Comparison of three seemingly similar lytic polysaccharide monooxygenases from \textit{Neurospora crassa} suggests different roles in plant biomass degradation

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Many fungi produce multiple lytic polysaccharide monooxygenases (LPMOs) with seemingly similar functions, but the biological reason for this multiplicity remains unknown. To address this question, here we carried out comparative structural and functional characterizations of three cellulose-active C4-oxidizing family AA9 LPMOs from the fungus \textit{Neurospora crassa}, NcLPMO9A (NCU02240), NcLPMO9C (NCU02916), and NcLPMO9D (NCU01050). We solved the three-dimensional structure of copper-bound NcLPMO9A at 1.6-Å resolution and found that NcLPMO9A and NcLPMO9C, containing a CBM1 carbohydrate-binding module, bind cellulose more strongly and were less susceptible to inactivation than NcLPMO9D, which lacks a CBM. All three LPMOs were active on tamarind xyloglucan and konjac glucomannan, generating similar products but clearly differing in activity levels. Importantly, in some cases, the addition of phosphoric acid–swollen cellulose (PASC) had a major effect on activity: NcLPMO9A was active on xyloglucan only in the presence of PASC, and PASC enhanced NcLPMO9D activity on glucomannan. Interestingly, the three enzymes also exhibited large differences in their interactions with enzymatic electron donors, which could reflect that they are optimized to act with different reducing partners. All three enzymes efficiently used \(\text{H}_2\text{O}_2\) as a cosubstrate, yielding product profiles identical to those obtained in \(\text{O}_2\)-driven reactions with PASC, xyloglucan, or glucomannan. Our results indicate that seemingly similar LPMOs act preferentially on different types of copolymeric substructures in the plant cell wall, possibly because these LPMOs are functionally adapted to distinct niches differing in the types of available reducants.

The filamentous fungus \textit{Neurospora crassa} is commonly found in nature where it grows on dead plant material, particularly grasses. Its cellulolytic potential has been known for decades (1–3). The \textit{N. crassa} genome encodes a similar number of glycoside hydrolase (GH) enzymes as \textit{Hypocrea jecorina}, the main industrial source of enzymes for biomass depolymerization (4). The \textit{N. crassa} genome also encodes 14 genes encoding lytic polysaccharide monooxygenases (LPMOs) belonging to CAZy family AA9 (5). Transcriptome studies done at a time when these LPMOs were erroneously thought to be glycoside hydrolases belonging to CAZy family GH61 showed transcripts for 13 of the AA9-encoding genes (6). LPMO-encoding genes tend to be abundant in the genomes of biomass-degrading fungi, indicating an important role in biomass degradation and raising questions as to the evolutionary drivers of this high multiplicity (7–11).

LPMOs are monocopper enzymes (12, 13) that were identified in 2010 (14) and that oxidatively cleave glycosidic bonds in polysaccharides. The discovery of LPMOs has been followed by extensive research related to their structure, function, mechanism, diversity, and industrial application (11, 15–20). LPMOs have become an important ingredient of commercial enzyme mixtures for industrial biomass conversion (21–24). Although LPMO action usually is considered in light of the enzymatic deconstruction of cellulose and chitin, some LPMOs act on other substrates, including xyloglucan and other (1,4)-linked \(\beta\)-glucans (25, 26), starch (27, 28), and xylan (29, 30). LPMO action requires reduction of the copper by an enzymatic or nonenzymatic electron donor (9, 31, 32). Subsequently, the enzymes use either molecular oxygen (14, 33) or hydrogen peroxide (34, 35) to generate an oxygen species at the copper site.

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The atomic coordinates and structure factors (code SF0H) have been deposited in the Protein Data Bank (http://wwpdb.org/).

This article contains Table S1 and Figs. S1–S10.

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that is capable of abstracting a hydrogen atom from the C1 or the C4 of the scissile glycosidic bond.

The biological reason for the multiplicity of LPMOs remains unresolved. For example, all of the nine biochemically characterized NcLPMOs (25, 36–38) are known to act on cellulose, seemingly varying only in terms of their oxidative regioselectivity: NcLPMO9F (NCU03328), NcLPMO9E (NCU08760), NCU02344, and NcLPMO9G (NCU00836) are strictly C1-oxidizing; NcLPMO9A (NCU02240), NcLPMO9C (NCU02916), and NcLPMO9D (NCU01050) are strictly C4-oxidizing; and NcLPMO9M (NCU07898) and NcLPMO9B (NCU07760) show mixed C1/C4 oxidation. For NcLPMO9C, activity on other β-glucans and soluble cellopectins has been demonstrated (25). Next to variation in oxidative regioselectivity, the LPMOs also vary in terms of the absence or presence of appended carbohydrate-binding modules (CBMs).

For more insight into potential differences between LPMOs, we have studied and compared the three strictly C4-oxidizing LPMOs of N. crassa, NcLPMO9A (NCU02240), NcLPMO9C (NCU02916) (39), and NcLPMO9D (NCU01050) (37). Of these, NcLPMO9A and NcLPMO9C contain a family 1 carbohydrate-binding module (CBM1; known for binding both crystalline and amorphous cellulose (40, 41)). To allow structural comparisons, we solved the crystal structure of the catalytic domain of NcLPMO9A. Functional analyses were inspired by the rich functional data set that was already available for NcLPMO9C (25, 39, 42, 43). We have assessed properties such as substrate binding and specificity, ability to recruit electrons from enzymatic and nonenzymatic electron donors, operational stability, the ability to generate H₂O₂ in the absence of substrate (36, 42), and potential differences between O₂-driven and H₂O₂-driven catalysis.

Results

Crystal structure and initial characterization of NcLPMO9A

The three-dimensional structure of NcLPMO9A with copper bound was solved to 1.6-Å resolution (Table S1) and revealed a typical LPMO structure, both with regard to the overall fold and the copper-containing active site (Fig. 1). Product analysis after incubation of NcLPMO9A with phosphoric acid–swollen cellulose (PASC) using chromatographic methods (44), MS/MS analysis (42), and identification of characteristic products obtained upon reduction (33) showed that NcLPMO9A exclusively oxidizes C4, confirming results of Vu et al. (38) (results not shown).

