

Topical supplementation with physiological lipids rebalances the stratum corneum ceramide profile and strengthens skin barrier function in adults predisposed to atopic dermatitis

Paul V Andrew¹, Samuel F Williams¹, Kirsty Brown¹, John Chittock¹, Abigail Pinnock¹, Anna Poyner¹, Michael J Cork^{1,2,3} and Simon G Danby¹

¹Sheffield Dermatology Research, Division of Clinical Medicine, School of Medicine and Population Health, The University of Sheffield, Sheffield, UK

²Sheffield Teaching Hospitals NHS Foundation Trust, The Royal Hallamshire Hospital, Sheffield, UK

³Sheffield Children's NHS Foundation Trust, Sheffield Children's Hospital, Western Bank, Sheffield, UK

Correspondence: Paul V. Andrew. Email: paul.andrew@sheffield.ac.uk

Abstract

Background People with atopic dermatitis (AD) suffer from dry, itchy skin with reduced skin barrier function that leaves it prone to irritant and allergen penetration. Alterations in the composition and structure of the stratum corneum (SC) lipid lamellae underpin this increase in permeability. A wide range of emollients is used to ameliorate the skin of patients with AD, but the majority have unclear effects on the lipid lamellae and barrier function.

Objectives To compare the effects of a multivesicular emulsion containing physiological lipids and glycerine (MVE+GL) with a commonly prescribed oil-in-water emulsion containing glycerine without physiological lipids (O/W+G).

Methods A double-blind intraparticipant-controlled study was undertaken in adults with a history of eczema. Participants applied MVE+GL to one forearm and lower leg and O/W+G to contralateral sites twice daily for 28 days. Skin properties were assessed before and after treatment. A detailed lipidomic profile was generated from SC samples, alongside *in vivo* attenuated total reflectance Fourier transform infrared spectroscopic analysis of its molecular composition.

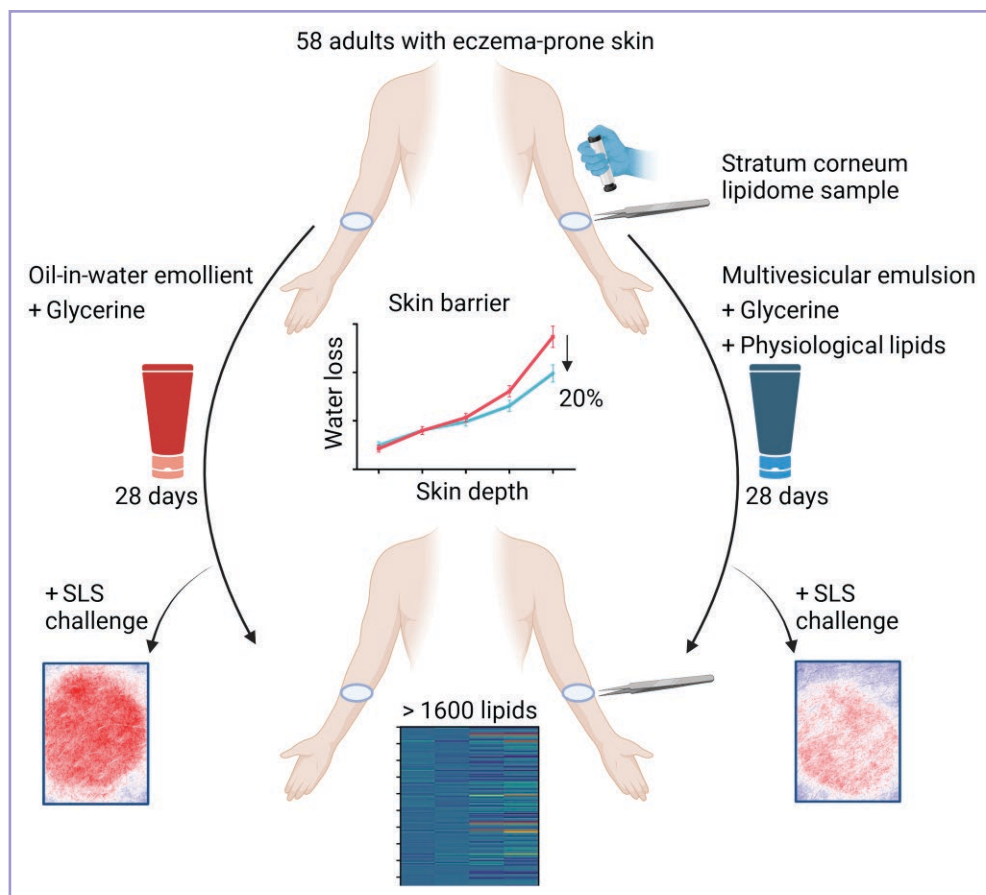
Results Fifty-eight people were included in the study [mean (SD) age 46 (21) years]. At sites treated with MVE+GL skin barrier integrity improved significantly [mean (SD) transepidermal water loss after 20 skin tape strips (TEWL₂₀) 38.02 (18.64) g m⁻² h⁻¹ pretreatment vs. 29.79 (13.47) g m⁻² h⁻¹ post-treatment; *P*<0.001], whereas O/W+G had no effect [35.6 (18.39) g m⁻² h⁻¹ vs. 37.4 (16.69) g m⁻² h⁻¹]. Concordantly, skin sensitivity to sodium lauryl sulfate (SLS) was significantly reduced by MVE+GL treatment [mean (SD) post-SLS TEWL 35.58 (15.43) g m⁻² h⁻¹ pretreatment vs. 29.54 (11.64) g m⁻² h⁻¹ post-treatment (*P*<0.001); erythema was also reduced]. Skin moisture increased more rapidly at sites treated with MVE+GL vs. O/W+G, leading to a more rapid reduction in visual skin dryness. Over 1600 lipid species were detected in the SC. Ceramide species NP (non-hydroxy-phytosphingosine) and AP (α -hydroxy-phytosphingosine) with 18-carbon sphingoid bases, both ingredients of the MVE+GL, increased significantly by 24% and 19%, respectively, following MVE+GL treatment. In contrast, changes of 9% for NP(18) and 6% for AP(18) were not statistically significant at sites treated with O/W+G. Increased abundance of NP(18) species relative to NdS (non-hydroxy-dihydrosphingosine) species was related to improvements in skin barrier integrity.

