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NMR Analysis on Molecular Interaction of Lignin with Amino Acid Residues of Carbohydrate-Binding Module from *Trichoderma reesei* Cel7A

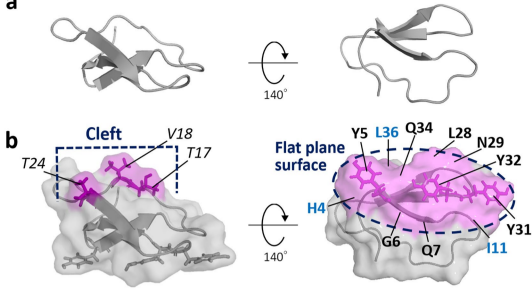
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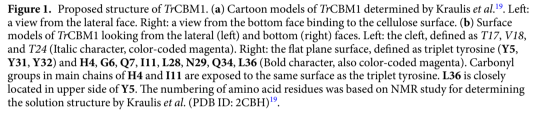
Lignocellulosic biomass is anticipated to serve as a platform for green chemicals and fuels. Nonproductive binding of lignin to cellulolytic enzymes should be avoided for conversion of lignocellulose through enzymatic saccharification. Although carbohydrate-binding modules (CBMs) of cellulolytic enzymes strongly bind to lignin, the adsorption mechanism at molecular level is still unclear. Here, we report NMR-based analyses of binding sites on CBM1 of cellobiohydrolase I (Cel7A) from a hyper-cellulase-producing fungus, *Trichoderma reesei*, with cellohexaose and lignins from Japanese cedar (C-MWL) and *Eucalyptus globulus* (E-MWL). A method was established to obtain properly folded *Tr*CBM1. Only *Tr*CBM1 that was expressed in freshly transformed *E. coli* had intact conformation. Chemical shift perturbation analyses revealed that *Tr*CBM1 adsorbed cellohexaose in highly specific manner via two subsites, flat plane surface and cleft, which were located on the opposite side of the protein surface. Importantly, MWLs were adsorbed at multiple binding sites, including the subsites, having higher affinity than cellohexaose. G6 and Q7 were involved in lignin binding on the flat plane surface of *Tr*CBM1, while cellohexaose preferentially interacted with N29 and Q34. *Tr*CBM1 used much larger surface area to bind with C-MWL than E-MWL, indicating the mechanisms of adsorption toward hardwood and softwood lignins are different.

Lignocellulosic biomass is the most abundant renewable carbon resource. It consists of structural polysaccharides, cellulose, and hemicelluloses coated with a heterogeneous aromatic polymer, lignin¹. Recently, the production of bio-based fuels and chemicals from lignocellulosic biomass has attracted increasing attention due to the depletion of fossil resources and environmental issues². To produce biofuels and chemicals by enzymatic saccharification and the fermentation of lignocelluloses, it is necessary to realize pretreatments exposing plant cell wall polysaccharides and subsequent hydrolysis of polysaccharides with a cellulolytic enzyme cocktail simultaneously or prior to fermentation. Highly efficient enzymatic saccharification of lignocellulose with cellulolytic enzymes in a hydrolytic process is a primary key step in achieving lignocellulosic biorefinery process. Typical fungal cellulolytic enzymes, such as cellobiohydrolase and endoglucanase, are composed of catalytic domain (CD) and carbohydrate-binding modules (CBMs) connected with highly glycosylated linker. CBMs play a role in bringing catalytic domains in close proximity to the substrate to improve enzymatic activity³. However, CBMs of polysaccharide hydrolases also bind to lignin. The efficiency of enzymatic saccharification, therefore, is strongly decreased⁴. Because the pretreated biomass is usually hydrolyzed by cellulolytic enzymes in the presence of lignin fragments, methods have been extensively explored⁵⁻⁸ for protecting enzymes from the unfavorable binding with lignin. The approaches include the addition of masking agents, such as bovine serum albumin⁵, polyethylene glycol⁶, and surfactants⁷, as well as the incorporation of ionic functional groups into lignin⁸. However,

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no fundamental theories have discussed how to alter the enzyme to avoid the unfavorable binding with lignin because the binding sites of lignin in enzymes are still not understood clearly.

Filamentous fungus Trichoderma reesei is known as a hyper producer of cellulolytic enzymes and widely used for commercial-scale production of cellulases and hemicellulases. Up to 60% of totally secreted cellulase is cellobiohydrolase I (TrCel7A) that bears family 1 CBM as the C-terminal domain (Fig. 1)9. Hence, there is a need for detailed understanding of the interaction between TrCBM1 and lignin to solve the nonproductive binding issue and to establish a low-cost, highly efficient enzymatic saccharification process. However, both homologous and heterologous expressions of TrCBM1 as well as its isolation are difficult due to its small molecular size (around 5-kDa). Because of these challenges, there has been no reports on the identification of amino acid residues of TrCBM1 that are involved in binding with lignin without using site-directed mutagenesis that may cause conformational changes of such a small protein. It should be noted that the comparison of intact and TrCBM1-deficient TrCel7A gives indirect information due to the interference of glycosylated linker¹⁰.

