

Identification of MEDIATOR16 as the *Arabidopsis* COBRA suppressor MONGOOSE1

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We performed a screen for genetic suppressors of *cobra*, an *Arabidopsis* mutant with defects in cellulose formation and an increased ratio of unesterified/esterified pectin. We identified a suppressor named *mongoose1* (*mon1*) that suppressed the growth defects of *cobra*, partially restored cellulose levels, and restored the esterification ratio of pectin to wild-type levels. *mon1* was mapped to the *MEDIATOR16* (*MED16*) locus, a tail mediator subunit, also known as *SENSITIVE TO FREEZING6* (*SFR6*). When separated from the *cobra* mutation, mutations in *MED16* caused resistance to cellulose biosynthesis inhibitors, consistent with their ability to suppress the *cobra* cellulose deficiency. Transcriptome analysis revealed that a number of cell wall genes are misregulated in *med16* mutants. Two of these genes encode pectin methyltransferase inhibitors, which, when ectopically expressed, partially suppressed the *cobra* phenotype. This suggests that cellulose biosynthesis can be affected by the esterification levels of pectin, possibly through modifying cell wall integrity or the interaction of pectin and cellulose.

cell wall | cellulose | freezing tolerance | pectin | transcription

Cellulose, the backbone of the primary plant cell wall, supports a complex polysaccharide-rich network formed of hemicelluloses and pectin (1). Unlike other cell wall polymers, cellulose is synthesized at the plasma membrane by the cellulose synthase complex, which synthesizes multiple β -1,4 glucan chains that hydrogen-bond to form cellulose fibrils (2–5). The proposed catalytic components of the cellulose synthase complex in higher plants are the CESA proteins. The stoichiometry of the cellulose synthase complex, as well as the exact number of glucan chains in individual cellulose fibrils, is unclear (3, 6, 7). Additional proteins involved in some aspect of the cellulose formation process have been implicated by analysis of transcriptional networks (8, 9). The *Arabidopsis* COBRA gene was found to be involved in cell expansion (10), and has been proposed to participate in cellulose synthesis (11).

During cellulose biosynthesis, other cell wall components can potentially affect the formation of fibrils by interacting with the nascent glucan chains or microfibrils. For example, in an *Arabidopsis* mutant that lacks xyloglucan, thicker cellulose fibrils have been observed (12), supporting the idea that xyloglucan prevents cellulose microfibril aggregation (13). Primary cell walls are also rich in pectin, which can bind cellulose with similar affinity to xyloglucan (14) and, in *Arabidopsis* primary cell walls, up to 50% of the cellulose is in direct contact with pectin (15). Thus, it may be anticipated that mutations that affect the structure or amount of pectin and other noncellulosic polysaccharides may impact cell wall assembly.

The factors that regulate cell wall composition are gradually being revealed (16). Several NAC transcription factors have been identified that control cell wall thickness (17, 18), and specific MYB transcription factors are known to be regulators of secondary cell wall biosynthesis (19, 20). Recently, a large transcriptional network that regulates secondary cell wall biosynthesis was elucidated, identifying tens of transcription factors and their role in a complex

regulatory network leading to organized secondary wall formation in xylem (21). It was recently discovered that disruption of the genes for MEDIATOR5a and 5b can suppress the growth defects of *Arabidopsis ref8*, a mutant defective in lignin biosynthesis (22).

The mediator transcriptional coactivator complex has been found to be a crucial component in promoting eukaryotic transcription, as it links transcription factor binding at promoters to the activity of RNA polymerase II (Pol II) (23). Mediator is a multisubunit protein complex comprising between 25 and 34 subunits depending on the species (24, 25), and plays a role in the transcription of both constitutively expressed and inducible genes (26). Mediator has been described as being organized into four submodules: the head, middle, tail, and kinase domains (27). The tail submodule is thought to associate directly with transcriptional activators and repressors and the head with Pol II (23). The *Arabidopsis* mediator complex was purified (28) and, in combination with subsequent bioinformatic analysis (25), 34 subunits were identified, a number of these specific to plants (29). In *Arabidopsis*, roles for Mediator subunits have been demonstrated in the transcriptional response to a number of biotic (30–32) and abiotic stress conditions (33, 34) as well as in plant development (35, 36).

