XAX1 from glycosyltransferase family 61 mediates xylosyltransfer to rice xylan

Dawn Chiniquy, Vaishali Sharma, Alex Schultink, Edward E. Baidoo, Carsten Rautengarten, Kun Cheng, Andrew Carroll, Peter Ulvskov, Jesper Harholt, Jay D. Keasling, Markus Pauly, Henrik V. Scheller, and Pamela C. Ronald

Xylan is the second most abundant polysaccharide on Earth and represents an immense quantity of stored energy for biofuel production. Despite its importance, most of the enzymes that synthesize xylan have yet to be identified. Xylans have a backbone of β-(1→4)-linked xylose residues with substitutions that include α-(1→2)-linked glucuronosyl, 4-O-methyl glucuronosyl, and α-1,2- and α-1,3-arabinofuranosyl residues. The substitutions are structurally diverse and vary by taxonomy, with grass xylan representing a unique component of dietary fiber in cereals (1). The amount of xylan as well as its chemical composition is an important element of bread formulation (2) and varied saccharification, probably reflecting a lower degree of dimeric cross-links. Activity assays with microsomes isolated from tobacco plants transiently expressing XAX1 demonstrated xylosyltransferase activity onto endogenous acceptors. Our results provide insight into grass xylan synthesis and how substitutions may be modified for increased saccharification for biofuel generation.

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Xylans are polysaccharides present in the cell walls of all plants. They comprise the major noncellulosic component of secondary walls of angiosperms. Hence, xylans are quantitatively second only to cellulose in biomass on earth. Xylans are valuable components of human and animal nutrition, constituting a major component of dietary fiber in cereals (1). The amount of xylan as well as its chemical composition is an important element of bread making, influencing dough yield, flavor, and shelf life (2). Xylan is of particular interest for the improvement of feedstocks for the generation of cellulosic biofuels, a currently expensive and inefficient process (3). Xylan inhibits access of cellulases to cellulose and is an additional substrate for cross-linking to lignin. Thus, genetic modification of xylan in crop species would rely on improving our knowledge of xylan biosynthesis.

Xylans play an essential structural role in plant cell walls, presumably through interactions with cellulose microfibrils and other components of the wall (4, 5). Although the organization of these various components in the cell wall is not well understood, the increased cross-linking apparently contributes to the recalcitrance of the cell wall (6, 7). Xylans are structurally diverse between taxonomic groups (4, 8). The most basal form of xylan is a main chain that lacks substitutions as seen in the green alga Caulerpa that has β-1,3-O-xylan in place of cellulose, and the red seaweeds Pulmonariales and Nemalilales that have a mixed linkage β-(1,3,4)-O-xyllose backbone (8). Xylans of embryophytes have a β-(1→4)-linked xylose backbone. Xylans found in dicots are mostly restricted to the secondary cell walls, and hence a main component of wood. Dicot xylans are commonly substituted with α-(1→2)-linked glucuronosyl and 4-O-methyl glucuronosyl residues (1). Xylans in birch, spruce, and Arabidopsis have been found to contain the reducing end oligosaccharide β-D-Xylose(1→3)-α-D-Rhap(1→2)-α-D-GalpA(1→4)-D-Xylose (9–11) which, interestingly, has not been found in the xylan of grasses. Comelinid monocot xylans are unique from those of dicots and other monocots. This group, which includes the grasses, contains xylan as the main noncellulosic component in both the primary and secondary cell walls (12). These xylans have very little glucuronosyl residues, but are mostly substituted with α-1,2 and α-1,3 arabinosyl residues. A unique feature of grass xylans is the ferulate and coumarate esters linked to the O-5 of some of the α-1,3 arabinosyl residues. These ferulate esters mediate intra- and intermolecular cross-linking, possibly increasing the strength of the cell wall (7). A common feature of grass xylans is the further substitution of the feruloyl-α-arabinofuranosyl residues with β-D-xyllose linked to O-2 of β-D-arabinose (13–17). Some of these xylosyl residues can be further substituted with β-(1→4)-α-galactosyl residues (18). Among these ferulate-containing xylan sidechain variants, 2-O-β-D-xylo-pyranosyl-(5-O-feruloyl)-α-arabinose has been found to be the most abundant, and in some cases as abundant as 5-O-feruloyl-α-arabinofuranose (13, 15, 19). During the past 6 y, there has been significant progress made toward identifying the glycosyltransferases (GTs) that synthesize xylan. The irregular xylan (irx) mutants, named for their collapsed xylem vessels due to their secondary cell wall deficiency, have been instrumental in elucidating the mechanisms of xylan biosynthesis in Arabidopsis thaliana (Arabidopsis). IRX9/IRX9L and IRX14/IRX14L from the GT43 family and IRX10/IRX10L from GT47 are thought to be responsible for elongation of the xylan backbone (20–25). IRX7 (FRAX8)/IRX7L (F8H) (from GT47), IRX8 (GAUT12) (from GT8), and PARVUS (from GT8) may be responsible for synthesizing the oligosaccharide found at the reducing end of some
dicot and conifer xylans (4, 21, 26, 27). GUX1 and GUX2 (from GT8) are thought to be responsible for adding both glucuronosyl and 4-O-methylglucuronosyl substitutions to the xylan backbone in Arabidopsis (28–30). Recently, rice and wheat GT61 family genes were found to be responsible for α-(1,3)-arabinosyl substitution on xylan (31). Thus, far, the enzymes that add the β-(1,2)-xylosyl residues to xylan have not been identified.