NMR titration analysis, such as chemical shift perturbation (CSP), is a powerful experimental strategy to identify substrate-binding sites of proteins at amino acid residue resolution¹¹. CSP enables comprehensive analysis of interaction sites on the proposed structure of a protein without crucial conformational change. This approach has been used previously for binding site analysis of CBMs with poly- and oligosaccharides, including the interaction site and binding specificity between CBM56 and β -1,3-glucan¹², CBM32 and chitosan oligosaccharides¹³, as well as CBM6 and xylohexaose¹⁴

In this study, we applied CSP to analyze the interaction sites of TrCBM1 against lignins from Japanese cedar and Eucalyptus globulus, using¹⁵N-labeled TrCBM1 prepared as a single protein with correct folding. In addition, interaction of TrCBM1 with cellohexaose was also analyzed by CSP to elucidate differences in the binding mechanisms of TrCBM1 between polysaccharides and lignin. Enhanced understanding these differential interactions will lead to fundamental theory to develop hydrolases having high specificities toward carbohydrates having decreased binding affinity to lignin.

Results

Expression and purification of ¹⁵N-labeled *TrCBM1*. ¹⁵N-labeled His tag-*TrCBM1*-GFP fusion protein was expressed using Escherichia coli BL21(DE3) (Fig. 2a). TrCBM1 was cleaved off from His tag and GFP by proteolytic cleavage using enterokinase and thrombin, respectively. Finally, TrCBM1 was purified to a single protein as demonstrated in SDS-PAGE (Fig. 3a) and MALDI-TOF-MS (Fig. 3b). The MALDI-TOF-MS spectrum gave the evidence that the obtained ¹⁵N-labeled *Tr*CBM1 possessed the correct molecular mass of 5255 expected for ¹⁵N incorporated protein (Fig. 2b).

The structures of ¹⁵N-labeled TrCBM1 were assessed by observing the signal patterns of 2D ¹H-¹⁵N SOFAST-HMQC spectra¹⁵. The ¹⁵N-labeled TrCBM1 sample that was prepared using E. coli whose competent cell was stocked for more than five months showed a mixture of 2D ¹H-¹⁵N SOFAST-HMQC spectra for both а

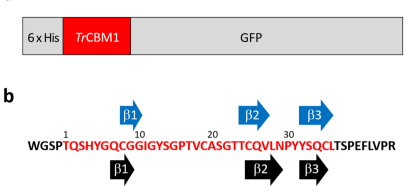


Figure 2. Schematics of His tag-*Tr*CBM1-GFP construct and the primary sequence of *Tr*CBM1 used in this study. (a) *Tr*CBM1 was expressed as a fusion protein with His tag and GFP at the N- and C-termini, respectively. (b) Amino acid sequences of *Tr*CBM1 (in red) and the residual regions at the N- and C-termini from the enterokinase and thrombin cleavage, respectively (in black). The secondary structures of *Tr*CBM1 predicted by TALOS+ software in this study (blue arrows) and that proposed by Kraulis *et al.*¹⁹ (black arrows) are also indicated.

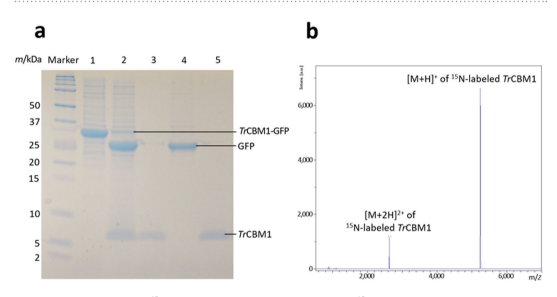


Figure 3. Purity analyses of ¹⁵N-labeled *Tr*CBM1. (**a**) SDS-PAGE of the ¹⁵N-labeled target proteins obtained in each purification step. Lanes 1 and 2: the protein fractions before and after cleavage of GFP using thrombin. Lanes 3 and 4: the protein fractions that passed through and were trapped in a benzamidine column, respectively. Lane 5: the *Tr*CBM1 obtained after the final cation exchange chromatography. (**b**) MALDI-TOF-MS spectrum of the purified ¹⁵N-labeled *Tr*CBM1. The corresponding full-length gel is shown in Supplementary Fig. S4.

folded and unfolded proteins (Fig. 4). The signals of the folded proteins appeared in the ¹H-chemical shift range

folded and unfolded proteins (Fig. 4). The signals of the folded proteins appeared in the ¹H-chemical shift range of 6.0–10.0 ppm. The signals of disordered proteins were observed only in the ¹H-chemical shift range of 8.0–8.5 ppm^{16,17}. We conclude that this sample contained folded as well as either partially or fully disordered forms, although the theoretical molecular mass for ¹⁵N-labeled *Tr*CBM1 was exhibited in MALDI-TOF-MS. ¹⁵N-labeled *Tr*CBM1 prepared using fresh competent cell gave merely the correctly folded protein signals. The correctly folded ¹⁵N-labeled, ¹³C/¹⁵N-labeled, and unlabeled *Tr*CBM1 with single molecular weight were used in this study.

Spectral assignments of ¹³**C**/¹⁵**N-labeled** *Tr***CBM1.** Spectral assignments of ¹³**C**/¹⁵**N-labeled** *Tr*CBM1 were achieved using a standard sequential assignment procedure. The ¹H-¹⁵N HSQC spectrum of ¹⁵N-labeled *Tr*CBM1 is shown in Fig. 5a with signal assignments. Backbone assignments of *Tr*CBM1 were 89% accomplished with the exception of eight residues. Their signals were not observed because of line broadening that is mainly related to their locations in flexible loop regions. The chemical shifts of backbone atoms (¹H^N, ¹⁵N, ¹³C^{α}, ¹³C^{β}, and ¹³C') of *Tr*CBM1 are listed in Supplementary Table S1. The amino acid residues that are in the secondary structures were predicted using TALOS + software based on Table S1¹⁸, (Fig. 2-b). As a result, ¹⁵N-labeled *Tr*CBM1 was predicted to have three β -strands: β 1 (C8 to G10), β 2 (T24 to V27), β 3 (Y32 to L36). Previously, Kraulis *et al.*

