The Unique Binding Mode of Cellulosomal CBM4 from
Clostridium thermocellum Cellobiohydrolase A

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The crystal structure of the carbohydrate-binding module (CBM) 4 Ig fused
domain from the cellulosomal cellulase cellobiohydrolase A (CbhA) of
Clostridium thermocellum was solved in complex with cellobiose at 2.11 Å
resolution. This is the first cellulosomal CBM4 crystal structure reported to
date. It is similar to the previously solved noncellulosomal soluble
oligosaccharide-binding CBM4 structures. However, this new structure
possesses a significant feature—a binding site peptide loop with a
tryptophan (Trp118) residing midway in the loop. Based on sequence
alignment, this structural feature might be common to all cellulosomal
clostridial CBM4 modules. Our results indicate that C. thermocellum
CbhA CBM4 also has an extended binding pocket that can optimally bind to
celldextrins containing five or more sugar units. Molecular dynamics
simulations and experimental binding studies with the Trp118Ala mutant
suggest that Trp118 contributes to the binding and, possibly, the orientation
of the module to soluble celldextrins. Furthermore, the binding cleft
aromatic residues Trp68 and Tyr110 play a crucial role in binding to
bacterial microcrystalline cellulose (BMCC), amorphous cellulose, and
soluble oligodextrins. Binding to BMCC is in disagreement with the
structural features of the binding pocket, which does not support binding to
the flat surface of crystalline cellulose, suggesting that CBM4 binds the
amorphous part or the cellulose “whiskers” of BMCC. We propose that
clostridial CBM4s have possibly evolved to bind the free-chain ends of
crystalline cellulose in addition to their ability to bind soluble celldextrins.

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cellulose degradation

Introduction

In the United States, The Energy and Security
Independence Act of 2007 mandated volumetric
targets (2015 and 2022) for the integration of
renewable transportation fuels to reduce reliance
on foreign sources of oil. A recent technoeconomic
analysis of the corn-stover-to-bioethanol process
points to the high cost of pretreatment and enzymes
as the key hurdle that needs to be overcome in the
path to commercialization.1 Although considerable
effort has been expended towards understanding
and improving the performance of free (nonaggre-
gated) cellulases, the search for new and improved
cellulose-degrading enzymes from nature is an
obvious parallel strategy for meeting this cost goal.2
Some bacteria and a few anaerobic fungi use large
macromolecular complexes of enzymes called cellu-
osomes, in addition to free enzymes in solution, to
hydrolyze plant cell walls. Efficient cellulose conversion by these microorganisms is known to occur in complex populations, such as in the rumina of ungulates and in the guts of plant-matter-degrading insects, leading to the hypothesis that cellulosomes may have high specific activities on biomass. Cellulosomes contain a number of glycoside hydrolase (GH) enzymes with different functionalities that are bound to a single protein scaffold. These enzymes work together as a complex on the surface of the cell to degrade plant biomass. In contrast, fungal cellulase systems rely on a reducing-end-specific cellobiohydrolase (GH family 7) to achieve efficient cellulose hydrolysis. Cellulosomes use enzymes from GH families 5, 9, and 48, and do not contain a family 7 enzyme. In cellulosomal systems, it appears that members of GH family 9 (GH9) play an important role in cellulose conversion. One of the dominant family 9 catalytic domains in Clostridium thermocellum has been shown to be part of the cellobiohydrolase A (CbhA) multidomain complex, which is one of the largest cellulosomal enzymes known to date. CbhA displays seven distinct domains starting from the N-terminus: carbohydrate-binding module (CBM) 4, an Ig-like domain, GH9, two X1 modules, CBM3b, and a type I dockerin.

The protein domains thought to bind to cellulose and often linked to GHs were originally termed cellulose binding domains. In more recent years, this unnecessarily specific term was replaced with the more general CBM. Today, CBMs have been classified into 59 specific families based on amino acid similarity (see CAZy Web site). Although some ambiguity exists in the classification of some CBM families, most belong to type A, type B, or type C. Type A CBMs bind on the surfaces of insoluble polysaccharides (primarily cellulose) and display binding surfaces usually defined by a series of planar amino acid side chains (Tyr, Trp, and Phe) interacting with the hydrophobic surface of cellulose (i.e., the 1,0,0 face of cellulose 1β). Type B CBMs bind to the solution form of polysaccharides and usually display open cleft structures that are able to accommodate highly flexible polymers. Type B CBMs are probably unable to translate on the targeted polysaccharide chain in contrast to type A CBMs (i.e., CBM1 from Trichoderma reesei cellobiohydrolase I). Type C CBMs interact with the ends of polysaccharide chains and contain binding sites that are pockets or short closed tunnels. In general, CBMs are thought to play three possible roles: substrate proximity effect, substrate targeting, and substrate disruption.

