To allow ready comparison between Avicel and cellobiose, the substrate consumption rate was determined in mM glucose equivalent/hour.

RESULTS

Comparison of growth rates of various mutants. As expected, the wild-type (WT) strain and the *cipA* deletion strains (DS11 and DS16) have similar substrate consumption rates when grown on cellobiose (Fig. 2). The metabolic burdens of plasmid maintenance and thiamphenicol inactivation do not have an effect on Avicel consumption, as can be seen by comparing the wild type to the empty-vector control (strain DS19) (Fig. 2). The effect of *cipA* overexpression can be seen by comparing the empty-vector control (strain DS19) with the *cipA* overexpression strain (DS22). The rate of Avicel consumption was unchanged (Fig. 2). Deletion of *cipA* (strains DS11 and DS16) resulted in a 100-fold decrease in Avicel consumption rate compared to that of the wild-type strain (Fig. 2). Both *cipA* deletion strains were able to consume 80% of the Avicel initially present after ~2,000 h (see Table S1 and Fig. S1 in the supplemental material). Fermentation products were similar to those of the parent strain (see Table S2 in the supplemental material), and there was no significant accumulation of glucose or cellobiose. To see if the rate of Avicel consumption could be im-

proved by adaptation, strain DS11 was subsequently passed two additional times on Avicel, but no change in rate was detected.

Transforming the *cipA* deletion strain (DS16) with the *cipA* expression plasmid (pDGO-40) (resulting in strain DS20) dramatically increased the rate of Avicel solubilization, although it was still only 1/3 as high as that for the empty-vector control (DS19) (Fig. 2; see Table S1 in the supplemental material). Interestingly, there was a 100-hour lag phase before the start of rapid cellulose solubilization. The technique of measuring Avicel concentration by TOCN resulted in a larger degree of measurement variation during the early parts of fermentation. It is therefore difficult to determine whether the slight negative trend observed in the first 100 h with strain DS20 represents a physical phenomenon or is simply an artifact of the measurement technique (Fig. 3).

Comparison of protein abundances in various mutants. Although the abundance was measured for all proteins (see Data Set S1 in the supplemental material), only those proteins thought to be able to participate in cellulose solubilization (due to the presence of a cohesin, dockerin, or CAZy domain) were analyzed. The effect of the metabolic burdens of plasmid maintenance and antibiotic inactivation can be determined by comparing the empty-vector control (strain DS19) to the wild type (WT) (Fig. 4, first column). Among the cohesin-containing proteins, none are significantly differentially expressed. Among the dockerin-containing proteins, only Cel9F is significantly differentially expressed. Among the other CAZy proteins, only LicA is significantly differentially expressed.

The effect of *cipA* overexpression is demonstrated by comparing the empty-vector control (strain DS19) with the *cipA* overexpression strain (strain DS22) (Fig. 4, second column). Among the cohesin-containing proteins, none of them were significantly different at a P value of 0.01. Although the increase in *cipA* expression was not significant at the 0.01 level ($P = 0.015$), when the values for the wild type are included as well, the significance increases to 0.0002, suggesting that the effect would likely be confirmed if we were to perform more replicates. *cipA* expression increased by 3-fold in strain DS22, although this was not significantly different from the mean *cipA* expression in the empty-vector control (strain DS19) at the 0.01 level ($P = 0.015$). Among the dockerin-containing enzymes, none were significantly differently expressed at the 0.01 level. Among the other CAZy proteins, LicA and Clo1313_0647 (CBM16, domain of unknown function) were significantly lower in abundance.

The effect of *cipA* complementation can be determined by comparing the empty-vector control (strain DS19) with the complemented *cipA* deletion strain (DS20) (Fig. 4, third column). Among the cohesin-containing proteins, OlpB showed significantly reduced expression and was 11-fold less abundant in the complemented deletion strain. Among dockerin-containing proteins, Cel9P and Clo1313_2861 (GH2-CBM6) showed increased expression. The significance of the Clo1313_2861 result is difficult to interpret because of the low number of spectra identified for this protein (≤ 7 for all samples). Among the other CAZy proteins, Clo1313_2460 (GH15) showed significantly increased abundance.

