Structure and Dynamics of *Brachypodium* Primary Cell Wall Polysaccharides from Two-Dimensional $^{13}$C Solid-State Nuclear Magnetic Resonance Spectroscopy

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ABSTRACT: The polysaccharide structure and dynamics in the primary cell wall of the model grass *Brachypodium distachyon* are investigated for the first time using solid-state nuclear magnetic resonance (NMR). While both grass and non-grass cell walls contain cellulose as the main structural scaffold, the former contains xylan with arabinose and glucuronic acid substitutions as the main hemicellulose, with a small amount of xyloglucan (XyG) and pectins, while the latter contains XyG as the main hemicellulose and significant amounts of pectins. We labeled the *Brachypodium* cell wall with $^{13}$C to allow two-dimensional (2D) $^{13}$C correlation NMR experiments under magic-angle spinning. Well-resolved 2D spectra are obtained in which the $^{13}$C signals of cellulose, glucuronoarabinoxylan (GAX), and other matrix polysaccharides can be assigned. The assigned $^{13}$C chemical shifts indicate that there are a large number of arabinose and xylose linkages in the wall, and GAX is significantly branched at the developmental stage of 2 weeks. 2D $^{13}$C−$^{13}$C correlation spectra measured with long spin diffusion mixing times indicate that the branched GAX approaches cellulose microfibrils on the nanometer scale, contrary to the conventional model in which only unbranched GAX can bind cellulose. The GAX chains are highly dynamic, with average order parameters of ~0.4. Biexponential $^{13}$C $T_1$ and $^1$H $T_1\rho$ relaxation indicates that there are two dynamically distinct domains in GAX: the more rigid domain may be responsible for cross-linking cellulose microfibrils, while the more mobile domain may fill the interfibrillar space. This dynamic heterogeneity is more pronounced than that of the non-grass hemicellulose, XyG, suggesting that GAX adopts the mixed characteristics of XyG and pectins. Moderate differences in cellulose rigidity are observed between the *Brachypodium* and *Arabidopsis* cell walls, suggesting different effects of the matrix polysaccharides on cellulose. These data provide the first molecular-level structural information about the three-dimensional organization of the polysaccharides in the grass primary wall.

Grasses have long contributed to human society as cereal crops for human diets and forage for animals. Recently, growing interest has been directed toward developing grasses as a major source of alternative, renewable energy. The polysaccharides stored in grass cell walls (CWs) can be extracted and converted into biofuels;1,2 thus, an understanding of the polysaccharide structures in the CWs is important for efficient degradation of this biomass.3 *Brachypodium distachyon* has been adopted as a model plant for structural and functional genomics of grasses, because it has a short lifecycle and a small genome, is self-pollinating, and is representative of many temperate grasses.4 Thus, a detailed understanding of the molecular structure and composition of the *Brachypodium* CWs in their native state is of significant interest.

The primary CW (PCW) of monocotyledonous (monocot) plants such as grasses differs from the PCW of dicotyledonous (dicot) plants as represented by *Arabidopsis thaliana*. In dicot PCWs, the main hemicellulose is xyloglucan (XyG), and the main pectins are linear homogalacturonan (HGA) and branched rhamnogalacturonan (RG), which contains arabinan, galactan (Gal), and arabinogalactan (AG) side chains.5,6 In

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grass PCWs, the main hemicelluloses are glucuronoarabinoxylans (GAX) and mixed-linkage glucan (MLG). GAX consists of a \((1 \rightarrow 4)\) β-linked xylose (Xyl) backbone substituted with arabinose (Ara) at the O3 position and glucuronic acid (GlcA) at the O2 position (Figure 1). Some of the Ara side chains are further esterified with ferulic acid (FA), which may cross-link multiple xylan (Xn) chains through dimerization. MLG is a copolymer of \((1 \rightarrow 4)\) and \((1 \rightarrow 3)\) β-linked glucose and is enriched in the walls of elongating plant cells. Grass PCWs also contain XyG, HGA, and RG, but their concentrations are much lower than in dicot PCWs. The amount of structural polysaccharides is more difficult to characterize because of the insoluble and amorphous nature of most of these polymers and the perturbing nature of chemical extraction. X-ray diffraction studies of plant CWs focused on the crystalline cellulose and not the amorphous matrix polysaccharides. Electron and atomic force microscopy of plant CWs require more concentrated base to be extracted. Scanning electron microscopy data of GAX-extracted cell walls showed a reduced matrix content between cellulose microfibrils, while less branched GAX and MLG lack the atomic resolution and chemical specificity. Solution nuclear magnetic resonance (NMR) studies require dissolution treatment of the CWs. One-dimensional \(^{13}\)C solid-state NMR has been used to study cellulose in various organisms but cannot easily resolve the signals of different polysaccharides in an intact CW.

While the composition of plant CWs can be obtained from chemical extractions followed by monosaccharide and linkage analysis, the three-dimensional structural arrangement of wall polysaccharides is more difficult to characterize because of the insoluble and amorphous nature of most of these polymers and the perturbing nature of chemical extraction. X-ray diffraction studies of plant CWs focused on the crystalline cellulose and not the amorphous matrix polysaccharides. Electron and atomic force microscopy of plant CWs lack the atomic resolution and chemical specificity. Solution nuclear magnetic resonance (NMR) studies require dissolution treatment of the CWs. One-dimensional \(^{13}\)C solid-state NMR has been used to study cellulose in various organisms but cannot easily resolve the signals of different polysaccharides in an intact CW.