Conclusions While the glycerine-containing emollient reduced skin dryness, it had no impact on barrier function. In contrast, MVE+GL improved the physical integrity of the barrier and reduced the sensitivity of the skin.

Accepted: 20 May 2025

© The Author(s) 2025. Published by Oxford University Press on behalf of British Association of Dermatologists. This is an Open Access article distributed under the terms of the Creative Commons Attribution License (<https://creativecommons.org/licenses/by/4.0/>), which permits unrestricted reuse, distribution, and reproduction in any medium, provided the original work is properly cited.

Graphical Abstract



Lay summary

Eczema is the most common chronic skin condition. It affects 1% to 10% of adults in Europe. The upper layers of our skin form a barrier that keeps water in our bodies and stops irritants from getting in. This skin barrier is made of cells called 'corneocytes'. These are surrounded by a tough protein envelope and embedded in a network of 'fatty' molecules (called 'lipids'). The composition of these fatty molecules is altered in the skin of people with eczema. The skin does not function effectively as a barrier as more water is lost, and irritants can more easily penetrate the deeper skin layers. Moisturizing creams are an effective treatment for eczema, but it is unclear which are most appropriate to use.

This study was carried out in the UK. We aimed to find out if a cream containing lipids found in skin could improve the skin's ability to act as a barrier. We compared this cream to one without these lipids. We asked 58 people with eczema to apply these creams separately on their forearms for a month and measured how the skin responded. We found that skin treated with the cream containing skin-identical lipids lost less water. It was also more resistant to physical damage and was less sensitive to irritation than before treatment. In contrast, the cream without lipids did not improve the skin barrier. Both creams improved dryness, but quicker results were found for the cream containing the lipids found in skin.

Our findings suggest that the addition of skin-identical lipids to moisturizing creams is a promising way of improving skin barrier function.

What is already known about this topic?

- Some emollients used to treat atopic dermatitis (AD) have little or no positive effect on the function of the skin barrier.
- The lipid profile of the stratum corneum (SC) differs in people with AD compared with the profile in healthy skin.
- Use of emollients can alter the arrangement of lipids in the skin barrier.
- Emollients containing physiological lipids have been used to treat AD, but whether the lipid composition is changed by treatment is unknown.

What does this study add?

- Glycerol as an emollient additive is insufficient to improve skin barrier function in eczema-prone individuals.
- The ceramide composition of the skin barrier is altered by emollient application.
- Changes in ceramide composition are identified which associate with improvements in skin barrier function.
- The carbon length of a ceramide's sphingoid base appears to determine its impact on skin barrier function.
- The abundance of non-hydroxy-dihydrosphingosine (NdS) ceramides negatively correlates with skin barrier function.

What is the translational message?

- Where skin sensitivity and skin barrier impairment are an issue, emollients must do more than resolve clinical signs of dryness.
- Topical supplementation with essential physiological lipids is an effective strategy to repair SC lipid matrices and leads to improved skin barrier function.
- The success of lipid supplementation depends on how applied lipid mixtures change the balance of ceramides in the skin at the species level.

