Structural Insights into the Affinity of Cel7A Carbohydrate-binding Module for Lignin*

Kathryn L. Strobel, Katherine A. Pfeiffer, Harvey W. Blanch, and Douglas S. Clark

From the Department of Chemical and Biomolecular Engineering, University of California Berkeley, Berkeley, California 94720

Background: Non-productive adsorption to lignin hinders cellulase efficiency.

Results: 31 alanine mutants of the carbohydrate-binding module (CBM) exhibit highly correlated cellulose and lignin affinity.

Conclusion: The cellulose-binding face of CBM is responsible for the majority of lignin affinity, although four mutations were identified that selectivity decrease lignin adsorption.

Significance: The mutations studied inform future engineering efforts of fungal CBMs.

The high cost of hydrolytic enzymes impedes the commercial production of lignocellulosic biofuels. High enzyme loadings are required in part due to their non-productive adsorption to lignin, a major component of biomass. Despite numerous studies documenting cellulase adsorption to lignin, few attempts have been made to engineer enzymes to reduce lignin binding. In this work, we used alanine-scanning mutagenesis to elucidate the structural basis for the lignin affinity of Trichoderma reesei Cel7A carbohydrate binding module (CBM). T. reesei Cel7A CBM mutants were produced with a Talaromyces emersonii Cel7A catalytic domain and screened for their binding to cellulose and lignin. Mutation of aromatic and polar residues on the planar face of the CBM greatly decreased binding to both cellulose and lignin, supporting the hypothesis that the cellulose-binding face is also responsible for lignin affinity. Cellulose and lignin affinity of the 31 mutants were highly correlated, although several mutants displayed selective reductions in lignin or cellulose affinity. Four mutants with increased cellulose selectivity (Q2A, H4A, V18A, and P30A) did not exhibit improved hydrolysis of cellulose in the presence of lignin. Further reduction in lignin affinity while maintaining a high level of cellulose affinity is thus necessary to generate an enzyme with improved hydrolysis capability. This work provides insights into the structural underpinnings of lignin affinity, identifies residues amenable to mutation without compromising cellulose affinity, and informs engineering strategies for family one CBMs.

Lignocellulosic biomass is an abundant, low-cost resource for the renewable production of fuels and chemicals. Unfortunately, lignocellulose is highly resistant to enzymatic degradation, necessitating high enzyme loadings that raise the cost of second-generation biofuels (1). The recalcitrance of biomass stems in part from the presence of lignin. Lignin, a major component of lignocellulosic biomass, inhibits the efficiency and recyclability of cellulase enzymes. Numerous studies have found that removing lignin from biomass increases the final hydrolysis yield (e.g. Refs. 2–7), and adding supplemental lignin decreases the final hydrolysis yield (e.g. Refs. 8–14). Lignin impedes hydrolysis through two main mechanisms: physically blocking cellulose access (5, 6) and non-productively binding enzymes (15), thus lowering the effective enzyme concentration. Cellulase adsorption to lignin has been well documented for both cellulase mixtures (10, 16) and individual enzymes (9, 17, 18), although the structural basis of the interaction is not fully elucidated.

Most fungal cellulases consist of two domains: a catalytic domain (CD) and a carbohydrate-binding module (CBM) connected by a highly glycosylated, flexible linker (19). The CBM is responsible for increasing the effective concentration of the catalytic domain on the surface of its substrate (20). Fungal CBMs are highly conserved, 30–40 amino acid domains with a planar cellulose-binding surface. Three aromatic amino acids, arranged such that they align with a sequence of every other glucose unit on the surface of cellulose, are crucial for cellulose binding (21). Polar amino acids on the same face of the CBM are also important for binding to cellulose, presumably through the formation of hydrogen bonds (21). Along with binding to cellulose, the CBM is also responsible for the majority of lignin affinity. Fungal cellulases expressed without the CBM, or with the CBM removed, exhibit greatly reduced lignin affinity (17, 22, 23), highlighting the role of the CBM in lignin adsorption.