Recently, we introduced multidimensional magic-angle-spinning (MAS) solid-state NMR spectroscopy for studying the structure of plant CW polysaccharides. By \(^{13}\)C labeling whole plants, we achieved the sensitivity necessary for two-dimensional (2D) and three-dimensional (3D) correlation NMR experiments, which in turn provide the resolution needed for understanding the structure and dynamics of multiple wall polysaccharides. The first organism we studied using this approach is the PCW of Arabidopsis thaliana. The 2D and 3D SSNMR spectra indicate that cellulose, XyG, and pectins form a single noncovalent network in the dicot PCW: both XyG and pectins show direct through-space cross peaks with cellulose, indicating spatial contacts within a nanometer. However, the sparseness of XyG–cellulose cross peaks indicates that XyG does not coat the microfibril surface extensively but may instead be embedded at specific locations in the microfibril. These results revise the prevailing model of PCWs, in which pectins and proteins are envisioned to form a network separate from the cellulose–hemicellulose network. The dicot wall polysaccharides are heterogeneously mobile: pectins are the most dynamic, and cellulose is the most rigid; hemicellulose exhibits intermediate mobility. A XyG-deficient Arabidopsis mutant exhibits enhanced mobilities for the remaining wall polysaccharides, consistent with the cross-linking function of XyG. In comparison, a partially depsectinated sample displays a much higher rigidity for the remaining polysaccharides. Loosening of this single-polysaccharide network by the protein expansin was achieved by expansin binding to cellulose, as shown using a sensitivity-enhanced NMR technique called dynamic nuclear polarization. Moreover, the expansin-binding site of cellulose is enriched in XyG, suggesting that the XyG-bound positions of cellulose may be mechanical hot spots for CW loosening.

Compared to that of dicot PCWs, the three-dimensional structure of grass PCWs has not yet been studied using multidimensional SSNMR. The existing model of grass PCW architecture depicts GAX as taking the role of XyG to cross-link cellulose microfibrils. The physical interactions between GAX and cellulose are believed to depend on GAX branching: linkage analyses and FT-IR data of sequentially extracted grass cell walls indicate that highly substituted GAX are extracted by dilute base, while less branched GAX and MLG require more concentrated base to be extracted. Scanning electron microscopy data of GAX-extracted cell walls showed a reduced matrix content between cellulose microfibrils. These results led to the proposal that highly substituted GAX forms the interstitial matrix between cellulose microfibrils, while less branched GAX hydrogen bonds with cellulose. However, in vitro binding assays indicate that none of the extracted GAX binds cellulose well: only 4–15% of GAX of any fraction binds to cellulose, in contrast to the tight binding between XyG and cellulose. Compared to GAX, MLGs are present at low concentrations in embryonal cell walls, but the concentrations increase significantly during growth. Because current understanding of the grass PCWs comes from extraction-based chemical and ultrastructural analysis, high-resolution structural...
studies by SSNMR under nonperturbing conditions will provide valuable and unique information.

In this study, we report the first comprehensive assignment of the $^{13}$C chemical shifts of Brachypodium primary wall polysaccharides and provide evidence of GAX–cellulose spatial contact in 2D $^{13}$C–$^{13}$C correlation spectra. We also investigated the mobilities of wall polysaccharides and show that GAX has larger-amplitude motions than its counterpart, XyG, in the dicot PCW. These results shed light on the 3D architecture of the grass PCW and establish a basis for future studies of grass cell walls.

**EXPERIMENTAL PROCEDURES**

**Plant Material.** Uniformly $^{13}$C-labeled *B. distachyon* PCW samples were prepared using a procedure similar to that reported previously. Briefly, plants were germinated and grown in the dark in a liquid containing $^{13}$C-labeled glucose (5 g/L) as the only carbon source. Growth in the dark prevents photosynthesis and the consequent dilution of $^{13}$C labels by unlabeled substrates produced from atmospheric CO$_2$. Two-week-old seedlings were harvested, and roots and leaves were separated and powered in liquid nitrogen, mixed with 80% (v/v) ethanol, heated at 80°C for 1 h, and cooled to room temperature. PCWs were pelleted by centrifugation at 12000 g for 20 min. The PCW material was never dried throughout the preparation. The wet pellet was washed with a chloroform/methanol (1:1) solution to remove nonpolar compounds and suspended in 50 mM sodium acetate buffer (pH 5.2) containing 1.5% SDS and 5 mM sodium metabisulfite to remove most intracellular proteins and low-molecular weight compounds. Starch was removed using α-amylase. These procedures are expected to have a minimal impact on the wall polysaccharides and structural proteins because of their low solubility and their interaction via hydrogen bonding or covalent cross-linking. Two samples were prepared from the roots and leaves of the plant. Approximately 76 mg of hydrated leaf CW and 58 mg of hydrated root CW were packed into two 4 mm MAS rotors for SSNMR experiments. A previously prepared $^{13}$C-labeled Arabidopsis PCW sample$^{30}$ was used to compare with the Brachypodium PCW samples.

**Solid-State NMR Spectroscopy.** All $^{13}$C NMR spectra were measured on a 600 MHz Avance II SSNMR spectrometer at 14.1 T using a 4 mm MAS probe. Typical radiofrequency (rf) field strengths were 62–71 kHz for $^1$H and 50 kHz for $^{13}$C. $^{13}$C chemical shifts were externally referenced to the Met Ce peak (14.0 ppm) in the model peptide formyl-Met-Leu-Phe-OH$^{29,30}$ on the TMS scale.

One-dimensional (1D) $^{13}$C MAS spectra were measured with either $^1$H–$^{13}$C cross-polarization (CP) or $^{13}$C direct polarization (DP) to create the initial transverse magnetization. CP spectra preferentially detect rigid molecules such as cellulose, while DP spectra with a short recycle delay (2 s) preferentially detect rigid molecules such as cellulose, while DP spectra with a short recycle delay (2 s) preferentially detect mobile polysaccharides. Quantitative spectra were measured using a DP experiment with a recycle delay of 15 s.

Several 2D $^{13}$C–$^{13}$C correlation experiments were conducted to resolve and assign the $^{13}$C chemical shifts of wall polysaccharides. The 2D J-INADEQUATE experiment$^{31,32}$ correlates the double-quantum (DQ) or sum chemical shifts of two directly bonded $^{13}$C spins with the single-quantum (SQ) chemical shifts of each $^{13}$C. The experiment used DP, $^{13}$C–$^{13}$C J coupling, and a recycle delay of 2 s to preferentially detect the signals of mobile matrix polysaccharides. The experiment was conducted at 293 K under 12 kHz MAS. A double-quantum-filtered (DQF) 2D $^{13}$C–$^{13}$C correlation spectrum complements the J-INADEQUATE experiment by detecting one-bond cross peaks of rigid polysaccharides. This experiment used SPC5-recoupled$^{33,34}$$^{13}$C–$^{13}$C dipolar coupling for polarization transfer and was conducted at 253 K under 7 kHz MAS. To probe intermolecular contacts, we measured 2D $^1$H-driven $^{13}$C spin diffusion (PDSD) spectra$^{22,34}$ with mixing times of 30 ms to 3.0 s. These PDSD experiments used either CP (at 253 K) or DP (at 293 K) to generate the initial $^{13}$C magnetization. A PDSD experiment with a short CP contact time of 35 μs was conducted at 293 K to selectively detect only the cellulose signals in the indirect dimension, whose cross peaks with GAX in the direct dimension can thus be better resolved in the 2D spectra.