In atopic dermatitis (AD), reduced function of the skin as a barrier leads to excessive water loss from, and increased irritant entry into, the viable skin layers. The barrier function of skin resides in the stratum corneum (SC), a structure composed of multiple layers of corneocytes embedded in a lipid-rich matrix called the lipid lamellae.¹ The major components of these lamellae are ceramides, cholesterol and fatty acids. Altered ceramide profiles in patients with AD,² and evidence from *in vitro* modelling of lipid mixtures,^{3–5} suggest that the ceramide profile of the skin is an important determinant of skin barrier function. Ceramides are composed of a sphingoid base (SB) linked through an amide bond to a fatty acid (the acyl chain). Over 1000 different ceramide species have been identified in the human SC.⁶ Attempts to describe the functional implications of this variability are an ongoing focus of research.

In AD, consistent use of some emollients can increase the time between flares and reduce the requirement for topical corticosteroid use.^{7,8} However, the effectiveness of emollients varies. Clear evidence to support the use of particular formulations or ingredients remains an unmet need. There is evidence that use of emollients enriched with physiological lipids can improve barrier function and clinical outcomes.^{9,10} However, it remains unclear how emollient use affects the lipid profile of the SC.

In this study, we compared an oil-in-water emollient with glycerine (O/W+G) to a multivesicular emulsion with physiological lipids and glycerine (MVE+GL). The aim was to determine the effects of these formulations on the skin barrier and the SC lipidome.

Patients and methods**Study design**

This was an interventional intraparticipant-controlled right/left randomized double-blind cohort study in adults with dry skin and a self-reported history of eczema, comparing the effects of twice-daily application of MVE+GL with O/W+G for 4 weeks. Skin condition was assessed at baseline, day 2, day 14 and at the end of treatment (EoT).

The study targeted recruitment of 58 participants in 3 cohorts of approximately equal size based on age (age groups: 18–39 years; 40–59 years; ≥60 years). The target for completion was 48 participants with 82% power to detect a difference of 6 g m⁻² h⁻¹ in transepidermal water loss (TEWL) after induced irritation (the primary outcome).

Intervention

Participants undertook a 7-day washout from leave-on topical products at the test sites on the volar forearm and the outer surface of the lower leg before baseline assessments. Application of O/W+G [Cetraben® (containing glycerine); Thornton & Ross, Huddersfield UK] and MVE+GL [CeraVe® (containing glycerine, capric triglyceride, ceramides NP, AP and EOP, phytosphingosine and cholesterol); L'Oréal, Clichy, France] started after these assessments. Participants were trained in the application of two fingertip units of emollient to each treatment area in accordance with the randomization schedule.

Outcomes

The primary outcome was the difference in TEWL between test sites after sodium lauryl sulfate (SLS)-induced irritation. Secondary outcomes included erythema following SLS exposure, skin barrier integrity measured with TEWL and skin tape-stripping (STS), barrier cohesion based on protein collection by STS, moisturization assessed by visual scoring and hydration (capacitance). Comparisons were made between test sites and the effect of each emollient vs. baseline. Exploratory outcomes included molecular composition of the SC using attenuated total reflectance Fourier transform infrared spectroscopy (ATR-FTIR) and STS, and mass spectroscopy of STS samples. Further detail is provided in Appendix S1 (see [Supporting Information](#)).

Statistical analysis

All data were analysed using Prism 10 (GraphPad, La Jolla, CA, USA). The significance threshold for the biophysical and ATR-FTIR analyses was $P < 0.05$. Results are presented as mean (SD) in the text, unless otherwise indicated.

Results

Fifty-eight adults with a self-reported history of eczema were recruited and began treatment between April 2022 and March 2023 (Figure S1, Table S1; see Supporting Information). Participants used the study emollients twice daily on separate test areas on opposite sides of the body for 28 (4) days. Similar amounts of each treatment were used: 3.66 (0.86) g daily of O/W+G and 4.04 (0.92) g daily of MVE+GL.

TEWL, an important measure of permeability barrier function, was within the expected range at baseline and was unaltered by treatment (Figure 1). STS removes cell layers from the SC leading to an increase in TEWL, a measure of skin barrier integrity. Before treatment there was no difference between test sites in TEWL following 20 STS (TEWL₂₀) or the amount of protein removed. At EoT, skin barrier integrity/cohesion was similar at sites treated with O/W+G [TEWL₂₀ 35.65 (18.40) g m⁻² h⁻¹ before vs. 37.38 (16.69) g m⁻² h⁻¹ after treatment (*P*=0.47), a difference of +5%; protein removed was 347.99 (110.18) µg cm⁻² before

vs. 328.95 (108.58) µg cm⁻² after treatment (*P*=0.18)]. At sites treated with MVE+GL, barrier condition was improved with a reduction in the amount of protein removed vs. baseline [353.57 (101.43) µg cm⁻² before vs. 282.85 (114.21) µg cm⁻² after treatment; *P*<0.001] and there was a 22% reduction in TEWL₂₀ [38.02 (18.64) g m⁻² h⁻¹ before vs. 29.79 (13.47) g m⁻² h⁻¹ after treatment; *P*<0.001]. The centre of gravity (COG) of the lipid peak (approximately 2850 cm⁻¹) in an ATR-FTIR mid-infrared spectrum of the skin indicates the chain conformation of the SC lipids,¹¹ with a higher position indicating an increase in conformational disordering associated with reduced barrier function.^{3, 12} Treatment with O/W+G significantly increased the COG of the lipid peak, whereas treatment with MVE+GL had no effect.