It is our long-term strategy to study the structure-function relationships governing the action of this large cellulosomal enzyme complex. The crystal structures of the CbhA Ig-GH9 construct and CBM3b have been solved recently. CBM4, in general, is considered to be a type B CBM, binding putatively to single (and soluble) polysaccharide chains. CBM4 was predicted to play an important role in the degradation of plant cell wall. In this study, we solved the structure of the CBM4-Ig construct from C. thermocellum CbhA and conducted a detailed comparison of this new structure with previously known CBM4 structures such as laminaranase 16A from Thermotoga maritima, cellulase 9B from Cellulomonas fimii, and xylanase 10A from Rhodothermus marinus. Finally, we analyzed computationally the ability of CbhA CBM4 to bind to different types of oligosaccharides and also tested the binding behavior of five site-directed CBM4 mutants.

### Results and Discussion

#### Crystal structure of CbhA CBM4-Ig

The crystal structure of the C. thermocellum CbhA CBM4-Ig construct, in complex with cellobiose, was refined to a resolution of 2.11 Å with R and Rfree of 0.161 and 0.224, respectively. The asymmetric unit contained one molecule, with a nine-residue-long completely ordered native linker region (between Pro178 and Pro186) connecting the CBM4 and Ig domains. The structure of the linker region was ordered due to intertwining with the neighboring symmetry-related molecule (Fig. 1). It should be noted that the threonine/proline-rich regions of some cellulosomal linker peptides are decorated by O-linked glycosylation with galactopyranose when expressed in C. thermocellum. However, the C. thermocellum CbhA CBM4-Ig linker peptide has no threonines, but there is one water-exposed threonine side chain close to the linker peptide at position 176 and three prolines in the linker at positions 178, 184, and 186.

The contacts between symmetry-related molecules, as well as between the CBM4 domain and the Ig domain, appear to be nothing more than just regular crystal contacts (the interface area between CBM4 and Ig is 379 Å2 out of 13,781 Å2 in total) and most likely are not related to any biological function. The Ig structure was almost identical (rmsd of 0.69 Å for Ca atoms) to the previously described C. thermocellum CbhA Ig-GH9 structure (1RQ5). The CBM4 domain has a typical β-sandwich fold with two β-sheets containing five antiparallel β-strands each. Found in the CBM4 domain are two metal ions, which have been identified as magnesium ions based on their coordination geometry, distances, and the presence of MgCl2 in the crystallization solution (Fig. 2). One of the magnesium ions is located on the surface, interacting with five water molecules and one amino acid side chain (Glu19).
and another one is partially buried, interacting with one water molecule and five amino acid residues (Thr21, Asp23, Asn53, Lys54, and Asp166). The location of this partially buried magnesium ion corresponds with the position of calcium ions reported in other CBM structures.\(^8\) We note that both magnesium ions are not in proximity to the binding site.

### The binding pocket of CbhA CBM4 and the mode of binding to cellulose

The binding pocket of CbhA CBM4 is located on the concave face of one of the \(\beta\)-sheets, with three loops (Gly104-Trp114, Glu34-Gly39, and Lys63-Ser69) forming the walls of the binding cleft. A cellulose molecule was found in this pocket competing for this site with a glucose-1-phosphate molecule (Fig. 3). We assumed that the glucose-1-phosphate molecule was trapped in the binding cleft during protein expression (i.e., it was not replaced by cellulose completely during cocrystallization). This cellulose molecule was fixed in place by ring stacking with the Tyr110 and Trp68 side chains; by hydrogen bonding to Gln71, Arg73, and Lys102; and by several water-mediated H-bonds with Cys33, Asp35, Arg75, Gln105, Met106, Glu146, and His150 (Fig. 4).

### Structure comparison

In order to elucidate the structure–function relationship of CbhA CBM4, we conducted a comparison of related structures. Twenty proteins structurally similar to CBM4 and 372 proteins similar to Ig were found in the Protein Data Bank (PDB\(^\dagger\)) using the VAST server.\(^{15}\) Three hundred twenty-nine structures with Z-scores of \(\geq 5\) for the CBM4 domain and 439 structures with Z-scores of \(\leq 5\) for the Ig domain were identified using the Dali\(^{16}\) search tool. Manual comparison of the CBM4 domain with other CBM \(\beta\)-sandwich folds\(^{17}\) yielded even more structurally similar PDB entries. The CBM4 structural homologues include many CBMs (families 2, 3, 4, 6, 11, 15, 16, 17, 22, 27, 28, 29, 30, 32, 35, 36, and 44), GHs (families 7, 16, 11, and 12), and various other proteins sharing the same \(\beta\)-sandwich fold.