Here, we report the characterization of a mutant xax1 in a rice GT61 family gene. Our results strongly indicate that the corresponding protein possesses β-1,2-xylosyl transferase activity, transferring xylose from UDP-xylose onto xylan. Mutant xax1 rice plants have a dwarfed phenotype and the leaves are deficient in xylose, ferulic acid, and coumaric acid, and have increased saccharification efficiency. These findings add further to our understanding of xylan biosynthesis and the mechanisms of ferulic acid addition to xylan and have important implications for biofuel production.

**Results**

*XAX1* Is a Previously Uncharacterized Xylan Biosynthesis Gene in Rice.

To discover genes important for grass xylan biosynthesis, we conducted a reverse genetics screen of 14 genes with insertional rice mutants that are highly expressed members of the GT61 family (Tables S1 and S2), a family previously identified through bioinformatic approaches as being expanded in grasses and containing grass-specific subgroups (29, 32, 33). Our screen prioritized lines with segregating phenotypes that had altered sugar composition, and from this screen we identified a mutant with a mutation in the third exon that knocked out Os02g22380 (Fig. S1), a gene member that is part of a grass-specific clade in the GT61 family (Fig. S2, clade C.IV). We identified this gene as a putative xylosyltransferase, based on its dwarfed phenotype and xylose deficiency in cell wall alcohol insoluble residue (AIR) in young rice leaves (Fig. 1A and B). Whereas the majority of all xylose in rice is derived from xylan, we confirmed that the deficiency was seen in the xylan-rich fractions (1 M KOH and 4 M KOH) of sequentially extracted cell wall material (Fig. S3 F and G). Based on this biochemical phenotype and subsequent data described below, we named the mutant *xylosyl arabinosyl substitution of xylan 1* (*xax1*).

To gain further insight into the nature of the xylose deficiency, AIR was subjected to partial hydrolysis in 0.1 M trifluoroacetic acid (TFA). This treatment preferentially cleaves arabinofuranosyl residues, leaving the disaccharide relatively more susceptible to hydrolysis than the main chain xylosyl residues (13, 34). A comparison of partial versus complete hydrolysis of AIR showed a stronger decrease in xylose (7.0%) than with total hydrolysis (2.6%), which indicates that the xylose deficiency may be a substitution rather than a xylan chain deficiency (Fig. S3 A and B). Extraction of saponified AIR revealed a strong decrease in both ferulic acid (59%) and coumaric acid (44%) (Fig. 1C), aromatic compounds known to be attached to arabinosyl residues in xylans that are substituted with xylosyl residues (13). The 4 M KOH extract of depectinated AIR was further characterized by enzymatic fingerprinting with M6 endoxylanase (35). The released oligosaccharides were separated and quantified by high performance anion exchange chromatography with electrochemical detection (HPAEC-PAD) (Fig. 2A). Notably, *xax1* showed an absence of one peak eluting at about 22 min, whereas the four other major peaks were found in both wild type and mutant. The peak, which was only detectable in wild type, “peak 1,” was isolated from and

