Xyloglucan Metabolism Differentially Impacts the Cell Wall Characteristics of the Endosperm and Embryo during Arabidopsis Seed Germination

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Cell wall remodeling is an essential mechanism for the regulation of plant growth and architecture, and xyloglucans (XyGs), the major hemicellulose, are often considered as spacers of cellulose microfibrils during growth. In the seed, the activity of cell wall enzymes plays a critical role in germination by enabling embryo cell expansion leading to radicle protrusion, as well as endosperm weakening prior to its rupture. A screen for Arabidopsis (Arabidopsis thaliana) mutants affected in the hormonal control of germination identified a mutant, xyl1, able to germinate on paclobutrazol, an inhibitor of gibberellin biosynthesis. This mutant also exhibited reduced dormancy and increased resistance to high temperature. The XYL1 locus encodes an α-xylosidase required for XyG maturation through the trimming of Xyl. The xyl1 mutant phenotypes were associated with modifications to endosperm cell wall composition that likely impact on its resistance, as further demonstrated by the restoration of normal germination characteristics by endosperm-specific XYL1 expression. The absence of phenotypes in mutants defective for other glycosidases, which trim Gal or Fuc, suggests that XYL1 plays the major role in this process. Finally, the decreased XyG abundance in hypocotyl longitudinal cell walls of germinating embryos indicates a potential role in cell wall loosening and anisotropic growth together with pectin de-methylesterification.

Seed germination is a complex process that begins with the absorption of water and ends when the radicle breaks through the seed coat (or testa). In Arabidopsis (Arabidopsis thaliana), as in most angiosperms, the embryo is surrounded by the triploid endosperm and the seed coat of maternal origin (Nonogaki et al., 2010; North et al., 2010). The completion of germination requires the growth potential of the embryo to overcome the resistance of the endosperm and testa layers, which is controlled by the hormonal balance between abscisic acid (ABA) and gibberellins (GAs). During seed development, ABA induces embryo growth arrest at the transition from embryogenesis to the phase of reserve accumulation and then induces primary dormancy, thus preventing vivipary and allowing seed dispersal in a dormant state. Dormancy delays germination until environmental conditions become favorable for seedling survival and growth (Finkelstein et al., 2008; Nambara et al., 2010; Graeber et al., 2012). Dormancy depth varies among plant species and between Arabidopsis accessions; however, seed dormancy of the most commonly used accession Columbia-0 (Col-0) is relatively low and can be released by a few weeks of afterripening (dry storage) or stratification (cold imbibition). Shortly after hydration, ABA is rapidly degraded in both dormant and nondormant seeds, but ABA catabolism is more active in nondormant seeds, leading to lower ABA levels and thus allowing GA activation of germination processes (Millar et al., 2006). GA increases the elasticity of the wall, thereby reducing the resistance of the endosperm while triggering the elongation of the hypocotyl (Nonogaki et al., 2010). Radicle protrusion through the micropylar endosperm is also stimulated by ethylene, which has an antagonist action with ABA on endosperm cap weakening (Linkies and Leubner-Metzger, 2012). Microarray analyses highlighted the importance of cell wall remodeling processes during germination in various species (Penfield et al., 2006; Morris et al., 2011; Endo et al., 2012; Martínez-Andújar et al., 2012; Dekkers et al., 2013). These studies provided compelling evidence that the tissue-specific expression of genes encoding cell wall biosynthesis or modification enzymes, and their differential response to hormonal
signals in the endosperm and embryo, influences the rate of germination.

Cell walls are constituted of crystalline cellulose microfibrils that are embedded in an amorphous matrix of complex polysaccharides: pectin and hemicelluloses. Xyloglucan (XyG) is the major hemicellulose polymer in the primary cell walls of gymnosperms and most angiosperms, and its binding to cellulose microfibrils by hydrogen bonding contributes to loosening or stiffening of the wall during cell elongation (Cosgrove, 2005). XyG chains can be cleaved and reconnected by endo-transglycosylases/hydrolases (XTH). Other families of proteins also act on XyG chains, such as expansins, which are thought to nonenzymatically modulate XyG interactions with cellulose microfibrils, thereby controlling the distance between the microfibrils. XyG has a backbone of (1→4)-linked β-D-glucopyranosyl residues, which can be substituted with α-D-xylpyranosyl residues at O-6 (Supplemental Fig. S1). The pattern of XyG substitutions is described using a single-letter nomenclature (Fry et al., 1993). The letter G is used for an unsubstituted Glc and X is described using a single-letter nomenclature (Fry et al., 1996; Nambara et al., 1998; North et al., 2007), and seed phenotypes were correlated with spatio-temporal XyG accumulation during seed development and germination. Comparative studies using mutants impaired in two other apoplastic glycosidases, BGAL10 and AXY8, indicate a major role for XYL1 in XyG remodelling processes that affect germination.

RESULTS

Identification of the Paclobutrazol-Resistant Mutant xyl1-4

A screen of an ethyl-methanesulfonate (EMS) mutagenized population was carried out for seeds able to germinate on paclobutrazol, and a novel mutant, later named xyl1-4, was isolated. This screen has previously been used to select ABA-deficient mutants (Léon-Kloosterziel et al., 1996; Nambara et al., 1998; North et al., 2007), and seed dormancy was reduced in xyl1-4 as reported for most ABA-deficient mutants (Fig. 1, A and B). ABA contents of dry seeds were, however, slightly higher than wild type, indicating that this mutant was not impaired in ABA biosynthesis (Fig. 1C). Furthermore, the germination rate of xyl1 mutants was slower than that of ABA-deficient mutants (Supplemental Fig. S2; North et al., 2007).

