Cuticular Wax Biosynthesis is Up-Regulated by the MYB94 Transcription Factor in Arabidopsis

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The aerial parts of all land plants are covered with hydrophobic cuticular wax layers that act as the first barrier against the environment. The MYB94 transcription factor gene is expressed in abundance in aerial organs and shows a higher expression in the stem epidermis than within the stem. When seedlings were subjected to various treatments, the expression of the MYB94 transcription factor gene was observed to increase approximately 9-fold under drought, 8-fold for ABA treatment and 4-fold for separate NaCl and mannitol treatments. MYB94 harbors the transcriptional activation domain at its C-terminus, and fluorescent signals from MYB94:enhanced yellow fluorescent protein (eYFP) were observed to increase approximately 2-fold in the leaves of the transgenic Arabidopsis roots. The total wax loads increased from MYB94:enhanced yellow fluorescent protein (eYFP) overexpression (MYB94-OX) lines, as compared with those of the wild type (WT). MYB94 activates the expression of WSD1, KCS2/DAISY, CER2, FAR3 and ECR genes by binding directly to their gene promoters. An increase in the accumulation of cuticular wax was observed to reduce the rate of cuticular transpiration in the leaves of MYB94-OX lines, under drought stress conditions. Taken together, a R2R3-type MYB94 transcription factor activates Arabidopsis cuticular wax biosynthesis and might be important in plant response to environmental stress, including drought.

Keywords: Arabidopsis – Cuticular wax – MYB94 – R2R3-type MYB transcription factor – Transcriptional regulation.

Introduction

The aerial part of plant surfaces is covered with a hydrophobic layer comprised of cutin polyesters and intra- and epicuticular waxes (reviewed by Bernard and Joube’s 2013, Yeats and Rose 2013). The cuticle prevents non-stomatal water loss and gaseous exchange, scatters UV radiation, and repulses pathogen spores and atmospheric pollutants, such as acid rain, ozone and carbon dioxide (reviewed by Shephered and Griffiths 2006, Bernard and Joube’s 2013). Cuticular waxes consist of solventsoluble lipids, which are made of very-long-chain fatty acids (VLCFAs; >20 carbons in length) and their derivatives (including alkanes, aldehydes, ketones, primary and secondary alcohols and wax esters) with smaller amounts of triterpenoids and phenylpropanoids (Jetter et al. 2006). Insoluble cutin layers are known to have a polyester structure that contains glycerols esterified with C16 and C18 fatty acids and their derivatives, such as hydroxy and dicarboxylic fatty acids, and phenylpropanoids, such as coumaric and ferulic acids (Kunst and Samuels 2009, Beisson et al. 2012).

Cuticular waxes are known to be synthesized in epidermal cells (Suh et al. 2005) where C16 and C18 fatty acids, which are synthesized de novo in the plastids, are further elongated into VLCFAs by an endoplasmic reticulum (ER)-localized fatty acid elongase complex (FAE). FAE consists of β-ketoacyl-CoA synthase (KCS), β-ketoacyl-CoA reductase (KCR), β-hydroxycarboxyl-CoA dehydratase (HCD) and enoyl-CoA reductase (ECR) (Kunst and Samuels 2009, Lee and Suh 2013, Li-Beisson et al. 2013). Recently CER2 and CER2-LIKE1/CER26, which have homology to BAHD acyltransferase, were reported to function in the synthesis of VLCFAs of >26 carbons in length (Haslam et al. 2012, Pascal et al. 2013). The synthesized VLCFAs are activated into VLCFA-CoAs by a long chain acyl-CoA synthetase (LACS) (Lu et al. 2009, Weng et al. 2010), and the VLCFA-CoAs are modified via the alkane-forming and alcohol-forming pathways (Kunst and Samuels 2009, Lee and Suh 2013, Li-Beisson et al. 2013). In the alkane-forming pathway, VLCFA-CoAs are catalyzed into alkanes by a CER1, CER3 and Cyt b6 complex (Bernard et al. 2012), and the alkanes are oxidized into secondary...
alcohols and ketones by a midchain alkane hydroxylase1 (MAH1) (Greer et al. 2007). In the alcohol-forming pathway, VLCFA-CoAs are reduced into primary alcohols by a fatty acyl-CoA reductase (FAR3/CER4) (Rowland et al. 2006), and the primary alcohols are esterified with C16-CoAs by a bifunctional wax synthase/acyl-CoAAdiaclyglycerol acyltransferase (WSD1) (Li et al. 2008). The wax precursors synthesized in the ER are exported to the extracellular space via both homodimers (ABCG11/WBC11) and heterodimers (ABCG11/WBC11 and ABCG12/CR5) of ATP-binding cassette (ABC) transporters (McFarlane et al. 2010). Glycosylphosphatidylinositol (GPI)-anchored LTPs (LTPGs), which are attached to the outer surface of the plasma membrane via a GPI anchor, were reported to be directly or indirectly involved in wax export (DeBono et al. 2009, Lee et al. 2009a, Kim et al. 2012). Wax transport from the ER to the plasma membrane was recently reported to require Golgi- and trans-Golgi network-mediated vesicle trafficking (McFarlane et al. 2014).

