In a three-dimensional reconstructed human epidermis filaggrin-2 is essential for proper cornification

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Atopic dermatitis is a chronic inflammatory skin disease with defects in the epidermal barrier. In a cohort of African-American children, a \( FLG2 \) nonsense mutation has been associated with the disease. In the epidermis of European patients, the expression of filaggrin-2, the filaggrin-related protein encoded by \( FLG2 \), is decreased. To describe the function of filaggrin-2 and evaluate the impact of its deficiency, its expression was downregulated using lentivirus-mediated shRNA interference in a three-dimensional reconstructed human epidermis (RHE) model. This resulted in parakeratosis and a compact \( stratum corneum \), presence of abnormal vesicles inside the corneocytes, increased \( \mathrm{pH} \) and reduced amounts of free amino acids at the RHE surface, leading to increased sensitivity to UVB radiation. The expression of differentiation markers was slightly modified. However, we observed reduced proteolytic processing of cornedesmosin, hornerin and filaggrin in parallel with reduced amounts of caspase-14 and bleomycin hydrolase. Our data demonstrated that filaggrin-2 is important for a proper cornification and a functional \( stratum corneum \). Its downregulation in atopic patients may be involved in the disease-associated epidermis impairment.

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Atopic dermatitis (AD; OMIM #603165), also known as atopic eczema, is a very common inflammatory skin disease.\(^1\,^2\) It is the result of complex interactions between genetic and environmental factors. The most robust and widely replicated genetic risk factor for the disease corresponds to nonsense mutations of the gene \( FLG \).\(^3\,^4\) This gene encodes filaggrin, an \( S100 \) fused-type protein essential for the epidermal barrier functions.\(^5\) Filaggrin deficiency is responsible for decreased amounts of free amino acids in the \( stratum corneum \) (SC),\(^6\) abnormal keratinocyte differentiation,\(^6,^{10} \) epidermal barrier defects and enhanced percutaneous sensitization.\(^6,^{7,9} \) All characteristics of the atopic skin. However, a significant number of Asian and European patients with AD do not display any of the numerous known \( FLG \) mutations,\(^3,4\) in particular in the South of Europe.\(^11,^{12} \) In addition, \( FLG \) nonsense mutations have not been detected in Ethiopian and South African populations,\(^13,^{14} \) and are not associated with AD in African-American patients.\(^15,^{16} \) Hence, the defect in epidermal barrier functions of patients without \( FLG \) mutations, including those of African ancestry, may be associated with other inherited/acquired abnormalities that compromise keratinocyte differentiation.\(^17\)

Recently, mutations in the \( FLG2 \) gene, in particular a nonsense mutation, were shown to be associated with persistent AD in a cohort of 60 US patients of African ancestry.\(^18\) \( FLG2 \) encodes filaggrin-2, another \( S100 \) fused-type protein.\(^19\) Filaggrin-2 is very similar to filaggrin in terms of protein structure, amino-acid composition, pattern of expression and of biochemical properties. It is synthetized by granular keratinocytes as a large precursor consisting of 23 homologous repeats and a \( S100 \)-homologous N-terminal domain. Filaggrin-2 and filaggrin are colocalized in keratohyalin granules in granular keratinocytes, and in the cytoplasmic matrix of the lower corneocytes.\(^20,^{21} \) In the upper SC, filaggrin-2 is deiminated and degraded by calpain 1.\(^22\) The role of filaggrin-2 in the SC remains to be discovered, but it may be similar to that of filaggrin.\(^22\) In addition, \( FLG2 \) is one of the genes that are the most downregulated after cholesterol depletion of keratinocytes, an experimental model of AD.\(^23\) Importantly, we and others have shown that filaggrin-2 expression is reduced, probably by pro-inflammatory cytokines, in the epidermis of European patients.\(^12,^{24} \)

To analyze the function of filaggrin-2 and understand the effect of its deficiency in AD in an immunological cell-free context, we downregulated its expression with shRNA technology in reconstructed human epidermis (RHE). This technology has previously been used with success to demonstrate the importance of filaggrin in the human epidermis.\(^6\) The research focused particularly on epidermal differentiation, SC properties and permeability barrier.