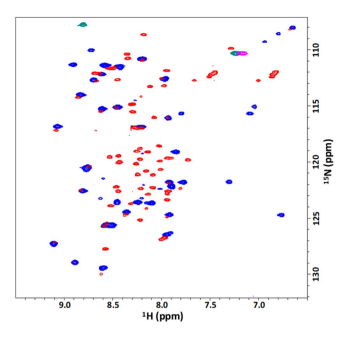


Figure 4. Overlay of 2D ¹H-¹⁵N SOFAST-HMQC spectra. Comparison of 100 μ M correctly folded *Tr*CBM1 (blue) and 20 μ M unfolded *Tr*CBM1 (red) which were expressed using older competent cell stocked for more than five months.

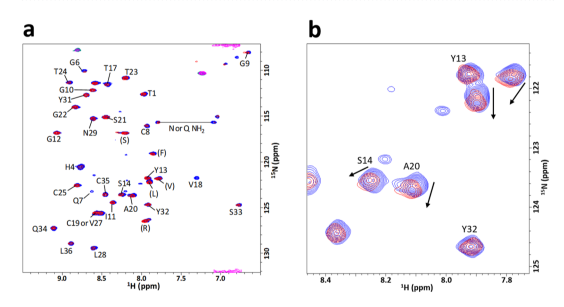


Figure 5. $2D \,{}^{1}H \,{}^{15}N$ SOFAST-HMQC spectra of NMR titration experiments using ${}^{15}N$ -labeled *Tr*CBM1 and C-MWL. (a) Superposition of $2D \,{}^{1}H \,{}^{15}N$ SOFAST-HMQC spectra of $100 \,\mu$ M *Tr*CBM1 in the presence (red) and absence (blue) of C-MWL (2695 μ M). The main chain resonances are labeled by corresponding residue number and amino acid type. The amino acid types in parentheses correspond to the amino acid residues in the residual regions from protease cleavage. (b) The close-up view of the region exhibited fairly large perturbations.

determined the three-dimensional solution structure of the unlabeled *Tr*CBM1 peptide (36 amino acid residues) that was chemically synthesized¹⁹. According to their report, *Tr*CBM1 has an anti-parallel β -sheet that comprised three β -strands: $\beta 1$ (Q7 to G9), $\beta 2$ (C25 to N29), and $\beta 3$ (Y32 to C35). The structures of *Tr*CBM1s prepared herein and by Kraulis *et al.*, therefore, are consistent. Accordingly, we used the solution structure of *Tr*CBM1 determined by Kraulis *et al.* to visualize the results of our NMR titration analyses.

Analysis of interaction sites of TrCBM1 with MWLs and cellohexaose. The interactions of *Tr*CBM1 with lignin and cellohexaose were comparatively analyzed by NMR titration experiments using ¹H-¹⁵N SOFAST-HMQC. We used highly purified milled wood lignins (MWLs) from a softwood, Japanese cedar (*Cryptomeria japonica*) (designated as C-MWL), and a hardwood, *Eucalyptus globulus* (E-MWL). Cellohexaose

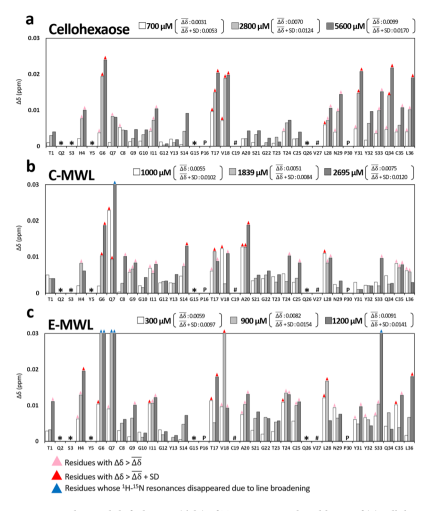


Figure 6. Chemical shift changes ($\Delta \delta s$) of *Tr*CBM1 upon the addition of (**a**) cellohexaose, (**b**) C-MWL, and (**c**) E-MWL. The $\Delta \delta s$ were calculated using Eq. (1) in three different concentrations of titrants. The residues that are not assigned are indicated in "*". Those residues that are overlapped are indicated in "#". Prolines are indicated in "P".

is an oligosaccharide having the minimum chain length recognizable by $TrCBM1^{20}$, which was used as a model compound of cellulose in the CSP analysis. The addition of excess amounts of MWLs resulted in the disappearance of NMR signals of TrCBM1, which hindered the assignments. Hence, the maximum concentrations of MWLs used for analyses were 2695 and 1200 μ M for C-MWL and E-MWL, respectively. The signals of TrCBM1 were still found at these concentrations. Incremental titration to the TrCBM1 solution was carried out using different concentrations of C-MWL (1000, 1839, and 2695 μ M), E-MWL (300, 900, and 1200 μ M), and cellohexaose (700, 2800, and 5600 μ M).