To date, there are pieces of several evidence indicating that cuticular wax biosynthesis is regulated at the transcriptional level (Bernard and Joube` s 2013, Li-Beisson et al. 2013). Arabidopsis WAX INDUCTER1/SHINE1 (WIN1/SHN1) encoding an AP2/EREBP-type transcription factor (TF) activates cuticular wax and cutin biosynthesis by inducing the expression of CER1, KCS1, CER2, LACS2, GATA4, CYP86A4, CYP86A7 and HTH-like genes, and the total amount of cuticular wax and cutin increases in transgenic Arabidopsis that overexpresses WIN1 (Broun et al. 2004, Kannangara et al. 2007). It was reported later that WIN1/SHN1 regulates cutin biosynthesis via direct binding to the promoter of the LACS2 gene (Kannangara et al. 2007). Oshima et al. (2013) have characterized the R2R3-type MYB16 and MYB106 TFs involved in cuticle development, and, in particular, MYB106 was identified to be a positive regulator of WIN1/SHN1 expression. In addition, MYB30, MYB41, CURLY FLAG LEAF1 (CFL1; containing WW domain protein) and HOMEODOMAIN GLABROUS1 (HDG1; classified in the Class IV HD-ZIP gene family) have been reported to function as positive regulators in cuticular wax biosynthesis or in cuticle deposition (Cominelli et al. 2008, Raffaele et al. 2008, Wu et al. 2011). In particular, MYB41 activates the synthesis and deposition of both aliphatic suberin-type polyesters and suberin-associated MYB transcription factor

**Results**

**Isolation of the MYB94 gene encoding the R2R3-type MYB transcription factor**

In order to characterize the novel TF that is involved in the activation of cuticular wax biosynthesis under drought stress, 147 TFs that showed higher expression in stem epidermal cells than in stem cells were isolated based on an Arabidopsis stem epidermis microarray analysis (Suh et al. 2005). They were further screened according to the induced expression patterns after treatments with ABA and/or drought stress using the Arabidopsis microarray database (www.arabidopsis.org). Sixteen MYB TFs, including a MYB96 TF, were selected (Supplementary Fig. S1A). Finally we isolated the MYB94 (At3g47600) gene that is up-regulated approximately 5-fold in stem epidermal cells relative to cells in the stems, and this was induced by approximately 10- and 2-fold in Arabidopsis seedlings after 10 μM ABA treatment and drought stress, respectively. In addition, MYB96 was observed to be very closely located in the phylogenetic tree of 126 R2R3-type MYB TFs (Supplementary Fig. S1B).

**MYB94 contains a transcriptional activation domain in its C-terminal region and the fluorescent MYB94:eYFP is localized to the nucleus.**

A MYB94 protein harboring 333 amino acid residues contains R2 (amino acid residues 9–63) and R3 (amino acid residues 69–117) domains with the conserved amino acid motifs R2 [Xm-]-W-(Xm-)-W-(Xm-)-W and R3 [Xm-]-W-(Xm-)-W that are known to be essential for DNA binding (Fig. 1A; Dubos et al. 2010). The transcriptional activation domain of the MYB94 protein was subsequently investigated via yeast-based transactivation assay (Park et al. 2009). The full-length (F; amino acid residues 1–333), N-terminal region (N; amino acid residues 1–123) and C-terminal region (C; amino acid residues 124–333)
of MYB94 were translationally fused to the GAL4 DNA-binding domain in the pGBKT7 plasmids (Fig. 1B; Clontech). When the constructed plasmids were induced in the yeast Y190 strain containing the GAL1 promoter and HIS3 and lacZ reporter genes, β-galactosidase activity was observed in yeast cells harboring MYB94-F or MYB94-C plasmids, but not in yeast cells containing pGBKT7 (V) or MYB94-N plasmids (Fig. 1C). The results indicate that the C-terminal region of MYB94 functions as a transcriptional activation domain.

To investigate the intracellular location of MYB94 TF, a full-length (999 bp) MYB94 without the termination codon (TAG) was translationally fused into the N-terminus of eYFP. The generated construct was introduced into tobacco leaf epidermal cells via Agrobacterium-mediated infiltration (Sparkes et al. 2006) or

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**Fig. 1** MYB94 is a transcriptional activator localized in the nucleus. (A) Amino acid sequences of the MYB94 protein. The two green boxes denote the conserved R2 and R3 domains required for DNA binding. The highly conserved amino acid residues are shown in red, and the nuclear localization signal motif is underlined. (B) Schematic diagram for the full-length protein and deletion constructs of MYB94 in pGBKT7 vector. F, the full-length MYB94 protein in the pGBKT7 vector; N, the N-terminal region of MYB94 protein in the pGBKT7 vector; C, the C-terminal region of MYB94 protein in the pGBKT7 vector. (C) Transactivation assay of MYB94 in yeast cells. The transformed yeast cells were grown on selective medium SD/–Trp (left) and SD/–Trp–His (middle) supplemented with 25 mM 3-amino-1,2,4-aminotriazole. An X-gal lift assay was carried out using an SD/–Trp–His plate (right). V, yeast transformed with the pGBKT7 vector; F, N and C, yeast transformed with the F, N and C constructs, respectively, as shown in (B). (D) Subcellular localization of the MYB94:eYFP in tobacco epidermis (upper) and a transgenic Arabidopsis root (lower). DAPI staining was performed to visualize the nucleus, and the images were acquired using a laser confocal scanning microscopy with a YFP or a UV filter. Scale bars = 10 μm.
was transformed into Arabidopsis via floral dip (Zhang et al. 2006). The fluorescent signals from the MYB94:eYFP construct were observed in the nucleus of tobacco leaf epidermal cells and in 2-week-old transgenic Arabidopsis roots, and these merged with the fluorescent signals from 4′,6-diamidino-2-phenylindole (DAPI) staining (Fig. 1D), indicating that the fluorescent MYB94:eYFP was localized in the nucleus.

