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Essential role of the cytochrome P450 CYP4F22 in the production of acylceramide, the key lipid for skin permeability barrier formation

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Short title: Skin barrier formation by CYP4F22
Abstract

A skin permeability barrier is essential for terrestrial animals, and its impairment causes several cutaneous disorders such as ichthyosis and atopic dermatitis. Although acylceramide is an important lipid for the skin permeability barrier, details of its production have yet to be determined, leaving the molecular mechanism of skin permeability barrier formation unclear. Here we identified the cytochrome P450 gene $CYP4F22$ as the long-sought fatty acid $\omega$-hydroxylase gene required for acylceramide production. $CYP4F22$ has been identified as one of the autosomal recessive congenital ichthyosis-causative genes. Ichthyosis mutant proteins exhibited reduced enzyme activity, indicating correlation between activity and pathology. Furthermore, lipid analysis of an ichthyosis patient showed a drastic decrease in acylceramide production. We determined that $CYP4F22$ was a type I membrane protein that locates in the endoplasmic reticulum (ER), suggesting that the $\omega$-hydroxylation occurs on the cytoplasmic side of the ER. The preferred substrate of the $CYP4F22$ was $\geq$C28 fatty acids. In conclusion, our findings demonstrated that $CYP4F22$ is an ultra long-chain fatty acid $\omega$-hydroxylase responsible for acylceramide production and provides important new insights into the molecular mechanisms of skin permeability barrier formation. Furthermore, based on the results obtained here, we proposed a detailed reaction series for acylceramide production.

Keywords: acylceramide; ceramide; lipid; skin; sphingolipid
**Significance statement**

The sphingolipid backbone ceramide is the major lipid species in the stratum corneum and plays a pivotal function in skin permeability barrier formation. Acylceramide is an important epidermis-specific ceramide species. However, the details of acylceramide production including its synthetic genes, reactions and their orders, and intracellular site for production, have remained unclear. In the present study, we identified the cytochrome P450 CYP4F22 as the missing fatty acid ω-hydroxylase required for acylceramide synthesis. We also determined that CYP4F22 is a type I ER membrane protein and that its substrate is ultra long-chain fatty acids. Our findings provide important new insights into the molecular mechanisms of not only acylceramide production but also skin permeability barrier formation.
Introduction

A skin permeability barrier protects terrestrial animals from inside the body, penetration of external soluble materials, and infection by pathogenetic organisms. In the stratum corneum, the outermost cell layer of the epidermis, multiple lipid layers (lipid lamellae) play a pivotal function in barrier formation (Fig. S1) (1-3). Impairment of the skin permeability barrier leads to several cutaneous disorders, such as ichthyosis, atopic dermatitis, and infectious diseases.