Motional amplitudes were measured using the dipolar-double 2D $^{13}$C–$^{13}$C dipolar chemical shift (DIPSHIFT) correlation experiment$^{35,36}$ under 7 kHz MAS. $^1$H homonuclear decoupling was achieved using the FSLG sequence, which has a theoretical scaling factor of 0.577. The rigid-limit one-bond C–H dipolar coupling, after doubling and scaling, was found to be 26.2 kHz on the crystalline model peptide formyl-MLF-OH$^{29,38}$.

The ratios between the measured couplings of the polysaccharides and this rigid-limit value gave the order parameters.

$^{13}$C $T_1$ relaxation times, which reveal nanosecond motions, were measured using the inversion recovery sequence. The initial $^{13}$C magnetization was excited using a single pulse, and a long recycle delay of 15 s was used to obtain quantitative spectra. $^{13}$C-detected $^1$H $T_{1ρ}$ relaxation times, which reflect microsecond motions, were measured using a Lee–Goldburg spin-lock sequence$^{39–41}$ in which $^1$H spin diffusion was suppressed during the spin-lock period and the CP period to obtain site-specific relaxation times of the protons that are directly bonded to a $^{13}$C spin. The tilted effective $^1$H spin-lock field was 50 kHz. Most relaxation curves were fit with a double-exponential function (Tables S1 and S2 of the Supporting Information).

**Monosaccharide Composition Analysis.** Approximately 30 mg of PCWs from the *Brachypodium* leaves and roots was hydrolyzed in 2 M trifluoroacetic acid for 2 h at 121 °C, after which the acid was removed when the sample was dried under air at 50 °C. The dry material was solubilized in 100 μL of ddH$_2$O, and the monosaccharide composition was determined using high-performance anion-exchange chromatography.

The cellulose content was estimated by treating 10 mg of CW material with acetic–nitric reagent (80% acetic acid and concentrated nitric acid, 10:1) for 30 min at 100 °C.$^{24}$ Undigested pellets were washed several times with deionized water and then acetone, air-dried, and weighed.

**Glycosyl Linkage Analysis.** The PCW samples were permethylated, depolymerized, reduced, and acetylated. The resulting partially methylated alditol acetates (PMAAs) were analyzed by gas chromatography and mass spectrometry (GC–MS) as previously described.$^{42}$ Briefly, a 1–2 mg sample was suspended in DMSO, permethylated by the addition of Hakomori base while being purged with nitrogen gas, and mixed for 7 h. Iodomethane was then added, and the sample was mixed overnight. The sample then passed through a C18 SEP-PAK column, dried, and reduced with lithium borodeuteride in THF, which was neutralized and evaporated later. The sample was then treated with NaOH and methyl iodide in dry DMSO.$^{43}$ The sample was subjected to 4 M NaOH for 15 min;
methyl iodide was added, and the mixture was left for 40 min to ensure complete methylation. The permethylated material was hydrolyzed using 2 M TFA, reduced with NaBD₄, and acetylated using acetic anhydride and TFA. The resulting PMAAs were analyzed by GC−MS. Separation was performed on a 30 m Restek 2330 or Supelco 2380 bonded phase fused silica capillary column.

### RESULTS

**Brachypodium Cell Wall Composition from Linkage Analyses and Quantitative ¹³C NMR.** We obtained ¹³C-labeled *Brachypodium* PCW by growing the plant for 2 weeks in the dark with ¹³C-labeled glucose as the only carbon source, harvesting the roots and leaves, and preparing the insoluble CW material as described previously except without drying. This cell wall preparation procedure was mild, without involving strong acids or bases. Previous comparison of *Arabidopsis* PCW samples obtained from plants grown in light without extraction versus plants grown in the dark with mild treatment showed similar polysaccharide ¹³C intensities, even though the *Arabidopsis* PCW has more pectins, which can be extracted more easily.

At 2 weeks, the *Brachypodium* PCW is expected to contain mainly cellulose and GAX. Linkage analysis (Table 1) shows that 4-Xyl, 2,4-Xyl, and 3,4-Xyl account for 19% of all linkages. Xyl O3 is 4 times more substituted than O2, indicating that Ara is predominantly linked to Xyl O3 in the grass PCW, and Ara dominates GlcA in the xylan side chains. The Xyl concentration from the linkage analysis is lower than that from the sugar composition analysis (Table 2). This discrepancy may result from the incomplete accessibility of GAX to methylation because of its low solubility. A significant amount of Ara was detected: t-Ara and 5-Ara can be assigned mostly to GAX, while the low levels of 2-Ara and 3-Ara may be assigned to arabinan and arabinoalactan proteins (AGPs). In addition to GAX, the grass PCW can also contain other hemicelluloses such as XyG, MLG, and glucomannan. Sugar and linkage analyses showed that XyG was indeed present (4,6-Glc) but no MLG can be detected, because no 3-Glc linkage was found (Table 1). This is consistent with the fact that the level of MLG is known to vary greatly with developmental stage. The common pectic sugars such as 2,4-Rha (6.5%), 4,6-GalA (2.1%), and 4-Gal (5.6%) occur at low concentrations, as expected.