The mutation affecting germination was mapped to a region on the lower arm of chromosome 1 between 24,699 and 25,665 Mb. Whole-genome sequencing identified a G to A substitution typical of EMS mutagenesis, which introduced a premature stop codon in the At1g68560 gene (Fig. 1D). This locus encodes an
The xyl1-2 and xyl1-4 alleles had similar phenotypes, consistent with the nomenclature first used by Sampedro et al. (2001), our mutant allele was named xyl1-4. Although XYL1 is the only α-xylanidosidase to be described that modifies XyG composition in vitro (Sampedro et al., 2001) and in vivo (Sampedro et al., 2010; Günl and Pauly, 2011), only mild vegetative phenotypes were observed for xyl1-4, as previously described for xyl1 alleles: reduced silique length and increased silique width (Supplemental Fig. S3). PAClobutrazol resistance, reduced dormancy, and ABA levels were analyzed for the T-DNA insertion allele xyl1-2/axy3-2 (Sampedro et al., 2010; Günl and Pauly, 2011) and showed that xyl1-2 and xyl1-4 alleles had similar phenotypes, confirming that these were due to defects in XYL1 function (Fig. 1).

**xyl1 Mutations Reduce the Relative Abundance of Fucosylated Residues in Seed Tissues**

Mutations in XYL1 have been shown to alter XyG accumulation in vegetative tissues (Sampedro et al., 2010; Günl and Pauly, 2011). To further compare xyl1-4 to xyl1-2 phenotypes and evaluate the effect of these mutations on XyG tissue-specific accumulation in seeds, XyG composition was analyzed by matrix-assisted laser-desorption ionization time of flight mass spectrometry. Measurement of relative contents of major oligosaccharide subunits in each extract was performed after endoglucanase digestion (Fig. 2). Seeds were dissected and separated into embryo and endosperm/testa fractions, either shortly after transfer of stratified seeds to 25°C in continuous light (3 h of imbibition) or at endosperm rupture (ER), when germination sensu stricto is completed and radicle protrusion is observed (after about 20 h of imbibition).

In both embryo and endosperm/testa fractions, a reduction in the relative abundance of fucosylated subunits (XXFG, XLFG) was correlated with an increased abundance of less substituted subunits (XXXG and XXLG) in both xyl1 alleles compared to wild type. Although XyG embryo composition was similar to that reported for seedlings (Günl and Pauly, 2011) or stems (Sampedro et al., 2010), highly substituted XLF subunits were more abundant in endosperm/testa (Fig. 2B) than in embryo fractions (Fig. 2A).

**Both xyl1-2 and xyl1-4 Impair XYL1 Protein Accumulation**

The xyl1-2 mutant contains a T-DNA insertion in the second exon of XYL1, and this has been shown to prevent transcript accumulation (Günl and Pauly, 2011). In xyl1-4, the mutation is located at the 3' end of the third exon, 200 nucleotides upstream of the stop codon (Fig. 1D), introducing a premature stop codon that would lead to the production of a truncated protein. To determine the impact of these two mutations on XYL1 protein accumulation, bidimensional (2D) gel electrophoresis was performed from dry mature seed protein extracts (Fig. 3). XYL1 protein accumulation has been previously observed during wild-type Arabidopsis seed germination but was not detected in the neosynthesised proteome (Galland et al., 2014). Comparative analysis of 2D gels indicated the absence, in dry seeds of both xyl1-2 and xyl1-4 mutants, of 4 protein spots detected in wild-type extracts (Fig. 3B-D). These 4 spots were analyzed by tandem mass spectrometry and confirmed to be the XYL1 protein (Supplemental Table S1). This highlighted the presence of multiple XYL1 isoforms; the shifts in mass and pI suggested that these isoforms might result from posttranslational modifications (Fig. 3B), which could not be determined by our mass spectrometry analysis. Nevertheless, in accordance with observed phenotypic alterations, our data provided evidence that neither xyl1-2 nor xyl1-4 seeds produce a full-length protein.

**Other Modifications of XyG Maturation Pathway Do Not Affect Germination**

In addition to XYL1, two other glycosidases, BGAL10 and AXY8, have been characterized that trim Gal and
To determine whether mutations of BGAL10 and AXY8 affect germination characteristics, we assessed dormancy, sensitivity to paclobutrazol, and thermoinhibition at 34°C of two T-DNA insertion alleles (bgal10-1 and bgal10-2; Sampedro et al., 2012), two EMS (axy8-1 and axy8-2), and two T-DNA insertion alleles (axy8-5 and axy8-6; Günl et al., 2011). Phenotypes of the xylosyltransferase double mutant xxt1 xxt2, which lacks XyG side chains (Cavalier et al., 2008), were also compared with xyl1.

In contrast to the increased tolerance to paclobutrazol and thermoinhibition and reduced dormancy of xyl1 mutant seeds, bgal10 and axy8 alleles displayed similar phenotypes to wild type (Fig. 4). Surprisingly, xxt1 xxt2 mutant seeds, which are defective for the addition of Xyl residues to XyG chains, displayed similar phenotypes to xyl1 mutants, which are affected in their removal. To evaluate whether the absence of bgal10 and axy8 seed phenotypes was correlated with the impact of these mutations on other organs, we measured silique size for two mutant alleles for each gene (Supplemental Fig. S3). As already described (Sampedro et al., 2010, 2012), silique length is reduced in xyl1 and bgal10 alleles. Although no vegetative phenotypes were previously observed in axy8 mutants (Günl et al., 2011),
reduced silique length was also detected for both axy8 alleles, axy8-1 being less affected than axy8-2. This was possibly due to differences in allele severity. Thus, in contrast to seeds, altered vegetative phenotypes were observed in all three XyG maturation mutants, together with the biosynthesis mutant xxt1 xxt2. Nevertheless, silique growth was less affected in bgal10 and axy8 than in xyl1 and xxt1 xxt2 mutants, suggesting either gene redundancy within BGAL10 and AXY8 families could compensate their defect in seeds or that XYL1 has a specific function in the control of seed dormancy and germination.