The major components of the lipid lamellae are ceramide (the sphingolipid backbone), cholesterol, and free fatty acid (FA). In most tissues, ceramide consists of a long-chain base (LCB; usually sphingosine) and an amide-linked FA with a chain length of 16-24 (C16-C24) (4, 5). On the other hand, ceramide species in the epidermis are strikingly unique (Fig. S2A). For example, epidermal ceramides contain specialized LCBs (phytosphingosine and 6-hydroxysphingosine) and/or FAs with α- or ω-hydroxylation (1-3). In addition, substantial amounts of epidermal ceramides have ultra long-chain (ULC) FAs (ULCFAs) with ≥C26 (4, 5). The most unique epidermal ceramides are acylceramides having C28-C36 ULCFAs, which are ω-hydroxylated and esterified with linoleic acid (EOS in Fig. S1; EODS, EOS, EOP, and EOH in Fig. S2A) (1-3, 6, 7). These characteristic molecules may be important to increase the hydrophobicity of lipid lamellae and/or to stabilize the multiple lipid layers. Linoleic acid is one of the essential FAs, and its deficiency causes ichthyosis symptoms due to a failure to form normal acylceramide (8). Ichthyosis is a cutaneous disorder accompanied by dry, thickened, and scaly skin; it is caused by a barrier abnormality. In atopic dermatitis patients, both total ceramide levels and the chain length of ceramides are decreased, and ceramide composition is also altered (9-11).
In addition to its essential function in the formation of lipid lamellae, acylceramide is also important as a precursor of protein-bound ceramide, which functions to connect lipid lamellae and corneocytes (Fig. S1) (12, 13). After removal of linoleic acid, the exposed ω-hydroxyl group of acylceramide is covalently bound to corneocyte proteins, forming a corneocyte lipid envelope. Acylceramides and protein-bound ceramides are important in epidermal barrier formation, and mutations in the genes involved in their synthesis, including CERS3, ALOX12B, and ALOXE3, can cause non-syndromic, autosomal recessive congenital ichthyosis (ARCI) (3, 14-16). The ceramide synthase CERS3 catalyzes the amide bond formation between an LCB base and ULCFA, producing ULC-ceramide, which is the precursor of acylceramide (Fig. S1 and Fig. S2B) (17). The 12(\(R\))-lipoxygenase ALOX12B and epidermal lipoxygenase-3 ALOXE3 are required for the formation of protein-bound ceramides (13, 18). Other ARCI genes include the ATP-binding cassette (ABC) transporter ABCA12, the transglutaminase TGM1, NIPAL4/ICHTHYIN, CYP4F22/FLJ39501, LIPN, and PNPLA1 (16, 19). The exact functions of NIPAL4, LIPN, and PNPLA1 are currently unclear. Causative genes of syndromic forms of ichthyosis also include a gene required for acylceramide synthesis: the FA elongase ELOVL4, which produces ULCFA-CoAs, the substrate of CERS3 (20).

Although acylceramide is essential for the epidermal barrier function, the mechanism behind acylceramide production is still poorly understood, leaving the molecular mechanisms behind epidermal barrier formation unclear. For example, acylceramide production requires ω-hydroxylation of the FA moiety of ceramide. However, the ω-hydroxylase responsible for this reaction has been heretofore unidentified (Fig. S1). Here, we identified the cytochrome P450 CYP4F22, also known as FLJ39501, as this missing FA ω-hydroxylase required for acylceramide production. CYP4F22 had been identified as one of the ARCI genes (21),
although its function in epidermal barrier formation remained unsolved. Our findings clearly demonstrate a relationship between ARCI pathology, acylceramide levels, and ω-hydroxylase activity.
Results

Identification of CYP4F22 as the FA ω-Hydroxylase Required for ω-Hydroxyseramide Production

Although researchers have long known ω-hydroxylation is essential for acylceramide formation, they have long puzzled over which gene is responsible for this reaction. To identify this gene, we first established a cell system that produced ULC-ceramides, a possible substrate of interest for ω-hydroxylase, since most cells cannot produce such extremely long ceramides. Human embryonic kidney (HEK) 293T cells overproducing the FA elongase ELOVL4 and/or the ceramide synthase CERS3 were labeled with [3H]sphingosine, and the chain lengths of ceramides were determined by reverse-phase thin-layer chromatography (TLC). Although overexpression of either ELOVL4 or CERS3 alone did not result in the production of ULC-ceramides, their co-overproduction caused generation of ULC-ceramides with ≥C26 (Fig. 1A). They migrated slower than long-chain (LC; C16-C20) ceramides and very long-chain (VLC; C22-C24) ceramides on reverse-phase TLC. Production of ULC-ceramides with chain length up to C36 was also confirmed by liquid chromatography-mass spectrometry (LC-MS) analysis (Fig. S3). When labeled lipids were separated by normal-phase TLC, ULC-ceramides were detected as a band at the adjacent, upper position of VLC-ceramides (Fig. 1B).