¹H-¹⁵N SOFAST-HMQC spectra of 100 μM ¹⁵N-labeled *Tr*CBM1 alone and in the presence of 2695 μM C-MWL were superimposed and are shown in Fig. 5. Upon incremental addition of the titrants to the solution of ¹⁵N-labeled *Tr*CBM1, several signals clearly exhibited perturbation with the reduction in signal intensity. Further perturbation of the signals was caused by increasing amounts of the titrant. Chemical shift change ($\Delta\delta$) calculated by the formula (1) is summarized in Fig. 6. The incremental addition of cellohexaose continuously increased $\Delta\delta$. ¹H-¹⁵N SOFAST-HMQC signals of **G6**, *T17*, *V18*, **Y31**, **Q34**, and **L36** exhibited large perturbation without line broadening. In addition, ¹H-¹⁵N SOFAST-HMQC signals of **G6**, **Q7**, S14, *T17*, *V18*, A20, and **L28** perturbed greatly upon the addition of C-MWL, while those of **H4**, **G6**, **I11**, *T17*, *V18*, *T24*, **L28**, C35, and **L36** perturbed when E-MWL was added. ¹H-¹⁵N SOFAST-HMQC signals of **G6** and S33 were line broadened in the presence of 900 and 1200 μM E-MWL, respectively, whereas the ¹H-¹⁵N SOFAST-HMQC signals were not line broadened in the presence of TrCBM1. When C-MWL and E-MWL were added with concentrations higher than 2695 and 700 μM, respectively, ¹H-¹⁵N SOFAST-HMQC signals of **Q7** resulted in line broadening. **Q7**, thus, was involved in direct or indirect interactions with MWLs.

The interaction sites of TrCBM1 revealed by NMR titration experiments were mapped on the solution structure of TrCBM1 determined by homonuclear NMR experiments (Fig. 7)¹⁹. As shown in Fig. 1, TrCBM1 has two major subsites, i.e., the flat plane surface and cleft. Triplet tyrosine (**Y5**, **Y31**, and **Y32**) of TrCBM1 is located on its flat plane surface, which plays a major role in the binding with cellulose. The triplet tyrosine is expected

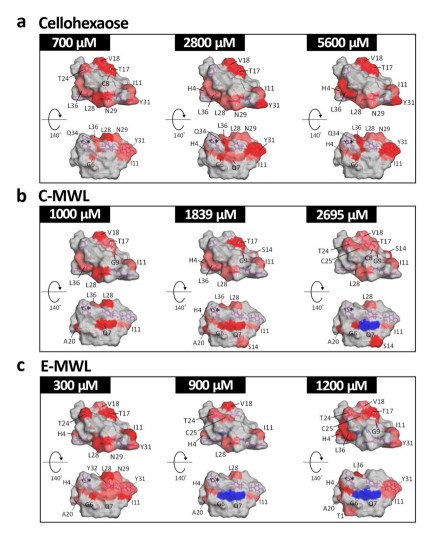


Figure 7. Mapping of cellohexaose and lignin binding sites identified by CSP on *Tr*CBM1. Binding sites of (a) cellohexaose, (b) C-MWL, and (c) E-MWL are shown on the *Tr*CBM1 surface. The residues that exhibited large $\Delta \delta s$ are color coded as described in Fig. 6. Two representative views, lateral and bottom faces of *Tr*CBM1, are shown with three concentrations of each titrant. Triplet tyrosine (**Y5**, **Y31**, and **Y32**) and cleft (*T17*, *V18*, and *T24*) are shown using purple stick. The residue **Y5** was not assigned.

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to be a main binding site with lignin due to its hydrophobic nature. However, the perturbations of the ¹H-¹⁵N SOFAST-HMQC signals of **Y31** and **Y32** became small, while the assignment of **Y5** was not accomplished. The small $\Delta\delta$ of the triplet tyrosine in the ¹H-¹⁵N SOFAST-HMQC spectra is due to the distant location between the aromatic ring in the side chain of tyrosine and ¹H-¹⁵N of the main chain, because specific detection of spin coupling of ¹H-¹⁵N in peptide bonds was monitored. Although the direct evidence of lignin binding via aromatic ring was not obtained, amino acid residues of the flat plane surface (**H4, G6, Q7, I11, L28, N29, Q34**, and **L36**) exhibited large $\Delta\delta$ upon the addition of MWLs and cellohexaose (Fig. 7). Thus, *Tr*CBM1 interacted with both MWLs and cellohexaose on the flat plane surface. We also found that **G6** and **Q7** were line broadened upon the addition of MWLs strongly bound to *Tr*CBM1 through the flat plane surface. The cleft composed of *T17, V18*, and *T24* also interacted with MWLs and cellohexaose. By extensive titration experiments, larger $\Delta\delta$ were consistently observed for *T17* and *V18* than *T24*.

Binding affinity of *Tr***CBM1 toward cellulose and lignin.** Adsorption experiment using Langmuir adsorption model was carried out using MWLs and Avicel. The latter is a commercially available cellulose rich in crystalline regions. A mixture of *Tr*CBM1 with 1%(w/v) of either MWLs or Avicel was incubated at 50 °C for 1 h. The amount of adsorbed *Tr*CBM1 was calculated by subtracting nonadsorbed *Tr*CBM1 from initial loading. The adsorption parameters that were calculated by the formula (2) are summarized in Table 1. The values of *Tr*CBM1 adsorption by Avicel are similar to the previously obtained values using synthesized *Tr*CBM1 analogs^{21,22}. Among these titrants, Langmuir affinity constant against *Tr*CBM1 was in the order of E-MWL>C-MWL>Avicel. Therefore, *Tr*CBM1 was found to possess higher affinity toward MWLs than Avicel. The highest Γ_{max} was given by Avicel, indicating that it has a wide surface area and MWLs aggregated in water solution.

	Langmuir affinity constant K _L (ml/mg)	Amount of adsorption at saturation $\Gamma_{max}\left(\mu g/mg\right)$
C-MWL	3.19	54.7
E-MWL	5.93	48.8
Avicel	2.65	63.7

Table 1. Adsorption parameters of *Tr*CBM1 for C-MWL, E-MWL, and Avicel determined by Langmuir adsorption isotherm.

