Arabidopsis Sphingolipid Fatty Acid 2-Hydroxylases (AtFAH1 and AtFAH2) Are Functionally Differentiated in Fatty Acid 2-Hydroxylation and Stress Responses1[OA]

Minoru Nagano, Kentaro Takahara, Masaru Fujimoto, Nobuhiro Tsutsumi, Hirofumi Uchimiya, and Maki Kawai-Yamada*

Graduate School of Biological Science, Nara Institute of Science and Technology, 8916–5 Takayama, Ikoma 630–0101, Japan (M.N.); Institute of Molecular and Cellular Biosciences, University of Tokyo, Bunkyo-ku, Tokyo 113–0032, Japan (K.T.); Graduate School of Agricultural and Life Sciences, University of Tokyo, Bunkyo-ku, Tokyo 113–8657, Japan (M.F., N.T.); Graduate School of Science, University of Tokyo, Bunkyo-ku, Tokyo 113–0033, Japan (M.F.); Institute for Environmental Science and Technology (H.U., M.K.-Y.) and Department of Environmental Science and Technology (M.K.-Y.), Saitama University, Sakura-ku, Saitama 338–8570, Japan; and Iwate Biotechnology Research Center, Kitakami, Iwate 024–0003, Japan (H.U.)

2-Hydroxy fatty acids (2-HFAs) are predominantly present in sphingolipids and have important physicochemical and physiological functions in eukaryotic cells. Recent studies from our group demonstrated that sphingolipid fatty acid 2-hydroxylase (FAH) is required for the function of Arabidopsis (Arabidopsis thaliana) Bax inhibitor-1 (AtBI-1), which is an endoplasmic reticulum membrane-localized cell death suppressor. However, little is known about the function of two Arabidopsis FAH homologs (AtFAH1 and AtFAH2), and it remains unclear whether 2-HFAs participate in cell death regulation. In this study, we found that both AtFAH1 and AtFAH2 had FAH activity, and the interaction with Arabidopsis cytochrome b5 was needed for the sufficient activity. 2-HFA analysis of AtFAH1 knockdown lines and atfah2 mutant showed that AtFAH1 mainly 2-hydroxylated very-long-chain fatty acid (VLCFA), whereas AtFAH2 selectively 2-hydroxylated palmitic acid in Arabidopsis. In addition, 2-HFAs were related to resistance to oxidative stress, and AtFAH1 or 2-hydroxy VLCFA showed particularly strong responses to oxidative stress. Furthermore, AtFAH1 interacted with AtBI-1 via cytochrome b5 more preferentially than AtFAH2. Our results suggest that AtFAH1 and AtFAH2 are functionally different FAHs, and that AtFAH1 or 2-hydroxy VLCFA is a key factor in AtBI-1-mediated cell death suppression.

Sphingolipids are a large class of lipids present ubiquitously in eukaryotic cells and involved in various cellular processes such as signal transduction, protein transport, and programmed cell death as structural components of the plasma membrane and tonoplast and as signaling molecules (Dunn et al., 2004; Pata et al., 2010). These numerous functions are based on the structural diversity of sphingolipids. Complex sphingolipids are largely formed by a variety of head groups and a ceramide, which is comprised of a long-chain base amide linked to a fatty acid. One of the main characteristics of sphingolipid fatty acids is that large number of sphingolipids contained very-long-chain fatty acids (VLCFAs; fatty acids with ≥ C20; Raffaele et al., 2009). Another feature is the 2-hydroxylation of fatty acids (Alderson et al., 2005). The 2-hydroxy fatty acid (2-HFA)-containing sphingolipids are present in most organisms, including yeast (Saccharomyces cerevisiae), vertebrates, and plants. Especially in higher plants, 2-HFAs are contained in more than 90% complex sphingolipids such as glucosylceramides and glycosylinositolphosphorylceramides (Imai et al., 1995; Pata et al., 2010).