It has been reported that the cytochrome P450 (CYP) inhibitor aminobenzotriazole inhibits the generation of ω-hydroxyseramide in cultured human keratinocytes (22). In humans, 57 CYP genes exist, and mammalian CYP genes are classified into 18 families and 43 subfamilies. Some CYP4F members are implicated in the ω-hydroxylation of long-chain FAs (23, 24), raising the possibility that certain CYP4F subfamily members are responsible for ω-hydroxylation of ULCFAs in acylceramide formation. To test this possibility, we
cloned all of the human CYP4F subfamily genes (CYP4F2, 3A, 3B, 8, 11, 12, and 22), and each was expressed as an N-terminally 3xFLAG-tagged protein in HEK 293T cells overproducing 3xFLAG-ELOVL4 and 3xFLAG-CERS3. All CYP4F subfamily proteins were expressed at similar levels (Fig. 2A). Among the CYP4F subfamily members, only CYP4F22 caused the disappearance of ULC-ceramide, which was concomitant with the production of a new band at the position of ω-hydroxyceramide (Fig. 2B and Fig. S4). LC-MS analysis determined that this band indeed represented ω-hydroxyceramides with C28-C36 (Fig. S5). Thus, CYP4F22 is the ω-hydroxylase required for ω-hydroxyceramide production.

**Correlation between CYP4F22 Activity and Ichthyosis Pathology**

Although CYP4F22 gene has been identified as one of the ARCI-causative genes (21, 25), its function in epidermal barrier formation has remained unsolved. Five missense mutations, which all cause amino acid substitutions (F59L, R243H, R372W, H435Y, and H436D), have been found in the CYP4F22 of ichthyosis patients. To examine their role in ichthyosis pathology, we introduced these mutations into CYP4F22 and examined the ω-hydroxylase activity of the resultant mutant proteins. All mutant proteins were expressed at equivalent levels to the wild type protein (Fig. 3A), and indeed all ω-hydroxylase activity of the mutant CYP4F22 proteins decreased to 4-20% of wild type protein activity (Fig. 3B). Therefore, protein activity and ichthyosis pathology were nicely correlated.

That being noted, ichthyosis due to CYP4F22 mutation is quite rare. In fact, a mere ~20 patients have been reported in Mediterranean populations (21), and only a single patient in Japan (25). The Japanese patient has compound heterozygous CYP4F22 mutations: one is a point mutation (c.728G>A) causing amino acid substitution (p.R243H; R243H) and the other
is a deletion (c.1138delG) causing a frame shift (p.D380TfsX2; D380TfsX2), in which Asp380 is substituted to Thr followed by a stop codon (25). The mutant R243H exhibited the decreased activity as described above (Fig. 3B). We also introduced the c.1138delG mutation into CYP4F22 and examined the ω-hydroxylase activity of its truncated protein product. D380TfsX2 (predicted molecular mass, 44.8 kDa) migrated faster than the wild type protein (62.0 kDa, Fig. 3A), and we found that it exhibited no activity (Fig. 3B).

To confirm our conclusion that CYP4F22 is involved in the production of acylceramide through ω-hydroxyceramide synthesis, we subjected the stratum corneum of the Japanese patient (Fig. 3C), as well as those of controls (her parents and a healthy volunteer), to LC-MS analysis and examined the levels of 11 major ceramide species. Although statistical analysis could not be performed due to the low number of samples, all three acylceramides containing sphingosine (EOS), 6-hydroxysphingosine (EOH), and phytosphingosine (EOP) were apparently decreased in the ARCI patient compared to the controls (to <1/10) (Fig. 3D and Table S1). Instead, the non-acylated ceramides NS (which are a combination of sphingosine and a non-hydroxylated FA) and AS (which are a combination of sphingosine and an α-hydroxylated FA) seemed to be increased (Table S1). These results confirm that ω-hydroxyceramide production by CYP4F22 is indeed required for acylceramide synthesis.