Overall structural comparison

Fig. 1 shows a sequence alignment and structural representations of the three C4-oxidizing LPMOs from N. crassa, NcLPMO9A, NcLPMO9C, and NcLPMO9D. The pairwise sequence identities of the catalytic domains range from 50 to 60%. Regions that are believed to be involved in substrate binding in family AA9 LPMOs are known as the L2, L3, LS, and LC loops (18, 45). Apart from the L3 loop, sequence conservation in these loops is similar to sequence conservation in the rest of the protein. The L3 loop shows higher sequence variability, and both the L3 and the LC loops are somewhat shorter in NcLPMO9A and NcLPMO9D compared with NcLPMO9C (Fig. 1A). Previous NMR studies on NcLPMO9C have shown that His-1, Ala-80, His-83, and His-155 (Fig. 1A, marked with * at positions 1, 78, 81, and 153, respectively) are involved in binding of both oligocellulose and polymeric xyloglucan (46), and three of these four residues are conserved among the three NcLPMOs. The residue at position 78 is predicted to take part in forming the −1 subsite (subsite prediction based on Courtaud et al. (46) and Frandsen et al. (47)) varies; it is an Ala (Ala-80) in NcLPMO9C and an Asp in NcLPMO9A (Asp-78) and NcLPMO9D (Asp-81). Superposition of the three NcLPMO structures with that of an AA9 LPMO from Lentinus similis (LsAA9A) in complex with cellohexaose (PDB code 5ACI (47)) (Fig. 2) shows that, apart from position 78, residues in subsites −1 to −3 are highly conserved among the three N. crassa enzymes, whereas more variation occurs in the +1 and +2 subsites as well as in the −4 subsite. These variations reside in the L2/L3 and LC regions (Fig. 1A) of the protein, respectively.

Courtaud et al. (46) showed that the L3 region is involved in the interaction with xyloglucan, and both the sequence alignment and the structures (Figs. 1 and 2) show considerable variation in this region. Although Asp-76 in the L3 loop of NcLPMO9C (position 74 in the alignment in Fig. 1A) is likely to interact with the C6-hydroxyl of the glucose unit in subsite +1, the corresponding residues in NcLPMO9A (Asp-74) and NcLPMO9D (Asp-77) seem not to be capable of such an interaction (Fig. 2). There is also considerable variation, i.e. at positions 67 and 68 in the L3 region and 25 and 26 in the L2 region. Position 68 in the L3 loop is of particular interest. The glutamate (Glu-65) in NcLPMO9C is likely to contribute to substrate binding at the +2 subsite (1.9–2.2 Å from the C1- and C2-hydroxyls), similarly to Asn-67 in LsAA9A (47). The side chains of the corresponding residues at this position in NcLPMO9A (Thr-68) and NcLPMO9D (Met-68) are less suitable to hydrogen bond with the substrate and seem to be located farther away (3.7–4.4 Å) from the glucose unit in the +2 subsite. Another clear difference concerns position 25 in the L2 region where both NcLPMO9A and NcLPMO9D have a tyrosine, whereas NcLPMO9C has an aspartate. A tyrosine could be engaged in stacking interactions with the substrate and may potentially extend the substrate-binding surface, perhaps forming a +3 subsite. The tyrosine at position 204 in the +3/+4 subsite is highly conserved in all AA9 LPMOs and well-known to be important for binding cello-oligomers (47) and xyloglucan (46). Interestingly, adjacent to this Tyr residue NcLPMO9D carries a Trp (Trp-207), which could contribute to substrate binding through aromatic stacking.

Binding to PASC

Fig. 3A shows that in the absence of ascorbic acid (AscA) binding of all three NcLPMOs reached equilibrium in 30 min after mixing with PASC. The strongest binding was observed for the two NcLPMOs containing a CBM1, NcLPMO9C and NcLPMO9A, with ~80% and ~50% binding, respectively. Binding of NcLPMO9D was much weaker, with only ~15% binding. The presence of AscA had an initial positive effect on binding observed for all three NcLPMOs, but during the next hour the bound fractions of all three LPMOs gradually decreased.
When equilibrium was reached, the fractions of bound protein were lower than those in the reactions without AscA. Analysis of the experiments shown in Fig. 3 by SDS-PAGE of nonbound protein showed similar trends (Fig. S1).

**Substrate specificity**

It has previously been shown that NcLPMO9A and -9D are active on PASC (38), whereas a broader substrate specificity has been demonstrated for NcLPMO9C, which acts on cellobiose (39), xyloglucan, glucomannan, and β-glucan (25, 42). To gain further information about substrate specificity of each of the three NcLPMOs, their activities were tested in the presence of AscA and single hemicelluloses but also on more complex substrates whereby PASC was premixed with different hemicelluloses. Product mixtures were analyzed by high-performance anion-exchange chromatography with pulsed amperometric detection (HPAEC-PAD) and MALDI-TOF MS.

When using PASC as a substrate, HPAEC-PAD analysis showed, as expected, only products oxidized at the C4 position for all three LPMOs (Fig. S2). Considerable amounts of native products were also detected, which is due in part to on-column conversion of C4-oxidized cellodextrins to native compound (48). Of note, native species emerge when LPMOs act close to chain ends of cellulose, which is more pronounced on amorphous substrates (44). In this case, the ratio of native-to-oxidized species is relatively high at low conversion levels as visible in Fig. S2.

Activity on tamarind xyloglucan (TXG) was detected for NcLPMO9C and NcLPMO9D but not for NcLPMO9A. The product profile generated by NcLPMO9C was as reported previously (Fig. 3B). When equilibrium was reached, the fractions of bound protein were lower than those in the reactions without AscA. Analysis of the experiments shown in Fig. 3 by SDS-PAGE of nonbound protein showed similar trends (Fig. S1).
Previously (25), NcLPMO9D yielded less intense peaks (Fig. S3A), indicating weaker activity of this LPMO on TXG or possibly reflecting that this LPMO is rapidly inactivated in reactions with TXG. MALDI-TOF MS spectra confirmed the TXG activity of NcLPMO9C and NcLPMO9D, showing characteristic clusters of products where each cluster contains products where the number of pentoses equals a multitude of 3 (Fig. 4A). According to the MS data, the products released by NcLPMO9C and NcLPMO9D were identical (Fig. 4B), and the clustered product profiles indicate that for both LPMOs the lytic reaction occurs at the nonsubstituted glucosyl unit.

