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The Maize Pathogen Ustilago maydis Secretes Glycoside Hydrolases and Carbohydrate Oxidases Directed toward Components of the Fungal Cell Wall

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ABSTRACT Filamentous fungi are keystone microorganisms in the regulation of many processes occurring on Earth, such as plant biomass decay and pathogenesis as well as symbiotic associations. In many of these processes, fungi secrete carbohydrate-active enzymes (CAZymes) to modify and/or degrade carbohydrates. Ten years ago, while evaluating the potential of a secretome from the maize pathogen Ustilago maydis to supplement lignocellulolytic cocktails, we noticed it contained many unknown or poorly characterized CAZymes. Here, and after reannotation of this data set and detailed phylogenetic analyses, we observed that several CAZymes (including glycoside hydrolases and carbohydrate oxidases) are predicted to act on the fungal cell wall (FCW), notably on β -1,3-glucans. We heterologously produced and biochemically characterized two new CAZymes, called UmGH16_1-A and UmAA3_2-A. We show that UmGH16_1-A displays β -1,3-glucanase activity, with a preference for β -1,3-glucans with short β -1,6 substitutions, and UmAA3_2-A is a dehydrogenase catalyzing the oxidation of β -1,3- and β -1,6-gluco-oligosaccharides into the corresponding aldonic acids. Working on model β -1,3-glucans, we show that the linear oligosaccharide products released by UmGH16_1-A are further oxidized by UmAA3_2-A, bringing to light a putative biocatalytic cascade. Interestingly, analysis of available transcriptomics data indicates that both UmGH16_1-A and UmAA3_2-A are coexpressed, only during early stages of U. maydis infection cycle. Altogether, our results suggest that both enzymes are connected and that additional accessory activities still need to be uncovered to fully understand the biocatalytic cascade at play and its physiological role.

IMPORTANCE Filamentous fungi play a central regulatory role on Earth, notably in the global carbon cycle. Regardless of their lifestyle, filamentous fungi need to remodel their own cell wall (mostly composed of polysaccharides) to grow and proliferate. To do so, they must secrete a large arsenal of enzymes, most notably carbohydrate-active enzymes (CAZymes). However, research on fungal CAZymes over past decades has mainly focused on finding efficient plant biomass conversion processes while CAZymes directed at the fungus itself have remained little explored. In the present study, using the maize pathogen *Ustilago maydis* as model, we set off to evaluate the prevalence of CAZymes directed toward the fungal cell wall during growth of the fungus on plant biomass and characterized two new CAZymes active on fungal cell wall components. Our results suggest the existence of a biocatalytic cascade that remains to be fully understood.

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lamentous fungi play a central regulatory role on Earth. Saprophytic fungi, through the decomposition of dead matter, are instrumental in the global carbon cycle while mycorrhizal fungi, ensure the survival of most plants via symbiotic associations (1). On the "dark side of the force," pathogenic fungi, which can cause dramatic crop losses or severe human diseases, also affect ecosystems balance. Significant advances have been achieved in the past decade, notably via ambitious genome sequencing programs (2), and postgenomic studies (3, 4), to understand the fungal strategies put in place in these various ecological contexts. This collective corpus of data clearly indicates that during their life cycle filamentous fungi deploy an extraordinary diversity of enzymes, encompassing notably a wide array of carbohydrate-active enzymes (CAZymes) (5–10). In a context largely dominated by the overarching goal of developing efficient plant biomass conversion processes for biorefinery purposes, the study of the enzymatic arsenal of filamentous fungi has logically focused on enzymes targeting plant components, notably the plant cell wall (PCW). Strikingly, the role of secreted enzymes potentially directed toward the fungus itself has remained under the radar of most studies. Developmental biology of fungi has taught us that to explore their environment, and eventually interact with and/or infect their host, filamentous fungi need to remodel their own cell wall (11). The fungal cell wall (FCW) can also serve as an emergency carbon source, via autophagy, in the case of external carbon source shortage (12). Deciphering the potential role of endogenous FCW-targeting enzymes, and their orchestration, is thus of utmost importance. Similar to lignocellulose, the FCW is an intricate multilayer of complex polymers and is depicted today as being composed of an inner layer of chitin, a middle layer of β -1,3-glucans and an outer layer of mannoproteins (13). Some fungal species are reported to also display galactoaminoglycans (14). Several FCW-targeting enzymes secreted by plants as defensive mechanism, notably β -1,3-glucanases and chitinases, have also been reported (15). However, the identity and role of FCW-active CAZymes produced by the fungus itself remains underexplored.

In the present study, we have used as a study model the maize biotrophic pathogen *Ustilago maydis*, also known as corn smut, causing major crop yield losses every year (16). *U. maydis* is a rather peculiar filamentous fungus among basidiomycetes as it is a dimorphic fungus (i.e., able to switch from yeast to filamentous state). Also, out of a total of 230 CAZymes-encoding genes (www.cazy.org; [17]), *U. maydis* possesses a rather poor set of lignocellulolytic CAZymes. Yet, its total secretome produced on maize bran was found to contain 86 proteins, including 23 CAZymes predicted to target the PCW (10). Of note, the latter study on the secretome of *U. maydis* was reported in 2012, i.e., before the extension of the CAZy database with auxiliary activities (AA; [18]) and fine sequence-function understanding of certain GH (Glycoside Hydrolase) families, such as GH16 (19). Today, the AA class comprises oxidoreductases that have gained significant importance as they target a wide range of oligonucleotide and poly-saccharides found in PCW (7) and/or FCW.

Here, we have reanalyzed the secretomic data published in 2012 (10) in light of today's knowledge and identified several enzymes potentially targeting the FCW rather than the PCW. We demonstrate that two of these enzymes, which belong to the GH16 and AA3 CAZy families, are active on β -1,3-glucans or compounds thereof. Our results suggest that both enzymes are most likely involved in a common biocatalytic cascade of importance for the fungus' lifestyle.

