Cell Wall Maturation of Arabidopsis Trichomes Is Dependent on Exocyst Subunit EXO70H4 and Involves Callose Deposition

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Arabidopsis (Arabidopsis thaliana) leaf trichomes are single-cell structures with a well-studied development, but little is understood about their function. Developmental studies focused mainly on the early shaping stages, and little attention has been paid to the maturation stage. We focused on the EXO70H4 exocyst subunit, one of the most up-regulated genes in the mature trichome. We uncovered EXO70H4-dependent development of the secondary cell wall layer, highly autofluorescent and callose rich, deposited only in the upper part of the trichome. The boundary is formed between the apical and the basal parts of mature trichome by a callose ring that is also deposited in an EXO70H4-dependent manner. We call this structure the Ortmannian ring (OR). Both the secondary cell wall layer and the OR are absent in the exo70h4 mutants. Ecophysiological aspects of the trichome cell wall thickening include interference with antiherbivore defense and heavy metal accumulation. Ultraviolet B light induces EXO70H4 transcription in a CONSTITUTIVE PHOTOMORPHOGENIC1-dependent way, resulting in stimulation of trichome cell wall thickening and the OR biogenesis. EXO70H4-dependent trichome cell wall hardening is a unique phenomenon, which may be conserved among a variety of the land plants. Our analyses support a concept that Arabidopsis trichome is an excellent model to study molecular mechanisms of secondary cell wall deposition.

Driven by selection pressure on reproductive success, the surface layers of land plants continually evolve to endure biotic and abiotic stresses. This has resulted in the dynamic evolution of different epidermis and derived structures, such as trichomes, across species and genotypes. Trichomes particularly can be of different forms, even within one individual plant. Because they are unessential for plant survival in laboratory conditions, trichomes became popular targets of genetic deletions, trichomes patterning/initiation and early morphogenetic processes. Nevertheless, trichome development has been shown to be modulated by abiotic stresses in Arabidopsis and other systems. Yamasaki et al. (2007), for instance, showed that UV-B irradiation was responsible for increasing the number of cells and the amount of polyphenolic compounds in trichomes. Roles in the drought and heat stresses (Ehleringer, 1982; Grammatikopoulos and Manetas, 1994; Espigares and Peco, 1995; Pérez-Estrada et al., 2000) and heavy metal detoxification and deposition (Salt et al., 1995; Pérez-Estrada et al., 2000; Servin et al., 2012; Jr and Kupper, 2014) also have been described. Additionally, different forms of trichomes have been linked to increased plant resistance against herbivores across different species of plants (for review, see Riddick and Simmons, 2014).

Arabidopsis trichomes are nonglandular hairs with epidermal origin. Unlike most plants with multicellular trichomes, Arabidopsis trichomes are unicellular (but undergo four endoreduplication cycles; Hülskamp et al., 1994). Despite unicellularity, Arabidopsis trichomes reach an extremely polarized shape, with bulged stalk and three to four branches. Development has been well characterized genetically (Hülskamp et al., 1994; Folkers et al., 1997; for review, see Marks et al., 1991), and it has been divided into six stages: initiation, polar expansion, branching, branch growth, diffuse growth, and maturation of cell wall (Szymanski et al., 1998). Nevertheless, interest has remained firmly focused on the first five stages, when trichome growth and formation occur, with rather less attention paid to the stage of trichome maturation. Only a few mutants with defective trichome maturation have been identified to date. For instance, the

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group of glassy mutants lacks surface papillae, rendering them a lustrous and transparent appearance. These include chablis, chardonnay, retsina (Hülskamp et al., 1994), *glassy hair1* (GLH1), GLH2, GLH3, GLH4, and GLH6 (Suo et al., 2013). Unfortunately, only phenotype description and chromosomal position mapping of glassy genes have been published. A similar situation prevails with mutant *underdeveloped trichome* (Haughn and Somerville, 1988) and constitutive expression of *PR genes5* (Brininstool et al., 2008). Mapped mutations affecting the trichome cell wall include several transcription factors and cell wall enzymes. *murus2* mutation is in the fucosyltransferase1, resulting in underdeveloped trichome papillae (Vanzin et al., 2008). Mapped mutants similar to *murus3* mutation in the xyloglucan galactosyltransferase (Madson et al., 2003). Mutants named *trichome bifringence* lack the birefringence of the trichomes, which is primarily caused by paracrystalline cellulose. The precise role of these proteins is, however, still unclear (Potikha and Delmer, 1995; Bischoff et al., 2010). Transcription factors involved in the trichome cell wall development include HOMEODOMAIN GLABROUS2 (Markus et al., 2009) and MYB DOMAIN PROTEIN106 (NOECK; Jakoby et al., 2008). Both of these mutants also have underdeveloped trichome papillae.