Remarkably, when incubating NcLPMO9A with a mixture of TXG and PASC, this LPMO showed activity on TXG, yielding products similar to the other two LPMOs (Figs. 4, C and D, and S3B). Although quantitative interpretation of MS data requires caution, it is worth noting the differences between the LPMOs that are visible in Fig. 4D: whereas the product spectrum for NcLPMO9C almost exclusively shows xylloglucan-derived products, the product spectra of the other two show considerable amounts of PASC-derived products.

HPAEC-PAD chromatograms showed that all three LPMOs generated were active on konjac glucomannan (KGM), with NcLPMO9D being the least efficient (Fig. 5A). Analysis of the reaction products by MALDI-TOF MS confirmed oxidative activity on KGM, as oxidized hexose oligomers as well as their acetylated and double acetylated forms (characteristic for KGM) appeared in the spectra (Fig. S4). HPAEC-PAD chromatograms further showed that the activity on KGM was promoted by the presence of PASC, in particular for NcLPMO9D (compare Fig. 5, A and B). Activity on cellopentaose was confirmed only for NcLPMO9C as reported before (42), whereas only trace activities were detected for both NcLPMO9A and NcLPMO9D (Fig. S5). No activity for any LPMO was observed toward birchwood xylan and ivory nut mannan, either in the absence or in the presence of PASC (data not shown). In the absence

Figure 2. Amino acids potentially involved in substrate binding in NcLPMO9A (green), NcLPMO9C (magenta), and NcLPMO9D (blue). Substrate binding was predicted based on superposition of the crystal structures of NcLPMO9A (PDB code 5FOH; this study), NcLPMO9C (PDB code 4D7U (39)), and NcLPMO9D (PDB code 4EIR (37)) with LsAA9A in complex with cellohexaose (PDB code 5ACI (47)); the picture shows the side chains of the three NcLPMO9s. Amino acids are numbered based on NcLPMO9A (see Fig. 1A). Fully conserved residues have black labels, whereas not fully conserved residues have multiple labels that are colored according to the corresponding LPMO. The numbers within the sugar units (shown in light gray) indicate subsites. For clarity, the catalytic copper of only NcLPMO9A and only one of the conformations for the side chains of the residues Tyr-25 and Ala-77 in NcLPMO9D are shown.

Figure 3. Time-dependent binding of C4-oxidizing NcLPMOs to PASC in the absence (A) and presence (B) of 1 mM AscA. The percentage of unbound LPMO was determined by measuring the reduction in the concentration of LPMO in solution over time. The experiments were carried out using 2 mg·ml⁻¹ PASC and 5 μM LPMO in 50 mM Bis-Tris (pH 6.5) at 45 °C and 1000 rpm. Error bars represent S.D. of triplicate samples. Analysis of the same experiments by SDS-PAGE showed similar trends (Fig. S1).
of an electron donor, no activity was detected for any substrate.

**Variation of electron donors**

The dependence of the activity of the NcLPMOs on the nature of the reductant was tested in the reactions with PASC using AscA as a nonenzymatic electron donor and cellobiose dehydrogenase from *Myriococcus thermophilum* (MtCDH) or pyrroloquinoline quinone (PQQ)-dependent pyranose dehydrogenase from *Coprinopsis cinerea* (CcPDH) as enzymatic electron donors.

Fig. 6 shows clear dose-response effects upon varying the concentration of AscA. Of note, in these reactions one molecule of AscA is expected to be consumed per substrate oxidation, and the amount of substrate oxidations may be underestimated, likely by a factor of approximately 2 (49), because only solubilized products are detected. In the reactions with 0.3 mM AscA, the maximum amount of released oxidized product was similar for all three LPMOs and in the order of 100 H9262 M (Fig. 6A). In this case, the observed termination of product formation is likely due to depletion of the reductant. Higher concentrations of reductant (Fig. 6, B–D) gave faster initial rates and higher
total yields of oxidized products but also increased enzyme inactivation. The latter is suggested by the fact that product formation stopped at product concentrations far below the concentration of AscA (e.g. at 0.6 mM in the reaction with NcLPMO9A and 10 mM AscA). Control experiments in which the reactions with 3.3 mM AscA were supplied with all possible combinations of fresh AscA, PASC, and/or LPMO after 240 min (i.e. at the end points shown in Fig. 6C) showed that only additions containing fresh enzyme led to restoration of product formation (Fig. S6). This confirms that the cessation of product formation indeed is due to enzyme inactivation.

In terms of inactivation, the three LPMOs showed clear differences (Fig. 6): although NcLPMO9D became rapidly inactivated and yielded low product levels, NcLPMO9A and, even more so, NcLPMO9C were less sensitive to increased AscA levels, stayed active for a longer time, and reached higher product levels.

MtCDH is known to be able to oxidize oligocellulose and cellulose, whereas the PQQ-dependent CcPDH preferably oxidizes monosaccharides with a 1C4 chair conformation such as L-fucose and D-arabinose (50, 51). Therefore, the reactions with CcPDH were fueled by addition of 1 mM L-fucose. Due to the ability of MtCDH to oxidize products released upon LPMO action on PASC, in the reactions with this electron donor it was not possible to calculate the concentration of products oxidized by LPMOs themselves. We could only assess the amount of C1-oxidized sites, which again reflects the degree of substrate solubilization by the LPMO.

Product formation in reactions with these enzymatic donors (Fig. 7) was remarkably different from the results obtained with AscA (Fig. 6). In all cases, more or less linear progress curves were obtained, and the enzyme performing best with AscA, NcLPMO9C, was clearly the least active in the reactions with MtCDH or CcPDH (Fig. 7). NcLPMO9D performed very well, and the reaction with CcPDH yielded product levels higher than in any of the AscA reactions without showing signs of enzyme inactivation. A control experiment using the conditions of Fig. 6C showed that the activity of NcLPMO9D was not improved by addition of 0.5 μM BSA (data not shown), indicating that the good performance of this enzyme in the presence of enzymatic electron donors is not due to the mere presence of additional protein. NcLPMO9A performed well with both AscA and the enzymatic electron donors.