To better understand the role of COBRA in cellulose synthesis, we identified six independent suppressors of the *cob-6* allele, named *mongoose1–6*. The *mongoose1* (*mon1*) mutation mapped to the *MEDIATOR16* [*SENSITIVE TO FREEZING6* (*SFR6*)/*MED16*] locus. Analysis of the effects of mutations in *MED16* on transcription identified two pectin methyltransferase inhibitors (PMEIs) that are regulated by *MED16*. Overexpression of these PMEIs causes partial

Significance

The *cobra* mutants of *Arabidopsis*, such as *cob-6*, have impaired growth associated with a defect in cellulose synthesis. Mutations in *MEDIATOR16* (*MED16*) reduce the number of misregulated genes in *cob-6* mutants and suppress the phenotypes. This observation implicates *MED16* in transcriptional responses to cell wall defects. Ectopic expression of two pectin methyltransferase inhibitors (PMEIs) identified in a suppressor screen partially suppressed the growth defect in the *cob-6* mutant. The results confirm that the PMEIs have significant *in vivo* activity, and provide evidence that pectin esterification can modulate cell wall properties.

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suppression of *cobra*, suggesting that pectin esterification is a significant factor in cell wall integrity.

Results

Isolation of *mongoose* Mutations. In an effort to understand the function of COBRA, we carried out a suppressor screen for restoration of root growth in plants homozygous for a *cobra* mutation. Because null mutations of *COBRA* are virtually sterile (37), we performed the screen using *cob-6*, a weak T-DNA allele that produces ~10% of functional transcripts (38). Approximately 100,000 seeds of a *cob-6* line were mutagenized with ethyl methanesulfonate (EMS), and seedlings were screened in the M2 generation. Six recessive *cob-6* suppressors, *mongoose1-6* (*mon1-6*), were identified (Fig. 1). Allelism tests revealed that the *mon* mutations represent six genes (Fig. S1). The *cobra* mutants are sensitive to increasing concentrations of sucrose (37), and all six suppressors restore normal growth of *cob-6* on high sucrose concentrations as well as normal hypocotyl elongation in dark-grown seedlings (Fig. 1).

Previously described suppressors of the *cob-6* mutation have been reported to act by increasing the transcript levels of *cobra* (39). In contrast, in all six *mon* lines, *cobra* transcript levels were similar to those in *cob-6* (Fig. S1), implying that suppression in these lines is not due to increased accumulation of functional *cobra* transcripts.

Cellulose Characterization of *mon1 cob-6*. In addition to the growth phenotype, we assessed whether the cellulose defects caused by *cob-6* were suppressed by *mon-1*. We analyzed the cellulose macrostructure and amounts in a line homozygous for *mon1* and *cob-6* (Fig. 2). Cellulose macrostructure was visualized using S4B staining (12). In *cob-6*, defects in cellulose macrostructure can be seen as early as in the division zone, at which point a fine network of thin fibrils can be seen in wild type. In *cob-6* the network appears more diffuse than in wild type, and in addition there are patches of bright staining in *cob-6* that are rarely observed in wild type. In *mon1 cob-6* the staining looks similar to wild type, however with a less well defined pattern than in wild type. In mature wild-type root cells, cellulose fibrils can be clearly seen. The staining in *cob-6* displays a higher variability between cells, with some cells exhibiting diagonal fibrils and some cells lacking stain completely. In *mon1 cob-6*, S4B staining of elongated cells in the root showed that the cellulose macrostructure was similar to wild type (Fig. 2A). Quantitative measurements of cellulose (Table 1) showed that there is an almost 50% reduction of

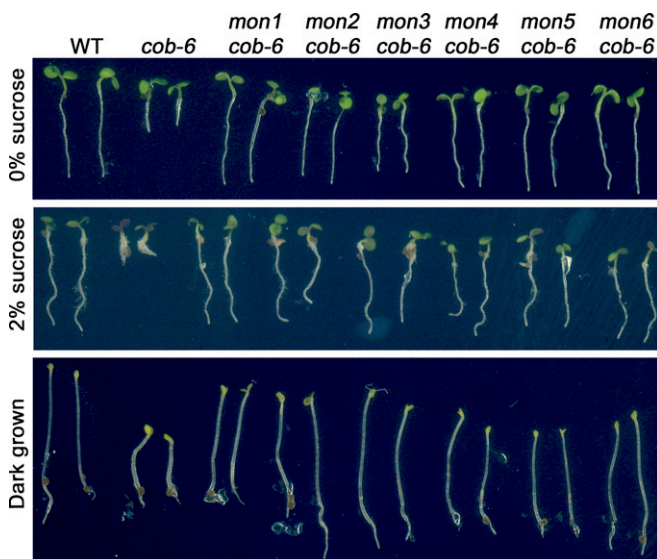


Fig. 1. Phenotypes of the *mongoose* mutants. All *mon* lines are also homozygous for the *cob-6* mutation.