Only recently has the potential of Arabidopsis trichomes as a model for cell wall biogenesis been recognized (Suo et al., 2013). In fact, most of the published work on cell walls formation was done on stem/xylem or whole-plant analysis and is related to cell wall component biosynthesis (Liepmann et al., 2010), with recent emphasis on transcriptional networks reconstructed from transcriptional analyses (Wang et al., 2012).

Exocytosis is a constituent process in cell wall formation, and we are interested in how delivery/targeting and tethering of secretory vesicles to the plasma membrane are involved in Arabidopsis trichome cell wall maturation. Exocyst is an evolutionarily conserved protein complex in all eukaryotes and plays an important role in polarized exocytosis (Elias et al., 2003; Munson and Novick, 2006; Cvrčková et al., 2012). The exocyst complex consists of eight subunits (SEC3, SEC5, SEC6, SEC8, SEC10, SEC15, EXO70, and EXO84). Although it has been initially described in yeast (*Saccharomyces cerevisiae*; TerBush et al., 1996), homologs of all of the subunits have been described in plants (Elias et al., 2003), where they also form a complex (Hála et al., 2008). The role of the exocyst complex is tethering and docking of the post-Golgi vesicles to plasma membrane. SEC3 and EXO70 proteins are believed to be spatial landmarks on the plasma membrane, marking the place of the vesicle fusion (Finger et al., 1998; He et al., 2007). This can be shown by artificial targeting of the SEC3 to the mitochondrial outer membrane, which is followed by relocalization of the remaining exocyst subunits and secretion into the mitochondrial (Luo et al., 2014). The vesicle is attached by the exocyst near to the acceptor target membrane, whereas SNARE proteins mediate the rest of the fusion process. Polarized exocytosis mediated by exocyst enables effective control of growth and final shape of plant cells and tissues. The role of polarized exocytosis in cell wall formation is crucial and makes possible the creation of extremely polarized cells, such as root hairs, or pollen tubes (Cole et al., 2005; Cole and Fowler, 2006; Synek et al., 2006; Zárský et al., 2009).

Interestingly, some of exocyst subunits are found in multiple copies in terrestrial plant genomes (Elias et al., 2003; Hála et al., 2008; Cvrčková et al., 2012). The most protruding is multiplication of subunit EXO70. For example, the genome of *Physcomitrella patens* encodes 13 paralogs of the EXO70 subunit, *Vitis vinifera* encodes 15 paralogs, and *Oryza sativa* encodes 47 paralogs (Cvrčková et al., 2012). The Arabidopsis genome encodes 1 copy of exocyst subunits SEC6 and SEC8, 2 copies of SEC3, SEC5, SEC10, and SEC15, 3 copies of EXO84, and strikingly, 23 copies of EXO70 (Elias et al., 2003; Hála et al., 2008; Cvrčková et al., 2012). It still remains unclear whether the functions of EXO70 paralogs are redundant or specialized, but new findings suggest that at least some EXO70 paralogs have entirely different functions than the most studied and most conserved EXO70A1. Early data show that the transcription of some EXO70 paralogs is up-regulated in specific tissues or under specific conditions. For example, EXO7OB2 and EXO7OHI are strongly up-regulated by pathogen elicitors (Pecenkova et al., 2011). Other recent works report the first evidence, to our knowledge, of functional diversification of some EXO70 paralogs, with possible roles in autophagy (Kulich et al., 2013). Formation of other double-membrane bodies has been reported for the EXO70E2 subunit (Wang et al., 2010). Overexpression of EXO70E2 also induces double-membrane structures in animal cells (Ding et al., 2014). This may reflect an exocyst-independent effect of EXO70 on the membrane curvature, which was reported in animal cells (Zhao et al., 2013).

Here, we focus on the EXO70H4 paralog, which is the 11th most up-regulated gene in the mature trichome (Jakoby et al., 2008). Arabidopsis trichomes functions are mostly associated with herbivore defense and UV protection (Yan et al., 2012). Corresponding with this, it was indicated that EXO70H4 is up-regulated by UV-B irradiation in a CONSTITUTIVE PHOTOMORPHOGENIC1 (COP1)-dependent manner (Oravecz et al., 2006) and that it is down-regulated by methyl jasmonate (MeJA; Hruz et al., 2008). MeJA is a volatile plant defense hormone, which is important, especially in response to herbivores and wounding. A recent transcriptome profiling study has identified the cucumber (*Cucumis sativus*) EXO70H4 paralog as also being highly up-regulated during the cucumber fruit trichome development, suggesting that similar mechanisms are present among various land plants (Chen et al., 2014).