The 2-hydroxylation of sphingolipid N-acyl chains is catalyzed in yeast and mammals by the enzyme sphingolipid fatty acid 2-hydroxylase (ScFAH1 and F2AH, respectively, and we refer to them as FAH in this report; Mitchell and Martin, 1997; Alderson et al., 2004; Hama, 2010). FAH is an endoplasmic reticulum (ER)-localized membrane protein composed of two domains: an N-terminal cytochrome b5 (Cb5)-like domain and a putative catalytic site with the His motif (HXXHH), which coordinates the nonheme diiron cluster at the active site of the enzyme and is highly conserved among membrane-bound desaturases and hydroxylases. Mammalian F2AH is reported to catalyze
the 2-hydroxylation of free fatty acid in vitro, which was dependent on a reconstituted electron transport system (Alderson et al., 2005). In mammals, FA2H-synthesized 2-HFAs abundantly accumulate in epidermal keratinocytes and the myelin-forming cells of the nervous system (Alderson et al., 2006; Uchida et al., 2007; Maldonado et al., 2008). Loss of FA2H in mouse altered the composition and physicochemical properties of sebum, causing alopecia (Maier et al., 2011). In addition, FA2H deficiency also causes a wide range of neurodegeneration phenotypes such as spastic paraplegia, leukodystrophy, and brain ion deposition (Schneider and Bhatia, 2010). Moreover, FA2H is a negative regulator of the cell cycle and facilitates dibutyryl-cyclic AMP-induced cell cycle exit in D6P2T schwannoma cells. In addition, FA2H-synthesized 2-hydroxy VLCFAs. These results allowed us to speculate that AtBI-1 interacts with AtFAHs via AtCb5 to activate the synthesis of 2-hydroxy VLCFA in Arabidopsis, which leads to the suppression of cell death. However, it remains unclear whether 2-hydroxy VLCFA is related to the regulation of cell death or stress responses. Furthermore, little is known about the function of AtFAH1 and AtFAH2 in 2-HFA synthesis of plant cells.

The results of this study confirmed that both AtFAHs are FAHs, and identified functional differences between AtFAH1 and AtFAH2 in fatty acid 2-hydroxylation in Arabidopsis. In addition, AtFAH1 modified the resistance to oxidative stress more strongly than AtFAH2, which was consistent with the preferential interaction with AtBI-1.

RESULTS

AtFAH1 Also Interacts with AtCb5-B in the ER of the Plant Cells

We showed previously that AtFAH2 is localized in the ER (Nagano et al., 2009). To investigate whether AtFAH1 has the same function as AtFAH2 or not, we first confirmed the subcellular localization of AtFAH1. Fusion constructs of AtFAH1 were generated using genes encoding GFP driven by the cauliflower mosaic virus 35S (CaMV35S) promoter. When bombarded in onion (Allium cepa) epidermal cells, AtFAH1-GFP was localized in the cell periphery, the perinuclear region, and the reticulum network (Fig. 1, A and D). In addition, AtFAH1-GFP was colocalized always with AtFAH2-RFP (Fig. 1, A–C), suggesting that AtFAH1 also localizes in the ER.

As described above, neither AtFAH1 nor AtFAH2 has any Cb5-like domains (Mitchell and Martin, 1997; Nagano et al., 2009). Alderson et al. (2004) demonstrated that the Cb5-like domain is essential for human FA2H activity, implying that AtFAH1 and AtFAH2 require an ER-localized AtCb5 to increase their activity. In fact, we have already demonstrated that both AtFAH1 and AtFAH2 interacted with all ER-localized AtCb5 in yeast cells. In addition, the interaction between AtFAH2 and AtCb5-B, which is one of four ER-type AtCb5s, was reported in plant cells using a fluorescence resonance energy transfer (FRET) analysis (Nagano et al., 2009). Accordingly, we attempted to verify the interaction between AtFAH1 and AtCb5-B in plant cells by using a bimolecular fluorescence complementation (BiFC) assay. The nYFP-pUGW0 vector contains nYFP (1–174 amino acids) in the N-terminal side of the target proteins, driven by CaMV35S promoter, whereas the cYFP-pUGW2 vector contains cYFP (175–239 amino acids) in the C-terminal side (Nakagawa et al., 2007; Fig. 1E). When nYFP-AtCb5-B and AtFAH1-cYFP were transiently coexpressed in onion epidermal cells, we observed BiFC fluorescence that was highly localized in the cell periphery and the perinuclear regions (Fig. 1, F and L). This signal was similar to that of nYFP-AtCb5-B/AtFAH2-cYFP in terms of localization and intensity (Fig. 1, F, H, L, and N). In contrast, no fluorescent signal was visible using empty vectors (nYFP/cYFP; Fig. 1J). In these studies, DsRed driven by CaMV35S promoter was used as an expression marker of particle bombardment (Fig. 1, G, I, K, M, and O). These results suggest that not only AtFAH2 but also AtFAH1 interacts with AtCb5-B in plant cells.

