Stratum corneum lipidomics analysis reveals altered ceramide profile in atopic dermatitis patients across body sites with correlated changes in skin microbiome

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Abstract

Background: Atopic dermatitis (AD) is driven by the interplay between a dysfunctional epidermal barrier and a skewed cutaneous immune dysregulation. As part of the complex skin barrier dysfunction, abnormalities in lipid organization and microbiome composition have been described. We set out to systematically investigate the composition of the stratum corneum lipidome, skin microbiome and skin physiology parameters at three different body sites in patients with AD and healthy volunteers.

Methods: We analysed tape strips from different body areas obtained from 10 adults with AD and 10 healthy volunteers matched for FLG mutation status for 361 skin lipid species using the Metabolon mass spectrometry platform. 16S rRNA data were available from all probands.

Results: Our study showed that the lipid composition differs significantly between body sites and between AD patients and healthy individuals. Ceramide species NS was significantly higher in AD patients compared to healthy volunteers and was also higher in AD patients with a FLG mutation compared to AD patients without a FLG mutation. The correlation analysis of skin lipid alterations with the microbiome showed that Staphylococcus colonization in AD is positively correlated with ceramide subspecies AS, ADS, NS and NDS.

Conclusion: This is the first study to reveal site-specific lipid alterations and correlations with the skin microbiome in AD.

KEYWORDS
Atopic dermatitis, Filaggrin, lipidome, metabolomics
1 | INTRODUCTION

Several pathophysiological mechanisms are thought to contribute to AD, but the two key drivers are a dysfunctional epidermal barrier and exaggerated type 2 T helper cell (TH2)-mediated immune responses. The two major epidermal barrier structures are the stratum corneum intercellular lipid lamellae and tight junction proteins in the nucleated epidermis. The function of the Stratum corneum critically depends on proper differentiation of keratinocytes and a coordinated activity of acid, lipid and enzyme constituents. In AD, epidermal barrier dysfunction is consistently observed in affected and unaffected skin and evidenced by elevated markers of epidermal permeability including elevated transepidermal water loss (TEWL), elevation of pH, increased permeability, reduced water retention and enhanced susceptibility to infection. At the molecular abundance, a reduced expression of epidermal structural proteins such as filaggrin, imbalances in protease-protease inhibitor interactions, a reduced expression of tight junction proteins and an altered skin microbiome configuration has been reported for both lesional and non-lesional skin. Further, several studies have also suggested that skin lipid metabolism, especially ceramide metabolism, is altered in AD. In particular, the chain length of ceramide-linked fatty acids was reported to be decreased, and it was shown that the lipid chain length reduction correlates with a compromised lipid envelope and thus an impaired barrier function. More recently, shifts in the epidermal lipid composition have been reported to correlate with increased Staphylococcus aureus colonization. Specifically, abundances of very long-chain ceramides (CER|EOH) C60 and CER|EOH|C68) were higher in AD patients without S. aureus colonization as compared to AD patients with S. aureus colonization. Alterations of single or multiple factors impacting barrier function can be mediated through primary mechanisms such as mutations in the gene encoding filaggrin (FLG) or indirectly through type 2 immune activity in the skin causing a secondary downregulation of skin barrier genes and stratum corneum lipids. It has been previously shown that knock-down of FLG in skin models leads to increased expression of secretory phospholipase A2, which converts phospholipids into fatty acids. In turn, free fatty acids accumulate and contribute to less ordered intercellular lipid lamellae and higher permeability in FLG knock-down skin constructs. However, whether filaggrin is indeed responsible for impaired lipid organization remains unclear, as several studies found no correlation between lipids and FLG mutations.

We here set out to systematically analyse the epidermal lipid composition at different body sites and in AD as compared to matched healthy individuals. We further took advantage of existing 16S data to evaluate its potential impact on the skin microbiome composition. To this end, we measured 361 skin lipid species at three different sites, namely the forehead (FH), the cubital fossa (CF) and the proximal lower forearm (LA) in 4 different groups matched for age, sex, presence/absence of AD and presence/absence of FLG haploinsufficiency.