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\(\dagger\) www.rscb.org
Fig. 3. The omit electron density map of cellobiose (CBI; purple carbons) and glucose-1-phosphate (GP; gray carbons). This $F_o - F_c$ map has been calculated at 3σ after 10 cycles of REFMAC5 refinement without ligands.

Fig. 4. Interactions of cellobiose in the binding pocket of CbhA CBM4. Top view (a) with important interactions, and side view (b) showing the shape of the binding cleft. Nearby residues and water molecules are labeled and shown as sticks. Note the position of water molecules 311, 343, 344, and 364 below the cellobiose molecule.
Naturally, the most closely related structures found were those from various CBM4 domains. For example, the core β-sandwich folds found in the structures of different CBMs were generally very similar, and even some proteins with unrelated functions, such as virus envelope proteins, have distinctly similar folds. Fourteen structures with ligands bound in the same pocket were available from other members of CBM4 (1GU3, 1GUI, and 1K42), CBM15 (1GNY and 1US2), CBM16 (e.g., 2ZEX), CBM27 (e.g., 1PMH), and CBM29 (e.g., 1GWM). From these 14 structures, nine structures belonging to families 16, 27, and 29 showed the ligands bound "flat" in the pocket (with approximately a 90° rotation of the sugar plane compared to the cellohexaose found in the CBM4-Ig structure). The remaining five structures from CBM families 4 and 15 displayed ligands that are bound in approximately the same orientation that we observed. Cellulase 9B CBM4-1 in complex with cellopentaose from Ce. fimi (CfCBM4, 1GU3; rmsd of 1.62 Å for all Cα atoms), laminarinase 16A CBM4-2 in complex with laminariheptaose from T. maritima (TmCBM4, 1GUI; rmsd of 1.58 Å for all Cα atoms), and the NMR structure of xylanase 10A CBM4-2 from R. marinus (RmCBM4, 1K42; modeled xylan, rmsd of 2.53 Å for all Cα atoms) showed binding modes very similar to that of the CbhA CBM4 reported here (i.e., with aromatic residues at almost identical positions). These aromatic residues are thought to facilitate ring stacking with the sugar units of their respective ligands (Fig. 5).

Fig. 5. Trp118 of CbhA CBM4. Comparison to other CBM4s (a) and details of the Trp118 environment (b). CbhA CBM4 ribbon and carbon atoms are shown in gray, 1K42 is shown in magenta, 1GUI is shown in green, and 1GU3 is shown in orange. The conserved binding site residues of 1K42, 1GUI, and 1GU3, as well as the cellohexaose ligands of 1GUI and 1GU3, are shown as sticks.
comparisons immediately suggested that the binding pocket of CbhA CBM4 was cleft shaped (as seen in other type B<sup>8</sup> CBMs) with a similar mode of ligand binding (Fig. 4).

**The unique binding site tryptophan of clostridial CBM4s and the extended binding pocket**

A unique feature of the CbhA CBM4 structure is the loop between Trp114 and Tyr121, which includes a tryptophan at position 118 (Fig. 5). While comparing CBM4 sequences and structures, we could not find any aromatic residues equivalent to Trp118 in similar proteins with known three-dimensional structures. However, our sequence comparison revealed that this tryptophan or another aromatic residue can be found at the same or equivalent position in many CBMs (Table S1). All available sequences with an identity of over 31% relative to CbhA CBM4 are cellulosomal CBM4 modules from clostridia. A number of sequences from various species with identities below 31% might also have an aromatic side chain at this position, but this finding remains uncertain because there are gaps in the sequence alignments in this region. All the above sequences were annotated as CBMs or domains from family 9 or family 16 cellulases, laminarinases, or endoglucanases. It seems that Trp118 and this loop might be dominant features of cellulosomal CBM4 modules from clostridia.

Closer inspection of the *C. thermocellum* CbhA CBM4 ligand binding interactions reveals a series of side chains starting from Trp118, continuing through the binding cleft described above, and ending with Arg73 and Arg75 on the opposite side of the binding cleft (Fig. 4). Sequence conservation of this extended pocket between CBMs shows some similarity between these CBM4 domains. Specifically, aromatic residues at positions 68 and 110 are conserved, but Trp118 is unique to the *C. thermocellum* CbhA CBM4 structure (Table S2).

**The Trp118 loop environment**

In the structure reported here, Trp118 seems to be stabilized and protected from water by forming contacts with the equivalent area of the neighboring symmetry-related molecule. Flexibilities from normal-mode analysis calculations, using the generalized Born theory, indicated that this loop is fairly stable. The other two binding site residues in loops directly involved in substrate binding, Tyr110 and Trp68, are only somewhat flexible (Fig. 6). Trp118 is located in a fairly stiff loop, whereas Tyr110 is located in a flexible one. The flexibility of this loop is correlated, in all probability, to the recognition of various substrates. The first three normal modes of CBM4 show extreme motion of the catalytic cleft (Fig. S1) possibly related to the mechanism by which the CBM acquires the oligodextrin chain. It is acknowledged that the first 5–10 normal modes are, in most cases, enough to describe the overall dynamics of a given protein. Usually, these low-energy modes have the largest contributions to atomic fluctuations or the overall dynamics of a given protein.