Studies of CBM lignin affinity have implicated hydrophobic and electrostatic interaction mechanisms. In a comparison of Trichoderma reesei Cel7A and Cel7B, Borjesson et al. (24) showed that Cel7B had higher affinity to lignin, perhaps due to the more hydrophobic surface on the flat face of the CBM together with an additional exposed aromatic residue on the rough face. Sammond et al. (18) found that the adsorption of enzymes to lignin surfaces correlates with solvent-exposed hydrophobic clusters. Increasing the carboxylic acid content, and therefore negative charge, of lignin has been shown to

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1 To whom correspondence should be addressed: 201 Gilman Hall, University of California, Berkeley, CA. Tel.: 510-642-2408; Fax: 510-643-1228; E-mail: clark@berkeley.edu.

2 The abbreviations used are: CD, catalytic domain; CBM, carbohydrate-binding module; Te, T. emersonii; Tr, T. reesei; Te-Tr, T. emersonii CD with T. reesei linker and CBM.
decrease the non-productive binding of cellulosomes and increase the enzymatic hydrolysis of cellulose (12), presumably due to increased repulsive electrostatic interactions between the enzymes and lignin. Adding negative charge to cellulosomes by succinylation has also been shown to reduce lignin inhibition (25). Improved hydrolysis of cellulose in the presence of lignin at high pH, along with reduced lignin binding at high pH, also demonstrates the importance of electrostatic interactions (26).

Despite the numerous studies documenting cellulosome adsorption to lignin, the structural basis has not been fully elucidated and few attempts have been made to engineer enzymes for reduced lignin affinity. Rahikainen et al. (26) showed that mutation of one aromatic amino acid on the flat face of the *T. reesei* Cel7A CBM from tyrosine to alanine reduced lignin affinity, whereas mutation from tyrosine to tryptophan increased lignin affinity. Interestingly, although the cellulose affinity of the alanine mutant was reduced to the level of the catalytic domain alone, the lignin affinity of the mutant remained greater than the catalytic domain, indicating that the mutated enzyme retained considerable lignin affinity.

In this work, we employed extensive alanine scanning mutagenesis of the entire *T. reesei* Cel7A CBM to identify residues involved in lignin adsorption and highlight locations where mutation may increase the selectivity for cellulose. Toward the overall goal of generating a lignin-resistant enzyme, we evaluated the most promising mutants for hydrolysis in the presence of lignin to investigate the extent to which lignin affinity must be reduced to decrease lignin inhibition.

**Experimental Procedures**

**Materials**—Microcrystalline cellulose PH-105 (Avicel), protease from *Streptomyces griseus*, 4-methylumbelliferone β-D-celllobioside, Tween 20, cellulase from *T. reesei* (Celluclast), and papain were purchased from Sigma. Accelerase 1500 was a gift from DuPont. *Aspergillus niger* β-glucosidase was purchased from Megazyme. Acid-pretreated *Miscanthus* was produced by two-step dilute acid pretreatment on the pilot scale by Andritz, Glens Falls, NY. Solids were washed with water to neutral pH and lyophilized before lignin preparation.

**Lignin Preparation**—Lignin residues were isolated from acid-pretreated *Miscanthus* by enzymatic hydrolysis followed by *S. griseus* protease treatment to remove bound enzyme. Pretreated, rinsed, lyophilized biomass was ball milled for 5 min (Kleco ball mill, Visalia, CA). Enzymatic hydrolysis was carried out at 50 °C, pH 5.0, with 4 weight % biomass and 0.25 ml of Accelerase 1500/g of biomass. The reaction mixtures were centrifuged and the supernatant replaced with fresh buffer and enzyme (0.25 ml of Accelerase/g original biomass) every 24 h. After 7 days of hydrolysis, the solids were separated by centrifugation and washed with acidified water (pH adjusted to 3.5 with HCl to avoid lignin solubilization). Bound protein was removed by an overnight protease treatment at 37 °C, pH 7.5, with 2 weight % solids and 0.3 mg/ml of protease. The solids were then washed with acidified water and heated in a boiling water bath for 60 min to deactivate any remaining protease. The solids were extensively washed with acidified water and freeze-dried.