**Results**

Filaggrin-2 knockdown in RHE leads to a thinner epidermis, parakeratosis, a compact SC and the presence of abnormal vesicles inside corneocytes. Filaggrin-2 expression knockdown in RHE was achieved using RNA interference in normal human primary keratinocytes that demonstrate the importance of filaggrin in the human epidermis.\(^6\) The research focused particularly on epidermal differentiation, SC properties and permeability barrier.

**Abbreviations:** AD, atopic dermatitis; K, keratin; RHE, reconstructed human epidermis; RT-qPCR, reverse transcription-quantitative PCR; SC, \( stratum corneum \)

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differentiated at the air–liquid interface for 10 days, as previously described.6,25 This was performed with keratinocytes obtained from three different donors. Two shRNAs targeting filaggrin-2 (shFLG2a and shFLG2b) were used separately.

After 10 days of culture at the air–liquid interface of either normal keratinocytes or keratinocytes transduced with a control shRNA that does not target any known human gene, morphologically differentiated RHEs were obtained. Proper expression of differentiation markers and a functionally competent SC were achieved, as reported before.6 In particular, filaggrin and filaggrin-2 were detected in the stratum granulosum with a granular pattern and in the SC (Supplementary Figure S1a and Figure 1c). Similar locations are known in the native epidermis, where the two proteins are located in the keratohyalin granules and the corneocyte matrix.19–21 The RHEs produced with keratinocytes transduced with either the control shRNA (shc-RHE) or shFLG2a (shFLG2-RHE) were analyzed by reverse transcription-quantitative PCR (RT-qPCR) and western blotting with a polyclonal antibody directed to filaggrin-2 (Figures 1a and b).

In shFLG2-RHE, filaggrin-2 mRNA and protein amounts were decreased by 85% and 78%, respectively, compared with shc-RHE. Although similar but not identical levels of filaggrin-2 expression were observed in the RHEs produced with keratinocytes from the three different donors (Supplementary Figures S1b and c), the efficacy of down-regulation was the same. Immunofluorescence labeling of RHE sections showed the expression of filaggrin-2 in the granular layers and SC. Confirming the efficacy of down-regulation, the staining of shFLG2-RHE was markedly reduced (Figure 1c). Similar results were obtained with shFLG2b (illustrated in Supplementary Figure S2).

At a morphological level, shFLG2-RHE was thinner than shc-RHE with a reduced thickness of the living keratinocyte layers, as shown by hematoxylin–eosin staining (Figures 2a and b). The SC thickness was identical, but parakeratosis was evident in the shFLG2-RHE. To try to understand this effect on epidermis thickness, the keratinocyte proliferation rate was analyzed. Before epidermal reconstruction, no modification was observed. However, in RHE at day 10, a significant increase in Ki67 staining was observed (Supplementary Figure S3). This indicated that the reduced thickness was not due to reduced proliferation of the shFLG2-treated keratinocytes.

Transmission electron microscopy analysis of shc- and shFLG2-RHE (Figure 2c) showed similar keratinocyte organization and structures, including desmosomes, keratohyalin granules and corneodesmosomes. However, at the lower magnification, parakeratosis of shFLG2-RHE was confirmed and the SC appeared denser without clear extracellular spaces. At the highest magnification, abnormal material was visible in the extracellular spaces, and the corneocytes displayed unusual, large, intracellular vesicles.

The amino-acid content and pH of the upper SC are disturbed in filaggrin-2-deficient RHE. To test for a possible effect of filaggrin-2 knockdown on SC permeability, we used the Lucifer yellow assay.26 At day 10, the hydrophilic fluorescent dye was applied to the external surface of shc- and shFLG2-RHE. As observed by fluorescence microscopy (Figure 3a), the green dye was retained in the SC of both control and filaggrin-2-deficient RHEs. In addition, Lucifer yellow was hardly detectable in the culture medium (Figure 3b). In contrast, large amounts of the dye were detected when RHEs were analyzed at day 4 as a control. This indicated that the outside-in permeability was not affected by the deficiency in filaggrin-2.