Next, we examined whether the skin's response to irritant exposure was affected by treatment. SLS was applied to the skin for 24 h under occlusion and the resulting inflammation assessed by quantifying erythema and TEWL. At baseline, there was a clear erythematous response to SLS at >95% of the test sites (median score 1, mild erythema). Erythema index (EI) increased from 38.77 (0.59) to 50.60

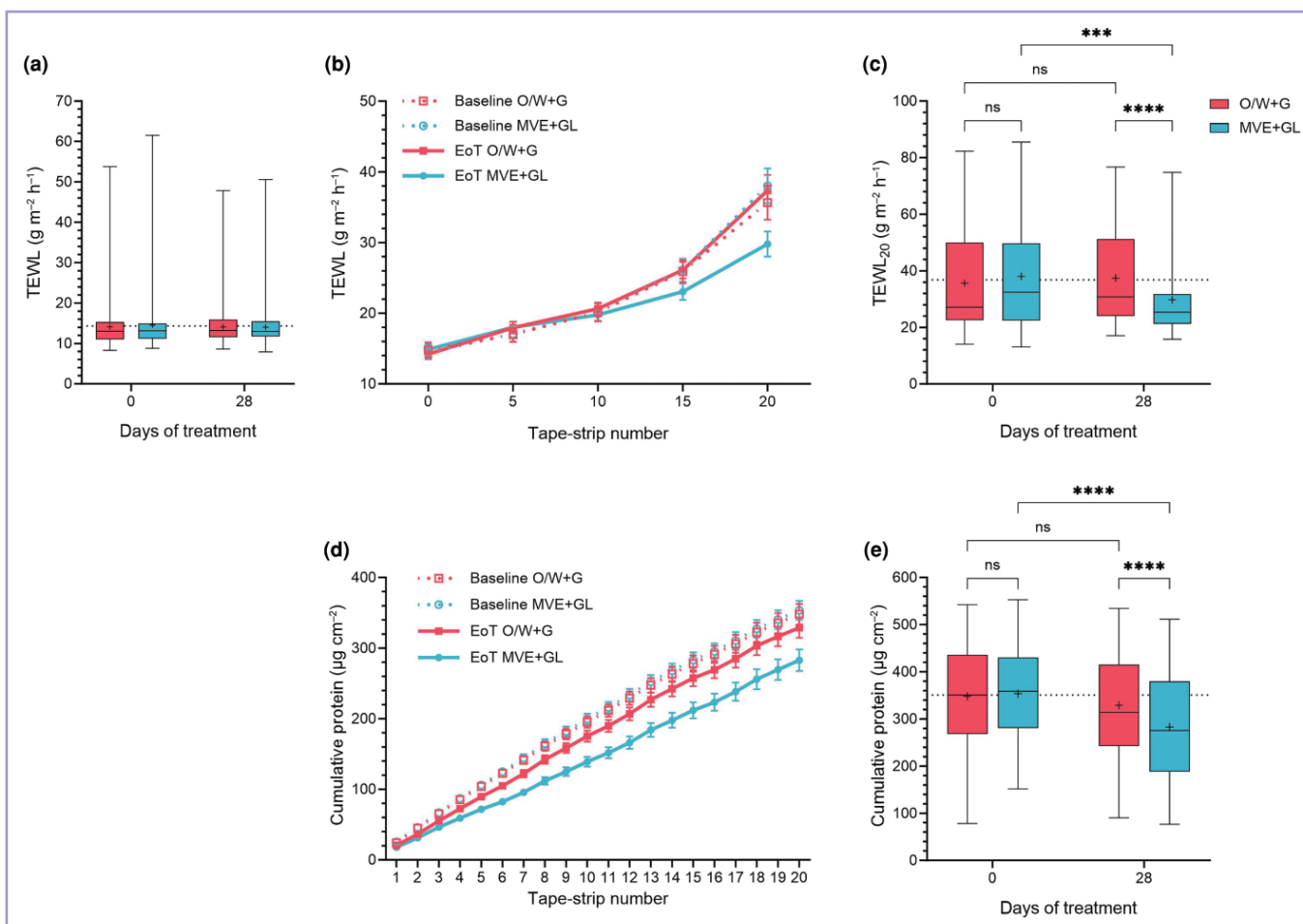


Figure 1 Skin barrier integrity and cohesion are affected by emollient treatment. (a) Basal transepidermal water loss (TEWL) on the forearm at baseline and at end of treatment (EoT). Mixed-effects analysis revealed no significant effect of treatment or difference between treatments. (b) TEWL and (d) protein removed at increasing depths through the stratum corneum (SC) and after skin tape-stripping (STS) (c, e) at baseline and EoT. Dotted line in (a), (c) and (e) indicates the day 0 mean. Line graphs in (b) and (d) show the mean (SEM). Boxes represent the interquartile range, whiskers show the range, median is indicated as a horizontal line and '+' denotes the mean. MVE+GL, multivesicular emulsion with physiological lipids and glycerine; ns, not significant (i.e. *P*>0.05); O/W+G, oil-in-water emollient with glycerine. ****P*<0.001, *****P*<0.0001 (Fisher's post-test following mixed-effects analysis).

