**Arabidopsis** membrane-associated acyl-CoA-binding protein ACBP1 is involved in stem cuticle formation

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Abstract

The membrane-anchored *Arabidopsis thaliana* ACYL-COA-BINDING PROTEIN1 (AtACBP1) plays important roles in embryogenesis and abiotic stress responses, and interacts with long-chain (LC) acyl-CoA esters. Here, AtACBP1 function in stem cuticle formation was investigated. Transgenic *Arabidopsis* transformed with an AtACBP1pro::GUS construct revealed β-glucuronidase (GUS) expression on the stem (but not leaf) surface, suggesting a specific role in stem cuticle formation. Isothermal titration calorimetry results revealed that (His)6-tagged recombinant AtACBP1 interacts with LC acyl-CoA esters (18:1-, 18:2-, and 18:3-CoAs) and very-long-chain (VLC) acyl-CoA esters (24:0-, 25:0-, and 26:0-CoAs). VLC fatty acids have been previously demonstrated to act as precursors in wax biosynthesis. Gas chromatography (GC)–flame ionization detector (FID) and GC–mass spectrometry (MS) analyses revealed that an acbp1 mutant showed a reduction in stem and leaf cuticular wax and stem cutin monomer composition in comparison with the wild type (Col-0). Consequently, the acbp1 mutant showed fewer wax crystals on the stem surface in scanning electron microscopy and an irregular stem cuticle layer in transmission electron microscopy in comparison with the wild type. Also, the mutant stems consistently showed a decline in expression of cuticular wax and cutin biosynthetic genes in comparison with the wild type, and the mutant leaves were more susceptible to infection by the necrotrophic pathogen *Botrytis cinerea*. Taken together, these findings suggest that AtACBP1 participates in *Arabidopsis* stem cuticle formation by trafficking VLC acyl-CoAs.

Key words: Acyl-CoA-binding protein, *Arabidopsis thaliana*, cuticle, cuticular wax, cutin, very-long-chain acyl-CoAs.

Introduction

The *Arabidopsis* cuticle is a lipophilic layer that consists of cutin and wax (Nawrath, 2002; Kunst and Samuels, 2009; Lee and Suh, 2013). Cutin is a polymer derived from hydroxy and epoxy-hydroxy C16 and C18 fatty acids (Nawrath, 2006). Cutin biosynthesis consists of sequential reactions including the activation of acyl chains to coenzyme A by long-chain acyl-CoA synthetase (LACS), hydroxylation and epoxidation catalysed by the cytochrome P450 family, and esterification

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to glycerol-3-phosphate by glycerol-3-phosphate acyltransferases (GPATs) (Li-Beisson et al., 2013). Waxes, including epicuticular waxes that cover the cuticle membrane and intracuticular waxes embedded in the cuticle membrane, are complex mixtures of alcohols, alkanes, aldehydes, ketones, and esters derived from long-chain fatty acids (Jenks et al., 2002). In plant epidermal cells, saturated very-long-chain fatty acids (VLCFAs), comprising acyl chains exceeding 20 carbons (>C20) generated by the extension of C16 and C18 fatty acids in the endoplasmic reticulum (ER), form precursors for the synthesis of alliphatic components of cuticular waxes (Kunst and Samuels, 2003).

Acyl-CoA-binding proteins (ACBPs) constitute a family of eukaryotic proteins that show conservation in an acyl-CoA-binding (ACB) domain with ability to bind long-chain acyl-CoA esters (Knudsen et al., 2000; Xiao and Chye, 2009, 2011a; Fan et al., 2010; Yurchenko and Weselake, 2011). In the model plants Arabidopsis thaliana and Oryza sativa, six genes designated as AtACBP1–AtACBP6 and OsACBP1–OsACBP6, respectively, encode ACBPs that bind acyl-CoA esters and phospholipids with varying affinities (Engeseth et al., 1996; Chye, 1998; Chye et al., 2000; Leung et al., 2004, 2006; Chen et al., 2008; Gao et al., 2009; Xiao et al., 2009; Meng et al., 2011, 2014; Du et al., 2013b). Variation in subcellular localization has also been observed (Chye et al., 1999; Li and Chye, 2003; Leung et al., 2006; Xiao et al., 2008b; Xiao and Chye, 2009; Meng et al., 2014). Thus, given the differences in subcellular localization and substrate preference, it appears that some Arabidopsis ACBPs perform distinct cellular functions in vivo while others with similar subcellular localization and binding affinities for acyl-CoA esters may share overlapping roles (Chye et al., 1999, 2000; Li and Chye, 2003; Leung et al., 2006; Xiao et al., 2008b; Xiao and Chye, 2009, 2011a). For example, AtACBP1 and AtACBP2 both show functions in embryogenesis (Chen et al., 2010) and seedling development (Du et al., 2013a, b), while AtACBP3 promotes starvation-induced leaf senescence, age-dependent leaf senescence (Xiao et al., 2010), and plant defence against Pseudomonas syringae (Xiao and Chye, 2011b; Zheng et al., 2012). AtACBPs are also associated with heavy metal/oxidative (Xiao et al., 2008a; Gao et al., 2009, 2010), freezing (Chen et al., 2008; Du et al., 2010; Liao et al., 2014), and drought (Du et al., 2013a) stresses. AtACBP1 (Du et al., 2013b) and AtACBP2 (Gao et al., 2009, 2010) both mediate protein–protein interactions by binding transcription factors and stress-responsive partners.