H2O2 production

Reduced LPMOs can carry out an oxidase reaction leading to the production of H2O2 (36, 42), which is a cosubstrate for LPMOs (35). It has been suggested that generation of H2O2 may be rate-limiting in most “standard” LPMO reactions (O2 and e.g. AscA) (11, 35), but this remains somewhat controversial (52). We tested the ability of the three NcLPMO9s to produce H2O2 in the presence of 50 μM AscA as reductant and absence of substrate. The measured initial H2O2 production rates were 1.13 ± 0.05, 0.60 ± 0.08, and 0.91 ± 0.02 min⁻¹ for NcLPMO9A, NcLPMO9C, and NcLPMO9D, respectively (Fig. S7). It is worth noting that these rates are in the same order as...
LPMO multiplicity relates to varying substrate preference

Figure 7. Time course for release of oxidized products during incubation of 1 μM NcLPMO with 2 mg ml⁻¹ PASC and 0.5 μM MtCDH (A) or 0.5 μM CcPDH (B). NcLPMO9A (black), NcLPMO9C (red), and NcLPMO9D (blue) were incubated with PASC and MtCDH or CcPDH in 50 mM Bis-Tris (pH 6.5) at 45 °C and 1000 rpm. In the reactions with MtCDH, the amount of released soluble products is expressed as the sum of the integrated peak areas (nC × min) for C1-oxidized (C1-ox) products only (in this case, C4-oxidized products cannot be detected as they are oxidized at C1 position by CDH). In the reactions with CcPDH, the released oxidized products for each sample were enzymatically converted to Glc4gemGlc using 1 μM TrCel7A. The concentrations of Glc4gemGlc in every sample were then determined by HPAEC-PAD using Glc4gemGlc standard prepared as described previously (22). Consequently, values on the y axis in A and B should not be compared; quantitative assessment is only valid for comparing the NcLPMOs within the same panel. Results from the reactions with CcPDH may be compared with the results obtained in the reactions with different AscA concentrations (Fig. 6). Error bars represent S.D. of triplicate samples.

Product release during the H₂O₂-driven degradation of PASC

Fig. 9 shows that under the conditions used above, i.e. gradual addition of H₂O₂, product formation from PASC was similar for all three LPMOs, suggesting that the amount of added H₂O₂ was limiting product formation under these reaction conditions. NcLPMO9A and -9C showed a linear progress curve throughout the reaction, whereas NcLPMO9D showed signs of inactivation after 3 h, in accordance with the apparent lower operational stability of this enzyme that was also observed in e.g. the dose-response study with AscA (Fig. 6). In the linear phase of the reactions, the ratio between the amount of oxidized products and the cumulative amount of added H₂O₂ was between 40 and 60%, which is compatible with close to stoichiometric incorporation of H₂O₂ in oxidized products because some 50% of oxidized sites likely remain in the insoluble material (49).

Discussion

N. crassa is a well-known model organism that also happens to be a proficient plant cell-wall degrader (3). It contains a large arsenal of different CAZymes involved in plant biomass degradation. Annotation of the genome shows that family AA9 LPMOs have the highest multiplicity of all CAZymes in N. crassa, with 14 genes predicted to encode such enzymes (6). For nine of these 14 LPMO9s, activity on cellulose has been demonstrated, and three of these (NcLPMO9A, -9C, and -9D) are strict C4 oxidizers. As structural comparisons and sequence alignments alone cannot provide a biological rationale for the multiplicity of lpmo9 genes, such as different roles during plant cell wall degradation, we carried out functional comparison of the three C4-oxidizing NcLPMOs.

Binding of LPMOs to their substrates may be enhanced by CBMs as in other fungal plant cell wall–degrading enzymes. LPMOs with reduced active-site copper and not bound to a substrate are prone to autocatalytic inactivation through non-productive reactions with O₂ or H₂O₂ (35), and it has indeed been shown that removal of the CBM not only reduces substrate affinity but also LPMO stability (49, 53). Accordingly, the present results show that stability of the three NcLPMO9s cor-
relates with substrate affinity. NcLPMO9D, without a CBM1 and showing the weakest substrate binding, became rapidly inactivated, yielded low product levels, and was more sensitive to even low AscA levels than NcLPMO9A and NcLPMO9C (both containing a CBM1). In contrast, NcLPMO9C, showing the strongest substrate binding, was the most stable of the three LPMOs. The high operational stability observed for NcLPMO9C under standard conditions (high AscA with O2) may also be due to a lower rate of H2O2 production (Fig. S7), which could reduce the chance of potentially destructive encounters of nonsubstrate-bound LPMO and H2O2. It is noteworthy that the considerable differences in substrate binding between the three LPMOs are not reflected in major differences in the initial catalytic rates, indicating the importance of other factors, such as enzyme mobility on the substrate surface, in enzyme catalysis.

Our results confirm previous observations that reduction of the active-site copper promotes substrate binding (43, 46). The gradual decrease in substrate binding over time, which is visible in Figs. 3 and S1, likely reflects oxidative damage of the LPMO catalytic site and not just reoxidation of the copper, as over time the bound fraction of LPMOs in the presence of AscA was less than in the reactions without AscA.