**Lignin Analysis**—Compositional analysis of pretreated biomass and lignin was performed as previously described, with sugar concentrations determined by HPLC (Dionex) (27). Protein content in the solid samples was determined by total nitrogen analysis. The analysis was carried out at the University of California Berkeley Microanalytical Facility using a Perkin-Elmer 2400 Series II combustion analyzer.

**Purification of T. reesei Cel7A**—Cel7A was purified from Celluclast using a Q-Sepharose anion exchange column followed by a MonoQ ion exchange column. Prior to injection to the MonoQ column, the crude Cel7A was treated with 0.1% Tween 20 to disrupt nonspecific interactions that were found to exist between Cel7A and Cel7B. Fractions from the MonoQ were checked for purity by SDS-PAGE and an assay for contaminant endoglucanase activity. Assays contained 2% carboxymethyl cellulose and roughly 10 μM enzyme in 100 mM sodium acetate, pH 4.5. Assays were incubated at 50 °C overnight and endoglucanase activity was assessed by adding 2,4-dinitrosalicylic acid, incubating at 90 °C for 5 min, and visually inspecting for the presence of brown color in the reactions. Fractions showing no endoglucanase activity were pooled and buffer exchanged into 50 mM sodium acetate, pH 5.0.

**Isolation of the T. reesei CD**—The *T. reesei* CD was cleaved from the linker and CBM using the protease papain. Papain was activated by incubating a 28 mg ml⁻¹ solution with 2 mM dithiothreitol and 2 mM EDTA in 100 mM sodium phosphate, pH 7.0, for 30 min at room temperature. Cleavage reactions contained a (w/w) ratio of 1:100 papain to Cel7A in 50 mM sodium acetate, pH 5.0, and were incubated overnight at room temperature with stirring. The CD was purified from the protease and CBM using a Q-Sepharose anion exchange column.

**Small-scale Enzyme Production**—The *Talaromyces emersonii* cel7A CD or a fusion of *T. emersonii* Cel7A CD and the *T. reesei* cel7A linker and CBM was cloned into a high-copy number plasmid (pCu424 (28)) and produced in YVH10 pmr1Δ Saccharomyces cerevisiae (29). Mutations were made by site-directed mutagenesis (30). For screening all mutants, 20 ml of SC-Trp medium was inoculated with *S. cerevisiae* containing the cel7A gene and grown for 3 days at 30 °C, 220 rpm. Cultures were spun down at 5,000 × g for 5 min and resuspended in SC medium supplemented with 500 mM CuSO₄ and then cultured for protein production for 3 days at 25 °C, 220 rpm. Supernatant was collected and buffer exchanged into 50 mM sodium acetate, pH 5.0, using Amplicon spin concentrators (Millipore).

**Large-scale Enzyme Production and Purification**—For largescale production, 1 liter of SC-Trp medium was inoculated with *S. cerevisiae* containing the cel7A gene and grown for 3 days at 30 °C, 220 rpm. Cultures were spun down at 5,000 × g for 5 min and resuspended in YPD medium supplemented with 500 mM CuSO₄ and cultured for 3 days at 25 °C, 220 rpm. Enzyme purification was carried out using a two-step procedure beginning with anion exchange chromatography. Active fractions were combined and further purified by hydrophobic interaction chromatography. Pure fractions (single band via SDS-PAGE) were combined and buffer exchanged into 50 mM sodium acetate buffer, pH 5.0.

**Enzyme Quantification**—Purified enzymes were quantified by absorbance at 280 nm using the calculated molar extinction coefficient.
coefficient of 76,240 M\(^{-1}\) cm\(^{-1}\) for the fusion enzyme and 71,120 M\(^{-1}\) cm\(^{-1}\) for the catalytic domain. Unpurified enzyme and enzyme remaining in the supernatant after binding assays were quantified by their activity on a soluble substrate, 4-methylumbelliferyl β-D-cellobioside. The specific activity of 10.2 μmol of MU (μmol of enzyme/min)\(^{-1}\) was measured for the purified wild type, catalytic domain, and mutant chimeras. Assays were conducted in a total volume of 100 μl with 1 mM 4-methylumbelliferyl β-D-cellobioside, pH 5.0, at 45 °C for 8 min, followed by a denaturation step at 98 °C for 2 min. Sodium hydroxide was added to a final concentration of 0.1 M and the fluorescence was measured on a Paradigm plate reader (Beckman Coulter) with excitation of 360 nm and emission of 465 nm.