XyG Immunolocalization Is Not Altered in xyl1 Developing Seeds

To investigate further the impact of the XYL1 defect on XyG accumulation and its possible link with dormancy and germination phenotypes, the pattern of XyG accumulation was studied in wild-type and xyl1 seeds by immunolocalization using three different antibodies. LM15, LM25, and CCRC-M1 bind specifically to xylosylated (XXXG), galactosylated (XXLG), or fucosylated (XXFG) XyGs, respectively (Puhlmann et al., 1994; Marcus et al., 2008; Pedersen et al., 2012). During seed development, two stages were examined: 10 DAP during the embryo growth phase after the completion of embryogenesis and 14 DAP during reserve accumulation after embryo growth arrest. These stages were chosen as they marked the beginning and end of the observed increase in XYL1 expression in embryos (Fig. 5). At 10 DAP, all three epitopes were abundant in testa and endosperm cell walls of wild-type seed cells (Fig. 6, A and B; Supplemental Fig. S4, A and B). In embryos, XyG were observed in intracellular vesicles and cell walls. Intracellular vesicle labeling (Fig. 6B) suggests a progressive deposition of XyGs in wild-type embryo cell walls during seed development following their synthesis in Golgi vesicles. Furthermore, the stronger labeling in vascular tissues, compared to other hypocotyl cell walls, probably indicates that XyG are deposited earlier in these tissues (Supplemental Fig. S4, B and C).
By 14 DAP, XyG appeared intensely labeled and ubiquitously distributed in embryo cell walls (Fig. 6C; Supplemental Fig. S4, C and D). Compared to embryos, endosperm cell walls were relatively more labeled at 10 DAP than at 14 DAP (Fig. 6; Supplemental Fig. S4), also suggesting a differential timing of XyG deposition in these tissues.

As shown in Supplemental Figures S5 and S6, at both stages during seed development, XyG localization in xyl1 mutant cell walls was similar to that of wild type for all seed tissues. Similar relative increases in XyG labeling in embryo cell walls and decreases in endosperm cell walls were observed. These results suggest that during development of both wild-type and xyl1 seeds, embryo growth arrest and transition to reserve accumulation are correlated with an important deposition of XyGs in embryo cell walls and a possible concomitant degradation of XyGs in endosperm cell walls. Nevertheless, there was no clear link between these changes in cellular and tissue-specific XyG distribution and XYL1 spatiotemporal expression (Fig. 5).

In Germinating Embryos, xyl1 Alters Asymmetric Distribution of XyGs But Not Demethylsterified Pectins

To further investigate the evolution of XyG distribution during germination, immunolocalization analyses were performed in mature stratified seeds of wild type and both xyl1 alleles, during imbibition after transfer to 25°C in continuous light, at two time points: 3 h after transfer and at ER. After 3 h, XyG localization in wild-type xyl1 seeds was quite similar in both embryo and endosperm to that observed at 14 DAP, suggesting that no major changes had occurred during late seed development (Fig. 6, C and D; Supplemental Figs. S7 and S8). Furthermore, similar antibody labeling profiles were observed for the three epitopes (XXXG, XXLG, and XXFG) in embryo cell walls. Seed imbibition induced the rupture of the seed coat epidermal cell walls and mucilage release. XyG that were highly accumulated in outer cell walls during seed development were still detected in cell wall fragments (Fig. 6; Supplemental Fig. S7).

Significant variations in epitope abundance were detected in seeds at ER compared to 3-h-imbibed seeds. XyG immunolocalization in germinating wild-type seeds displayed an asymmetric distribution of the three epitopes in hypocotyl cells, with a more intense labeling in transversal than longitudinal cell walls (Fig. 7A). The reduced epitope labeling of longitudinal cell walls was detected only in elongating hypocotyl cells, whereas it remained uniform in cotyledon or root tip cell walls. Furthermore, an accumulation of XyG was also visible in intracellular vesicles (Fig. 7C). In contrast to wild type, the analysis of XyG localization in xyl1
mutants did not show an anisotropic epitope distribution in hypocotyl cell walls (Fig. 7B). Epitope abundance was low in both transversal and longitudinal cell walls, and intense epitope labeling was detected only in intracellular vesicles (Fig. 7D). Although the endosperm layer was not easily visible at this stage due to its rupture, no clear difference in XyG labeling was detected between \textit{xyl1} and wild type (Fig. 7). Radicle protrusion was therefore correlated with the disappearance of XyG epitopes in longitudinal cell walls of wild-type hypocotyls and in both transversal and longitudinal cell walls in \textit{xyl1}.

We hypothesize that the decreased abundance of XyG observed in longitudinal cell walls at ER (Fig. 7) might be correlated to asymmetric cell wall loosening, leading to hypocotyl directional growth. Selective pectin demethylesterification in longitudinal cell walls has been recently reported to trigger growth asymmetry in etiolated hypocotyls of dark-grown seedlings (Peaucelle et al., 2015). To determine whether pectin demethylesterification also occurred in elongating hypocotyls at ER, germinating seeds of wild-type and \textit{xyl1} were examined after immunolocalization using the 2F4 antibody that recognizes Ca\textsuperscript{2+}-crosslinked homogalacturonan with a low degree of methylesterification (Liners et al., 1989). More intense epitope labeling was observed in longitudinal compared to transversal cell walls of both wild-type and \textit{xyl1} hypocotyls (Fig. 7, E and F). However labeling was more heterogeneous than XyG epitope labeling, with stronger labeling at cell corners.