AtACBP1 was subcellularly localized to the ER and the plasma membrane (PM) (Li and Chye, 2003), and immunoelectron microscopy using anti-AtACBP1 antibody revealed that the AtACBP1 protein accumulates in developing embryos (Chye et al., 1999). The roles of AtACBP1 in embryo development were confirmed by phenotypic and biochemical studies using the acbp1 T-DNA insertional mutant (Chen et al., 2010). Alterations in membrane lipid composition and acyl-CoA content in the acbp1 siliques were observed. In addition, the observation of arrest of early embryo development in the acbp1acbp2 double mutant suggested that AtACBP1 and AtACBP2 are essential during early embryogenesis in Arabidopsis (Chen et al., 2010), most probably in lipid transfer because (His)$_2$-tagged recombinant AtACBP1 (rACBP1) and AtACBP2 bind acyl-CoA esters and both demonstrated preference for unsaturated over saturated long-chain acyl-CoA esters (Chye, 1998; Chye et al., 2000; Leung et al., 2006; Gao et al., 2009).

AtACBP1 has been observed to accumulate in the outer integument cells of the developing seed coat and has been previously proposed to be involved in the biosynthesis of cutin and cuticular waxes (Chye et al., 1999). A membrane-associated ACBP from Agave americana showing 62% amino acid identity to AtACBP1, AaACBP1, was enriched in the epidermis of mature leaves (Guerrero et al., 2006). Besides the presence of the conserved ACB domain, AaACBP1 contains ankyrin repeats which potentially mediate protein–protein interactions (Michaely and Bennett, 1992) similarly to AtACBP1 (Xiao and Chye, 2011a). The expression of AaACBP1 in the epidermal cells (Guerrero et al., 2006) and observation of AtACBP1 localization at the endomembranes (Chye et al., 1999) support the feasibility of these ACBPs as candidates involved in the biosynthesis of cuticular lipids (Chye et al., 1999; Kunst and Samuels, 2003; Xiao and Chye, 2011a; Li-Beisson et al., 2013). Interestingly, in mice, ACB has been reported to be essential in the formation of an epidermal barrier (Bloksgaard et al., 2012), while Xia et al. (2012) demonstrated that AtACBP3, AtACBP4, and AtACBP6 function in cuticle formation in Arabidopsis. To address the role of AtACBP1 in cuticle formation, it was first demonstrated that (His)$_2$-tagged rACBP1 can bind very-long-chain (VLC) acyl-CoA esters in vitro. Subsequently, investigations on the acbp1 T-DNA insertional mutant showed that it displayed reduction in cuticular wax and cutin monomer composition in Arabidopsis stems, suggesting that AtACBP1 functions in stem cuticle formation.

**Materials and methods**

**Plant materials and growth conditions**

Seeds of wild-type A. thaliana (ecotype Col-0), the acbp1 mutant (SAIL-653 B06, ecotype Col-0; Xiao et al., 2008a), and acbp1-1COM (Xiao et al., 2008a) were surface-sterilized, cold-stratified, and germinated on Murashige and Skoog medium (MS medium) (Murashige and Skoog, 1962) supplemented with 2% sucrose for 10 d under cycles of 8 h dark (21 °C) and 16 h light (23 °C). Plants were transferred into soil and were grown in a growth chamber under a 16 h light/8 h dark cycle. Stems were harvested from 6-week-old plants for gas chromatography (GC) analysis following Lee et al. (2009b).

**Expression and purification of rACBP1**

(His)$_2$-AtACBP1 recombinant protein was expressed in the soluble fraction of Escherichia coli BL21(DE3), and was purified through Ni-NTA agarose (Qiagen, Valencia, CA, USA) affinity columns as previously described (Chye, 1998).

**Isothermal titration calorimetry (ITC) measurements**

ITC experiments were performed using an isothermal titration calorimeter (MicroCal iTC$_{200}$ system) from MicroCal Inc. (USA). Long-chain and VLC acyl-CoA esters used in this study were purchased from Avanti Polar Lipids (http://www.avantilipids.com). The acyl-CoA concentration (250 μM) in the titration syringe was 25-fold
higher than the protein concentration (10 μM) in the cell. Acyl-CoA solutions and rACBP1 protein were degassed under vacuum and stirred immediately before use. The experiments were performed at 30 °C, and injections were initiated after equilibration to baseline stability. Each injection was made up to a volume of 1.5 μl and lasted 10 s, with an interval of 240 s between injections. The syringe was rotated at 1000 rpm during the assay to ensure immediate mixing. Raw data were integrated, corrected for non-specific heat, and analysed using the ORIGIN software supplied with the instrument by the General Electric Company. The dissociation constant ($K_D$) was calculated by non-linear regression fitting the isotherm.