(0.25) arbitrary units (AU) and TEWL increased from 13.42 (0.25) to 35.36 (0.31) $\text{g m}^{-2} \text{h}^{-1}$ (areas combined). Following treatment, the skin's response to SLS was unchanged at sites treated with O/W+G (Figure 2). In contrast, sensitivity to SLS was reduced at sites treated with MVE+GL vs. baseline; more sites had no or barely perceptible erythema than moderate or strong erythema (Table S2; see Supporting Information). EI was reduced [47.30 (8.53) vs. 50.77 (8.90) AU; $P=0.002$] and TEWL was lower [29.54 (11.64) vs. 35.58 (15.43) $\text{g m}^{-2} \text{h}^{-1}$; $P<0.001$]. At EoT, average TEWL after SLS exposure was 5.04 $\text{g m}^{-2} \text{h}^{-1}$ lower at sites treated with MVE+GL vs. O/W+G ($P<0.001$).

At baseline, all participants had visible dryness on the lower leg [median score 2; scaling, slight roughness (Table S3; see Supporting Information)]. Hydration measured using the capacitance method indicated very dry skin [25.37 (0.07) AU, both areas]. Skin moisturization improved with both treatments; median skin dryness decreased to 1 (faint scaling/roughness) at sites treated with O/W+G and to 0.5 at sites treated with MVE+GL after 2 days of treatment. Most

sites had no dryness by day 14 (Table S4; see Supporting Information). SC hydration increased concordantly: sites treated with MVE+GL were more hydrated than sites treated with O/W+G after 2 days [O/W+G 31.70 (10.14) vs. MVE+GL 33.32 (9.16) AU; $P=0.02$] and 14 days [O/W+G 35.64 (8.51) vs. MVE+GL 37.53 (8.57) AU; $P=0.003$] of treatment. Both sites had similar levels of hydration at EoT ($P>0.99$; Figure 3). To directly measure the amount of water in the SC, ATR-FTIR spectroscopy with STS was used. FTIR measurements are normalized to protein, inferred from the amide peak. Both treatments increased SC water levels: sites treated with MVE+GL exhibited 25% more water across the SC than sites treated with O/W+G at EoT (Fisher's $P<0.001$; Figure 4 d). ATR-FTIR spectroscopy was also used to determine the amount of glycerine through quantification of hydroxyl groups of polyols. Polyols increased from baseline with both treatments [at EoT: O/W+G 0.73 (0.11) vs. MVE+GL 0.87 (0.20) AU; $P<0.001$]. Polyol abundance was significantly associated with skin barrier integrity (TEWL₂₀: $r=-0.51$) and hydration (capacitance: $r=0.45$).