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*Fig. 1. XAX1 T-DNA insertional rice mutant. (A) Dwarfed phenotype of xax1 plants at 5 wk. (B) Monosaccharide composition of total plant cell wall preparations and acid-catalyzed depolymerization of carbohydrate-free residues. (D) Saccharification of cell wall after 120 °C pretreatment and cellulase and hemicellulase enzymatic digestion; quantified by DNS. Error bars represent SD with at least three biological replicates. *Significantly different from WT, t test: P < 0.05.*
is primarily expressed in younger tissue, including seedlings with a Table S4 and AraXyl β
and Fig. S5 October 16, 2012 OH showed a 56% decrease in arabinofuranose and | no. 42 14
| 35 of arabinofuranose 7
residue from the nonreducing end (Fig. 2 hydrolyzes terminal α acceptors in the presence of UDP-
C-xylose. Peak activity as determined by NMR spectroscopy.

To con
Absence of Xylose Substitution Improves Saccharification. Cell walls from the xax1 mutant exhibited a 62% increase in the total sugars released compared with wild type after a 24-h treatment with an enzyme mixture (Ctec2; Novozymes) that contained cellulase, β-glucosidase, and hemicellulases (Fig. 1D). The cellulose content of the mutant was the same as wild type (Fig. S4A), which indicates that the increased extractability is not due to increased carbohydrate content in the mutant. Additionally, CDTA and Na2CO3 pectin-rich fractions, which contained ~2.5% of the xylose content in wild type (Fig. S3H), showed that the xylose was more easily extracted in the mutant, also indicating a more easily extractable hemicellulose (Fig. S3 D and E). The improved saccharification may be due to a decrease in the diferulic cross-links (45–47) or because a more easily extracted xylan allows cellulase enzymes better access to cellulose (48).

Discussion

Evolution of a Grass-Specific Xylan Substitution. XAX1 is a member of the xylosyltransferase family GT61, and we have shown it to be essential for the grass specific β-(1,2) xylosyl substitution of α-(1,3) arabinosyl residues, a disaccharide substitution on the xylan chain that is unique to grasses. The GT61 family was recently identified through bioinformatic analysis to be highly grass diverged (32, 33, 49). Enzymes are grouped into the GT61 family on the basis of a transmembrane domain and a DUF563 protein sequence [carbohydrate-active enzyme (CAzy) database] (50) (Fig. S1B). We have found a single DUF563 sequence in the brown algae Ectocarpus siliculosus, which suggests that this family originated around the time brown algae evolved or before; we have used this brown algae sequence to root our GT61 tree. There are at least three distinct clades identifiable in the GT61 phylogenetic tree (Fig. S2, clades A, B, and C) (49). Clades A and B, which both include moss genes, are represented relatively evenly between grasses and dicots. Clade B genes have diverged far less than other genes in the family, and are generally represented by only a single sequence per genome (At5G55500 and Os08g39380 in Arabidopsis and rice, respectively).

One member of this clade, At5g55500, has been previously characterized to have β-(1,2)-xylosyltransferase activity, transferring xylose to N-linked glycans (51). There is nothing yet known about clade A, but it does contain five Selaginella representatives. We found Selaginella leaves to have no detectable arabinose on their xylan (Fig. S4E) and propose that this clade does not contain arabinosyl transferases involved in xylan biosynthesis. Most likely clade A has a function similar to that of clade B or an as yet uncharacterized function. However, clade C, which contains Os02g22380 (XAX1), is dramatically expanded in the grasses, which is particularly true for groups III and IV, which have no known dicot orthologs.