Spatial and temporal expression patterns of MYB94 in various Arabidopsis organs and in seedlings after treatments with ABA, drought, salt and osmotic stress

To investigate the expression levels of the MYB94 gene in various Arabidopsis organs and in seedlings after treatment with ABA and abiotic stresses, quantitative real-time reverse transcription–PCR (qRT–PCR) was performed. The MYB94 transcripts were ubiquitously expressed in germinating seeds, rosette and cauline leaves, flower buds, open flowers, stems and developing siliques, but they were not or were very weakly observed in pollen and roots. In addition, the expression of the MYB94 gene was approximately five times higher in stem epidermal peels than in the stems (Fig. 2A). The expression of the MYB94 gene was also analyzed in 10-day-old Arabidopsis seedlings after treatment with drought stress, 100 μM ABA, 200 mM NaCl and 200 mM mannitol. When the expression of the rd29A gene was induced as a control of water-deficit response (Liu et al. 1998), the levels of MYB94 transcripts increased by approximately 9-fold in response to drought, 8-fold in response to ABA and 4-fold for NaCl and mannitol treatments, separately (Fig. 2B).

β-Glucuronidase assay (GUS) was performed in transgenic Arabidopsis plants expressing the GUS gene under the control of the MYB94 promoter (approximately 4.2 kb) to examine the spatial and temporal expression of the MYB94 gene. More than 30 independent transgenic lines were screened for GUS activity, and the expression of the GUS gene was observed in the epidermal part of a 7-day-old seedling, stem epidermis, stem cortex, phloem and xylem cells of stems, leaves including trichomes, sepals, anther and anther filaments, the upper portion of the stigma and the silique walls, but not in the root and the developing seeds (Fig. 2Ca–g).

The amounts of cuticular waxes increased in the leaves of transgenic Arabidopsis plants overexpressing MYB94

In order to investigate the function of the MYB94 gene in planta, the Arabidopsis transgenic plants overexpressing MYB94 (MYB94 OX) were generated under the Cauliflower mosaic virus (CaMV) 35S promoter control (Fig. 3A). The MYB94 transcript levels in the leaves of MYB94 OX lines were elevated by approximately 15–500 times that of the WT (Fig. 3B). Although no significantly altered phenotype in the MYB94 OX lines was observed during growth and development under long-day conditions (16 h/8 h; light/dark), as compared with the WT, scanning electron microscopy analysis showed that the deposition of epicuticular wax crystals was detected on the leaves of MYB94 OX lines, but was not observed on the leaves of the WT (Fig. 3Ca–d).

In addition, transgenic Arabidopsis plants expressing the MYB94 gene under the control of the β-estradiol-inducible promoter were generated and, after treatment with β-estradiol, epicuticular wax crystals formed on the leaves of transgenic Arabidopsis expressing MYB94 but not on the leaves of WT or mock-treated leaves of transgenic Arabidopsis expressing MYB94 (Fig. 3D).

The amount and the composition of cuticular waxes were subsequently measured from leaves and stems of MYB94 OX lines and from WT plants using gas chromatography (GC). The total wax loads increased by approximately 1.7- and 1.3-fold in the leaves of the MYB94 OX-1 line and the OX-5 line, respectively, but was not significantly altered in the leaves of MYB94 OX-3 relative to the WT (Fig. 4A), suggesting that an increase in the total wax loads is correlated with the up-regulated expression of the MYB94 gene. In addition, the levels of C28 and C30 VLCFAs, C28, C30 and C32 aldehydes, and C29, C31, C33 alkanes synthesized by the alkane-forming pathway and the amounts of C26, C28, C30 and C32 primary alcohols produced by the alcohol-forming pathway were significantly increased in MYB94 OX-1 and in OX-5 lines compared with the WT (Fig. 4B). However, no significant increase in the total wax loads was observed in the stems of MYB94 OX-1 and OX-5 lines compared with those of the WT (Supplementary Fig. S2).

To examine if overexpression of MYB94 affects cutin biosynthesis, the amounts and composition of cutin monomers were measured in the leaves of MYB94 OX-1 and WT plants using GC–mass spectrometry (GC/MS). The total amounts of cutin monomers increased by approximately 18% in MYB94 OX-1 relative to the WT (Supplementary Fig. S3A). In particular, the amounts of hydroxy fatty acids, including C18:1 and C18:2 α-hydroxy fatty acids, and dicarboxylic fatty acids, including C18:2 dicarboxylic fatty acids, increased by 39% and 24% in MYB94 OX-1 relative to the WT (Supplementary Fig. S3B).

The expression of genes involved in cuticular wax biosynthesis was up-regulated in leaves of transgenic Arabidopsis plants overexpressing MYB94

To investigate the target genes of the MYB94 TF, total RNAs were extracted from the leaves of the WT and the MYB94 OX-1 lines under the control of the CaMV 35S promoter and were subjected to a microarray analysis using the Arabidopsis Affymetrix ATH1 GeneChip, which contains >22,500 probe sets. About 230 genes were up-regulated by approximately 1.5-fold in MYB94 OX-1 relative to the WT. The list of genes that were up-regulated by >1.5-fold in MYB94 OX-1 relative to the WT, and that are involved in cuticular wax biosynthesis, is shown in Fig. 5A. Interestingly, the expression of genes involved in cutin synthesis was not significantly up-regulated in MYB94 OX-1 as compared with the WT (Supplementary Table S1). qRT-PCR analysis showed that the MYB94, WSD1, KCS2/DAISY, CER1, CER2, FAR3 and ECR transcript levels were up-regulated by approximately 2.5- to 30-fold, but no significant changes in
the levels of KCS1 transcripts were observed in MYB94 OX-1 relative to the WT (Fig. 5B).