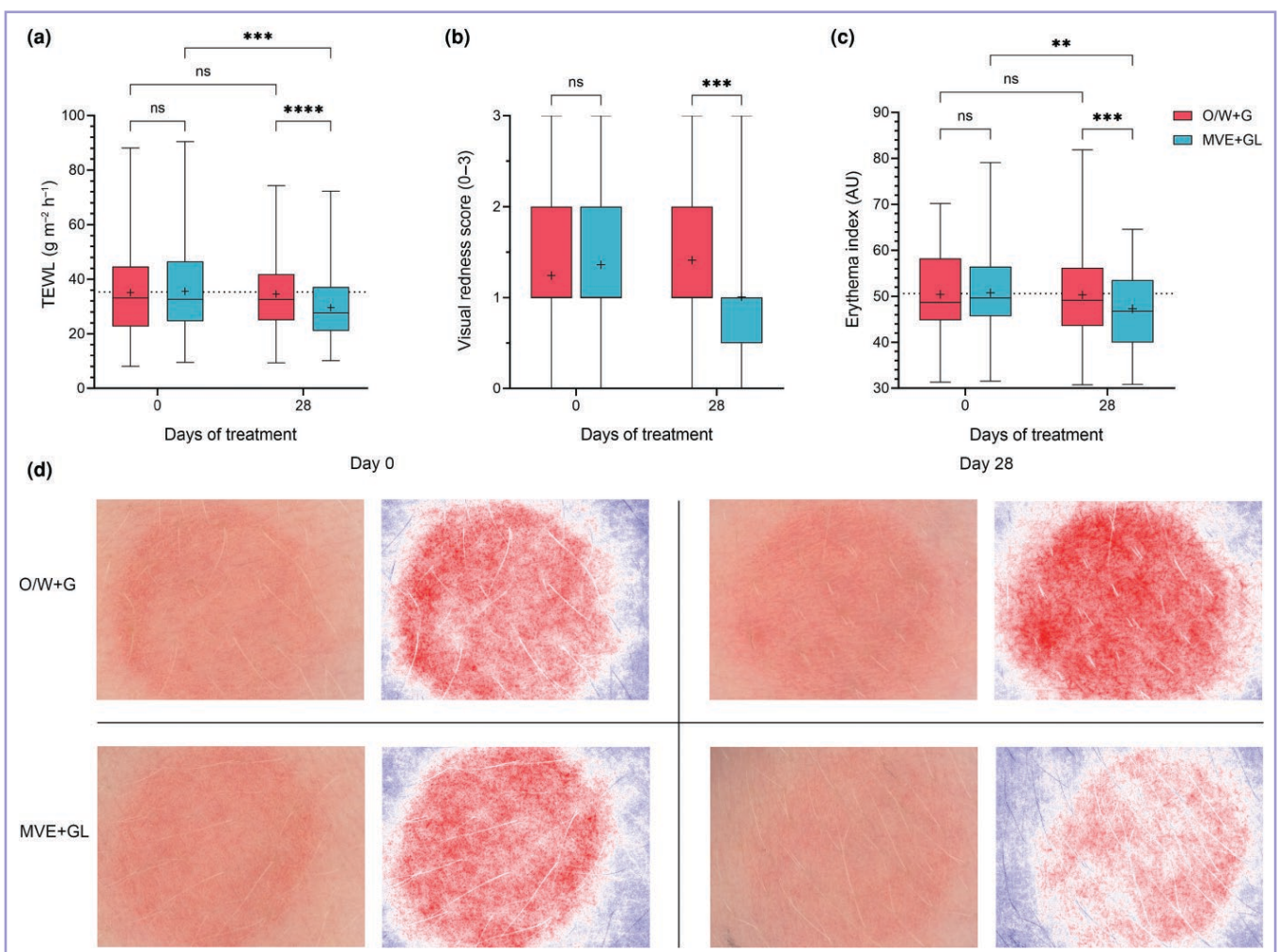


Figure 2 Skin irritation following exposure to sodium lauryl sulfate (SLS) is affected by emollient treatment. (a) Transepidermal water loss (TEWL), (b) visual redness score and (c) erythema index on the forearm 24 h after SLS exposure, at baseline and at end of treatment. (d) Representative images of erythema: rendered panels show erythema index scoring. Boxes represent the interquartile range, whiskers show the range, median is indicated as a horizontal line and '+' denotes the mean. AU, arbitrary unit; MVE+GL, multivesicular emulsion with physiological lipids and glycerine; ns, not significant (i.e. $P>0.05$); O/W+G, oil-in-water emollient with glycerine. ** $P<0.01$, *** $P<0.001$, **** $P<0.0001$ [(a, c) Fisher's test following mixed-effects analysis; (b) Wilcoxon signed rank test].