In this report, we studied and compared Arabidopsis trichomes maturation between the wild type and EXO70H4 subunit mutants. Our work has resulted in the description of a component and structure in Arabidopsis trichomes, which we named Ortmannian ring (OR). Additionally, our data highlight that EXO70H4 plays a role in Arabidopsis trichome cell wall maturation...
as well as plant response to UV-B irradiation and herbivore attack.

RESULTS

Two Arabidopsis SALK insertional mutants in the EXO70H4 single-exon gene were used for this study: exo70H4-1 and exo70H4-3 (SALK_023593 and SALK_003200, respectively). Annotated position of the insertions was verified by sequencing of the flanking regions. Both lines have transfer DNA insertion in the single exon of the EXO70H4 gene, and recessive homozygotes exhibit identical phenotypic deviations. exo70H4-1 has a tandem insertion, disrupting the coding sequence (CDS) 47 bp from the start codon, and exo70H4-3 has a single insertion, disrupting CDS 580 bp from the start codon (Fig. 1A). According to reverse transcription (RT)-PCR, both mutations prevent synthesis of the full-length RNA, with partial transcripts present (Fig. 1B). Because the phenotypes were identical, most of the following work has been done on the exo70H4-1 allele. Transformation of the exo70H4-1 mutant with the genomic fragment, including the promoter (895 bp) and the C-terminal fluorescent marker tagRFP (Entacmaea quadricolor red fluorescent protein), has restored the wild-type phenotype of the trichomes (Supplemental Fig. S1).

To test whether EXO70H4 is an exocyst subunit, we performed yeast two-hybrid analysis, which has resulted in interactions with SEC5a, SEC6, and the C terminus of EXO84b and also, a weak interaction with SEC15a (Supplemental Fig. S2).

Overall plant size, structure, and trichome morphogenesis of mutant plants are not affected. However, mutant plants have leaf trichomes that lack rigidity and are more flexible and bendable compared with the wild type. For example, in a blinded study, we were able to precisely distinguish 100% of the mutants from the wild types by touching the leaves (n = 40). This difference is because of the apparently thinner cell walls of mutant trichomes compared with the wild type (Fig. 1C). These observations imply that deposition of cell wall components is defective in exo70H4 mutants during the trichome maturation phase. At this phase, wild-type trichome cell wall undergoes massive secondary thickening, which progressively fills up trichome inner space. This thickening is mostly pronounced in the branch tips (Fig. 1C) and can be easily measured (Fig. 1D). This extensive deposition results in cytoplasm reduction and recession toward the stalk.

OR, Cell Wall Structure of the Mature Arabidopsis Trichome

During our observation of the trichome cell wall thickening, we noticed a distinct circular thickening of cell wall at the basal region of the stalk (Fig. 1C). This ring, present above the surrounding epidermal cells, appears to divide the trichome into two different domains. Staining with decolorized aniline blue has revealed that the ring is a callose-rich structure (Fig. 2A).

To show that such circular formation is a normal part of Arabidopsis trichome maturation, we captured photographs of whole 24-d-old rosettes stained by aniline blue (Fig. 2B) by automated stitching of individual microscopic frames in three focal depths. With this resolution, we could detect the presence of the callose ring in all leaf trichomes. First, callose rings appear in the oldest trichomes and grow on the apex. Second, a wave of maturation continues, and rings also appear in the leaf base. Because this ring was not studied before, we propose naming it OR after author J.O., who first spotted it.

The OR acts as a boundary and apparently divides the trichome cell wall into two clearly distinct domains—apical (branches and stalk) and basal (bulge). The cell wall in the apical domain shows strong autofluorescence upon UV-B excitation (Fig. 3), it has surface papillae, and it undergoes massive secondary thickening.
thickening during the maturation phase. In contrast, the basal domain lacks autofluorescence (Fig. 3A) and papillae (Fig. 1C), and its cell wall remains thin. The \textit{exo70H4-1} mutant lacks the OR and most of the callose in the vast majority of the trichomes (more than 95%). \textit{exo70H4-1} trichomes either lack the callose entirely or it can be detected within the cytoplasm in the form of ectopic fibrous bodies (Fig. 2A). No major differences in callose deposition were spotted elsewhere (plasmodesmatal plugs after wounding and vasculature; data not shown). Papillae do develop in the \textit{exo70H4} mutant; however, their distribution is not limited by any clear border area (Fig. 1C).