Avicel and Lignin Binding Screen—Unpurified, buffer-exchanged mutants were normalized to the same concentration of active Cel7A. Binding to Avicel and lignin was measured in 50 mM sodium acetate buffer, pH 5, using 15 mg/ml of Avicel or lignin and 125 mM enzyme. Experiments were conducted in a total volume of 70 μl in a rotating shaker (300 rpm) at room temperature. An initial time course with the wild type enzyme was measured to determine the time required for binding to reach equilibrium. Equilibrium was reached after ~30 min. Subsequent assays were performed for 1 h. After equilibration, solids were separated by centrifugation and the supernatant was analyzed for free, active enzyme using 4-methylumbelliferyl β-D-cellobioside. The amount of bound enzyme was calculated from the difference between added enzyme and enzyme remaining in solution. The values plotted in Fig. 2 are bound enzyme divided by enzyme remaining in solution. The values reported in duplicate samples and the errors are calculated as standard deviation.

The one-site Langmuir isotherm at low free protein concentrations can be written as,

\[
\Gamma = \Gamma_{\text{max}} K_A C = \alpha C \tag{Eq. 1}
\]

where \(\Gamma\) is the amount of adsorbed protein, \(\Gamma_{\text{max}}\) is the amount of adsorbed protein at saturation, \(K_A\) is the Langmuir affinity constant, \(C\) is the concentration of free protein, and \(\alpha\) is the partition coefficient.

Adsorption Isotherms—Adsorption isotherm measurements were performed in 50 mM sodium acetate buffer, pH 5, using 10 mg/ml of Avicel or lignin and initial enzyme concentrations ranging from 0.015 to 4.0 μM. Experiments were conducted using the same conditions as the binding screen. A one-site Langmuir adsorption isotherm (Equation 2) was fit to the binding data by non-linear least squares optimization to determine \(K_A, \Gamma_{\text{max}}\), and their corresponding standard deviations.

\[
\Gamma = \frac{\Gamma_{\text{max}} K_A C}{1 + K_A C} \tag{Eq. 2}
\]

### Results

#### Isolation and Characterization of Lignin from Pretreated Miscanthus

Lignin used in this study was isolated from acid-pretreated Miscanthus using a commercial cellulase mixture (Accelerase 1500, DuPont) to ensure that the lignin was as similar as possible to lignin present in an industrial hydrolysis reaction. Excess enzyme and long digestion times were used to remove all accessible carbohydrates. The solid residue was then treated with S. griseus protease to remove bound enzymes. The compositions of the starting biomass and resulting isolated lignin are presented in Table 1. The Klasson lignin, remaining carbohydrates, and protein content are similar to protease-treated lignin used in previous studies (9–11, 17, 26).
Lignin Inhibition of Cel7A and Isolated Catalytic Domain—We compared the lignin inhibition of natively expressed T. reesei Cel7A to that of the chimera used for engineering in this study, recombinantly expressed T. emersonii Cel7A catalytic domain fused to the T. reesei Cel7A linker and carbohydrate-binding module (Te-Tr chimera). Fig. 1 shows the hydrolysis of Avicel, Avicel in the presence of supplemental lignin, and acid-pretreated Miscanthus by each enzyme and the corresponding isolated CD.

The Te-Tr chimera was less inhibited by lignin than T. reesei Cel7A. The addition of lignin to Avicel decreased the glucose generated by ~15% for the Te-Tr chimera, and 73% for the full-length T. reesei enzyme. The T. emersonii CD was not inhibited by supplemental lignin, indicating that the inhibition of the Te-Tr chimera is due entirely to the linker and CBM, our mutagenesis target. The T. reesei CD generated 38% less sugar in the presence of lignin. Although both CDs were less inhibited by lignin, they generated significantly less sugar than the corresponding full enzyme.

