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**FgLPMO9A from *Fusarium graminearum* cleaves xyloglucan independently of the backbone substitution pattern**

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**Abstract**

Lytic polysaccharide monooxygenases (LPMOs) are important for the enzymatic conversion of biomass and seem to play a key role in degradation of the plant cell wall. In this study, we characterize an LPMO from the fungal plant pathogen *Fusarium graminearum* (FgLPMO9A) that catalyzes the mixed C1/C4 oxidative cleavage of cellulose and xyloglucan, but is inactive towards other (1,4)-linked β-glucans. Our findings indicate that FgLPMO9A has unprecedented broad specificity on xyloglucan, cleaving any glycosidic bond in the β-glucan main chain, regardless of xylosyl substitutions. Interestingly, we found that when incubated with a mixture of xyloglucan and cellulose, FgLPMO9A efficiently attacks the xyloglucan, whereas cellulose conversion is inhibited. This suggests that removal of hemicellulose may be the true function of this LPMO during biomass conversion.

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Keywords: AA9; cellulose; *Fusarium graminearum*; lytic polysaccharide monooxygenase; xyloglucan.

Abbreviations: AA, auxiliary activity; DP, degree of polymerization; GH, glycoside hydrolase; LPMO, lytic polysaccharide monooxygenase; MALDI-ToF MS, matrix-assisted laser desorption ionization time-of-flight mass spectrometry; PAD, pulsed amperometric detection; PASC, phosphoric acid swollen cellulose; SDS-PAGE, SDS polyacrylamide gel electrophoresis; TXG, tamarind xyloglucan.

INTRODUCTION

Plants have evolved multiple defense mechanisms against microbial pathogens, varying from defense systems triggered by recognition of microbe-associated molecular patterns, to physical barriers, such as the plant cell wall [1,2]. The plant cell wall is the major structural defense barrier against fungal and bacterial pathogens. The composite secondary cell wall consists of cellulose microfibrils embedded in a matrix of hemicellulose and lignin, with cross-linking between the polymers. The primary cell wall, i.e. the thin outer layer of plant cell walls, also contains pectins and structural proteins, which are much less abundant in the secondary walls [3–5]. Many plant pathogenic fungi produce enzymes that allow them to degrade and penetrate this complex plant cell wall barrier [6,7]. *Fusarium graminearum* is a highly destructive pathogen causing *Fusarium* head blight disease on wheat and barley. In addition, this devastating fungus produces mycotoxins, which pose a threat to human and animal health [8]. The genome of *F. graminearum* contains almost 600 predicted carbohydrate-active enzymes (CAZymes). These CAZymes include a variety of glycoside hydrolases (GHs) as well as 18 putative lytic polysaccharide monooxygenases (LPMOs) [9]. *F. graminearum* is an efficient degrader of plant biomass, secreting cell wall degrading enzymes, including cellulases, xylanases, pectinases and LPMOs [10,11].

LPMOs are copper-dependent redox enzymes that oxidatively cleave polysaccharides using molecular oxygen and an electron donor [12–15]. These enzymes are classified as auxiliary activities (AAs) in the Carbohydrate-Active enZyme database, comprising four families, AA9, AA10, AA11, and AA13 (CAZy; http://www.cazy.org) [16]. LPMOs cleave polysaccharides while oxidizing one of the new chain ends at the C1 or C4 position, thus contributing to substrate depolymerization while increasing accessibility of the substrate to conventional GHs [12,15,17,18]. Since their discovery in 2010 [12], LPMOs with preference for various plant polysaccharides, such as cellulose [19,20], xyloglucan and other (1,4)-linked β-glucans [21], starch [22] and xylan [23] have been identified. LPMOs have become an
important ingredient in commercial enzyme cocktails for industrial biomass conversion, such as Cellic CTec2 [24], owing to their synergy with GHs in improving saccharification yields [25,26].

In the present study, we cloned and characterized an AA9 LPMO from *F. graminearum*, hereafter referred to as *Fg*LPMO9A, which is the first *F. graminearum* LPMO to be characterized. We studied the action of *Fg*LPMO9A on various plant polysaccharides including cellulose and xyloglucan as well as mixtures of these two. We show that *Fg*LPMO9A possesses unprecedented broad specificity when acting on xyloglucan and we speculate that hemicellulose degradation may be the enzyme’s true biological function.