Our results thus suggest an involvement of both XyG removal and pectin demethylesterification in longitudinal cell wall loosening during hypocotyl elongation at ER. Furthermore, although XyG labeling was ubiquitously absent in \textit{xyl1} hypocotyls, asymmetric growth was not impaired. Indeed, radicle protrusion was synchronously observed after 20 h of imbibition at 25°C in the light in the stratified seeds of both \textit{xyl1} and wild type used here for immunolabeling, and the absence of XyG in transversal cell walls did not notably affect directional hypocotyl growth.

\textbf{XYL1 Activity in Endosperm Controls Seed Germination}

Since XyG alterations in \textit{xyl1} embryos were not associated with hypocotyl growth modifications, we investigated whether the permeability or mechanical resistance of the testa or endosperm layers was affected. Increased seed coat permeability could allow increased solute leakage from mutant seeds at imbibition. Despite ABA contents in mutant dry seeds being higher than those of wild type (Fig. 1C), \textit{xyl1} seeds exhibited a reduced GA requirement for germination (Fig. 1A); this suggested that ABA might be released upon imbibition. Measurement of ABA contents in dry and imbibed seeds found that similar amounts were leached into the imbibition medium (Supplemental Figure S9). This corresponded to a loss of 15% and 12% of dry seed ABA content on imbibition for wild type and mutant seeds, respectively; further decreases could also occur through ABA catabolism.

Reciprocal crosses enable the maternal determinism of altered germination characteristics to be investigated. Since the testa is derived from the ovule integuments, heterozygous F1 \textit{zygotes} (embryo and endosperm) are surrounded by a maternal testa. Germination rates for freshly harvested seeds showed that F1 seeds, harvested from a mutant \textit{xyl1} mother plant fertilized with wild-type pollen, displayed the same phenotype as wild type and were dormant (Fig. 8). Thus, the altered germination of \textit{xyl1} seeds did not result from defective XYL1 in testa layers reducing their mechanical resistance.

To determine whether XYL1 activity in the endosperm may be involved in the control of its rupture during seed germination, we produced \textit{xyl1}-2 primary transfectants that expressed the XYL1 gene under the control of the \textit{EPR1} promoter, which was previously
shown to be endosperm specific during seed development and germination (Dubreucq et al., 2000). Germinating T2 seeds of most lines showed reduced resistance to paclobutrazol compared to xyl1-2. To facilitate subsequent analyses, we selected transgenic lines, segregating with a 3:1 ratio for hygromycin resistance in T2 seeds as predicted for a single T-DNA insertion. Further analysis of germination characteristics of seeds from four independent homozygous transgenic lines showed that the dormancy of freshly harvested seeds and the resistance of stratified seeds to paclobutrazol and thermoinhibition were similar to wild type (Fig. 9).

To determine whether the differential XYL1 expression in embryo and endosperm may affect germination rates, we analyzed the timing of radicle protrusion of after-ripened seeds in which dormancy was released. In the absence of stratification treatment, wild-type and xyl1 seeds exhibited similar germination rates, while in transgenic seeds radicle protrusion was delayed (Fig. 10). This suggested that when endosperm resistance is increased due to XYL1 overexpression, the growth potential of the XYL1-deficient embryo is less effective at overcoming the mechanical restraint imposed by the endosperm.

To confirm that pEPR:XYL1 transformants exhibit differential XYL1 expression in endosperm and embryo, the XyG composition was analyzed. XyG profiles for transgenic embryos were very similar to xyl1-2, at both stages of imbibition, whereas in testa/endosperm fractions they were similar to wild type (Fig. 11). EPR1-driven endosperm-specific XYL1 expression in xyl1-2 mutants was thus able to restore dormancy of freshly harvested seeds and sensitivity of stratified seeds to germination inhibitors and even delay germination of nondormant seeds.

**DISCUSSION**

XyG, the most abundant hemicellulose in dicot cell walls, is thought to play a crucial role in cell elongation and also impact cell wall rigidity (Scheller and Ulvskov, 2010). Nevertheless, little is known about its precise contribution to dormancy and germination processes. In this study, we have isolated a xyl1 mutant allele able to germinate on high concentrations of the GA biosynthesis inhibitor paclobutrazol (Fig. 1), thus indicating its lower GA requirement for seed germination. The mutated locus,
XYL1, encodes a previously described glycoside hydrolase, which modifies XyGs by removing Xyl residues from XyG side chains (Sampedro et al., 2001). This gene has previously been reported to be expressed in vegetative tissues and involved in cell wall remodeling processes (Sampedro et al., 2010; Günl and Pauly, 2011). Here, we observed that \textit{XYL1} is differentially expressed throughout embryo and endosperm tissues during seed development and imbibition (Fig. 5). Accordingly, in addition to plant growth defects, such as shorter and thicker siliques, \textit{xyl1} mutants also exhibit altered germination characteristics, such as reduced dormancy and increased tolerance to germination inhibitors and thermoinhibition (Figs. 1 and 4; Supplemental Figs. S2 and S3).