2 | METHODS

2.1 | Study population

The study population has been described previously and consisted of four groups of German adults recruited through the POPGEN population-based cohort: (a) subjects with AD carrying a single FLG mutation (AD FLGmut), (b) subjects with AD without FLG mutations (AD FLGwt), (c) subjects with no history of chronic skin or allergic disorders carrying a single FLG mutation (controls FLGmut) and (d) subjects with no history of chronic skin or allergic disorders without FLG mutations (controls FLGwt). FLG mutation status was available from prior studies. Characteristics of the probands are reported in Table S1. The follow-up examination included a skin examination by an experienced dermatologist blinded to the genotype. AD was defined by the American Academy of Dermatology diagnostic criteria. The selected individuals per group were matched by age and sex, and selected AD patients had to display at least one unaffected antecubital crease. None of the participants had received systemic immunosuppressants or systemic antibiotics in the preceding 3 months. All participants were asked to avoid bathing and application of any topical agent 24 hours prior to the examination visit. All skin biophysical measurements were done in standardized environmental conditions (room temperature, 22 to 25°C; humidity abundances, 30% to 35%). Skin pH was measured with a Skin-pH-Meter® PH 905 (Courage & Khazaka Electronic, Cologne, Germany), and TEWL was measured with the Binx Aquaflex AF200 closed condenser chamber device (Binx Systems Ltd, London, UK) according to international guidelines. The mean of 3 separate measurements was used for analysis.

The study was approved by the ethical review board of the Medical Faculty of the University Kiel, Germany, and written informed consent was obtained from all study participants.

2.2 | Lipid analysis

A total of 348 ceramides (CER), 12 free fatty acids (FFA) and cholesterol sulphate were determined by SFC-MS/MS using the TrueMass® Stratum Corneum Metabolon Lipid Panel (Waters Corporation, Milford, MA/Sciex, Framingham, MA; Metabolon, Inc, Durham, NC) at three body sites (CF, FA and LA).

Four consecutive tape strips were taken using the D-squame pressure instrument from non-lesional skin sites, frozen and stored at −80°C prior analysis. We obtained a total of 60 samples consisting of four pooled D-Squame discs per subject and sampling site. All D-Squame discs were extracted in a batch together with four additional negative control samples using a polar and a non-polar organic solvent after addition of a known amount of surrogate standard solution (C16 ceramide-d31 ([S(C16)16:0]-d31), cis-10-heptadecenoic acid (FA17:1n7) and cholesteryl sulphate-d7). The organic extracts are combined and evaporated to dryness. The dried extract is reconstituted, and an aliquot is analysed on a Waters UPC2/Sciex.
2.3 | 16S rRNA gene data processing

Skin samples were subjected to 16S rRNA amplicon sequencing of the hypervariable regions V1 and V2 as described previously. In summary, bacterial genomic DNA was extracted using the PowerSoil DNA Isolation Kit (MoBio Laboratories, Carlsbad, Calif) and the hypervariable regions V1 and V2 of 16S rRNA gene were sequenced on the MiSeq platform (Illumina, San Diego, Calif). Stringent demultiplexing was carried out by allowing no mismatches in either index sequence, and a series of quality steps (dereplication, quality trimming, chimera filtering) were applied before resulting OTUs were classified into bacterial taxa.

2.4 | Statistical analysis

All data are represented as mean ± standard deviation unless otherwise indicated. To evaluate differences between groups, we used the Welch test on relative abundance of lipid species, log-transformed pH and log2-transformed TEWL values. Differences between sites were evaluated by the paired t test. The balance between proportions of lipid classes was calculated by the Shannon index implemented in the “vegan” (v2.4) package. Correlations of taxa of different abundances (eg phylum and genus) were calculated by the spearman correlation coefficient.

Overall correlation between abundances of lipid species and bacterial abundance across all samples was calculated by sparse canonical correlation analysis with feature selection using the R package “mixOmics” (v6.8.5).

All analyses have to be interpreted as exploratory, and presented P-values are unadjusted if not otherwise stated. If adjustment for multiple testing was applied, false discovery rate (fdr) for univariate analysis and the step-down resampling procedure (mvabund package) for multivariate analysis were used and respective P-values are indicated.

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In contrast, FFA composition was not significantly different between the 3 sites, apart from the saturated fatty acid 24:0, which significantly differed between the FH and the CF and the FH and the proximal LA (Figure 2B).