**MATERIALS AND METHODS**

**Cloning and expression of the *F. graminearum* LPMO**

The gene encoding *Fg*LPMO9A [UniProt: I1REU9], including its native signal sequence, was codon optimized for *Pichia pastoris* (GenScript, NJ, USA). The synthetic gene was inserted into the pPink-GAP vector, which was then transformed into *P. pastoris* PichaPink™ Strain 4 cells (Invitrogen, CA, USA), as described earlier [27]. Transformants were screened for protein production in BMGY medium (containing 1% (v/v) glycerol).

The best-producing transformant was pre-grown in 20 ml of BMGY medium (containing 1% (v/v) glycerol) in a 100-ml shake flask at 29 °C and 200 rpm for 16 hours. This pre-culture was used to inoculate 0.5 l BMGY medium (containing 1% (v/v) glycerol) in a 2-l baffled shake flask. After 24 hours of incubation at 29 °C and 200 rpm, the medium was supplemented with 1% (v/v) glycerol. After a total incubation time of 48 hours, the cells were harvested by centrifugation at 7,000 \( \times g \) for 15 min. The supernatant was filtered through a 0.2-µm polyethersulfone membrane (Millipore, MA, USA), dialyzed against 50 mM Bis-Tris buffer (pH 6.5), and concentrated to 50 ml with a VivaFlow 50 tangential crossflow concentrator (MWCO 10 kDa, Sartorius Stedim Biotech, Germany).

**Purification and Cu(II) saturation of the enzyme**

*Fg*LPMO9A was purified using a two-step purification protocol, starting with hydrophobic interaction chromatography (HIC) followed by size exclusion chromatography (SEC). The concentrated culture supernatant was prepared for HIC by slow addition of solid ammonium sulfate to 2.03 M final concentration, at 20 °C, followed by centrifugation (15,000 \( \times g \), 15 min, 4 °C). The sample was loaded

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onto a 5-ml HiTrap Phenyl FF column (GE Healthcare, Sweden) equilibrated with 50 mM Bis-Tris buffer (pH 6.5), containing 2.03 M ammonium sulfate. Proteins bound to the column were eluted using a 25-ml linear gradient from 2.03 M to 0 M ammonium sulfate in 50 mM Bis-Tris buffer (pH 6.5). Collected fractions were analyzed by sodium dodecyl sulfate–polyacrylamide gel electrophoresis (SDS–PAGE), and the fractions containing FgLPMO9A were pooled and concentrated using Amicon Ultra centrifugal filters (MWCO 10 kDa, Millipore). The sample was saturated with Cu(II) by incubating the enzyme with an excess of CuSO₄ (at 3:1 molar ratio of copper:enzyme) for 30 min at room temperature as described previously [28]. Subsequently, the sample was loaded onto a HiLoad 16/60 Superdex 75 size exclusion column (GE Healthcare) in 50 mM Bis-Tris buffer (pH 6.5), using a flow rate of 1.0 ml/min. Fractions containing pure protein were identified using SDS-PAGE and subsequently pooled and concentrated using Amicon Ultra centrifugal filters (MWCO 10kDa, Millipore). Protein concentrations were determined using the Bradford assay (Bio-Rad).

NcLPMO9C from Neurospora crassa [UniProt:Q7SHI8] was expressed in P. pastoris X-33 and purified as described previously [29].