Similar to filaggrin, filaggrin-2 has been suspected to be involved in the acidification of the SC and in the formation of natural moisturizing factor. Therefore, we measured the pH at the external surface of the shc- and shFLG2-RHE (Figure 3c). Filaggrin-2 inhibition induced a statistically significant pH increase (mean value from 6.38 ± 0.25 to 7.1 ± 0.15; P = 0.0075). Then, we quantified the contents of the urocanic acid and pyrrolidine carboxylic acid in the RHE. HPLC analysis of RHE lysates (Figure 3d) revealed a statistically significant urocanic acid decrease in filaggrin-2-knockdown RHE as compared with the control samples (from 5.90 to 4.80 mmol/ml). Concerning the pyrrolidine carboxylic acid concentration, a similar trend was observed (from 47.66 to 40.68 mmol/ml) but the decrease did
Filaggrin-2 deficiency alters the epidermal differentiation program. In order to test for a possible effect on keratinocyte differentiation, the expression of a panel of genes differentially expressed in the various layers of the epidermis was explored at the mRNA level by quantitative real-time PCR (Figure 4a). As compared with shc-RHE samples, mRNA levels of loricrin, involucrin and E-cadherin were slightly reduced in filaggrin-2-deficient RHE. The levels of mRNAs encoding keratin (K)14, K10, desmoglein 1, desmocollin 1 and claudin 1 were not modified. To determine whether the modifications at the mRNA level translated to the protein level, western blotting was performed and the detected bands were quantified (Figure 4b). Compared with controls, the detection of loricrin was decreased in shFLG2-RHE. Interestingly, the immunodetected claudin 1 amount was increased by a factor of 3. The expression of K14, K10, involucrin, desmoglein 1/2, desmocollin 1 and E-cadherin was not modified.

It has been suggested that the expressions of filaggrin-2, filaggrin and hornerin are coregulated. Therefore, we investigated the expression of the last two S100 fused-type proteins (Figure 5). No changes were detected at the mRNA level, confirming the specificity of the shRNAs used (Figure 5a). When shc- and shFLG2-RHE extracts were analyzed by western blotting, different patterns of immunodetection were

![Figure 2](image_url)
observed (Figure 5b). Profilaggrin was shown to accumulate whereas filaggrin monomers were not detected and intermediates decreased by 52%. Similar results were obtained concerning hornerin with an increase in the detection of the entire protein (prohornerin) and a decrease of 60% in the smaller bands. To test whether the proteolytic processing of other epidermal proteins was affected by the knockdown of filaggrin-2, we analyzed corneodesmosin in the same way (Figures 5a and b, lower part). Whereas the corneodesmosin mRNA amounts were similar, the entire 55 kDa protein increased and the 40 kDa processed form decreased by 90%.

When indirect immunofluorescence analysis was performed (Figure 5c), the labeling patterns obtained with the anti-filaggrin and anti-hornerin antibodies were modified. Filaggrin and hornerin were not detected in the SC of shFLG2-RHE but only in the stratum granulosum, with a granular pattern suggesting

Figure 3  Filaggrin-2 knockdown leads to impaired stratum corneum properties. (a and b) A solution of Lucifer yellow was applied to the shc- and shFLG2-RHEs. After 6 h of incubation, location of the dye was investigated using a fluorescence microscope (a) and dye concentration in the culture medium was quantified (b). As a control, a normal RHE at day 4 was used (RHE-D4). (c) The pH was measured at the surface of the RHEs. Each indicated value corresponds to the mean pH of a different RHE (n=5 shc-RHEs and n=6 shFLG2-RHEs). The RHEs were produced with keratinocytes from two different donors. (d) Pyrrolidone carboxylic acid (PCA) and urocanic acid (UCA) amounts were quantified in lysates of shc-RHEs (n=8) and shFLG2-RHEs (n=9). RHEs were produced with keratinocytes from the same two different donors. (e) Active caspase-3 (red staining; arrows) was detected without and with UVB irradiation, as indicated. Please note the nonspecific labeling of the polycarbonate membrane (under the thin line). (f) The number of active caspase-3-positive cells was quantified and is indicated per RHE length unit. Differences in pH, in amino-acid amounts and in number of cells were analyzed using Student’s t-test. Only statistically significant differences are indicated.