\textbf{exo70H4 Mutants Lack a Highly Autofluorescent Internal Cell Wall Layer That Is BORDERED by the OR in the Wild Type}

In the trichomes of the wild type, the internal cell wall layer can be easily recognized. This layer shows strong autofluorescence, especially upon excitation with 405 nm and emission maxima at 485 nm (Fig. 3B). Interestingly, this high autofluorescence is found only in the upper part of the trichome above the OR (Fig. 3A).

Autofluorescence of the inner cell wall layer is rapidly increased by trichome breakage or damage (Fig. 3, C and D). Because the damaged trichomes were killed by their breakage, gain of autofluorescence was likely caused by oxidation of the inner cell wall layer. This statement is supported by treatment with reducing agents (40 mM mercaptoethanol), which suppressed the autofluorescence buildup after breakage (Fig. 3E). With or without physical damage, \textit{exo70H4} mutants show dramatically decreased autofluorescence compared with the wild type because of the absence of the whole inner cell wall layer. The remaining autofluorescence was more equally distributed within the mutant cell wall (Fig. 3A). The absence of the inner trichome cell wall layer also explains differential calcofluor white (CW) staining of isolated \textit{exo70H4-1} trichome versus the wild type. Whereas in wild-type trichome, only the trichome base was stained, in the mutant, the apical domain was also stained (Supplemental Fig. S3). Hence, the callose-rich inner trichome cell wall layer likely insulates the apical trichome domain and prevents cellulose from staining.

To further characterize EXO70H4-dependent cell wall thickening, we have observed transmission electron microscopy (TEM) sections in isolated trichomes. Our sectioning confirmed that the wild-type trichome contains two clearly distinct cell wall layers—an electron-dense external layer and an internal layer. The internal layer is limited to the apical part of the trichome above the OR, where it forms a bulge of thick cell wall. This second internal cell wall layer also explains differential calcofluor white (CW) staining of isolated \textit{exo70H4-1} trichome versus the wild type. Whereas in wild-type trichome, only the trichome base was stained, in the mutant, the apical domain was also stained (Supplemental Fig. S3). Hence, the callose-rich inner trichome cell wall layer likely insulates the apical trichome domain and prevents cellulose from staining.

By extensive observation of trichomes in different stages of their maturation, we were able to summarize the thickening progress. The secondary cell wall thickening is accompanied by growth of the OR, which is U-shaped on the cross section and widens in the latest stages of the trichome maturation (Fig. 5).
UV-B-Induced EXO70H4 Expression Promotes Cell Wall Thickening and OR Biogenesis

According to several published microarray experiments (Hruz et al., 2008), EXO70H4 transcription is induced by UV-B irradiation. This induction is, moreover, COP1 dependent, linking expression of EXO70H4 with an already known UV response pathway (Oravecz et al., 2006). However, MeJA (a major regulator of necrotrophic pathogen and herbivore response; Jander and Howe, 2008) inhibits EXO70H4 transcription (Hruz et al., 2008). Therefore, we have designed experiments to test whether these environmental factors do affect trichome maturation in an EXO70H4-dependent manner.

We have isolated trichomes from fifth and sixth leaves of 24-d-old plants that were treated by UV-B irradiation and/or MeJA for 5 d. Isolated trichomes were then stained with decolorized aniline blue to visualize the OR. As can be seen in Figure 6A, UV irradiation clearly stimulates the callose ring formation, whereas MeJA suppresses it. If both treatments were applied simultaneously, their effects subtracted, resulting in OR abundance similar to in wild-type plants.

We have also examined the effect of UV-B on cell wall thickening in the trichome stalk above the OR. Despite the large defects of cell wall thickening in the trichome apices (Fig. 1C), the differences were rather small but highly significant; UV-B treatment stimulated wild-type thickening by about 10% in contrast to no stimulation in the exo70H4 mutant. Combined UV and MeJA treatment caused the 10% thickening in both the wild type and exo70H4, indicating that this particular mutant phenotype is complemented by an unknown MeJA-dependent mechanism under these combined treatment conditions (Fig. 6B).