Quantitative ¹³C MAS spectra of the *Brachypodium* leaf and root PCWs (Figure 2a) show identical ¹³C chemical shifts and very similar intensity distributions, indicating that roots and leaves have similar polysaccharide compositions at 2 weeks of development.
Subtle differences in the relative intensities of Ara and Xn peaks are found between the two tissues: the leaf sample has slightly higher Ara intensities (108 ppm for AC1 and 82 ppm for AC2/C4) but lower Xn intensities (102 ppm for XnC1), indicating that the cell wall of leaves contains slightly more substituted GAX than the cell wall of leaves. The 13C CP spectra (Figure S1 of the Supporting Information) are also very similar between the two samples, indicating that the polysaccharides have similar mobilities in the two tissues.

When 1D 13C DP spectra were measured using a short recycle delay to preferentially detect the mobile polysaccharides, the spectrum differed from the quantitative 1D spectrum by precisely the chemical shifts of crystalline and amorphous cellulose (Figure 2b,c), as verified by comparison with the 13C spectrum of Avicel cellulose (Figure 2d). However, the grass PCW spectrum has much higher intensities of the surface amorphous cellulose than the Avicel spectrum, consistent with the microcrystalline nature of Avicel. Overall, the main rigid polysaccharide in the Brachypodium PCW is cellulose, while hemicelluloses and pectins are mobile.

To investigate how the PCW compositions differ between Brachypodium and Arabidopsis, we compared the quantitative 1D 13C spectra of the two plants (Figure 3). The Brachypodium sample shows protein (10−50 and 170−180 ppm) and pectin signals are the 101 ppm peak of galacturonic acid (GaIA) and rhamnose (R) and the 53.5 ppm peak of methyl ester.

Figure 3. 1D 13C quantitative MAS spectra of Brachypodium (black) and Arabidopsis (red) primary cell walls. (a) Spectra plotted with the same total integrated intensities. (b) Spectra plotted to have the same ic4 peak intensity at 89 ppm. The Brachypodium spectrum shows higher xylose (Xn), arabinose (A), and ferulic acid (FA) intensities and lower protein and pectin intensities. Representative pectin signals are the 101 ppm peak of galacturonic acid (GaIA) and rhamnose (R) and the 53.5 ppm peak of methyl ester.

Figure 4. 2D 13C J-INADEQUATE spectrum of the Brachypodium PCW, measured at 293 K under 12 kHz MAS. (a) Polysaccharide region of the 2D spectrum. (b) Selected 1D cross sections along the single-quantum (ω1) dimension. Peaks are assigned using the abbreviations shown, and superscripts distinguish different subtypes of the same monosaccharide. For example, eight sets of arabinose C1−C2 cross peaks are identified and indicated by superscripts a−h.
intensities lower than those of the Arabidopsis sample, both relative to the total spectral intensities and relative to the 89 ppm iC4 peak. For example, the 101 ppm peak of GalA and Rha C1 and the 80 ppm peak of GalA C4 and Rha C2 are weaker in the Brachypodium spectrum. The 53.5 ppm pectin methyl ester peak in the Arabidopsis spectrum is replaced by a 56 ppm ferulic acid (FA) methyl ester peak in the Brachypodium spectrum. However, the Brachypodium PCW has stronger Ara C1 (108 ppm) and Xn C1 (102 ppm) peaks, consistent with the fact that GAX primarily exists in grass.

13C Resonance Assignments of Brachypodium Cell Wall Polysaccharides by 2D Correlation NMR. To fully resolve the signals of all polysaccharides, we measured a 2D J-INADEQUATE spectrum31,49 at ambient temperature under conditions that favor the observation of the signals of mobile polysaccharides while disfavoring cellulose.18 The resulting 2D spectrum (Figure 4) shows narrow line widths of 0.4−0.9 ppm, indicating the highly dynamic nature of the matrix polysaccharides. Several regions of the 2D spectrum are of particular interest. First, the regions with DQ chemical shifts of 132−156 and 172−182 ppm contain the Xyl peaks and are more crowded than the corresponding region of the Arabidopsis spectrum (Figure 5) because of the presence of both GAX and XyG in the grass PCW. Five different sets of Xn peaks are identified and denoted a−e. Types a and b are characterized by a C2 chemical shift of 77−78 ppm, which can be assigned to GlcA O2-substituted Xyl. This assignment is supported by the fact that the Xn6 and Xn5 peaks have intensities similar to those of the GlcA peaks (Figure 4a). Type c was assigned to terminal Xyl based on literature chemical shifts.50,51 These Xn peaks are absent in the Arabidopsis spectrum.10 In comparison, terminal Xyl signals that originate from XyG are stronger in the Arabidopsis spectrum. These α-Xyl signals of XyG differ from the β-Xyl signals of GAX because of the different configurations of the anomeric carbon.

Eight sets of Ara C1−C2 cross peaks (denoted a−h) are resolved in the 2D J-INADEQUATE spectrum based on the characteristic Ara chemical shifts of 107−110 ppm21,52 (Figure 6), while nine sets of Ara C4−C5 cross peaks can be resolved. Unfortunately, the C2−C3 and C3−C4 cross peaks of Ara partially overlap (Figure 6b) and cannot be assigned unambiguously. The successive levels of ambiguity in the C2−C3 and C3−C4 assignments are denoted by symbols such as A2c2/f2 and A3c2/f2, respectively. For example, while the C2 chemical shifts of Ara-c, -d, and -f are resolved in the C1−C2 region, they become overlapped in the C2−C3 region, giving rise to ambiguous A2c2/f2 and A3c2/f2 peaks. Subsequently, the Ara-c and Ara-f C3−C4 peaks acquire another 2-fold assignment ambiguity, as denoted by symbols such as A3c2/f2 and A4c2/f2'. These assignment ambiguities create 17 possible combinations of Ara spin systems (Table 3), but only nine different Ara structures should exist in the sample, because only nine sets of C4−C5 cross peaks are resolved in the 2D spectrum. The Ara structural diversity in the grass PCW is understandable, because Ara exists not only in GAX side chains9 but also in pectins and AGPs. The GAX side chain Ara can be substituted with FA, while the pectic arabinans can have linkages at O5, O2, and O3.9,53 Finally, while the protein concentration in the grass CW is low, AGPs are ~90%
carbohydrate and thus can still contribute significant intensities in the spectra.\textsuperscript{45,46}