β-Glucuronidase (GUS) histochemical assays

GUS histochemical assays were carried out on AtACBP1::GUS Arabidopsis transformed with construct pAT352 according to Du et al. (2013b). The standard 5-bromo-4-chloro-3-indolyl-β-d-glucuronide (X-Gluc) solution (100 mM sodium phosphate buffer, pH 7.0, 0.1% Triton X-100, 1 mg ml$^{-1}$ X-Gluc) with the addition of 2 mM potassium ferricyanide and 2 mM potassium ferrocyanide was used. Leaves and stems from 4-week-old AtACBP1::GUS transgenic Arabidopsis were vacuum-infiltrated in X-Gluc solution for 30 min and kept at 37 °C until a blue colour developed. Samples were destained in 70% ethanol and photographed. The controls in the GUS assays were samples from Col-0 and transgenic Arabidopsis transformed with vector pBI101.3, and they were not stained blue during the same incubation period. The GUS-stained stems and leaves were embedded in Paraplast for sectioning according to Sin et al. (2006).

Quantitative real-time polymerase chain reactions (qRT-PCRs)

Total RNA was isolated from stems of five 5-week-old Arabidopsis plants using the RNeasy Plant Mini Kit following the protocol provided by QIAGEN. RNA (3.5 μg) was reverse-transcribed into cDNA using the SuperScript First-Strand Synthesis System (Invitrogen). PCR was conducted on a StepOne Plus Real-time PCR system using SYBR Green (Invitrogen). PCR was performed with gene-specific primers, and Arabidopsis ACTIN2 (At1g48300), At1g49430 (At4g00360), At2g19450 (At2g19450), At3g51520 (At3g51520), At3g53630 (At3g53630), At4g00360 (At4g00360), At4g04930 (At4g04930), At4g42450 (At4g42450), At5g13640 (At5g13640), At5g52310 (At5g52310), and At5g52540 (At5g52540) were used as internal controls (Supplementary Table S1 available at JXB online). The relative expression of the targeted gene was normalized using the ACTIN2 control as described (Xiao et al., 2011b).

Scanning electron microscopy (SEM)

Stems of the first internodes above the rossette from 6-week-old wild-type Arabidopsis and the acbp1 mutant were used. Samples were treated with 1% osmium tetroxide (OsO$_4$) for 24 h and then air-dried for 3 d, followed by mounting onto standard aluminum stubs and sputter coating with gold particles using six 30-s bursts according to Chen et al. (2003). The coated samples were viewed with a Hitachi S3400 scanning electron microscope.

Transmission electron microscopy (TEM)

The ultrastructure of 6-week-old stems from wild-type Arabidopsis and the acbp1 mutant were prepared for TEM following Siebet et al. (2000) with some modifications. Samples were fixed using 2.5% glutaraldehyde in cacodylate buffer (0.1 M sodium cacodylate-HCl buffer, pH 7.4) for 4 h at 4 °C, followed by post-fixation treatment with 1% osmium tetroxide in cacodylate buffer for 4 h at 4 °C. After gradient dehydration with ethanol, samples were infiltrated overnight in an epoxy resin/proplyene oxide 1:1 mixture. This was followed by infiltration overnight in epoxy resin. Samples were subsequently embedded in epoxy resin and polymerized overnight at 60 °C. Ultrathin (60 nm) sections were prepared and stained with 2% uranyl acetate and lead citrate (Li et al., 2008), and subsequently examined using a Phillips CM100 transmission electron microscope.

Wax analysis

Cuticular waxes were extracted by immersing two stem segments (each 10-cm in length) in 5 ml of chloroform for 30 s. The internal standards used were C28 alkane (n-octacosane), C22 fatty acid (docosanoic acid), and C23:0 fatty alcohol (1-tricosanol). The solvent was then removed by heating (40 °C) under a mild stream of nitrogen. Derivatization was performed by adding 100 μl of pyridine and 100 μl of bis N,N-(trimethylsilyl) trifluoroacetamidat (Sigma) to the dried extract and incubating for 30 min at 90 °C. The qualitative composition was then evaluated by capillary GC–mass spectrometry (GC–MS; GCMS-QP2010; Shimadzu). GC-MS conditions were as follows: injection at 220 °C, maintenance of the temperature at 220 °C for 4.5 min, followed by an increase to 290 °C at a rate of 3 °C min$^{-1}$. The temperature was then maintained at 290 °C for 10 min, after which it was raised to 300 °C at a rate of 2 °C min$^{-1}$ and held for 10 min (Lee et al., 2009b). Analysis of quantitative wax materials was performed using a capillary GC program with a flame ionization detector (FID) using the same conditions as in GC-MS. Compounds were quantified relative to the corresponding internal standards by integrating the peak areas.