CBM Engineering Approach—We used alanine-scanning mutagenesis to identify residues that contribute to lignin affinity. The isolated T. emersonii catalytic domain, wild type Te-Tr chimera, and 31 Te-Tr mutant chimeras, each with one position in the CBM changed to alanine, were produced in S. cerevisiae strain YVH10 pmr1Δ. This strain includes an overexpressed protein-disulfide isomerase for increased Cel7A titers and a knock-out of the PRM1 gene for reduced hyperglycosylation (29). Each residue in the CBM was mutated to alanine except cysteine residues involved in disulfide bonds and native alanine residues.

Binding Screen—The Cel7A mutants were secreted into defined medium, buffer exchanged, and screened for cellulose and lignin binding without purification. Binding was quantified by measuring the Cel7A activity remaining in the supernatant using a soluble substrate, 4-methylumbelliferyl β-D-cellobioside, which permitted specific detection of Cel7A among other secreted proteins. Each mutant was screened for binding at low enzyme concentration, in the range where the Langmuir isotherm is linear, permitting direct calculation of the partition coefficient from the amount of bound and free enzyme (Equation 1 under “Experimental Procedures”).

Impact of Point Mutations on CBM Adsorption to Avicel—The cellulose partition coefficient of each mutant is shown in Fig. 2 (light bars) and the CBM structure is shown in Fig. 3. The CD alone has significantly less affinity for cellulose than the chimera containing the wild type CBM, as expected. All mutants have an Avicel partition coefficient in the range between the wild type and CD alone. As previously reported, mutating the three tyrosines on the planar face of the CBM...
(Tyr-5, Tyr-31, and Tyr-32) as well as polar residues on the same face (Asn-29, Gln-34) to alanine significantly reduced cellulose binding (21). Mutation of Gln-7, Gly-10, Tyr-13, Gly-15, Gly-22, and Leu-36 to alanine also significantly reduced binding to cellulose (<30% of wild-type affinity).

Impact of Point Mutations on CBM Adsorption to Lignin—The lignin partition coefficient of each mutant is shown in Fig. 2 (dark bars). The T. emersonii CD has negligible lignin affinity when compared with the fusion containing the T. reesei linker and CBM. We found that mutation of Tyr-5, Gln-7, Gly-10, Tyr-13, Gly-15, Gly-22, and Leu-36 to alanine also significantly reduced binding to cellulose (210%).

Impact of Point Mutations on CBM Selectivity—For hydrolysis of lignocellulose, it is important to consider the selectivity of adsorption to cellulose over lignin. As seen in Fig. 4, lignin and cellulose affinity of the mutants were highly correlated ($R^2 = 0.92$), although a few outliers exhibited a selective reduction in cellulose or lignin adsorption. Q34A, I11A, and N29A reduced cellulose selectivity, whereas Q2A, H4A, V18A, and P30A increased cellulose selectivity.

Adsorption Isotherms of Selected Mutants—The wild type enzyme, CD alone, and the alanine mutants with increased selectivity toward Avicel were characterized further. The partition coefficient calculated from the binding screen does not provide enough information to predict binding behavior at higher concentrations, where the Langmuir isotherm is no longer linear. Therefore, full Langmuir isotherms were measured and fit using non-linear least squares regression to determine the adsorption at saturation ($\Gamma_{max}$) and the Langmuir adsorption constant ($K_A$). The isotherms are shown in Fig. 5 and the model parameters are listed in Table 2.

Qualitatively, the mutations decreased lignin adsorption more than Avicel adsorption, as seen in Fig. 5. The partition coefficients to Avicel derived from the Langmuir isotherms ($K_A / \Gamma_{max}$) are similar for the wild type and mutant enzymes (Table 2), as expected based on data from the binding screen. The saturating amount of enzyme adsorbed to lignin, $\Gamma_{max}$, is lower for each of the mutants than the wild type enzyme, although the affinity constants and partition coefficients are within error of the wild type enzyme.