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Discussion

T. reesei is one of the most important industrial microorganisms for producing cellulolytic enzymes due to its high productivity and high activity for the produced enzymes. Using the cellulolytic enzyme system of *T. reesei*, the production of CBHI (Cel7A) reaches up to 60% of the total enzymes⁹. CBHI plays a major role in the catalysis. Its molecular functions including *Tr*CBM1, therefore, have been studied extensively⁹. In cellulose hydrolysis, *Tr*CBM1 plays a crucial role in bringing enzyme close to the substrate, cellulose. However, due to the difficulties of expressing small proteins in *E. coli*, the molecular functions of *Tr*CBM1 have been studied using a chemically synthesized analog or as fusion proteins between *Tr*CBM1 and catalytic domain of *T. reesei* or other microbes, such as *Talaromyces emersonii* and *Melanocarpus albomyces*^{4,23,24}. The exceptions are the studies of Guo and Arslan. They studied the affinities of *Tr*CBM1 to various cellulose substrates²⁵ as well as the binding behavior of *Tr*CBM1 to lignocellulosic substrates using an atomic force microscope²⁶. These reports described the expression and purification of *Tr*CBM1. However, the molecular mass of the obtained *Tr*CBM1 and whether the obtained *Tr*CBM1 was correctly folded were not presented. These are crucial points, because we found that expression conditions greatly affected the correct folding of *Tr*CBM1. In this study, we focused on the experimental scheme that the binding behavior of *Tr*CBM1 at the molecular level was analyzed using a correctly folded single protein, *Tr*CBM1, as revealed by MALDI-TOF-MS and 2D ¹H-¹⁵N SOFAST-HMQC (Figs 3 and 4).

In general, point mutation has been extensively applied for protein-ligand interaction analysis. Indeed, this approach enabled us to identify the key amino acid residues involved in either ligand binding or catalytic activity. In some cases, the substitution of amino acid residues caused undesired changes in the conformation of proteins either partially or entirely. These unwanted structural changes may distort understanding of the actual roles of amino acid residues, especially when the protein of interest is small, such as in the case of *Tr*CBM1²¹. The use of stable isotope labeled proteins in combination with the adapted NMR titration experiments in this study is extraneous from such a disadvantage, giving direct information on the ligand and protein interaction at a molecular level in amino acid resolution.

Our NMR experiments indicated that two subsites of *Tr*CBM1 were the major interaction sites with cellohexaose and MWLs, i.e., the flat plane surface and cleft, (Fig. 7). Previous studies based on site-directed mutagenesis suggested that pyranose rings of cellulose and aromatic rings of lignin bound to *Tr*CBM1 through their triplet tyrosine, i.e., **Y5**, **Y31**, and **Y32**, that are exposed in the flat plane surface by hydrophobic interaction, CH- π , and π - π stacking, respectively, although substitution of the tyrosine residues affected alignment of neighboring amino acid residues^{23,24,27}. Our NMR study without the mutagenesis clearly indicated that the amino acid residues around triplet tyrosine (H4, G6, Q7, II1, L28, N29, Y31, Q34, and L36) constituting the flat plane surface exhibited large $\Delta\delta$. This CSP is explained by changes in shielding effects caused by the interactions of the tyrosine and neighboring amino acid residues with adsorbed cellohexaose or MWLs. Aliphatic OH groups in cellohexaose as well as both aliphatic and phenolic OH groups in MWLs are also the potential binding sites with *Tr*CBM1 through hydrogen bonding and electrostatic interaction²⁸. Our NMR study showed that large $\Delta\delta$ was observed in the 2D ¹H-¹⁵N SOFAST-HMQC signals of H4, Q7, and II1 that are located on the flat plane surface as well as *T17* in the cleft, suggesting that the main chain of H4 and II1 as well as the main chain and side chain of Q7 and *T17* participated in the binding with cellohexaose and MWLs via hydrogen bonding and electrostatic interaction.

Interestingly, a differential binding pattern was observed between cellohexaose and MWLs. N29 and Q34 showed large $\Delta\delta$ upon the addition of cellohexaose and thereby were identified as the interaction sites for cellohexaose. This result is consistent with a previous report by Mattinen *et al.*²⁰. It was reported that the substitution of N29 and Q34 to alanine reduced the affinity toward cellulose over lignin, indicating that N29 and Q34 interacted with cellulose more effectively than lignin²³. In our NMR study, the interactions of N29 and Q34 with MWLs were much less remarkable. The hydrophilic side chains of these amino acid residues, therefore, participated in the specific electrostatic interactions and hydrogen bonding with cellulose chains. G6 and Q7 in the flat plane surface of *Tr*CBM1 were line broadened upon the addition of MWLs. This phenomenon, however, was not observed for cellohexaose. When MWLs were added to *Tr*CBM1 solution, line broadening was mainly caused by both (1) on- and off-rates of the complex formation and (2) diverse binding states due to the heterogeneity of lignin, which is a good indicator of binding²⁹. Additionally, irregular increasing and decreasing of CSP (Fig. 6b,c) support the diverse binding states between lignin and *Tr*CBM1. Therefore, the line broadening of G6 and Q7 suggests that the flat plane surface of *Tr*CBM1 played a central role in the binding with lignin.

Cellohexaose bound to the flat plane surface and cleft with high specificity (Fig. 8a). In comparison, MWLs bound to various surface sites, including the flat plane surface and cleft, from much lower concentrations of titrants (Fig. 8b). The cumulative binding of MWLs on multiple exposed sites increased the overall binding affinity to the lignin although the observed CSPs at each site are small (Table 1).