RESULTS

Reassessment of *U. maydis* secretome on corn bran reveals the presence of putative FCW-active enzymes. The secretome of *U. maydis*, cultivated on corn bran, was first reported in 2012 (10). At that time, the identified top enzymes were arabinoxylandegrading enzymes (GH10, GH27, GH51, GH62). Here, taking advantage of progresses

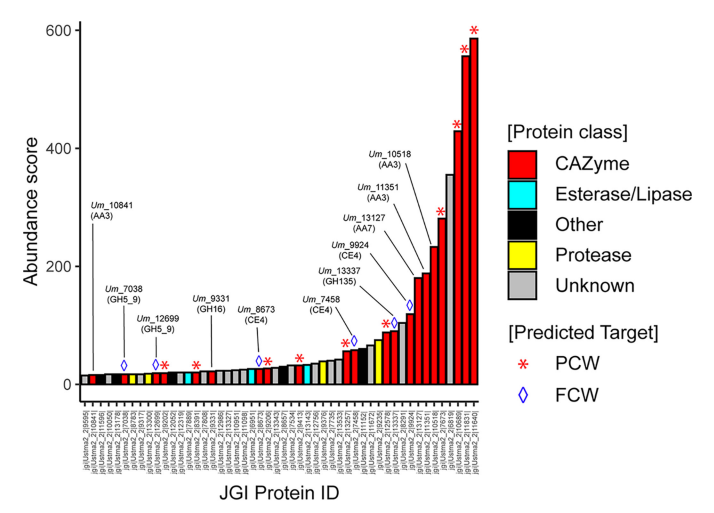


FIG 1 Reannotation of the Top50 proteins secreted by *U. maydis* when cultivated on corn bran. The enzymes are classified according to their abundance in the secretome (after 7 days growth on maize bran; [10]) and a color code identifies the class of protein (see legend in the figure, "Other" refers to all other types of detected proteins). The protein number that is provided corresponds to the JGI protein ID (*U. maydis* 521 v2.0 strain).

made in the CAZy field since then, and notably the creation (18) and enrichment of the AA class (20), we set off to reannotate and evaluate the enzymes deployed by *U. maydis* during the conversion of corn bran. Out of the top 50 proteins, 21 are CAZymes (13 GHs, one expansin, three CE4 and four AAs) (Fig. 1). Among them, 10 can clearly be predicted as active on PCW (Table S1), whereas the roles/targets of the 11 others (two GH5_9s, one GH16, one GH135, three CE4s, three AA3_2s, and one AA7) are not so obvious (all the corresponding UMAG_ID can be found in the legend of Fig. S1).

In the present study, our selection of enzymes was guided by (i) the substrate targeted, and (ii) the likeliness of biological interplay between enzymes. After carrying out phylogenetic analyses, we decided to focus on enzymes predicted to target the main component of the FCW, i.e., the β -1,3-glucans.

Among the putative FCW-active enzymes detected in the secretome (Table S1), the three CE4 enzymes (*Um*_7458, *Um*_9924 and *Um*_8673; the number corresponds to the JGI protein ID) are directed toward the chitin fraction, as they were biochemically characterized as chitin deacetylases in a recent study by Rizzi et al. (in which they were, called, respectively, *Um*CDA1, *Um*CDA3 and *Um*CDA4) (21). These CE4 enzymes were notably shown to be necessary for development and virulence of *U. maydis*. The enzyme from GH135 family (*Um*_13337) is predicted to be active on galactoaminogalactan (GAG), a polysaccharide of the extracellular matrix covering the cell wall.

Regarding the remaining enzyme candidates, to help us in the selection of the most relevant ones for biochemical validation and interplay studies, we searched for

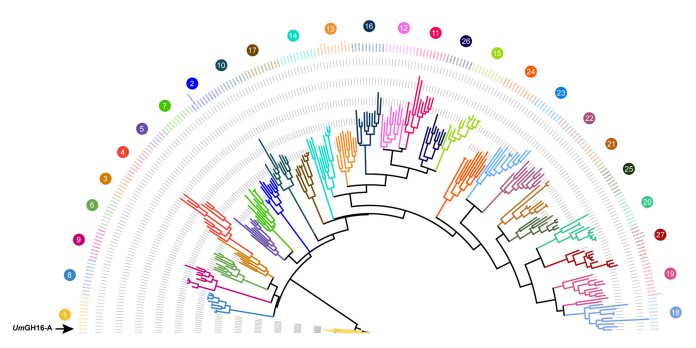


FIG 2 Phylogenetic analysis of GH16 family. Phylogenetic clades, as defined by Viborg et al. (19), are indicated with colored numbers. UmGH16_1-A (indicated by a black arrow) falls within the GH16_1 clade. The tree was inferred using RAxML (100 bootstraps) on the basis of a MSA made with MAFFT.

hints from biological conditions. To this end, we parsed available data reporting the transcriptomic profiling of the entire life cycle of *U. maydis* on maize (22) (Fig. S1A). Figure S1B shows the differential transcription along the pathogenic cycle of the genes coding for the 11 CAZymes mentioned above with putative activity on FCW (18). We observed that one of the AA3_2 (JGI ID 10841/UMAG_03551), the GH16 (JGI ID 9331/UMAG_02134) and one of the CE4 (JGI ID 7458/UMAG_0638) are the only ones to display a similar expression profile: they are expressed at relatively low levels, very early in the cycle (0.5 to 1 dpi) and are clearly downregulated during the plant infection cycle. We underscore that the CE4 is known to act on chitin (21), whereas the GH16_1 and AA3_2 are shown in the present study to act on β -1,3 glucans and oligomers thereof.

Phylogenetic analysis of the taxonomically broad GH16 family revealed that, out of 27 subfamilies (19), the GH16 *Um_*9331 belongs to the GH16_1 subfamily (Fig. 2) and is henceforth called *Um*GH16_1-A (as it is the first GH16 from *U. maydis* to be biochemically characterized). The GH16_1 subfamily is composed of almost exclusively fungal sequences, with the following reported activities: mainly *endo-* β -(1, 3)-glucanases (EC 3.2.1.39), *endo-* β -(1, 3)/(1, 4)-glucanases (EC 3.2.1.35) (23), and *exo-* β -(1, 3)-glucosyltransferase/elongating β -transglucosylase (EC 2.4.1.–) (24). *Um*GH16_1-A is thus potentially active on β -1,3-glucans components found in the FCW.

The AA3 family is a rather broad family divided into four subfamilies and composed of FAD-dependent oxidases (i.e., main electron acceptor is O_2) and dehydrogenases (organic electron acceptor) that oxidize various types of electron donors (25). Phylogenetic analysis revealed that the three AA3s (Um_10518 , Um_10841 and Um_11351) detected in the secretome of U. maydis all fall within the AA3_2 subfamily (Fig. 3A). A closer look at the AA3_2 subfamily (Fig. 3B) reveals that Um_10518 and Um_11351 belong to undefined groups. Interestingly, Um_10841 (henceforth called $UmAA3_2-A$) clusters together with the AA3_2 from the white-rot fungus *Pycnoporus cinnabarinus* hitherto called *Pc*GDH (26), and recently renamed oligosaccharide dehydrogenase (ODH) after it was shown to be active on laminaribiose (G3G; Glc- β -1,3-Glc) and on mixed β -1,3/ β -1,4 trimers (G3G4G; Glc- β -1,3-Glc- β -1,4-Glc) (27). Of note, *Pc*ODH appeared to be much more active on G3G than on glucose, the initially reported substrate. This recent finding highlights that the phylogenetic functional

