Elucidating the roles of three β-glucuronosyltransferases (GLCATs) acting on arabinogalactan-proteins using a CRISPR-Cas9 multiplexing approach in Arabidopsis

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Abstract

Background: Arabinogalactan-proteins (AGPs) are one of the most complex protein families in the plant kingdom and are present in the cell walls of all land plants. AGPs are implicated in diverse biological processes such as plant growth, development, reproduction, and stress responses. AGPs are extensively glycosylated by the addition of type II arabinogalactan (AG) polysaccharides to hydroxyproline residues in their protein cores. Glucuronic acid (GlcA) is the only negatively charged sugar added to AGPs and the functions of GlcA residues on AGPs remain to be elucidated.

Results: Three members of the CAZy GT14 family (GLCAT14A-At5g39990, GLCAT14B-At5g15050, and GLCAT14C-At2g37585), which are responsible for transferring glucuronic acid (GlcA) to AGPs, were functionally characterized using a CRISPR/Cas9 gene editing approach in Arabidopsis. RNA seq and qRT-PCR data showed all three GLCAT genes were broadly expressed in different plant tissues, with GLCAT14A and GLCAT14B showing particularly high expression in the micropylar endosperm. Biochemical analysis of the AGPs from knock-out mutants of various glcat single, double, and triple mutants revealed that double and triple mutants generally had small increases of Ara and Gal and concomitant reductions of GlcA, particularly in the glcat14a glcat14b and glcat14a glcat14b glcat14c mutants. Moreover, AGPs isolated from all the glcat mutants displayed significant reductions in calcium binding compared to WT. Further phenotypic analyses found that the glcat14a glcat14b and glcat14a glcat14b glcat14c mutants exhibited significant delays in seed germination, reductions in root hair length, reductions in trichome branching, and accumulation of defective pollen grains. Additionally, both glcat14b glcat14c and glcat14a glcat14b glcat14c displayed significantly shorter siliques and reduced seed set. Finally, all higher-order mutants exhibited significant reductions in adherent seed coat mucilage.

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Background
Arabinogalactan-proteins (AGPs) are one of the most complex protein families in plants due to the diversity of the core proteins and the heterogeneity of the glycan chains attached to these protein cores. AGPs are present on the plant cell surface including the plasma membrane via GPI anchors, cell wall, and intercellular spaces of plants. AGPs are implicated in an array of plant growth and development processes including cell expansion, somatic embryogenesis, root and stem growth, salt tolerance, hormone signaling, programmed cell death, male and female gametophyte development, and wounding/defense [1, 2]. While the diversity of AGPs is manifested by the existence of different protein domains in the protein core, multiple glycans (AG chains) decorating each protein core, and the heterogeneity of glycosylation, they are unified by a common structure. Specifically, AGPs are O-linked glycoproteins characterized by the presence of type II (β-1,3 and β-1,6) AGs attached to hydroxyproline (Hyp) residues present in the protein core. More simply, the Hyp residues in AGP core proteins are modified by the addition of β-1,3-galactose sugar backbones, which are further modified by the addition of multiple, branching β-1,6 galactose side chains, which are further modified by the addition of arabinose (Ara) residues and less extensively by other sugars, such as rhamnose (Rha), fucose (Fuc), xylose (Xyl) and glucuronic acid (GlcA) [3, 4].

Several glycosyltransferases (GTs), each of which is responsible for transferring individual monosaccharides to synthesize the AGP glycans, are localized mainly in the Golgi. They were isolated and identified using both biochemical and bioinformatics approaches. Previously, our lab discovered five β-1,3-galactosyltransferases (GALT2, GALT3, GALT4, GALT5, and GALT6) [5, 6]. Three additional GALTs were subsequently discovered namely HPGT1, HPGT2, and HPGT3 [7]. All eight enzymes are from the Carbohydrate Active Enzyme (CAZy) GT31 family and function in catalyzing the addition of the first sugar, galactose, onto hydroxyproline residues contained in the AGP core proteins in Arabidopsis. Single genetic mutants of the GALT family showed subtle or no ontotypic changes compared to wild type (control) plants. However, the hpgt triple mutants exhibited longer root hairs, smaller leaves, and reductions in plant height and seed set [7]. In contrast, the galt2 galt5 double mutant demonstrated phenotypes such as shorter root length and root swelling phenotype under salt stress condition, as well as a reduction in seed coat mucilage [8]. Another β-1,3-galactosyltransferase named KNS4/GALT14, which is also found in the CAZy GT31 family, was recently characterized in Arabidopsis [9]. Mutant analyses found that ksn4 had an irregular and collapsed pollen phenotype, which was due to abnormal formation of the exine layer in the developing microspore [9]. AtGALT29A and AtGALT31A from the CAZy GT29 and GT31 families, respectively, represent two β-1,6-galactosyltransferases that were also identified in Arabidopsis [10]. In addition, a β-arabinofuranosyltransferase, RAY1, was also characterized in Arabidopsis. The ray1 mutant showed a reduction of arabinofuranose (Araf) in its AGPs [11]. Although heterologous expression of RAY1 in tobacco microsomes demonstrated β-Araf activity, β-1,3-linked Araf has not been found in plants. Moreover, two α-1,2–fucosyltransferases, AtFUT4 and AtFUT6, were also identified through heterologous expression in tobacco BY2 cells [12]. AtFUT4 and AtFUT6 were able to catalyze the transfer of fucose to AGPs in vitro [12]. Further research found that double mutants of fut4 fut6 contained no fucose and exhibited reduced root length under salt stress conditions compared to wild-type (WT) [13]. Finally, three β-glucuronosyltransferases (GLCATs) in the CAZy GT14 family, GLCAT14A, GLCAT14B, and GLCAT14C were identified through co-expression analyses with AGPs in Arabidopsis and verified by heterologous expression and an in vitro enzyme assay [14, 15]. So far only GLCAT14A has been functionally characterized to some extent by the use of mutants [14]; however, no such work has been done for GLCAT14B and GLCAT14C.

Due to gene redundancy among the GT families, a comprehensive understanding of the functions of the diverse sugars decorating the AGP protein core requires the generation of higher-order mutants in order to substantially reduce or eliminate addition of a particular sugar residue and observe its functional consequences. The traditional (T-DNA based) method of generating multiple gene knockout mutants is both labor-intensive and time-consuming. Moreover, genetic linkage, which is commonly seen in GTs and cell wall families, also...
makes it more challenging to generate higher-order mutants by conventional crossing method.

Consequently, we have chosen to utilize CRISPR-Cas9 gene editing technology to address this issue of gene redundancy in the GT families. CRISPR-Cas9 allows for the production of mutations in multiple genes in a single transformation event by the design of multiple guide RNAs (gRNAs) targeting multiple GT genes. Here we report on the successful use of CRISPR-Cas9 gene editing technology to target three GLCAT genes, known to be involved in the transfer of GlcA residues to AGPs as a test case. The resulting mutants not only demonstrate the feasibility of this approach, but also provide new evidence for the function of GlcA residues in AGPs with respect to plant growth and development.