**CYP4F22 Is a Type I Endoplasmic Reticulum Membrane Protein**

We next examined the subcellular localization of CYP4F22 by subjecting 3xFLAG-tagged CYP4F22 to indirect immunofluorescence microscopy (Fig. 4A). 3xFLAG-CYP4F22 exhibited a reticular localization pattern and was co-localized with calnexin, HA-ELOVL4, and HA-CERS3, all of which are endoplasmic reticulum (ER) proteins, indicating CYP4F22 is localized in the ER. A hydropathy plot showed that CYP4F22 contains a highly
hydrophobic region at the N-terminus as well as some weak hydrophobic stretches (Fig. 4B). To reveal the membrane topology of CYP4F22, we introduced a N-glycosylation cassette, which is N-glycosylated when exposed to the lumen of the ER, into several positions of CYP4F22, the N-terminus, E85/K (between Glu85 and Lys86 residues), H155/R, A285/L, C361/R, D455/N, and R508/K. Among these fusion proteins, only CYP4F22 containing the N-glycosylation cassette at the N-terminus received glycosylation, as the molecular weight shift was observed upon treatment with endoglycosidase H (Fig. 4C). This result indicates that CYP4F22 spans the ER membrane once. Furthermore, it oriented its N-terminus to the ER lumen and the large, hydrophilic C-terminal domain containing the active site to the cytosolic side of the ER membrane. The same membrane topology was determined for other CYP members by the detection of N-terminal N-glycosylation (26, 27).

When the N-terminal hydrophobic region was removed, the resulting CYP4F22ΔN became distributed throughout the cytoplasm (Fig. 4D). CYP4F22ΔN was fractionated into both the soluble and membrane fractions by ultracentrifugation, in contrast to full-length CYP4F22, which was detected only in the membrane fraction (Fig. 4E). These results confirmed that CYP4F22 is a type I ER membrane protein. CYP4F22ΔN could not produce ω-hydroxyceramide (Fig. 4F), suggesting that anchoring to the ER membrane, where all of the reactions of acylceramide synthesis occur, is crucial for the CYP4F22 function.

**ULCFAs Are Substrates of CYP4F22**

It was still unclear whether CYP4F22 introduces a ω-hydroxyl group into ULCFAs before or after formation of ceramide. Therefore, we examined ω-hydroxy FA levels using LC-MS in the presence of the ceramide synthase inhibitor fumonisin B1. If ω-hydroxylation occurs prior to ceramide production, it was expected that free ω-hydroxy FA levels should be increased.
with fumonisin B₁ treatment. We found that ω-hydroxy FA levels with C26-C36 were significantly increased by addition of fumonisin B₁ (Fig. 5A), suggesting that ω-hydroxylation occurs prior to ceramide production. Thus, it is highly likely that the substrates of CYP4F22 are not ceramides but rather FAs, the same type of substrate as catalyzed by other CYP4F family members.

To confirm that the substrates of CYP4F22 are FAs, we performed an in vitro analysis utilizing yeast, which has no endogenous FA ω-hydroxylase activity. When C30:0 FA was used as a substrate, the total membrane fractions prepared from yeast bearing a vector plasmid exhibited FA ω-hydroxylase activity only at the background levels (Fig. 5B). On the other hand, the ectopic expression of human CYP4F22 resulted in the production of ω-hydroxy FA in an NADPH-dependent manner (Fig. 5B). The hydroxylation reactions by CYP generally require O₂ and NADPH. These results indicated that the substrates of CYP4F22 are indeed FAs.

Acylceramide specifically contains ULCFA (mostly C28-C36) as its FA component. It is possible that the substrate preference of CYP4F22 determines the FA chain length of acylceramides. To examine this possibility, we prepared HEK 293T cells producing different sets of ceramides with specific chain lengths by introducing particular combinations of ceramide synthase and FA elongase (ELOVL1 and CERS2, C22-C24 ceramides; ELOVL1 and CERS3, C26 ceramide; and ELOVL4 and CERS3, ≥C26 ceramides) (Fig. 5C) (28, 29). Mammals have six ceramide synthases (CERS1-6) and seven FA elongases (ELOVL1-7), and each exhibits characteristic substrate specificity (Fig. S2B) (2, 4, 5, 28). When CYP4F22 was expressed in these cells producing different ceramide species, ω-hydroxyceramides were produced in cells producing C26 ceramide (ELOVL1/CERS3 combination) and ≥C26 ceramides (ELOVL4/CERS3 combination) but not in cells producing C22-C24 ceramides.
(ELOVL1/CERS2 combination) (Fig. 5D). These results suggest that CYP4F22 can ω-hydroxylate ULCFAs (≥C26) but not VLCFAs (C22 and C24). The levels of ω-hydroxyceramide produced in ELOVL1/CERS3 cells were similar to those in ELOVL4/CERS3 cells, although the levels of non-hydroxyceramide substrates were much higher in ELOVL1/CERS3 cells. Thus, CYP4F22 exhibits especially high activity toward ULCFAs with ≥C28.
Discussion