Recently, we found that lignin-binding peptides that can recognize lignin specifically changed their conformation upon the addition of softwood and hardwood lignins to adopt their molecular shapes along with the surface of lignins³⁰. Differences in the absorptivity toward softwood and hardwood lignins were also observed for

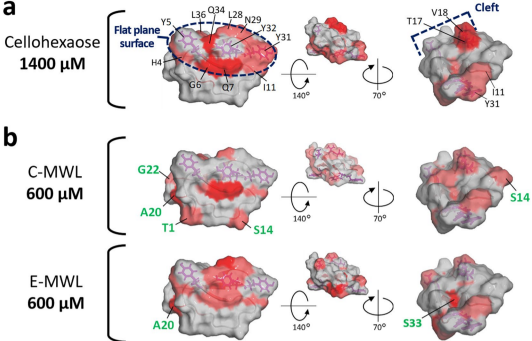


Figure 8. Comparison of interaction property between cellohexaose and MWLs. (a) Cellohexaose specifically bound to the flat plane surface and cleft. (b) Both MWLs bound to multiple binding sites, some of which are included in the flat plane surface and cleft even in low concentration of titrant. These non-specific binding sites are labeled by green characters.

TrCBM1. In addition to the flat plane surface and cleft, C-MWL interacted with the surface of TrCBM1 comprising amino acid residues of T1, S14, A20, and S22. E-MWL interacted with A20 and S33, which are also outside of the flat plane surface and cleft (Fig. 8). Therefore, we conclude that TrCBM1 recognized structural differences of softwood and hardwood lignins having similar weight-average molecular weights (C-MWL: 6254, E-MWL: 5776).

The binding of cellulase to lignin is affected by the structures of exposed surfaces of residual lignin, which results from structural differences of original biomass and pretreatment methods^{31,32}. Guo et al. compared lignins from six different plants species and concluded that low S/G ratio induced high adsorption capacity, which is consistent with our results (Table 1)³³. Moreover, high hydrophobicity, phenolic OH groups, and condensed structure of lignin increased adsorption capacity of cellulase, whereas aliphatic OH groups decreased adsorptivity^{28,34,35}. Our NMR titration experiments indicated that TrCBM1 bound to lignins through various outer surfaces of the protein, including the flat plane surface and cleft. We found the differences of binding sites between softwood and hardwood lignins. Structural differences caused by pretreatments should also affect the binding behaviors. Thus far, the involvement of triplet tyrosine in lignin binding has been suggested by a combination of point-mutation and adsorption experiments^{23,24,27}. These studies suggest the participation of the triplet in lignin binding; however, the role of other protein surfaces in the lignin binding cannot be analyzed and the point mutation may cause conformational changes of the protein. Our CSP study enables comprehensive analysis of interaction sites between the proposed structure of a protein and lignin without crucial conformational change.

Understanding of the flexible molecular recognition mechanism of TrCBM1 bound to polysaccharides and lignins from pretreated biomass softwood and hardwood could contribute to the molecular design of cellulolytic enzymes having controlled affinity to lignin and polysaccharides. The molecular design is indispensable for enzymatic saccharification with the minimum enzyme dosage.

Conclusion

Nonproductive binding of cellulolytic enzymes to lignin has been a serious issue for enzymatic saccharification of lignocellulosics. Understanding of the adsorption mechanism at the molecular level, however, is still limited. In the present study, we analyzed the interaction sites of correctly folded ¹⁵N-labeled TrCBM1 with MWLs and cellohexaose through NMR titration experiments. TrCBM1 bound to cellohexaose through the flat plane surface comprising triplet tyrosine as well as cleft with high site specificity. In high contrast, the interaction sites of TrCBM1 with MWLs were spread on the protein surface including the flat plane surface and cleft. Line broadening of G6 and Q7 suggests that the flat plane surface of TrCBM1 strongly interacted with MWLs, while hydrophilic amino acid residues, N29 and Q34, interacted with cellohexaose preferentially. The NMR approach using stable isotope labeling could lead to the development of a fundamental theory to design hyper enzymes that preferentially bind to polysaccharides without inactivation by coexisting lignin.

Materials and Methods

Materials. *E. coli* BL21 (DE3) was purchased from Merck (Darmstadt, Germany). The pRSET-EmGFP vector was obtained from Thermo Fisher Scientific (Waltham, MA, USA). Enterokinase and thrombin were purchased from New England Bio Labs (Ipswich, MA, USA) and GE Healthcare (Chicago, IL, USA), respectively. Cellohexaose was obtained from Toronto Research Chemicals (Toronto, Canada). Other laboratory reagents were purchased from Sigma-Aldrich (St. Louis, MO, USA), Wako Pure Chemical Ltd. (Osaka, Japan), nacalai tesque (Kyoto, Japan), and Cambridge Isotope Laboratories (Tewksbury, MA, USA).