\textbf{XYL1 Activity in Seeds May Require Posttranslational Modifications}

In accordance with mutant germination phenotypes and \textit{XYL1} transcript accumulation, the \textit{XYL1} protein is accumulated in dry seeds. Proteome analysis by 2D gel electrophoresis of wild-type seed extracts detected four polypeptide spots corresponding to \textit{XYL1}, which were absent in both the T-DNA allele \textit{xyl1-2} and the EMS allele \textit{xyl1-4} (Fig. 3; Supplemental Table S1), thus indicating that at least 4 different isoforms of the enzyme are present at this stage. Previous cell wall proteomic studies have also reported the presence of several \textit{XYL1} isoforms in vegetative tissues. The \textit{XYL1} protein was detected in extracts from stems and etiolated hypocotyls, and the observed mass shifts were predicted to be due to glycosylation at a number of sites (Minic et al., 2007; Albenne et al., 2009; Zhang et al., 2011). Moreover, a pI shift has also been observed due to protein phosphorylation. Such phosphorylation would lead to activation of the protein (Kaida et al., 2010), whereas the glycosylation that takes place in the Golgi apparatus would mediate activity modifications and induce protein secretion to cell walls (Ruiz-May et al., 2012). It is likely, therefore, that the mass and pI shifts detected in dry seed extracts correspond to \textit{XYL1} polypeptides targeted to the wall after glycosylation processes in the Golgi as well as differentially phosphorylated isoforms. These protein isoforms most probably represent different states of the translational modification process leading to the mature protein, whose activity has been detected in the apoplastic fluid of Arabidopsis seedlings (Sampedro et al., 2001). A regulatory role for these posttranslational modifications in cell wall remodelling during embryo growth and ER during germination is probable but would require further investigation.

\textbf{XyG Composition Modulates Endosperm Resistance}

Loss of \textit{XYL1} function caused alterations in XyG composition (Fig. 2). The defect in Xyl removal caused a relative increase in galactosylated residues at the expense of fucosylated residues in both embryo and testa/endosperm, as previously reported in \textit{xyl1} mutant stems and seedlings (Sampedro et al., 2010; Günl...
and Pauly, 2011; Sampedro et al., 2012). It has been suggested that the sequential removal of substituted sugars is most probably dependent on the previous enzymatic hydrolysis of neighbor residues. In xyl1, the removal of the galactosyl residue may be subject to a previous hydrolysis by XYL1, therefore resulting in an excess of galactosyl residues (Günl et al., 2011).

The accumulation of XyG that links cellulose microfibrils has been suggested to increase cell wall stiffness (Hayashi and Kaida, 2011). Variations in the complexity of XyG side chains may modulate endosperm weakening by hydrolases (XTH, glucanases, endo-β-mannanases) or extensibility by expansins, since these activities have been widely reported to facilitate radicle protrusion (Nonogaki et al., 2010; Park and Cosgrove, 2012; Shi et al., 2015). Upon imbibition, the micropylar endosperm has been shown to stretch before its rupture due to embryo growth (Lee et al., 2012). A clear link between germination characteristics and XyG composition in the endosperm has been demonstrated here. A reduction in the relative abundance of highly ramified fucosylated residues in the xyl1 endosperm was correlated with decreased dormancy and GA requirement for germination, whereas XYL1 expression under the control of the endosperm-specific EPR1 promoter in transgenic mutant seeds restored wild-type XyG profiles and germination phenotypes (Figs. 1, 2, 9, and 10). Since the double mutant xxt1 xxt2 exhibited the same phenotypes as xyl1 (Fig. 4), it further suggests that endosperm cell wall resistance is reduced when XyG are either absent or contain less branched side-chains that potentially could influence interactions with cellulose microfibrils (Zhao et al., 2014; Park and Cosgrove, 2015).

The bgal10 and axy8 mutants did not display similar seed phenotypes to those observed for xyl1 (Fig. 4). Nevertheless, transcriptome analyses have revealed that transcript abundance of AXY8, BGAL10, and XYL1 increased during seed imbibition, although XYL1 was more strongly expressed (Nakabayashi et al., 2005; Winter et al., 2007). Moreover, BGAL10 and XYL1 displayed similar expression patterns in response to dormancy cycling, indicating that BGAL10 is also highly regulated in seeds (Cadman et al., 2006). The milder effect of mutating BGAL10 and AXY8 might result from functional redundancy, since residual galactosidase and fucosidase activities, respectively, were detected in these mutants, which contrasts with the complete absence of xylooxidase activity in xyl1 (Sampedro et al., 2010, 2012; Günl et al., 2011). Nonetheless, a reduction in silique length was observed in all three mutants (Supplemental Fig. S3), suggesting that if functional redundancy occurs for galactosidase and fucosidase, it is differentially regulated during plant and seed development. In axy8 vegetative tissues, increased amounts of the fucosylated oligosaccharide XXFG were observed with a concomitant decrease in XXLG (Günl et al., 2011). In bgal10 mutants, the relative abundance of XXFG was lower than in wild type, but higher than in the bgal10 xyl1 double mutant (Sampedro et al., 2012). These data further indicate that XyG side chain complexity is negatively correlated with germination potential in these mutants, as observed in pEPR:XYL1 overexpressors.

Figure 11. Matrix-assisted laser-desorption ionization time of flight analysis of XyGs in germinating seeds of wild type (black bars), xyl1-2 (white bars), and four independent pEPR1:XYL1 homozygous transgenic lines (gray bars, from left to right: T1 to T4). Analysis was performed on 3-d-stratified seeds during imbibition at 25°C in the light. Embryos were dissected from testa and endosperm 3 h after sowing and at ER. Relative abundance of major subunits is shown as means of 4 biological replicates with SE. Minor subunits (<5% relative abundance) are not shown. Mean values represent acetylated and nonacetylated subunits of XXLG, XXFG, and XLFG combined.
Nevertheless, other explanations are possible. An increased accumulation of unbound XyG oligosaccharides was previously measured in xyl1 mutants compared to wild type (Sampedro et al., 2010; Günl et al., 2011), and it is possible that these unbound oligomers interfere with XTH activities, thereby impacting XyG remodeling and germination properties. Alternatively, as described in earlier studies and recently reviewed by Scheller and Ulvskov (2010), XyG oligomers might act as signal molecules. Nevertheless, the accumulation of these oligomers in xyl1 seeds is unlikely to play a major role in the germination phenotypes observed here, as they are not detected in xxt1 xxt2 mutants that have similar phenotypes to xyl1.