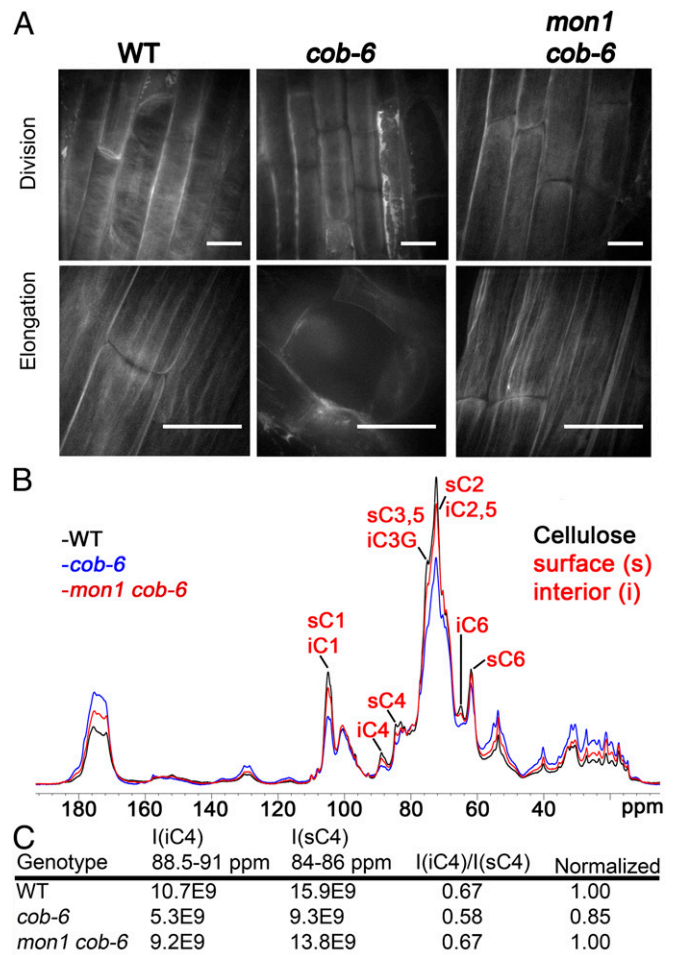


Fig. 2. Cellulose macrostructure and amount in *mon1 cob-6* mutant. (A) Cellulose in root cells stained with S4B. In *cob-6*, staining is reduced and less homogeneous, with some cells exhibiting almost a complete lack of fluorescence. Fibrils that can be detected are not as defined as in wild type, and are not as regularly oriented (Upper, *cob-6*). In *mon1 cob-6*, fibrils were similar to wild type, more so in elongated cells. (Scale bars, 15 μ m.) (B) One-dimensional ssNMR analysis. Quantitative ^{13}C direct polarization (DP)-MAS ssNMR spectra of wild-type, *cob-6*, and *mon1 cob-6* cell walls. Additional annotation is shown in Fig. S8. (C) Relative intensities of interior and surface cellulose C4 signals from ^{13}C DP-MAS spectra.

cellulose in *cob-6*. The *mon1 cob-6* lines showed a significant increase of cellulose levels compared with *cob-6*, albeit still below that of wild type (Table 1). The *mon1* mutation also reversed changes in other polysaccharides in the *cob-6* mutant, as indicated by changes in cell wall sugar composition of the *mon1 cob-6* line (Table 1).

Further analysis of the molecular structure of *mon1 cob-6* cell walls compared with *cob-6* and wild type was obtained using magic-angle-spinning (MAS) solid-state NMR (ssNMR) spectroscopy (Fig. 2C). We applied quantitative ^{13}C ssNMR by direct polarization experiments using a long recycle delay. There were clear differences between wild type and *cob-6* revealed in the 1D spectrum due to a significant reduction in cellulose and relative increase in pectin and glycoprotein. However, the 1D ^{13}C spectrum of *mon1 cob-6* resembles the wild-type spectrum, indicating a significant recovery of cellulose and decrease in pectin and glycoprotein. Furthermore, the ratio between the intensity of the interior C4 peak (iC4) and surface C4 peak (sC4) of cellulose was measured and, whereas *cob-6* showed 15% lower crystallinity relative to wild type, *mon1 cob-6* showed no difference relative to wild type (Fig. 2D). These results correlate with the cell wall analysis results (Table 1),