Here we identified CYP4F22 as an ULCFA ω-hydroxylase involved in acylceramide production. Acylceramide is quite important for epidermal barrier formation, and impairment of its production such as by ELOVL4 and CERS3 mutations causes ichthyosis (15, 20). CYP4F22 was first identified as an ARCI-causative gene by Fischer and her co-workers (21). They proposed that CYP4F22 and most other ichthyosis-causative genes are involved in a metabolic pathway producing 12-lipoxygenase products (hepoxilins and trioxilins) from arachidonic acid by analogy to the 5-lipoxygenase pathway creating leukotrienes. In their scenario, arachidonic acid is first converted to 12(R)-hydroperoxycicosatetraenoic acid (12(R)-HPETE) by one of the ichthyosis gene products, ALOX12B, and then to 12(R)-hepoxilin A3 by another ichthyosis gene product, ALOXE3. 12(R)-Hepoxilin A3 is further converted to a triol compound, 12(R)-trioxilin A3. CYP4F22 was proposed to be involved in the metabolism of 12(R)-trioxilin A3 by converting 12(R)-trioxilin A3 to 20-hydroxy-12(R)-trioxilin A3. However, the exact roles of hepoxilins and trioxilins in epidermal barrier formation and/or keratinocyte differentiation are still unclear. Furthermore, recent findings demonstrated that ALOX12B and ALOXE3 are rather involved in the reactions necessary for conversion of acylceramide to protein-bound ceramide, i.e., peroxidation of the linoleate moiety and subsequent epoxyalcohol derivatization (13). In addition, ALOX12B was proven not to be involved in hepoxilin/trioxilin production (30). These findings suggested that while hepoxilin/trioxilin metabolism may not be relevant to the pathogenesis of ichthyosis, the impairment of acylceramide/protein-bound ceramide formation causes ichthyosis. Based on these recent findings, some researchers have predicted that CYP4F22 is involved in acylceramide generation (3, 31), but their suppositions have lacked experimental evidence — until now.
We determined the membrane topology of CYP4F22 (Fig. 4C), which indicates that the large C-terminal hydrophilic domain including catalytic residues is located in the cytosol. Therefore, ω-hydroxylation of the ULCFA portion of acylceramide must occur on the cytosolic side of the ER membrane. Based on this finding, we propose a working model for the process of acylceramide production in the ER membrane as follows (Fig. S6). Elongation of palmitoyl-CoA to ULCFA-CoA occurs on the cytoplasmic side of the ER membrane. Since lipids comprising the ER membrane are mostly C16 and C18, ULCFA (C28-C36) portions of ULCFA-CoAs should be bent in the cytosolic leaflet of the ER membrane (Fig. S6) or be penetrated into the luminal leaflet. Although the latter possibility cannot be excluded, we prefer the former possibility, since in the latter model ULCFA must ‘flip-flop’ at least three times in the ER membrane in the course of acylceramide production. Since the substrates of CYP4F22 are ULCFAs (Fig. 5), ULCFA-CoAs should be converted to ULCFAs before ω-hydroxylation. After ω-hydroxylation of ULCFAs by CYP4F22, the resulting ω-hydroxy-ULCFA is converted to ω-hydroxy-ULCFA-CoA by acyl-CoA synthetase. ACSVL4/FATP4 is the candidate acyl-CoA synthetase for this reaction, since Acsvl4 mutant mice exhibited skin barrier defects (32). CERS3 catalyzes formation of ω-hydroxyceramide from ω-hydroxy-ULCFA-CoA and LCB. An unknown acyltransferase then introduces linoleic acid into the ω-hydroxy group of ω-hydroxyceramide, generating acylceramide.