Figure 4  Filaggrin-2 deficiency affects keratinocyte differentiation. (a) Expression of keratin (K)14 and differentiation-associated genes was analyzed by qRT-PCR in shc-RHEs (black bars) and shFLG2-RHEs (gray bars). The expression of the TATA box-binding protein gene was used for normalization. *P=0.041; **P=0.005; ***P=0.0036. (b) Expression of the corresponding proteins was analyzed by western blotting, quantified and normalized to glyceraldehyde 3-phosphate dehydrogenase (GAPDH) immunoreactivity. **P=0.0039; ***P=5×10−4. The mRNA and protein amounts corresponding to shc-RHE were arbitrarily set at 100. Mean values and S.D. (error bars) were calculated from independent experiments performed with keratinocytes from three different donors, each being duplicated (n=6). Data were compared using Student’s t-test. Only statistically significant differences are indicated. LOR, loricrin; IVL, involucrin; DSG1, desmoglein 1; DSC1, desmocollin 1; CLDN1, Claudin 1; CDH1, E-cadherin.
an accumulation of their respective proforms in the keratohyalin granules. The distribution of corneodesmosin was not altered.

Procaspase-14 and bleomycin hydrolase expression is reduced in shFLG2-RHE. Several proteases are known to be involved in the processing of filaggrin, hornerin and corneodesmosin, including caspase-14, bleomycin hydrolase, calpain 1 and kallikrein 7. To study the protease deficiency in shFLG2-RHE, the expression of these enzymes was analyzed by western blotting (Figure 6a). No differences were observed for calpain 1 or kallikrein 7. Procaspase-14 and bleomycin hydrolase protein levels were reduced by 81% and 69%, respectively, in shFLG2-RHE compared with shc-RHE (Figure 6a). In the shFLG2-RHE, in addition to the downregulation of procaspase-14, a reduction was observed in the processing of the proform to the active form of the protease (mean factor of $1.3 \pm 0.6; n=4$). To determine whether the observed modifications of enzyme expression were because of changes in the corresponding mRNA levels, a quantitative real-time PCR analysis was performed (Figure 6b). Compared with control RHE samples, mRNA levels of caspase-14 and bleomycin hydrolase were significantly reduced in filaggrin-2-deficient RHE, whereas mRNA levels of calpain and kallikrein 7 were not modified.

Discussion

Since the first description of the expression of FLG2 in differentiated keratinocytes, little has been done to understand the role of the encoded protein, filaggrin-2. However, the close relationships between filaggrin and filaggrin-2 suggested that the two proteins could play similar roles in the epidermal barrier function. The recent descriptions of an association between FLG2 nonsense mutations and disease persistence in AD patients of African ancestry, and of a reduced expression of the protein in the epidermis of Caucasian patients, prompted us to look for filaggrin-2 function using RHE as an experimental model. We report here that filaggrin-2 knockdown in RHE led to alterations in the keratinocyte differentiation program. This culminated in changes of SC morphology and functions. More particularly, we observed changes in the amounts of loricrin and claudin 1, defects in the proteolytic processing of filaggrin, hornerin and corneodesmosin, and a compact SC with parakeratosis, abnormal intracellular vesicles, a higher external surface pH...
Figure 6 Expression of proteases is decreased in the absence of filaggrin-2. (a) Expression of caspase-14 (CASP14), bleomycin hydrolase (BLMH), calpain 1 (CAPN1) and kallikrein-7 (KLK7) in shc- and shFLG2-RHE was analyzed by western blotting and quantified. Act C14, active processed form of CASP14. **P = 0.0094; ***P = 7.8 × 10^-10; ****P = 8.09 × 10^-5; ***P = 1.09 × 10^-5. The mRNA and protein amounts corresponding to shc-RHE were arbitrarily set at 100. The error bars correspond to the S.D. calculated from independent experiments performed with keratinocytes from three different donors, one being duplicated for qRT-PCR (n = 4) and each being duplicated for western blotting (n = 6). Data were analyzed using Student’s t-test. Only statistically significant differences are indicated.

and decreased urocanic acid content. These data demonstrate the importance of filaggrin-2 in cornification and suggest its implication in AD pathophysiology.