AtFAHs Exhibit FAH Activity in Yeast Cells

To clarify whether AtFAH1 and AtFAH2 are FAHs, we performed a complementation test using yeast ScFAH1-deletion variant (Δfah1), which cannot produce any 2-HFAs. In these experiments, AtFAH1 or AtFAH2
FAH activity. However, these levels were quite less than those of the wild type and Δfah1 expressing ScFAH1 (Fig. 2). This finding may be due to the structural difference between ScFAH1 and AtFAHs with or without Cb5-like domain, respectively (Mitchell and Martin, 1997; Nagano et al., 2009). Transformation of AtCb5-B into AtFAH1- or AtFAH2-expressing Δfah1 resulted in a marked increase in the amounts of 26h:0 (Fig. 2). AtCb5-B itself exhibited no FAH activity, because no 26h:0 was detected in AtCb5-B-expressing Δfah1 (Fig. 2). These results indicate that AtCb5 is a key factor for the activity of AtFAH1 and AtFAH2.

Different Roles for AtFAH1 and AtFAH2 in 2-HFA Synthesis in Arabidopsis

To elucidate the roles of AtFAH1 and AtFAH2 in 2-HFA synthesis pathway and the physiological functions in Arabidopsis, we searched for AtFAHs knockout lines. However, no suitable T-DNA insertion line for AtFAH1 is available among the existing T-DNA mutant collections. We therefore generated AtFAH1 knockout transgenic lines (AtFAH1-KD) by RNA interference with 100 bp of 3′ untranslated region in AtFAH1 cDNA as a trigger (Fig. 3A). Reverse transcription (RT)-PCR analysis confirmed that AtFAH1 was markedly reduced in AtFAH1-KD-1 and -2 lines (Fig. 3C). In contrast, we obtained T-DNA-tagged Arabidopsis line in AtFAH2 gene (SALK_011293, atfah2), which has a T-DNA insertion in third exon (Fig. 3B). As shown in Figure 3C, not only the full-length AtFAH2 mRNA (primer a-b), but also the upstream region of T-DNA insertion site in AtFAH2 gene (primer a-c) was completely deleted. AtFAH1-KD and atfah2 plants showed no apparent phenotype compared with the 3-week-old wild-type (Columbia-0) plants under normal growth condition (Fig. 3D).

We first dissected 2-HFA composition in these transgenic plants. Total lipids including total sphingolipids were extracted from the aboveground parts in 3-week-old plants according to Markham et al. (2006), and these 2-HFA methyl esters were detected with GC-MS after derivatization. In this study, we analyzed five major 2-HFAs (16h:0, 22h:0, 24h:0, 24h:1, and 26h:0) of 13 2-HFAs in Arabidopsis (Imai et al., 2000). In the AtFAH1-KD lines, the levels of 22h:0, 24h:0, and 26h:0 were considerably lower than in wild type, whereas the level of 16h:0 was marginally decreased (Fig. 4, A and B). In contrast, the level of 16h:0 was markedly reduced by 22.9% in atfah2 mutant, whereas the levels of other 2-HFAs were similar to those in wild type (Fig. 4, A and B). Thus, these results suggest that VLCSA is 2-hydroxylated by AtFAH1, whereas palmitic acid is 2-hydroxylated by AtFAH2 and marginally AtFAH1.

2-HFAs Are Implicated in Resistance to Oxidative Stress and 2-Hydroxy VLCFAs Respond to Oxidative Stress

Our previous report suggested that FAH is related to AtBI-1-mediated suppression of cell death (Nagano et al., 2009). As shown in Figure 2, 26h:0 accumulated in AtFAH1- or AtFAH2-expressing Δfah1, but not in Δfah1, suggesting that both AtFAH1 and AtFAH2 have

Figure 1. AtFAH1 interacts with AtCb5-B in the ER of plant cells. A to D, Distribution of AtFAH1-GFP (A and D) and AtFAH2-RFP (B). Onion epidermal cells transiently coexpressing AtFAH1-GFP and AtFAH2-RFP were examined by confocal laser-scanning microscopy. The merged images of A and B are shown in C. Scale bars in A, B, and C = 100 μm. Scale bar in D = 5 μm. Arrow indicates the perinuclear region. E, Schematic representation of BiFC constructs. The coding regions of AtCb5 and AtFAH1 or AtFAH2 were fused with the N-terminal (nYFP, 1–174 amino acids) and the C-terminal (175–239 amino acids) region of YFP, respectively, p35S, CaMV 35S promoter; tNOS, terminator of the nopaline synthase (NOS) gene. F to O, BiFC visualization in onion epidermal cells transiently coexpressing AtFAH1-GFP and AtFAH2-RFP (nYFP-AtCb5-B and cYFP-AtFAH). Images were examined by confocal laser microscopy. Scale bars = 100 μm. Arrows in A indicate the perinuclear region.
et al., 2009). Because AtBI-1 is reported to suppress oxidative-stress-induced cell death (Kawai-Yamada et al., 2004; Ishikawa et al., 2010), we tested the sensitivity of AtFAH1-KD and atfah2 plants to oxidative stress by using ROS-inducing chemicals, hydrogen peroxide (H$_2$O$_2$), and menadione (MD; Yoshinaga et al., 2006). Treatment of leaf discs obtained from 3-week-old plants with 100 mM H$_2$O$_2$ or 60 μM MD for 24 h significantly increased electrolyte leakage in AtFAH1-KD and atfah2 plants compared with wild type (Fig. 5A). Electrolyte leakage measurement is the assay often used to examine irreversible membrane damage by oxidative stress or during cell death. In addition, chlorophyll contents in AtFAH1-KD and atfah2 plants were lower than wild type (Fig. 5B), indicating that AtFAH1-KD and atfah2 plants have higher sensitivity to oxidative stress. Moreover, AtFAH2 gene controlled by its native promoter (pAtFAH2::AtFAH2, Fig. 5C) complemented the atfah2 phenotype for oxidative-stress-induced ion leakage and chlorophyll contents (Fig. 5, A and B). These results suggest the involvement of 2-HFAs in the resistant response to oxidative stress.