Anisotropic XyG Intracellular Distribution May Contribute to Cell Wall Loosening

Immunolocalization of xylosylated, galactosylated, and fucosylated residues indicated that XyG accumulation is developmentally regulated in seeds, but no major alterations were observed prior to ER (Figs. 6 and 7). At ER, a striking difference was observed in growing hypocotyl cells of germinating wild-type embryos with XyG labeling detected in transversal cell walls, but absent from longitudinal walls, suggesting an anisotropic XyG secretion in transversal walls or a specific hydrolysis and/or remodeling from longitudinal walls. The latter seems more likely, since ubiquitous cell wall labeling was observed at earlier stages and decreased XyG abundance correlated with concomitant labeling of intracellular vesicles. In accordance, inhibitors of endocytosis have previously been shown to significantly delay germination, indicating that internalization of extracellular molecules plays an important function (Pagnussat et al., 2012). Interestingly, the differential XyG accumulation was observed in a region just above the radicle tip previously described as the embryo elongation zone during germination (Slawinska et al., 2009). The XyG localization correlates well, therefore, with cell elongation and may be implicated in cell wall loosening prior to the directional growth of hypocotyls that leads to radicle emergence.

In dark-grown elongating hypocotyls, an asymmetric loosening of longitudinal walls has been observed by atomic force microscopy prior to changes in cortical microtubule orientation (Peaucelle et al., 2015). The selective demethylsterification of cell wall pectin in longitudinal walls has been reported to be responsible for this mechanical asymmetry, which is required for growth symmetry breaking. In accordance, an asymmetrical accumulation of homogalacturonans with a low degree of methylesterification was also observed here in elongating hypocotyls at ER, which was inversely correlated with XyG abundance (Fig. 7). Nevertheless, we cannot exclude that decreased XyG abundance in longitudinal cell walls results from epitope masking, since removal of pectic homogalacturonan has previously been shown to increase LM15 labeling (Marcus et al., 2008). However, almost identical labeling patterns were observed here with three different antibodies (LM15, LM25, and CCRC-M1), implying that if epitope masking occurs, it would affect the recognition of distinct XyG epitopes equally. Furthermore, the intense homogalacturonan labeling of cell corners with 2F4 was not an exact match of the uniform reduction of XyG labeling in longitudinal cell walls. This rule out a direct negative effect of pectin demethylesterification on XyG epitope detection, although remodeling of other cell wall components might still be involved. Nonetheless, the specific spatiotemporal regulation of XyG abundance in hypocotyl cells suggests that decreased XyG abundance contributes to longitudinal cell wall loosening together with pectin demethylsterification.

XYL1 Activity Is Essential for Anisotropic XyG Accumulation in Hypocotyl Elongating Cells

While no differences were detected between xyl1 and wild type from seed maturation to early imbibition, at ER XyG were absent from the longitudinal and transversal walls of xyl1 hypocotyl cells, although they were still detected within intracellular vesicles (Fig. 7). This indicates that alterations to XyG structure in xyl1 mutants may affect polarized XyG secretion or remodeling. Moreover, labeled intracellular vesicles observed in xyl1 hypocotyl cells may contain the excess, unbound XyG detected in previous studies (Sampedro et al., 2010; Günl and Pauly, 2011). In accordance, a recent study on the biosynthesis mutant mur3 that produces less substituted XyG due to defective galactosylation reported that intracellular polysaccharide aggregates were observed in mur3, which were attributed to altered secretion of Gal-depleted XyG (Kong et al., 2015).

Intriguingly, although mutation of XYL1 prevented the establishment of an anisotropic XyG distribution in mutant hypocotyls (Figs. 6 and 7), this did not negatively impact hypocotyl elongation and radicle protrusion (Figs. 1 and 4). Hence, even if decreased XyG contents in hypocotyl longitudinal cell walls are necessary for cell elongation to occur, their anisotropic distribution is not essential for asymmetrical growth. Other mechanisms could generate this asymmetry, such as the degree of pectin demethylsterification, prior to microtubule reorganization (Peaucelle et al., 2015). Yet size reduction in siliques was correlated with increased width in both xyl1 and xxt1 xxt2 and to a lesser extent in bgl110 and axy8, strongly suggesting a more predominant role of XyG metabolism in directional growth in these organs than hypocotyls (Supplemental Fig. S3; Sampedro et al., 2010; Günl and Pauly, 2011).

To conclude, our study provides further evidence that endosperm has a pivotal role as a mechanical barrier to embryo growth in the control of seed dormancy and germination. It also reveals that fine-tuning...
of XyG hydrolysis in cell walls is essential in the control of microfibrillar ER in response to exogenous and endogenous signals, such as temperature and GAs. The spatio-temporal regulation of XyG intracellular accumulation was shown to be tightly correlated with hypocotyl cell elongation; however, the mechanisms regulating XyG anisotropic abundance and possible roles in cell wall loosening still remain to be clarified.