| TABLE 2 |
| --- |
| Langmuir adsorption parameters |
| | Avicel | | Lignin | |
| | $K_A^a$ | $\Gamma_{max}$ | $\alpha$ | $K_A^a$ | $\Gamma_{max}$ | $\alpha$ |
| | $\mu$mol^{-1}$ | $\mu$mol/g | liter/g | $\mu$mol^{-1}$ | $\mu$mol/g | liter/g |
| Wild type | 1.10 ± 0.11 | 0.27 ± 0.01 | 0.29 ± 0.03 | 1.07 ± 0.21 | 0.21 ± 0.02 | 0.23 ± 0.05 |
| CD | 1.34 ± 0.17 | 0.059 ± 0.003 | 0.08 ± 0.01 | 2.52 ± 0.77 | 0.013 ± 0.001 | 0.03 ± 0.01 |
| Q2A | 1.74 ± 0.27 | 0.18 ± 0.01 | 0.32 ± 0.05 | 1.26 ± 0.33 | 0.15 ± 0.02 | 0.19 ± 0.05 |
| H4A | 1.90 ± 0.26 | 0.16 ± 0.01 | 0.31 ± 0.05 | 1.46 ± 0.32 | 0.11 ± 0.01 | 0.16 ± 0.04 |
| V18A | 0.82 ± 0.21 | 0.28 ± 0.04 | 0.23 ± 0.06 | 1.19 ± 0.30 | 0.17 ± 0.02 | 0.20 ± 0.05 |
| P30A | 0.92 ± 0.17 | 0.24 ± 0.02 | 0.22 ± 0.05 | 2.49 ± 0.75 | 0.08 ± 0.01 | 0.21 ± 0.07 |

$^a$ $K_A$, Langmuir adsorption constant; $\Gamma_{max}$, adsorption at saturation; $\alpha$, partition coefficient ($\alpha = K_A / \Gamma_{max}$) indicated with their standard deviation.
Hydrolysis of Avicel in the Presence of Lignin and Acid-pre-treated Miscanthus—To examine the impact of increased cellulose specificity on hydrolysis in the presence of lignin, the wild type, CD, and mutant enzymes were tested for their ability to hydrolyze Avicel, Avicel with added lignin, and acid-pre-treated Miscanthus (Fig. 6). Although each of the mutants displayed increased selectivity toward binding cellulose, none generated more glucose than the wild type enzyme.

Predicting the Amount of Enzyme Bound to Cellulose—We sought to explain the hydrolysis results and predict levels of binding selectivity necessary for hydrolytic improvement by using the Langmuir adsorption model. The first step of cellulose hydrolysis is adsorption to the surface. Predicting the amount of enzyme adsorbed to cellulose can therefore lend insight into the performance of enzymes with different affinities. Using the Langmuir parameters (\(K_A\) and \(\Gamma_{max}\)) shown in Table 2, the approximate amount of enzyme bound to Avicel under the hydrolysis conditions employed in this study was calculated for the wild type chimera, CD, and the four selected mutants. This prediction is restricted to hydrolysis of pure cellulose with supplemental lignin, due to the use of Langmuir parameters measured on the pure substances.

As shown in Fig. 7, none of the mutants are predicted to have significantly more enzyme bound to Avicel than the wild type, which explains why none of the mutants generated more glucose during the hydrolysis assays. The catalytic domain is predicted to bind in similar amounts to cellulose in the presence and absence of lignin, as expected given that it performed equally well in hydrolysis assays with and without lignin (Fig. 6). The full-length wild type enzyme and each of the mutants are predicted to have between 3 and 55% less enzyme bound to cellulose when lignin is present, which covers the range of lignin inhibition observed in the hydrolysis assays.

Discussion

Non-productive adsorption of cellulases to lignin reduces the effective enzyme concentration during biomass hydrolysis, increasing the overall cost of producing biofuels. Although it is well known that the CBM drives this non-productive adsorption, the mechanisms of CBM lignin affinity have not been fully elucidated. We employed alanine-scanning mutagenesis of the \(T. reesei\) Cel7A CBM to reveal the structural basis of lignin affinity and identify residues where mutation increases the selectivity for cellulose over lignin. Our studies focused on \(T. reesei\) Cel7A because it is the most abundant enzyme in many commercial cellulase mixtures. Each CBM mutant was produced with a linker and CD to ensure realistic binding conditions and enable hydrolysis studies. We used the \(T. reesei\) Cel7A CBM and linker, and the \(T. emersonii\) Cel7A CD. This CD is homologous to \(T. reesei\), and unlike the \(T. reesei\) enzyme, expresses well in yeast, making enzyme production and screening tractable.