Table 1. Monosaccharide and cellulose analysis

Component	WT	<i>cob-6</i>	<i>mon1 cob-6</i>
Man	6.2 ± 0.2 ^A	7.1 ± 0.2 ^B	6.5 ± 0.1 ^A
Fuc	4.1 ± 0.1 ^A	3.8 ± 0.2 ^A	4.2 ± 0.2 ^A
Ara	19.4 ± 0.6 ^A	37.5 ± 0.8 ^B	21.3 ± 0.3 ^C
Glu	9.8 ± 0.2 ^A	11.1 ± 0.2 ^B	10.0 ± 0.3 ^A
Xyl	20.9 ± 0.5 ^A	21.1 ± 0.4 ^A	20.5 ± 0.7 ^A
Rha	11.1 ± 0.3 ^A	11.4 ± 0.4 ^A	11.1 ± 0.3 ^A
Gal	45.8 ± 2.1 ^A	74.3 ± 4.2 ^B	50.1 ± 3.5 ^A
Cellulose	122 ± 3.2 ^A	65 ± 4.7 ^B	105 ± 2.4 ^C

Values are μg per mg of alcohol-insoluble residue. Superscripts represent statistical differences of $P < 0.05$ by two-way analysis of variance coupled to Tukey test. The high level of galactose (Gal) and arabinose (Ara) in *cob-6* is due to the high level of pectin in the mutant.

and support the conclusion that suppression of *cob-6* by *mon1* involves restoration of cell wall composition and structure.

Mapping of *mon1* to *MED16/SFR6*. We used a bulk segregant approach with genomic DNA sequencing to identify the *mon1* mutation (40). Two hundred segregants from a *mon1 cob-6* line backcrossed to *cob-6* were pooled for whole-genome sequencing. The analysis showed that the causative mutation was located on the upper arm of chromosome 4 (Fig. S2). In the region with single nucleotide polymorphism (SNP) frequencies over 80% there were 26 mutations in genes, but only 5 were missense mutations. One of the five candidate SNPs was a C-to-T mutation at position 5679 in AT4G04920, which leads to a serine-to-phenylalanine substitution at position 889 (Fig. 3A).

To test whether this was the causative mutation, we crossed *cob-6* with a previously characterized T-DNA mutation in AT4G04920, *sfr6-3*. *sfr6-3 cob-6* seedlings exhibited normal growth under light (Fig. 3B) and dark (Fig. 3C) conditions. Cellulose analysis (Fig. 3D) revealed that *sfr6-3 cob-6* cellulose levels were similar to that in *mon1 cob-6*—significantly higher than *cob-6* but lower than wild type. *COBRA* transcript levels in *sfr6-3 cob-6* were the same as in *cob-6* and *mon1 cob-6* (Fig. S3). These results confirmed that the suppression in *mon1 cob-6* is due to a mutation in AT4G04920.

SFR6 was originally identified in a screen for plants that were sensitive to freezing after cold acclimation (41). *sfr6-1* was mapped to AT4G04920, and additional T-DNA alleles, *sfr6-2* and *sfr6-3*, were characterized (42). We performed freezing experiments for the different lines using acclimated and nonacclimated plants (Fig. S4). *sfr6-3* was sensitive to freezing despite cold acclimation, as expected. With no cold acclimation, wild type was sensitive to freezing, as were *sfr6-3*, *mon1 cob-6*, and *sfr6-3 cob-6*. However, *cob-6* was resistant to freezing, even when the plants were not acclimated (Fig. S4). We hypothesized that this might be due to altered expression of genes that respond to the cell wall damage of *cob-6* that are also involved in cold acclimation. To test this, we analyzed the expression of *CBF1*, a transcription factor gene, known to be up-regulated during cold acclimation (43). The results showed that in *cob-6* plants, *CBF1* is up-regulated without cold acclimation (Fig. S4).

Characterization of *med16* Mutations as Suppressors of Cellulose Deficiency. Seedlings of *sfr6-3* are larger than wild type (Fig. 3), but there is no other obvious phenotype under normal growth conditions. However, *sfr6* mutants were shown to be hypersensitive to freezing, as well as to osmotic stress (44). To test the response of *sfr6-3* to perturbations in cellulose synthesis, we performed cellulose biosynthesis inhibitor assays (Table 2) using inhibitors with different effects on the cellulose synthase complex (45).

To test the response to the different cellulose biosynthesis inhibitors, seedlings were grown on 1/2 Murashige and Skoog (MS) plates with increasing concentrations of the drugs for 7 d. For each line, 50 seedlings were measured and growth inhibition (GI50)

was calculated. The results were similar for all three inhibitors (Table 2). *cobra* is hypersensitive to all of the inhibitors, although the response to isoxaben is more dramatic compared with dichlorobenzonitrile (DCB) and indaziflam. In contrast, *sfr6-3* is significantly more resistant than wild type to all of the inhibitors. Although *mon1 cob-6* and *sfr6-3 cob-6* exhibit normal root length under control conditions or even when grown with 2% (wt/vol) sucrose, which enhances the *cobra* phenotype (Fig. 1), they were found to be more sensitive than wild type to the inhibitors, although still significantly more resistant than *cob-6*. The results in Table 2 demonstrate that the mutation in *MED16* (*sfr6-3*) causes resistance to cellulose synthesis perturbation.