Trp118 is indeed very well stabilized even in the absence of crystal contacts. It is positioned on the tip of the loop between residues Trp114 and Tyr121 (Fig. 4). A proline at position 120 appears to lock the C-terminal “hinge,” and a helical turn formed by residues 115–119 (with a hydrogen bond between OD1 of Asn115 and the main-chain nitrogen of Trp118) appears to further rigidify this peptide loop. The Trp118 side chain is firmly anchored in place by contacts with the side chains of Tyr100, Glu112, and Asn115, further increasing the stability (or inflexibility) of this loop.

**Docking and molecular dynamics simulations**

To confirm the existence of the extended binding pocket formed by Trp118, Tyr110, Trp68, Arg73, and Arg75, we conducted docking experiments and molecular dynamics (MD) simulations with different oligodextrins. The docking experiments with cellodextrin and oligoxylan ligands showed that this CBM4 can bind to both types of oligosaccharides. The best results were obtained when four water molecules (331, 343, 344, and 364) positioned between the ligand and protein molecules were included in the calculations. This result suggested an
important structural role for these water molecules. It is possible that the hydrogen bonds formed by these water molecules are important for ligand binding or specificity. However, Pell et al., in their studies of CBM15 domains that are structurally similar to CBM4 modules, reported that individual water molecules are not necessary for binding. The best ICM scores with a clear margin were obtained for cellopentaose (−27.55) and xylotetraose (−19.34) (Table S3). Together with manual inspection of the binding pocket area, this result indicated that the optimum number of interactions by CbhA CBM4 can be achieved with oligodextrins four to five sugar units in length. Another interesting observation is the predominance of the highest-scoring cellodextrins (larger than cellotriose) oriented with their nonreducing ends to the Trp118 side of the binding cleft.

Further investigation of two binding scenarios using MD simulations revealed stable stacking configurations (several nanoseconds during each simulation) between cellodextrin chains and Trp118 (Fig. 7; Figs. S2 and S3). Importantly, Trp118 displays a side chain in the same plane as the protein surface adjacent to the binding cleft (Fig. 5). This observation confirms the possible role of Trp118 and the existence of the extended binding pocket formed by Trp118, Tyr110, Trp68, Arg73, and Arg75 that can accommodate oligodextrins four to six sugar units long.

Fig. 7. MD simulation snapshot of CbhA CBM4 with cellohexaose. Docking experiments and MD simulation demonstrated that the binding pocket of CbhA CBM4 can bind to oligodextrins up to six sugar units long. The carbons of cellohexaose are shown in magenta.

Fig. 8. Binding of various mutants to different insoluble polysaccharides revealed by SDS-PAGE. (a) Before binding. (b) Binding to BMCC. (c) Binding to amorphous cellulose. (d) Binding to insoluble xylan of birchwood. The gel figures have been cropped to remove unnecessary parts or results.
Site-directed mutagenesis and binding assays

In order to elucidate the mechanism of the binding of CBM4 to oligodextrins, we generated five mutants. These mutants included three single-residue mutants (Trp68Ala, Tyr110Ala, and Trp118Ala), one double-residue mutant (Trp68Ala+Tyr110Ala), and one triple-residue mutant (Trp68Ala+Tyr110Ala+Trp118Ala). With these mutations, we probed the importance of binding pocket aromatic residues. These mutants were designed to evaluate the possible binding ability of Trp118 compared to the binding cleft aromatic residues Trp68 and Tyr110.

The binding of the mentioned mutants to bacterial microcrystalline cellulose (BMCC; a cellulose standard material that contains a high crystalline cellulose content19) and amorphous cellulose (Avicel treated by phosphoric acid20) was determined by affinity electrophoresis. The results showed that C. thermocellum CbhA CBM4 can bind to BMCC and amorphous cellulose (Fig. 8b and c). In comparison to wild-type CBM4, only the Trp68Ala and Tyr110Ala mutants showed reduced binding to both BMCC and amorphous cellulose. Thus, both Trp68 and Tyr110 are crucial for CbhA CBM4 binding to these forms of cellulose. The binding strength of both the double-residue mutant (Trp68Ala+Tyr110Ala) and the triple-residue mutant (Trp68Ala+Tyr110Ala+Trp118Ala) was also considerably reduced compared to wild type (became essentially nonexistent). This result further suggests that Trp68 and Tyr110 are essential for CbhA CBM4 binding to cellulose, but that Trp118 is not strictly required.