In addition, significant up-regulation of MYB94, WSD1, KCS2/DAISY, CER1, CER2, FAR3 and ECR genes was observed in the leaves of transgenic Arabidopsis expressing MYB94 under the control of the β-estradiol promoter, but not in the WT, at 2, 6, 12 and 24 h after treatment with 10 μM β-estradiol. However, the expression of those genes was not significantly altered in either mock-treated WT or transgenic leaves (Fig. 5C). These results indicate that the MYB94 TF activates cuticular wax biosynthesis via up-regulation of genes involved in wax biosynthesis.
Fig. 3 Constitutive or \(\beta\)-estradiol-inducible expression of \(MYB94\) in transgenic Arabidopsis plants. (A) Schematic diagram of \(MYB94\) in the pPZP212 vector under the control of the CaMV 35S promoter. 35S-P, CaMV 35S promoter; 35S-T, CaMV 35S terminator; rbs-T, terminator of ribulose 1,5-biphosphate carboxylase/oxygenase small subunit; RB, T-DNA right border sequences; LB, T-DNA left border sequences. (B) The relative expression level of \(MYB94\) genes in the leaves of WT and \(MYB94\) overexpression (\(MYB94\) OX) lines. The total RNA was isolated from the leaves of WT and \(MYB94\) OX lines and was subjected to qRT–PCR. The \(PP2A\) (At1g13320) gene was used to determine the quantity and quality of the cDNAs. Each value is the mean of triplicate experiments. (C) Scanning electron microscopy (SEM) images of epicuticular wax crystals on the leaves. The rosette leaves of 3-week-old WT (a, b) and \(MYB94\) OX (c, d) plants grown in soil were used to perform a SEM analysis. (b, d) White boxes in (a) and (c) are shown at higher magnifications. Scale bars = 2.5 \(\mu\)m (a, c), 1.5 \(\mu\)m (b, d). (D) SEM analysis of epicuticular wax crystals on the leaves of transgenic Arabidopsis after induction of \(MYB94\). Ten-day-old plants grown in soil were sprayed every 2 d with dimethylsulfoxide (DMSO; mock) or 10 \(\mu\)M \(\beta\)-estradiol solution. Deposition of epicuticular wax crystals was examined 3 weeks after induction. Scale bars = 2 \(\mu\)m.

Fig. 4 Cuticular wax composition and amount in the leaves of WT and \(MYB94\) OX lines. Cuticular waxes were extracted from 3- to 4-week-old leaves (A, B) of Arabidopsis plants. Each value is the mean of five independent measurements. Bars indicate the SE. \(\dagger\) and * denote statistical differences with respect to the WT using Student’s \(t\)-test (\(\dagger\) \(P < 0.05\), * \(P < 0.01\)).
MYB94 activates the expression of wax biosynthetic genes via direct binding of their promoters

To examine if the MYB94 TF regulates the expression of wax biosynthetic genes directly, transcriptional activation assays in tobacco leaf protoplasts and electrophoretic mobility shift assays (EMSAs) were performed. To make reporter constructs, each promoter region of *WSD1*, *KCS2/DAISY*, *CER1*, *CER2*, *FAR3* and *ECR* genes was isolated and transcriptionally fused to the upstream region of the luciferase gene. In addition, the effector constructs with or without a MYB94 gene under the control of the CaMV 35S promoter were generated (Fig. 6A). Then the reporter and effector constructs were co-transformed into tobacco protoplasts, and the luciferase activity was measured.

A vector construct harboring the *GUS* gene was included to monitor the transformation efficiencies, and the co-transformation of the p35-MYB94 effector construct and each reporter construct elevated the luciferase activity by approximately 3.5- to 96-fold (Fig. 6B). In contrast, co-transformation with the p35 effector construct and with each reporter construct did not significantly elevate the expression of the reporter gene, indicating that MYB94 functions as a transcriptional activator of the *WSD1*, *KCS2/DAISY*, *CER1*, *CER2*, *FAR3* and *ECR* genes.

We next isolated the MYB binding consensus sequences in the promoter regions of the *WSD1*, *KCS2/DAISY*, *CER1*, *CER2*, *FAR3* and *ECR* genes (as shown in *Supplementary Fig. S4A*; Urao et al. 1993, Abe et al. 2003). The oligonucleotides containing the MYB binding consensus sequences (BS) and their mutant oligonucleotides (mBS) were synthesized and end-labeled with [γ-32P]dATP. The radiolabeled oligonucleotides were incubated with recombinant MYB94:maltose-binding protein (MBP) purified from *Escherichia coli*, and then the reaction mixtures were electrophoresed on 6% native PAGE gels.