**Substrates used for enzyme specificity analysis**

The following substrates were used for exploring enzymatic activities of FgLPMO9A: phosphoric acid swollen cellulose (PASC) prepared from Avicel as described by Wood [30]; xyloglucan (XG) from tamarind seed; xyloglucan-heptamer (XG7 or XXXG, where X stands for glucose, G, substituted at the C6 position with xylose); xyloglucan oligomers (XG-oligomers) derived from tamarind xyloglucan mainly containing XXXGXXGXG, with 0–3 galactose substitutions (L stands for X where the xylose is substituted at the C2 position with a galactose; it is not known which X is galactosylated); a reduced 14-mer (XG14OH) mainly containing XXXGXXGXGOH and variants containing one or more L instead of X (the reducing end D-glucose is reduced to a D-glucitol). FgLPMO9A was also screened against cello-oligosaccharides (Glc₃–₆), birchwood xylan, wheat arabinoxylan, konjac glucomannan, ivory nut mannann, barley β-glucan, lichenan from Icelandic moss, starch, and deacetylated galactoglucomannan from Norway spruce. Birchwood xylan was purchased from Sigma-Aldrich (MO, USA), starch was purchased from Merck (Darmstadt, Germany). All other substrates were purchased from Megazyme (Ireland).

**Enzyme activity assays**

Reaction mixtures with single substrates contained 2 mg/ml substrate, while the reactions with substrate mixtures contained 2 mg/ml PASC and 2 mg/ml of the other polysaccharide. Reaction mixtures
Additionally contained 2 µM FgLPMO9A or NcLPMO9C and 1 mM ascorbic acid in 20 mM Bis-Tris buffer (pH 6.5) in a total volume of 100 µl, in 2-ml Eppendorf tubes. Samples were incubated at 15 or 45 °C in an Eppendorf Thermomixer (Eppendorf AG, Germany) with shaking at 900 rpm. After incubation, the reaction mixtures were placed on ice and the reactions were immediately stopped by boiling for 10 min, before separating soluble and insoluble fractions using a 96-well filter plate (Millipore) operated by a Millipore vacuum manifold. Control reactions were performed using identical conditions, but in the absence of ascorbic acid.

**Analysis of enzyme products**

Products generated by the LPMOs were analyzed using MALDI-ToF mass spectrometry (MS) and high-performance anion exchange chromatography (HPAEC) as follows. MALDI-ToF MS analysis was performed with an Ultraflex MALDI-ToF/ToF instrument (Bruker Daltonics, Germany) equipped with a Nitrogen 337 nm laser beam as described by Vaaje-Kolstad et al. [12]. The instrument was operated in positive acquisition mode and controlled by the FlexControl 3.3 software package. The data were processed with mMass software [31]. Baseline correction and Gaussian smoothing (window size 0.3 m/z) were applied to all spectra. Prior to MALDI-ToF MS analysis, the samples were saturated with sodium by mixing 5 µl sample with 5 µl 50 mM sodium acetate, followed by 30 min incubation, sample spotting and drying.

HPAEC analysis was performed on a Dionex ICS-5000 system (Dionex, CA, USA) equipped with pulsed amperometric detection (PAD) and a CarboPac PA1 analytical column (2 × 250 mm) with a CarboPac PA1 guard column (2 × 50 mm). The system was operated at a flow rate of 0.25 ml/min and kept at 30 °C while running a 75-min stepwise gradient as described by Agger et al. [21]. The data were collected and analyzed using Chromeleon 7.0 (Dionex).

**RESULTS**

**Heterologous expression of FgLPMO9A from F. graminearum**

Recombinant FgLPMO9A was successfully expressed by growing *P. pastoris* in BMGY medium for 48 hours. The enzyme was purified in two chromatographic steps to ≈95% purity, as confirmed by SDS-PAGE (see Fig. S1). The protein band representing FgLPMO9A appears as a broad band in the 60–70 kDa range, whereas the theoretical molecular mass calculated from the amino acid sequence is 32 kDa. The observed mass difference is likely due to O- and/or N-glycosylation, which are predicted (using the
NetOGlyc and NetNGlyc servers available at http://www.cbs.dtu.dk/services) to possibly occur at Ser/Thr 234, 235, 238, 240, 241, 243, 245, 247, 249, 251, 257, 262, 287, 292 and Asn211, respectively. Such mass differences are not uncommon although in the present case the difference is rather large, possibly due to a large number of O-glycosylations in the C-terminal region of low complexity and unknown function (residues 226-314). Notably, these putative O-glycosylations are likely to affect the substrate-binding surface of the LPMO. Sequence alignments and structural modelling indicated that the only potential N-glycosylation site (Asn211) is located far away from the catalytic center and that the side chain of this residue points away from the substrate-binding surface. Attempts to deglycosylate the enzyme by treatment with PNGase F from Flavobacterium meningosepticum (New England Biolabs, Ipswich, MA) or α-mannosidase from Canavalia ensiformis (Sigma-Aldrich) were not successful, although a small shift in electrophoretic mobility was observed with the latter enzyme (results not shown).