To understand binding to xylan in lignocellulose, we tested wild-type CBM4 and the five CBM4 mutants for their binding to soluble and insoluble forms of xylan. There was no apparent binding of the wild-type and mutant modules to (insoluble) birchwood xylan (Fig. 8d). However, the electrophoretic mobility of CBM4 changed in Native PAGE when soluble xylan from birchwood and switch grass was added (Fig. 9). Therefore, CBM4 can bind to the soluble xylan fraction in birchwood and switch grass. The Trp118Ala mutant showed a similar performance in binding to the wild-type module, implying again that Trp118 is not imperative for binding to soluble xylan. However, the electrophoretic mobility of the Trp68Ala and Tyr110Ala mutants was not significantly retarded by electrophoresis in the presence of soluble xylan, suggesting that Trp68 and Tyr110 are crucial for CBM4 binding to soluble xylan.

Binding to soluble oligodextrins

To assess the ability of CbhA CBM4 to bind to soluble oligodextrins, we performed experiments using fluorescence emission spectrum measurements. The effects of carboxymethyl cellulose
(CMC), xylobiose, xylohexaose, cellobiose, cellotetraose, cellopentaose, and cellohexaose were probed. Measurements with the native CBM4 showed binding only with CMC, cellopentaose, and cellohexaose (Fig. 10). Also, the ability of CBM4 to bind to these ligands was abolished if Trp68, Tyr110, or Trp118 was mutated to an alanine. This clearly demonstrates that Trp118 contributes to the binding of cellodextrins at least five sugar units in length.

The binding of ligands to Trp118 results in a strong signal, as shown by a comparison of the fluorescence emission spectra of native CBM4 and the Trp118Ala mutant. The fact that mutating any one of the binding cleft aromatic residues (Trp68 or Tyr110) also completely abolishes binding shows that Trp118 on its own cannot bind to cellodextrins; the strong “sandwich” binding interaction afforded by Trp68 and Tyr110 is required. This result is consistent with the outcome of affinity electrophoresis experiments, which showed that Trp118 is not needed for binding to cellodextrins.

The lack of binding signal with cellobiose, cellotetraose, xylobiose, and xylohexaose is inconsistent with the affinity electrophoresis experiments and our knowledge of the three-dimensional structure of the binding pocket. We believe that the reason for the apparent lack of signal in the case of these ligands is twofold. Firstly, from our structure, we can see that we have a tightly bound contaminant in the binding cleft with a high occupancy (∼50%) that could not be fully removed during protein purification or even cocrystallization with 100 mM cellobiose. This could have easily removed at least half of the signal by already binding Trp68 when what was meant to be an unliganded emission spectrum was measured. Secondly, it is possible that Trp68 is in an environment where emission changes do not occur upon ligand binding.

The role of Trp118

We have suggested that Trp118 is a common feature in clostridial CBM4 modules and that its side chain interacts with the sugar rings of cellodextrins. The orientation of its side chain (normal to the planar side of the binding pocket) is indeed similar to the orientation of critical aromatic residues in type A CBMs that are known to bind to the crystalline cellulose surface. This implies two possible roles for Trp118 in substrate binding: (1) binding to the crystalline cellulose surface and/or (2) guiding or stabilizing a soluble cellodextrin in the binding cleft. At first glance, the first hypothesis appears to have been supported by experimental results showing that both C. thermocellum CbhA CBM4 and cellulase K CBM4 (with a 76% sequence similarity to CbhA CBM4 and has the predicted unique tryptophan-containing loop) can bind to BMCC.\(^{21, 22}\)

We note, however, that BMCC is not purely crystalline cellulose.

With the use of targeted mutations, our results demonstrated that Trp118 is not required for the binding of CbhA CBM4 to BMCC, amorphous cellulose, or soluble xylan. Therefore, the binding of CbhA CBM4 to these substrates can be achieved by only two aromatic residues situated in the binding cleft (i.e., Trp68 and Tyr110). This appears to be in contradiction to what we have observed in the structural analysis of the binding pocket. The shape of the cleft, as well as the orientation of the aromatic side chains in it, does not support binding to the planar crystalline cellulose surface. Binding to crystalline cellulose requires a planar interactive protein modular face, as found in type A CBM modules, as well as a direct presentation of the interactive surface aromatic side chains to the cellulose surface.\(^{8, 23}\) Indeed, BMCC, as well as many other cellulose substrates, contains a significant amorphous content even though it is the most crystalline of the commonly used cellulose materials (crystalline content is 73.1–95.2%, depending on the measurement technique\(^ {19}\)). We suggest that the CBM4 interaction with BMCC occurs primarily through the amorphous regions of BMCC (which presumably expose cellodextrins), as well as through cellulose “whiskers” (i.e., free-chain ends extending from the crystalline core to the solution), rather than through the planar crystalline surfaces of the cellulose microfibril itself.