Next, the expression levels of AtFAH1 and AtFAH2 mRNAs in response to H$_2$O$_2$ were examined. Total mRNAs were extracted from 10-d-old wild-type plants treated with 100 mM H$_2$O$_2$ for 1, 3, 5, 7, and 24 h. RT-PCR analysis showed that AtFAH1 mRNA increased rapidly at 1 h and subsequently remained at higher levels than no treatment, although they were decreased slightly with time (Fig. 6A). AtFAH2 mRNA diminished gradually from 3 h and was only expressed weakly at 24 h. Incidentally, AtBI-1 mRNA continued to increase gradually in response to H$_2$O$_2$ until 24 h. These results imply that AtFAH1 promptly responds to oxidative stress, whereas AtFAH2 does not.

To clarify the roles of 2-HFAs in the response to oxidative stress, we examined changes in fatty acids in response to oxidative stress. Four-week-old Arabidopsis wild-type plants were treated with H$_2$O$_2$ for 24

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**Figure 2.** AtFAH1 and AtFAH2 exhibit FAH activity. Complementation test of AtFAH1 and AtFAH2. Yeast fah1-deletion variant was transformed with plasmids possessing ScFAH1, AtFAH1, AtFAH2, or AtCb5-B, and total lipids were extracted from each. 26h0 methyl ester was analyzed by GC-MS. Data are mean ± so of four experiments.

**Figure 3.** AtFAH1-KD lines and AtFAH2 T-DNA insertion mutant. A, Structure of the AtFAH1 gene. Red line represents the region which was used as a trigger of RNA interference. B, Structure of the AtFAH2 gene and designated locations of the T-DNA. Arrows indicate the sites of primers used for PCR. C, RT-PCR analysis of AtFAH1, AtFAH2 (a-b), AtFAH2 (a-c), and Actin 2 (ACT2) expression in 10-d-old seedlings of wild-type, AtFAH1-KD, and atfah2 plants. Representative results of five experiments with similar findings. D, Phenotypes of 3-week-old AtFAH1-KD and atfah2 plants.
h, and their derivatized 2-HFA methyl esters and nonhydroxy fatty acid (non-HFA) methyl esters were analyzed by GC-MS. As shown in Figure 6B, the total non-HFA content was significantly lower at 24 h after treatment with H₂O₂. In contrast, total 2-HFA content was higher at 24 h after treatment with H₂O₂ (Fig. 6C). However, the amount of 16h:0 was not affected by H₂O₂ (Fig. 6D), whereas the levels of total 2-hydroxy VLCFAs were especially elevated (Fig. 6E). These results suggest the importance of 2-hydroxy VLCFAs under oxidative stress condition.

To further confirm the relationship between AtFAHs and oxidative stress, we analyzed the levels of oxidative-stress-related metabolites in the leaves of 3-week-old AtFAH1-KD and atfah2 plants grown under normal growth conditions by using a capillary electrophoresis (CE)-MS. As shown in Figure 7, reduced glutathione (GSH) level in AtFAH1-KD was significantly lower than wild-type plants, whereas oxidized glutathione (GSSG) was markedly higher, resulting in reduced rate of GSH in total glutathione in spite of the lower total glutathione in AtFAH1-KD lines (Fig. 7A–D). In addition, reduced ascorbate was markedly lower in AtFAH1-KD lines (Fig. 7E). Moreover, γ-aminobutyric acid (GABA) was markedly increased compared with wild-type plants (Fig. 7F). In contrast, the levels of glutathione, reduced ascorbate, and GABA in atfah2 were similar to those in wild type (Fig. 7, A–F).