Importantly, in-depth functional characterization of the three C4-oxidizing NcLPMO9s revealed clear differences in substrate preferences. Only NcLPMO9C is active on shorter cello-oligosaccharides, whereas only NcLPMO9A and NcLPMO9D are active on TXG. Interestingly, NcLPMO9A did show activity on TXG in reactions that also contained cellulose. It is likely that hemicelluloses such as TXG associate with cellulose (54–56), creating junction zones between cellulose and hemicelluloses that enable the LPMO to act also on the now “stabilized” hemicellulose (see also Ref. 30). Further illustrating differences between the LPMOs, in reactions with PASC–TXG mixtures, PASC was still the preferred substrate for NcLPMO9A and NcLPMO9D, whereas NcLPMO9C primarily acted on TXG. Addition of cellulose also promoted activity of the LPMOs on KGM, in particular for NcLPMO9D, which was hardly active on KGM alone. It is worth noting that in

Figure 8. Reaction products generated by C4-oxidizing NcLPMOs from PASC or TXG in reactions with H2O2 as a cosubstrate. A and B, HPAEC-PAD analyses of products generated by 1 μM NcLPMO9A (black line), 1 μM NcLPMO9C (red line), or 1 μM NcLPMO9D (blue line) in reactions with 2 mg·ml⁻¹ PASC (A) or 2 mg·ml⁻¹ TXG (B) in 50 mM Bis-Tris (pH 6.5) at 45 °C and 1000 rpm with addition of ~45 μM H2O2 to the reactions every 15 min for 4 h. Prior to every addition of H2O2, ~12 μM AscA was added to ensure reduction of the LPMO. Control reactions (dashed lines) were done in the absence of H2O2, meaning that only ~12 μM AscA was added every 15 min for 4 h. The labeling of oxidized cello-oligosaccharides in A is based on previous work (42); the identity of the peaks in B is unknown due to lack of standards (but see Ref. 25 for some indications). C, MALDI-TOF MS spectra of oligosaccharides with DP6 and DP7 released from PASC by NcLPMO9A (black line), NcLPMO9C (red line), or NcLPMO9D (blue line). Peaks representing sodium adducts in the DP6 cluster are labeled, and this cluster shows the native product, (Glc)₆, and the oxidized hexamer in its keto form, Glc⁺⁺⁺(Glc)₅, and in its gem-diol form, Glc⁺⁺⁺(Glc)₅. D, MALDI-TOF MS spectra of products generated from TXG by NcLPMO9A (black), NcLPMO9C (red), and NcLPMO9D (blue). Clusters of products where the number of pentoses equals a multitude of 3 are indicated.
LPMO multiplicity relates to varying substrate preference

![Graph](image)

Figure 9. Time course for release of oxidized products during incubation of 2 mg·mL⁻¹ PASC with 1 μM C4-oxidizing NcLPMO in reactions with 

- **H₂O₂** (dashed lines)
- **NcLPMO9A** (black squares)
- **NcLPMO9C** (red squares)
- **NcLPMO9D** (blue squares)

The products were measured and that the nonsoluble material contains a significant fraction of the oxidized sites. The cumulative amount of 

$\text{H}_2\text{O}_2$ that was added to the reactions is indicated by green asterisks, and the concentration values are shown on the right $y$ axis.

Reactions with TXG the effect of adding cellulose was largest for NcLPMO9A, whereas it was largest for NcLPMO9D in reactions with KGM. Although further studies on true copolymeric substrates, such as intact plant cell walls, are needed, the present findings clearly show that the three seemingly similar NcLPMO9s have different substrate specificities that may relate to the complex structure of the plant cell wall.

Structural comparisons do not provide obvious explanations for the differences in substrate specificity. In accordance with previous studies on the binding of polymeric substrates (13, 46, 57), it may seem that a larger part of the putative substrate-binding surface is involved in determining substrate preferences. It is known that the L3 region is involved in binding xylolucan, and Figs. 1 and 2 show that the three LPMOs exhibit relatively much variation in this region, affecting e.g. subsite +2. It is worth noting that the LPMO without a CBM, NcLPMO9D, contains relatively many aromatic residues near the substrate-binding surface (Tyr at positions 25 and 67, subsite +2; Trp at position 205, subsite −4; see Fig. 1A) that may add to substrate affinity and could thus, perhaps, compensate for the lack of a CBM. However, mutagenesis studies on the proposed residues will need to be performed before conclusions on the role of these residues in substrate binding can be drawn.

It has previously been shown that LPMOs differ when it comes to their interactions with small-molecule reductants (58, 59). Here, we analyzed both a small-molecule reductant, AscA, and two enzymatic electron donors, MtCDH and CcPDH. The three LPMOs showed remarkable differences in operational stability and ability to recruit electrons from these varying electron donors. Slower, controlled activation of the LPMOs with an enzymatic donor increased the operational stability of NcLPMO9D, whereas AscA was the better electron donor for NcLPMO9C. Interestingly, in the reactions with enzymatic electron donors, the efficiency of the LPMOs was very different from that in reactions with AscA and inversely correlated to the LPMOs’ ability to bind substrate. For NcLPMO9C, it has been shown that the presence of substrate reduces the ability of CDH to reduce the LPMO (46), which could explain why tight substrate binders are less easily reduced by an enzymatic electron donor. In line with this, Várnai et al. (60) have shown that difference in the activation of different LPMOs by CcPDH can be abolished by addition of a small redox mediator, which can more easily access the active site of an LPMO bound to a substrate. The operational stability of the LPMO in the presence of various reductants is determined in part by the extent to which the LPMO is close to the substrate as it becomes reduced. The results with NcLPMO9D seem to support the notion that LPMO reduction close to the substrate is favorable: this LPMO, showing weak substrate binding and low operational stability when reduced by AscA, fully maintained operational stability when fueled by the CBM-containing and thus likely substrate-bound CcPDH.

The results presented above lend further support to the recent claims that $\text{H}_2\text{O}_2$ is a preferred, if not the only, cosubstrate of LPMOs (34, 35). All three NcLPMO9s could efficiently use added $\text{H}_2\text{O}_2$ as a cosubstrate, whereas their activity under standard conditions (1 mM AscA with no added $\text{H}_2\text{O}_2$) was correlated with their ability to produce $\text{H}_2\text{O}_2$. It has been proposed that $\text{H}_2\text{O}_2$-driven LPMO reactions are less specific than $\text{O}_2$-driven reactions because oxidative damage in the LPMO, claimed to only occur in the $\text{H}_2\text{O}_2$-driven reaction, would compromise enzyme specificity (52). Although reduction of specificity due to oxidative damage in the active site certainly is conceivable, we would argue that this possible process is not related to the nature of the cosubstrate used. Rather, the degree of oxidative damage, for example caused by a surplus of $\text{H}_2\text{O}_2$ reacting with nonsubstrate-bound reduced LPMOs, is due to an imbalance between the various reactants that leads to nonproductive and potentially damaging LPMO reactions. Here, we show that in well-controlled reactions $\text{O}_2$- and $\text{H}_2\text{O}_2$-fueled reactions yield essentially identical product profiles for all three LPMOs with PASC, xylolucan, and glucomannan. Fig. 9 shows that under controlled conditions one may achieve stoichiometric incorporation of $\text{H}_2\text{O}_2$, whereas the enzymes show good stability. These results show that the specificity of the LPMO is independent of the nature of the cosubstrate used to drive the reaction.