Fluorescence emission spectrum measurements clearly demonstrate that Trp118 contributes to the binding of CBM4 to soluble cellodextrins at least five sugar units in length. This experimental result is in good agreement with our computational results, where consistent and simultaneous ring stacking
between the soluble cellodextrins and Trp68, Tyr110, and Trp118 can be observed (Fig. 7; Figs. S2 and S3). For all three contacts to be possible at the same time, only certain helical conformations with the right orientation of the cellodextrin chain are feasible. Thus, computer simulations and fluorescence emission spectrum measurements support our second hypothesis for the function of Trp118 (i.e., to direct soluble cellodextrins with specific helical conformations to the binding cleft or towards the catalytic domain). For example, Simpson et al. demonstrated with mutational studies of *Ce. fimi* xylanase 11A CBM2 that the orientation of the aromatic side chains at the binding site can be important for distinguishing between substrates.  

By a single-point mutation, they succeeded in changing the orientation of one tryptophan side chain 90° from planar to a perpendicular orientation with respect to the CBM surface. This change was enough to make *Ce. fimi* xylanase 11A CBM2 specifically bind the helical conformation of xylan instead of its original substrate. This observation is consistent with our proposed function for Trp118—stabilizing a specific helical conformation of a cellodextrin substrate.

**The function of clostridial CBM4s**

Does Trp118 bind to soluble cellodextrins or can it also bind the flat surface of crystalline cellulose even though this amino acid is not necessary for binding to BMCC? Binding to cellulose “whiskers” still looks like the most probable binding mode for the CBM4 “cleft” aromatics Trp68 and Tyr110. Furthermore, binding of the module to the free ends of cellulose chains would presumably bring Trp118 close to the crystalline surface. Perhaps CbHA CBM4 has evolved to bind to these cellodextrin “whiskers” at a point near the crystalline cellulose surface, where Trp118 can bind to this planar surface. This scenario leads us to two possible binding modes for *C. thermocellum* CbhA CBM4: (1) binding of cleft residues to soluble cellodextrins both in solution and projecting from amorphous regions of cellulose, and (2) (weaker) binding of primarily Trp118 to the crystalline cellulose surface. Our experiments with CMC, cellohexaose, and BMCC confirm the first hypothesis, but do not disapprove the second hypothesis. Both are probably true. This result raises the possibility that clostridial CBM4 modules have evolved to bind to the free-chain ends of crystalline cellulose in addition to their ability to bind to soluble cellodextrins.

**Conclusions**

In this article, we have described the X-ray-diffraction-derived structure of the *C. thermocellum* CbhA CBM4-Ig domain in complex with cellobiose. The CBM4 module constitutes a β-sandwich fold and a cleft-like binding site typical of type B\(^6\) CBM modules. CBM4 thus would seem to prefer soluble substrates to crystalline cellulose. A unique feature of this and other clostridial CBM4 modules not found in any other CBM4 structure reported to date is a peptide loop displaying a strategically placed tryptophan residue (Trp118) at the tip of the loop. The side chain of Trp118 is located close to the binding pocket and seems to be in position to facilitate binding. Our results show that *C. thermocellum* CbhA CBM4 has an extended binding pocket formed by Trp118, Tyr110, Trp68, Arg73, and Arg75 that can optimally bind cellodextrins containing five or more sugar units. Site-directed mutagenesis and binding studies of this module demonstrated that Trp68 and Tyr110 are critical for the binding of CbhA CBM4 to soluble sugars. This observation, together with the structural analysis of the binding cleft and the known amorphous contents of the commonly used cellulose substrates,\(^3\)\(^\text{3}\)\(^\text{2}\)\(^\text{4}\)\(^\text{4}\)\(^\text{5}\) suggests that CBM4 binds the amorphous regions and/or the cellulose “whiskers” of BMCC. Our experiments indicate that clostridial CBM4 modules may have evolved to bind to the free-chain ends of crystalline cellulose in addition to soluble oligodextrins.