Results
GLCAT14A, GLCAT14B, and GLCAT14C belong to the CAZY GT14 family
This study focused on functional characterization of GLCAT14A (At5g39990), GLCAT14B (At5g15050), and GLCAT14C (At2g37585) found in the CAZY GT14 family, which includes 11 proteins in Arabidopsis. The GT14 family was identified by a branch domain (Pfam 02485), which is named as a GLCAT domain [16]. Following heterologous expression of these 11 Arabidopsis GT14 members in *Pichia pastoris*, only three members (GLCAT14A, GLCAT14B, and GLCAT14C) exhibited GlcA transferase activity when using UDP-[14C]-GlcA as the sugar donor and AGP substrate acceptors [14, 15]. While all three enzymes demonstrated GlcA transferase activity with β-1,3 galactan and β-1,6 galactan AGP substrate acceptors, GLCAT14A and GLCAT14B preferred the former acceptor, while GLCAT14C preferred the later acceptor [14, 15]. Furthermore, co-expression analysis by GeneCAT found that GLCAT14A was co-expressed with AtGALT31A [14, 17].

Among the three GLCATs, GLCAT14A and GLCAT14B share 73.3% amino acid similarity, but only share 43.4 and 48.3% amino acid similarity with GLCAT14C, respectively (Additional file 1: Figure S1). GLCAT14A and GLCAT14B are both located on chromosome 5, whereas GLCAT14C is found on chromosome 2.

Expression profiles of GLCAT14A, GLCAT14B, and GLCAT14C
Our qRT-PCR and publicly available RNA-seq data from the Klepikova Atlas [18] were used to examine expression levels of the three GLCATs (Fig. 1, Additional file 1: Figure S2). Both the qRT-PCR and RNA-seq data showed that GLCAT14A and GLCAT14C were expressed at a higher level than GLCAT14B in most tissues. In addition, qRT-PCR results showed that all three GLCATs exhibited high expression in 36 h water-imbibed seeds and siliques than other tissues. GLCAT14B also showed a high expression in stems. Publicly available RNA-seq data additionally showed that GLCAT14A exhibited higher expression in seedling roots, leaf petioles, and siliques, GLCAT14B was mostly expressed in mature leaves, internodes, and leaf petioles, and GLCAT14C was highly expressed in flowers, seeds and siliques. The high expression values for both GLCAT14A and GLCAT14C in siliques and for GLCAT14B in internodes based on RNA-seq data were consistent with our qRT-PCR results. In addition, transcriptome data from developing seeds as displayed in the eFP browser [19] found that GLCAT14A and GLCAT14B were very highly expressed in the micropylar endosperm at the preglobular and globular stages of seed development in contrast to GLCAT14C which was not expressed at this location (Additional file 1: Figure S3). Both GLCAT14A and GLCAT14C also had moderate expression in the seed coat, whereas GLCAT14B was exclusively found in the micropylar endosperm. Given the high expression levels in micropylar endosperm for GLCAT14A and GLCAT14B, and in water-imbibed seeds for GLCAT14A and GLCAT14C as shown by qRT-PCR, the three GLCATs were speculated to play a role in seed development and germination.

Generation of GLCAT higher-order mutants using a CRISPR-Cas9 multiplexing approach
To date, only GLCAT14A in the Arabidopsis GT14 family was subjected to mutant analysis and partial characterization. Two allelic T-DNA insertion mutants for GLCAT14A, named glcat14a-1 and glcat14a-2, were obtained and showed a reduced level of β-GLCAT
activity as well as a 12% increase of Gal and 12% decrease of Ara in seedling AGPs. Physiologically, both allelic mutants displayed a 30% increase in root length and a 30% increase of hypocotyl length when grown in the dark [14]. However, as GLCAT14A and GLCAT14B likely redundant, a single genetic mutant such as glcat14a is often not sufficient to uncover fully the biological functions of GlcA residues. Furthermore, it is not ideal to use a traditional crossing method to create a double mutant for two genetically linked genes such as GLCAT14A and GLCAT14B. Therefore, this study utilized a multiplexing CRISPR-Cas9 gene editing strategy to create and examine higher-order knockout mutants for all three GLCATs simultaneously.

Six gRNAs (A1, A2, B1, C1, C2, and C3) were designed to target GLCAT14A, GLCAT14B, and GLCAT14C at different loci (Fig. 2A), and they were assembled into two CRISPR/Cas9 multiplexing constructs using the pHEE401E binary vector, which also contains a Zea mays codon-optimized Cas9 gene under the control of an Arabidopsis egg-cell specific promoter (E.C1.1) fused with an egg-cell specific enhancer (E.C1.2) [20]. Previous studies using pHEE401E successfully created higher-order homozygous CRISPR mutants in a single generation in Arabidopsis [21]. With our two CRISPR constructs, a number of single and double glcat were created, and we chose one Cas9-free line for each of the resulting mutants: glcat14b, glcat14c, glcat14a glcat14b, and glcat14b glcat14c for functional characterization. In addition, one glcat14a glcat14b glcat14c triple glcat CRISPR mutant line was identified and used for further characterization.

Three glcat homozygous mutants, namely glcat14b, glcat14a glcat14b, and glcat14a glcat14b glcat14c were generated in the T1 generation using the construct shown in Fig. 2B (1). The glcat14b mutant contained a 1 bp insertion at B1, while the glcat14a glcat14b mutant (line #17–5) contained 1 bp insertions at both A1 and B1 (Fig. 3). These mutations occurring in glcat14b and glcat14a glcat14b resulted in pre-mature stop codons for both genes. Cas9-free mutants were obtained for these two mutant lines in the T2 generation. Moreover, glcat14c and glcat14b glcat14c mutants were generated using the gene construct shown in Fig. 2B (2). Both mutant lines contained a 188 bp deletion between C2 and C3, while the glcat14b glcat14c double mutant line (line #14–4) additionally contained a 1 bp insertion at B1 (Fig. 4). Homozygous Cas9-free glcat14c and glcat14b glcat14c mutants were obtained in the T2 generation. Furthermore, one glcat14a glcat14b glcat14c triple mutant (line #52–4) was identified in the T2 generation using the construct in Fig. 2B (1). This mutant had a 12 bp deletion at A1, a 1 bp insertion at B2, and a 12 bp deletion at C1 (Fig. 5). In this triple mutant, gene editing events in gRNA A1 and B2 resulted in pre-mature stop codons, while the mutation at C1 caused the loss of four amino acids from its protein sequence (Fig. 5).

**Quantification of β-D-Gal-Yariv precipitated AGPs in the various glcat mutants**

β-D-Gal-Yariv is a synthetic dye that is often used for AGPs purification as it can bind to the β-1,3-galactose backbone on AGPs; thus, it is an ideal reagent to facilitate characterization of the glycan structure of AGPs [22]. Here, β-D-Gal-Yariv was used to precipitate AGPs from rosette leaf, stem, and silique tissues of various glcat mutants (Fig. 6). The single mutants and the glcat14b glcat14c double mutant did not show much difference in the AGP content except for glcat14c, which showed an ~25% increase in AGP content only in silique but not in other organs. The glcat14a glcat14b double mutant and the glcat14a glcat14b glcat14c triple mutant, however, showed increases in AGP content in all three organs examined, namely rosette leaf, stem, and silique. Specifically, glcat14a glcat14b showed ~60%, ~50%, and ~25% increases in AGP content in rosettes, stems, and silique, respectively; whereas glcat14a glcat14b glcat14c showed ~50%, increases in AGP content in all three organs.