All in all, the present study shows that the C4-oxidizing cellulose-active LPMOs of *N. crassa* exhibit a variety of functional differences that may relate to their biological roles. Most importantly, although the true natural substrates of these LPMOs, such as, possibly, certain substructures of the plant cell wall, still need to be discovered, our results clearly show that the three enzymes have different substrate specificities. Another major difference concerns the interplay...
with varying reductants, which is interesting because the availability of both enzymatic and nonenzymatic electron donors will vary during plant cell wall degradation both in space and time (9, 61). Thus, it is conceivable that these three LPMOs have distinctive roles during degradation of the complex cell wall, which could explain the multiplicity of LPMOs in plant cell wall–degrading fungi.

**Experimental procedures**

**Chemicals**

Bis-Tris buffer salt was purchased from VWR (Radnor, PA); hydrogen peroxide (30% aqueous solution) was purchased from Merck; ascorbic acid, l-fucose, Avicel® PH-101, xylan from birch wood, horseradish peroxidase, and 10-acetyl-3,7-dihydroxyphenoxazine (AmplexRed) were purchased from Sigma-Aldrich. Cellopentaose (purity >95%), xylglucan from tamarind seeds (purity ~95%), konjac glucomannan (purity >98%), and ivory nut mannan (purity >98%) were purchased from Megazyme (Bray, Ireland). PASC was prepared from Avicel PH-101 as described before (62).

**Enzymes**

MtCDH and PQD-dependent CcPDH were expressed in *Pichia pastoris* and purified as reported previously (63, 64). The enzyme concentrations of MtCDH and CcPDH preparations were determined by measuring absorbance at 280 nm and using molar extinction coefficients of 159,000 and 146,500 M⁻¹·cm⁻¹, respectively, determined using the ExPASy ProtParam tool (65).

**Cloning, expression, and purification of LPMOs from *N. crassa***

The gene encoding *Nc*LPMO9A with its native signal peptide was codon-optimized for expression in *P. pastoris* and ordered from GenScript (Piscataway, NJ) in a pUC57 vector. A Kozak sequence (GAAACG) was inserted upstream of the start codon, and EcoRI and Acc65I restriction sites were introduced for cloning in the 5' and 3' ends, respectively. The pUC57 vector containing the codon-optimized sequence was digested by EcoRI and Acc65I, and the gene fragment was ligated into EcoRI/Acc65I-digested pPINK-GAP-TaCel5A (66), yielding pPINK-GAP_2240. The expression vector was transferred into *Escherichia coli* TOP10 (Invitrogen), and transformants were selected on brain–heart infusion agar with 200 µg·ml⁻¹ ampicillin. The pPINK-GAP_2240 plasmid was purified, linearized with AfIII, and transformed into electrocompetent cells of *P. pastoris* PichaPink™ Strain 4. Electrocompetent *P. pastoris* cells were prepared by following the manufacturer’s procedure. The electrocompetent cells were electroporated using a Bio-Rad Gene Pulser II electroporation unit (Bio-Rad Laboratories) at 1.8 kV, 25 microfarads, 200 ohms. The transformed cells were incubated in yeast extract–peptone–dextrose medium containing 1 M sorbitol overnight and spread on *Pichia* adenine-dropout selection plates, which were incubated at 30 °C.

For purification, the production strain was grown in 20 ml of buffered complex glycerol medium containing 1% (v/v) glycerol in a 100-ml shake flask at 29 °C and 200 rpm for 16 h. Subsequently, this preculture was used to inoculate 0.5 liter of buffered complex glycerol medium containing 1% (v/v) glycerol in a 2-liter shake flask followed by incubation at 29 °C and 200 rpm for 48 h. After 24 h, the culture was supplemented with 1% (v/v) glycerol. The cells were removed by centrifugation at 7,000 × g for 15 min at 4 °C. The supernatant was collected and dialyzed against 50 mM Bis-Tris buffer (pH 6.5) and concentrated to 100 ml using a VivaFlow 200 tangential cross-flow concentrator (molecular mass cutoff, 10 kDa; Sartorius Stedim Biotech GmbH, Germany). Ammonium sulfate was added to the concentrated supernatant to a final concentration of 1.42 M after which the solution was loaded onto a 5-ml HiTrap Phenyl FF column (GE Healthcare) equilibrated with 50 mM Bis-Tris buffer (pH 6.5) containing 1.42 M ammonium sulfate. Proteins bound to the column were eluted using a 25-ml linear gradient from 1.42 to 0 M ammonium sulfate in 50 mM Bis-Tris buffer (pH 6.5) using a flow rate of 1 ml·min⁻¹. Collected fractions were analyzed by SDS-PAGE, and the fractions containing *Nc*LPMO9A were pooled and subsequently concentrated down to 2 ml using Amicon Ultra centrifugal filters (molecular mass cutoff, 10 kDa; Merck Millipore). The protein solution was loaded onto a HiLoad 16/60 Superdex 75 size-exclusion column (GE Healthcare) in 50 mM Bis-Tris buffer (pH 6.5) containing 150 mM NaCl using a flow rate of 0.75 ml·min⁻¹. Fractions containing pure *Nc*LPMO9A were identified using SDS-PAGE and subsequently pooled and concentrated using Amicon Ultra centrifugal filters (molecular mass cutoff, 10 kDa).

*Nc*LPMO9C (UniProt accession number Q7SHI8) was cloned in *P. pastoris* (67) and expressed and purified following a previously published protocol (36). *Nc*LPMO9D (UniProt accession number Q1K8B6) was cloned as described earlier (23) using a protocol similar to that described above for *Nc*LPMO9A and a synthetic gene that encoded the protein with its native signal sequence. The enzyme was purified as described above for *Nc*LPMO9A.