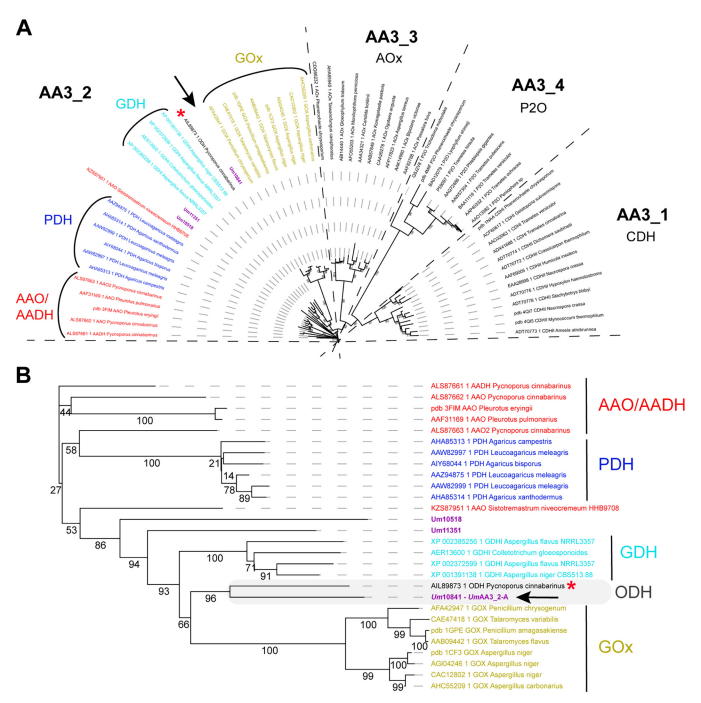


FIG 3 Phylogenetic analysis of the AA3 family (A) and zoom-in view on the AA3_2 subfamily (B). The AA3s identified in the secretome of *U. maydis* are shown in purple. The new oligosaccharide dehydrogenase clade, including *Um*AA3_2-A (indicated by a black arrow), characterized in the present study, and the *Pc*ODH (red asterisk), is framed in gray. The tree was inferred using PhyML (bootstrap values, as percentages, are shown on the branches).

annotation and biological role of AA3_2s is far from being firmly established and that *Pc*ODH and *Um*AA3_2-A may form a new, intermediate clade in between GOX and GDH activities. While this analysis suggests that *Um*AA3_2-A could be a candidate for the oxidation of β -1,3-glucans components, its rather dissimilar sequence (46% sequence identity with *Pc*ODH) called for biochemical investigations.

UmGH16_1-A is a β -1,3-glucanase with preference for β -1,3-glucans branched with short β -1,6 substitutions. UmGH16_1-A was heterologously expressed in *Pichia pastoris* and purified to homogeneity (Fig. S2), but the protein yield was very low (0.175 mg/liter of culture), preventing extensive characterization. Sequence and

structure comparisons between UmGH16_1-A (Alpha-fold2 model; [28]) and its closest structural homologue (RMSD of 0.473 Å and sequence identity of 38%), the GH16_1 from Phanerochaete chrysosporium (PcGH16_1; also called Lam16A; PDB id 2W52; [29]), indicated the presence in UmGH16_1-A of an extra 57 amino acid C-terminal extension with no predictable canonical fold (Fig. S3). A protein BLAST search on NCBI against the nr database showed the occurrence of orthologs of UmGH16_1-A bearing similar C-terminal extensions in a broad range of Ustilaginomycotina fungi (data not shown). We hypothesized that this extension could pose heterologous production issues and found that, indeed, upon its deletion, the production of the catalytic domain (cd) of UmGH16_1-A, henceforth called UmGH16_1-A_cd, was increased by ca. 40-fold (ca. 7 mg/liter of culture). We underscore that the role of the C-term extension could prove important in vivo, for instance in regulation processes or substrate anchoring. However, the role of such extensions in CAZymes remains nearly unaddressed (30) and clearly deserves further attention. Wondering about the possibility of maturation of UmGH16_1-A by proteolytic cleavage, we looked at predicted cleavage sites using the PROSPER online tool (31). We found one metalloprotease and one serine protease cleavage site near the beginning of the C-term extension (Fig. S4A). The latter could be a plausible site since 4 serine peptidases are found in the TOP100 secreted proteins (Fig. S4B).

Screening of the substrate specificity of *Um*GH16_1-A_cd showed the release of oligosaccharides from different β -1,3-glucans, with the largest amounts of products detected for laminarin, followed by yeast β -glucan and then pachyman (Fig. 4A and B). In order to further understand this substrate preference, we carried out linkage analysis of those three substrates (Fig. 4C, Fig. S5A and B). We confirm that Pachyman is a linear β -1,3-glucan and show that the structure of laminarin and yeast β -glucans is somewhat different from the suppliers' descriptions. Indeed, laminarin appears to be a linear β -1,3-glucan with low frequency (ca. 3%) of single glucose units branched in β -1,6 (Fig. 4C). Interestingly, yeast β -glucan appears to have a similarly low substitution frequency, but longer branches (on average four β -1,6-linked glucose units on each branch).

We underscore that HPAEC-PAD and LC-MS analyses did not show the release of β -1,6/1,3-gluco-oligosaccharides. This is in contrast with the product profile of its ortholog *Pc*GH16_1 acting on laminarin, for which the release of G6G3G3G has been shown by NMR (32). Yet, the presence of short β -1,6 substitutions on the β -1,3-glucan main chain appears to significantly increase the activity of *Um*GH16_1-A_cd (Fig. 4A and B). We propose that the presence of those substitutions may either help the enzyme to bind to the targeted β -1,3-chain and/or alter the physicochemical properties of the polymer improving reactivity with the enzyme.

Furthermore, one can observe with both HPAEC-PAD (Fig. 4A) and MALDI-ToF MS (Fig. S6) analyses the release from laminarin by *Um*GH16_1-A_cd of a series of secondary peaks adjacent to the Lam series. LC-MS analysis of these peaks showed that they correspond to C1-reduced cello-oligosaccharides, already present in the initial laminarin suspension (Fig. S7). This modification most probably occurred during laminarin extraction/preparation by the supplier.