Figure 6d shows nine possible Ara linkages,\textsuperscript{54} many of which can be matched to the spin systems in the 2D spectra. For example, Ara-a can be assigned to 2,5-Ara (structure 7) because its downfield C2 and C5 chemical shifts of 88−90 and 69−71 ppm indicate O2 and O5 substitutions, respectively. Similarly, Ara-b can be assigned to 2-Ara. Both Ara-a and Ara-b peaks are relatively weak, consistent with the linkage analysis that reveals that the 2-Ara concentration is low. Ara-e and Ara-g have downfield C3 chemical shifts of 85 ppm, indicating O3 substitution such as in structure 3 or 4. Ara-d, -c1, and -f1 have upfield C5 chemical shifts of ∼62 ppm, indicating that they are t-Ara. Among these three types, Ara-d has high intensities; thus, it is likely the common t-Ara in the GAX side chain (structure 9). This is also consistent with the fact that Ara-d C3−C4 cross peaks are absent in the Arabidopsis PCW (Figure S2 of the Supporting Information), which does not contain GAX. Ara types c2, f2, c3, and f3 can be assigned to 5-Ara because of their downfield C5 chemical shift of 64−68 ppm.

In addition to the polysaccharide region, the 2D INADEQUATE spectrum also shows two pairs of carboxyl−methyl cross peaks (Figure S3 of the Supporting Information). These can be assigned to the acetyl groups of O2- and O3-acetylated xylan,\textsuperscript{55} because the other known acetylated polysaccharides, pectins and XyG, are present at very low levels in the Brachypodium PCW. The presence of O-acetylated GAX in the 2-week-old cell wall is interesting because O-acetylation is known to inhibit enzymatic degradation of wall polymers.\textsuperscript{56}

Panels a and b of Figure 7 show the aromatic region of the 30 ms 2D PDSD spectrum, which exhibits multiple FA signals,\textsuperscript{9,54} assigned from the unique FA methyl ester peak at 56 ppm. These measured FA 13C chemical shifts do not give strong evidence of the presence of diferulic acid, which is expected to exhibit more downfield chemical shifts for the quaternary carbons. Interestingly, the FA cross peaks have asymmetric intensities with respect to the diagonal, with the top left peaks having much higher intensities than the bottom right peaks. This can be attributed to the different 1H densities in the phenylene ring: the 1H-bonded carbons such as C2 and C8 resonate in the top left half of the spectrum, whereas quaternary and carbonyl carbons resonate in the bottom right. A clear FA−Ara cross peak at (148, 108) ppm is observed, confirming the position of FA in the GAX side chain.

GlcA is present as a minor substituent at position O2 of the xylan backbone. We observed a carbonyl cross peak at (72.6, 177.5) ppm, which can be assigned to GlcA C5/2−C6 correlation (Figure 7c). The GlcA signals of the grass PCW are also observed in the 2D INADEQUATE spectrum. These
GlcA peaks are readily distinguished from the 171 and 175 ppm GalA C6 peaks of the *Arabidopsis* PCW, which result from COOCH3 and COOH. The lack of these GalA signals in the *Brachypodium* sample is consistent with the absence of the 53.5 ppm methyl ester peak and indicates the low abundance of pectins in the grass PCWs.

In addition to GAX, the 2D J-INADEQUATE spectrum also resolved six different cellulose spin systems (Figure S4 of the Supporting Information). Cellulose is more rigid than GAX, as shown by the fact that its intensities are higher than those of the Xn, GlcA, and Ara signals in the 2D 13C DQF correlation spectrum (Figure S5 of the Supporting Information).

Compared to the *Brachypodium* spectrum, the *Arabidopsis* 2D DQF spectrum has much lower Ara and Xn intensities, consistent with a lack of GAX in the dicot PCW. On the other hand, the Gal C1−C2 cross peak at (101, 69) ppm is weaker in the *Brachypodium* spectrum, confirming that pectin concentrations are lower in the grass PCW.

None of these 2D spectra showed any signal near (86, 69) ppm, which are the expected frequencies of the C3−C4 cross peak for 3-Glc. Thus, MLG is below the detection limit in the 2-week-old grass PCW.