### Table 1

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**Fig. 5** Up-regulation of cuticular wax biosynthetic genes in MYB94 OX-1. (A) List of wax biosynthetic genes up-regulated in MYB94 OX. Genes up-regulated >1.5-fold in MYB94 OX compared with the WT are listed. The P-values were corrected for multiple testing using FDR (false discovery rate) methodology. AGI, Arabidopsis Genome Initiative number; FC, fold changes. (B, C) qRT–PCR analysis of wax biosynthetic genes. Total RNAs were extracted from the leaves of 3- to 4-week-old WT and MYB94 OX-1 (B) and from β-estradiol-treated or non-treated leaves of 2-week-old transgenic Arabidopsis (C). Two-week-old transgenic Arabidopsis plants were incubated in MS liquid medium supplemented with DMSO or 10 μM β-estradiol. The total RNA was isolated from whole plants, which were harvested at the indicated time points (hours) after application of DMSO or 10 μM β-estradiol, and were subjected to qRT–PCR analysis. The data from three independent experiments were averaged, and the bars indicate the SE (t-test, *P* < 0.05).
As shown in Fig. 6C, the MYB94:MBP was bound to the MYB binding consensus motifs of the WSD1, KCS2/DAISY, CER2, FAR3 and ECR gene promoters, but MBP alone did not bind to these. On the other hand, the MYB94:MBP binding was remarkably reduced in the presence of excess amounts of unlabeled BS fragments, and it was reduced to a lesser degree by the mutated BS fragments (mBS), indicating that MYB94 specifically binds to the BS sequences in the WSD1, KCS2/DAISY, CER2, FAR3 and ECR gene promoters. However, no specific binding of the MYB94:MBP to the BSs in the CER1 promoter was observed (Supplementary Fig. S4B). These observations revealed that MYB94 activates the expression of wax biosynthetic genes such as WSD1, KCS2/DAISY, CER2, FAR3, and ECR via direct binding to their promoters.

Cuticular transpiration occurred more slowly in the leaves of the MYB94 OX lines compared with those of the WT

To examine if enhanced accumulation of cuticular waxes in MYB94 OX lines affects water loss through the cuticle, cuticular transpiration assays were carried out. Four-week-old MYB94 OX and WT plants grown under either normal or drought stress conditions were dark acclimated for 12 h to ensure stomatal closure and were soaked in water for 1 h in the dark. The leaves were then used for water loss measurements. As shown in Fig. 7, no significant differences were observed in terms of the cuticular transpiration between MYB94 OX and WT leaves which were grown under normal conditions. However, cuticular transpiration occurred more slowly in drought-treated MYB94 OX leaves, but more rapidly in drought-treated WT leaves, indicating that the elevated accumulation of cuticular waxes contributes to a reduction in water loss through the cuticle under water-deficit conditions.

Discussion

The plant surface is covered with a cuticle that is comprised of cutin polyesters and cuticular wax layers. Since the cuticle is the
first barrier between the plants and their environment, the regulation of cuticle biosynthesis is important for optimal plant growth and for normal plant development. In this study, we identified a novel MYB94 TF that activates Arabidopsis cuticular wax biosynthesis, which is supported by the following evidence: (i) the expression levels of MYB94 are higher in stem epidermal peels than in stems and are increased significantly by ABA, drought, salt and osmotic stresses; (ii) the MYB94 TF harbors the conserved R2R3 DNA-binding motif at the N-terminus and a transcriptional activation domain at the C-terminus; (iii) MYB94ceYFP was localized in the nucleus in tobacco epidermis and in transgenic Arabidopsis roots; (iv) overexpression of MYB94 causes the activation of cuticular wax biosynthesis via up-regulation of cuticular wax biosynthetic genes, such as WSD1, KCS2/Daisy, CER1, CER2, FAR3 and ECR; and (v) the recombinant MYB94:MBP specifically binds to the MYB consensus motifs in the promoters of the WSD1, KCS2/Daisy, CER2, FAR3 and ECR genes. These findings provide a fundamental insight into the understanding of the regulatory network of cuticular wax biosynthesis and might be useful in the generation of transgenic crops with enhanced tolerance to drought stress.

Several R2R3-type MYB transcriptional activators, including MYB30, MYB41, MYB96, MYB16 and MYB106, have been reported to be involved in cuticular wax biosynthesis or in cuticle development (Cominelli et al. 2008, Raffaele et al. 2008, Seo et al. 2011, Oshima et al. 2013). MYB94, as well as MYB30, 96, 16 and 106, were expressed approximately 8-fold more in stem epidermal peels than in stems (Supplementary Fig. S1A; Suh et al. 2005). A particularly strong expression of MYB94 was observed in the aerial organs of Arabidopsis (Fig. 2A, C), and the transcript levels of MYB94 as well as MYB96 and 41 were significantly elevated in response to drought stress and ABA treatments (Fig. 2B; Supplementary Fig. S1A; Cominelli et al. 2008, Seo et al. 2011). Therefore the expression patterns of MYB94 indicate that MYB94 might play a role in an increased accumulation of cuticular wax under drought conditions.

It was reported that MYB96 and MYB30 activate VLCFA biosynthesis via up-regulation of genes encoding KCSs and/or KCR1, which are comprised of an FAE complex (Raffaele et al. 2008, Seo et al. 2011). In particular, the contents of most wax components, including aldehydes and alkanes, were remarkably increased in myb96-ID relative to the WT, which corresponds to an enhanced synthesis of VLCFAs as precursors for wax biosynthesis (Seo et al. 2011). Similarly, the amount of VLCFAs, primary alcohols and alkanes increased by approximately 35, 140 and 85%, respectively, in MYB94 OX leaves, as compared with the WT (Fig. 4). An increase in the specific wax components is closely associated with an elevated expression of KCS2/DAISY, ECR and CER2 genes in VLCFA synthesis (Zheng et al. 2005, Lee et al. 2009b, Haslam et al. 2012), a CER1 gene in the alkane-forming pathway (Bourdenx et al. 2011) and a FAR3/CER4 gene in the alcohol-forming pathway (Rowland et al. 2006), which are target genes of a MYB94 TF.