Although the Te-Tr chimera was less inhibited by lignin than the native \(T. reesei\) enzyme, the inhibition of the chimera was entirely due to the CBM, making it a well suited construct for CBM engineering. Surprisingly, the \(T. reesei\) Cel7A CD was also inhibited by supplemental lignin. This result is unexpected given the general consensus that the CBM is responsible for the majority of non-productive adsorption to lignin. Engineering both domains of \(T. reesei\) Cel7A may be required to fully alleviate lignin inhibition. The highly homologous \(T. emersonii\) CD was not inhibited by lignin, perhaps due to this higher melting temperature of the enzyme (31, 32). Engineering the CD for greater thermal stability may therefore be a promising strategy to reduce lignin inhibition.

The wild type Te-Tr chimera, isolated \(T. emersonii\) CD, and 31 mutants, each with one position in the CBM mutated to alanine were produced and screened for binding to microcrystalline cellulose (Avicel) and lignin. The isolated CD had dramatically reduced affinity for lignin, consistent with previous reports that the CBM is responsible for the majority of lignin binding (17, 22).

Mutation to alanine of the three tyrosines (Tyr-5, Tyr-31, and Tyr-32) on the planar face of the CBM significantly reduced cellulose and lignin binding. These tyrosine residues are known to be essential for cellulose affinity (21), although only Tyr-32
Structural Basis of Cellulase Lignin Affinity

has been evaluated previously for lignin binding (26). According to NMR data, mutation of Tyr-31 and Tyr-32 to alanine does not cause significant structural changes, and mutation of Tyr-5 to alanine causes only minor structural changes (21), indicating that the reductions in binding are due to diminished interactions between these residues and the lignin surface. The three tyrosine residues likely contribute to lignin adsorption through Pi stacking interactions with phenolic groups on the lignin surface. This result supports the hypothesis that the planar, cellulose-binding face of the CBM contributes significantly to lignin affinity.

Mutation of Gln-7 on the cellulose-binding face of the CBM also greatly reduced the binding to cellulose and lignin. This residue has been predicted to interact with cellulose through hydrogen bonding (33), although its impact on affinity had not been previously assessed experimentally. Loss of these hydrogen-bonding interactions likely causes the decrease in cellulose affinity and may also cause the reduction in lignin affinity. However, the decrease in affinity to both cellulose and lignin may also be due to structural alterations in the CBM caused by this mutation.

Mutation of an internal tyrosine (Tyr-13) to alanine also significantly decreased affinity for both cellulose and lignin. Tyr-13 has been suggested to undergo a conformational change when the CBM is bound to cellulose, moving away from the protein interior to the wedge surface to establish van der Waals contact with the cellulose chain (34, 35). Mutation of Tyr-13 to alanine may decrease affinity due to loss of these van der Waals contacts, although disruptive structural changes (caused by mutation of a large, internal hydrophobic residue) may also play an important role.

Interestingly, Q34A, which had greatly reduced cellulose affinity, retained lignin affinity similar to that of the wild type enzyme. This residue is predicted to interact with the surface of cellulose through hydrogen bonding. NMR data indicate that the Q34A mutation results in only small conformational perturbations (21). Therefore, the selective reduction in binding to cellulose implies that this hydrogen bond is important for cellulose, but not lignin, affinity. Asn-29 and Ile-11, also predicted to interact with cellulose through hydrogen bonding (35), also retained a greater fraction of their lignin affinity than their cellulose affinity. The N29A mutation has been shown to disrupt the stability of the loop containing Tyr-31 and Tyr-32 (21), and the I11A mutation has not been assessed for structural perturbations. Therefore, their role in interacting with the surface of cellulose or lignin is more difficult to ascertain. Nevertheless, these results suggest that hydrogen bonding is more important for binding to cellulose than binding to lignin. The hydrogen-bonding residues at specific positions in the CBM are able to align with the uniformly spaced hydroxyls in crystalline cellulose. In contrast, the free-radical polymerized lignin structure lacks the precise repeating structures of crystalline cellulose, making specific hydrogen bonding on the CBM surface less likely.