Preparation of MWLs. Japanese cedar (*Cryptomeria japonica*) and *Eucalyptus globulus* woods were used for the preparation of C-MWL and E-MWL, respectively. The wood meal was extracted using a toluene and ethanol (2:1, v/v) mixture via a Soxhlet extractor at reflux temperature for 10 h. The extracted wood meal was dried at 105 °C for 12 h and finely divided in a vibratory ball mill having constant cooling water under a nitrogen atmosphere for 48 h. The milled wood was extracted using 96% aq. dioxane at room temperature for 24 h. The extract was allowed for solvent evaporation and then freeze dried. The crude MWL was dissolved in 90% aq. acetic acid and then precipitated from distilled water. The precipitates were washed using distilled water and dissolved in a 1,2-dichloroethane and ethanol (2:1, v/v) mixture before they were added to diethyl ether. The precipitates were washed using petroleum ether and allowed for solvent evaporation to give MWL fractions. Molecular weight of MWLs was determined by gel permeation chromatography on three TSK gel supermultipore HZ-M columns (Tosho, Tokyo, Japan) using a Shimadzu instrument equipped with an LC-20AD pump, an SPD M20A diode array detector (Kyoto, Japan). Tetrahydrofuran was used as the eluent at a flow rate of 0.35 ml/min at 40 °C.

Plasmid construction. The gene of *Tr*CBM1 in Cel7A from *T. reesei* and thrombin recognition site was inserted into pRSET-EmGFP vector. The vector map of His tag-*Tr*CBM1-GFP expression plasmid is shown in supplementary (Fig. S1).

Expression and purification of ¹⁵**N-labeled** *Tr***CBM1.** *E. coli* BL21 (DE3) was transformed by heat shock with His tag-*Tr*CBM1-GFP expression plasmid. The transformant was inoculated to a ¹⁵N-labeling M9 medium (10 ml), containing ¹⁵N-NH₄Cl as the sole nitrogen source and 100 µg/ml ampicillin, before it was precultured at 37 °C for 18 h with shaking at 200 rpm. The culture was used to inoculate ¹⁵N-labeling M9 medium (750 ml) and further incubated at 37 °C with shaking at 200 rpm, until OD₆₀₀ reached 1.2. Protein expression was induced using 1 mM isopropyl β-1-thiogalactopyranoside (IPTG) at 37 °C for 5 h with shaking at 200 rpm as well.

After centrifugation at 7000 rpm for 15 min (HITACHI, himac CR21GII, R13A rotor), the cells were resuspended by a buffer containing 50 mM sodium phosphate (pH 7.5) and 500 mM NaCl to make a 10% weight per volume solution. This suspension was sonicated, centrifugated at 12000 g for 60 min, and filtered through a 0.45 μ m filter. Then, cOmplete His Tag Purification Resin (Roche, Basel, Switzerland) equilibrated with the same buffer was mixed into the solution. The mixture was gently shaken for 15 min on ice and loaded into an open column (Bio Rad, CA, USA). The target protein containing His tag-*Tr*CBM1-GFP was eluted with the same buffer but containing 250 mM imidazole. The fractions containing His tag-*Tr*CBM1-GFP were collected and diluted by tenfold using a 20 mM Tris-HCl buffer (pH 8.0), before they were applied to a 5 ml Hi Trap Q FF column (GE Healthcare, IL, USA) equilibrated with the same buffer. The protein was then eluted from the column using a 0–500 mM NaCl gradient in 20 mM Tris-HCl buffer (pH 8.0) on AKTA prime (GE Healthcare, IL, USA).

Using 10-kDa molecular weight cut off (MWCO) Vivaspin turbo ultrafiltration devices (Sartorius, Göttingen, Germany), the target fraction including His tag-*Tr*CBM1-GFP was concentrated to 1.0 mg/ml in a buffer containing 20 mM Tris-HCl and 50 mM NaCl (pH.8.0). The obtained His tag-*Tr*CBM1-GFP solution was treated by enterokinase (33 U/mg of protein) at 23 °C for 20 h without shaking to cleave the His tag. To remove the cleaved His tag, the reaction mixture was diluted by fivefold using 50 mM sodium phosphate buffer (pH 7.5) containing 500 mM NaCl and incubated with cOmplete His Tag Purification Resin for 15 min on ice. The obtained *Tr*CBM1-GFP solution was then centrifuged at 500 g for 1 min, before the supernatant was filtered through a 0.45 μ m filter. Subsequently, the solution was concentrated using 10-kDa MWCO ultrafiltration devices as well to obtain 2.6 mg/ml *Tr*CBM1-GFP dissolved in PBS (-) buffer. The *Tr*CBM1 was separated from GFP by treating with thrombin (50 U/mg of protein) at 22 °C for 20 h without shaking.

The reaction mixture was applied on a 1 ml Hi Trap Benzamidine FF column (GE Healthcare, IL, USA) to remove GFP and thrombin, which were bound to the column. The run-through fraction containing *Tr*CBM1 was collected and buffer exchanged into 20 mM citric acid buffer (pH 3.0) using 3-kDa MWCO ultrafiltration device. The obtained solution was then applied to a 1 ml Hi Trap SP HP column (GE Healthcare, IL, USA) on AKTA prime, before *Tr*CBM1 was eluted using a 0–1 M NaCl gradient in a 20 mM citric acid buffer (pH 3.0). Finally, the obtained *Tr*CBM1 was buffer exchanged into a 100 mM citric acid buffer (pH 5.0) using 3-kDa MWCO ultrafiltration devices.

We used LB medium, a M9 medium containing only ${}^{15}NH_4Cl$ (99%, Cambridge Isotope Laboratories), and a M9 medium containing [U- ${}^{13}C$] glucose (99%, Cambridge Isotope Laboratories)/ ${}^{15}NH_4Cl$, respectively, to obtain each of the nonlabeled, ${}^{15}N$ -labeled, and ${}^{13}C/{}^{15}N$ -labeled *Tr*CBM1s.