**Substrate specificity**

In order to determine substrate specificity, FgLPMO9A was incubated with a wide range of oligo- and polysaccharides in the presence of an electron donor. Studies with insoluble substrates showed that the enzyme is capable of cleaving cellulose (Fig. 1), but not chitin (results not shown). No activity was detected towards shorter cellobiose saccharides (Glc₃₋₆, results not shown). Among the tested hemicelluloses, FgLPMO9A was active on tamarind xyloglucan and longer xyloglucan oligosaccharides (see below for details). No activity was observed towards the xyloglucan-heptamer (XXXG), birchwood xylan, wheat arabinoxylan, konjac glucomannan, ivory nut mannan, β-glucan from barley, lichenan from Icelandic moss, starch, and spruce galactoglucomannan (data not shown).

**Activity on amorphous cellulose (PASC)**

When using PASC as a substrate, HPAEC analysis of the products generated by FgLPMO9A showed native as well as C1- and C4-oxidized cellobiose saccharides (Fig. 1A). In the absence of an electron donor, no LPMO activity was detected. Oxidative cleavage of cellulose and the C1/C4 mixed oxidation pattern were confirmed by MALDI-ToF MS analysis of the released products (Fig. 1B–C). Notably, both types of oxidized products may occur in the non-hydrated and the hydrated form, the latter being a gemdiol for C4-oxidized products and an aldonic acid for C1-oxidized products. For the C4-oxidized compounds, the non-hydrated ketoaldose form is favored (m/z 1335.7, for DP 8). On the other hand, C1-oxidized compounds largely occur as the hydrated aldonic acid form (m/z 1353.6 for DP 8), which at neutral conditions forms a diagnostic sodium salt (m/z 1375.5 for DP 8). Other species visible in Fig. 1C are the native oligomer (m/z 1337.7), a double-oxidized oligomer with one hydration (m/z 1351.6), and a
double-oxidized oligomer with two hydrations (m/z 1369.6). The appearance of native oligosaccharides and double-oxidized oligosaccharides confirms C1/C4 mixed oxidation. All product clusters, ranging in DP from 5 to 10 (Fig. 1B), showed adduct distributions similar to that shown for the DP 8 cluster in Fig. 1C.

**Activity on tamarind xyloglucan**

During the course of this study, we noted that the activity of FgLPMO9A was considerably higher at 15 °C compared to 45 °C; hence reactions with TXG were carried out at both temperatures. FgLPMO9A was capable of cleaving tamarind xyloglucan (TXG), and HPAEC analysis (Fig. 2) revealed a more complex product profile compared to the product profile generated by NcLPMO9C [21]; see below). FgLPMO9A produced a mixture of oligosaccharides, some of which corresponded to the native xyloglucan heptasaccharide (XXXG), eluting at 26 min. Some species eluted before XXXG, which are likely to be shorter native fragments, such as XXX, eluting at 22 min (see [21]). The elution profile in the 50-55 min range is similar to the profile previously published for NcLPMO9C [21]. In addition, a broad range of products appears at 26-50 min that has not been observed before, not for NcLPMO9C, nor for the other xyloglucan-active LPMOs described so far, PoLPMO9H and TtLPMO9E [21,32,33]. Although some of these peaks may correspond to longer native XG oligomers (see black line in Fig. 2), most of the products released from TXG remain unidentified due to the heterogeneity of the substrate and lack of standards.