The protein concentrations of *Nc*LPMO9A, *Nc*LPMO9C, and *Nc*LPMO9D preparations were determined by measuring absorbance at 280 nm and using molar extinction coefficients of 45,000, 46,900, and 35,660 M⁻¹·cm⁻¹, respectively, determined using the ExPASy ProtParam tool (65). The enzymes were saturated with Cu(II) by incubating with an excess of Cu(II)SO₄ (at an ~3:1 molar ratio of copper:enzyme) for 90 min in 50 mM Bis-Tris (pH 6.5) at room temperature as described previously (68). The Cu(II)-loaded sample of *Nc*LPMO9A was buffer-exchanged to 50 mM Bis-Tris buffer (pH 6.5) using Amicon Ultra centrifugal filters (molecular mass cutoff, 3 kDa). The resulting solution with purified protein was filtered through a 0.22-µm Millex®-GV filter (Merck Millipore) and stored at 4 °C.

**Crystallization and X-ray data collection**

Purified *Nc*LPMO9A was cleaved with papain (papain from papaya latex, Sigma-Aldrich) at an LPMO:papain ratio of 33:1 (w/w) to remove the CBM1 that is appended to the catalytic LPMO9 domain. The reaction was carried out in 100 mM sodium acetate buffer (pH 5.0) for 48 h at 37 °C, and its outcome was assessed by SDS-PAGE. For the isolation of the cleaved

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*LPMO multiplicity relates to varying substrate preference*
catalytic domain, size-exclusion chromatography was applied using a Superdex 75 16/600 column (GE Healthcare) equilibrated in 50 mM MES (pH 6.5) and 150 mM NaCl. The purified catalytic domain (NCU02240-N) was buffer-exchanged into 20 mM MES (pH 6.5) and concentrated to 15 mg/ml−1 for crystallization trials. A wide range of crystallization conditions were tested using a Mosquito crystallization robot (TTP Labtech, UK) and commercially available 96-well kits. Crystals of NCU02240-N grew at room temperature in the presence of 20% PEG3350 and 0.2 M Li2SO4 using the sitting-drop vapor-diffusion technique. Before being flash frozen in liquid N2, a single crystal was cryoprotected by transferring to mother liquor containing 35% PEG3350. X-ray diffraction data were collected to 1.6 Å resolution on beamline ID23-2 (European Synchrotron Radiation Facility, Grenoble, France) under cryogenic conditions. The wavelength of the X-ray beam was 0.8726 Å, and the oscillation range was 0.05°. The resulting data set was processed using XDS (69), and the structure was solved by molecular replacement using Phaser (70). The molecular replacement search model was produced by CHAINSAW (71) using NcLPMO09D (PDB code 4EIR) (37) as the template structure. Iterative rounds of model building and refinement of the structure were performed using Coot (72) and REFMAC (73) from the CCP4i program suite. Solvent molecules were added using Coot and checked manually. The quality of the final structure model was evaluated using MolProbity (74). The final structure model and the structure factors are deposited at the PDB under accession code 5FOH. Data collection and refinement statistics are summarized in Table S1. All structure figures were prepared using the UCSF Chimera package (75).

**Binding of LPMOs to PASC**

Binding studies with PASC were performed as described before (76) in the presence or absence of 1 mM AscA. The reaction mixtures contained 2 mg/ml−1 PASC or 2 mg/ml−1 PASC premixed with 1 mM AscA and 5 μM enzyme in 50 mM Bis-Tris buffer (pH 6.5) and were incubated at 45 °C with shaking at 1000 rpm in an Eppendorf ThermoMixer. At various time points (5, 15, 30, 60, and 120 min), samples were taken, and insoluble substrate and substrate-bound enzyme were removed by filtering using a 96-well filter plate (Merck Millipore) operated by a Millipore vacuum manifold. The concentration of the enzyme in the filtrate was determined by measuring A280 (Eppendorf BioPhotometer). Furthermore, the filtrates from each time point were mixed with an equal volume of SDS sample buffer and analyzed by SDS-PAGE by loading exactly 5 μl of such prepared samples per well.

**LPMO reactions with AscA**

Unless otherwise stated, reaction mixtures contained 2 mg/ml−1 substrate, 1 μM NcLPMO, and 1 mM AscA in 50 mM Bis-Tris buffer (pH 6.5). Reactions were performed in 2-ml Eppendorf tubes containing 100-μl total reaction volume and incubated at 45 °C with shaking at 1000 rpm in an Eppendorf ThermoMixer. Reactions were stopped by boiling for 10 min. Subsequently, separation of soluble and insoluble fractions was done by centrifugation at 11,000 × g for 10 min. As controls, reactions without added reductant were also analyzed.

**Detection of oxidized products**

Oxidized products were analyzed using HPAEC-PAD and by MALDI-TOF MS. HPAEC was performed on a Dionex ICS5000 system equipped with a CarboPac PA1 analytical column (2 × 250 mm) and a CarboPac PA1 guard column (2 × 50 mm) using a 50-min gradient (44) for cellulose and a 75-min gradient (25) for hemicellulosic substrates. Chromatograms were recorded with Chromeleon and analyzed using Origin 9.1 software (OriginLab, Northampton, MA).

MALDI-TOF MS was performed on an Ultraflex MALDI-TOF/TOF instrument (Bruker Daltonik GmbH, Bremen, Germany) equipped with a nitrogen 337 nm laser. The instrument was operated in positive-linear acquisition mode. The samples were sodium-saturated by mixing 5 μl of sample with 5 μl of 50 mM sodium acetate. 1 μl of the saturated sample was spotted on an MTP 384 ground steel MALDI target and mixed immediately with an equal volume of 20 mg/ml−1 2,5-dihydroxybenzoic acid in 30% acetonitrile and 0.1% (v/v) TFA. The data were analyzed using mMass software (77). Baseline correction and Gaussian smoothing (window size, 0.3 m/z) were applied to all spectra.

**Quantitative assessment of products released from PASC with varying electron donors**

Reaction mixtures (600 μl) with nonenzymatic electron donors contained 2 mg/ml−1 PASC, 1 μM NcLPMO, and 0.33, 1, 3.3, or 10 mM AscA in 50 mM Bis-Tris buffer (pH 6.5). Control reactions to check for substrate, enzyme, and reductant depletion were set up using the same conditions with 3.3 mM AscA in 100 μl; after 240 min, 100 μl of the reaction mixtures were supplemented with 50 μl of buffer (50 mM Bis-Tris (pH 6.5)) or with 50 μl of the same buffer containing various combinations of 2 mg/ml−1 substrate, 1 μM LPMO, and 3.3 mM AscA followed by further incubation for 120 min.