Control experiments showed no (for DP2-DP4) or extremely low (for DP5-DP6) activity on β -1,3-gluco-oligosaccharides (Fig. S8A&B). This observation suggests that the enzyme requires more than six carbohydrate units to be active. Furthermore, the concomitant release from laminarin of oligosaccharides with both low and high DP by UmGH16_1-A_cd, even at very early time points (Fig. S9), suggests that the enzyme would act with both *exo* and *endo* modes. However, given the putative low degree of polymerization of laminarin (25 units on average) (33), we could also expect a strict *endo*-active enzyme to release simultaneously short and medium length oligosaccharides from this polysaccharide upon cleavage in a random fashion. This question would deserve further investigations. Additional control experiments showed that no activity

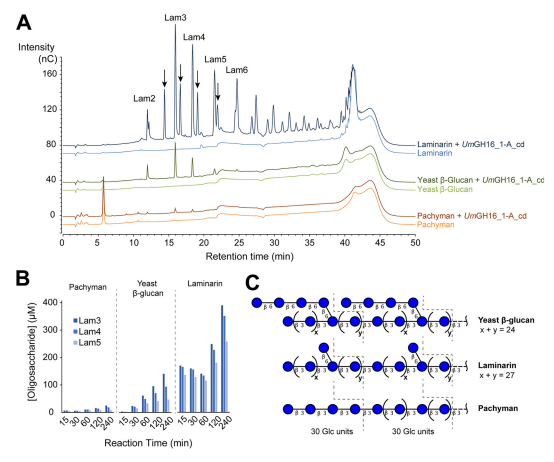


FIG 4 Activity of *Um*GH16_1-A_cd on β -1,3 glucans. (A) The graphs show HPAEC-PAD chromatograms of reaction products released from laminarin, yeast β -glucan and pachyman (10 mg.mL⁻¹ final concentration) by *Um*GH16_1-A_cd (10 nM). Black arrows indicate reduced β -1,3-gluco-oligosaccharides (see Fig. S7). All reaction mixtures were incubated during 4 h, in citrate phosphate buffer (50 mM, pH 5.5), in a thermomixer (30°C, 1,000 rpm). All experiments were carried out in triplicate but for the sake of clarity, only one replicate is shown. See Fig. S8 for additional control experiments. (B) Time-course release of Lam3-Lam5 oligosaccharides from laminarin, yeast β -glucan and pachyman (same reaction conditions as in panel A; n = 1). (C) Proposed chemical structure of the three tested polymers on the basis of carbohydrate linkage analysis (see Fig. S5 for more details). β 3 and β 6 represent β -(1,3) and β -(1,6) linkages, respectively.

could be detected on any of the tested polysaccharides with β -1,4 linkages (cellulose and α -chitin) or mixed β -1,3/1,4 linkages (lichenan) (Fig. S8C).

Overall, we conclude that *Um*GH16_1-A_cd is a β -1,3-glucanase with a marked preference for β -1,3-glucans substituted with single β -1,6-branched glucose units.

*Um*AA3_2-A is a dehydrogenase active on β-1,3 and β-1,6-gluco-oligosaccharides. *Um*AA3_2-A was heterologously produced with success in *P. pastoris* and purified to homogeneity (ca. 5 mg/liter of culture). We underscore that SDS-PAGE analysis was not trivial, most likely due to excessive protein instability under denaturing conditions (main band with apparent MW ~50 kDa), proteolysis (band at ~25 kDa) and oligomerization mediated by intermolecular disulfide-bonds (Fig. S10A). During the preparation of the manuscript, Wijayanti et al. reported the production and preliminary characterization of several AA3_2s, including *Um*AA3_2-A (called there *Um*GDHIII; [34]), for which they observed the same atypical, cleavage and polymerization pattern under SDS-PAGE denaturing conditions. We carried out size exclusion chromatography and showed a unique, monodisperse peak, corresponding to an estimated monomeric size of 48 kDa (Fig. S10B).

*Um*AA3_2-A substrate specificity was assessed (Fig. 5A) by measuring dehydrogenase activity (with DCIP as an electron acceptor) on various (oligo)saccharides (Fig. S11). This analysis revealed that β -D-Glcp-(1,6)-D-Glc (G6G, gentiobiose), followed by β -D-Glcp-(1,3)-D-Glc (G3G; laminaribiose) and G3G3G (laminaritriose), were the preferred

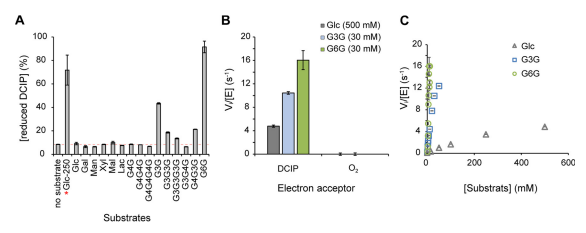


FIG 5 Activity of *Um*AA3_2-A. (A) Substrate specificity screening monitored as the reduction of DCIP (400 μ M) by *Um*AA3_2-A (14 nM) in the presence of various substrates (2.5 mM for all, 250 mM when marked with a red star) after 3 h of incubation (see Fig. S11 for substrate nomenclature). (B) Dehydrogenase *versus* oxidase activity was measured, as, respectively, the reduction of DCIP (400 μ M) *versus* O₂ (250 μ M) by *Um*AA3_2-A (110 nM) in the presence of Glucose (500 mM), G3G (30 mM) or G6G (30 mM). (C) [Glucose], [G3G] and [G6G]-dependency of *Um*AA3_2 initial rate. All reactions were carried out in citrate-phosphate buffer (50 mM, pH 5.5), at 30°C. Data are presented as average values (n = 3, independent biological replicates) and error bars show s.d.

substrates. These results are in good agreement with those reported by Cerutti et al. (27) and Wijayanti et al. (34). As in those two studies, no activity on either cellobiose (G4G), cellotriose (G4G4G) or the trisaccharide G3G4G could be measured, whereas some activity was retained on the mixed trisaccharide G4G3G containing a β -(1, 3) glycosidic bond between the reducing end glucose unit and the adjacent unit (Fig. 5A). We determined an optimum pH of 5.5 for the dehydrogenase activity, on glucose (Fig. S12A), G3G (Fig. S12B) and G6G (Fig. S12C). Of note, we also probed the ability of *Um*AA3_2-A to reduce O₂ in the absence of organic electron acceptor by measuring the production of H₂O₂ using either G3G, G6G or glucose as electron donor (Fig. 5B). No oxidase activity could be detected, confirming thereby the strict dehydrogenase nature of the enzyme as previously observed in hereinbefore mentioned studies (26, 27, 34).

We then determined the dehydrogenase kinetic parameters of *Um*AA3_2-A, at optimum pH, for glucose, G3G and G6G (Fig. 5C). For these three substrates, substrate saturation could hardly be reached within a reasonable concentration range. Yet, determination of the initial slopes on the V_i = f(S) plot allowed us to approximate the catalytic efficiencies, yielding values of k_{cat}/K_M of 697, 636, and 18 M⁻¹.s⁻¹ for G6G, G3G, and glucose, respectively (see Table S2 for the full set of approximate kinetic parameters).