Interestingly, MYB94 and MYB96 genes might be generated from a duplication of an ancestral gene (Supplementary Fig. S1) and were shown to have a 98% deduced amino acid sequence identity in their DNA-binding domains, but their target genes were found to be distinguishable, except for the KCS2/DAISY gene. The target genes of MYB96 were reported to be KCS1, KCS2, KCS6 and KCR1 (Seo et al. 2011), whereas the direct target genes of MYB94 were WSD1, KCS2/DAISY, CER2, FAR3 and ECR (Fig. 6), suggesting that each TF independently and/or synergistically contributed to the activation of cuticular wax biosynthesis. According to Lee et al. (2009b), the levels of KCS2 transcripts were induced by approximately 30- and 65-fold after treatments with ABA and drought, respectively. We found that MYB96 and MYB94 were bound to the same cis-element (TAATCTAACATA) of the KCS2 promoter by EMSA and/or chromatin immunoprecipitation (ChIP) assay (Seo et al. 2011). Therefore the ABA- or drought-inducible expression of KCS2 is probably controlled by the synergistic regulation of MYB94 and MYB96. It would be interesting to understand how two different regulatory factors share a cis-acting element. It is possible that heterodimers of MYB96 and MYB94 may interact with the cis-element of the KCS2 promoter based on the evidence that two closely related R2R3-MYB proteins, MYB21 and MYB24, form homo- and heterodimers (Song et al. 2011).

In MYB94 OX leaves compared with the WT, the total amounts of cutin monomers were elevated, but the expression of genes involved in cutin synthesis was not significantly altered (Supplementary Fig. S3; Table S1). Although there is a possible explanation whereby unidentified genes involved in cutin synthesis are up-regulated by MYB94, the molecular mechanism underlying cutin synthesis in MYB94 OX plants is still unclear. Kannangara et al. (2007) suggested that the regulation of cutin and wax production by WIN1/SHN1 is a two-step process where the rapid induction of cutin synthesis is followed by the up-regulation of wax biosynthesis. Oshima et al. (2013) observed that MYB106 activates the expression of WIN1/SHN1 and both MYB106 and WIN1/SHN1 regulate cutin biosynthesis and wax accumulation in a co-ordinated manner.
The levels of MYB96, 106 and WIN1/SHN1 transcripts were observed to have not been significantly altered in MYB94 OX as compared with the WT, indicating that MYB94 might not be an upstream component of MYB96, MYB106 and WIN1/SHN1. Therefore, the relationship between MYB94 and the other TFs in the regulatory networks of cutin and wax biosynthesis remains to be further investigated.

One interesting point is that an increase in total wax loads was observed in leaves of MYB94 OX, but not in stems of MYB94 OX. In previous report, we observed that overexpression of MYB96 increased total wax loads by approximately 8.6-fold in leaves, but only 1.6-fold in stems (Seo et al. 2011). Kosma et al. (2014) also observed the similar phenotype that overexpression of MYB41 caused enhanced accumulation of suberin monomers and suberin-associated waxes in leaves, but the amount and composition of suberin monomers were not altered in the roots and seed coats that already contain a lot of suberin monomers. Therefore those results suggest that it might be difficult to observe a higher level of increased cuticular wax loads in stems, where total wax loads are already sufficient.

R2R3-type MYB transcription factors contain the conserved R2R3 domains required for DNA binding at the N-terminus and a highly variable transactivation domain at the C-terminus (Dubos et al. 2010). R2R3 domains are known to bind specifically to the consensus sequence motifs 'YAACKG' and 'CNGTTR' (Urao et al. 1993, Abe et al. 2003). For example, MYB96 directly interacts with MYB consensus motifs that are present in the promoter regions of KCS1, KCS2, KCS6, KCR1 and CER3 genes (Seo et al. 2011). The recombinant MYB94:MBP also specifically binds to the MYB consensus motifs in the WSD1, KCS2/DAISY, CER2, FAR3 and ECR promoters (Fig. 6, Supplementary Fig. S4). In addition, the deletion of the C-terminal region (amino acids 234–323) of MYB30 leads to the inactivation of pKCS1:GFP:GUS and pDHGFP:GUS reporter expression, compared with each reporter and intact MYB30 in the transient assays in N. benthamiana leaves (Raffaële et al. 2008), indicating that the transactivation domain of MYB30 is present at the C-terminus of MYB30. Similarly, MYB94 was observed to contain the transactivation domain in its C-terminus (Fig. 1C), and we also found that MYB94 harbors the glutamine- and the proline-rich C-terminal region, with characteristics that are frequently associated with transcription activation domains (Tanaka et al. 1994).

TFs were reported to have nuclear localization signals (NLSs) required for their targeting into the nucleus (Boulilka 1994). The fluorescent signals from the MYB94eYFP construct were observed to be localized in the nucleus in tobacco epidermis and in transgenic Arabidopsis roots (Fig. 1D). When the NLS motifs of MYB94 were searched for using a cNLS Mapper (http://nls-mapper.iab.keio.ac.jp/), the region of MYB94 (amino acids 62–92; RPKGKRGNFTHEEKMLHLHQALLGN RWAAI) that overlaps with the DNA-binding domain was detected to be a putative NLS motif. The region is also conserved in other MYB TFs, including MYB16, 30, 41, 96 and 106. In particular, the conserved arginine (R) and lysine (K) residues in the NLSs are probably essential for the import of MYB TFs (Klucevsek et al. 2007, Kosugi et al. 2008).