Analysis of the reaction products with MALDI-ToF MS revealed that FgLPMO9A indeed produces a much wider range of native and oxidized xyloglucan-oligosaccharides (Fig. 3A), compared to NcLPMO9C (Fig. 3B). Both enzymes released a cluster of Hex$_n$Pen$_3^{ox}$ products with m/z 1083.7 (n=4), m/z 1245.5 (n=5), and m/z 1407.5 (n=6) and a cluster of Hex$_n$Pen$_6^{ox}$ products with m/z 2452.1 (n=10), m/z 2614.1 (n=11), and m/z 2776.2 (n=12), where Hex stands for a hexose (glucose or galactose, 162.14 Da) and Pen for a pentose (xylose, 132.11 Da). Only FgLPMO9A released a wide variety of additional products where the number of pentoses does not equal a multitude of 3. As an example, Fig. 3C shows signals for Hex$_7$Pen$_4^{ox}$, showing a double oxidized species (m/z = 1699.5), a single oxidized species (m/z = 1701.5), the native species (m/z = 1703.6), and hydrated single (m/z = 1719.6) and double (m/z = 1717.5) oxidized species. Taken together, the data in Figs. 3A-C show that while NcLPMO9C only cleaves at the unsubstituted glucose unit and exclusively oxidizes C4 [21], FgLPMO9A oxidizes both C1 and C4 and cleaves TXG at any position in the β-glucan backbone, i.e. also between two substituted units (Fig. 3D).
Activity on mixtures of hemicelluloses and PASC

It has previously been shown that certain LPMOs can cleave xylan, only if the xylan is complexed with cellulose [23]. In order to gain further insight into the potential role of FgLPMO9A in plant cell wall degradation, we mixed cellulose (PASC) with different hemicelluloses and treated the resulting polysaccharide mixtures with FgLPMO9A, in the presence or absence of ascorbic acid as electron donor. Mixing PASC with birchwood xylan or ivory nut mannan reduced PASC conversion, while mixing with konjac glucomannan completely inhibited activity on PASC. We were not able to detect cleavage of these three hemicelluloses (data not shown). Upon mixing TXG and PASC, activity on PASC was greatly reduced, whereas the activity on TXG was hardly affected (Fig. 2; note the lack of native cello-oligomers in the red chromatograms, relative to the green chromatograms).

It is conceivable that mixing TXG with PASC leads to complexation where the hemicellulose forms a physical barrier on the surface of the cellulose. To investigate this further, we carried out studies with a cellulose-active LPMO that is inactive towards xyloglucan. These studies showed that the activity of the cellulose-active LPMO was inhibited by adding TXG to the PASC and that this inhibition could be relieved by adding an enzyme degrading TXG (for details, see Fig. S2 and supporting discussion).

Activity on a reduced xyloglucan-oligosaccharide (XG14)

To gain a deeper understanding of the FgLPMO9A activity towards TXG, we studied the degradation products generated by the enzyme when incubated with the reduced form of a relatively pure xyloglucan oligosaccharide with DP14 (XG14OH). The sequence of this oligomer is XXXGXXXGOH, where GOH represents a D-glucitol, reduced glucose, at the reducing end, leading to an m/z shift of +2 Da compared to a native species. The product profile was analyzed using HPAEC and MALDI-ToF MS (Fig. 4). Only in the presence of ascorbic acid, FgLPMO9A generated a variety of products, including the native XXXG peak eluting at 26 min and several additional compounds eluting before XXXG, such as the native XXX at 22 min. These latter compounds also appeared in the chromatograms obtained upon degradation of TXG by FgLPMO9A (Fig. 2) and are likely to be shorter native fragments. In addition, some of these compounds may be reduced oligosaccharides (with a D-glucitol at the end), which also elute earlier than their corresponding native oligosaccharides [21]. At the same time, several product species eluting after the substrate, at 44-48 min and at 56-64 min, were visible similar to products generated by NcLPMO9C from this same substrate [21].