Ecophysiological Aspects of Trichome Cell Wall Thickening

Because trichomes have been previously reported to also play a role in anti-herbivore protection, we next asked whether thick secondary walls of wild-type trichomes facilitate plant resistance against insect pests. For this, we performed a weight gain experiment using
a specialist herbivore (cabbage butterfly [*Pieris brassicae*]) and a generalist herbivore (Egyptian cotton leafworm [*Spodoptera littoralis*]). First instar caterpillars were placed on both wild-type and *exo70H4* plants for 7 d, after which dry weight gain was measured. For both herbivore species, caterpillars grew less on the *exo70H4* mutants compared with on wild-type plants (Supplemental Fig. S4A). In accordance, we also observed that endogenous MeJA content in *exo70H4* mutants was generally higher than in wild-type plants (Supplemental Fig. S4B). Because jasmonic acid (JA) induction has been linked to increased herbivore resistance in Arabidopsis, we suggest that the influence of trichome stiffness is hazed by other effects, likely MeJA-induced glucosinolate synthesis and overall resistance.

Copper Is Accumulated in the Cell Wall of the OR and above It

Trichomes of other species are well known to accumulate heavy metals, so we were interested if they could be detected in the secondary cell wall and the OR. We have observed second and third true leaves of 24-d-old plants watered by 2 mM CuSO₄ and water. Then, copper was stained using dithiooxamide. The copper mainly accumulated in the OR and the cell wall band right above it, whereas it was not detected in the samples watered by regular water. Also, copper was not detected in the *exo70H4*-1 mutant, where the whole secondary cell wall is absent (Supplemental Fig. S5).

OR Is Not Likely a Rudiment of Cytokinesis

Arabidopsis trichomes are of unicellular origin; however, DNA endoreduplication suggests their multicellular origin (Hülskamp et al., 1994). In the multicellular trichomes of cucumber, EXO70H4 paralogs are also highly up-regulated (Chen et al., 2014). In these trichomes, the two apical cells also accumulate big amounts of callose, and callose rings are common at the cell divisions (Supplemental Fig. S6).

### Figure 4

TEM micrographs of wild-type and *exo70H4*-1 trichomes. A, Wild-type cell wall above the OR has two distinct layers. B, Wild-type cell wall at the OR. Above the OR, two layers can be recognized, but only one layer is visible beneath the OR. C, *exo70H4*-1 cell wall at the place corresponding to A. Only one cell wall layer is visible. D, *exo70H4*-1 cell wall at the place corresponding to B. No OR is visible. E, The blue dotted line highlights the region that forms the OR according to the immunogold labeling. The orange square represents the approximate position of F. F, Detail on the immunogold labeling by the anticallose antibody. The position of F is highlighted by the orange square in E. G, Position and orientation of micrographs depicted for better orientation. Bars = 2 μm (A–E) and 100 nm (F).

### Figure 5

Development of the OR. A, Stages of the trichome maturation accompanied by the secondary cell wall thickening and OR growth. Bar = 20 μm. B, Graphical portrayal of A. Cyan, The OR; purple, the secondary cell wall above the OR; brown, the primary cell wall (respective cellulose-rich cell wall).
Hence, we asked whether Arabidopsis OR is a rudiment of cell plate formation. To address this question, we used the microtubules reporter MICROTUBULE ASSOCIATED PROTEIN4-GFP (MAP4-GFP)-expressing Arabidopsis plant. A band of microtubules distantly resembling a preprophasic band was observed in many trichomes during our observations, and we spotted a correlation between the nucleus position and OR biogenesis (Fig. 7, A–D). Previous studies concluded the final position of the nucleus at the point of the third branching (Hülskamp et al., 1994). We carefully monitored position dynamics of the nucleus in maturing trichomes and realized that, in finally matured/thickened wild-type trichomes, the nucleus, in fact, moved back to basal position. In wild-type trichomes without OR, the nucleus was mostly spotted at the branching point. However, the position of the nucleus was much more basal in the trichomes with OR (Fig. 7, A–C). To test the possibility that OR is a rudiment of cytokinesis, we observed OR in the Siamese (sim) mutant (GABI_170C01). SIM encodes an Interactors of Cdc2 kinase/Kip-related protein cell cycle inhibitor, and mutation in this gene results in ectopic trichome cell divisions (Walker et al., 2000). Surprisingly, these trichomes did not divide in the OR domain (Fig. 7E). Moreover, additional ORs were present in newly divided branches. Hence, this observation did not support the hypothesis that the OR would be a simple rudiment of preprophase or cytokinesis.