Cuticle biosynthesis is essential not only for plant growth and development but also for resistance to environmental stress. However, the regulatory mechanisms of cuticle biosynthesis are still largely unknown. In this study, we have identified a novel MYB94 TF that activates cuticular wax biosynthesis via direct transcriptional up-regulation of wax biosynthetic genes. The findings provide fundamental knowledge useful in understanding the molecular mechanisms underlying the regulation of cuticular wax biosynthesis. Considering the results of previous reports where elevated accumulation of cuticular waxes conferred drought tolerance (Aharoni et al. 2004, Zhang et al. 2005, Zhang et al. 2007, Seo et al. 2011), the overexpression of MYB94, which did not affect plant growth or development, could be applied to generating drought-tolerant transgenic crops.

Materials and Methods

Plant materials and growth conditions

The seeds of transgenic Arabidopsis (the background is Columbia-0, Col-0) were washed with 70% ethanol for 1 min, 20% bleach for 5 min, and sterilized water. They were then germinated on half-strength Murashige and Skoog (1/2 MS) medium including 1% sucrose and 0.7% phytagar. The germinated plants were planted on mixed soil (perlite : vermiculite : soil, 1 : 2 : 3) and were grown under long-day conditions (16 h/8 h, light/dark) in a growth room at 24 ± 2°C. Ten-day-old Arabidopsis plants grown in 1/2 MS medium were transferred to MS liquid medium containing 100 μM ABA, 200 mM NaCl or 200 mM mannitol for 1, 2 and 6 h for hormone and stress treatments. To treat drought stress, 2-week-old Arabidopsis plants were air-dried for 1 h and then for 2 h on filter paper.

Construction of binary vectors and Arabidopsis transformation

To generate MYB94 OX lines, MYB94 cDNA (1,002 p) was amplified from 2-week-old Arabidopsis plants using the gene-specific primers MYB94_cDNA_F1 (Smol) and MYB94_cDNA_R1 (BamHI), as shown in Supplementary Table S2. The amplified products were digested with Smol and BamHI, and were then inserted into the pZPF212 binary vector (Hajdukiewicz et al. 1994). For transient induction of MYB94 under the control of the β-stradiol-inducible promoter, MYB94 cDNA was amplified using MYB94_inducible_F1 (XhoI) and MYB94_inducible_R1 (Xhol). The amplified cDNA was cloned into the same restriction enzyme site of the pER8 vector (Zuo et al. 2000).

The generated constructs were transformed to Agrobacterium strain GV3101 using a freeze–thaw method. The transformed Agrobacterium cells were then transformed to Arabidopsis plants (Col-0) using vacuum infiltration. The seeds harvested from the transgenic plants were surface sterilized and were germinated on 1/2 MS medium. Each medium includes 25 μg ml−1 (w/v) kanamycin and 30 μg ml−1 (w/v) hygromycin. The selected transgenic plants (T1 or T2 plants) were transferred to mixed soil and were used for further experiments.

Transactivation assay in yeast

The transactivation assay was followed as described in Park et al. (2009). The MYB94-F, MYB94-N and MYB94-C constructs were produced via PCR using MYB94 cDNA and gene-specific primers (Supplementary Table S2). The amplified PCR products were cloned with pGEM T-easy vectors (Promega) and were digested with NdeI and BamHI. The digested constructs were cloned with the same restriction enzyme site of the pGBK7 vector (Clontech). The resultant constructs were transformed to yeast strain Y190 (MATa, HIS3, lacZ, rpl1, leu2, cyhr2) and then the transformant was selected from selection medium (5DL–Trp) including 25 mM 3-amino-1,2,4-aminitriazole (3-AT). The selected transformant was cultured in selection medium (5DL–Trp–His) including 25 mM 3-AT for 1 d, and a filter-lift assay was performed for...
blue color development in accordance with the method described in Breeden and Nasmyth (1985).

**Subcellular localization**

To develop the MYB94:eYFP recombinant plasmids, MYB94 cDNA was amplified using the primers described in Supplementary Table S2, digested with Smal restriction enzyme and then cloned into the same restriction enzyme site of the pCPZ212 binary vector. The developed construct was transformed to an Agrobacterium strain GV3101 and was then injected into the abaxial side of tobacco (*N. benthamiana*) leaf epidermis or transformed to Arabidopsis plants (Col-0). At 36–48 h after injection into tobacco leaves, the eYFP fluorescence was observed under a TCS SPS AOBS/Tandem laser confocal microscopic scope (Leica). In the case of Arabidopsis, the transgenic seeds were surface sterilized and germinated on the 1/2 MS agar medium including 25 μg·ml⁻¹ (w/v) kanamycin. Then, the expression of fluorescent protein in the root of 2-week-old transgenic plants (T₀) was observed. The eYFP fluorescence was detected with a YFP filter (excitation, 500 nm; emission, 535 nm). In the case of DAPI staining, the tissues expressing fluorescent protein were stained with DAPI solution (5 μg·ml⁻¹ in phosphate-buffered saline (PBS)) for 5 min at room temperature, and the signals were observed with a UV filter (excitation, 355 nm; emission, 450 nm) (Yao et al. 2013).