The recently characterized rice and wheat α-(1,3)-arabinosyltransferases are part of clade C.I and clade C.II (XAT1, XAT2, TaXAT1, and TaXAT2) (31). Clade C.II contains only the dicot orthologs of Clade C (Fig. S2) (31). Arabinose substitutions on xylan are abundant in grasses, but not unique to them. Whereas O-3 substituted arabinose is characteristic of the grasses, many dicot species have O-2 substituted arabinose (38, 41, 42). In addition, flax and psyllium seed mucilage have been shown to contain α-(1,2)- α-(1,3) disubstituted or α-(1,3) monosubstituted arabinose on xylan, respectively (39, 40). Thus, having dicot orthologs in Clade C.II is not surprising. That XAX1 is located in the grass-unique clade C.IV, appears correct, because the β-Xyl-α-Ara disaccharide substitution is a grass-specific trait in xylan and associated with ferulate esters, which are also unique features of grass xylans.

Role of XAX1 in the Context of Ferulic Acid Esters. The decreased content in feruloyl and coumaroyl esters in the xax1 mutant is interesting and somewhat unexpected. It is not clear how hydroxycinnamates are added to xylan, but it is known to take place in the Golgi, and there is evidence that acyltransferases belonging to the BAHD family are involved (17, 52). The BAHD enzymes are cytoplasmic, suggesting that they do not directly ferulate xylan, but add a feruloyl residue to an intermediate. A likely intermediate that has been proposed is UDP-arabinofuranose (53), which is also synthesized in the cytoplasm—again somewhat surprisingly because the precursor, UDP-arabinopyranose, is largely synthesized inside the Golgi lumen (54). However, if UDP-feruloyl-arabinose were the substrate for feruloylation of xylan, we would not expect to see a decrease in ferulate ester content in xax1. The decreased hydroxycinnamate in the mutant suggests that ferulic acid is added subsequently to the addition of xylose to the unferulylated arabinose residues. In that case, the intermediate for feruloylation would be another as yet unidentified compound. Alternatively, hydroxycinnamates attached to arabinose in xax1 are more susceptible to being removed by ferulic acid esterases. If the latter is the case, it would suggest a role of β-(1,2)-xylosyl substitutions on grass xylans in stabilizing the ferulic acid esters.

Rice Xylan Lacking β-(1,2) Xyl Substitution Have Improved Release of Sugars. Mutant xax1 plants exhibit an increased saccharification efficiency (Fig. 1D) and an increased extractability of xylan (Fig. S3 D and E), most likely due to the lower ferulic cross-links (Fig. 1C), but it may also be due to increased solubility of the hemicelluloses and hence possible increased access of cellulosomes to cellulose. There does appear to be a slight increase in arabinose substitutions (Fig. 1B) possibly as a compensatory mechanism due to the xax1 mutation. Based on the linkage analysis (Table S3), there is an increase in disubstituted xylose (2,3,4-Xyl) presumably with α-(1,2) and α-(1,3) arabinose. This increase in arabinose would account for a degree of the increased extractability of xylan, but we propose that, whereas this ~10% increase in arabinose may account for a fraction of the 62% increase in saccharification efficiency, it most likely does not account for the majority of the increased saccharification observed in xax1. We find it more likely that the increased
saccharification is largely a consequence of the 59% decrease in ferulic acid (Fig. 1C).

In conclusion, mutant xax1 plants show a dwarfed phenotype and increased extractability that demonstrates the importance of the β-1,2-xylosyl substitutions on arabinoxylan. These findings also provide us with new insights into xylan synthesis and a unique capability to modify ferulic cross-links by targeting a glycosyltransferase. As ferulic cross-links inhibit sugar release as well as microbial fermentation, this is a potentially important biotechnological tool for grass biofuel feedstocks.

Materials and Methods

Details of plant material, reagents, quantitative RT-PCR, microscopy, construction of GT61 phylogeny, liquid chromatography (LC)/TOF mass spectrometry, NMR, and primers are described in SI Materials and Methods.