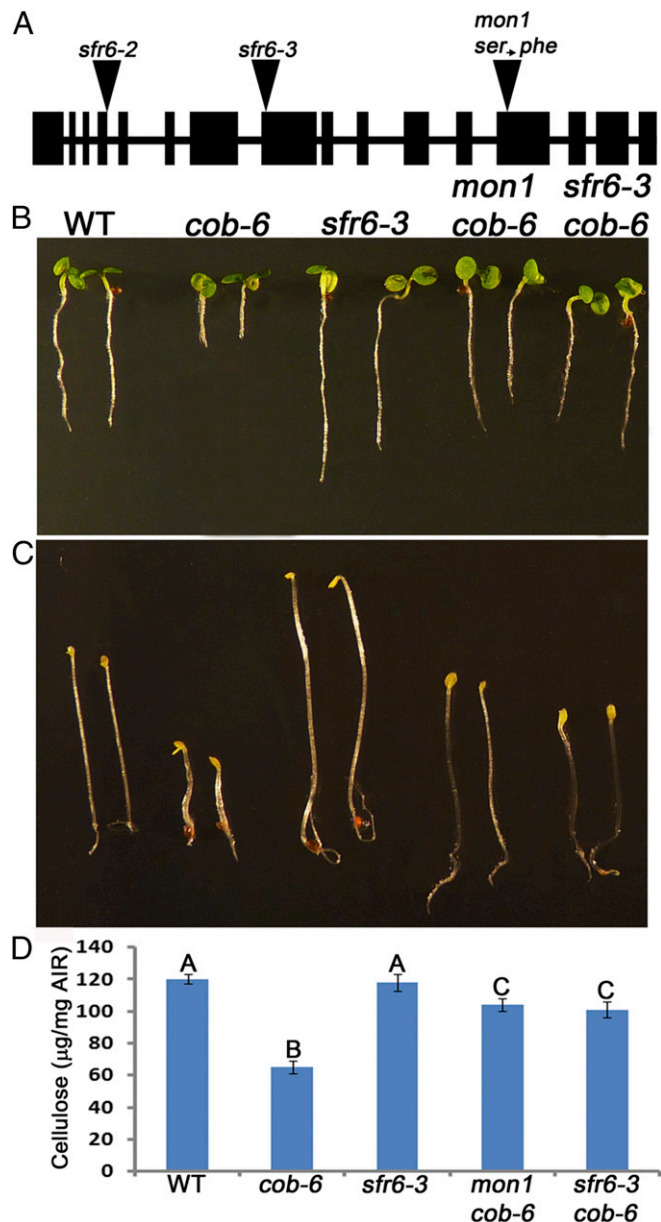


Fig. 3. Properties of *med16* mutants. (A) Gene structure of AT4G04920 (*MED16*) showing T-DNA insertion sites and the *mon1* mutation. (B and C) Growth phenotype of 7-d light-grown seedlings (B) and 5-d dark-grown hypocotyls (C) showing suppression of the *cob-6* phenotype by *mon1* and *sfr6-3*. (D) Cellulose measurement showing suppression of *cob-6* cellulose deficiency. Values are means \pm standard deviation. Letters above the bars indicate significant differences based on one-way ANOVA and Tukey's test, $P < 0.05$. AIR, alcohol-insoluble residue.

Table 2. Mutations in *MED16* cause resistance to cellulose biosynthesis inhibitors

Line	Growth inhibition		
	Isoxaben, nM	DCB, nM	Indaziflam, pM
WT	2.81 ± 0.21 ^A	102 ± 8 ^A	220 ± 10 ^A
<i>cob-6</i>	0.53 ± 0.07 ^B	57 ± 2 ^B	167 ± 7 ^B
<i>sfr6-3</i>	3.11 ± 0.15 ^C	138 ± 8 ^C	274 ± 11 ^C
<i>mon1 cob-6</i>	2.58 ± 0.14 ^D	92 ± 5 ^D	198 ± 9 ^D
<i>sfr6-3 cob-6</i>	2.51 ± 0.12 ^D	88 ± 7 ^D	191 ± 7 ^D

Root length of seedlings grown on vertically oriented agar plates with various concentrations of the inhibitors for 7 d was measured. GI50 is the concentration of drug that leads to 50% inhibition of root length relative to the growth of the line on agar without inhibitors. For statistical analysis, we performed ANOVA-coupled Tukey test on the raw data. The values are the means ± SE ($n = 50$).