For comparison purposes, Cerutti et al. reported for *Pc*ODH k_{cat}/K_{M} values of 777 $M^{-1}.s^{-1}$ and 47 $M^{-1}.s^{-1}$ for G3G and glucose, respectively (27). Wijayanti at al. reported similar kinetic parameters for *Um*AA3_2-A as those we present here (see Table S2). For both *Pc*ODH and *Um*AA3_2-A, the presence of a β -1,3 linkage between the reducing and first nonreducing D-Glc units is thus clearly crucial for the activity.

Here, in addition to the commonly used DCIP-based assay, we used LC and MS methods to characterize the product profile of this enzyme. Using mass spectrometry, we first verified that the reaction catalyzed by *Um*AA3_2-A on G3G, G3G3G and G6G yielded oxidized species, as shown by the presence of simple and double sodium adducts of M + 16 species (Fig. S13). To establish whether these species are geminal-diols (i.e., oxidized on nonreducing end carbon) or aldonic acids (i.e., oxidized on the C1 carbon of substrate reducing end) we carried out UPLC-MS using positive and negative ionization mode (Fig. S14-S16). For conversion reactions of G3G (Fig. S14), G3G3G (Fig. S15), and G6G (Fig. S16), one oxidized species was observed, in negative mode only, which is indicative of the formation of the corresponding aldonic acid.

Altogether, these results are consistent with a two-electron oxidation of the

oligosaccharide at the C1 carbon, yielding a lactone, which is known to undergo a spontaneous hydrolysis leading to aldonic acids as end products.

To gain insight into the structure-function relationship underlying UmAA3_2-A mode of action, we compared a homology model (generated using AlphaFold 2) to the X-ray structure of the PcODH-G3G complex (PDB id: 6XUV; [27]) (Fig. S17A). This analysis shows very similar structure and active site architecture between UmAA3_2-A and PcODH, with a wider active site entrance than the one observed for AnGOX and AfGDH that accommodate monosaccharides (Fig. S17B). In particular, Y64, F416 and W430, as well as F421 from the flexible "substrate binding loop" described for PcODH, are held in optimal position to bind the reducing and nonreducing end, respectively, of G3G, by CH- π interactions. This observation correlates with better dehydrogenase activity detected on β -1,3-oligosaccharides than on glucose. In line with this, out of three residues involved in hydrogen bonding to glucose hydroxyl groups in AfGDH and AnGOX, and lacking in PcODH, only one residue (Asp446) is conserved in UmAA3_2-A (Fig. S18). Remarkably, this residue interacts with glucose O4 hydroxyl in AfGDH (Glu435) and seems to be strictly conserved in type I GDH and in GOx enzymes (Asp424 in AnGOx), as well as in most ODH and ODH-like enzymes, with a few exceptions, such as PcODH (Val428) (27). This comparison also revealed the presence of an additional loop (residues 173 to 192) in UmAA3_2-A (Fig. S18), that seems conserved in ODH-like proteins as previously described (27). The role of these structural differences in potential biocatalytic differences and biological functions remains to be investigated.

UmGH16_1-A and UmAA3_2-A interplay on fungal β -1,3 glucans. As shown above, UmGH16_1-A and UmAA3_2-A are active on β -1,3/ β -1,6-glucans and oligosaccharides thereof, respectively. Therefore, we set off to investigate the interplay between both enzymes. Using a fraction of U. maydis fungal cell wall (UmFCW) enriched in β -1,6/ β -1,3glucans, we could detect the release of β -1,3-gluco-oligosaccharides by the versatile, commercial *Tsp*GH16_3 but could not detect any activity when using *Um*GH16_1-A_cd, despite several attempts (Fig. S19). We suspect that the nature and branch length of substitutions present in UmFCW β -glucans hamper UmGH16_1-A-cd activity, underlining the necessity to finely characterize the FCW fraction. Other upstream glucanases activities could be necessary to unlock UmGH16_1-A_cd access to the substrate. In this perspective, it could be worth testing the recently identified GH51 β -1,3 glucanase (called Erc1) from U. maydis, active on laminarin and laminarihexaose (35). Working with a better characterized glucan polymer (i.e., laminarin), we demonstrated that the in vitro combination of UmGH16_1-A-cd and UmAA3_2-A led to a functional biocatalytic cascade where β -1,3gluco-oligosaccharides released by $UmGH16_1-A-cd$ (DP2 to > DP6) were further oxidized by UmAA3_2-A (Fig. 6 and 7).

To get further insights into the relevance of this potential interplay, we tested two hypotheses. In our first hypothesis, we tested whether product inhibition of the GH16 enzyme by its oligosaccharide products could be alleviated upon their oxidation by the dehydrogenase. A similar scenario has been observed for the cellobiose hydrolase/cellobiose dehydrogenase pair, where cellobiose released from cellulose by the cellobiose dehydrogenase (36). However, here, *Um*GH16_1-A_cd was neither inhibited by G3G (Fig. S20A), nor by G6G (Fig. S20B). Conversely, in our second hypothesis, oxidized oligosaccharides, generated by *Um*AA3_2-A, could be inhibitors of *Um*GH16_1-A. The addition of G3G^{ox} or G6G^{ox} to a reaction of *Um*GH16_1-A_cd on laminarin did not show any significant inhibitory effect (Fig. S21). Thus, our results rule out any product-based regulatory interplay between both enzymes.

DISCUSSION

In the present study, by reanalyzing previously published data (10) in the light of today's knowledge, we have revealed that the secretome of the plant pathogen

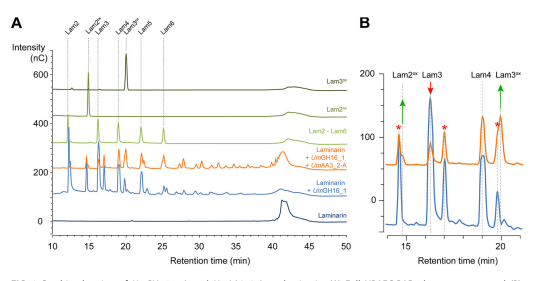


FIG 6 Combined action of *Um*GH16_1-A and *Um*AA3_2-A on laminarin. (A) Full HPAEC-PAD chromatograms and (B) zoom-in view on the 11–21 min region comparing products released from laminarin by *Um*GH16_1 alone (blue line) or in combination with *Um*AA3_2-A (orange line). The red stars indicate peaks of reduced oligosaccharides already present in the laminarin (see main text).

U. maydis grown on corn bran contains a significant fraction of CAZymes predicted to be active on the FCW, including several hydrolases and carbohydrate oxidases that may act in concert. On the basis of phylogenetic analyses and after interrogating published transcriptomic studies, we have selected and biochemically characterized *Um*GH16_1-A and *Um*AA3_2-A, which proved to be active on β -1,3/1,6-glucans and oligosaccharides thereof, respectively.