### Table 3. Assigned 13C Chemical Shifts of *Brachypodium* Primary CW Polysaccharides from 2D J-INADEQUATE Spectra

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<td>78.6</td>
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<td>These peaks are absent in <em>Arabidopsis</em> CW.</td>
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<td>Xn⁺</td>
<td>t-Xyl of GAX</td>
<td>102.4</td>
<td>73.9</td>
<td>76.7</td>
<td>70.2</td>
<td>66.0</td>
<td>ND</td>
<td>These peaks are absent in <em>Arabidopsis</em> CW.</td>
</tr>
<tr>
<td>Xn⁺</td>
<td>t-Xyl of GAX</td>
<td>102.4</td>
<td>73.9</td>
<td>76.7</td>
<td>74.6</td>
<td>63.6</td>
<td>ND</td>
<td>Consistent with Hollmann et al.</td>
</tr>
<tr>
<td>Xn⁺</td>
<td>4-Xyl of GAX</td>
<td>102.4</td>
<td>71.7</td>
<td>75.0</td>
<td>77.4</td>
<td>63.8</td>
<td>ND</td>
<td>Consistent with Roubroeks et al.</td>
</tr>
<tr>
<td>Xn⁺</td>
<td>t-Xyl of XyG</td>
<td>99.6</td>
<td>72.5</td>
<td>74.2</td>
<td>70.4</td>
<td>62.6</td>
<td>ND</td>
<td>α-Xyl C1 differs from β-Xyl in GAX by 2−3 ppm.</td>
</tr>
<tr>
<td>A⁺ (w)</td>
<td>2,5-Ara</td>
<td>107.1</td>
<td>89.8</td>
<td>76.8</td>
<td>78.6</td>
<td>70.7</td>
<td>ND</td>
<td></td>
</tr>
<tr>
<td>A⁺ (w)</td>
<td></td>
<td>107.3</td>
<td>87.8</td>
<td>72.6</td>
<td>ND</td>
<td>ND</td>
<td>ND</td>
<td></td>
</tr>
<tr>
<td>A⁺</td>
<td>t-Ara of GAX</td>
<td>109.2</td>
<td>82.1</td>
<td>78.2</td>
<td>85.7</td>
<td>62.4</td>
<td>ND</td>
<td>These peaks are absent in <em>Arabidopsis</em> CW.</td>
</tr>
<tr>
<td>A⁺</td>
<td>t-Ara</td>
<td>110.0</td>
<td>82.2</td>
<td>77.7</td>
<td>84.9</td>
<td>62.3</td>
<td>ND</td>
<td>Consistent with Dick-Perez et al.</td>
</tr>
<tr>
<td>A⁺</td>
<td>5-Ara</td>
<td>108.4</td>
<td>81.7</td>
<td>83.2</td>
<td>79.5</td>
<td>67.9</td>
<td>ND</td>
<td></td>
</tr>
<tr>
<td>A⁺</td>
<td>5-Ara</td>
<td>108.4</td>
<td>81.7</td>
<td>83.0</td>
<td>79.5</td>
<td>64.0</td>
<td>ND</td>
<td></td>
</tr>
<tr>
<td>A⁺</td>
<td>5-Ara</td>
<td>108.4</td>
<td>81.7</td>
<td>82.5</td>
<td>79.5</td>
<td>67.5</td>
<td>ND</td>
<td></td>
</tr>
<tr>
<td>A⁺</td>
<td>5-Ara</td>
<td>110.0</td>
<td>82.2</td>
<td>83.3</td>
<td>79.5</td>
<td>62.3</td>
<td>ND</td>
<td>C3 chemical shifts consistent with Tan et al.</td>
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<tr>
<td>A⁺</td>
<td>3,5-Ara</td>
<td>109.6</td>
<td>80.8</td>
<td>85.3</td>
<td>83.3</td>
<td>62.3</td>
<td>ND</td>
<td></td>
</tr>
<tr>
<td>A⁺</td>
<td>3,5-Ara</td>
<td>108.9</td>
<td>80.8</td>
<td>85.3</td>
<td>83.3</td>
<td>62.3</td>
<td>ND</td>
<td></td>
</tr>
<tr>
<td>A⁺</td>
<td>3,5-Ara</td>
<td>108.9</td>
<td>80.8</td>
<td>85.3</td>
<td>83.3</td>
<td>62.3</td>
<td>ND</td>
<td></td>
</tr>
<tr>
<td>A⁺</td>
<td>3,5-Ara</td>
<td>108.9</td>
<td>80.8</td>
<td>85.3</td>
<td>83.3</td>
<td>62.3</td>
<td>ND</td>
<td></td>
</tr>
<tr>
<td>A⁺</td>
<td>3,5-Ara</td>
<td>108.4</td>
<td>80.2</td>
<td>ND</td>
<td>ND</td>
<td>ND</td>
<td>ND</td>
<td></td>
</tr>
<tr>
<td>R (w)</td>
<td></td>
<td>100.8</td>
<td>79.1</td>
<td>ND</td>
<td>69.2</td>
<td>17.8</td>
<td>61.7</td>
<td>C5−C6 peaks from the 30 ms PDSD spectrum</td>
</tr>
<tr>
<td>Gal⁺</td>
<td></td>
<td>105.0</td>
<td>72.5</td>
<td>74.4</td>
<td>78.4</td>
<td>75.3</td>
<td>61.7</td>
<td>from Dick-Perez et al.</td>
</tr>
<tr>
<td>Gal⁺</td>
<td>103.4</td>
<td>74.0</td>
<td>74.9</td>
<td>73.8</td>
<td>74.8</td>
<td>61.7</td>
<td>61.7</td>
<td>from t-Gal (L) of Dick-Perez et al.</td>
</tr>
<tr>
<td>Gal⁺</td>
<td>103.9</td>
<td>71.9</td>
<td>73.8</td>
<td>69.5</td>
<td>75.8</td>
<td>61.8</td>
<td>61.8</td>
<td>Consistent with <em>Arabidopsis</em> and <em>Brachypodium</em> CWs</td>
</tr>
<tr>
<td>Unk (w)</td>
<td>unknown pectin</td>
<td>104.0</td>
<td>70.9</td>
<td>81.0</td>
<td>69.3</td>
<td>ND</td>
<td>ND</td>
<td>present in both <em>Arabidopsis</em> and <em>Brachypodium</em> CWs</td>
</tr>
<tr>
<td>GaLa/Man</td>
<td></td>
<td>100.2</td>
<td>69.3</td>
<td>ND</td>
<td>ND</td>
<td>ND</td>
<td>ND</td>
<td>C1 and C2 shifts consistent with Dick-Perez et al. and Jarvis et al.</td>
</tr>
<tr>
<td>99.6</td>
<td>69.0</td>
<td>101.3</td>
<td>69.8</td>
<td>ND</td>
<td>ND</td>
<td>ND</td>
<td>ND</td>
<td></td>
</tr>
</tbody>
</table>

**The downfield chemical shifts that reflect the linkages of arabinoses are shown in bold. Unless specifically noted, most assignments are *de novo* based on the connectivity patterns in the 2D spectra. Weak signals are denoted (w). ND means not determined. Tentatively assigned 13C chemical shifts are italicized and underlined.**

GlcA peaks are readily distinguished from the 171 and 175 ppm GalA C6 peaks of the *Arabidopsis* PCW, which result from COOCH3 and COOH. The lack of these GalA signals in the *Brachypodium* sample is consistent with the absence of the 53.5 ppm methyl ester peak and indicates the low abundance of pectins in the grass PCWs.

In addition to GAX, the 2D J-INADEQUATE spectrum also resolved six different cellulose spin systems (Figure S4 of the Supporting Information). Cellulose is more rigid than GAX, as shown by the fact that its intensities are higher than those of the Xn, GlcA, and Ara signals in the 2D 13C DQF correlation spectrum (Figure S5 of the Supporting Information). Compared to the *Brachypodium* spectrum, the *Arabidopsis* 2D DQF spectrum has much lower Ara and Xn intensities, consistent with a lack of GAX in the dicot PCW. On the other hand, the Gal C1−C2 cross peak at (101, 69) ppm is weaker in the *Brachypodium* spectrum, confirming that pectin concentrations are lower in the grass PCW.