Reaction mixtures (600 μl) with an enzymatic electron donor contained 2 mg/ml−1 PASC, 1 μM NcLPMO, and 1 μM MtCDH (63) or 1 μM CcPDH (64). The reactions with CcPDH also contained 1 mM l-fucose, acting as a substrate for CcPDH.

All reactions were incubated at 45 °C with shaking at 1000 rpm in an Eppendorf ThermoMixer. At various time points (20, 40, 60, 120, and 240 min), 120-μl samples were taken and boiled for 10 min. Separation of soluble and insoluble fractions was achieved by centrifugation at 11,000 × g for 10 min. Prior to product quantification, soluble fractions (32 μl) were mixed with 31 μl of 150 mM sodium acetate buffer (pH 4.5) and 1 μl of TrCel7A (~1 μM) and incubated for 16 h at 37 °C. The treatment with TrCel7A converts oligomeric products to a mixture of glucose, cellobiose, and Glc4-gemGlc (C4-oxidized cellobiose) as the only oxidized product. Soluble fractions treated in this way were subsequently analyzed using HPAEC-PAD. The amount of released oxidized products was quantified using 4-hydroxy-β-d-xyl-β-d-Glc p (Glcc4-gemGlc), prepared as described previously (22), as a standard. In the reactions with MtCDH, the sum of integrated peak areas of C1-oxidized products (nC × min) was used as a proxy for LPMO activity.
LPMO reactions with H$_2$O$_2$

Unless otherwise stated, reaction mixtures (200 µl) contained 2 mg·ml$^{-1}$ substrate and 1 µM NcLPMO in 50 mM Bis-Tris buffer (pH 6.5). Reactions were initiated by adding 2 µl of 1.5 mM AscA immediately followed by addition of 2 µl of 5 mM H$_2$O$_2$. Every 15 min, an additional 2 µl of 1.5 mM AscA and 2 µl of 5 mM H$_2$O$_2$ (in this order) were added to the reaction mixtures (15 additions in total over a period of 4 h). Reactions were stopped by boiling for 10 min followed by separation of soluble and insoluble material by centrifugation at 11,000 × g for 10 min. Control reactions without added H$_2$O$_2$ were also performed.

Quantitative assessment of products released from PASC in reactions with H$_2$O$_2$

Reaction mixtures with H$_2$O$_2$ (600-µl final volume after initiation of the reaction) contained 2 mg·ml$^{-1}$ PASC and 1 µM NcLPMO in 50 mM Bis-Tris buffer (pH 6.5). Reactions were initiated by adding 6 µl of 1.5 mM AscA (final concentration, 15 µM) followed by addition of 6 µl of 5 mM H$_2$O$_2$ (final concentration, 50 µM). After exactly 15 min, an additional 6 µl of 1.5 mM AscA and 5 mM H$_2$O$_2$ were added. 30 min after the reactions were initiated, a 122-µl sample was taken from the reaction mixture and boiled for 10 min. To the rest of the reaction mixture (490 µl), 5 µl of 1.5 mM AscA (15 µM) and 5 µl of 5 mM H$_2$O$_2$ (50 µM) were added. The same volumes of 1.5 mM AscA and 5 mM H$_2$O$_2$ were also added at 45 min after reaction initiation. At 60 min, a 118-µl sample was taken and boiled for 10 min. To the rest of reaction mixture (392 µl), 4 µl of 1.5 mM AscA (15 µM) and 4 µl of 5 mM H$_2$O$_2$ (50 µM) were added. The same volumes of 1.5 mM AscA and 5 mM H$_2$O$_2$ were also added at 75, 90, and 115 min after the reactions were initiated. After 120 min, 130-µl samples were taken and boiled for 10 min. To the rest of reaction mixture (294 µl), 3 µl of 1.5 mM AscA (15 µM) and 3 µl of 5 mM H$_2$O$_2$ (50 µM) were added. The same volumes of 1.5 mM AscA and 5 mM H$_2$O$_2$ were added at 135, 150, and 165 min after reaction initiation. After 180 min, 122-µl samples were taken and boiled for 10 min. To the rest of the reaction mixture (196 µl), 2 µl of 1.5 mM AscA (15 µM) and 2 µl of 5 mM H$_2$O$_2$ (50 µM) were added, and this was repeated at 195, 210, and 225 min after reaction initiation. After 240 min, the reactions were stopped by boiling for 10 min.

Separation of soluble and insoluble materials was achieved by centrifugation at 11,000 × g for 10 min. Samples of the soluble fractions (32 µl) were mixed with 31 µl of 150 mM sodium acetate buffer (pH 4.5) and 1 µl of TrCeF7A (−1 µM) and incubated for 16 h at 37 °C. The oxidized products were subsequently detected and quantified using HPAEC-PAD and Glc4gemGlc as a standard as described above.

H$_2$O$_2$ production

Measurement of H$_2$O$_2$ production was done as described previously (78). A reaction mixture (180 µl) containing 1 µM LPMO, 5 units·ml$^{-1}$ horseradish peroxidase, and 100 µM AmplexRed in 50 mM Bis-Tris buffer (pH 6.5) was incubated for 5 min at 40°C in a 96-well microtiter plate in a plate reader (Multiskan™ FC microplate photometer, Thermo Fisher Scientific, Bremen Germany). The reaction was initiated by the addition of 20 µl of 500 µM AscA (50 µM final concentration) in each well, and the production of resorufin was monitored at 540 nm. Control reactions in the absence of LPMO were carried out to obtain the LPMO-independent resorufin production rate. This control reaction provided a background signal equal to 0.3% of the LPMO-catalyzed reaction and was subtracted from the latter. An H$_2$O$_2$ standard curve was prepared using the same conditions (without AscA and LPMO). The reactions were monitored for 45 min, and H$_2$O$_2$ production rates were derived from data points in the linear region between 0 and 8.5 min.

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