Cutin analysis

Five- to six 6-week-old primary stems of Arabidopsis were used for cutin analysis according to Lee et al. (2009a). The internal standards used were C17:0 methyl ester (methyl heptadecanoate) and C15:0 cycloketone (o-pentadecaketone) (Sigma). Polymers from dried solvent-extracted residues of stems (wax-free) were depolymerized by hydrolysis with methanolysis using sodium methoxide. The products recovered after hydrolysis were dried and derivatized as mentioned above and separated and quantified by GC-MS. The GC-MS protocol was as follows: injection at 110 °C, elevation by 2.5 °C min$^{-1}$ to 300 °C, and holding for 3 min at 300 °C. The mass-to-charge ratios (m/z) used to diagnose the cutin compounds are shown in Supplementary Table S2 at JXB online.

Inoculation of plants with Botrytis cinerea

The necrotrophic fungus B. cinerea was maintained on a potato dextrose agar plate (BD Difco) at room temperature. Collection of conidia and plant infection assays (Botrytis suspension concentration of 2 × 10$^5$ spores ml$^{-1}$) were carried out as previously described (Li et al., 2008; Xiao and Chye, 2011b). Photographs were taken at 0 and 6 days after infection (DAI).

Accession numbers

Sequence data from this article can be found in the Arabidopsis Genome Initiative database under the following accession numbers: At5g53470 (ACBP1), At3g18780 (ACTIN2), At2g47240 (CER8), At1g67730 (CKR1), At1g55360 (ECR), At1g68530 (CUT1/KCS6), At1g49430 (LACS2), At4g00360 (CYP86A2), At1g01600 (CYP86D4), At4g04930 (GPAT8), At2g19450 (DGAT1), At3g51520 (DGAT2), At1g48300 (DGAT3), At5g13640 (PDAT1), At1g20440 (COR47), At5g52310 (LTT8), At4g25490 (CBF7), and At3g26744 (ICL1).

Results

Recombinant AtACBP1 binds VLC acyl-CoA esters (C24:0-, C25:0-, and C26:0-CoA) in vitro

(His)$_6$-tagged rACBP1 was shown to bind long-chain acyl-CoA esters (C18:1-, C18:2-, and C18:3-CoA) in Lipidex
assays (Leung et al., 2006) and gel-binding assays (Chye, 1998). As rACBPs have not been reported to bind VLC acyl-CoA esters, rACBP1 was tested using commercially available VLC acyl-CoA esters (C24:0-, C25:0-, and C26:0-CoA) by ITC to determine the $K_D$ values. As controls, long-chain acyl-CoA esters (C18:1-, C18:2-, and C18:3-CoA) were included. Analysis of calorimetric data by the ORIGIN software (General Electric Company, USA) indicated that the binding isotherms fitted well with a model of a single binding site (Fig. 1). Consistent with the results from Lipidex assays (Leung et al., 2006) and gel-binding assays (Chye, 1998), rACBP1 interacted with long-chain acyl-CoA esters including C18:1-, C18:2-, and C18:3-CoAs with high affinities in ITC measurements (Fig. 1; Table 1). Furthermore, rACBP1 was also shown to bind to VLC acyl-CoA esters (C24:0-, C25:0-, and C26:0-CoA), although with lower affinities (Fig. 1; Table 1). The affinities as reflected by the $K_D$ values of rACBP1 for VLC acyl-CoA esters supported their participation in fatty acid elongation during the biosynthesis of VLCFAs.

AtACBP1 is expressed in stem epidermis

AtACBP1 mRNA has been previously reported to be expressed in all plant organs (Chye, 1998; Chen et al., 2010). The microarray database e-FP Browser (Winter et al., 2007; http://www.bar.

Table 1. The dissociation constants ($K_D$) of recombinant AtACBP1 (rACBP1) binding to acyl-CoA esters of different acyl chain lengths

The values are means ± SD ($n=3$).

<table>
<thead>
<tr>
<th>Acyl-CoA esters</th>
<th>$K_D$ (μM)</th>
</tr>
</thead>
<tbody>
<tr>
<td>C18:1</td>
<td>0.76 ± 0.15</td>
</tr>
<tr>
<td>C18:2</td>
<td>0.83 ± 0.04</td>
</tr>
<tr>
<td>C18:3</td>
<td>0.44 ± 0.01</td>
</tr>
<tr>
<td>C24:0</td>
<td>2.14 ± 0.13</td>
</tr>
<tr>
<td>C25:0</td>
<td>1.69 ± 0.11</td>
</tr>
<tr>
<td>C26:0</td>
<td>1.94 ± 0.12</td>
</tr>
</tbody>
</table>

Fig. 1. Binding isotherms of recombinant AtACBP1 titrated with C18:1-, C18:2-, C18:3-, C24:0-, C25:0-, and C26:0-CoA esters at 30 °C using isothermal titration calorimetry. Top panels show raw data of 300 μl of 10 μM recombinant AtACBP1 titrated with 20 injections of 1.5 μl of 250 μM acyl-CoA ester solution. Bottom panels show the integrated area of each injection and the plotted graph. Parameters of the dissociation constant ($K_D$) are given in Table 1.
utoronto.ca/efp/cgi-bin/efpWeb.cgi) revealed that *AtACBP1* was highly expressed in the top and bottom of stems, in comparison with roots and rosette and cauline leaves, and that this expression localized in the epidermal peels (Fig. 2A).