The distinct differences in CER lipid classes of the FH compared to the LA and the CF could further be shown by the differences in the Shannon index (Table S7). The CER diversity measured by the Shannon index is significantly reduced in AD patients compared to controls at the CF (P = .0021) and the proximal LA (P = .0185), but not at the FH (Table S7). This corresponds with an increased proportion of the CER subclasses NS, AS and ADS and a decreased proportion of CER subclasses NP, NH and EOS in AD patients (Figure 3) as previously reported.\cite{32}

### 3.3  Stratum corneum lipids are altered in AD patients

Increased total FFA abundance was mainly associated with a global upregulation of all FFAs, and no association was found with a significant upregulation of specific FFA proportions. We did not observe an association of AD with FFA proportions (Shannon entropy) or average FFA chain length (Figure S6). However, in line with previous reports at all 3 sites examined, the average CER chain length was lower in AD as compared to healthy individuals (P = .00049) (Figures S7-S9).

Detailed analysis of ceramide and FFA class abundance between patients with AD and healthy individuals showed that for ceramides, the abundance of NS class was significantly higher in AD patients, but no differences in FFAs were observed (Figure S10). Among the AD patients, no ceramide subclass was found to be significantly altered in FLG mutation carrier vs. non-mutation carrier (Figure S11, Figure S12A). However, abundance of most FFAs was higher in FLG mutation carrier than in non-mutation carrier, but only FFA 22.1 and 24.1 were significantly increased (Figure S12B).

### 3.4  SC lipid composition between body sites shows significant differences in abundance of ceramide subclasses and abundance of the saturated fatty acid 24:0

As the data showed that both arm sample sites are similar and in order to increasing the power of analysis, we pooled the data of both the

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**FIGURE 1** Heatmap of ceramides in healthy individuals and patients with AD. The coloured bar at the top indicates ceramide subclass composition. The coloured bar at the side indicates patients with AD with FLG mutation (red), patients with AD without FLG mutation (orange), healthy individuals without FLG mutation (black) and healthy individuals with FLG mutation (grey). Hierarchical clustering indicates that FLG is not a major component of variance across data.
LA and the CF to generate a forearm (FA) sample group. Differential analysis of the FH and the FA clearly shows that the proportion of CER[NS] and CER[NH] are regulated differently (Figure 4A). Comparison of healthy individuals and AD patients shows that the CER[NS] group is upregulated in the FH in both healthy and AD patients. The observed increase of ceramides in sun exposed areas is in line with previous findings that ceramide production is induced after sun exposure.[33]

The increase of CER[NS] in the FH compared to the FA is greater in healthy individuals than in AD patients, showing that in AD patients there is a reduced induction of ceramide production at the FH (Figure 4). We also detected a site-specific and AD-specific lipid dysregulation in the CER[NH] subclass. The CER[NH] abundance was lower in the FH as compared to the FA. Of note, healthy individuals showed a significantly greater reduction than AD patients (Table S8). These results show that different ceramide subclasses are not only differentially regulated between sites, but that site-specific regulation may differ in AD patients.

To assess whether the site-specific differences in the proportions of the CER subclasses is due to the regulation of all measured ceramides or whether the variance can be explained by a few specific ceramides, we looked at all ceramides in the CER[NS] and CER[NH] group and calculated the p-value, as well as the FDR value of their differential abundance between FH and FA (Figure 4B). Interestingly, only 5 ceramides in both ceramide subclass group were responsible for the site-specific differences in ceramide class proportions. Out of the CER[NS] group, S(C18)16:0, also known as N-(hexadecanoyl)-sphing-4-enine, was the ceramide with the lowest p-value.

**FIGURE 2** Ceramide and free fatty acid distribution in healthy controls at three different sites (A) ceramides: blue = α-hydroxy CERs, red = non-hydroxy CERs and brown/orange = esterified ω-hydroxyl CERs; proportion of CER[NP] is significantly higher in the cubital fossa compared to the forehead, and proportion of CER[AH] is significantly higher in the proximal lower forearm compared to the forehead. (B) Free Fatty Acids: Blue = saturated FFAs, red = MUFAs and brown/orange = DUFAs. Free fatty acid proportions are not significantly altered at the different sites with the exception of saturated fatty acid 24:0, which is significantly increased at the cubital fossa as well as the proximal lower forearm compared to the forehead.
S(C18)16:0 has been associated with AD before.\cite{15} In the CER[NH] group, H(C16)26:0 was the ceramide with the lowest p-value.