Since OlpB contains 7 type II cohesins, a dramatic decrease in OlpB levels could result in a decrease in type II cohesin availability. Type II cohesin availability was calculated by multiplying the

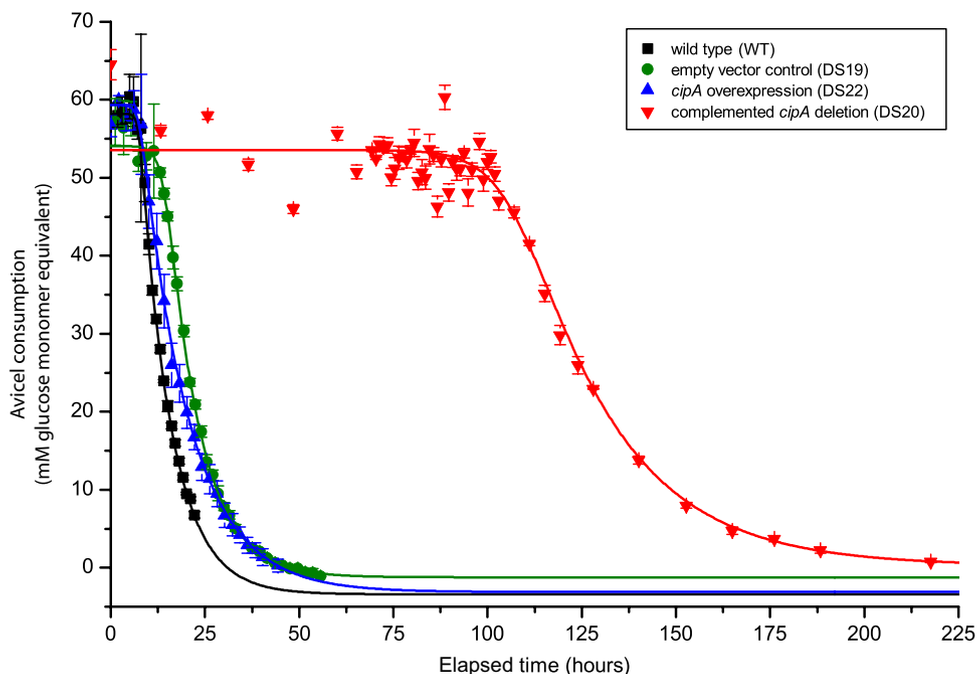


FIG 3 Avicel consumption by 4 strains of *C. thermocellum* growing on 10 g/liter Avicel. In order to allow subsequent comparison with growth rates on cellobiose, the rate was reported in mM glucose equivalents per hour. Based on an assumed monomer mass of 162 g/mol and a 5% moisture content of Avicel, 58.6 mM glucose equivalents were present initially. Avicel consumption was measured by elemental analysis of the pellet fraction of fermentation broth corrected for cell carbon. Error bars represent one standard deviation ($n = 3$) for Avicel measurement of a representative fermentation. Solid lines represent the best fit of a 5-parameter logistic equation. Equation parameters are given in Table S1 in the supplemental material.

abundance of each anchor scaffoldin (SdbA, Orf2p, and OlpB) by the number of type II cohesins it contains (Table 2).

DISCUSSION

In agreement with Zverlov et al., we have shown that *cipA* is essential for rapid solubilization of crystalline cellulose (13). However, contrary to what was reported previously, we observed that *cipA* deletion strains are able to solubilize Avicel microcrystalline cellulose (which is similar to the MN300 microcrystalline cellulose used by Zverlov et al.). Furthermore, since the solubilization of Avicel resulted in the production of lactate, acetate, and ethanol and this ability was maintained despite serial transfer, it appears that the strain was able to grow on Avicel. What is the explanation for residual ability of the *cipA* deletion strains (DS11 and DS16) to solubilize crystalline cellulose? One possibility is that components of the noncomplexed cellulase system (i.e., Cell and CelY), which have been shown to synergistically solubilize crystalline cellulose (31), can compensate for the expected loss of activity. Since each enzyme has its own CBM, there is no need for CipA to mediate binding with the cellulosic substrate. These enzymes were found at very low levels in all strains ($<0.01\%$ of cell protein) as determined by nSpC values (see Data Set S1 in the supplemental material), which reduces support for this explanation. Another possibility is that the cellulosomal components are bound directly to the cell surface via OlpA, which contains both a type I cohesin (for binding a cellulase enzyme containing a type I dockerin) and an S-layer homology (SLH) binding domain (for binding to the cell surface). Levels of OlpA were about 40% higher in the complemented *cipA* deletion strain (DS20) than in the empty-vector control (DS19), which supports this hypothesis.