The expression of *AtACBP1* in leaves and stems was investigated in transgenic *Arabidopsis* lines expressing *AtACBP1pro::GUS* (Fig. 2B–I). *AtACBP1pro::GUS* was expressed in the leaf vasculature (Fig. 2B–D) and on the stem surface (Fig. 2E) including the trichomes (Fig. 2F). The cross- and longitudinal-sections of the stem showed *AtACBP1pro::GUS* expression in the epidermis, the cortex, and the vascular bundles (Fig. 2G–I). Stem and leaf sections from control *Arabidopsis* transformed with pBI101.3 were not stained.

The *acbp1* mutant shows defects in epicuticular wax crystallization and cuticle membrane structure

When SEM was used to investigate wax crystallization patterns on the stem surface of the *acbp1* mutant, the occurrence of epicuticular wax crystals was significantly reduced (Fig. 3A) in comparison with the wild type (Fig. 3B). Upright rod-, tube-, and umbrella-shaped wax crystals were arrayed in an orderly manner on the wild type, but not the *acbp1* mutant stem. In contrast, the mutant had fewer crystals (Fig. 3).

When TEM was used to examine the fine structural changes of the stem cuticle, the cuticle membrane was intact in Col-0 (Fig. 4A) but not in the *acbp1* mutant (Fig. 4B). Instead, in the mutant, a ruptured and discontinuous cuticle membrane was observed (Fig. 4B). Absence of expression of *AtACBP1* culminated in an aberrant cuticle membrane, suggesting that *AtACBP1* is involved in stem cuticle formation.

The *acbp1* mutant shows reduction in stem cuticular wax constituents and cutin monomers and down-regulation of cuticular biosynthetic genes

To evaluate further the roles of *AtACBP1* in cuticle formation, GC-FID and GC-MS were employed to determine the amount and composition of cuticular waxes from stems of the wild type (Col-0), the *acbp1* mutant, and the *acbp1*-complemented (*acbp1-COM*; Xiao et al., 2008a) lines.

In stems, a 16% reduction of total wax was observed in *acbp1* in comparison with the wild type (Fig. 5A). In particular, the levels of C29 alkane, C28 and C30 primary alcohols, and C29 secondary alcohol and ketone were significantly reduced in the *acbp1* mutant (Fig. 5A). Their percentage reductions were 16% (C29 alkane), 15% (C28 primary alcohol), 21% (C30 primary alcohol), 21% (C29 secondary alcohol), and 16% (C29...
The normal amounts of total wax and various wax species were recovered in stems of *acbp1-COM* (Fig. 5A), confirming that the decrease in stem cuticular wax in the *acbp1* mutant resulted from knockout of *AtACBP1* expression. Subsequently, when the expression of wax biosynthetic genes was determined by qRT-PCR analysis (Fig. 6A), *CER8*, *KCRI*, *ECR*, and *CUT1/KCS6* significantly decreased in the *acbp1* stem in comparison with the wild type (Fig. 6A), and these decreases in gene expression were recovered in the *acbp1-COM* plants (Supplementary Fig. S1A at *JXB* online).

GC-MS was also used to determine cutin monomer composition and amount in the stems of the *acbp1* mutant, *acbp1-COM*, and wild type (Fig. 5B). The amounts of total cutin monomer were altered in stems of the *acbp1* mutant (Fig. 5B) in comparison with the wild type. Levels of C18:1 and C18:2 ω-hydroxyl fatty acids, as well as C18:1 and C18:2 dicarboxylic fatty acids, were significantly reduced in stems of the *acbp1* mutant (Fig. 5B). Their percentage reductions were 45% (C18:1 ω-hydroxyl fatty acid), 38% (C18:2 ω-hydroxyl fatty acid), 24% (C18:1 dicarboxylic fatty acid), and 31% (C18:2 dicarboxylic fatty acid). The chemical change in stems of the *acbp1* mutant was recovered in the *acbp1-COM* line (Fig. 5B), confirming that reduction in the amounts of stem cutin monomer in the mutant resulted from knockout of *AtACBP1* expression. Subsequently, on qRT-PCR, expression of some genes involved in cutin synthesis (*LACS2*, *CYP86A2*, *CYP86A4*, and *GPAT8*) (Supplementary Fig. S1C) showed a significant decrease in the stems of *acbp1* in comparison with wild-type *Arabidopsis* (Fig. 6B), which could be recovered in *acbp1-COM* plants (Supplementary Fig. S1B at *JXB* online). However, the expression of triacylglycerol biosynthetic genes (*DGAT1*, *DGAT2*, *DGAT3*, and *PADT1*) (Supplementary Fig. S1D) and cold-related genes (*COR47*, *LTI78*, *CBF1*, and *ICE1*) (Supplementary Fig. S1D), which are not implicated in cuticle formation, was not affected in the *acbp1* mutant and *acbp1-COM* plants in comparison with the wild type.