Filaggrin-2 is composed of an NH2-terminal domain homologous to S100 proteins and by two central domains formed by repeated subunits. Because (1) the amino-acid composition of filaggrin-2 B-type repeats is similar to that of filaggrin, (2) filagrin and filaggrin-2 B-type repeats are concomitantly degraded in the upper SC and (3) both proteins are substrates of calpain 1, it has been suggested that the breakdown of filaggrin-2 into free amino acids contributes to the production of natural moisturizing factor components, control of the pH at the SC surface and photoprotection of the epidermis.20,21 Supporting these hypotheses, the SC pH was increased in filaggrin-2 knockdown RHE as compared with control RHE, and the urocanic acid and pyrrolidone 5-carboxylic acid contents were decreased. The decrease was not so pronounced as by the FLG loss-of-function mutations (i.e., one mutation results in ∼50% reduction) but is consistent with their relative contribution to the risk for AD. In addition, we observed an increased UVB-induced apoptosis of keratinocytes, urocanic acid being a major chromophore in the skin protecting from UVB radiation.9

Many features indicated that the keratinocyte differentiation program is perturbed when filaggrin-2 expression is downregulated. Such perturbations include a reduced expression of loricrin at both the mRNA and protein levels and an apparent stabilization of claudin 1. Many of them suggested premature cornification: a reduced thickness of spinous/granular layers without reduction of the keratinocyte proliferation rate, persistence of nuclei (parakeratinosis) and abnormal vesicles in the corneocyte matrix and reduced proteolytic processing of the late differentiation proteins (hornerin, profilaggrin and corneodesmosin). An increased epidermal turnover has been suggested to induce incomplete removal of organelles because the cellular proteolytic machinery necessary for protein/nucleic acid degradation is overfed.29 In an alternative explanation, the expression of procaspase-14, the precursor form, and of the processed form of caspase-14, the active form of the protease, was highly reduced in shFLG2-RHE. Caspase-14 deficiency has been reported to promote parakeratosis in skin equivalent models treated with siRNA and in knockout mice.27,28 Interestingly, parakeratosis has also been described in some AD patients,29,30 in particular in skin regions where caspase-14 was not detected.28

In order to understand the above-mentioned reduced proteolytic processing of profilaggrin and corneodesmosin, we investigated the expression of the respectively suspected proteases, calpain 1 and kallikrein 7. The expression of both proteases was similar in filaggrin-2 knockdown and control RHE. Therefore, alterations in their activity and/or localization could be supposed. As the pH was increased in the filaggrin-2 knockdown epidermis, and as kallikrein 7 has been reported to be more active at neutral than acidic pH, difference in the pH is probably not the explanation. Another possibility concerning kallikrein 7 is altered secretion. Loss of filaggrin-2 led to a compact SC with apparently abnormal extracellular spaces and the presence of intracellular large vesicles, suggestive of abnormal secretion of lamellar body content. It has been discussed how mutations in FLG, FATP4 and TMEM79 genes and increase in SC pH converge to produce defective lamellar body and/or to impair the secretion of these structures in patients with AD.17 Mutations in FLG2 and reduced expression of filaggrin-2 may well be involved.

When we compared the effects of filaggrin deficiency and those of filaggrin-2 deficiency in RHE, some similarities were observed (Figure 7 and Table 1), confirming that the proteins are related. This induces a certain complexity in the interpretation of the results. For example, are the reductions in the level of natural moisturizing factor components and in the expression of loricrin, bleomycin hydrolase and caspase-14 directly because of the downregulation of filaggrin-2 or are they a consequence of the reduced production of filaggrin monomer or both? The increase of claudin-1, a tight junction protein, suggests a compensatory mechanism to overcome SC defects. However, the functions of filaggrin and filaggrin-2 are probably not identical as differences were also noted. In particular, whereas filaggrin deficiency in RHEs and in a three-dimensional skin model leads to enhanced penetration of foreign molecules,6,31 filaggrin-2 knockdown appeared not to disturb outside-in SC permeability: Lucifer yellow was retained in the SC of shFLG2- as well as shc-RHE. Modification of the
SC superficial pH was only observed when FLG2 was downregulated. Hypogranulosis was only evidenced when filaggrin was downregulated, even if filaggrin-2 is also a component of keratohyalin granules, suggesting that the former is much more abundant than the second. Accordingly, the copy number of filaggrin mRNA per 10 ng of total RNA is $\sim 10^5$, and that of filaggrin-2 is between 6 and 30.