MATERIALS AND METHODS

Plant Materials and Growth Conditions

Arabidopsis (Arabidopsis thaliana; Col-0 accession) wild-type and mutant seeds were surface sterilized, sown in petri dishes containing Arabidopsis Gamborg B5 medium (Duchefa, http://www.duchefa.com) supplemented with 30 mM Suc, and stratified at 4°C in the dark for 3 d. Petri dishes were then placed for 4 d in a growth chamber (16-h photoperiod, 50-μmol m⁻² s⁻¹ light intensity, 18°C, 60% relative humidity). Germinated seedlings were transferred to soil (Tref Substrates, http://www.trefgroup.com) and, unless otherwise stated, grown in a glasshouse with a minimum photoperiod of 13 h assured by supplementary lighting.

Mutant Isolation and Map-Based Cloning

A mutant screen was performed as previously described for the axl1 mutant identified using the same phenotype of germination tolerance to paclobutrazol (North et al., 2007). For mapping, the mutant (Col-0 accession) was crossed with wild-type plants (Landsberg erecta accession), and 148 homozygous xyl1-1 plants were selected from the F2 progeny based on germination phenotypes. Genomic DNA was extracted from flower buds of selected plants and an approximate genome position determined based on recombination percentages with simple sequence length polymorphisms, as described previously (North et al., 2007). Whole-genome resequencing was carried out on xyl1-1 DNA isolated from 3 g of flower buds with CTAB buffer (2% [w/v] N-cetyl-N,N,N-trimethyl-ammonium bromide, 1.4 μL NACl, 20 mM EDTA, 100 mM Tris-HCl, pH 8.0, 0.2% [v/v] β-mercaptoethanol) prior to extraction with an equal volume of chloroform:isooamyl alcohol (24:1) and precipitation with 0.7 volumes of isopropanol. After supernatant removal and pellet drying, the pellet was resuspended with 10 mM Tris-HCl, 1 mM EDTA, pH 8.0. DNA was then purified with DNeasy Plant MidiKit (Qiagen) from step 13 including an RNase treatment. Sequencing was performed by Genoscreen (France) using an Illumina HiSeq2000 sequencer.

Germination Experiments

For dormancy assays, freshly harvested seeds were sown in triplicate in Petri dishes containing 0.5% (w/v) agarose and then placed in a growth chamber (continuous light, 25°C, 70% relative humidity). Germination was scored each day based on radicle protrusion. For paclobutrazol resistance tests, surface-sterilized seeds were sown on 0.5% (w/v) agarose supplemented with 30 mM Suc, and separated using immobilized pH gradients gel strips (Immobiline, Whatman, http://www.whatman.com) prior to extraction with an equal volume of chloroform:isooamyl alcohol (24:1) and precipitation with 0.7 volumes of isopropanol. After supernatant removal and pellet drying, the pellet was resuspended with 10 mM Tris-HCl, 1 mM EDTA, pH 8.0. DNA was then purified with DNeasy Plant MidiKit (Qiagen) from step 13 including an RNase treatment. Sequencing was performed by Genoscreen (France) using an Illumina HiSeq2000 sequencer.

ABA Content Determination

One hundred milligrams of dry seeds was analyzed directly or imbibed in 0.8 M L-water, and after 1 or 5 h of imbibition in the light at room temperature with gentle agitation, the medium around imbibed seeds was recovered. Dry seeds, imbibed seeds, and imbibition medium were frozen in liquid nitrogen and then freeze-dried. Dry and imbibed seeds were ground in 2 mL of extraction solvent (acetone/water/ acetic acid, 80/19/1, v/v/v) to which 10 ng of deuterated ABA (purchased from Irisa Zaharia, Plant Biotechnology Institute, National Research Council Canada, http://www.nrc-cnrc.gc.ca) was added as an internal standard. Samples were centrifuged and the supernatant recovered, the pellet was then resuspended in a further 1 mL of extraction solvent by sonication, recentrifuged, and the supernatant combined. The extraction solvent was then evaporated and the residue resuspended by sonication in 0.2 mL of chromatography mobile phase (acetoniitrile/water/acetic acid, 50/50/0.05, v/v/v). Dried imbibition medium was directly resuspended in 0.2 mL chromatography mobile phase with 10 ng deuterated ABA by mixing. Finally, all samples were filtered through a 1.6-mm GFA filter (Whatman, http://www.whatman.com/). ABA was quantified using an LC-ESI-MS-MS system (Quattro LC; Waters, http://www.waters.com).

Cloning and Plant Transformation

A 0.8-kb promoter fragment for the EPR1 gene (At5g27380) was amplified from wild-type Col-0 genomic DNA by PCR using hpf Ultra DNA polymerase (Stratagene, Amsterdam, The Netherlands) and the following primers: forward 5′-CTAATCGGCCCTGATCATATAG-3′ and reverse 5′-GGGGGGCCCAACTTGGTAAGTATGA-3′. The PCR product was then recombined into pDONR207 vector with BP clonase (Invitrogen; http://www.invitrogen.com/) according to the manufacturer’s instructions and transformed into the Escherichia coli strain DH10B. The XYL1 coding sequence (At1g68560) was amplified from wild-type Col-0 genomic DNA by PCR using the following primers containing attB1 and attB2 recombination sequences: forward primer 5′-GGGACGAGTGGTGA-3′; reverse primer 5′-GGGACGAGTGCGCTAGTAC-3′. The resulting PCR product was then recombined into pDONR207 vector with BP clonase (Invitrogen; http://www.invitrogen.com/) according to the manufacturer’s instructions and transformed into the binary Gateway Destination vector pMD132 (Life Technologies) after digestion with the restriction enzymes HindIII and AccI.