Oxidative cleavage of XG14OH by FgLPMO9A was confirmed by analysis of the products with MALDI-ToF MS (Fig. 4B). A single cleavage of XG14OH with C1-oxidation generates two products that differ with
Δm/z ±2 from the native ones: the non-reducing end product carrying the C1-oxidation (Δm/z=-2) and the reducing end product carrying the reduced glucose at the “reducing” end (Δm/z=+2). On the other hand, a single cleavage with C4-oxidation generates two products with masses identical to those of native species (Δm/z=0): the non-reducing end product is a native oligosaccharide and the reducing end product carries a keto-group from the C4-oxidation (Δm/z=-2) and the reduced D-glucitol at the “reducing end” (Δm/z=+2), (Fig. 4C). The mass spectrum shows a wide range of xyloglucan oligosaccharides with masses corresponding to native species or differing from that with Δm/z±2, showing that both C1 and C4 oxidation occurred and that the enzyme can cleave the substrate at several positions. The only two C1-oxidized products detected were Hex$_3$Pen$_3^{ox}$ (m/z = 921.5; Δm/z=-2) and Hex$_4$Pen$_3^{ox}$ (m/z = 1083.7; Δm/z=-2), which are indicative of C1 oxidation at the non-substituted glucose, perhaps only on the non-reducing side of this glucose (due to varying galactosylation, both Hex$_3$Pen$_3^{ox}$ and Hex$_4$Pen$_3^{ox}$ could be products). The occurrence of a wide variety of products with a number of pentoses not equaling a multitude of 3 shows that cleavage between two substituted units occurred. The fact that these products emerged both as reduced species (Δm/z=+2) and native species (Δm/z=0) shows that both C1- and C4-oxidation occur during cleavage in between substituted units (e.g. m/z 1115.6 and 1117.6 for Hex$_3$Pen$_3$, or m/z 1541.8 and 1543.7 for Hex$_6$Pen$_4$).

**DISCUSSION**

The discovery of LPMOs has been a significant breakthrough in understanding how plant biomass is degraded in nature, and has also had major technological implications, as illustrated by the fact that, today, these enzymes are important components of enzyme cocktails for industrial biomass conversion [34]. LPMOs are produced not only by saprophytic fungi but also by plant pathogens, suggesting a putative role of these enzymes in plant cell wall degradation during pathogenesis [10]. *F. graminarum* displays both a saprophytic and a pathogenic lifestyle and to what extend *Fg*LPMO9A plays a role in each of these is not yet known. Notably, it has been shown that most of the putative LPMOs encoded in the genome of *F. graminearum*, including *Fg*LPMO9A, are upregulated during infection of barley and wheat [35]. Although to date the CAZy database harbors more than 400 fungal LPMOs, only about 20 of these enzymes have been characterized, and these are all from saprophytic fungi. To our knowledge, *Fg*LPMO9A is the first LPMO from a plant pathogen to be characterized.
FgLPMO9A has mixed oxidative regioselectivity, oxidizing both C1 and C4 in insoluble (cellulose) and soluble (xyloglucan) substrates. The enzyme produces relatively large amounts of native cello- and xyloglucan-oligosaccharides (Figs. 1–3). Native products are commonly observed for mixed activity LPMOs, and could emerge when a single polysaccharide chain is cleaved twice, once with C1 and once with C4 oxidation, with the C1 oxidation happening upstream of the C4 oxidation. Notably, native products are also commonly observed with HPAEC analysis for strictly C4 oxidizing LPMOs, as C4-oxidized products are converted to native cello-oligomers after losing the 4-ketoglucose at the non-reducing end (upstream end), promoted by the conditions of standard HPAEC analysis, which has recently been shown \[36\]. Hence, some of the native oligosaccharides seen in the HPAEC profile are likely degradation products of the corresponding C4-oxidized compounds. Control reactions, where ascorbic acid was excluded from the reaction, showed no formation of oxidized or native xyloglucan- or cello-oligomers, verifying that the FgLPMO9A preparation was free of contaminating background endo-β-1,4-glucanase activity.