Ascorbate (vitamin C) is the most important reducing substrate for H₂O₂ detoxification in plant cells, and the series of ROS-scavenging reactions accompanied by glutathione is called the ascorbate-glutathione cycle (Nakano and Asada, 1987; Mehlhorn et al., 1996; Noctor and Foyer, 1998). When plants are exposed to ROS-mediated stresses, glutathione-ascorbate cycle participates in detoxification of ROS at the expense of electrons from the GSH pool, causing a transient increase in GSSG (Noctor and Foyer, 1998). In addition, GABA is a four-carbon nonprotein amino acid that is rapidly and markedly produced in response to biotic and abiotic stress (Kois and Turano, 2000; Bouché and Fromm, 2004). Therefore, these results indicate that AtFAH1-KD lines are exposed to some stress even under nonstress conditions. Taken together, the results indicate that 2-HFAs seem important in the
AtBI-1 preferentially interacts with AtFAH1 via AtCb5-B

To reinforce the link between AtFAH and AtBI-1-mediated suppression of cell death, we examined the three-way interaction among AtBI-1, AtCb5, and AtFAH in plant cells using the BiFC-based FRET assay in plant cells. This assay detects ternary complexes in living cells by the combination of BiFC and FRET assays (Shyu et al., 2008a, 2008b; Kwaaitaal et al., 2010; Taoka et al., 2011). When the combination of AtBI-1-CFP, nYFP-AtCb5-B, and AtFAH2-cYFP, and the combination of AtBI-1-CFP, nYFP-AtCb5-B, and AtFAH1-cYFP were transiently expressed in onion epidermal cells, CFP fluorescence and BiFC signal were completely merged in both combinations (Fig. 8, A–F). We next examined the interactions between AtBI-1 and AtCb5-B/AtFAH1 or AtCb5-B/AtFAH2 by FRET-acceptor photobleaching analysis. The mean FRET efficiency was 17.4% ± 5.58% for the pair of AtBI-1-CFP and nYFP-AtCb5-B/AtFAH1-cYFP (Fig. 8G). In contrast, the pair of AtBI-1-CFP and nYFP-AtCb5-B/AtFAH2-cYFP showed a lower-level FRET efficiency (6.8% ± 2.72%), although the efficiency was significantly higher than control experiments (CFP + YFP, 3.19% ± 2.8%; Fig. 8G). These results suggest preferential ternary interactions among AtBI-1, AtCb5, and AtFAH1-KD and atfah2 plants. A and B, Leaf discs of wild-type, AtFAH1-KD, atfah2, and atfah2/pAtFAH2::AtFAH2 plants were exposed to H2O2 or MD for 24 or 6 h, respectively. A, Relative ratios of ion leakages to total ion. B, Relative ratios of chlorophyll contents to total chlorophyll contents. Data in A and B are mean ± SD of four biological replicates. *P < 0.05, **P < 0.01, compared with wild type (t test). C, RT-PCR analysis of AtFAH2 and ACT2 expression in 10-d-old seedlings of wild-type, atfah2, and atfah2/pAtFAH2::AtFAH2 plants. Representative results of three experiments with similar findings.

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AtFAH1, and strongly support the significance of AtFAH1 in AtBI-1-mediated cell death regulation (Fig. 9).

**DISCUSSION**

**Importance of AtCb5 in AtFAH Activity**

It has been reported that ScFAH1 and mammalian FA2H contain a Cb5-like domain at their N terminus, and that the truncated FA2H lacking the Cb5-like domain has only a low level of FAH activity, indicating that the Cb5-like domain is required for efficient hydroxylation activity (Mitchell and Martin, 1997; Alderson et al., 2004). However, the Arabidopsis AtFAH1 and AtFAH2 lack the Cb5-like domain. This study demonstrated that both AtFAH1 and AtFAH2 were activated by AtCb5-B through their interaction. In Arabidopsis, there are five Cb5s (AtCb5-A to -E), and AtCb5-B, -C, -D, and -E localized in the ER membrane (Maggio et al., 2007; Nagano et al., 2009). Cb5 is a ubiquitous eukaryotic electron transfer protein, and is related to a number of ER-linked redox enzyme systems such as hydroxylation, desaturation, and elongation (Mathews, 1985; Schenkman and Jansson, 2003). In addition, AtFAHs also possess several His motifs to receive electrons like ScFAH1 and FA2H. Therefore, AtFAHs would also need electrons from ER-localized Cb5 to work sufficiently, although there is a great difference among them in existence or lack of the Cb5-like domain.