**Materials and Methods**

**Expression, purification, and crystallization**

CBM4-Ig (the dual domain of CBM4 and Ig) of *C. thermocellum* CbhA was amplified by PCR (the template is *C. thermocellum* genomic DNA) using the primers TCCGTGCAATGTTAGAAGATAATTCTTCGACT and CATCIGCTCGAGGATCTTGCTTTTGGCAATGTCATTG. The side-directed mutants of CBM4\(_{\text{CbhA}}\) were synthesized by GenScript (Piscataway, NJ). The PCR fragment of CBM4-Ig or synthesized genes was inserted into the plasmid pET22b (Novagen, Madison, WI) via NdeI and XhoI to generate expression plasmids. CBM4-Ig was overexpressed in *Escherichia coli* (BL21) (Stratagene, La Jolla, CA) with induction of 0.3 mM IPTG. The recombinant CBM4-Ig containing a C-terminal His-tag (6\(^\text{6}\) histidine) was purified with the QIAexpress Ni-NTA protein purification system (Qiagen, Valencia, CA), followed by size-exclusion chromatography with HiLoad Superdex 75 (26/60) (GE Healthcare, Piscataway, NJ) in buffer A [20 mM Tris–HCl (pH 7.0) containing 100 mM NaCl, 1 mM ethylenediaminetetraacetic acid, and 1 mM sodium azide] using the protocol recommended by the manufacturer. The purified fusion protein was concentrated with a Vivaspin 5K concentrator (Vivaproducts, Littleton, MA), and the protein concentration was measured by a NanoDrop UV spectrophotometer (NanoDrop, Wilmington, DE). Diffraction-quality crystals were obtained by sitting-drop vapor diffusion using a 96-well plate with Crystal Screen (Hampton Research, Aliso Viejo, CA). Fifty microliters of well solution was used with drops
containing 1 μL of well solution and 1 μL of protein solution. The recombiant CBM4-Ig protein was crystallized at 25 °C in buffer B [0.1 M Tris–HCl (pH 8.5), 0.2 M MgCl2, and 30% wt/vol polyethylene glycol 4000] containing 21 mg/mL protein kept in buffer A and supplemented with 100 mM cellbiose.

X-ray diffraction and structure determination

Before data collection, the CBM4-Ig crystal was soaked in 50%/50% (vol/vol) paraffin/paratone oil and flash frozen in a cold nitrogen gas stream at 100 K. Data collection was performed using a Bruker X8 MicroStar X-ray generator with Helios mirrors and Bruker Platinum 135 CCD detector. Data were indexed and processed with the Bruker Suite of programs (version 2008.1-0; Bruker AXS, Madison, WI). Intensities were converted into the Bruker Suite of programs (version 2008.1-0; Bruker 135 CCD detector. Data were indexed and processed with X-ray generator with Helios mirrors and Bruker Platinum collection was performed using a Bruker X8 MicroStar frozen in a cold nitrogen gas stream at 100 K. Data in 50%/50% (vol/vol) paraffin/paratone oil and flash containing 21 mg/mL protein kept in buffer A and

Table 1. X-ray data collection and refinement statistics

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<td></td>
<td>60.58, 94.32, 113.49</td>
</tr>
<tr>
<td></td>
<td></td>
<td>a, b, c (Å)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>90.0, 90.0, 90.0</td>
</tr>
<tr>
<td>Wavelength (Å)</td>
<td>1.54178</td>
<td></td>
</tr>
<tr>
<td>Temperature (K)</td>
<td>100</td>
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</tr>
<tr>
<td>Resolution (Å)</td>
<td>25.0–2.11</td>
<td>(2.15–2.11)</td>
</tr>
<tr>
<td>Rint a</td>
<td>0.161 (0.192) / 0.224 (0.273)</td>
<td></td>
</tr>
<tr>
<td>Observed reflections</td>
<td></td>
<td>77,757 (2171)</td>
</tr>
<tr>
<td>Unique reflections</td>
<td>19,105 (807)</td>
<td></td>
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<tr>
<td>Redundancy</td>
<td>4.07 (2.69)</td>
<td></td>
</tr>
<tr>
<td>I/σ(I)</td>
<td>10.83 (2.33)</td>
<td></td>
</tr>
<tr>
<td>Completeness (%)</td>
<td>99.7 (98.1)</td>
<td></td>
</tr>
</tbody>
</table>

Statistics for the highest-resolution bin are in parentheses, and the number of atoms includes alternative conformations.

Rint = ∑|I – ⟨I⟩| / ∑⟨I⟩, where I is the intensity of an individual reflection, and ⟨I⟩ is the mean intensity of a group of equivalents.

Ramachandran plot statistics (%) were calculated using MolProbity.