**Monosaccharide composition analyses**

Previous monosaccharide composition analysis of glcat14a from 14-day-old seedlings found that there was a slight increase of Gal in 3-, 6- and 3,6-linkages in the AGPs extracted from glcat14a compared to that of WT, while the amount of GlcA in glcat14a was similar to that of WT [14]. Here, monosaccharide composition analyses were done by HPAGEC-PAD for AGPs extracted from 40-day-old glcat mutants and WT. As seen in Table 1, single glcat mutants did not show notable differences in the amounts of most sugars, including GlcA, compared to WT with the exception of glcat14c, which showed an elevated amount of Gal and a reduced amount of GlcA. In contrast, the higher-order glcat mutants, namely glcat14a glcat14b, glcat14b glcat14c, and glcat14a glcat14b glcat14c, exhibited 27, 12, and 40% reductions in the amount of GlcA, respectively. Interestingly, these higher order mutants also displayed increased amounts of Ara and Gal compared to WT. Furthermore, the glcat14a glcat14b glcat14c mutant exhibited more pronounced reductions of GlcA than the double mutants; nonetheless, the GlcA content in the triple mutant was not completely eliminated.

Previous research detected calcium associated with certain AGPs, including gum arabic and AGPs isolated from cell cultures of Arabidopsis, broccoli, carrot, tobacco, and tomato [23]. Lamport hypothesized that the negatively charged GlcA residues on AGPs may serve as sites for such calcium binding [24]. To examine
whether calcium binding by AGPs is altered in our glcat mutants, we extracted AGPs from the aerial portion of Arabidopsis plants and measured the amount of associated calcium using a colorimetric assay [25]. As shown in Fig. 7, WT Arabidopsis contained 1.5% Ca\(^{2+}\) per \(\mu\)g AGP, whereas all the glcat mutants contained 0.5% Ca\(^{2+}\) per \(\mu\)g AGP. Interestingly, the single glcat mutants had reductions in calcium binding comparable to the higher-order mutants. These results indicate that GlcA on AGPs is responsible for binding calcium and loss of any of the three GLCATs resulted in major loss of calcium binding of AGPs.

glcat14a glcat14b and glcat14a glcat14b glcat14c exhibited delayed germination
A significant delay in germination was observed in both post-harvested (> 3 months) and freshly-harvested seeds of glcat14a glcat14b and glcat14a glcat14b glcat14c seeds. Only ~ 40% of glcat14a glcat14b and ~ 30% of glcat14a glcat14b glcat14c seeds germinated by the 2nd day after sowing compared to ~ 75% seed germination for the other genotypes (Fig. 8a). By the 3rd day most of the WT, glcat14b, glcat14c, and glcat14b glcat14c seeds germinated, while only ~ 60% and ~ 55% of glcat14a glcat14b and glcat14a glcat14b glcat14c seeds respectively germinated. Overall, the germination time of N 50 (50% germination rate) for most glcat mutants and WT was ~ 1.5 days compared to 2.5 days to 3 days for the glcat14a glcat14b and glcat14a glcat14b glcat14c mutants, respectively. This delayed germination phenotype for these glcat mutants, however, did not affect their ultimate ability to germinate, as most seeds eventually germinated (Fig. 8a).
Given that GLCAT14A and GLCAT14B are both highly expressed in the micropylar endosperm and the weakening of the endosperm cell wall is negatively regulated (i.e., delayed) by plant hormones such as ABA, we germinated glcat mutant seeds in the presence of 0.5, 1, and 2 μM ABA. The germination of WT seeds shared a similar trend to most glcat mutants; all of them germinated faster than glcat14a glcat14b and glcat14a glcat14b glcat14c in the presence of 0.5 μM ABA (Fig. 8b). In the presence of 1 μM ABA, glcat14c and glcat14b glcat14c germinated later than WT along with a more severe delay of germination observed for glcat14a glcat14b and glcat14a glcat14b glcat14c (Fig. 8c). In the presence of 2 μM ABA, all the glcat mutants demonstrated delayed germination compared to WT (Fig. 8d).

glcat14a glcat14b and glcat14a glcat14b glcat14c are hypersensitive to stratification conditions

Stratification at cold temperature is a common strategy to release dormancy of Arabidopsis seeds. Stratification can especially speed up the germination process of freshly harvested seeds, which contain a higher ABA content than post-harvested (> 3 months) seeds. To investigate whether the delayed germination phenotype observed for glcat14a glcat14b and glcat14a glcat14b glcat14c is related to stratification conditions, freshly harvested seeds from the glcat mutants and WT were treated with normal stratification (water imbibition for 72 h, 4 °C), stratification at room temperature (water imbibition for 72 h, 22 °C), and no stratification (no imbibition) before sowing. Germination rates were then measured 48 h after sowing. While stratification at room temperature and no stratification reduced germination percentages in all seeds compared to normal stratification, the glcat14a glcat14b and glcat14a glcat14b glcat14c seeds were especially sensitive to these two treatments and exhibited significant delays in germination (Figs. 9 and 10). These experiments indicate that the GLCATs, especially GLCAT14A and GLCAT14B together, play a role in the transition of dormant seeds to non-dormant seeds and is promoted by cold stratification.
Higher-order glcat mutants showed less trichome branching, reduced root hair length, and reduced plant height

Mutants that have both GLCAT14A and GLCAT14B knocked out also showed defects in trichome branching and reduction in root hair growth. Both gcat14a gcat14b and gcat14a gcat14b gcat14c had smaller trichomes and reduced trichome branching compared to WT, whereas single gcat mutants and gcat14b gcat14c phenocopied WT in trichome development (Fig. 11a). Moreover, root hair length was reduced in gcat14b, gcat14a gcat14b, and gcat14a gcat14b gcat14c compared to WT (Fig. 11b and d). Furthermore, the gcat double and triple mutants demonstrated reductions in plant height (Fig. 11c). The above mutant phenotypes indicate that GlcA residues on AGPs likely contribute to cell differentiation and tip-focused growth.

gcat14a gcat14b and gcat14a gcat14b gcat14c contain defective pollen

In vitro pollen germination experiments were done to examine pollen phenotypes of the various gcat mutants. Only gcat14a gcat14b gcat14c exhibited a lower pollen germination rate compared to WT, whereas the pollen tube lengths of all gcat mutants were not significantly different from WT (Fig. 12a and b). In addition, gcat14a gcat14b and gcat14a gcat14b gcat14c mutants exhibited a significant number of defective pollen (Fig. 12c). The defective pollen was much smaller in size compared to normal pollen and failed to germinate (Fig. 13).

Disruptions of the GLCATs resulted in shorter silique lengths and reduced seed set

Only the gcat14b gcat14c and gcat14a gcat14b gcat14c mutants exhibited shorter silique lengths and reductions in seed set compared to WT (Fig. 14). GLCAT14C showed the highest expression in silique with GLCAT14A and GLCAT14B showing more modest expression based on our RT-qPCR data (Fig. 1).

Higher-order gcat mutants had less adherent seed coat mucilage

Another notable phenotype of gcat mutants is the significant reduction in adherent seed coat mucilage. Seeds of gcat14a gcat14b, gcat14b gcat14c, and gcat14a gcat14b gcat14c showed dramatic reductions in...
Fig. 5 Sequencing results of three target sites (A1, B1, and C1) in the glcat14a glcat14b glcat14c triple mutant (line #52–4). GLCAT14A contained a 12 bp deletion, which resulted in pre-mature stop codon; GLCAT14B contained a 1 bp insertion, which resulted in a pre-mature stop codon in its protein sequence; GLCAT14C contained a 12 bp deletion, which resulted in the loss of four amino acids (red). Sequences of the mutation sites are indicated by red arrows on the chromatograms on the right.