Together with previously published work (27, 34), we show that both *Pc*ODH and *Um*AA3_2-A appear to form an evolutionarily distinct subclade within the AA3_2 GOx/GDH clade, associated with a new substrate specificity. Enzyme kinetics tell us that G6G and G3G are more than one order of magnitude better substrates than glucose. Yet, the measured rates still remain low compared to other AA3 oxidoreductases, indicating that the biologically relevant substrate may be more complex, potentially harboring some ramifications. Beyond the structure of the natural substrate, there are also open questions regarding the fate of the electrons extracted from G3G or G6G by *Um*AA3_2. Analogous enzymatic systems active on β -(1,4)-glucans (cellulose and cellooligosaccharides) have shown that the extracted reducing power could feed down-stream enzymatic activities such as LPMOS (37–39). Provided it exists, a similar cascade remains to be found for β -(1,3)-glucans.

Furthermore, while apparent activity on β -1,3/1,6 glucans led us to focus on FCW, it is worth mentioning that β -1,3 glucans can also be found in the cell walls of cereals (including maize) as mixed-linkage $(1\rightarrow 3),(1\rightarrow 4)-\beta$ -D-glucans, mostly concentrated in the endosperm (40, 41). Although we cannot rule out these PCW components as potential target for the enzymes studied herein, several facts rather support the hypothesis of FCW-directed activities: (i) UmGH16_1-A being active on β -1,3/1,6-glucans but not on lichenan (a 1,3/1,4-linked polymer), and UmAA3_2-A being most active on G6G, together with (ii) β -1,3/1,6-glucans being mainly present in FCWs (as well as in some seaweeds and bacteria) (42), and virtually absent from plants, and (iii) their coexpression during early stages of U. maydis infection cycle and repression at later stages during plant infection cycle by U. maydis. While additional accessory activities still need to be uncovered to fully understand the putative biocatalytic cascade at play (Fig. 7), we believe UmGH16_1-A and UmAA3_2-A could play a role in FCW remodeling, during which released fungal oligosaccharides, known to act as elicitors of plant immunity (43), may be oxidized to evade the host immune response. Of note, a similar hypothesis has been recently proposed for another U. maydis β -1,3 glucanase (Erc1)

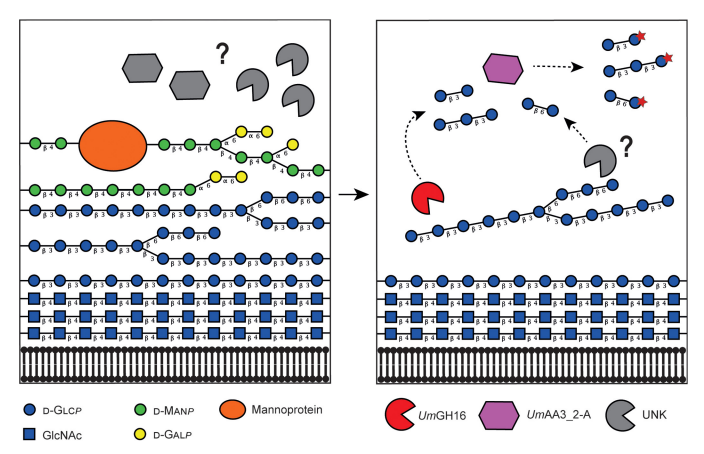


FIG 7 Proposed reaction scheme illustrating the combined action of *Um*GH16_1-A and *Um*AA3_2-A on FCW. Putative enzymatic activities secreted by *Ustilago maydis* to target its own cell wall. The legend key and the glycosidic linkage between each carbohydrate unit are indicated in the figure (for instance, " β 4" indicates a β -1,4 linkage). In the left-hand side panel, hypothetical enzymatic activities (in gray) degrade the galactomannan and mannoproteins, allowing access to the lower layer of β -1,3/ β -1,6-glucans. In the right-hand side panel, the uncovered glucans can act as potential substrate for *Um*GH16_1-A (shown in red) and other hypothetical hydrolytic activities (in gray), releasing β -1,3 and β -1,6-oligosaccharides oxidizable by *Um*AA3_2-A (in purple). Another scenario where *Um*GH16_1-A and *Um*AA3_2-A would access their substrate by simple diffusion through the FCW, rather than after extracellular secretion, is also possible. In such scenario, extracellular degradation of the first layer of FCW would not be required. For now, it is impossible to settle on the actual trajectory of these enzymes, but, for the sake of clarity, the first layer of mannoprotein and galactomannan is not shown in the right-hand panel.

whose activity toward laminarihexaose has been shown to suppress laminarihexaoseinduced ROS burst in plant leaves (35). All in all, it is clear that the fate and role of FCW/PCW-derived oxidized products released by fungal oxidative enzymes is an emerging matter of utmost importance (38, 44–47).

Conclusions. Recent-omics studies and biochemical characterization have enriched our knowledge over the plethora of activities that constitute the fungal enzymatic arsenal. The various questions that emerge from our study underscore the need for a deeper integration of enzymology, cellular biology and microbial ecology to better understand the genuine activities, biological role, and potential biotechnological interest of CAZymes, and most notably of oligosaccharide oxidases. We believe that the diversity and roles of FCW-active enzymes only starts to unfold, promising important discoveries to be made in the coming years.

MATERIALS AND METHODS

Materials. Most chemicals were purchased from Sigma-Aldrich. Oligosaccharide substrates and polysaccharides (Yeast β -glucan, reference P-BGYST, batch number: 180808a/pachyman, reference P-PACHY, batch number: 10301/Lichenan reference P-LICHN, batch number: 70901b), as well as the *endo*-1,3- β -D-glucanase *Tsp*GH16 (reference E-LAMSE), were purchased from Megazyme (Wicklow, Ireland). Laminarin was purchased from Merk (reference L9634).

Enzymes cloning, production, and purification. The gene encoding *Um*AA3_2-A (Uniprot ID A0A0D1DW37, Gene ID UMAG_03551) was PCR amplified from the genome of *Ustilago maydis* BRFM 1093 strain, with the following primers containing EcoRI and Xbal restriction sites (underlined):