Our results presented here demonstrated for the first time that CYP4F22 is a bona fide ULCFA ω-hydroxylase required for acylceramide production. Our findings provide important new insights into the molecular mechanisms of skin permeability barrier formation. Future development of compounds that strengthen the skin permeability barrier by increasing acylceramide-synthetic proteins such as ELOVL4, CERS3, and CYP4F22 may be useful for treatment of cutaneous disorders including ichthyosis and atopic dermatitis.
Materials and Methods

Detailed materials and methods used for all procedures are available in SI Text.

[^3H]Sphingosine Labeling assay

Cells were labeled with 2 μCi [3-[^3H]sphingosine (20 Ci/mmol; PerkinElmer Life Sciences, Waltham, MA) for 4 h at 37 °C. Lipids were extracted as described previously (28, 33) and separated by normal-phase TLC and reverse-phase TLC.

Lipid Analysis Using LC-MS

FAs and ceramides prepared from cultured cells were analyzed by reversed-phase LC/MS using ultra-performance liquid chromatography (UPLC) coupled with electrospray ionization (ESI) tandem triple quadrupole MS (Xevo TQ-S; Waters, Milford, MA). FAs were analyzed after derivatization to N-(4-aminomethylphenyl)pyridinium (AMPP) amides using the AMP^+ Mass Spectrometry Kit (Cayman Chemical, Ann Arbor, MI). Stratum corneum ceramides were analyzed by reversed-phase LC/MS using Agilent 1100 Series LC/MSD SL system (Agilent Technologies, Santa Clara, CA).

Immunoblotting

Immunoblotting was performed as described previously (34, 35) using anti-FLAG M2 (1.85 μg/mL; Sigma, St. Louis, MO), anti-calnexin 4F10 (1 μg/mL; Medical & Biological Laboratories, Aichi, Japan), or anti-GAPDH 6C5 (1 μg/mL; Ambion, Life Technologies, Carlsbad, CA) antibody as a primary antibody and an HRP-conjugated anti-mouse IgG F(ab’)_2 fragment (1:7500 dilution; GE Healthcare Life Sciences, Buckinghamshire, UK) as a
secondary antibody. Labeling was detected using Pierce ECL Western Blotting Substrate (Thermo Fisher Scientific, Waltham, MA).
Footnotes

Y.O. designed and performed the experiments and analyzed the data. S.N., A.O., and N.K. performed the experiments. K.S. and M.A. prepared stratum corneum samples from controls and an ARCI patient, and A.N., H.T., U.Y., and J.I. analyzed their ceramide compositions. A.K. planned the project, designed the experiments, and wrote the manuscript.

The authors declare no conflict of interest.
Acknowledgments

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References


**Figure Legends**

**Fig. 1.** Overproduction of ELOVL4 and CERS3 causes generation of ULC-ceramides. HEK 293T cells were transfected with plasmids encoding 3xFLAG-ELOVL4 and 3xFLAG-CERS3, as indicated. Cells were labeled with [3H]sphingosine for 4 h at 37 °C. Lipids were extracted, separated by reverse-phase TLC (A) or normal-phase TLC (B), and detected by autoradiography. Cer, ceramide; GlcCer, glucosylceramide; SPH, sphingosine; SM, sphingomyelin.