Why does the *cipA* deletion and complementation strain (DS20) grow more slowly than the wild-type strain and have a longer lag phase? *cipA* expression does not seem to be a likely explanation, since minor variations in CipA abundance do not appear to be correlated with growth rate (Fig. 5). On the other hand, OlpB levels were unexpectedly low in this strain. Compared to the empty-vector control strain (DS19), the *cipA* deletion and complementation strain (DS20) had 30% fewer type II cohesins, since it seems to have partly compensated for the reduction in *olpB* expression with higher levels of *sdbA* and *orf2p* expression. Furthermore, the wild-type strain and the *cipA* overexpression strain (DS22) both had 15% fewer type II cohesins, and this change in type II cohesin number did not have a substantial effect on fermentation performance; thus, it seems unlikely that the change in type II cohesin number is the full explanation. Another possibility is that the anchor scaffoldins (OlpB, Orf2p, and SdbA) are not, in fact, interchangeable. For example, if Orf2p and SdbA are primarily used during cellulosome assembly (as has been suggested for ORFXp in *Clostridium cellulolyticum* [32]) and OlpB is the final destination for the assembled cellulosome, then a change in the abundance of OlpB might have a greater impact on cellulosome function than would be indicated simply by the overall change in type II cohesin availability. Further investigations will be required to determine the exact molecular roles of the various type II cohesin-containing proteins in cellulosome assembly.

Why is *olpB* expression changed in the *cipA* deletion strain? Although *cipA* and *olpB* have been reported to be transcribed independently (33), they may, in fact, be cotranscribed. Even if *cipA* and *olpB* are expressed from individual promoters, the 1-kb region upstream of *cipA* may contain other regulatory elements that