### 3.5 FLG deficiency is not associated with altered stratum corneum lipid composition

Analysis of the lipid class proportions with respect to the presence/absence of FLG haploinsufficiency revealed that lipid composition is FLG independent. Comparing all FLG mutation carriers vs. non-mutation carriers regardless of their AD status, we could not observe significant difference in either ceramide balance or CER subclass proportions at any body site (Table S9). These results are in line with previous reports,\cite{16} where no significant difference in lipid proportions due to FLG could be shown. Hierarchical clustering analysis also revealed that FLG mutation had no significant impact on sample variability nor on ceramide subclass composition at different sample sites (Figure 1 and Figure S11). There was also no association of FFA average chain length, relative frequencies of specific FFAs or FFA proportions with FLG (Figure S6). CER chain length was also not associated with FLG at all sample sites (Figures S7-S9).

### 3.6 Microbial colonization is shifted towards Staphylococcus in patients with atopic dermatitis

Analysis of the skin microbiome composition at the sites of lipid analysis by 16S sequencing showed a decreased abundance of Bacteroidetes and Actinobacteria and an increased abundance of Firmicutes in AD at all sites (Figure 5A). This is in accordance with previous findings where sampling of separate skin sites showed the same trend.\cite{23} A more detailed analysis revealed a decreased abundance of the genera Chryseobacterium, Propionibacterium and Kocuria and an increased abundance of the genera Staphylococcus and Lactobacillus (Figure 5A), indicating that in AD there is a shift in the bacterial community favouring Staphylococcus and Lactobacillus. A correlation analysis between bacterial and lipid abundances showed a strongly positive correlation of Staphylococcus and ceramide subclasses AS, ADS, NS and NDS in patients with AD, but not in healthy subjects (Table 1). In contrast, the abundances of Propionibacteria, Rothia, Actinomycices, Lactobacillus, Acinetobacter, Haemophilus and Neisseria were negatively correlated with the ceramide species AS, ADS, NS and NDS in patients with AD, but not in healthy subjects (Table 1). In healthy subjects, only Chryseobacteria showed a strong positive correlation with the ceramide species AS, ADS, NS and NDS. In healthy subjects, only Chryseobacteria showed a strong positive correlation with the ceramide species AH, AS and AP, while other bacterial genera showed only weak positive correlations (Table 1). These results indicate a correlation of ceramide species with an increased bacterial colonization with Staphylococcus and a decreased colonization with other bacterial genera in AD, but not in healthy subjects. Further, in AD patients, the abundances of Rothia, Actinomycices, Lactobacillus, Acinetobacter, Haemophilus and Neisseria showed a strongly positive correlation with abundances of saturated short-chain FFAs, such as FA16:0 and FA18:0. The short-chain FFAs FA16:0 and FA18:0 were also negatively correlated with Staphylococcus in AD patients, thus showing both a negative correlation with Staphylococcus in AD and a positive correlation with bacterial genera other than Staphylococcus in healthy volunteers.

### 4 DISCUSSION

Our systematic skin lipidome analysis indicates that while across body sites total ceramide, FFA and cholesterol sulphate abundances are increased in AD patients, the relative abundance of several ceramide subclasses (NH, NP, EOS, EOH and EOP) is reduced in AD patients. This might be surprising, as most studies have reported a decrease of ceramides in AD.\cite{34,35} However, decrease of ceramides in AD is due to the decrease of long-chain ceramides with a concomitant increase in short-chain ceramides.\cite{16,35} Out lipid panel did not...
cover long-chain ceramides (Figure S10), which explains our finding of increased total ceramide levels as only the increase in short-chain ceramides is detected. In line with previous findings, we also found an increase in the relative abundance of ceramide subclass NS\(^{[17,36]}\). Further, several ceramide species correlated with an increased colonization with Staphylococcus and a decreased colonization.