The protein concentration was determined by reading the absorbance at 280 nm and using extinction coefficient (11960 M^{-1} cm⁻¹), proposed by Pace *et al.*³⁶. Molecular mass and purity of the purified ¹⁵N-labeled *Tr*CBM1 were analyzed by SDS-PAGE and MALDI-TOF-MS using Autoflex III (Bruker Daltonics, MA, USA), respectively. The structures of the purified *Tr*CBM1s were evaluated using 2D ¹H-¹⁵N SOFAST-HMQC¹⁵.

NMR spectroscopy and spectral assignment of *Tr***CBM1.** For NMR experiments, we used the ${}^{13}C/{}^{15}N$ -labeled *Tr*CBM1 of 150 μ M dissolved in 45 mM sodium acetate buffer (pH 5.0), containing 10% D₂O and 20 μ M 2,2-dimethyl-2-silapentane-5-sulfonic acid (DSS). All NMR spectra were recorded at 298 K on a Bruker

Avance III 600 spectrometer equipped with a cryogenic probe and Z-gradient (Bruker BioSpin, MA, USA). NMR spectra were processed by NMRPipe/NMRDraw³⁷. Spectral analysis was performed by MagRO^{38,39} working with NMRView⁴⁰, following the methods described previously⁴¹. The assignments of the backbone ¹HN, ¹⁵N, ¹³C^{α}, ¹³C³, and ¹³C' resonances of *Tr*CBM1 were made using ¹H-¹⁵N HSQC, HNCO, HN(CA)CO, HNCACB, CBCA(CO) NH. The secondary structural elements of *Tr*CBM1 were identified using the TALOS+ software¹⁸.

NMR chemical shift perturbation analysis. ¹⁵N-labeled *Tr*CBM1 of 100 μ M was dissolved in a 100 mM citric acid buffer (pH 5.0), 90% H₂O/10% D₂O, and 20 μ M DSS. Three different titrants, C-MWL, E-WML, and cellohexaose, were individually titrated into the ¹⁵N-labeled *Tr*CBM1 solution with incremental concentrations, i.e., C-MWL (600, 1000, 1839, 2695 μ M), E-MWL (300, 600, 900, 1200 μ M), and cellohexaose (700, 1400, 2800, 5600 μ M). To identify the amino acid residues of ¹⁵N-labeled *Tr*CBM1, which were involved in binding, chemical shift change $\Delta\delta$ (ppm) for each amino acid was calculated using the following equation⁴²

$$\Delta\delta(\text{ppm}) = \sqrt{(0.17\Delta\delta\text{N})^2 + (\Delta\delta\text{NH})^2} \tag{1}$$

where $\Delta\delta N$ and $\Delta\delta NH$ are chemical shift changes in ¹⁵N-axis and ¹H-axis, respectively. Because MWL titrants were dissolved in d_6 -DMSO, a control titrant containing d_6 -DMSO without MWL was also prepared. The chemical shift changes obtained for MWL titrants were subtracted by those obtained for control titrants to obtain the actual $\Delta\delta s$ of *Tr*CBM1 residues for MWLs. The amino acid residues that showed $\Delta\delta$ values larger than the average value ($\overline{\Delta\delta}$) were mapped on the proposed *Tr*CBM1 solution structure, which were color coded in pink. The amino acid residues that showed $\Delta\delta$ values larger than the sum of the $\overline{\Delta\delta}$ and the standard deviation of $\Delta\delta$ were mapped in red. The amino acid residues whose signals disappeared upon the addition of titrants were mapped in blue. Three-dimensional solution structure of the *Tr*CBM1 was shown using molecular graphics software, PyMOL (Schrödinger, NY, USA).

Adsorption experiment. Adsorption experiment was employed to evaluate the affinities of *Tr*CBM1 with each of the MWLs and Avicel using Langmuir adsorption isotherm. Sample solutions contained *Tr*CBM1 and one of 1% (w/v) C-MWL, E-MWL, and Avicel in 50 mM citric acid buffer (pH 5.0). The concentration of *Tr*CBM1 was varied as 40, 80, 160, 320, 640, 1280, and 2000 μ g/ml with the total volume of 50 μ l in a 1.5 ml micro tube. The sample solutions were incubated at 50 °C and were shaken at 1000 rpm for 60 min using thermomixer comfort (Eppendorf, Hamburg, Germany). Subsequently, the sample solutions were centrifuged at 12000 g for 10 min. The free *Tr*CBM1 content in the supernatant was quantified based on the Bradford method using Bio-Rad Protein Assay (Bio-rad, CA, USA). The amount of adsorbed *Tr*CBM1 was calculated by subtracting the amount of free *Tr*CBM1 from that of the initially loaded *Tr*CBM1. C-MWL, E-MWL, or Avicel (1%, w/v) without *Tr*CBM1 were used as a blank. Experiments were carried out at least two times and the results were expressed as average values. Langmuir affinity constant was calculated by the following formula.

$$\Gamma_{\rm C} = \Gamma_{\rm max} \frac{K_{\rm L}C}{1 + K_{\rm L}C} \tag{2}$$

where $\Gamma_{\rm C}$ is the amount of adsorbed *Tr*CBM1 and $\Gamma_{\rm max}$ is the amount of adsorbed *Tr*CBM1 at saturation to MWLs and Avicel. K_L is the Langmuir affinity constant to MWLs and Avicel. C is the concentration of free *Tr*CBM1 in the supernatant.

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Author Contributions

Designed the experiments: Y.T., T.N., M.K., T.W. Performed the experiments: Y.T., T.S. Analyzed and interpreted the NMR data: Y.T., K.K., T.N., M.K. Contributed protein expression and purification: S.O. Wrote the paper: Y.T., T.W. T.N., M.K.

Additional Information

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