Fig. 6 Quantification of AGP content of the various glcat mutants in different organs of 40-day-old plants. Single mutants did not show much change in their AGPs across different tissues except for glcat14c that contained more AGPs in the siliques. Both glcat14a glcat14b and glcat14a glcat14b glcat14c showed increases in AGP content in rosette leaves, stems, and siliques compared to other single mutants and WT across tissues, AGPs were measured as microgram per gram of fresh weight. Statistical differences were determined by two-way ANOVA, followed by the Tukey’s honestly significant difference test (* P < 0.05; ** P < 0.01; *** P < 0.001).
adherent seed mucilage compared to WT as determined by staining with ruthenium red staining, which binds to acidic biopolymers such as pectin (Fig. 15).

Discussion
CRISPR-Cas9 approach to generate higher-order GLCAT mutants

Gene redundancy is common for genes encoding enzymes that function in cell wall biosynthesis including glycotransferases (GTs). In order to better understand the function of redundant gene families, higher order knockout mutants, mutants in which multiple genes in the family are inactivated, need to be generated and studied. Traditional genetic crossing/breeding approaches to obtain higher order mutants, however, are both time-consuming and complicated by genetic linkage. Compared to traditional breeding, CRISPR-Cas9 gene editing has been recently used to study redundant gene families in several plant species such as rice, wheat, cabbage, tomato and Arabidopsis [21, 26–29]. The research reported here, however, represents the first application of CRISPR/Cas9 to understand GTs families in Arabidopsis. Here, we show that CRISPR/Cas9 gene editing was successfully used to create both single glec14 mutants as well as higher order glec14 mutants. Without this approach, it would have been extremely difficult to generate glec14a glec14b and higher order mutants, as the two genes are approximately 12 cM apart on chromosome 5. As discussed below, the single glec mutants (glec14b and glec14c) largely phenocopied WT and did not show changes in the monosaccharide composition on their AGPs, whereas disruption of two or more GLCATs had effects on both the AGP sugar composition as well as several significant developmental/physiological phenotypes in Arabidopsis.

Two multiplexed CRISPR constructs (Fig. 2B) were created to target the three known GLCAT gene family members, namely, GLCAT14A, GLCAT14B, and GLCAT14C. Following transformation with these constructs, a number of single, double, and triple glec mutants were generated. Indel mutations created in glec14b, glec14c, glec14a glec14b, and glec14b glec14c all resulted in pre-mature stop codons (Figs. 3, 4). In the glec14a glec14b glec14c triple mutant, mutations in GLCAT14A and GLCAT14B resulted in pre-mature stop codons, whereas mutation in GLCAT14C caused the loss of four amino acids in the GLCAT Pfam domain (Pf02485) (Fig. 5).

To evaluate potential off-target mutations caused by CRISPR-Cas9, CRISPR-P 2.0 software [30] was used to

**Table 1** Monosaccharide composition analysis of AGPs extracted from aerial part of 40-day-old glec CRISPR mutants and WT (Col-0)*

|                | Fuc        | Rha        | Ara        | Gal        | Glu        | Xyl        | Man        | GalA       | GlcA       |
|----------------|------------|------------|------------|------------|------------|------------|------------|------------|------------|
| WT             | 2.8 ± 0.7  | 4.8 ± 1.0  | 26.3 ± 2.4 | 39.1 ± 0.9 | 4.2 ± 0.3  | 7.0 ± 1.4  | 3.1 ± 0.6  | 5.0 ± 0.8  | 6.0 ± 1.3  |
| glec14b        | 2.7 ± 0.4  | 5.1 ± 0.6  | 24.6 ± 0.9 | 38.2 ± 0.4 | 3.7 ± 2.2  | 8.5 ± 1.2  | 3.3 ± 0.6  | 7.6 ± 1.4  | 6.1 ± 0.9  |
| glec14c        | 2.1 ± 0.4  | 4.4 ± 0.7  | 25.7 ± 2.1 | 41.5 ± 0.1 | 4.5 ± 0.2  | 6.7 ± 1.0  | 2.8 ± 0.4  | 7.2 ± 0.5  | 5.4 ± 0.9  |
| glec14a glec14b| 2.2 ± 0.2  | 3.4 ± 0.5  | 30.5 ± 3.8 | 42.3 ± 1.0 | 3.2 ± 1.0  | 5.6 ± 0.4  | 2.8 ± 0.2  | 4.5 ± 0.0  | 4.4 ± 0.4  |
| glec14b glec14c| 2.4 ± 0.2  | 4.1 ± 1.0  | 27.5 ± 0.4 | 44.7 ± 0.4 | 4.1 ± 0.1  | 7.0 ± 0.2  | 2.8 ± 0.0  | 3.8 ± 0.2  | 5.3 ± 0.4  |
| glec14a glec14b glec14c | 2.8 ± 0.4 | 4.0 ± 1.2  | 30.7 ± 1.6 | 41.2 ± 1.0 | 3.6 ± 0.3  | 7.0 ± 0.7  | 2.7 ± 0.5  | 4.1 ± 1.3  | 3.6 ± 1.2  |

*Values are relative to total sugar composition (expressed as mol %) of triplicate assays ± SE

Fig. 7 Quantification of calcium content of AGPs extracted from the aerial portion of 40-day-old plants. All the glec mutants showed significant reductions of calcium binding. Statistical differences were determined by one-way ANOVA, followed by the Tukey’s honestly significant difference test (*** P < 0.001)
identify off-target candidates. Four sites located in the coding regions and showed off-target scores between 0.1 and 0.2 were identified (Additional file 1: Table S4). These sites were amplified and sequenced in the mutants. No off-target mutations were identified in any of our glcat mutants.

CRISPR-induced GLCAT mutations alter AGP sugar composition and calcium binding

AGP monosaccharide composition analyses of mature plants showed that glcat14a glcat14b, glcat14b glcat14c, and glcat14a glcat14b glcat14c exhibited approximately 27, 12, and 40% reductions of GlcA, respectively (Table 1). In addition, glcat14c, glcat14a glcat14b, glcat14b glcat14c, and glcat14a glcat14b glcat14c exhibited more Gal (~5–12%) and more Ara (~15%) compared to WT. This is in general agreement with previous work on a glcat14a T-DNA insertion mutant, which detected 12% more Gal and 12% less Ara in AGPs isolated from young seedlings, although differences in GlcA levels were not detected by monosaccharide composition analysis this study [14]. Our findings are also consistent with the suggestion that GlcA residues on AGPs may serve to terminate further elongation of β-1, 3- and β-1,6-galactan sugar additions to AGPs [14], such that reduced levels of GlcA lead to additional elongation and incorporation of Gal and Ara residues (Table 1).