**DISCUSSION**

Here, we show that the final stages of trichome development (i.e. trichome maturation) are characterized by the deposition of an additional layer of the cell wall. Because it is clearly distinguishable and deposited in the last stage of trichome development, we refer to this layer as the secondary cell wall, although biochemical analysis suggested that it has primary wall-like composition (Marks et al., 2008). Strong argument for this cell wall being secondary is the fact that this layer is only formed in the apical part of the trichome and not in the base. The secondary cell wall layer is bordered by a previously undescribed structure, which we propose to name the OR. OR not only borders the secondary cell wall but also, limits the surface papillae distribution, which is much more delimited in the exo70H4 mutant.

The scenario of OR being a phylogenetic relic of preprophase is unlikely, although our observations do not rule out any other link to the cell cycle. Other phenomena observed can also be explained by different mechanisms. It was shown previously in multiple cases that the nucleus moves toward places with massive secretion (for example, during plant-microbe interactions; Grif et al., 2014). Also, microtubular structures are often present in active secretory plasma membrane domains, such as the mucilage secretion domain of the outer layer of Arabidopsis seed coat (McFarlane et al., 2008).

OR has been anecdotally spotted before, however, without any additional attention. In figure 1m in Potikha and Delmer (1995), a callose ring was noticed in the trichome, but it was interpreted as a circular wound response. Here, we show that the OR constantly appears with trichome maturation, resulting in nearly 100% of the mature trichomes having the OR. In sum, we propose that, as the trichome matures, most of its volume is filled up by the callose-rich cell wall, and this thickening is accompanied by the growth of the OR. It should be also noted that callose patterns in the secondary cell wall may vary and are not always as we described here as general pattern. The most common patterns above the OR are small speckles spreading down from the branch tips. Other than the OR, a wide callose band repeatedly appears on the stalk. Interestingly, sometimes not only one OR but two or more are formed. Controlling mechanisms and conditions of these patterns remain unclear and have yet to be studied.

**Role of the OR and the Autofluorescent Secondary Cell Wall Layer**

The precise role of OR is yet unknown; however, because it borders the autofluorescent and callose-rich inner layer of the cell wall, it likely has a structural
The role of the inner layer may be more general and important for many ecophysiological aspects, including UV protection and heavy metal detoxification. Because many toxic compounds, including secondary metabolites, are expected to be deposited in the trichome cell wall, the OR may limit their localization from spreading into other tissues. For example, sequestration of heavy metals in trichomes has been described in both hyperaccumulating and nonaccumulating plant species (Martell, 1974; Salt et al., 1995; Krämer et al., 1997; MacFarlane and Burchett, 1999; Servin et al., 2012; Jr and Kupper, 2014). Arabidopsis halleri is a well-known heavy metal hyperaccumulator, with strong evidence for accumulation of zinc, cadmium, and other heavy metals in trichomes. Subcellular localization is in accordance with our results with striking cellular distribution in a narrow ring or thick band at the trichome base (Küpper et al., 2000; Zhao et al., 2000; Sarret et al., 2002; Fukuda et al., 2008), corresponding to the secondary cell wall above the OR. There is also evidence of cadmium hyperaccumulation in trichomes of Arabidopsis (Ager et al., 2002). Jakoby et al. (2008) have shown that zinc transporters and copper chaperones are on the list of the 5% most up-regulated genes in the mature trichome. Hence, the Arabidopsis trichome secondary cell wall may be a permanent storage for toxic heavy metals, which we show here with copper. In this regard, Arabidopsis simple trichomes are analogous to the
glandular trichomes of other species, which also usually have cell wall domains specialized for phytochemical deposition (Turner and Croteau, 2004; Ramirez et al., 2012). Heavy-metal stress has been shown previously to up-regulate the secretory pathway genes in *O. sativa*, including EXO70FX14 and EXO70FX15, which were induced by the copper stress. Similarly, *Nicotiana benthamiana* EXO70 is up-regulated by the copper stress and required for the heavy metal-induced reactive oxygen species production (Lin et al., 2013).

We do not yet know the nature of the autofluorescent compounds stored in the trichome secondary cell wall. We have tried multiple lignin stainings, which all gave us negative results, with both healthy and broken trichomes (data not shown).

Interestingly, the whole process of callose accumulation, cell wall thickening, and OR development is enhanced by UV-B irradiation. Trichomes, indeed, play a role in the UV stress response, and their density is increased in plants exposed to UV-B (Yan et al., 2012). UV-B-induced trichome cell wall thickening is in accordance with these results. Induction of EXO70H4 by UV light, resulting in trichome cell wall thickening, indicates a possible function in apical meristem protection against damaging ROS production, although we have not yet performed experiments to confirm this.