The remaining mutants with severely reduced affinity for cellulose and lignin are G10A, G15A, G22A, and L36A. It is unlikely that these glycine residues interact with the surface of cellulose or lignin. Gly-10 and Gly-22 are completely conserved in an alignment of homologous family one CBMs and Gly-22 is conserved as either a glycine or proline. Therefore, the decreased affinity is most likely due to structural changes caused by mutation to alanine. Mutation of Leu-36, an internal hydrophobic residue, also likely results in structural perturbations that are responsible for decreased affinity. Although these mutants had significantly reduced lignin affinity compared with the wild type chimera, they retained much greater lignin affinity than the isolated CD. This indicates that the mutated CBMs have residual lignin affinity or that the linker contributes to affinity. Linker mutations that decrease lignin inhibition are described in a patent (36), indicating that the linker does participate in non-productive adsorption to lignin.

Four mutants with increased selectivity for cellulose in the binding screen (Q2A, H4A, V18A, and P30A) were further characterized by constructing full Langmuir isotherms. Although the Langmuir model does not elucidate the complex mechanisms involved in protein adsorption to solid surfaces, the model describes the adsorption data well and is a useful tool for evaluating cellulase adsorption (37).

As expected from the binding screen, the Te-Tr wild type and mutants had similar Langmuir affinity and saturation constants for cellulose. The mutant enzymes exhibit similar lignin binding compared with the wild type enzyme at low enzyme concentration (leading to similar values for $K_a$) but decreased binding at the higher concentrations, leading to reduced values for $\Gamma_{\text{max}}$. Given the heterogeneous structure of the lignin, it is likely that there are multiple binding sites on the lignin surface with different affinities (23). Mutations in the CBM may impact interactions with one type of site more than others, resulting in decreased binding only over certain enzyme concentration ranges. The initial concentrations of enzyme employed here (up to 0.4 $\mu$mol (24 mg) Cel7A enzyme/g of cellulose) cover the range of typical enzyme loadings used in hydrolysis of lignocellulosic biomass, and therefore describe the binding behavior under industrially relevant conditions.

We next studied the impact of increased cellulose specificity on hydrolysis of Avicel in the presence of lignin and pretreated Miscanthus. None of the mutants outperformed the wild type enzyme, likely because lignin affinity was not reduced enough to impact hydrolysis. To test this hypothesis, we used the Langmuir model to predict the amount of each enzyme bound to cellulose under the hydrolysis conditions. Because the first step of hydrolysis is adsorption to cellulose, the calculated binding should reflect the relative hydrolytic performance of similar enzymes with different affinities. This analysis assumes that the binding to both Avicel and lignin is reversible, the amount of Avicel and lignin is constant, and that the Langmuir isotherms collected at room temperature accurately describe the binding behavior at 60 °C, the hydrolysis temperature. Although limited, this approach enables comparison of enzymes with different affinities, and estimates the magnitude of change in specificity necessary to generate an enzyme with less lignin inhibition. None of the mutants were predicted to bind in larger quantities to cellulose than the wild type enzyme, supporting the conclusion that lignin affinity must be reduced further to improve hydrolysis.
A central conclusion from the combined results of this study is that improving cellulose-hydrolyzing activity in the presence of lignin will require an even greater reduction in CBM/lignin affinity accompanied by a high level of cellulose binding. Although the mutants’ cellulose and lignin affinity were highly correlated, outliers with increased specificity toward either cellulose or lignin were identified that gave insights into the mechanisms of adsorption. The four residues where mutation to alanine increased selectivity (Gln-2, His-4, Val-18, and Pro-30) are consequently better hydrolytic performance in the presence of lignin.

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Kathryn L. Strobel, Katherine A. Pfeiffer, Harvey W. Blanch and Douglas S. Clark

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