Knocking out any of the GLCAT genes also led to significantly less calcium binding by AGPs compared to WT (Fig. 7), suggesting that GlcA, as the only negatively charged sugar on AGPs, plays a key role in calcium binding by AGPs. This provides the first direct experimental
In order to examine whether the amount of glycosylated AGPs was altered in the various organs of the glcat mutants, we used β-Gal Yariv reagent to quantitate glycosylated AGPs from rosette leaves, stems, and siliques (Fig. 6). Both the glcat14a glcat14b and the glcat14a glcat14b glcat14c triple mutant exhibited ~30–60% increases of AGPs from rosettes, stems and siliques.
compared to WT. In addition, the glcat14c single mutant exhibited a ~25% increase of AGPs in the siliques. Such increases in AGP content are consistent with the idea that the loss of GlcA residues results in the elongation of β-1,3-galactose backbone and the addition of β-1,6-galactose side chains in AGPs, as revealed by β-Gal Yariv reagent and corroborated by monosaccharide analysis (Table 1). These results indicate that GLCAT14A and GLCAT14B play redundant roles in catalyzing the transfer of GlcA to AGPs in all three organs.
GLCAT14A and GLCAT14B function redundantly in regulating seed germination

Among the glcat mutants, only glcat14a glcat14b and glcat14a glcat14b glcat14c exhibited 1–1.5 day delays in seed germination compared to WT (Figs. 8 and 9). Such delays in germination were exacerbated in seeds without normal (cold) stratification. This indicates that GLCAT14A and GLCAT14B function redundantly in regulating seed germination and such regulation most likely involves enhancing the degradation of the cell wall in the micropylar endosperm, where both genes are highly expressed. Thus, cold temperature imbibition likely loosens the micropylar endosperm cell wall to facilitate germination. A number of other cell wall enzymes were shown to be involved in seed germination such as expansin, β-1, 3-glucanase, endo-β-1,4 mannanase, pectin methyltransferase, endotransglycosylase/hydrolases (XTH), α-xilosidase, and xylosyltransferases, all of which are also highly expressed and many of which are specifically expressed in the micropylar endosperm [31–36]. These enzymes likely play key roles in seed germination, as supported by expression analysis and by genetic mutant analysis. For example, knockout mutants of genes encoding endo-β-1,4 mannanase including man5, man6, and man7 all exhibited a delayed germination phenotype in Arabidopsis. Moreover, a man7 homolog also accumulated in the micropylar endosperm in Lepidium sativum, a close Arabidopsis relative [37].

In addition to the germination delays under normal conditions, all the glcat mutants also demonstrated even greater delays in the presence of 1–2 μM ABA. ABA inhibits only the 2nd phase of germination, which involves radical protrusion. The observed germination delay exhibited here by the glcat mutants in response to ABA indicates that GlcA residues on AGPs may function as signaling molecules in such ABA-mediated germination.
GLCAs involved in vegetative tip-focused growth in Arabidopsis

Among the glcat mutants, both glcat14a glcat14b and glcat14a glcat14b glcat14c displayed reductions in root hair length, trichome size, and trichome branching (Fig. 11). The role of AGPs in cell growth and expansion has long been shown by applying β-Gal-Yariv reagent to plant culture and observe growth inhibition [38]. β-Gal-Yariv, which specifically binds to the β, 1,3-galactose sugar backbone on AGPs, can inhibit root cell expansion and proliferation in vitro [39]. The galt2 galt5 knockout mutant, affecting two GALTs involved in adding the first galactose residues to hydroxyproline residues on AGPs, also demonstrated reductions in root hair length [8]. Root hair and trichome formation are two examples of tip-focused growth driven by a cytoplasmic calcium gradient generated at the tip [40]. Recent studies found that AGPs can bind and release calcium in a pH-dependent fashion, presumably by their negatively charged GlcA residues [23]. Consequently, it was suggested that AGPs serve as extracellular calcium capacitors and are involved in calcium-mediated signaling, such that lower pH levels generated by actin mediated H⁺-ATPase disassociates calcium from extracellular AGPs and releases it into the cytosol, generating a cytoplasmic calcium influx [24]. This process may thus be important for various developmental processes, including root hair and trichome growth. A calcium-binding assay performed here for glcat14a glcat14b and glcat14a glcat14b glcat14c indeed found significant less calcium binding to AGPs compared to WT (Fig. 7). Therefore, the shorter root hair lengths and deformed trichome growth observed in our glcat mutants may be caused by reduced calcium influx in these tissues as brought about by the reduction of GlcA levels, and hence calcium levels, in the extracellular AGPs in these tissues.

There are other potential mechanisms of action for AGPs that could account for impaired root tip growth observed in the glcat mutants, as well as for the other mutant phenotypes reported in this study. For example, AGPs may control growth by interacting with other cell wall components. AGPs are often associated with pectin and in some cases covalently interact with them [41]. Studies on SOS5/FLA4, which is a chimeric AGP, found that sos5 showed significant reduction in the seed mucilage content, and suggested that SOS5 interacts with and mediates pectin network assembly in seed mucilage [42]. In another study, AGPs were shown not only to covalently interact with pectin, but also to covalently interact with arabinoxylan (hemicellulose) [43]. More precisely, this study demonstrated that GlcA residues in AGPs were covalently linked to Rha residues in pectin [43]. Such structural interactions could thus be involved in controlling cell wall rigidity and extensibility. Another recent study on UPEX1, which is an AGP-specific glycosyltransferase in the GT31 family, also suggested that AGPs and xylan interact during exine cell wall deposition during pollen development [44]. Similar to the idea that un-methyl esterified pectin can form “egg-box” structures through Ca²⁺ binding that has been known to control cell wall integrity, the recently proposed AGP-Ca²⁺ model makes the negatively charged GlcA residues...
other potential Ca$^{2+}$ binding sites on the cell surface to regulate cell wall integrity.

AGPs may also act as sensors and/or signaling molecules. The presence of AGPs at the plasma membrane/cell wall interface by virtue of the GPI anchoring domains present on many AGPs position them at an ideal location for sensing cell wall perturbations and transmitting information to associated signaling molecules such as receptor-like kinases. Physical interaction between AGPs and receptor like kinases (RLKs) is likely; this is based largely on genetic data. Specifically, sos5/fla4 mutants and two Leu-rich repeat RLKs mutants, fei1 and fei2, exhibited similar root swollen phenotypes as did sos5 fei1 fei2 triple mutants, indicating nonadditive genetic interaction [45]. Thus, it was proposed that SOS5 is in the same genetic pathway as FEI1 and FEI2 and that SOS5 is a ligand for FEI1 and FEI2.

**GLCATs contribute to reproductive growth in Arabidopsis**

Phenotypic analyses of the glcat mutants also indicates a role for the GLCATs in plant reproduction. Both glcat14a glcat14b and glcat14a glcat14b glcat14c displayed significant percentages of defective pollen that are much smaller than normal pollen and fail to germinate and glcat14a glcat14b glcat14c also exhibited a significantly lower pollen germination rate (Figs. 12 and 13). Furthermore, glcat14b glcat14c and glcat14a glcat14b glcat14c exhibited a reduction in si- lique length and seed set (Figs. 14). These observations relate to previous studies on AGPs. One of which found that agp6 single mutants, agp11 single mutants, and agp6 agp11 double mutants display reduced pollen germination, pollen tube elongation, and pollen release [46]. The other study showed that AGP4, which is only expressed in the inflorescence, functions to prevent multiple pollen tubes from entering into one embryo sac in Arabidopsis, as agp4 loss of function mutants failed to block multiple pollen tubes from entering the embryo sac [47]. Recent studies in *Torenia fournieri* identified a disaccharide sugar, β-methyl-glucuronosyl galactose (4-Me-GlcA-β-1,6-Gal) present on an AGPs secreted from the ovary that makes pollen tubes competent for ovule targeting/guidance [48]. Thus, GlcA residues on AGPs could play roles in ovule guidance and/or polytubey in Arabidopsis, but further research is needed to support these potential roles. As the only negatively charged sugar on AGPs, GlcA is likely responsible for calcium-binding by AGPs. Intracellular calcium signatures were detected at every step of the double fertilization process in Arabidopsis [49]. As such, GlcA residues on AGPs could function in fertilization via calcium signaling.