Analysis of the Effect of *MED16* on Gene Regulation in the *cobra* Background. *SFR6* was identified as *MED16*, a component of the MEDIATOR transcriptional coactivator complex (31), and is required for RNA polymerase II recruitment of genes regulated by the CBF transcription factor (34). More generally, *MED16* has been shown to regulate expression of multiple genes associated with a variety of biological functions (31, 34). To identify potential targets of *MED16* that are involved in *cobra* suppression, we performed RNA-sequencing (seq) analysis on 7-d-old seedlings of wild type, *cob-6*, *sfr6-3*, and *sfr6-3 cob-6* (Fig. 4A, GEO accession no. GSE75199). Using a 1.5-fold difference in expression value with $P < 0.05$ as the cutoff parameter, we identified 277 misregulated genes in *cob-6* (180 ↑, 97 ↓), 302 in *sfr6-3* (29 ↑, 273 ↓), and 677 in *sfr6-3 cob-6* (201 ↑, 476 ↓) (Fig. 4A). To have a broad look at the misregulated genes in the three mutants, we analyzed the data based on Gene Ontology (GO) annotation (Table S1). The largest difference was found to be in the signal transduction and DNA-dependent transcription classes, where, as expected, there is a larger portion of misregulated genes in these classes for *sfr6-3* and *sfr6-3 cob-6*.

To learn more about the suppression mechanism, we focused on the comparison between *cob-6* and *sfr6-3 cob-6* (Fig. S5, GEO accession no. GSE75199 and Table S1). First, there are 148 genes that are misregulated in *cob-6* and are not misregulated in *sfr6-3 cob-6*. It is difficult to evaluate the contribution of these genes to the *cob-6* phenotype, because genes in this list are spread across cellular localization and function. To identify potential targets for further analysis, we raised the cutoff to twofold, keeping $P < 0.05$. The 20 most strongly misexpressed genes (10 highest and 10 lowest) were either unknown genes or genes with no obvious connection to cell wall biosynthesis. However, further down the list, 48 cell wall-related genes were differentially expressed twofold or greater between *cob-6* and *sfr6-3 cob-6* (14 ↑, 34 ↓). The largest subgroup within this list consisted of 11 genes involved in the modification of pectin. Characterization of other cellulose-deficient mutants has revealed that the pectin fraction is highly altered in response to reduction in cellulose (46), so we focused additional studies on this group.

The Role of *MED16* in Pectin Esterification. Pectin is deposited in the apoplast in a highly esterified form, and esterification level decreases during development. Nonesterified pectin can form calcium bridges that affect its rigidity (47). The degree of pectin esterification was decreased in *cob-6* compared with wild type (Fig. 4B). However, the degree of esterification in *cob-6 mon1* or *cob-6 sfr6-3* was similar to wild type. This suggests that the suppression mechanism involves the restoration of pectin esterification, at least in part.

We identified 11 misregulated pectin-related genes in *sfr6-3 cob-6* compared with *cob-6*: 4 pectin lyase-like, 2 pectin methylsterases

(PMEs), and 5 PMEIs. We focused on the two most highly misexpressed PMEIs, AT3G17130 (*PMEI8*) and AT1G62770 (*PMEI9*).

PMEIs can bind to PMEIs and inhibit their activity, and have been shown to participate in growth control (48). *PMEI8* is expressed at relatively low levels during development, whereas *PMEI9* expression is higher in seedlings compared with the rest of the developmental stages [based on expression profiles from Genevestigator (49)]. We tested the expression of both *PMEI8* and *PMEI9* in the different lines using quantitative (q)RT-PCR (Fig. 4C). Consistent with the RNA-seq data, both *PMEI8* and *PMEI9* were up-regulated in *mon1 cob-6* and *sfr6-3 cob-6* compared with wild type (Fig. 4A), the *PMEI9* increase being more striking. *PMEI8* and *PMEI9* expression was slightly reduced in *cob-6* single mutants, and slightly higher in *sfr6-3*, but to a lesser extent. The differences in expression between *sfr6-3* and *mon1 cob-6* suggested that *PMEI8* and *PMEI9* expression is exaggerated when there are cell wall defects. To test this, we analyzed *PMEI8* and *PMEI9* expression in *sfr6-3* after treatment with isoxaben. The results (Fig. S6) showed that, indeed, *PMEI8* and *PMEI9* are up-regulated significantly when *sfr6-3* seedlings are treated with isoxaben, supporting the idea that regulation of *PMEI8* by *sfr6-3* is affected by an additional stress. To directly test the effect of these two PMEIs on the *cob-6* phenotype, we expressed the genes under control of the 35S promoter in the *cob-6* background (Fig. 4D). The results demonstrate that overexpression of *PMEI8* or *PMEI9* causes partial suppression of the *cob-6* phenotype.