**Fig. 2.** CYP4F22 is involved in ω-hydroxyceramide synthesis. HEK 293T cells were transfected with plasmids encoding 3xFLAG-ELOVL4, 3xFLAG-CERS3, and 3xFLAG-CYP4F subfamily members, as indicated. (A) Total lysates prepared from the transfected cells were separated by SDS-PAGE, followed by immunoblotting with anti-FLAG antibodies. (B) Cells were labeled with [3H]sphingosine for 4 h at 37 °C. Lipids were extracted, separated by normal-phase TLC, and detected by autoradiography. Cer, ceramide; GlcCer, glucosylceramide; SPH, sphingosine; SM, sphingomyelin; ω-OH, ω-hydroxy.

**Fig. 3.** Hydroxylase activity of CYP4F22 is impaired by ichthyosis-causing mutations. (A and B) HEK 293T cells were transfected with plasmids encoding 3xFLAG-ELOVL4, 3xFLAG-CERS3, and 3xFLAG-CYP4F22 (wild type (WT) or mutant), as indicated. (A) Total cell lysates prepared from the transfected cells were separated by SDS-PAGE and subjected to immunoblotting with anti-FLAG antibodies. (B) The transfected cells were labeled with [3H]sphingosine for 4 h at 37 °C. Extracted lipids were separated by normal-phase TLC and detected by autoradiography. Cer, ceramide; GlcCer, glucosylceramide; SPH, sphingosine; SM, sphingomyelin; ω-OH, ω-hydroxy. (C) Representative clinical feature of an ARCI
patient (at 2 years old). Leaf-like flakes presented on the extensor side of the left lower limb prior to tape stripping. (D) Acylceramide (EOS, EOH, and EOP) levels in stratum corneum of a control (WT/WT), carriers (WT/R243H, ichthyosis patient’s father; WT/D380T fs2X (fs2X), patient’s mother), and an ARCI patient (R243H/D380T fs2X) were measured by LC-MS.

Fig. 4. CYP4F22 is a type I ER membrane protein. (A, D, and E) HeLa cells were transfected with plasmids encoding HA-ELOVL4, HA-CERS3, 3xFLAG-CYP4F22, and 3xFLAG-CYP4F22ΔN (CYP4F22 lacking 54 N-terminal amino acids), as indicated. (A and D) Cells were subjected to indirect fluorescence microscopic observation. Calibration bar, 10 μm. (B) The hydrophilicity of CYP4F22 was analyzed by MacVector software (MacVector, Cary, NC) using the Kyte and Doolittle algorithm (window size, 15). (C) HEK 293T cells were transfected with pCE-puro 3xFLAG-CYP4F22, pCE-puro 3xFLAG-CYP4F22 (N-term, insertion of the N-glycosylation cassette to the N-terminus), pCE-puro 3xFLAG-CYP4F22 (E85/K, insertion of the cassette between Glu-85 and Lys-86), pCE-puro 3xFLAG-CYP4F22 (H155/R), pCE-puro 3xFLAG-CYP4F22 (A285/L), pCE-puro 3xFLAG-CYP4F22 (C361/R), pCE-puro 3xFLAG-CYP4F22 (D455/N), or pCE-puro 3xFLAG-CYP4F22 (R508/K). Lysates (3 μg) prepared from transfected cells were treated with or without endoglycosidase H (Endo H) and separated by SDS-PAGE, followed by immunoblotting with anti-FLAG antibodies. (E) Total cell lysates (10 μg) were centrifuged at 100,000 x g for 30 min at 4 ºC. The resulting supernatant (soluble fraction; S) and pellet (membrane fraction; M) were subjected to immunoblotting using anti-FLAG, anti-calnexin (membrane protein marker), or anti-GAPDH (soluble protein marker) antibodies. IB, immunoblotting. (F) HEK 293T cells transfected with plasmids encoding 3xFLAG-ELOVL4, 3xFLAG-CERS3, and 3xFLAG-
CYP4F22 (wild type (WT) or CYP4F22ΔN (ΔN)), as indicated, were labeled with [3H]sphingosine for 4 h at 37 °C. Lipids were extracted, separated by normal-phase TLC and detected by autoradiography. Cer, ceramide; GlcCer, glucosylceramide; SPH, sphingosine; SM, sphingomyelin; ω-OH, ω-hydroxy.