Structure analysis

The programs Coot, PyMOL, and ICMs were used for comparing and analyzing structures. Ramachandran plot statistics were calculated using MolProbity. The contact area between the CBM4 domain and the Ig domain was calculated using the program ICM after cutting the linker region between residues 177 and 185. Accessible surface was determined using a probe with a 0.7-Å radius to calculate the individual and combined surface areas. Sequence similarity searches were performed using NCBI BLAST. Structural similarity searches were completed using the VAST server and Dali. Rmsds were calculated using ICM (version 3.6-1e; Molsoft LLC, La Jolla, CA). Docking studies were performed using the virtual ligand screening module of the program ICM. After initial docking with oligodextrin (G10; oligodextrin with n number of glucose units; G2–G10) and oligoxylan (X10; oligoxylan with n number of xylose units; X2–X10) ligands, the binding pocket area was refined, with the most representative docking result obtained with cello-pentaose (G5) and four water molecules (331, 343, 344, and 364) deemed structurally relevant to ligand binding. Final docking studies were repeated three times, using the refined receptor with the four water molecules and a ligand library with oligodextrin and oligoxylan ligands containing G2–G6 and X2–X6, as described above. This approach was used in consideration of the initial docking results, which indicated that longer oligosaccharides could not be accommodated in the binding pocket area. In all docking experiments, a thoroughness of 5 was used, and calculations were repeated three times. The sequence conservation of the binding pocket was analyzed by taking the docking result with xylohexaose (X6) and selecting all residues with atoms inside a 7-Å sphere from it. This residue selection was then combined with a sequence alignment of similar structures that were structurally superimposed onto CBM4-Ig.

MD simulations

The CBM with bound cellulose was modeled with the CHARMM27 force field using CMAP correction for the protein. The new CHARMM force field for carbohydrates was applied to cellulose, the TIP3P water model was used for the solvent, and particle mesh Ewald summation was used for electrostatics with a nonbonded interaction cutoff of 10 Å. The system was
initially equilibrated for 1 ns in an MD simulation with an NVT ensemble and a time step of 2 fs. We performed six 15-ns MD simulations with an NPT ensemble and a time step of 2 fs with different random seeds starting from two initial cellohexaose configurations.

**Normal-mode analysis**

Normal-mode analyses were carried out with the molecular mechanics program package NAB46,47 (now part of Amber 10),48,49 using the parameter set parm99SB,50,51 and the pairwise approach of Hawkins et al. for the GB model.52,53 The monomeric CBM4 structure without the Ig module was minimized using the Limited-Memory Brodyen-Fletcher-Goldfarb-Shanno Truncated Newton Conjugate minimization technique to obtain an rms gradient below $1 \times 10^{-5}$ kcal/mol Å. This level of convergence is necessary to avoid contamination from translational and rotational modes into true internal modes. Diagonalization of the Hessian matrix was performed using ARPACK54 routines, in combination with Cholesky decomposition and inversion of the Hessian matrix, therefore providing better separation of eigenvalues to enhance convergence.

**Fluorescence emission spectrum measurements**

Steady-state fluorescence emission spectrum measurements were performed on a Jobin Yvon Fluoromax3 spectrofluorometer.55 Excitation of the protein was selective for tryptophan at 295 nm, and spectra were collected from 310 to 400 nm. The CBM samples had a concentration of 0.1 mg/mL in 20 mM Tris buffer (pH 7.0). Various ligands were added at the following concentrations: CMC of 0.1 mg/mL in 20 mM Tris buffer (pH 7.0). Various ligands were added at the following concentrations: CMC, 0.01%; and cellobiose, cellotetraose, cellopentaose, cellohexaose, xylbiose, and xyllohexaose were added to a final concentration of 1 mM. The contribution of buffer and ligand addition was measured and subtracted from the final results.

**Binding assays**

The binding assays followed the procedure of Xu et al.,56 where 0.3 mg of purified recombinant proteins was mixed with 1 mg of BMCC (a gift from the Cornell University, prepared in accordance with Jung et al.57), 0.8 mg of amorphous cellulose (treated with phosphoric acid58; Avicel PH101; FMC Corporation, Philadelphia, PA), and 1 mg of insoluble birchwood xylan (washed with water to remove partially soluble xylan; Sigma, St. Louis, MO). The suspension was brought to a final volume of 1.0 mL with 50 mM Tris (pH 8.0) buffer with 150 mM NaCl. These mixtures were kept at room temperature for 20 min with gentle rotation and centrifuged at 18900g for 6 min to separate the polysaccharide containing the bound protein from the unbound protein in the supernatant. The cellulose particles with bound protein were washed three times with vortexing in 1 mL of the previously mentioned buffer. Finally, the polysaccharides were resuspended in 50 and 100 μL of loading buffer and heated in a boiling water bath for 10 min. After centrifugation, 20 μL of the supernatant was subjected to SDS-PAGE. Gel retardation assays with insoluble ligands were performed by running a normal 12% Native PAGE supplemented with 0.12% soluble xylan. The native sample buffer and protein standard were purchased from Invitrogen (Carlsbad, CA). Soluble xylan from birchwood and switchgrass was prepared in accordance with Naran et al.56

**PDB accession number**

Atomic coordinates and structure factors have been deposited in the PDB with accession number 3K4Z.

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**Supplementary Data**

Supplementary data associated with this article can be found, in the online version, at doi:10.1016/j.jmb.2010.07.028

**References**