Discussion

Isolation of *cobra* Suppressors. The phenotype of *cobra* mutants includes reduced growth and swollen organs that are associated with defects in cellulose synthesis and deposition (10, 37). Suppression of the swollen root phenotype, but not the root elongation defect, of a *cobra* mutant was observed in the *IAA-Alanine Resistant 4* (*iar4*) mutant (50). The mechanism was suggested to be related to the role of auxin in regulating cell wall loosening (50). Another cellulose-deficient mutant, *procuste*, was suppressed

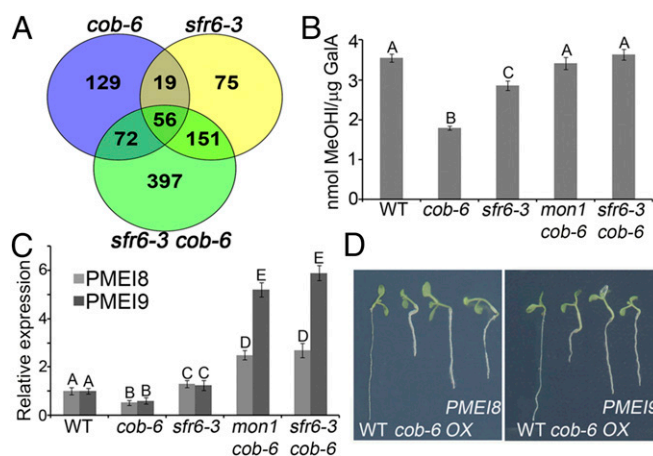


Fig. 4. Increased transcription of pectin methylsterase inhibitors and the restoration of pectin esterification to wild-type levels in *mon1 cob-6* and *sfr6-3 cob-6*. Bar graphs indicate means ± standard deviation. Letters above the bars indicate significant differences based on one-way ANOVA and Tukey's test, $P < 0.05$. (A) Venn diagram for RNA-seq analysis of *cob-6*, *sfr6-3*, and *sfr6-3 cob-6* shows all genes that are up-regulated and down-regulated compared with wild type with 1.5-fold change or above. For further analysis, we looked at genes that are misexpressed between *sfr6-3 cob-6* and *cob-6*, and identified 48 cell wall-related genes. (B) Pectin esterification levels in the different lines. Values were normalized to the total amount of galacturonic acid (Fig. S7A). (C) Steady-state mRNA levels of two pectin methylsterase inhibitor genes, AT3G17130 (*PMEI8*) and AT1G62770 (*PMEI9*), in various genotypes. (D) *PMEI8* and *PMEI9* were overexpressed (OX) in the *cob-6* background. Ten independent lines were analyzed, all demonstrating partial suppression of the *cob-6* phenotype.

by mutation in a receptor-like kinase, *theseus1* (51). However, *theseus1* does not suppress *cobra* (Fig. S7B), suggesting that not all cellulose deficiencies are equal.

The relatively large number of *cobra* suppressors reported here raises the possibility that some or all of the *mon* mutations may be in a common pathway for what is obviously a dispensable function under laboratory conditions. Additional research will be required to characterize the functions of the other *MONGOOSE* genes, and to identify any other mutations that can be recovered by expanding the screen. Whatever the case, identification of *med16* (*sfr6/mon1*) as a *cob-6* suppressor contributes to our knowledge of the transcriptional regulation of genes involved in cell wall biosynthesis or remodeling.

mediator16 as a *cobra* Suppressor. Mediator subunits, particularly those comprising the tail submodule, interact with transcription factors to control gene expression (52). However, no MED16-interacting transcription factors from plants have been identified to date. Lines carrying a mutation in *MED16* (EMS line *sfr6-1*, T-DNA insertion lines *sfr6-2*, *sfr6-3*, and *yid1*) do not exhibit major phenotypic changes under normal conditions. *sfr6* mutants are slightly larger than their wild-type counterparts at the seedling stage (Fig. 3) (42), and exhibit pale, chlorotic leaves (42, 53), which was recently found to be attributable to iron deficiency (53). The largest phenotypic differences observed between *med16* mutants and wild-type plants are seen under abiotic and biotic stress conditions; *med16* mutants are sensitive to freezing, osmotic stress, pathogen attack, and iron deficiency (42, 44, 53). RNA-seq data show that in *sfr6-3*, 302 genes are misregulated compared with wild type, whereas 677 genes are misregulated in an *sfr6-3 cob-6* double mutant.