Seedlings of *acbp1* are more susceptible to *Botrytis cinerea* infection

To examine whether a reduction of wax and cutin in the *acbp1* mutant confers an altered response to the necrotrophic fungal pathogen *B. cinerea*, 3-week-old seedlings of the *acbp1* mutant and wild type were inoculated with *Botrytis* spores. As shown in Fig. 7, the *acbp1* mutant seedlings displayed enhanced susceptibility after spraying with *Botrytis* suspension. At 6 DAI, chlorosis and necrosis were observed in the *acbp1* mutant, but not in the wild type (Fig. 7). Measurement of leaf wax and cutin from the *acbp1* mutant in comparison with the wild type revealed a significant decrease in wax but not in cutin (Supplementary Fig. S2 at *JXB* online). In particular, C31 and C33 alkanes, C28 fatty acids, and total wax load significantly declined (Supplementary Fig. S2B). This is not surprising because *AtACBP1pro::GUS* was expressed more in stem epidermis than in leaf epidermis (Fig. 2). These results suggest that a reduction in wax content in the *acbp1* mutant could have caused greater susceptibility to *Botrytis* infection.

Discussion

In plants, fatty acids are synthesized in the plastids by the addition of two-carbon units to a growing acyl chain facilitated by the acyl carrier protein (ACP) during *de novo* fatty
acid synthesis (Ohlrogge and Browse, 1995). Subsequently, 16:0-ACP and 18:0-ACP are exported to the ER for the biosynthesis of other lipids including cutin, suberin, and cuticular waxes (Post-Beittenmiller, 1996; Jenks et al., 2002; Nawrath, 2002). Cutin and wax are synthesized exclusively in the epidermis (Nawrath, 2002; Suh et al., 2005; Samuels et al., 2008). AtACBP1pro:GUS is expressed in the embryos, lateral root primordia, vascular bundles, stigmas, and ovaries (Du et al., 2013b). In this study, transgenic Arabidopsis expressing AtACBP1pro:GUS showed strong GUS expression in stem epidermis, in agreement with the corresponding expression analysis of KCS20 and KCS2/DAISY genes involved in VLCFA elongation (Lee et al., 2009b). It is noteworthy that the GUS stain was not detected in leaf epidermis (Fig. 2D), suggesting a putative function for AtACBP1 in stem cuticle formation.

In wax biosynthesis, C18 fatty acyl-CoAs are the predominant precursors for production of VLCFAs in four sequential reactions catalysed by a membrane-bound multiple enzyme system consisting of KCS, β-ketoacyl-CoA reductase (KCR), β-hydroxyacyl-CoA dehydratase (HCD), and enoyl-CoA reductase (ECR) (Kunst and Samuels, 2009). After several cycles of condensation of malonyl-CoA with long-chain acyl-CoAs, reduction to β-hydroxyacyl-CoA, dehydration to an enoyl-CoA, and reduction of the enoyl-CoA, VLCFAs with different acyl chains ranging from C20 to C34 are generated and subsequently converted to various wax components through decarbonylation and acyl reduction (Kunst and Samuels, 2003; Samuels et al., 2008). Through multiple steps of hydroxylation and epoxidation, 16:0 and 18:X fatty acyl-CoAs are converted into cutin monomers (Schnurr et al., 2004). ITC analysis from the present study revealed that rACBP1 binds not only long-chain acyl-CoA esters (C18:1-, C18:2-, and C18:3-CoA) but also saturated VLC acyl-CoA esters (C24:0-, C25:0-, and C26:0-CoA). The reduced binding affinity of rACBP1 for VLC acyl-CoA esters may be attributed to either the longer acyl chain length or unsaturation of the acyl chain. In measurements of rACBP1 interaction with VLC acyl-CoA esters, only C24:0 to C26:0 were tested because acyl-CoAs with acyl chains longer than C27:0 are not commercially available (http://www.avantilipids.com/). The present analysis demonstrated that AtACBP1 is able to bind C18 fatty acyl-CoAs and VLC acyl-CoAs, and can potentially transport these precursors for cutin and wax biosynthesis during stem cuticle formation (Fig. 8). A reduction in wax crystal density and observations on the irregularity of the cuticle membrane on the stems of the acbp1 mutant suggest that a defective cuticle had resulted from functional loss of AtACBP1. Interestingly, the T-DNA insertional mutants of AtACBP3 also showed a highly irregular outermost cell wall surface (Xia et al., 2012). These phenotypes are not only evident in AtACBP mutants but have also been observed in other mutants in cuticle development. Wax crystals were also absent on the stem surfaces of the cer1, cer2, and cer6 mutants (Millar et al., 1999). In addition, the lacs2 mutant showed a reduction in plant size, seed set, and seedling establishment (Schnurr et al., 2004). Furthermore, in the wax2, fdh, and lcr mutants, severe organ
Fusions occurred and pollen fertility was affected (Lolle et al., 1998; Wellesen et al., 2001; Chen et al., 2003). These drastic phenotypic changes accompanied by severe reduction in wax or cutin load were caused by mutation in the genes of the wax and cutin pathways (Millar et al., 1999; Wellesen et al., 2001; Chen et al., 2003; Lü et al., 2009).