So far, only four LPMOs have been reported to cleave xyloglucan, namely NcLPMO9C from Neurospora crassa, PaLPMO9H from Podospora anserina, AN3046 LPMO from Aspergillus nidulans and TtLPMO9E from Thielavia terrestris; and the cleavage pattern of NcLPMO9C and PaLPMO9H has been reported [21,32,33,37]. Both enzymes cleave xyloglucan exclusively adjacent to the unsubstituted glucosyl unit, with NcLPMO9C only yielding C4-oxidized products, whereas PaLPMO9H yields a mixture of C1- and C4-oxidized products with the latter being dominant [32]. Considering the oxidative mechanism, FgLPMO9A is more similar to PaLPMO9H since it oxidizes both C1 and C4. However, FgLPMO9A is fundamentally different from both enzymes described above, since FgLPMO9A is not restricted to cleaving by the unsubstituted glucosyl, thereby generating a broader product spectrum. The lack of activity on cellohexaose further separates FgLPMO9A from NcLPMO9C and PaLPMO9H, the two well-characterized xyloglucan-active LPMOs, and indicates that the catalytic site of FgLPMO9A needs to accommodate a β-1,4-glucan chain with more than six glucose residues for catalysis to happen.

Interestingly, the XG-cleaving properties of FgLPMO9A are not commonly found amongst XG-active enzymes, as the majority of GHs cleaving this polymer are selective for non-substituted backbone glucose residues. The only GHs known to cleave XG between substituted backbone residues are found in the GH74 family [38,39]. Thus, our findings provide a new activity to the XG enzymatic toolbox. The F. graminearum genome only contains one GH74 enzyme, which has not yet been characterized and thus has an unknown function (the GH74 family also harbors enzymes with endoglucanase and exo-
xyloglucanase activity). It may be that \textit{Fg}LPMO9A plays a crucial role in the XG degradative machinery of the fungus by providing the ability to cleave between substituted backbone residues.

The effect of temperature on \textit{Fg}LPMO9A activity is remarkable and has several potential explanations. Control experiments showed that the enzyme is thermally stable under the conditions used (see Fig. S3 and supporting discussion), excluding temperature-induced denaturation as an explanation. In this qualitative study of the activity of \textit{Fg}LPMO9A, we routinely used 16-hour incubation times. A closer examination of product formation over time (Fig. S4) revealed that enzyme activity ceases already after 1–4 hours. Higher incubation temperatures lead to faster initial rates but more rapid inactivation of the enzyme, the net effect being that incubation at the lower temperature gives higher final product yields. Enzyme inactivation is often observed in LPMO reactions but has hardly been discussed and remains unexplained. Inactivation could be due to the production of \text{H}_2\text{O}_2 [29] and more powerful oxidative species (such as hydroxyl radicals) that may damage the LPMO molecule. A further discussion of this issue is beyond the scope of this paper, but the present data show that this issue deserves more attention.

\textit{Fg}LPMO9A did not show detectable activity on xylan, arabinoxylan, glucomannan, mannan, \(\beta\)-glucan, or lichenan. Mixing PASC with several hemicelluloses did not reveal hemicellulolytic activities beyond activity on xyloglucan but did show inhibition of activity on PASC, likely because hemicellulose-PASC associations shield the cellulose. This observation was explored further for xyloglucan, which is thought to associate with cellulose to form load-bearing networks in plant primary cell walls [3,40,41]. Interestingly, when using a mixture of xyloglucan and cellulose as substrate, the activity of \textit{Fg}LPMO9A on xyloglucan was maintained, whereas activity on cellulose was strongly reduced. Control experiments with a cellulose-active LPMO that is not active on xyloglucan (see Supporting information) showed that cellulose degradation is inhibited by the addition of xyloglucan and that such inhibition may be relieved by also adding a xyloglucanase. All in all, these data suggest that xyloglucan shields cellulose from being degraded by \textit{Fg}LPMO9A and that one natural function of this LPMO could be to make cellulose accessible.

LPMOs are thought to have a big impact on the depolymerization of organic matter in nature [42]. It is conceivable that \textit{Fg}LPMO9A, with its unique ability to randomly cleave xyloglucan, contributes to overcoming the plant cell wall barrier by removing hemicellulose and by making other polymers such as cellulose more accessible for enzymatic degradation.
SUPPORTING INFORMATION

Additional supporting information, including Figs. S1–S4 may be found in the online version of this article at the publisher’s web site.