**Fatty Acid 2-Hydroxylation through AtFAH1 and AtFAH2 in Arabidopsis**

Mammalian FA2H had no substrate specificity because FA2H is a single gene and the protein can generate almost all kinds of 2-HFAs when overexpressed (Alderson et al., 2004). However, analysis of 2-HFAs in Arabidopsis demonstrated that AtFAH1 2-hydroxylated all kinds of fatty acid, especially VLCFA, whereas AtFAH2 2-hydroxylated only palmitic acid in Arabidopsis cells. This finding suggests that
AtFAH1 and AtFAH2 are used differently depending on the length of fatty acids in Arabidopsis.

The synthetic pathway of VLCFA is quite different from that of long-chain fatty acid (LCFA). LCFA, such as palmitic acid and stearic acid, are produced in plastids, whereas VLCFAs are generated in the ER membrane by a special elongase system comprising a four-step reaction cycle: condensation by 3-ketoacyl-CoA synthase; reduction by 3-ketoacyl-CoA reductase; dehydration by 3-hydroxyacyl-CoA dehydratase; and further reduction by trans-2-enoyl-CoA reductase (Bach and Faure, 2010). Therefore, it may be reasonable that the substrates of AtFAHs are differentiated between LCFA and VLCFA in Arabidopsis cells.

However, there might be no substrate specificity between AtFAH1 and AtFAH2, because the complementation assay showed that AtFAH2 could produce 26h:0 in yeast cells. One can speculate that VLCFAs are not supplied into AtFAH2 as substrates in Arabidopsis cells, although AtFAH2 could 2-hydroxylate VLCFAs. Alternatively, AtFAH1 might be associated also with the elongase system in the ER membrane of Arabidopsis cells. Further studies are needed to investigate the relationship between AtFAHs and elongase system or transportation of fatty acids. In addition, it is interesting to examine substrate specificities of AtFAHs in vitro.

On the other hand, above differentiation in Arabidopsis might not be applicable to other plants such as *Oryza sativa* and *Zea mays*, because they have only a few 2-hydroxy LCFA, including 2-hydroxy palmitic acid in spite of possessing two FAH homologs (Imai et al., 1995; Pata et al., 2010). Therefore, it is also interesting to analyze the substrate of FAHs in a large number of plants and understand their differences.

The Relationship between AtFAH and AtBI-1 in Cell Death Suppression

Our previous studies showed that AtBI-1 required FAH function to suppress cell death, implying that spherolipids with 2-HFAs play an important role in cell death regulation (Nagano et al., 2009). In this study, AtFAH1-KD and atfah2 plants were more sensitive to H2O2 and MD than wild type, suggesting that 2-HFAs provide tolerance to oxidative stress, regardless of their length.

2-HFAs are considered to function as a part of spherolipids, because they are mostly contained within sphingolipids in higher plants. A number of previous studies demonstrated the relationship between sphingolipids and cell death regulation in not only animals but also plants (Taha et al., 2006; Hannun and Obeid, 2008; Pata et al., 2010). Focused on the sphingolipid fatty acids, non-HFA ceramide induces apoptosis, whereas 2-HFA ceramide is less proapoptotic in mammalian U937 cells (Ji et al., 1995). In addition, Uchida et al. (2007) proposed that 2-hydroxylation of sphingolipid fatty acids can protect mammalian keratinocytes from apoptosis. In keratinocytes, 2-HFA ceramide and 2-HFA glucosylceramide are markedly increased during differentiation. Interestingly, differentiated keratinocytes are more resistant to apoptosis induced by ultraviolet irradiation and by tumor necrosis factor related apoptosis-inducing ligands (Chaturvedi et al., 1999; Magnoni et al., 2002; Jansen et al., 2003; Uchida et al., 2003). Similarly, the uptake of 2-HFA ceramide did not affect programmed cell death in Arabidopsis suspension cells, whereas non-HFA ceramide as well as C2-ceramide induced cell death (Townley et al., 2005). Therefore, 2-HFA spherolipids may be important to cell death suppression or stress resistance in plant cells.