AIR Preparation and Analysis. AIR was prepared and analyzed with a HPAEC-PAD as described (21) and in SI Materials and Methods. The extraction of aromatic compounds from AIR was performed as described (55) and in SI Materials and Methods. For the saccharification assay, 5 mg destarched AIR was autoclaved at 120 °C for 1 h. A 500-μL enzyme mixture containing 50 mM citrate buffer pH 6.2, 1.6% (wt/vol) tetracycline, 2 μL CteC2 enzyme mixture (Novozymes), which contains cellulases, β-glucosidases, and hemicellulases, was added and incubated at 50 °C for 24 h with shaking. After enzyme treatment, samples were pelleted, and released sugars in the supernatant were measured using DNS reagent [1% (wt/vol) 3,5-dinitrosalicylic acid, 30% (wt/vol) potassium sodium tartrate, 400 mM NaOH] reading absorbance at 540 nm.

Enzymatic Fingerprinting of Xylan. Xylan was extracted from AIR essentially as previously described (56), following 1,2-Cyclohexylenedi nitro-tetracetic Acid (CDTA) and Na2CO3 extraction, 5 mg AIR was extracted with 4 M KOH, neutralized with glacial acetic acid, diazylated extensively against water, and incubated with 1 unit endoxylanase (Megazyme) in 0.05 M ammonium acetate buffer (pH 4) at 24 °C for 24 h. The reaction mixture was heated at 100 °C for 10 min to stop the reaction and centrifuged before HPAEC analysis. In some experiments, the endoxylanase-treated extract was further incubated with 0.25 units α-L arabinofuranosidase from Aspergillus niger (Megazyme) in 0.05 M sodium acetate buffer (pH 4) at 4 °C for 24 h. The reaction was stopped by boiling at 100 °C for 10 min and centrifuged. The oligosaccharides released after enzymatic digestion were separated by HPAEC on a Dionex DX 600 system equipped with a pulsed amperometric detector (PAD). A Carbopac PA200 column (3× 250 mm) eluted with a gradient of 0–250 mM sodium acetate in 50 mM NaOH over 40 min at 0.4 mL/min was used. Xylose, arabinose, and xylooligosaccharides (xylobiotriose, xylotriose, xylotetraose, xylopentaose, and xylohexaose) (Megazyme) were run as standards.

Xyf Activity Assay. Infiltration of 4-wk-old N. benthamiana leaves was done using Agrobacterium tumefaciens strain C58 (OD = 1), following the method described in ref. 57. For details on the plasmid constructs, see SI Materials and Methods. Leaves were collected 5 d after infiltration, flash frozen, ground in a mortar with pestle in 15 mL buffer (50 mM Hepes pH 7.0, 40 mM MgSO4, 1 mM PMSF, 1% (wt/vol) PVPP, protease inhibitor mixture). The buffer with plant material was filtered through Miracloth mesh and centrifuged at 3,000 × g for 10 min. The supernatant was centrifuged at 50,000 × g for 1 h (Beckman Ultracentrifuge). The pellets containing the microsomes were resuspended in a 50 mM Hepes pH 7.0, 400-mM sucrose buffer and stored at –80 °C. Protein concentration was determined using the Bradford method. Microsomes corresponding to 100 μg protein were incubated with 50 mM Hepes (pH 7.0), 40 mM sucrose, 5 mM MnCl2, 20 mM UDP-[14C]xylose (American Radiolabeled Chemicals) in a total reaction volume of 50 μL. After incubation at 24 °C for various times, the reaction was stopped by adding 70% ice-cold ethanol and 100 μg dextran as a carrier. The products were resuspended and washed in 70% ethanol in a sonicator bath (VWR) at 30 °C for 30 min, vortexed, precipitated at 20 °C for 30 min, and repeated the wash until no counts were found in the wash solution. The pellet was resuspended in water, an equal volume of scintillation fluid was added (National Diagnostics), and the amount of activity was determined using a scintillation counter set to measure 1 μC counts for 2 min (Beckman LS 6300).

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