Forward: <u>GAATTC</u>GCCATCGTCACAGATG

Reverse: TCTAGACCCCTGGCGAGAATGGTGT

The amplicon and TOPO vector were subsequently used to cotransform E. coli DH5 α competent cells according to the TOPO Cloning reaction protocol (Invitrogen). Positive transformants were selected on LBagar-ampicillin (50 μ g.mL⁻¹). Plasmidic DNA was extracted, purified and the expected size was verified by agarose electrophoresis. Then, the pPICZ α A vector and TOPO-UmAA3_2-A vectors were digested with EcoRI and Xbal, gel-purified and a ligation of linearized pPICZ α A and UmAA3_2-A insert was carried out. The ligation product was then transformed in *E.coli* DH5 α for plasmid production. After plasmid extraction the final construct pPICZ α A-UmAA3_2-A was sequenced before transformation in P. pastoris. The intron-free sequence of the gene coding for UmGH16_1-A (Uniprot ID A0A0D1E047, Gene ID UMAG_02134) was synthesized after codon optimization for expression in *P. pastoris* and inserted into a modified pPICZ α C vector using Xhol* and Notl restriction sites in frame with the α secretion factor at N-terminus (i.e., without native signal peptide) and with a (His)₆-tag at the C terminus (without c-myc epitope) (Genewiz, Leipzig, Germany). For both enzymes, transformation of competent P. pastoris X33 was performed by electroporation with Pmel-linearized pPICZ α C recombinant plasmids. P. pastoris strain X33 and the pPICZ α C vector are components of the P. pastoris Easy Select Expression System (Invitrogen), all media and protocols are described in the manufacturer's manual (Invitrogen). Zeocin-resistant P. pastoris transformants were screened for protein production as described by Haon et al. (48). The best-producing transformants were grown in 2 L flasks. The proteins of interest were expressed and secreted upon methanol induction and purified from the supernatant by IMAC, according to a previously described protocol (49).

Enzyme concentrations were determined by measuring UV absorbance at 280 nm using a Nanodrop ND-200 spectrophotometer (Thermo Fisher Scientific, MA, USA) and extinction coefficient determined with ProtParam (Expasy) for $UmGH16_{-1}$ -A ($\varepsilon^{280} = 83,350 \text{ M}^{-1}.\text{cm}^{-1}$) and $UmAA3_{-2}$ -A ($\varepsilon^{280} = 85,830 \text{ M}^{-1}.\text{cm}^{-1}$).

Phylogenetic analyses. To build the phylogenetic tree of AA3s, we used 57 sequences of experimentally characterized enzymes of fungal origin (ascomycetes and basidiomycetes) belonging to the four different subfamilies (i.e., AA3_1 to AA3_4), together with the three AA3_2s from *U. maydis* secretome (JGI ID 10518, 10841 and 11351). For the GH16s tree, 264 sequences (including *Um*GH16-A) representing the 27 subfamilies described in the work of Viborg et al., were provided by the CAZy team (AFMB, Marseille), (19). Of note, the variable C-terminal regions of the GH16s sequences were cut using BioEdit (50) in order to keep the catalytic domain only. Both AA3s and GH16s sequences batches were aligned using MAFFT-DASH (L-INS-i method) (51), which include structural data input. The resulting multiple sequence alignments were used to infer the phylogenetic trees via the MAFFT online platform for AA3s and the RAxML software for GH16s. A neighbor-joining method (NJ, on the basis of conserved sites) or a Maximum Likelihood method (ML) was used for AA3s and GH16s, respectively. In both cases, the Whelan and Goldman (WAG) amino acid substitution model was selected (52). Branch support was calculated by 500 (for the AA3s tree, values displayed in percent on the tree) or 100 (for the GH16s tree) bootstrap repetitions. The trees were visualized in iTOL (53) and edited in Illustrator.

Fungal cell wall extraction. The U. maydis strain 521, which was provided by the CIRM-CF collection (strain BRFM1093) (54) was grown in 100 mL of yeast extract (10 g.L⁻¹)/BactoPeptone (20 g.L⁻¹)/ Dextrose (20 g.L⁻¹) (YPD medium) for 48 h at 28°C in 250 mL-baffled Erlenmeyer flask under orbital agitation (150 rpm). Cells were then harvested and washed once in H₂O by centrifugation (1,500 g, 10 min), counted and stored at 10^7 cells/mL in 20% glycerol at -80° C as a working cell bank for long term preservation. In order to produce material for sequential extraction 20 250 mL-baffled Erlenmeyer flasks containing 100 mL of YPD medium were inoculated at 10⁵ cells/mL with U. maydis cells from the frozen cell bank and incubated for 24h at 28°C under orbital agitation (150 rpm). Cells were then harvested and washed three times with H₂O by centrifugation cycles (8,000 g, 4°C, 20 min). The washed cell pellet was lyophilized (ca. 7 g). Five grams of this material was resuspended in 500 mL H₂O, homogenized using utra-turax (2 min, 13,500 rpm) and boiled for 4 h. After a centrifugation (8,000 g, 20 min), the supernatant was filtered on 0.7 μ m glass microfibers and stored at 4°C. The pellet was resuspended in 500 mL of 1.25 M NaOH solution for 4 h at 60°C. After centrifugation (8,000 g, 20 min), the supernatant was filtered on 0.7 μ m glass microfibers and stored at 4°C. Polysaccharides extracted in H₂O and NaOH were further precipitated in 50% ethanol at 4°C for 16 h under stirring. Precipitated polysaccharides were washed five times with 50 mL of 50% ethanol and lyophilized. Alkali insoluble material was washed in H₂O until pH reached 7 and kept in suspension to enable pipetting.

Dehydrogenase activity assay. The dehydrogenase activity was monitored by measuring spectrophotometrically the decolorization upon reduction of the cosubstrate 2,6-dichlorophenolindophenol (DCIP), at 520 nm ($\varepsilon_{s20} = 6,800 \text{ M}^{-1}\text{.cm}^{-1}$). Most experiments were carried out at the optimal pH value of 5.5. Substrate specificity was assessed by screening 14 different substrates. Unless stated otherwise, reactions (100 μ L final reaction volume) were carried out in 96-wells transparent microtiter plates (Corning Costar, Corning, NY, USA) and contained *Um*AA3_2-A (110 nM) and DCIP (0.4 mM) in citratephosphate (50 mM, pH 5.5). The mixtures were incubated during 2 min at 30°C before the reaction was initiated by the addition of substrate (250 mM final for glucose and 2.5 mM for other substrates, including glucose). The absorbance was monitored over 10 min using a Tecan Infinite M200 (Tecan, Switzerland) plate reader. All reactions were carried out in triplicate. Initial rates, determined at various substrate concentrations, were used to calculate the kinetic parameters according to the standard Michaelis-Menten equation for G3G or using a modified model accounting for excess-substrate inhibition in the case of glucose. SigmaPlot 12.0 was used to fit the experimental data.

Glycoside hydrolase activity assay. The activity of *Um*GH16_1-A was evaluated by monitoring the release of gluco-oligosaccharides from various glucans by high-performance anion-exchange chromatography (HPAEC) coupled to pulsed amperometric detection (PAD) (see below). Unless stated otherwise, reactions

(500 μ L final reaction volume) were carried out in 2 mL Eppendorf tubes and contained the substrate (10 g.L⁻¹) in citrate-phosphate buffer (50 mM, pH 5.5). The mixtures were incubated during 2 min at 30°C in a Thermomixer (1,000 rpm) and the reactions were initiated by the addition of *Um*GH16_1-A_cd (10 nM). For each time point (15 min to 4 h), one sample (500 μ L) is sacrificed by boiling for 10 min, centrifuged (12,000 g, 2 min, 4°C), and diluted 10-fold in milliQ H₂O before injection on the HPAEC column. Reactions using FCW extract were incubated overnight (16 to 18 h) and the supernatant was injected without prior dilution.