On the other hand, oxidative stress induced a rapid increase in *AtFAH1* mRNA and production of a large amount of 2-hydroxy VLCFAs. In addition, knockdown of *AtFAH1* affected the amounts of oxidative-stress-related metabolites such as ascorbate, glutathione, and GABA. Moreover, the BiFC-FRET assay demonstrated that AtFAH1 interacted with AtBI-1 via AtCb5-B more strongly than AtFAH2. These results are in agreement with our previous data that overexpression of *AtBI-1* in Arabidopsis is associated with increased 2-hydroxy VLCFAs (Nagano et al., 2009). These findings suggest that AtFAH1 is involved in AtBI-1 function more than AtFAH2, although atfah2 are also hypersensitive to oxidative stress like AtFAH1-KD lines. It is possible that the presence of 16h:0 in the membranes of plant cells seems important, because the expression of *AtFAH2* mRNA were gradually decreased and no obvious changes were noted in the amount of 16h:0 and oxidative-stress-related metabolites in atfah2 mutant. One can speculate that membrane permeability is probably enhanced in atfah2 mutant. Further studies are necessary to clarify the difference in the effects of 2-HFA length on cell death regulation and the significance of 2-hydroxy VLCFA. In addition, 2-HFAs might be essential for plant development, because we could not obtain AtFAH1-KD/atfah2 double mutants until now. We firmly believe that more vigorous efforts using AtFAH1 and AtFAH2 can shed more information on
the meaning of 2-HFA, because Arabidopsis has two functionally different FAHs.

MATERIALS AND METHODS

Plant Materials

The Columbia ecotype of Arabidopsis (Arabidopsis thaliana) was used in all experiments. The T-DNA insertion mutant of AIFAH2 was obtained from the SALK Institute (SALK_011293). To generate AIFAH1-KD and aflibl/2mAIFAH2-AIFAH2 lines, pK7GWiWG2(S)-AIFAH1 and pMDK9-AIFAH2 were constructed, respectively. Briefly, the 3′-untranslated region of AIFAH1-cDNA (amplified by the primers: 5′-ACAAGACCGACGAAAAGGAAGGTTTC-3′ and 5′-AGAAGGATGATTTAACACT-3′), or putative native promoter and gene region of AIFAH2 (amplified by the primers: 5′-GTGAGGAACCTTACATCGC-3′ and 5′-CCGGAATCAGAAGAAATGGAATCT-3′) were introduced into the pK7GWiWG2(S) or pMDK9 plasmids, respectively (Kanini et al., 2002; Curtis and Grossniklaus, 2003), by the Gateway system. The resulting plasmid was transferred to A. thaliana by particle bombardment with helium-driven particle accelerator (GIE-3 IDERA; TANAKA). The T-DNA insertion mutant of AIFAH2 was also cobombarded to the cells as an expression marker. CaMV35S-DsRed was also cobombarded to the cells as an expression marker.

Confocal laser microscopy was carried out with Nikon TE-2000U (Nikon) and 570-nm long-pass filter for RFP.

Oxidative Stress Treatment

For measurement of ion leakage and chlorophyll content, and for RT-PCR analysis under oxidative stress condition, three leaf discs (5.5 mm in diameter) obtained from 3-week-old Arabidopsis plants were submerged in 1 mL of distilled water. After adding H2O2, or MD (Sigma) to become 100 mM or 60 μM, respectively, the leaf discs were vacuumed for 5 min. The floating discs were incubated at 23°C under continuous light (60 μmol m−2 s−1) without shaking for several times.

For fatty acid analysis, 250 mg of whole rosettes of Arabidopsis were submerged in 20 mL of 100 mM H2O2 (Wako), which was diluted with distilled water, for 24 h, and washed in distilled water before extraction of total lipids.

RT-PCR

Total RNA was extracted from Arabidopsis leaf discs of seedlings using the RNeasy plant mini prep kit (Qiagen) and 1 μg was used as a template for RT-PCR with specific primers for AIFAH1 (amplified by the primers: 5′-ATGTGGCTGCAAGGATCAC-3′ and 5′-TGCTCTTCCTTCGCGCTT-3′), AIFAH2 (amplified by the following primers: 5′-ATGTGGCTGAGAAGGATA-3′ and 5′-GCTCTTCCTCCAGGCCGCTT-3′), AtIB1 (amplified by the primers: 5′-ATGGATGCTCTTCTTCGCTT-3′ and 5′-GTTCTCTTCTTTCTTCT-3′), Actin2 (amplified by the primers: 5′-GAGATTCACATGGCAGAATGCTG-3′ and 5′-ACTCTGCCATCATACCTGCGC-3′), and AtAct8 (amplified by the primers: 5′-TGACGCTCCGCCATGTCG-3′ and 5′-TCTCCTTGCCTGATCGCA-3′). The conditions for direct amplification with Ready-to-Go RT-PCR beads (GE Health Care) were one cycle at 42°C for 25 min and 95°C for 5 min; 25 cycles at 95°C for 30 s, at 55°C for 30 s, and at 72°C for 1 min.