**GLCATs play an essential role in seed coat mucilage extrusion**

Seed coat mucilage represents a useful model for studying genes involved in cell wall biosynthesis. As much as 90% of this mucilage is composed of pectin polysaccharides and less than 10% of the mucilage is made up of other cell wall component such as cellulose, galactoglucomannan, xylaglucan, xylan, and AGPs [50]. Disruptions of genes involved in cell wall biosynthesis often results in a reduction or loss of the seed mucilage [51]. Knocking out two of the AGP-specific GALTs, galt2 galt5, also resulted in a reduction of seed mucilage [8]. Mutations of other cell wall proteins such as csla2, mum2, and fly1 also result in reductions of seed coat mucilage formation [52–54]. This study found that gcat double and triple mutants exhibited significant reductions in adherent seed mucilage as detected by ruthenium red staining (Fig. 15). There is insufficient information on role of GlcA in seed coat mucilage composition. In Arabidopsis, mucilage analyses either failed to detect or only detected less than 0.5 mol% GlcA [50, 51]. This study, for the first time, implicates that GlcA residues on AGPs play a substantial role in Arabidopsis seed coat mucilage formation as disruptions of two or more of the GLCATs resulted in a considerable loss of adherent seed mucilage. These GlcA residues, although minor components of the mucilage, may be important for covalent cross-linking with other cell wall components in the mucilage such as pectin, and specifically RG-1, as recently indicated by biochemical analysis of an extracellular AGP-pectin-arabinoxylan polymer, or alternatively, by ionic interactions with other negatively charged polymers (e.g., other AGPs or pectins) via calcium bridges [38, 40]. Reduction of GlcA on the AGPs may interfere with the normal structure of the primary cell wall thus impair the release or retention of the pectin-rich seed coat mucilage.

**Conclusions**

The pioneering CRISPR-Cas9 approach used in this study has proven to be an efficient and effective way to characterize higher-order knockout mutants of the three membered GLCAT gene family. Our biochemical and phenotypic analyses indicate that GLCAT14A and GLCAT14B play a particularly redundant role in adding GlcA to AGPs in Arabidopsis. Nonetheless, mutant analyses showed that in fact all three GLCATs shared some degree of redundancy, as double and particularly triple mutants of GLCATs resulted in more severe phenotypic defects than single mutants. All the GLCATs, by virtue of their ability to add GlcA residues, confer the ability of AGPs to bind calcium. The concept that intracellular calcium originates by the influx of calcium ions from the plant cell wall/extracellular matrix is often overlooked in favor of the release from intracellular reservoirs (ER, vacuole) [55]. It is hypothesized that AGPs can serve as pH-regulated calcium capacitors and the source for intracellular calcium signals required for many plant
growth and development processes [24]. Further studies will be required to examine potential calcium-related signaling pathways revealed by the glcat mutants. The observation that GlcA content was only reduced by ~50% on the AGPs in the glcat14a glcat14b glcat14c triple mutant indicates that additional GLCATs are likely involved in catalyzing the transfer of GlcA to AGPs. Thus, two important avenues of research to pursue in this area are: 1. Discovering which of the other members of the CAZy GT14 family may be responsible for adding GlcA residues to AGPs and 2. Understanding the exact mode or mechanism of action whereby GlcA residues and AGPs control specific developmental/physiological phenotypes, whether it be calcium-mediated signaling, altered cell wall structural interactions, and/or other AGP sensing/signaling scenarios.

Methods

Plant material

Arabidopsis thaliana (Columbia ecotype) was used as the WT, and it was originally obtained from the Arabidopsis Biological Research Center (ABRC), Columbus, Ohio, USA. We generated all the CRISPR mutants in this study using this Col-0 background.

In silico analysis of GLCAT14A, GLCAT14B, and GLCAT14C

Genomic DNA, protein sequences, and gene locations of GLCAT14A, GLCAT14B, and GLCAT14C were obtained from the TAIR website (https://www.arabidopsis.org/). Protein domain searches were done in Pfam (https://pfam.xfam.org/). Expression profiles of GLCATs based on RNA-seq data were obtained from the TRAVA database [18]. The transcriptomic data of seed development was obtained from the eFP browser (https://bar.utoronto.ca/efp/cgi-bin/efpWeb.cgi) [56].

Guide RNA design and vector construction

All gRNAs were selected using the online software CRISPR-P 2.0 (http://crispr.hzau.edu.cn/CRISPR2/). Several rules regarding sequence and structural requirements for choosing a gRNA were as described [57]. First, the gRNA should have a high on-target and low off-target score based on the algorithm of CRISPR-P 2.0. Also, the gRNA should contain no less than 40% GC content for strong binding to the template DNA and have a G or C at the position 20 before the PAM sequence. The vector construction strategy was adopted from a published method [21]. Three template plasmids (pCBCT-DT1, pCBCT-DT2DT3, and pCBCT-DT3DT4) and a binary vector (pHEE401E) were gifts from Dr. Qi-Jun Chen’s lab (Addgene plasmid #50590; #50591; #50592; #71287). Each CRISPR/Cas9 multiplexing construct, consisting of an individual gRNA cassette, which includes a gRNA sequence, a gRNA scaffold, an AtU6 promoter, and an AtU6 terminator, was first amplified from a specific template plasmid; see Additional file 1: Tables S1-S3 for specific gRNA and primer sequences. A detailed procedure for primer design and PCR reaction steps were described previously [21]. For the CRISPR/Cas9 construct containing three gRNAs: A1, B1, and C1, the gRNA A1 cassette was amplified using primers DT1-A1_F, DT1-A1_F0, and DT0-R2; while the B1 and C1 cassettes were amplified using primers DT2-B1_F, DT2-B1_F0, DT3-C1_R0, and DT3-C1_R. After amplifying each gRNA fragment, they were cloned into BsaI site of the pHEE401E binary vector by the Golden Gate cloning method [58]. For the CRISPR/Cas9 construct containing four gRNAs: A1, B1, C2, and C3, the gRNA C3 cassette was amplified using primers DT1-C3_F, DT1-C3_F0, and DT0_R2; the gRNA A2 cassette was amplified using primers DT2-A2_F2, DT2-A2_F0, and DT0-BsR3; the gRNA B1 and C2 cassettes were amplified using primers DT3-BsF3, DT3-B1-F0, DT4-C2-R0, and DT4-C2-BsR. The four gRNA fragments were then cloned into the BsaI site of the pHEE401E binary vector by the Golden Gate cloning method. The pHEE401E binary vector contains a maize codon optimized Cas9 gene driven by egg cell-specific promoters (E.C.1.1 and E.C.1.2) [20]. The final constructs were transformed into Agrobacterium strain GV3101, which was then used for transforming Arabidopsis Col-0 ecotype plants via the floral dip method [59]. A PCR-based method was used for initially detecting indel mutations, which were subsequently confirmed by Sanger sequencing [60].