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FIGURE LEGENDS

Figure 1. Reaction products generated by FgLPMO9A from PASC. (A) HPAEC-PAD profiles of reaction mixtures containing FgLPMO9A and PASC, with (red line) and without (yellow line) ascorbic acid; the grey line shows cello-oligosaccharide standards with DP 2-6. The produced oxidized and native cello-oligosaccharides are labeled in the figure and annotations are based on previous work [17,18]. (B) MALDI-ToF MS spectrum of oligosaccharides released from PASC by FgLPMO9A, ranging from DP 5-10, after Na⁺-saturation. (C) Close-up of the DP 8 cluster showing the sodium adduct of native and oxidized products; single or double oxidation is denoted with “ox and 2-ox”; single or double hydration is indicated by * or **. See text for a discussion of the various peaks.

Figure 2. HPAEC-PAD chromatograms showing products obtained after FgLPMO9A action on tamarind xyloglucan (TXG, blue), PASC (green) or 1:1 TXG-PASC mixtures (red). Samples were incubated at 15 °C (dashed lines) or 45 °C (solid lines) with ascorbic acid (AA) for 16 hours. In addition, chromatograms for a xyloglucan-oligosaccharide standard (black line), a xyloglucan heptamer (XXXG) standard (grey line) and TXG (yellow line) are shown. Control reactions showed that neither native nor oxidized xyloglucan oligosaccharides were generated by FgLPMO9A in the absence of an electron donor.

Figure 3. Reaction products generated from tamarind xyloglucan (TXG). (A,B) MALDI-ToF MS spectra showing product profiles generated from TXG by (A) FgLPMO9A and (B) NcLPMO9C with oxidized product peak annotations. Note the much more complex product profile obtained with FgLPMO9A. (C) Close-up of the Hex₇Pen₄ cluster showing the sodium adduct of native and oxidized products; single or double oxidation is denoted with “ox and 2-ox”; hydration is labeled with “**”. See text for a discussion of the various peaks. Product distributions in other DP clusters were similar. Abbreviations: Hex, hexose (+ 162 Da); Pen, pentose (+ 132 Da); ox, oxidized. (D) Illustration of the structure of a fragment of TXG (blue circle, glucose; orange star, xylose; yellow circle, galactose [43]; the position and number of galactosyl units vary) and possible cleavage sites for FgLPMO9A (black arrows) and NcLPMO9C (red arrows; see also ref. [21]).
**Figure 4.** Analysis of products generated from XG14\(^{\text{OH}}\). **(A)** HPAEC-PAD chromatograms showing products generated upon incubation of XG14\(^{\text{OH}}\) with *Fg*LPMP09A (blue line), as well as non-treated XG14\(^{\text{OH}}\) (black line) and the xyloglucan heptamer (XG7, XXXG) standard (orange line). **(B)** MALDI-ToF MS spectrum showing the sodium adduct of products generated from XG14\(^{\text{OH}}\) by *Fg*LPMP09A (blue line) and the non-treated XG14\(^{\text{OH}}\) (black line); the indicated \(m/z\) values represent mass average. Numbers in parenthesis show \(\Delta m/z\) compared to the mass of the native XG-oligosaccharide products, as explained in panel C; blue peaks are either from the substrate or the result of C1 oxidation which yields oxidized (\(\Delta m/z = -2\), “ox”) and reduced (\(\Delta m/z = +2\), “red”) oligomers; green peaks are the result of C4 oxidation (\(\Delta m/z = 0\) for all species, hence no further indication of oxidation and reduction; see panel C). The peaks at \(m/z\) 1999.9, 2132.0 and higher represent the XG14\(^{\text{OH}}\) substrate (see black line), which contains a mixture of oligomers with varying degrees of galactosylation (see panel C). Reactions without ascorbic acid showed only these peaks. Due to high background signals in the lower mass range, the spectra are shown from \(m/z\) 800. **(C)** The substrate, XG14\(^{\text{OH}}\) (blue circle, glucose; orange star, xylose; yellow circle, galactose [43]), and comparison of the C1- and C4-oxidation products and their masses relative to the masses of native species. Parenthesis surrounding galactosyl-units signify that the number and position of these units vary. Abbreviations: Hex, hexose (+ 162.14 Da); Pen, pentose (+ 132.11 Da); red, reduced; ox, oxidized.