The inner trichome cell wall layer also contains a lot of callose, and its absence in *exo70H4* mutants does not cause a change in the trichome birefringence (data not shown). Therefore, the birefringent cell wall layer is likely the outer one, which is also more electron dense on the TEM and contains dense cellulose fibers. This observation is also supported by our CW staining, in which the inner cell wall layer was not stained by the CW in contrast to the *exo70H4* mutant. The presence of ectopic fibrous callose patches in *exo70H4* mutants suggests that callose rather than callose synthase is one of EXO70H4-dependent cargoes. Because the whole internal cell wall layer is callose rich, its development may be callose dependent. Occasional presence of the OR in the *exo70H4* mutant is likely caused by existence of a very close paralog EXO70H3, which is a twin gene next to EXO70H4, similar to SEC10a and SEC10b (Vukašinović et al., 2014). EXO70H3 shows a pollen-specific expression; however, like many other genes, there may be some ectopic expression in the sporophyte. In Arabidopsis pollen, more EXO70 subunits of the H group are expressed, which is probably why *exo70H3* has viable and functional pollen with normal size of the callose plugs (data not shown).

To our surprise, simultaneous application of both UV-B and MeJA induced cell wall thickening in both the *exo70H4* mutant and the wild type without the induction of the OR biogenesis. This may be caused by up-regulation of another EXO70 paralog (for example, EXO70H7, which is stimulated by MeJA on the transcriptional level; Hruz et al., 2008) or another exocyst-independent mechanism.

Because our larval weight experiment did not reveal any importance of *exo70H4*-mediated cell wall thickening for herbivore resistance, we speculate that EXO70H4 is essential for default UV-B-induced thickening, which may be modulated upon herbivore attack. Trichome stiffness itself, however, may still be important to protect against another herbivore species (for example, slugs), and it may be a more general way to improve plant fitness.

**Exocyst and Trichome: Questions of the Future Cell Wall Biogenesis Investigation**

According to the transcriptomic data (Oravecz et al., 2006; Jakoby et al., 2008), EXO70H4 is one of the most up-regulated genes in the mature trichome. Surprisingly, this high expression was not noticed in GUS reporter promoter activity assays (Li et al., 2010). This may be because of early observations of too young trichomes combined with very low permeability of old ones.

The EXO70H4 paralog up-regulation in cucumber trichomes (Chen et al., 2014) is a clue that mechanisms of the cell wall thickening in Arabidopsis described here may be a more general phenomenon for land plants that also include multicellular trichome types.

Other (core) exocyst subunits are not dramatically up-regulated in the trichome. Hence, there is a question of the stoichiometry of the complex. Recently, mammalian EXO70 was shown to also function independently from the rest of the exocyst complex in a way analogous to Bin–Amphiphysin–Rvs domain proteins functioning in a membrane deformation (Zhao et al., 2013). We cannot yet exclude the possibility of exocyst-independent function; however, it is worth noting that EXO70H4 interacts with other exocyst subunits in the yeast two-hybrid assays (Supplemental Fig. S2). The stoichiometry question could be explained by post-transcriptional regulation of the EXO70H4 level, which is the case in many other Arabidopsis EXO70s (Stegmann et al., 2012), or possibly, high doses of this subunit are important to compete with the other EXO70s for the rest of the complex, thus modulating its function as proposed in Žárský et al., 2013. Our work aims to answer some of these questions and includes more general analysis of exocyst-dependent callose deposition processes in defense responses.

**MATERIALS AND METHODS**

**Plant Growth and Conditions**

T-DNA insertion lines of Arabidopsis (*Arabidopsis thaliana*) were obtained from the Nottingham Arabidopsis Stock Centre (Scholl et al., 2000). These lines were crossed with MAP4-GFP lines (Marc et al., 1998). Plants were grown in Jiffy soil pellets under long-day conditions with illumination of 100 μm m⁻² s⁻¹ photosynthetically active radiation (OSRAM L 58W/930). Plants were planted, placed into paper boxes, and sealed from above by filter foils. Rosco 226 (UV filter) and Rosco 130 (control filter) were used.

Subsequently, plants were exposed to UV-B radiation for 5 d under LT 36W/958 T8 BIOVITAL NARVA fluorescent tubes (which contain UV-B emission peaks) and/or treated with MeJA by installation of slices of filter.
papers soaked in the MeAq amount corresponding to a concentration of 0.8 μL L⁻¹ solution (box volume also for 5 d).