Some of the effects induced by filaggrin-2-deficiency reproduced some AD-related epidermis alterations (Table 1), as already mentioned, including parakeratosis, an apparently reduced degradation of cornodesmosome components including corneodesmosin, reduced amounts of bleomycin hydrolase, and natural moisturizing factor, altered profilaggrin processing, and the presence of vesicles inside corneocytes. Similarly, the AD-associated increase in the SC superficial pH could be because of the decrease in filaggrin-2 amount, as it has not been observed in filaggrin-null mice, filaggrin knockdown RHEs or patients with FLG nonsense mutations.

In conclusion, this study demonstrates that filaggrin-2 is essential for normal keratinocyte differentiation, and that filaggrin-2 downregulation may be responsible for reported risk to AD.

Table 1

| Filaggrin-2 knockdown | Filaggrin knockdown | AD |
|-----------------------|---------------------|----|
| Normal stratum granulosum | Hypogranulosis | Hypogranulosis |
| Parakeratosis and a compact stratum corneum | Thin stratum corneum | Some parakeratosis |
| Vesicles in corneocytes | No vesicle in corneocytes | Vesicles in corneocytes |
| Normal corneocyte matrix | Disturbed corneocyte matrix | Increased outside-in SC permeability |
| Normal outside-in SC permeability | Increased outside-in SC permeability | Reduced NMF level |
| Reduced NMF level | Reduced NMF level | Reduced NMF level |
| Increased pH | Unaffected pH | Increased pH |
| Decreased amounts of bleomycin hydrolase | Decreased amounts of bleomycin hydrolase | Decreased amounts of bleomycin hydrolase |
| Reduced proteolysis of filaggrin and hornerin precursors | Reduced expression of filagrin and hornerin | Reduced expression of filagrin and hornerin |
| Reduced degradation of corneodesmosin | Increased expression of corneodesmosin | Normal (pro)filaggrin proteolysis |
| Reduced expression of loricrin | Reduced expression of loricrin | Persistence of corneodesmosomes |
| Abnormal expression of tight junction proteins | Abnormal expression of tight junction proteins | Reduced expression of loricin |

Abbreviation: NMF, natural moisturizing factor

See "Pendaries et al" and "Oyoshi et al" and Gutman-Yassky et al and references in the text

SC superficial pH could be because of the decrease in filaggrin-2 amount, as it has not been observed in filaggrin-null mice, filaggrin knockdown RHEs or patients with FLG nonsense mutations.

In conclusion, this study demonstrates that filaggrin-2 is essential for normal keratinocyte differentiation, and that filaggrin-2 downregulation may be responsible for reported risk to AD.

Materials and Methods

shRNA lentiviral particles. For FLG2 knockdown in keratinocytes, we used MISSION pLKO.1-puro vector based lentiviral particles containing a purmycin resistance gene and a shRNA insert under the U6 promoter (Sigma-Aldrich, St. Louis, MO, USA). Two shRNAs targeting exon 3 of FLG2 gene (shFLG2) were tested (shFLG2a sequence: 5′-GATGARATCAAGCTGACTAT-3′, nucleotide position 301–321 on the mRNA sequence (NCBI reference NM_001014342.2); shFLG2b sequence: 5′-GATGARATCAAGCTGACTAT-3′, nucleotide position 301–321 on the mRNA sequence (NCBI reference NM_001014342.2));
shFLG2b sequence: 5′-CAGTGGAGAAAGGAAAGATT-3′, nucletide position 763–783. A non-target shRNA that does not target any known human gene (shc) was used as a control (5′-CCGGCAACAAGTAGAAGACGACCACTC-3′).

Keratinocyte culture and transduction. Primary normal human keratinocytes were obtained from abdominal dermolepctomy of three different healthy subjects who had given their informed consent. They were cultured and transduced with the lentivirus containing either shFLG2a, shFLG2b or shc at a multiplicity of infection of 10 in the presence of 4 μg/ml of proteamine sulfate, as described previously. After selection in the presence of puromycin, keratinocytes were used to produce RHE on polycarbonate filters as described previously. Briefly, cells were harvested by trypsinization, and 350,000 cells in ice-cold Epilife medium (Invitrogen Life Technologies) containing 1.5 mM calcium were seeded on polycarbonate culture inserts (area of 0.62 cm² with pores 0.4 μm in diameter; Merck Millipore, Bedford, MA, USA). After 24 h of incubation at 37°C in a humidified atmosphere containing 5% CO₂, cells were exposed to the air–liquid interface, and 50 μg/ml vitamin C (Sigma-Aldrich) and 10 ng/ml keratinocyte growth factor (Sigma-Aldrich) were added to the medium in the lower compartment. The medium was renewed every 2 days during the 10 days of air–liquid interface culture.