None of these 2D spectra showed any signal near (86, 69) ppm, which are the expected frequencies of the C3−C4 cross peak for 3-Glc. Thus, MLG is below the detection limit in the 2-week-old grass PCW.

**GAX−Cellulose Intermolecular Contact from 2D 13C PDSD Spectra.** To determine if cellulose and GAX have spatial contact on the nanometer scale, we measured 2D 13C−13C PDSD spectra with mixing times from 30 ms to 3.0 s (Figure 8 and Figure S5 of the Supporting Information). At 30 ms, intraresidue cross peaks such as the Xn C1 correlations at 102 ppm and the Ara C1 correlations at 108 ppm are observed. When the mixing time increased to 1.5 s, multiple
intermolecular GAX—cellulose cross peaks were detected, for example, between XnC1 and iC4 at (102, 89) ppm, between AC1 and iC4 at (108, 89) ppm, and between FA and cellulose at (89, 56) ppm. Thus, both GAX backbone and side chains are in molecular contact with the cellulose microfibrils. The 1D cross sections illustrate some of these cellulose—GAX cross peaks more clearly (Figure 8c).

To further verify the cellulose—GAX interactions, we measured an edited 2D correlation spectrum in which the cellulose signals are detected in the direct dimension while all polysaccharides’ signals are detected in the direct dimension after a mixing time. The resulting 2D spectra (Figure 9) show unambiguous cellulose—Xn and cellulose—Ara cross peaks after samples have been mixed for 3.0 s. However, these intermolecular cross peaks are weak, indicating that only a small fraction of cellulose is in close contact with GAX. Indeed, the cellulose cross sections have intensity patterns quite different from the GAX cross sections after samples have been mixed for 1.5 s (Figure 8c), indicating that the two polysaccharides are not homogeneously mixed on the 1 nm scale that is relevant for 13C spin diffusion. This is not surprising, because the 3–5 nm diameter of the cellulose microfibril presents a significant spin diffusion barrier to the surrounding matrix polysaccharides. In comparison, all GAX cross sections have similar intensity patterns, indicating that 13C magnetization is equilibrated between the xylan backbone and the Ara and FA side chains.

**Polysaccharide Mobilities in the Brachypodium Primary Cell Wall.** To investigate polysaccharide mobilities in the Brachypodium PCW, we measured 13C—1H bond order parameters (S_{CH}) and 13C T1 and 1H T1ρ relaxation times. Order parameters provide information about the motional amplitudes, while relaxation times reflect motional rates. The C—H order parameters were measured using the 2D DIPSHIFT experiment, which yields time-dependent intensity decays indicative of the dipolar coupling strength. The matrix polysaccharides and cellulose show different dipolar decay rates, which allow their signals to be better resolved at chosen dipolar dephasing times. For example, with 41 μs dipolar dephasing, the 105 ppm peak of cellulose and Gal C1 was completely suppressed while the Ara (108 ppm) and Xn (102 ppm) C1 signals remain (Figure 10a). In the time domain dipolar cross sections (Figure 10b), cellulose exhibits much deeper intensity decays than GAX, indicating stronger dipolar couplings or higher order parameters. The cellulose S_{CH} values are 0.7–0.9, similar to the values of the Arabidopsis PCW sample (Figure 10c), and correspond to root-mean-square (rms) amplitudes of 15–26°. GAX exhibits much smaller S_{CH} values of ~0.4, indicating larger motional amplitudes. Interestingly, the GAX order parameters are noticeably smaller than the XyG order parameters in the Arabidopsis PCW and also smaller than the pectin S_{CH} values, indicating that GAX in the grass PCW is more mobile than most matrix polysaccharides in the dicot PCW.

The 13C T1 and 1H T1ρ relaxation times are shown in Figure 11. Most 13C sites exhibit double-exponential T1 decays (Figure S6 of the Supporting Information), with a short T1 component and a long T1 component of <0.5 and >3 s, respectively. The long T1 component is longer for cellulose (4–5 s) than for GAX (2–4 s). In addition, GAX contains a higher population of the faster-relaxing component while cellulose is dominated by the long T1 component. These results are consistent with the order parameter data in indicating that GAX is much more mobile than cellulose on the nanosecond time scale. Because 13C T1 data contain spin diffusion effects, to obtain more site-specific relaxation times and to probe motion on a slower time scale, we measured the 1H T1ρ values using a Lee–Goldburg
Similar to the $^{13}$C $T_1$ data, GAX exhibits much shorter $^1$H $T_{1p}$ values than cellulose, indicating faster motions. The short and long $T_{1p}$ components are similarly populated in GAX, while the long $T_1$ component dominates (~80%) in cellulose. Taken together, these data indicate that GAX undergoes faster motion than cellulose on both the nanosecond and microsecond time scales.

Comparing the Brachypodium and Arabidopsis samples, we find the relaxation times of GAX and XyG are similar but the fractions of the dynamic components are higher in GAX than in XyG. Cellulose in the two cell walls exhibits subtle differences: the slow-relaxing components in Brachypodium have longer relaxation times than in Arabidopsis. This result, together with the larger amplitude dynamics of GAX, suggests that cellulose–GAX interactions in the grass PCW are weaker than the cellulose–XyG interactions in the dicot PCW.

**DISCUSSION**

Brachypodium has recently been proposed as a model plant for grasses, but structural characterization of the Brachypodium PCW has been scarce. The study presented here establishes benchmark polysaccharide $^{13}$C chemical shifts for the PCW of this model grass. The samples analyzed here are never dried; thus, the polysaccharide interactions and mobilities closely resemble those under the native condition. At 2 weeks of age, the roots and leaves have very similar quantitative $^{13}$C spectra, indicating similar compositions of the CWs. Monosaccharide composition and linkage analyses (Tables 1 and 2) and $^{13}$C SSNMR spectra indicate that the Brachypodium PCW mainly...
The high-resolution 2D $^{13}$C correlation spectra of the never-dried grass PCW sample resolved nine Ara spin systems, five Xn spin systems, and six cellulose $^{62}$ spin systems. The peak multiplicities indicate the complex sugar linkages, hydrogen bonding patterns, and conformations of these polysaccharides. Only some of these $^{13}$C chemical shifts had been reported in the literature, mostly based on studies of extracted polysaccharides. $^{17,50,53,63-65}$ Our connectivity-based de novo assignments are consistent with peak intensities, with carbons in the same monosaccharide unit showing similar intensities and line widths. The unique chemical shifts of Xn (102 ppm) and FA (56 ppm) are particularly useful for the assignment of GAX peaks.