Yeast Strain and Lipid Extraction

The yeast (Saccharomyces cerevisiae) gene disruption mutant (ΔΔib1Δ) was purchased from Euroscarf. Plasmid vectors for overexpression in yeast cells were kindly provided by Dr. Ralph Panstruga and Dr. LE. Somssich. To generate pMet-AIFAH1 and pMet-AIFAH2, AIFAH1 and AIFAH2 open reading frames with their stop codons were combined with the destination vector pMet-Cub-Ura3 by the Gateway system, respectively. The construction of pMet-ScFAH1 was described in our previous work (Nagano et al., 2009). The plasmid DNA was introduced into yeast (Saccharomyces cerevisiae) (Curtis and Grossniklaus, 2003), by the Gateway system. The resulting plasmid was transferred to S. cerevisiae by the oral-dip method (Bechtold and Pelletier, 1998). Growth of yeast was performed in Edinburgh’s medium (EM) supplemented with 2% glucose or 2% raffinose, and 0.67% yeast nitrogen base (YNB), or 0.67% yeast nitrogen base, without amino acids (YNB). YNB without amino acids was supplemented with 5% glucose.

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Plant Lipid Extraction

Total lipids including sphingolipids were extracted from whole rosettes of Arabidopsis (300 mg fresh weight) according to the method described by Markham et al. (2006). Total other lipids mainly including nonhydroxyl fatty acids were extracted from rosettes of Arabidopsis (1 g fresh weight) according to the method described by Bligh and Dyer (1959).
Fatty Acid Analysis

2-HFA methyl esters with 13kDa methyl ester as an internal standard, or non-HFA methyl esters with margaric acid (17:0) methyl ester as an internal standard were prepared and separated by silica gel TLC using n-hexane/diethyl ether (4:1, vol/vol) as the mobile phase as described by Imai et al. (2000).

Derivatized 2-HFA samples were analyzed by GC (GC-2010, Shimadzu Scientific) as described by Nagano et al. (2009). Similarly, derivatized non-HFA samples were analyzed by GC with the injector set in split mode. The analytes were fractionated on a Restek RTX-5 column (5% diphenyl, 95% dimethyl polysiloxane; 0.25 mm i.d., 0.25 µm, 50 m, 30 m), and the injection port and the transfer line were maintained at 250°C. The column temperature was programmed from 100°C (2-min hold) to 280°C at 4°C/min and maintained at 280°C for another 30 min. The absolute values of non-HFA were computed like those of 2-HFA as described by Nagano et al. (2009).

Ion Leakage Measurement

Electrolyte leakage was monitored using an electrical conductivity meter (Horiba, B-173) at several time points, and expressed as a relative value against total ion leakage, which was measured in the autodissolved samples (Kawai-Yamaeda et al., 2009).

Measurement of Chlorophyll Content

Three leaf discs from 3-week-old plants treated with various chemicals were soaked overnight in 1 mL of dimethyl formamide at 4°C. The chlorophyll content was then determined spectrophotometrically as described previously (Porra et al., 1989).

Analysis of Various Metabolites

For measurements using CE-MS, metabolites were extracted using methanol-MilliQ water (1:1) as described by Saito et al. (2004). Leaf samples were obtained, immediately frozen in liquid nitrogen, and stored at −80°C until further analysis. The metabolites were extracted by rapid grinding of tissues in liquid nitrogen followed by immediate addition of 10 volumes of ice-cooled methanol (10 µL µg⁻¹ fresh weight) including 50 µM internal standards; Met sulfone was mixed by a vortex mixer for 1 min at 4°C and equal volume of Milli-Q water was added to the sample mixture. After centrifugation for 5 min at 15,000 × g, the supernatant was ultrafiltered through a 5-kD cutoff filter (Amicon). The filtrate was analyzed using the CE-MS method.

Separation and determination of metabolites were performed using the CE/MS system (Agilent Technologies). For the determination of anionic compounds, separations were carried out using fused-silica capillary (50 µm i.d. × 80 cm total length) filled with 50 µs ammonium acetate (pH 9.0) as the electrolyte according to the pressure-assisted CE-MS method (Harada et al., 2006). Cationic compounds were separated in an uncoated fused-silica capillary (50 µm i.d. × 100 cm total length) using 1 µM formic acid (pH 1.9) as the electrolyte (Soga and Heiger, 2000; Miyagi et al., 2010; Takahara et al., 2010).

Sequence data from this article can be found in the Arabidopsis Genome Initiative or GeneBank/EMBL databases under the following accession numbers: A1fAH1, A1g2c4770; A1fAH2, A1g2c2870; A1cB5-8, A1g2c3220; A1B1-1, A1g5c47120.

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LITERATURE CITED


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