Immunostaining. At day 10, RHEs were fixed with 4% formaldehyde-containing buffer, and paraffin embedded. After deparaffinization and hydration, sections were blocked with PBS containing 2% bovine serum albumin and incubated with primary antibodies (Supplementary Table S1). After incubation with the corresponding secondary antibody (Alexa Fluor 555 anti-Rabbit IgG, Alexa Fluor 555 or 488 anti-Mouse IgG, Alexa Fluor 555 or 488 anti-Goat IgG, Invitrogen Life Technologies), nuclei were stained with DAPI (Sigma-Aldrich) and the slides were observed with a Nikon Eclipse 80i fluorescence microscope equipped with a Nikon DXM 1200 digital camera (Nikon, Tokyo, Japan).

Morphological analysis. Epidermal thickness was assessed using images of hematoxylin–eosin-stained sections of paraffin-embedded RHEs counting three fields per slide. Experiments were performed with keratinocytes from the three volunteers. Transmission electron microscopy analysis was carried out as previously described. Briefly, samples were fixed with 2.5% glutaraldehyde–formaldehyde solution, washed with Sorensen phosphate buffer, post-fixed with 1% OsO4, dehydrated and embedded in Embed 812 resin (Electron Microscopy Sciences, Hatfield, PA, USA). Ultrathin sections were mounted on 100-mesh collodion-coated copper grids and post-stained with 3% uranyl acetate and 8.5% lead citrate before observation with a Tecnai T20 (FEI, Netherlands) or a Tecnai T12 (FEI, Netherlands).

Western blotting. At day 10, RHEs were lysed in 100 μl of Laemmli buffer. Total epidermal proteins were separated on acrylamide gels and immunodetected with the primary antibodies and peroxidase-conjugated secondary antibodies (Goat anti-rabbit IgG-HRP and swine anti-goat IgG-HRP, SouthernBiotech, Birmingham, AL, USA; goat anti-mouse IgG-HRP, Bethyl Laboratories, Montgomery, TX, USA). Reaction products were detected by chemiluminescence with the ECL kit (Pierce, Thermo Scientific, Rockford, IL, USA). Quantity One 1-D Analysis Software (Bio-Rad Laboratories, Hercules, CA, USA) was used to quantify immunoreactive bands on western blot films after scanning. Signals were normalized to glyceraldehyde 3-phosphate dehydrogenase immunodetection.

Reverse transcription-quantitative PCR. Reverse transcription (RT) was performed as previously described. The primers used are listed in the Supplementary Table S2. Only Ct values between 17 and 26 were considered. Relative levels of gene expression among samples were determined with the ΔΔCt method. TATA box binding protein (TBP) gene expression was used as an endogenous control.

Determination of the pH at the RHE surface. At day 10, the pH at the RHE surface was determined with a pH electrode (Skin-pH Meter 905, Courage +Khazaka Electronic GmbH, Köln, Germany).

Determination of the urocanic acid and pyrrolidone carboxylic acid contents. At day 10, RHE specimens were lysed in 0.1 M KOH and amino acids were analyzed by HPLC as previously described.

Statistical analysis. Statistical differences were determined with Student’s t-test. A value of P<0.05 was considered statistically significant.

Conflict of interest
The authors declare no conflict of interest.

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Determination of the pH at the RHE surface. At day 10, the pH at the RHE surface was determined with a pH electrode (Skin-pH Meter 905, Courage +Khazaka Electronic GmbH, Köln, Germany).

Determination of the urocanic acid and pyrrolidone carboxylic acid contents. At day 10, RHE specimens were lysed in 0.1 M KOH and amino acids were analyzed by HPLC as previously described.

Statistical analysis. Statistical differences were determined with Student’s t-test. A value of P<0.05 was considered statistically significant.

Conflict of interest
The authors declare no conflict of interest.

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