**FIGURE 4** Analysis of ceramide subclass composition in healthy individuals and patients with AD across different sample sites. A, Abundance of CER[NS] and CER[NH] in healthy controls and patients with atopic dermatitis at the sun forehead (FH) and the sun arm. Comparison of healthy individuals and AD patients shows that the CER[NS] group is upregulated in the FH in both healthy and AD patients. B, Differential abundance of the single ceramides of the CER[NS] and CER[NH] subclass in healthy controls and patients with atopic dermatitis at forehead (FH) compared to the arm. The fold change is shown for ceramides with a significant false discovery rate. Only 5 ceramides in both ceramide subclass groups were responsible for the site-specific differences in ceramide class proportions. + = \(P < .1\), * = \(P < .05\), ** = \(P < .005\), *** = \(P < .0005\)
with other bacterial genera in AD patients. This is in line with previous reports, which also confirmed an increased colonization of Staphylococcus on the skin of AD patients.\textsuperscript{[23,37-39]} In healthy patients, a positive correlation of Rothia, Actinomyces, Lactobacillus, Acinetobacter, Haemophilus and Neisseria with abundances of saturated short-chain FFAs, such as FA16:0 and FA18:0, was observed, while in AD patients FA16:0 and FA18:0 negatively correlated with Staphylococcus. The short-chain FFAs FA16:0 and FA18:0 thus show an overall inverse correlation effect to bacterial colonization as observed by lipid species correlation. However, this result has to be interpreted with caution, as many topical formulations contain C16:0 and C18:0. While patients were advised to not use any topical treatment for 24 hours before samples were taken, the washout period is short and it is possible that residues of C16:0 and C18:0 remained, which could explain increased levels of these FFAs in AD patients. Additionally, the glue of tape strips often contains C16:0 and C18:0 as well, but as negative samples did not reveal upregulated levels we conclude that the elevated levels of C16:0 and C18:0 were not due to sampling methods. Another concern regarding the interpretation of this data is that the first tape strip could contain high levels of C16:0 and C18:0 from sebum. A recent study analysed the difference in sebum fatty acid profiles in AD patients and concluded that C18 ceramides were decreased in AD patients compared to healthy subjects.\textsuperscript{[36]} Thus, although the correlation of C16:0 and C18:0 levels with skin microbiome changes has to be interpreted with caution, it seems that the elevated levels in AD patients are not due to sampling methods or differences in sebum. It is tempting to speculate that these saturated short-chain FFAs could play an important role in reversing lipid deregulation in AD, but further studies are warranted to analyse beneficial properties of these FFAs.

Our analysis also revealed significant changes between body areas, especially with regard to abundances of ceramide subclasses. The lipid profile at different sites in healthy skin has previously been studied,\textsuperscript{[40,41]} but no analysis of different skin sites in AD has been conducted. Ludovici et al could show that sebaceous gland secretion plays a role in lipid composition and that lipid composition differs at different body sites in respect to sebaceous gland density.\textsuperscript{[41]} While NS ceramides are upregulated at the FH in both healthy and AD patients, NH ceramides are downregulated. In both classes, we can see that the site-specific regulation is different between healthy individuals and patients with AD. While the differences in NS class are mainly due to alterations in the abundances of S(C18)16:0, the differences in NH class are primarily due to changes in H(C16)26:0. While S(C18)16:0 is a well-known ceramide already identified to be implicated in AD pathogenesis, the biological role of H(C16)26:0 is still unknown. Further studies will be needed to elucidate the biological function of H(C16)26:0 and the possible implications for the pathogenesis in AD. Analysis of ceramide abundances at the FH shows that ceramide abundances are much more similar between healthy individuals and patients with AD, as compared to the FA where ceramide abundances differ more greatly. This indicates a complex dysregulation of skin lipid metabolism in AD, but further studies will be needed to verify our findings.

Other than sebaceous gland density and differences in skin site physiology, filaggrin could also play a role in lipid composition regulation. Filaggrin monomers can be incorporated into the lipid envelope in the stratum corneum, and mutation in FLG can thus lead to impaired skin barrier function. One possible mechanism by which FLG mutation can lead to altered lipid organization is via secretory phospholipase A2 (sPLA2), which converts phospholipids to fatty acids.
## TABLE 1
Analysis of microbiome core taxa at all skin sample sites. Correlation analysis of skin lipids and microbiome core taxa. The abundance of various ceramide species correlated strongly positive with the abundance of Staphylococcus and strongly negative with Propionibacterium and Acinetobacter in AD patients, but not in healthy subjects.