The assigned $^{13}$C chemical shifts allow us to detect, for the first time, intermolecular cross peaks between cellulose and GAX in the grass PCW (Figures 8 and 9). The cellulose cross peaks to FA and Ara are of particular interest, because they indicate that substituted GAX can approach the cellulose microfibrils on the $\sim$1 nm scale, which is the approximate distance upper limit of $^{13}$C spin diffusion. This result counters the notion that highly substituted GAX cannot bind cellulose. $^9$

The quantitative 1D $^{13}$C spectra (Figure 2) indicate an Ara/Xn intensity ratio of $\sim$0.78. This means that approximately three of four Xyl units are decorated with Ara, if we make the approximation that all arabinoses are located in GAX. Although some Ara exists in pectins and AGPs, these two polymers have very low concentrations in the wall; thus, the degree of GAX substitution should not be much lower than the estimate given by the $^{13}$C spectra. The 2D $^{13}$C correlation spectra of the Brachypodium PCW show larger amounts of xylan and smaller amounts of XyG and pectins. $^9,10$

Supplementary Figure 9. 2D $^{13}$C PDSD correlation spectra of the Brachypodium PCW. The spectra were measured with a short CP contact time of 35 $\mu$s at 293 K to select cellulose signals in the indirect dimension and detect their cross peaks with matrix polysaccharides in the direct dimension. Mixing times are (a) 30 ms and (b) 3.0 s. (c) Representative cellulose cross sections as a function of mixing time, where increasing GAX intensities are observed.
above. Therefore, the 2D $^{13}$C spin diffusion spectra indicate that $\sim$75% substituted GAX has molecular contact with cellulose. This result to some extent is not surprising, because in the absence of MLG, GAX is the dominant polysaccharide to cross-link cellulose, but because highly branched xylan chains cannot easily stack onto the cellulose surfaces, these intermolecular cross peaks imply that GAX surrounds the cellulose microfibrils in a disordered fashion. Moreover, the cellulose-interacting GAX domain must be a small fraction of all GAX, because the cellulose–GAX cross peaks are much weaker than the GAX–GAX and cellulose–cellulose cross peaks, and even with a mixing time of 1.5 s, the cellulose and GAX cross sections are far from equilibrated in their intensities (Figure 8).

The fraction of GAX that is in molecular contact with cellulose can be estimated from the biexponential relaxation data, which show that $\sim$60% of GAX has very short $^{13}$C $T_1$ and $^1$H $T_{1ρ}$ relaxation times (Figure 8). This means that the rigid GAX domain, which may be responsible for cross-linking cellulose, accounts for up to 40% of all GAX.

The Brachypodium PCW sample is more dynamic than the Arabidopsis sample, as manifested qualitatively by the many sharp signals in the 2D J-INADEQUATE spectrum, and quantitatively by the smaller C–H order parameters ($\sim$0.4) and shorter relaxation times of GAX compared to those of XyG in the Arabidopsis PCW. GAX shows large amplitude motions on both the nanosecond ($^{13}$C $T_1$) and microsecond ($^1$H $T_{1ρ}$) time scales, similar to XyG, but the fractions of the mobile component in GAX are larger than the mobile fractions of XyG in Arabidopsis. Compared to cellulose, the rigid GAX domain, which potentially interacts with cellulose, is still more mobile, as seen by its shorter relaxation times. This mobility difference is not inherently contradictory, because substantial dynamic gradients can occur within several bonds, as shown by $^{13}$C-resolved $^1$H line shapes of synthetic polymers and proteins. Overall, the main hemicellulose in the Brachypodium PCW has more heterogeneous dynamics than the main hemicellulose, GAX, in the Arabidopsis PCW.

Cellulose is rigid in both Brachypodium and Arabidopsis cell walls, with high order parameters of 0.7–0.9 for non-C6 groups, but the cellulose rigidity is not identical. The Brachypodium sample has a smaller C6 order parameter but longer $^{13}$C $T_1$ and $^1$H $T_{1ρ}$ relaxation times than the Arabidopsis sample. The former likely results from the influence of the highly dynamic GAX, which may promote larger amplitude motion of the protruding hydroxymethyl group. The latter observation, which implies faster motions of the Arabidopsis...
cellulose, is not fully understood. In general, the nanosecond motions that induce $T_1$ relaxation are fast torsional fluctuations, while the microsecond motions that induce $T_1\rho$ relaxation are likely collective motions of several sugar units. We hypothesize that the slightly faster motions of the dicot cellulose may be caused by the highly dynamic pectins, which are now known to interact with cellulose.6,22

In conclusion, this study provides the first comprehensive set of $^{13}$C chemical shifts of the polysaccharides in the near-native Brachypodium PCW. At 2 weeks of age, the cell wall contains cellulose and GAX as the major polysaccharides, while pectins, XyG, and structural proteins are present at concentrations much lower than those in the Arabidopsis PCW at the same developmental stage. The GAX in the grass PCW is $\sim$75% substituted, and a small portion of this highly branched GAX contacts cellulose on the nanometer scale based on 2D correlation spectra. Bond order parameters and relaxation times indicate that GAX is heterogeneously dynamic: a dominant dynamic domain, which may fill the intercellular space, coexists with a minor rigid domain, which may be responsible for cross-linking the cellulose microfibrils. Thus, in the grass PCW, GAX takes on the dual structural and dynamical characteristics of XyG and pectins in the dicot PCW. A large number of arabinose structures and linkage patterns are observed, indicating that this monosaccharide acts as a versatile building block in the grass PCW.

**ASSOCIATED CONTENT**

Supporting Information
Additional NMR spectra and tables. This material is available free of charge via the Internet at http://pubs.acs.org.

**REFERENCES**

The path forward for biofuels and biomaterials. Science 311, 484–489.