**Fig. 5.** CYP4F22 hydroxylates ULCFAs. (A) Keratinocytes were differentiated for seven days in the presence or absence of 10 μM fumonisin B1. Lipids were extracted, treated with an alkali and derivatized to AMPP amides. Derivatized FAs were analyzed by a Xevo TQ-S LC/MS system and quantified by MassLynx software. Statistically significant differences are indicated (*p<0.05, **p<0.01; t-test). hC26:0 stands for hydroxy C26:0 FA. (B) Total membrane fractions (50 μg) prepared from BY4741 bearing the pAK1017 (vector) or pNS29 (His6-Myc-3xFLAG-CYP4F22) plasmids were incubated with 10 μM C30:0 FA and 1 mM NADPH as indicated for 1 h at 37 °C. Lipids were extracted, derivatized to AMPP amides, and analyzed as in (A). The values represent the amount of each FA ω-hydroxylase activity relative to that of the vector/-NADPH sample. The value of the CYP4F22/+NADPH sample was statistically different from those of all other samples (**p<0.01; t-test). hC30:0 stands for hydroxy C30:0 FA. (C and D) HEK 293T cells were transfected with the plasmids encoding the indicated ELOVL (ELOVL1 or ELOVL4), CERS (CERS2 or CERS3), and CYP4F22. Transfected cells were labeled with [3H]sphingosine for 4 h at 37 °C. Extracted lipids were separated by reverse-phase TLC (C) or by normal-phase TLC (D) and detected by autoradiography. Cer, ceramide; GlcCer, glucosylceramide; SPH, sphingosine; SM, sphingomyelin; ω-OH, ω-hydroxy.
Figure 2

A

B
Figure 3

A

B

C

D

ELOVL4 + CERS3

ULC-Cer
VLC-Cer
LC-Cer
α-OH-Cer
Ω-Cer
Sph
SM
CYP4F22: WT
FSL
R243H
H438Y
H435K
H352D

ELOVL4 + CERS3

WT/WT
WT/R243H
WT/ΔS2
R243H/ΔS2

α-Phosphorylation (ng/mg protein)

0.7
0.6
0.5
0.4
0.3
0.2
0.1
0

ECS
EOC
EOP
Figure 4

A 3xFLAG-CYP4F22

Cainexin Mering

3xFLAG-CYP4F22 HA-ELOVL4 Mering

3xFLAG-CYP4F22 HA-CERS3 Mering

B

Hydropathy

3.0

2.0

1.0

0.0

-1.0

-2.0

100

200

300

400

500

Amino acid number

C

Insertion: N-term E85K H155R A286L C361R D458N R508K

Endo H: kDa

75

50

D

CYP4F22 WT CYP4F22 ΔN

E

WT ΔN

IB: FLAG

50

100

75

37

IB: Cainexin

IB: GAPDH

F

ULC-Cer

VLC-Cer

LC-Cer

α-OH-Cer

GloCer

ELOVL4 + CERS3

WT ΔN
Figure 5

A

B

C

D

Graph A shows the levels of fatty acids in different samples with and without Fumonisin B₁. Graph B illustrates the PA₂-hydroxylase activity in various conditions. Graph C presents a gel showing the presence of different ceramide species. Graph D displays a gel with additional annotations indicating the presence of various ceramide species and other lipid classes.
Figure S1
Figure S2

A

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<th>LCBs</th>
<th>FAs</th>
<th>Non-hydroxy FA [N]</th>
<th>α-hydroxy FA [A]</th>
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<td>ADS</td>
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<td>AH</td>
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</tr>
</tbody>
</table>

B

Diagram showing different ceramide structures and their conversion pathways with arrows indicating elongation, ceramide synthesis, and desaturation.

Legend:
- Arrows with two heads indicate elongation.
- Arrows with one head indicate ceramide synthesis.
- Arrows with no heads indicate desaturation.
Figure S3

A

B

C
Figure S5

A

B

C