**Gene expression analysis**

Total RNAs were isolated from various Arabidopsis tissues and from 2-week-old Arabidopsis plants treated with ABA, drought and osmotic stress using an RNEasy® Plant Mini Kit (QIAGEN). Reverse transcription was performed using GoScript™ Reverse Transcriptase (Promega), and qRT–PCR was performed using SYBR Green I master mix (KAPA) in a 20 μL volume. Gene-specific primers, as described in Supplementary Table S2, and a Bio-Rad CFX96 Real-Time PCR system were used for PCR analysis. The PP2A (At1g13320) gene was used for normalization.

**GUS staining and microscopy**

The MYB94 promoter region (approximately 4.2 kb) was amplified using the MYB94_promoter_F3 and MYB94_promoter_R3 primers and was cloned into the HindIII and Smal site of the pCAMBIA1391Z binary vector (Supplementary Table S2). The produced construct was transformed to the *Agrobacterium* strain GV3101 and transformed to Arabidopsis plants (Col-0). The transgenic plants were selected from 1/2 MS medium including 30 μg·ml⁻¹ hygromycin and were used for histochemical analysis of GUS activation with the gas chromatograph (GC-2010, Shimazu; column 60 μm H-P, 0.32 mm i.d., df = 0.25 mm, Agilent) for quantitative element analysis. The analysis conditions are as follows: injection at 220 °C for 4.5 min, an increase of 3 °C min⁻¹ to 290 °C, maintained for 10 min at 290 °C, an increase of 2 °C min⁻¹ to 300 °C, and maintained for 15 min at 300 °C.

**Microarray assays**

Affymetrix GeneChip® Arabidopsis Genome ATH1 arrays were carried out to determine the gene expression analysis. The method is as follows: the total RNA was isolated from the leaves of WT and MYB94 OX-1 (3- to 4-week-old) plants, and the contaminated DNA was removed using RNase-free DNase I (QIAGEN). Synthesis of biotin-labeled cRNA from total RNA, hybridization, detection and scanning were performed according to the manufacturer’s procedure. The raw.cel files were used for further analysis. The expression data were generated using Expression Console software (version 1.1). The Affymetrix microarray suite 5 (MASS) algorithm was used for normalization. To delete the noise during the process of significant gene searching, the probes that did not determine a Present call (>50%) among the MASS detection calls were excluded from the analysis. The probe set that showed a >1.5-fold change compared with the control were selected as differentially regulated genes for the next analysis. The web analysis tool (DAVID, Database for Annotation, Visualization, and Integrated Discovery) was used for biological interpretation analysis of differentially regulated genes, and each functional group was classified by gene function information from the KEGG, Gene ontology database (http://www.ebi.ac.uk/arrayexpress). The microarray data were deposited in ArrayExpress with accession number E-MTAB-2870 at https://www.ebi.ac.uk/arrayexpress.

**Transcriptional activation assays**

The transcriptional activation assays were performed as described in Go et al. (2014). In order to investigate the MYB94 transcriptional activator function, the construct used for development of the overexpression plant (p3SS-MYB94) was used as an effector plasmid. The promoter region of the cuticular wax biosynthetic genes was fused with luciferase as a reporter plasmid (Supplementary Table S2). The reporter and effector plasmid DNA were co-transfected into tobacco leaf protoplasts and were placed under dark conditions for 16 h. The protoplasts were used for the luciferase activity assay and for the GUS enzymatic assay using a dual-luciferase reporter assay system luminescence reader (GROMAX-20/20; Promega). At this stage, the luciferase gene expression was equilibrated by the GUS gene expression, showing the relative scores.

**Electrophoretic mobility shift assays**

The MYB94 N-terminal region (370 bp) was amplified using MYB94-Bam-F and MYB94_yeast_NR1 primer (Supplementary Table S2) to perform the EMSAs, and then it was inserted into the BamHI site of the pMAL-c2X E. coli expression vector (NEB), including an MBP-coding sequence. The MYB94-MBP protein was extracted using a pMAL™ Protein Fusion and Purification System (New England Biolabs) method. The synthesized double-stranded oligomers were end-labeled with [γ-32P]ATP using T4 polynucleotide kinase (TAKARA) and were then desalted using chloroform. The extracted recombinant proteins (5 μg) were reacted at room temperature with labeled probes for 30 min in a binding buffer (10 mM Tris–HCl, pH 7.6, 50 mM NaCl, 1 mM Na₂EDTA, 5 mM dithiothreitol (DTT), 5% glycerol) including 100 ng of poly(dI–dC) under conditions with or without competitors. The reaction mixtures were then electrophoresed by 6% native PAGE, and the gel was dried on paper for scanning in a Typhoon FLA 7000 phosphorimager (GE Healthcare Life Science).

**Cuticular transpiration assay**

The rate of water loss through the cuticle was estimated using well-watered leaves and drought-treated leaves of 4-week-old plants. Two-week-old plants were subjected to drought stress by withholding water for 2 weeks and were...
then rehydrated for 1 d. The rehydrated plants were acclimated in the dark for 12 h and were then soaked in water for 1 h. The weight of the rosette leaves was measured using a micro balance in the dark under a green light (wavelength of approximately 510 nm).

Supplementary data

Supplementary data are available at PCP online.

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Disclosures

The authors have no conflicts of interest to declare.

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