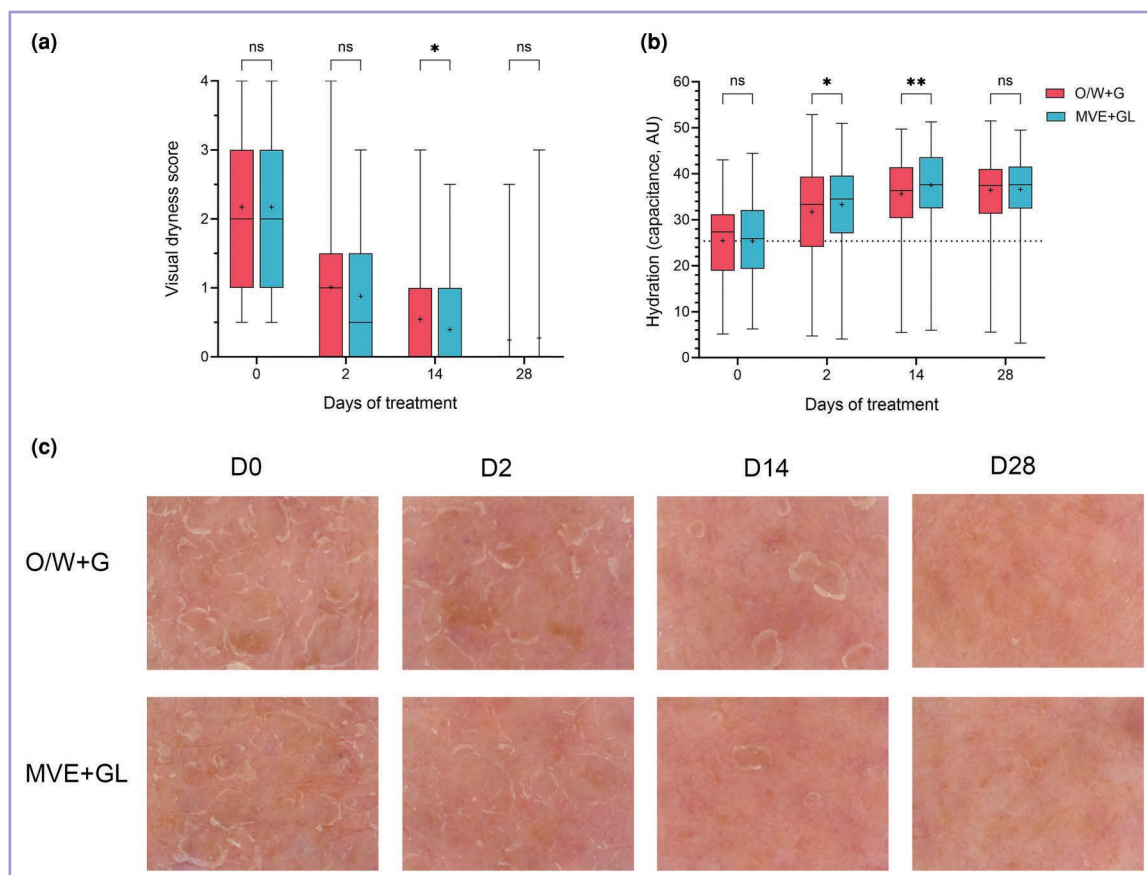


Figure 3 Skin moisturization is affected by emollient treatment. (a) Visual dryness scores, (b) hydration and (c) representative images of the lower leg at baseline and throughout treatment. In (b), mixed-effects analysis identified significant effects of treatment and a difference between treatments; however, post-test comparisons were made between treatments only. Boxes represent the interquartile range, whiskers show the range, median is indicated as a horizontal line and '+' denotes the mean. The dotted line indicates the day 0 mean. AU, arbitrary unit; D, day; MVE + GL, multivesicular emulsion with physiological lipids and glycerine; ns, not significant (i.e. $P > 0.05$); O/W + G, oil-in-water emollient with glycerine. * $P < 0.05$, ** $P < 0.01$ [results of multiple (a) Wilcoxon tests and (b) Sidak's post-test].

Finally, we undertook MS-based profiling of SC ceramides, cholesterol and glycerides obtained from STS in tandem with ATR-FTIR spectroscopy at baseline and EoT. Total lipids, including physiological and nonphysiological lipids (based on CH_2 groups), were increased from baseline by 49% following O/W + G and by 54% following MVE + GL treatment (Figure 4 g), whereas lipid esters predominantly from physiological lipids, including triglycerides (included in MVE + GL), were only increased after MVE + GL treatment [49% ($P = 0.008$); Figure 4 h]. Total lipids and lipid esters were associated with skin dryness ($r = -0.46$ and $r = -0.34$, respectively). In the five lipid groups analysed by MS, the proportion of ceramides relative to total measured lipids increased with emollient use (Figure 5 a); however, this change was not reflected when normalized to protein (Figure S2; see Supporting Information). Analysis of the ceramides by subclass revealed few changes in response to treatment [Figure 5 b, c (relative to lipid class); Figure S2b, c (normalized to protein)]. Stratification of ceramides by their constituent chain lengths revealed striking treatment effects on the relative abundance of ceramides with 18-carbon SBs, and 18- and 24-carbon acyl chains. Regarding the 18-carbon SBs, MVE + GL treatment increased the relative abundance of six of eight subclasses [AH(18) 12%, AP(18)

19%, AS(18) 27%, NP(18) 24%, NS(18) 16%, NdS(18) 18% (Figure 5 h), five of which were consistent with significant changes normalized to protein (Figure S2 d)]; O/W + G increased the levels of three of eight subclasses [AS(18) 14%, NS(18) 11%, NdS(18) 18%, where only AS was significantly elevated when normalized to protein]. NP(18), AP(18) and AS(18) ceramides were significantly more abundant at sites treated with MVE + GL compared with O/W + G (relative to lipid class or normalized to protein). We observed a trend for lower TEWL_{20} (improved skin integrity) in skin with increasing proportions of ceramides with shorter SBs [< 20 carbons; Figure 5 e, Figure S3 (see Supporting Information)]. A relative increase in the proportion of 18-carbon SB ceramides within most subclasses was associated with better skin barrier integrity. We did not find a relationship between acyl or total chain length and skin barrier integrity in this analysis [Figure 5 g; Figure S4 (see Supporting Information)].