Based on expression data from Genevestigator (49), *MED16* expression does not change dramatically under perturbation conditions. Due to its position in the yeast mediator complex, where Sin4 (a yeast MED16 homolog) links the so-called triad of tail subunits to the rest of the complex (54), it has been suggested that some of the phenotypes associated with loss of MED16 in *Arabidopsis* may be attributable to loss of other tail subunits that require MED16 to tether them to the complex (34).

Transcriptome analysis has been carried out previously for *sfr6* using microarrays on 7- to 8-d-old seedlings (34). These studies had implicated MED16 as a factor in the transcriptional regulation of some aspects of cell wall biosynthesis or remodeling. Here we performed RNA-seq analysis on *sfr6-3* as well as *sfr6-3 cob-6*. Three hundred and two genes are misregulated in *sfr6-3*, whereas 677 genes are misregulated in *sfr6-3 cob-6*. One hundred and forty-eight genes that are misregulated in *cob-6* are not misregulated in *sfr6-3 cob-6*. It is not obvious whether this is a cause or an effect of the suppression of *cob-6* by *sfr6-3*. It is possible that the elevation in cellulose levels in *mon1 cob-6* causes suppression of the *cob-6* phenotype. On the one hand it is possible that the *cob-6* mutation triggers a cascade of gene expression changes that cause the phenotypes, and that a phenotypically important part of that cascade is blocked by the *sfr6-3* mutation. Alternatively, the phenotypes might be caused directly by the loss of COBRA function in *cob-6*, the usual working assumption (37), and the *sfr6-3* mutation suppresses those effects by altering expression of genes that encode compensating functions. Unfortunately, the large number of misregulated genes is a barrier to a simple explanation for the mechanistic basis for the suppression of *cob-6* by mutations in *MED16*. However, the

observation that ectopic constitutive expression of two MED16-regulated genes (*PME18*, *PME19*) causes partial suppression of the *cob-6* phenotype suggests that a significant component of the suppression effect is via pectin modification.

Pectin Esterification and Freezing Tolerance. Both pectin amounts and degree of esterification increase after cold acclimation (55). Therefore, the regulation of pectin content and degree of esterification that is partially mediated by MED16 might contribute to the established role of MED16 in freezing tolerance.

In *cob-6* plants the pectin is highly nonesterified, and the plants are resistant to freezing (Fig. S4). However, *CBF1* is up-regulated in *cob-6* even when plants are not cold-acclimated. This fact is likely to be the most significant factor in the freezing tolerance of non-acclimated *cob-6*. Assuming that the primary effect of the *cob-6* mutation is a defect in cellulose synthesis, the implication seems to be that altered cell wall structure can induce *CBF1*.

Pectin Esterification and Cellulose Biosynthesis. We have shown that the pectin fraction in the *cobra* mutant is highly nonesterified (Fig. 4B). Introduction of *PME18* and *PME19* under control of the 35S promoter increases the amount of pectin esterification in the *cob-6* mutant to essentially wild-type levels and partially suppresses the *cobra* phenotype. This effect runs counter to the notion that increased Ca²⁺-mediated cross-linking of nonesterified pectin may compensate for a defect in cellulose synthesis by strengthening the cell wall. Thus, it is unclear why increased nonesterified pectin partially suppressed the *cobra* phenotype. One possibility arises from the observation that changes in pectin esterification can trigger brassinosteroid signaling, resulting in cell wall remodeling (56). Perhaps such remodeling compensates for the cell wall defects in *cobra* mutants. Another, highly speculative, possibility is that pectin methylation may affect the binding of pectin to nascent cellulose microfibrils during cellulose synthesis. Pectin has been shown to bind cellulose (57). Perhaps methylated pectin is less disruptive or inhibitory to cellulose crystallization, a process that appears to be defective in the *cobra* mutants, or to some other aspect of cell wall assembly.

Materials and Methods

Arabidopsis ecotype Columbia was used for all experiments, and was grown as described (12). Seeds were mutagenized as described (35). The *mon1* mutation was identified using bulk segregant analysis of 200 homozygous *mon1* F2 segregants from a backcross to the *cob-6* mutant (40). Freezing experiments were carried out as described (42). Spinning-disk confocal microscopy was as described (12). Uniformly ¹³C-labeled primary cell walls were prepared and analyzed by NMR (57, 58). Cellulose biosynthesis inhibitor assays (45) and monosaccharide and cellulose analyses were performed as previously described (59). Pectin esterification was measured as previously reported (60). More detailed materials and methods are available in *SI Materials and Methods*.

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