| Actinobacteria | Firmicutes | Proteobacteria |
|----------------|------------|----------------|
| Controls | | |
| Propionibacterium | Corynebacterium | Kocuria | Rothia | Staphylococcus | Streptococcus | Anaerococcus | Finegoldia | Acinetobacter | Haemophilus | Neisseria | Chryseobacterium |
| AH | 0.310 | 0.436 | 0.026 | -0.291 | -0.081 | -0.087 | 0.207 | 0.322 | 0.424 | -0.086 | -0.430 | 0.634 |
| As | 0.407 | 0.471 | -0.033 | -0.154 | 0.060 | 0.068 | 0.391 | 0.416 | 0.389 | 0.118 | -0.231 | 0.387 |
| AP | 0.337 | 0.468 | 0.019 | -0.302 | -0.086 | -0.084 | 0.224 | 0.343 | 0.452 | -0.082 | -0.452 | 0.671 |
| NP | 0.179 | 0.287 | 0.081 | -0.263 | -0.121 | 0.126 | 0.233 | 0.307 | -0.127 | 0.350 | 0.487 |
| FA16:0 | -0.436 | -0.555 | 0.067 | 0.260 | 0.089 | -0.010 | -0.280 | 0.379 | -0.498 | 0.000 | 0.444 | -0.701 |
| FA18:0 | -0.429 | -0.552 | 0.054 | 0.272 | 0.090 | 0.023 | -0.276 | -0.381 | 0.500 | -0.014 | 0.454 | -0.710 |
| FA18:2 | 0.206 | 0.095 | -0.339 | 0.320 | 0.032 | 0.325 | 0.101 | -0.039 | 0.357 | 0.265 | -0.233 |
| FA20:1 | 0.586 | 0.529 | -0.490 | 0.226 | -0.023 | 0.415 | 0.336 | 0.226 | 0.290 | 0.464 | 0.021 | 0.209 |
| FA20:2 | 0.631 | 0.622 | -0.430 | 0.104 | -0.049 | 0.343 | 0.372 | 0.313 | 0.404 | 0.387 | -0.128 | 0.403 |
| FA22:1 | 0.634 | 0.573 | -0.529 | 0.244 | -0.026 | 0.449 | 0.364 | 0.246 | 0.315 | 0.501 | 0.021 | 0.228 |
| FA24:1 | 0.600 | 0.557 | -0.472 | 0.189 | -0.031 | 0.393 | 0.347 | 0.253 | 0.325 | 0.441 | -0.025 | 0.268 |

| Cases | | |
|----------------|------------|----------------|
| Propionibacterium | Corynebacterium | Actinomyces | Rothia | Staphylococcus | Lactobacillus | Acinetobacter | Haemophilus | Neisseria | Enhydrobacter |
| AS | -0.360 | 0.118 | -0.847 | -0.640 | 0.700 | -0.808 | -0.835 | -0.911 | -0.706 | -0.477 |
| ADS | -0.357 | 0.112 | -0.887 | -0.670 | 0.723 | -0.839 | -0.867 | -0.944 | -0.743 | -0.496 |
| NS | -0.370 | 0.117 | -0.902 | -0.681 | 0.738 | -0.855 | -0.884 | -0.963 | -0.754 | -0.506 |
| NDS | -0.416 | 0.157 | -0.812 | -0.617 | 0.710 | -0.803 | -0.832 | -0.909 | -0.669 | -0.471 |
| FA16:0 | -0.080 | 0.158 | 0.904 | 0.661 | -0.496 | 0.680 | 0.692 | 0.737 | 0.807 | 0.423 |
| FA18:0 | -0.066 | 0.143 | 0.853 | 0.625 | -0.474 | 0.646 | 0.658 | 0.701 | 0.761 | 0.402 |
| FA20:0 | -0.254 | 0.240 | 0.699 | 0.503 | -0.280 | 0.452 | 0.454 | 0.474 | 0.647 | 0.292 |
| FA22:0 | -0.296 | 0.267 | 0.721 | 0.517 | -0.271 | 0.452 | 0.453 | 0.472 | 0.671 | 0.295 |
| FA24:0 | -0.232 | 0.241 | 0.825 | 0.596 | -0.367 | 0.559 | 0.564 | 0.593 | 0.755 | 0.357 |
| FA20:2 | 0.184 | -0.211 | -0.822 | -0.596 | 0.392 | -0.576 | -0.583 | -0.615 | -0.747 | -0.365 |
| FA24:1 | -0.268 | 0.098 | -0.544 | -0.413 | 0.470 | -0.534 | -0.553 | -0.604 | -0.450 | -0.313 |

*Bold and italic values indicate nominal significant associations (P<.05).*
acids. In 3D skin constructs, a FLG knock-down led to increased activation of sPLA2 and to an accumulation of free fatty acids.\[20\]

In this study, we have comprehensively profiled lipid species in correlation with the microbiome. We have utilized the non-invasive method of tape stripping to analyse skin lipids in 10 patients with AD and 10 healthy subjects. Although our study cohort is small, we have a balanced study design, and by utilizing the complex lipid panel, we were able to measure all three lipid classes, including ceramides, but also cholesterol sulphate and free fatty acids. Previous studies have already utilized tape stripping and lipidomics analysis to investigate lipid composition in AD.\[15\] The strength of our study is that we have not only analysed ceramides, but also cholesterol sulphate and free fatty acids. Moreover, to the best of our knowledge we are the first to directly compare the lipid profile at different skin sites in parallel with analysing the microbiome. The limitation of our study is the small study cohort as well as missing longitudinal data to investigate lipid ratios during the disease course. Another limitation of our study that by using a targeted lipidomics approach, we might have not analysed lipids that are of importance (eg very long-chain FFAs or long-chain ceramides).