The wax species that were significantly lower in the acbp1 mutant stem included not only alkanes, but also primary alcohols, C29 secondary alcohol, and ketones (Fig. 5A). This is in agreement with the down-regulation in the acbp1 mutant stem of wax biosynthetic genes CER8, KCR1, ECR, CUT1/KCS6, LACS2, CYP86A2, CYP86A4, and GPAT8 decreased in stems of acbp1 in comparison with the wild type (Col-0). Asterisks denote significant differences from the wild type (*P<0.05; **P<0.01). Values are means ±SE (n=3).

The broad range of wax species altered in the acbp1 mutant stem implies that the function of AtACBP1 in wax biosynthesis lies upstream in the pathway, involving VLCFA elongation (1). AtACBP1 may participate in general stem wax biosynthesis and could affect both the decarboxylation and acyl reduction pathways (2, 3). Changes in stem cutin monomer content in the acbp1 mutant suggest that AtACBP1 participates in stem cutin synthesis (4), ACP, acyl carrier protein; LC-CoA, long-chain acyl-CoA; VLC-CoA, very-long-chain acyl-CoA; KCS, β-ketoacyl-CoA synthase; KCR, β-ketoacyl-CoA reductase; HCD, β-hydroxyacyl-CoA dehydratase; ECR, enoyl-CoA reductase; VLCFA, very-long-chain fatty acid; ER, endoplasmic reticulum; PM, plasma membrane; CW, cell wall.

Fig. 6. qRT-PCR analysis of wax (A) and cutin (B) biosynthetic genes in stems of the acbp1 mutant and Col-0. Expression of CER8, KCR1, ECR, CUT1/KCS6, LACS2, CYP86A2, CYP86A4, and GPAT8 decreased in stems of acbp1 in comparison with the wild type (Col-0). Asterisks denote significant differences from the wild type (*P<0.05; **P<0.01). Values are means ±SE (n=3).

Fig. 7. Response of acbp1 to Botrytis cinerea infection. Three-week-old wild-type (Col-0) and acbp1 plants were sprayed with B. cinerea (2 × 10^5 spores ml^-1). Photographs were taken at 0 and 6 days after inoculation (DAI). The experiments were repeated twice with consistent results. Arrows indicate chlorosis and necrosis of leaves.

Fig. 8. Proposed function of AtACBP1 in the wax and cutin synthesis pathway. The broad range of wax species altered in the acbp1 mutant stem implies that the function of AtACBP1 in wax biosynthesis lies upstream in the pathway, involving VLCFA elongation (1). AtACBP1 may participate in general stem wax biosynthesis and could affect both the decarboxylation and acyl reduction pathways (2, 3). Changes in stem cutin monomer content in the acbp1 mutant suggest that AtACBP1 participates in stem cutin synthesis (4). ACP, acyl carrier protein; LC-CoA, long-chain acyl-CoA; VLC-CoA, very-long-chain acyl-CoA; KCS, β-ketoacyl-CoA synthase; KCR, β-ketoacyl-CoA reductase; HCD, β-hydroxyacyl-CoA dehydratase; ECR, enoyl-CoA reductase; VLCFA, very-long-chain fatty acid; ER, endoplasmic reticulum; PM, plasma membrane; CW, cell wall.
is a fatty acid ω-hydroxylase in the synthesis of hydroxy fatty acids (Xiao et al., 2004), while CYP86A4 and GPAT8 catalyse ω-hydroxylation and esterification to glycerol, respectively, during cutin synthesis (Li et al., 2007; Li-Beisson et al., 2009). Knockout of AtACBP1 probably adversely affected the accumulation of long-chain and VLC acyl-CoA essential for stem wax and cutin biosyntheses (Fig. 8). Furthermore, the lack of substrates for wax and cutin biosyntheses led to a decrease in the expression of both stem wax and cutin biosynthesis genes, which will reduce wax or cutin production.