Quantitative RT-PCR

Total RNA was extracted from various tissues (imbibed seeds, germinated seeds, roots, rosettes, stems, open flowers, siliques) of Col-0 Arabidopsis using Trizol (Life Technologies, Grand Island, NY, USA). RNA (1 μg) was used for first-strand cDNA synthesis along with an oligo (dT20) primer and SuperScript III reverse transcriptase (Thermo Scientific). The qPCR was performed using PerfeCTa SYBR Green SuperMix (Quanta Biosciences). A 20 μl reaction mixture was set up which contained 10 μl PerfeCTa SYBR Green SuperMix, 4 μl 10x diluted cDNA, and 0.6 μl forward/reverse primer (10 μM). The qPCR was run on an AriaMx Real-time PCR machine (Ohio University Genomics Facility: http://www.dna.ohio.edu/). PCR conditions were according to the PerfeCTa SYBR Green SuperMix protocol. Expression levels were calculated relative to the Arabidopsis ACTIN2 gene. The qPCR primers are listed in Additional file 1: Table S5.

AGPs quantification by β-D-gal-Yariv

AGPs were extracted from 40-day-old glcat mutants and WT using the β-D-Gal-Yariv precipitation method
described by Lamport [61]. Briefly, tissues were harvested from rosette leaves, stems, and siliques of 40-day-old glcat and WT plants and pulverized using a mortar and pestle in the presence on liquid nitrogen. Approximately 0.3 g of pulverized tissues were mixed with 1 mL 2% CaCl₂ and shaken at 150 rpm for 2 h. The ground tissues were then separated from the supernatant by centrifugation at 13,000 g for 10 min. Around 500 μL of supernatant was transferred to a new 1.5 mL centrifuge tube and mixed with 200 μL β-D-Gal-Yariv dissolved in 2% CaCl₂ (1 mg/mL). Meanwhile, 500 μL 2% CaCl₂ was mixed with 200 μL β-D-Gal-Yariv as control. After 2 h of β-D-Gal-Yariv precipitation, the pellet was obtained by centrifugation at 13,000 g for 10 min and then dissolved in 20 mM NaOH. The dissolved AGPs were quantified by measuring absorbance at OD₅₂₀. Different concentrations of gum arabic (G9752, Sigma-Aldrich, St. Louis, MO, USA) were used for making the standard sugar mixture (fucose, rhamnose, arabinose, galactose, glucose, xylose, mannose, galacturonic acid, and glucoronic acid) of various concentrations (5 nM, 2.5 nM, and 1.25 nM) were used for monosaccharide composition analysis by high performance anion exchange chromatography with pulsed amperometric detection (HPAEC-PAD)

AGPs were extracted from the aerial part of 40-day-old glcat mutants and WT using the protocol developed by Lamport [61]. Pulverized plant tissue was mixed with 2% NaCl in 1:4 (w/v) ratio and shaken at 200 rpm for 3 h followed by centrifugation at 13,000 g for 30 min. Then 2 mL of 1 mg/mL β-D-Gal-Yariv reagent was mixed with the supernatant and allowed it to precipitate overnight. The next day, the precipitated AGPs were collected by centrifugation at 2000 g for 10 min, washed with 2% NaCl once, then resuspended in 2 mL H₂O. The β-D-Gal-Yariv reagent was dissociated from the AGPs by adding ~25 mg sodium dithionite to each 2 mL AGPs sample and incubated at 50 °C for 15 min until the solution became yellow. A PD-10 column (GE Healthcare) was used for desalting, and the eluate was freeze-dried. Approximately 100 μg of AGPs dissolved in Milli-Q water were used for monosaccharide composition analysis. AGPs were hydrolyzed using 300 μL 2 N TFA at 121 °C for 90 min followed by removal of TFA using N₂ gas. The sample was washed with isopropanol three times before dissolving it in 500 μL Milli-Q water. A standard sugar mixture (fucose, rhamnose, arabinose, galactose, glucose, xylose, mannose, galacturonic acid, and glucoronic acid) of various concentrations (5 nM, 2.5 nM, and 1.25 nM) were used for making the standard curve. Monosaccharide compositions were calculated as molar percentages (mol %). All samples and standards were subjected to high-performance anion-exchange chromatography with pulsed amperometric detection (HPAEC-PAD) on a Dionex PA-20 (Thermo Fisher Scientific, Sunnyvale, CA, USA) essentially as described by [62].

**Calcium binding assay**

AGPs were extracted from the aerial part of 40-day-old glcat mutants and WT plants as described above and dissolved in water. Approximately 1–2 mg/mL AGPs were dissolved in ultrapure water and the concentration was determined by β-D-Gal-Yariv precipitation method at OD₄₂₀. A commercial calcium colorimetric assay kit (MAK022, Sigma-Aldrich, St. Louis, MO, USA) was used for calcium measurement following the manufacturer’s protocol. In this assay, calcium ions from the AGP extract forms a complex with the o-cresolphthalein in the assay kit and cause a color change from transparent to pink. The amount of calcium was determined using a UV spectrometer at OD₄₇₅ and a standard curve made with different concentrations of CaCl₂.

**Germination experiment**

Both freshly harvested and two-month post-harvested seeds of glcat and wild-type plants were used for germination experiments. Sterilized seeds were treated three ways: without stratification, stratification at 4 °C for 3 d, and 22 °C for 3 d in the dark. Seeds were sowed onto ½ MS and 1% sucrose agar plates. Germination percentages were counted from 1 to 5 d after sowing. Approximately 50 seeds were sown for each genotype with three replicates. For germination under ABA treatment, two-month post-harvested seeds were sterilized and stratified at 4 °C for 3 d. Seeds were sown onto ½ MS and 1% sucrose agar plates containing 0–2 μM ABA. Germination percentages were counted from 4 to 10 d after sowing. Approximately 50 seeds were sown for each genotype with three replicates.

**Root hair and trichome morphology**

Seeds from glcat mutants and WT were sterilized and kept at 4 °C in the dark for 3 d for stratification before being sown onto ½ MS and 1% sucrose agar plates. Four-day-old seedlings were transferred onto ½ MS agar plates and kept in a growth chamber at 22 °C, 16 h light/20 °C, 8 h dark photoperiod. Root hairs were measured 5 mm from the root tip, 4 days after transfer. Approximately 300 root hairs were measured for each genotype with three replicates. For trichome morphology, trichomes from rosettes of 40-day-old glcat mutants and WT were used for imaging at 20x magnification with a Nikon SMZ1500 stereomicroscope.

**Evaluation of silique length and seed set**

Siliques were collected from 40-day-old glcat mutants and WT for measurement of silique lengths and seed set. Seed set was measured after clearing the siliques.
color with 70% ethanol overnight. Ten plants for each genotype were used for measurement with three replicates.

In vitro pollen germination assay
Flowers from 35-day-old gcat mutants and WT were used for an in vitro pollen germination assay. The pollen germination medium contained 10% sucrose, 0.01% boric acid, 1 mM CaCl$_2$, 1 mM Ca (NO$_3$)$_2$, 1 mM KCl, 0.03% casein enzymatic hydrolysate, 0.01% myo-inositol, 0.1 mM spermidine, 10 mM GABA, 500 μM methyl jasmonate, and 1% low-melting agarose. Pollen was taken from five flowers for each genotype and was incubated on pollen germination medium. Pollen shape, pollen germination rate, and pollen tube length were measured 3 h after incubation with a Nikon photlab2 microscope at 50x magnification. Around 300 pollen and pollen tubes were measured in an individual experiment with three replicates.