We conclude that site-specific sampling is important for the holistic analysis of AD lipids and that lipid dysregulation in AD is dependent on exogenous factors.

CONFLICT OF INTEREST

The authors declare that they have no relevant conflicts of interest.

AUTHOR CONTRIBUTIONS

HE, ER, and SW designed the study. FT, DS, and IH performed biosampling. ER performed sample processing and analysis. HE and HB analysed the data and wrote the manuscript. ER, EP and SW revised the manuscript critically.

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SUPPORTING INFORMATION

Additional supporting information may be found online in the Supporting Information section.
**Figure S1.** Impact of sex on total abundance of ceramides, cholesterol and free fatty acids.

**Figure S2.** Correlation of age with total abundance of ceramides, cholesterol and free fatty acids.

**Figure S3.** Correlation of TEWL with total abundance of ceramides, cholesterol and free fatty acids.

**Figure S4.** Correlation of pH with total abundance of ceramides, cholesterol and free fatty acids.

**Figure S5.** Heatmap of ceramides in healthy individuals and patients with AD. The colored bar at the top indicates ceramide subclass composition. The colored bar at the side indicates sample site: Forehead (red), forearm (blue), antecubital fossa (darkblue).

**Figure S6.** Free fatty acid chain length in FLGmut and FLGwt carrier, as well as in healthy individuals and patients with AD.

**Figure S7.** Average chain length of CER (left) and FFA (right) at the forehead. Comparison between Ctrl and AD (upper row), FLGwt and FLGmut (middle row) and all 4 study groups (lower row).

**Figure S8.** Average chain length of CER (left) and FFA (right) at the cubital fossa. Comparison between Ctrl and AD (upper row), FLGwt and FLGmut (middle row) and all 4 study groups (lower row).

**Figure S9.** Average chain length of CER (left) and FFA (right) at the proximal lower arm. Comparison between Ctrl and AD (upper row), FLGwt and FLGmut (middle row) and all 4 study groups (lower row).

**Figure S10.** Ceramide (A) and FFA (B) abundance in AD patients vs healthy individuals. (A) and in AD patients with FLG mutation carrier vs. non-carrier (B). Significant differences are indicated by (*).

**Figure S11.** Ceramide abundance in FLG mutation carrier vs. non-carrier at three different sites (A) forehead, (B) cubital fossa and (C) proximal lower arm. Blue = ω-hydroxy CERs, red = nonhydroxy CERs and brown/orange = esterfied ω-hydroxy CERs.

**Figure S12.** Ceramide (A) and FFA (B) abundance in AD patients comparing FLG mutation carrier vs. non-carrier. Significant differences are indicated by (*).

**Table S1.** Proband characteristics.

**Table S2.** Ceramide composition in healthy volunteers without FLG mutations measured by MLP = Metabolon Lipid Panel.

**Table S3.** Correlation of age and sex on lipid composition.

**Table S4.** TEWL and pH measurements in healthy individuals and patients with AD as well as FLG competent and FLG deficient individuals. Fh: forehead, Fa: forearm.

**Table S5.** Correlation of TEWL and pH on lipid composition.

**Table S6.** Ceramide species (pmol/disk) significantly different at a threshold of 0.01 between AD patients and controls in at least one of the three sites: forehead, cubital fossa, proximal lower arm. FC = fold change, CI = confidence interval.

**Table S7.** Balance of ceramides and free fatty acids measured by the Shannon Index.

**Table S8.** Differential analysis of CER[NS] and CER[NH] in healthy individuals and patients with AD.

**Table S9.** Ceramide proportions (relative abundance) and their 95% CIs in FLG wildtype (FLGwt) and FLG mutation carriers (FLGmut).

**Table S10.** Ceramide anaytes of the Metabolon Lipid Panel.

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