Waxes are known to be synthesized in the epidermis (Samuels et al., 2008), and the leaf wax content was lower in the acbp1 mutant in comparison with the wild type (Supplementary Fig. S2A at JXB online). However, AtACBP1pro::GUS was not observed to be expressed in the leaf epidermis (Fig. 2D), suggesting that AtACBP1 may not participate directly in leaf wax biosynthesis. Possibly, leaf wax changes may have been affected by the dramatic alterations observed in stem cuticular contents. Although the AtACBP1pro::GUS-transformed plants did not express detectable GUS activity in the leaf epidermal cells (Fig. 2C, D), reductions in several compounds (i.e. C31 and C33 alkanes, C28 fatty acid, and total wax load) in leaf wax but not leaf cutin (Supplementary Fig. S2) in the acbp1 mutant may be attributed to a systematic change in the expression of wax-related genes and/or the activities of their gene products as a result of the defective stem cuticle. As the acbp1 mutant exhibited lesions in stem cuticle formation, this could have potentially affected the status of the plant as a whole (e.g. water loss, susceptibility to pathogens, etc.), and could have culminated in an indirect effect on wax synthesis in the leaf epidermis. It is well documented that cuticular wax biosynthesis is sensitive to diverse environmental cues, and several transcription factors have been identified to play a role in its biosynthesis and accumulation (Aharoni et al., 2004; Zhang et al., 2007; Seo et al., 2011; Cominelli et al., 2008; Lü et al., 2009). Xia et al. (2012) showed that in the leaves of both acbp3 and acbp4 mutants, the cutin monomers were greatly reduced, with pronounced reduction in C16:0, C18:1, and C18:2 dicarboxylic fatty acids, but no change in most cutin monomers was evident in the acbp6 mutant. In comparison, the present analysis revealed that stem cutin monomer levels also declined in the acbp1 mutant, confirming its role in cutin biosynthesis (Fig. 8). In particular, C18 species (C18:1 and C18:2 ω-hydroxy fatty acids and dicarboxylic fatty acids) of cutin were more affected in the acbp1 mutant. This corresponds well to ITC data that showed that rACBP1 binds long-chain acyl-CoA esters (C18:1-, C18:2-, and C18:3-CoAs) with a greater affinity (i.e. smaller dissociation constant, K_d) than VLC acyl-CoAs (C24:0-, C25:0-, and C26:0-CoA). AtACBP1 is localized in the PM and the ER, but AtACBP3 is targeted to the extracellular space, while AtACBP4 is a cytosolic protein. Although they show differential subcellular localization, they all affect cutin biosynthesis, suggesting that the binding and trafficking of precursors in cutin synthesis transverse across subcellular compartments.

The acbp1 mutant in TEM showed an aberrant cuticle membrane in stems and was more susceptible to infection caused by B. cinerea possibly by entry through the aberrant cuticle, suggesting that alteration of cuticle constituents in this mutant impaired its basal defence responses. These results are consistent with the reduction in fungal pathogen resistance observed in the acbp3, acbp4, and acbp6 mutants which were also cuticle-defective (Xia et al., 2012). Previous findings have also revealed that AtACBP3 overexpression constitutively activated salicylic acid accumulation, PR gene expression and cell death, and increased resistance to the virulent bacterial pathogen P. syringae DC3000 (Xiao and Chye, 2011b).

Leaf susceptibility of the acbp1 mutant to B. cinerea infection arising from a decline in leaf wax (but not leaf cutin) suggested that this decrease affected the leaf cuticle membrane through extrapolating the observations of altered cuticle in stems including significant decreases in both stem wax and cutin loads (Fig. 4). Lee et al. (2009a) reported that the ltp1 mutant showed a reduction in the C29 alkane in stems but not leaves, and they did not see any significant changes in total wax in neither stem nor leaf. Although ltp1 mutant leaves showed increases in three cutin constituents, they were more susceptible to Alternaria brassicicola (Lee et al., 2009a). These findings support that changes in wax and cutin loads in stem and leaf may affect the cuticle barrier which is known to protect the plant against pathogen infection (Jenks et al., 1994). Perhaps changes in cuticular content may also have altered the leaf surface structure which then enhanced susceptibility of the acbp1 mutant to B. cinerea. Li et al. (2007) have reported that a change in the thickness or the structure of the pavement cells and the guard cells in the gpat4gpat8 double mutant made it more susceptible to A. brassicicola. Similarly, in the ltp1 mutant which was more susceptible to A. brassicicola, Lee et al. (2009a) observed alterations in the structure of the cuticular layer, a protrusive cytoplasm, and disorganized grana and stroma lamellae in the chloroplasts.

In summary, using phenotypic and biochemical analyses of the acbp1 mutant, it is demonstrated that AtACBP1 is involved in stem cuticle formation. Previous studies have suggested that plasma membrane-localized glycosylphosphatidylinositol-anchored lipid transfer proteins function in cuticular lipid transport (DeBono et al., 2009; Lee et al., 2009a; Kim et al., 2012). It is illustrated herein that ER- and PM-associated AtACBP1 also participates in stem wax and cutin biosynthesis, probably as a carrier protein, as supported by ITC data (Fig. 1; Table 1).

**Supplementary data**

Supplementary data are available at JXB online.

**Figure S1.** Expression analysis of wax and cutin biosynthetic genes, and genes with no implication on cuticle formation (triacylglycerol biosynthetic genes and cold-related genes) in stems of Col-0, the acbp1 mutant, and the acbp1-COM line.

**Figure S2.** Cuticular wax and cutin monomer composition and amount in leaves of Col-0 and the acbp1 mutant.

**Table S1.** Sequences of gene-specific primers for qRT-PCR.

**Table S2.** Mass-to-charge ratios (m/z) of cutin compounds used in mass spectrometry.
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