Ruthenium red staining
Seeds of gcat mutants and WT were hydrated with 1 mL water and shaken at 200 rpm for 1 h to remove non-adherent mucilage. The water was removed with a pipet and replaced with 1 mL of 0.01% ruthenium red followed by shaking at 200 rpm for 30 min. Following removal of the stain with a pipet, seeds were rinsed with 1 mL water three times before being observed at 20x magnification under a Nikon SMZ1500 stereomicroscope.

Supplementary information
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arabinoarabinoxylan-protein glycosylation in tobacco and Arabidopsis. Plant Physiol. 2010;154:632–42.

6. Basu D, Liang Y, Liu X, Himmel Dick K, Falk A, Kieliszewski M, et al. Functional identification of a hydroxyproline-O-galactosyltransferase specific for arabinoarabogalactan protein biosynthesis in Arabidopsis. J Biol Chem. 2013;288:10132–43.

7. Ogawa-Omishi M, Matsuyashi Y. Identification of three potent hydroxyproline-O-galactosyltransferases in Arabidopsis. Plant J. 2015;81:736–46.

8. Basu D, Tian L, Wang W, Robbs S, Herock H, Travers A, et al. A small multigene hydroxyproline-O-galactosyltransferase family functions in arabinoarabogalactan-protein glycosylation, growth and development in Arabidopsis. BMC Plant Biol. 2015;15:295.

9. Suzuki T, Nascimento JO, Zeng W, van de Meene A, Yasutomi M, Takemura S, et al. KNS4/UXPK1: a type II arabinoarabogalactan-(1,3,4)-galactosyltransferase required for pollen exine development. Plant Physiol. 2017;173:183–205.

10. Dilkopifl A, Poulsen CP, Vereb G, Kaneko S, Schulz A, Geshi N. Galactosyltransferases from Arabidopsis thaliana in the biosynthesis of type II arabinoarabogalactan: molecular interaction enhances enzyme activity. BMC Plant Biol. 2014;14:190.

11. Gille S, Sharma V, Baidoo EK, Keating JD, Scheller HV, Paul M. Arabinoarabogalactan of a Yariv-precipitable cell wall polymer impacts plant growth as exemplified by the Arabidopsis glycosyltransferase mutant ray1. Mol Plant. 2013;6:369–72.

12. Wu Y, Williams M, Bernard S, Driouich A, Showalter AM, Faik A. Functional identification of two nonredundant Arabidopsis α-(1→2)-fucosyltransferases specific to arabinoarabogalactan proteins. J Biol Chem. 2010;285:13638–45.

13. Tryfon A, Theys TE, Wagner T, Stott K, Kegstra K, Dupree P. Characterisation of FUT4 and FUT6 α-(1→2)-fucosyltransferases reveals that absence of root arabinoarabogalactan fucosylation increases Arabidopsis root growth salt sensitivity. PLoS One. 2014;9:e92932.

14. Knob E, Dilkopifl A, Tryfon A, Poulsen CP, Xiong G, Hafthó J, et al. β-Galactosyltransferase from Arabidopsis thaliana involved in biosynthesis of type II arabinoarabogalactan has a role in cell elongation during seedling growth. Plant J. 2013;79:1016–29.

15. Dilkopifl A, Geshi N. Arabidopsis thaliana thigalactosyltransferase in family GT14. Plant Signal Behav. 2014;9:e28891.

16. Ye CY, Li T, Tuskan GA, Tschaplinski TJ, Yang X. Comparative analysis of GT14/GT14-like gene family in Arabidopsis, Oryza, Populus, Sorghum and Vitis. Plant Sci. 2011;181:688–95.

17. Mutwil M, Obro J, Willats WGT, Persson S. GeneCAT–novel webtools that combine BLAST and co-expression analyses. Nucleic Acids Res. 2008;36(Web Server issue):W320–6.

18. Kleipiova AV, Kasianov AS, Gerasimov ES, Logacheva MD, Penin AA. A high resolution map of the Arabidopsis thaliana developmental transcriptome based on RNA-seq profiling. Plant J. 2016;88:1037–50.

19. Le BH, Cheng C, Bai AQ, Wagmaister JA, Henry KP, Pelletier J, et al. Global analysis of gene activity during Arabidopsis seed development and identification of seed-specific transcription factors. Proc Natl Acad Sci U S A. 2010;107:8063–70.

20. Sprunck S, Rademacher S, Vogler F, Gheyelinck J, Grossniklaus U, Desselhaus T. Egg cell-secreted EC1 triggers sperm cell activation during double fertilization. Science. 2012;338:1093–7.

21. Wang Z-P, Xiong H-L, Dong L, Zhang H-Y, Han C-Y, Wang X-C, et al. Egg cell-specific promoter-controlled CRISPR/Cas9 efficiently generates homozygous mutants for multiple target genes in Arabidopsis in a single generation. J Integr Biol. 2015;16:144.

22. Kitazawa K, Tryfon A, Toshimi Y, Hayashi Y, Kawauschi S, Antonov L, et al. β-Galactosyl Yariv reagent binds to the β-(1,3)-galactan of arabinoarabogalactan proteins. Int J Mol Sci. 2015;16:3452.
52. Yu L, Shi D, Li J, Kong Y, Yu Y, Chai G, et al. CELLULOSE SYNTHASE-LIKE A2, a glucomannan synthase, is involved in maintaining adherent mucilage structure in Arabidopsis seed. Plant Physiol. 2014;164:1842–56.
53. Dean GH, Zheng H, Tewari J, Huang J, Young DS, Hwang YT, et al. The Arabidopsis MUM2 gene encodes a beta-galactosidase required for the production of seed coat mucilage with correct hydration properties. Plant Cell. 2007;19:4007–21.
54. Voiniciuc C, Dean GH, Griffiths JS, Kirchsteiger K, Hwang YT, Gillett A, et al. Flying saucer1 is a transmembrane RING E3 ubiquitin ligase that regulates the degree of pectin methylesterification in Arabidopsis seed mucilage. Plant Cell. 2013;25:944–59.
55. Hepler PK, Kunkel JG, Rounds CM, Wnship LJ. Calcium entry into pollen tubes. Trends Plant Sci. 2012;17:32–8.
56. Winter D, Vinegar B, Nahal H, Ammar R, Wilson GV, Provart NJ. An "electronic fluorescent pictograph" browser for exploring and analyzing large-scale biological data sets. PLoS One. 2007;2:e718.
57. Gagnon JA, Valen E, Thyme SB, Huang P, Ahkmetova L, Pauli A, et al. Efficient mutagenesis by Cas9 protein-mediated oligonucleotide insertion and large-scale assessment of single-guide RNAs. PLoS One. 2014;9:e98186.
58. Engler C, Marillonnet S. Golden Gate cloning. Methods Mol Biol. 2014;1116:119–31.
59. Clough SJ, Bent AF. Floral dip: a simplified method for agrobacterium-mediated transformation of Arabidopsis thaliana. Plant J Cell Mol Biol. 1998;16:735–43.
60. Hua Y, Wang C, Huang J, Wang K. A simple and efficient method for CRISPR/Cas9-induced mutant screening. J Genet Genomics. 2017;44:207–13.
61. Lamport D. Preparation of arabinogalactan glycoproteins from plant tissue. BIO-Protoc. 2013:3. https://doi.org/10.21769/BioProtoc.918.
62. ØBro J, Harholt J, Scheller HV, Orfila C. Rhamnogalacturonan I in Solanum tuberosum tubers contains complex arabinogalactan structures. Phytochemistry. 2004;65:1429–38.

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