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Proteomic Analysis

Total soluble protein extracts were prepared from 50 mg of dry mature seeds, and the concentrations of total soluble proteins in the lysates were estimated using the Bradford protein assay, as previously described (Rajou et al., 2011). All samples were stored at −20°C prior to analysis. Isoelectric focusing was performed with 150 μg of protein for each sample. Proteins from the various extracts were separated using immobilized pH gradient gels strips (Immobile DryStrip, nonlinear pH gradient 3-11, 24 cm; GE Healthcare). Immobilized pH gradient gels strips were incubated for 14 h at 22°C in rehydration buffer (18 mM Tris-HCl, 14 mM Tris base, 7 μL urea, 2 μL thiorurea, 4% [w/v] CHAPS, 20 mM dithiothreitol, 2% [v/v] Triton X-100, and 1% [v/v] pharmalyte pH 3-10 carrier ampholutes) containing 150 μM protein, and then recombined into the modiﬁed binary vector pMDC32 containing the EPR1 promoter fragment. The resulting binary vector was transformed into Agrobacterium tumefaciens C58CipMP90 by electroporation prior to stable transformation of wild-type or xyl1-2 plants.
XyG and Pectin Immunolocalization

Developing siliques were staged by tagging flowers on the day of pollination, in the same way as for reporter gene expression analysis, and harvested at 10 or 14 DAP. Seeds were also sown on filters, placed at 4°C for 3 d in the dark, and then placed in a growth chamber (continuous light, 25°C, 70% relative humidity) prior to harvest after 3 h of illumination. Seeds that had undergone ER were selected at the start of radicle protrusion, about 20 h after transfer to 25°C. Samples were fixed in a solution of 4% (w/v) paraformaldehyde and 0.1% Triton X-100 under vacuum for 1 h. Samples were then washed three times in phosphate-buffered saline (PBS), dehydrated through a graded ethanol series (30, 50, 70, 90, and 97% [v/v] in PBS), and then incubated in a mixture of 100% wax and 97% ethanol (1:1, v/v) at 40°C overnight and embedded in 100% wax (30, 50, 70, 90, and 97% [v/v] in PBS). At each step, the PBS buffer was used for LM15, LM25, and CCRC-M1 epitope detection, and the T/Ca/S buffer (20 mM Tris-HCl, pH 8.2, 0.5 mM CaCl₂, and 150 mM NaCl) for 24F epitope detection. Samples were blocked with 1% (w/v) bovine serum albumin in PBS or T/Ca/S buffer. Samples were then incubated with antibodies diluted in their respective blocking solution. After three washes of 5 min each in PBS or T/Ca/S buffer without bovine serum albumin, slides were incubated with antirat IgG or antimouse IgG labeled with Alexa Fluor 488 (Molecular Probes, Eugene, OR) in blocking solution. Following antibody labeling, the samples were washed in PBS or T/Ca/S and counterstained for 10 min with Calcofluor White M2R (fluorescent brightener 28; Sigma-Aldrich) at 0.25 mg ml⁻¹ in water. After washing with PBS or T/Ca/S, slides were sealed. As controls, sections were incubated without the primary antibodies and with Alexa Fluor 488-labeled secondary antibodies. Immunofluorescence was observed using a spectral confocal laser-scanning microscope (Zeiss LSM710) at 405-nm excitation and 410- to 490-nm emission for Calcofluor White M2R and at 488-nm excitation and 495- to 550-nm emission for Alexa Fluor 488.

XyG Analysis

XyG analysis was based on the rapid phenotyping method using enzymatic oligosaccharide fingerprinting, previously described (Lerouxel et al., 2002). After 3 d of stratification (T0), seeds were transferred to 25°C in continuous light. Embryos were separated from testa and endosperm by dissection after 3 h at 25°C and at ER. Samples were conserved in ethanol. After ethanol removal and rehydration, XyG oligosaccharides were generated by treating samples with endoglucanase in 50 mM sodium acetate buffer, pH 5, overnight at 37°C. And rehydration, XyG oligosaccharides were generated by treating samples with endoglucanase in 50 mM sodium acetate buffer, pH 5, overnight at 37°C. After washing with PBS or T/Ca/S, slides were sealed. As controls, sections were incubated without the primary antibodies and with Alexa Fluor 488-labeled secondary antibodies. Immunofluorescence was observed using a spectral confocal laser-scanning microscope (Zeiss LSM710) at 405-nm excitation and 410- to 490-nm emission for Calcofluor White M2R and at 488-nm excitation and 495- to 550-nm emission for Alexa Fluor 488.

Accession Numbers

The AGI locus identifier for XYL1 is At1g68560.

Supplemental Data

The following supplemental materials are available.

Supplemental Figure S1. Nomenclature and schematic representation of XyG.

Supplemental Figure S2. Germination rates of freshly harvested seeds of Arabidopsis wild type, xyl1-2, and xyl1-4.

Supplemental Figure S3. Silique length of xyl1, bgal10, bgv1, and xvt1 xvt2 mutants compared to wild type (WT).

Supplemental Figure S4. Immunolocalization of XyGs in Arabidopsis wild-type developing seed sections at 10 and 14 DAP.

Supplemental Figure S5. Immunolocalization of XyGs in Arabidopsis xyl1 developing seed sections at 10 DAP.

Supplemental Figure S6. Immunolocalization of XyGs in Arabidopsis xyl1 developing seed sections at 14 DAP.

Supplemental Figure S7. Immunolocalization of XyGs in Arabidopsis wild-type imbibed seed sections at 3 HAL.

Supplemental Figure S8. Immunolocalization of XyGs in Arabidopsis xyl1 imbibed seed sections at 3 HAL.

Supplemental Figure S9. ABA levels in seeds and imbibition medium.

Supplemental Table S1. XYL1 protein identification after 2D gel electrophoresis and peptide analysis by tandem mass spectrometry.

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LITERATURE CITED


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