In high-light conditions, plants were grown under 700 μm m⁻² s⁻¹ photosynthetically active radiation with Philips HPI-T Plus 400W/645 E40 light bulbs (also with UV-B peak).

Genotype Analysis

The following primers were used for genotyping: exo70H4-1, left primer (LP), TGCGATTCCTTTCAACATGTC and right primer (RP), AATCCTCATCAGCACAGCC. From exo70H4-1, LP, AACACACCTGGAACCCCGAG and RP, GGTCGTAAGTCGAGGAGG; SaklLb 1.3, ATTTGGAGCTTGGGCGAC; GABI 1700C1, LP, ATGGATGAACTGCGAGTGG and RP, TGGAAATG-TACCTCTTGGCAC; and GABI LB 08409, ATATTGACATCATACATATTG.

The following primers were used for cloning fragments for genomic complementation: EXO70H4 Pst1_for, AAGCTGCAGGCTACAGGAGGCTGGATCATC; and EXO70H4 Kpn1_rev, TCAGTCTACCGACATCGATGCTTCCGAGCC.

The following primers were used for testing its function: H3seq3 for, CCGGAGATCATTAATACTAAAA; and TagRFPRev, ATCAAACCTTCACCTTACCTCCCC.

This fragment has been subcloned into the TagRFP AS-N vector (Evrogen), transferred to the binary vector pBGW (Karimi et al., 2002) by the Clontech LR clonase recombination, and sequenced.

Transcript Detection and Semiquantitative RT-PCR

For detection of transcript in exo70H4-1 and exo70H4-3 mutants, RNA was isolated from 100 mg of 7-d-old seedlings grown on Murashige and Skoog plates. RT-PCR was done with the Fermentas First-Strand cDNA Synthesis Kit isolated from 100 mg of 7-d-old seedlings grown on Murashige and Skoog series and embedded in Epon-Araldite.

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The main effects of the three different genotypes (the wild type, exo70H4-1, and exo70H4-3), the two herbivore species (Egyptian cotton leafworm and cabbage butterfly), and their interaction were analyzed with two-way ANOVA.

JA content in leaves was analyzed by collecting one leaf per plant. The extraction was performed by grinding 200 mg of fresh leaves to a powder and mixing with 990 μL of extraction solvent (ethyelacetate/formic acid at 99:0.5:0.5) and 10 μL of internal standards (containing isotopically labeled d5-JA), and evaporation of the supernatant, the residue was resuspended in 100 μL of 70% (v/v) MeOH. Five microliters of solution was injected for ultra-high-performance liquid chromatography-mass spectrometry/mass spectrometry analysis following the same conditions as in Glauser et al. (2014). The final concentration of JA is expressed in nanograms per gram times fresh weight. The main effects of the three different genotypes (the wild type, exo70H4-1, and exo70H4-3), the herbivore treatment (Egyptian cotton leafworm, cabbage butterfly, and control), and their interaction on JA concentrations were analyzed with two-way permutation ANOVA using the lmperm package in R (Wheeler, 2010).

Yeast Two-Hybrid Assays

The yeast (Saccharomyces cerevisiae) two hybrid was performed as described previously (Hála et al., 2008). We used the Matchmaker GAL4 Two-Hybrid System 3 (Clontech) following the manufacturer’s instructions. We used constructs prepared by Hála et al. (2008), Fendrich et al. (2010), and Peenková et al. (2011), and EXO70H4 cds was amplified from genomic DNA and subcloned into the pGBKKT vector between NcoI and SalI using these primers: For_EXO70H4_NcoI, TTTCATGAGTGGAAGAAGACGAT; and For_EXO70H4_SalI, TTTCGACTACGACAGGATGCTTGAG. From this vector, EXO70H4 cds was recloned into pGADT7 using Xfel and Xhol restriction sites.

Supplemental Data

The following supplemental materials are available.

Supplemental Figure S1. Genetic complementation of the exo70H4-1 mutant.

Supplemental Figure S2. Two hybrid interactions of EXO70H4 and exocyst subunits.

Supplemental Figure S3. Calcofluor white staining of the wild-type and exo70H4-1 mutant trichomes.

Supplemental Figure S4. Herbivore studies on exo70H4 mutants.
Supplemental Figure S5. Copper accumulation in wild-type and exo70H4-1 trichomes.

Supplemental Figure S6. Callose in the cucumber trichome.

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