Reactions combining UmGH16_1-A_cd and UmAA3_2-A were carried out under similar conditions as described above with the addition of UmAA3_2-A (1 μ M) and DCIP (400 μ M).

HPAEC-PAD analyses. The detection of soluble oligosaccharides is performed using HPAEC-PAD (DIONEX ICS6000 system, Thermo Fisher Scientific, Waltham, MA, USA). The system is equipped with a CarboPac-PA1 guard column (2 × 50 mm) and a CarboPac-PA1 column (2 × 250 mm) kept at 30°C. Elution was carried out at a flow rate of 0.25 mL.min⁻¹ and 25 μ L of sample was injected. The eluents used were 100 mM NaOH (eluent A) and NaAc (1 M) in 100 mM NaOH (eluent B). The initial conditions were set to 100% eluent A, and the following gradient was applied: 0 to 10 min, 0 to 10% B;10 to 35 min, 10 to 35% B (linear gradient);35 to 40 min, 30 to 100% B (curve 6);40 to 41 min, 100 to 0% B;41 to 50 min, 100% A. Integration was performed using the Chromeleon 7.2.10 software based on commercially available standards: laminari-oligosaccharides and G6G. G3G^{ox} and G6G^{ox} standards were prepared by incubating, respectively, G3G and G6G (1 mM each) with $UmAA3_2-A$ (1 μ M) and DCIP (2 mM) in citrate phosphate buffer (50 mM, pH 5.5), in a thermomixer (30°C, 1,000 rpm) during 24 h.

Linkage analyses. Polysaccharides (laminarin, pachyman and yeast β -glucans) were prepared at a concentration of 1 mg.mL $^{-1}$ in dimethyl sulfoxide (DMSO) and left overnight at 60°C under constant agitation. Methylation (method adapted from [55]) was performed with 500 μ L of each sample by adding in the following order: 500 μ L of NaOH-DMSO reagent and sonicate the tubes during 10 min, 100 μ L of methyl iodide and sonicate the tubes during 10 min (twice) and 200 μ L of methyl iodide and sonicate the tubes during 5 min. The reaction was stopped by the addition of H₂O (2 mL) and the methylated products were extracted with chloroform (500 μ L). The solutions were vigorously vortexed before a brief centrifugation, which allowed a strict separation of two phases. The aqueous supernatant phase was removed by aspiration. The organic phase was washed three times with H₂O (2 mL). Methylated carbohydrates were hydrolyzed with 2 M trifluoroacetic acid in the presence of an internal standard (mvo-inositol) and converted to the corresponding alditol acetates. The partially methylated alditol acetates were analyzed by GC-MS (TRACE-GC-ISQ, Thermo) on a nonpolar thermo scientific TraceGOLD TG-1MS GC Column (30 m x 0.25 mm x 0.25 μ m), carrier gas H₂ at 1.5 mL.min⁻¹. The sample was injected at 240°C and the oven temperature was maintained for 5 min at 60°C and increased up to 315°C (3°C/min), and further maintained at 315°C for 2 min. The gas flow rate was set at 1.5 mL.min⁻¹. The ion source temperature of the electron impact (EI) mass spectrometer was 230°C. Masses were acquired with a scan range from m/z 100 to 500. Identification of partially methylated alditol acetates was based on their retention time and combined with confirmed by mass spectra fragmentation and compared to a home-made library. Quantitative detection was performed at 220°C with a flame ionization detector (FID).

Matrix assisted laser desorption/ionization (MALDI)-time of flight (TOF) analysis. MALDI-TOF-MS spectra were acquired on a Rapiflex TissueTyper mass spectrometer (Bruker Daltonics, Bremen, Germany), equipped with a Smartbeam II Laser (355 nm, 10 kHz) and reflector detection. Samples were diluted in H₂O (100 μ g.mL⁻¹) and directly mixed on a polished steel MALDI target plate with a solution of ionic liquid matrix DMA-DHB (2,5-dihydroxybenzoic acid 100 mg.mL⁻¹ in H₂O/ACN (50:50 vol/vol) with an addition of 0.2% of *N*,*N*-dimethylaniline (56). Spectra were recorded in the *m/z* range 350 to 3200 using FlexControl and processed using FlexAnalysis (Bruker Daltonics, Billerica, MA, USA). Mass spectra were acquired in positive ionization mode.

Ultra high-performance liquid chromatography (UHPLC)-electrospray (ESI)-ion trap (IT) analysis. UHPLC-ESI-IT acquisitions were performed on an amaZon SL 3D ion trap mass spectrometer (Bruker Daltonics, Bremen, Germany) coupled with an Acquity H-Class UHPLC (Waters, Wilmslow, UK). Samples were diluted in a solution of H₂O/ACN (95.5:4.5) at 10 μ g.mL⁻¹. 10 μ L of each sample was injected on an Hypercarb column (100 × 1 mm, particle size 3 μ m, Thermo-Fisher Scientific, Courtaboeuf, France) heated at 80°C with a flow rate settled at 0.165 mL.min⁻¹. A binary gradient was performed. The gradient started with 8 min at 95.5% of A (H₂O) and then ramped linearly to 80% of B (ACN) in 22 min and stayed at 80% of B during 12 min; initial conditions were restored during the last 5 min. The ESI source parameters were the following: capillary voltage: 4.5 kV; nebulizer gas: 7.3 lb/in²; and dry gas: 4 L.min⁻¹ (80°C). Mass spectra were recorded in the *m*/z range 350 to 2,200 in the positive ionization mode. Acquisitions were performed using TrapControl 8.0 and Compass HyStar 4.1 (Bruker Daltonics, Bremen, Germany). Data were processed using Data Analysis 4.4 (Bruker Daltonics, Bremen, Germany).

SUPPLEMENTAL MATERIAL

Supplemental material is available online only. **SUPPLEMENTAL FILE 1**, PDF file, 3.8 MB.

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J.-L.R., M.H., O.T., and S.L.G. carried out the enzymology experiments. D.R. and S.L.G. carried out mass spectrometry analyses. J.-L.R., D.R., S.L.G., G.S., J.-G.B., and B.B. interpreted the data. J.-G.B. and B.B. conceptualized the study, designed the experiments, and supervised the work. B.B. wrote the first draft and finalized the manuscript. All authors contributed to the writing of the manuscript, with main contributions from J.-L.R. and J.-G.B. All authors reviewed and approved the final version of the manuscript.

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