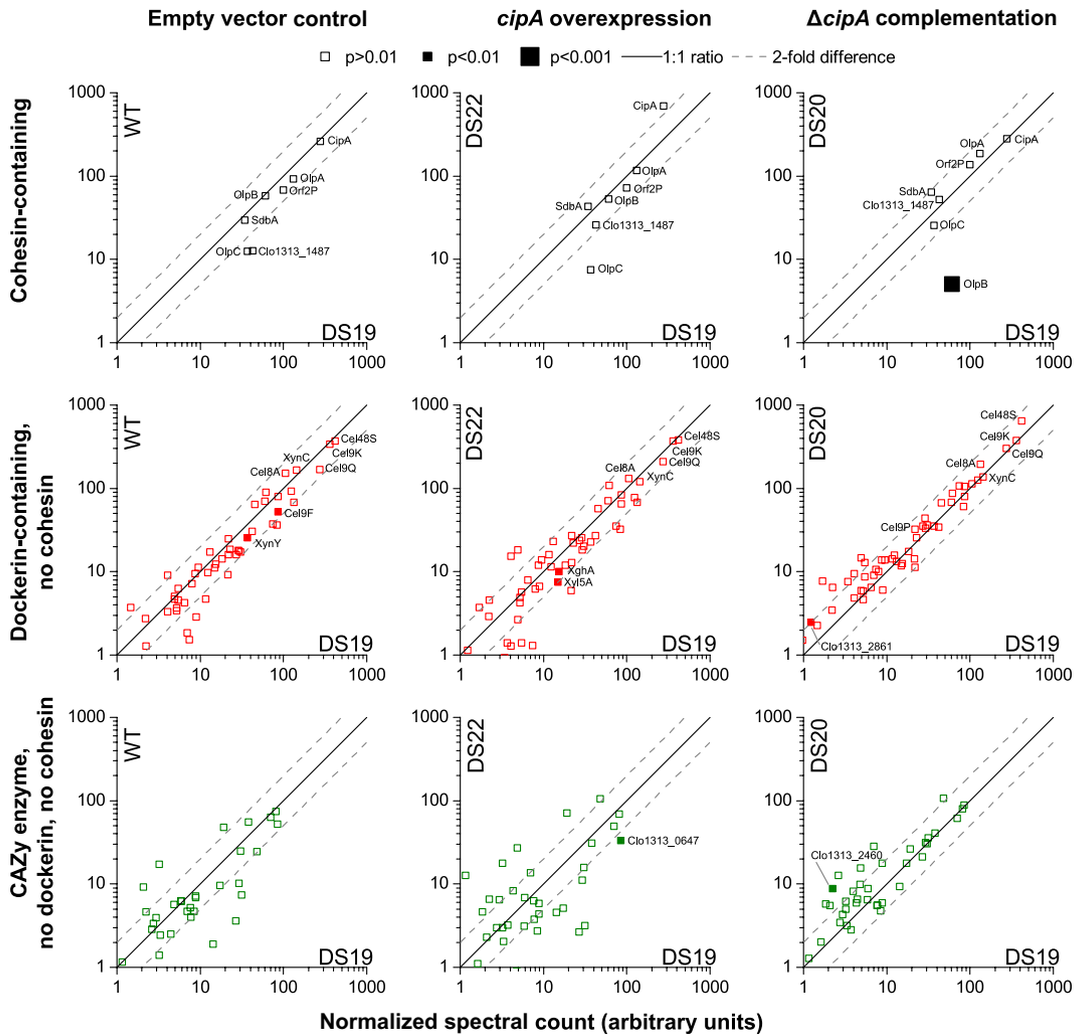


FIG 4 Comparison of protein abundance as determined by normalized spectral abundance factor (nSpC) from tandem mass spectrometry measurements of fermentation broth (combined cells and supernatant) at the end of Avicel fermentations. nSpC measurements were taken from biological duplicate experiments. Pairwise comparisons were made, and proteins with significant changes ($P < 0.01$) are indicated by filled symbols. Other proteins are indicated by unfilled symbols. The presence of cohesins, dockerins, and carbohydrate binding modules (CBMs) was determined by searching the Pfam database (29).

affect *olpB* expression. Replacing the native *cipA* promoter on plasmid pDGO-40 with a different promoter would allow *cipA* to be expressed from a replicating plasmid in a *cipA* deletion strain where the native *cipA* promoter has been left on the chromosome (e.g., strain DS11).

The *cipA* deletion and complementation system described here

TABLE 2 Abundances of type II cohesins

Protein	No. of type II cohesins per molecule	Protein abundance (nSpC) in:			
		DS19	WT	DS20	DS22
SdbA	1	35.9	30.1	67.7	46.2
OlpB	7	44.1	42.0	3.8	38.7
Orf2p	2	106.3	70.8	148.1	77.9
Total type II cohesins		557.4	466.0	390.8	473.5

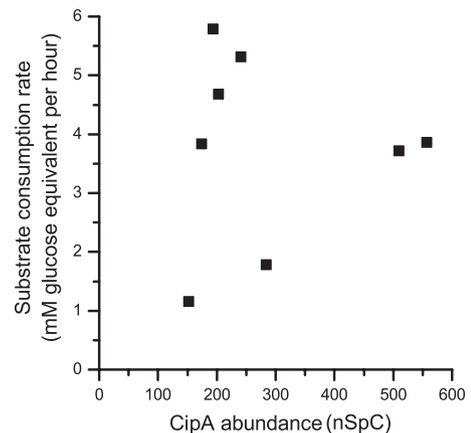


FIG 5 Comparison of substrate consumption rate with the abundance of the CipA scaffolding protein for duplicate fermentations with strains DS1, DS19, DS20, and DS22 grown on 10 g/liter Avicel.

will be useful for systematic understanding of the cellulolytic capabilities of *C. thermocellum*. The ability to express *cipA* from a replicating plasmid will enable the rapid exploration of the roles of its subcomponents, including elucidation of the function of individual modules of *cipA*, exploration of alternative cellulosomal architectures, and characterization of its noncomplexed cellulase system.

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