Of the 1633 molecular species detected, 364 were present in $> 70\%$ of samples in one of the treatment groups. We compared the abundance of these species before and after treatment. A similar number of species significantly changed in abundance with both treatments (O/W + G, $n = 87$; MVE + GL, $n = 83$), including 52 identical

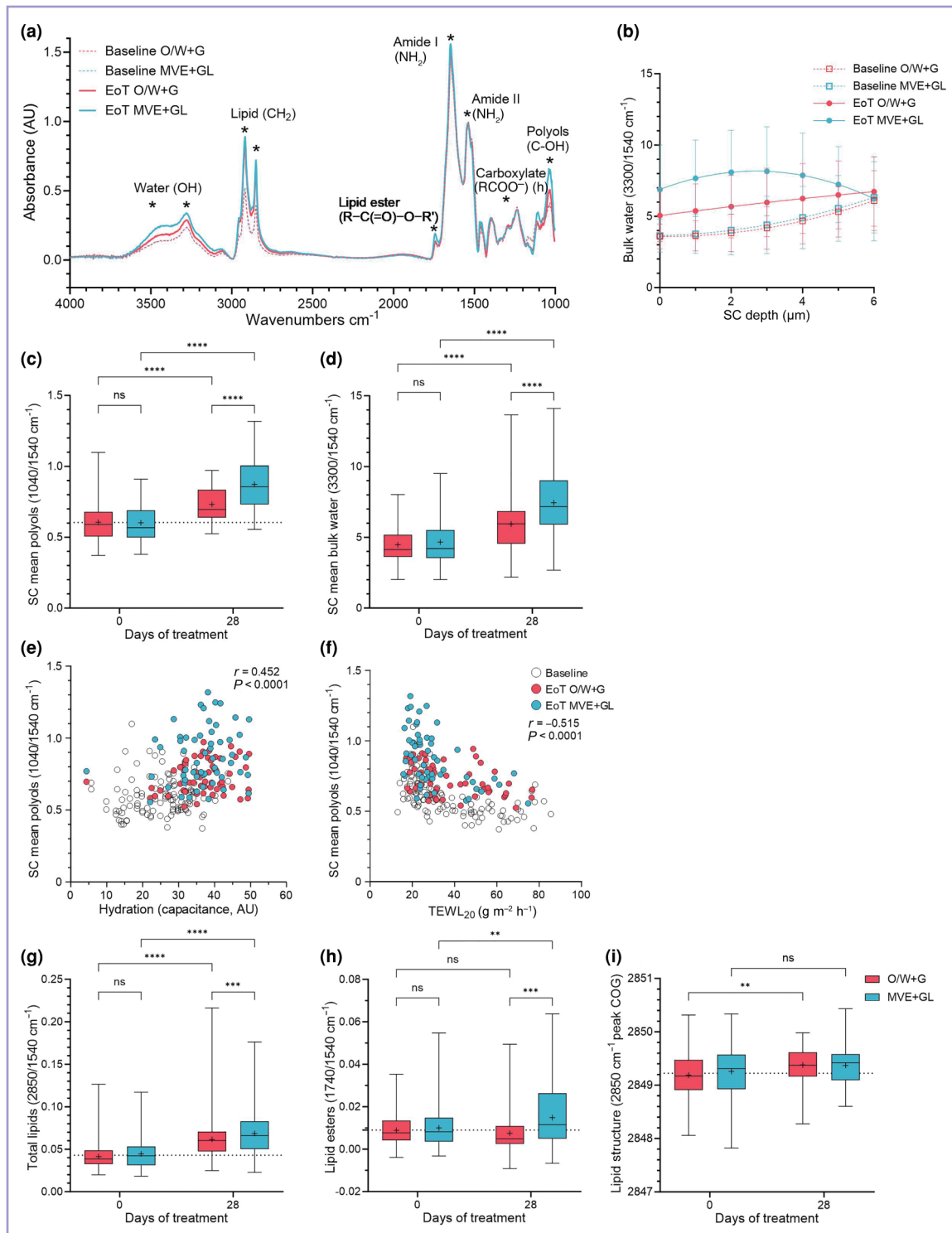


Figure 4 Emollient treatment affects the molecular profile of the stratum corneum (SC). (a) Average mid-infrared spectrum collected using an attenuated total reflectance Fourier transform infrared spectrometer from the skin surface of the forearm. (b) Interpolated water values from 0 to 6 μm within the SC constructed using measurements collected after 5, 10, 15 and 20 tape strips. SC (c) polyols (glycerine) and (d) water. (e) Correlation between polyol (glycerine) abundance and hydration, and (f) barrier integrity. (g) Total lipid, (h) lipid esters and (i) lipid structure. Quantification from absorbance peak area (mean value 0–6 μm) normalized to amide II-associated peak (c, d, g, h) and (i) peak position. AU, arbitrary units; COG, centre of gravity; EoT, end of treatment; MVE+GL, multivesicular emulsion with physiological lipids and glycerine; ns, not significant (i.e. $P > 0.05$); O/W+G, oil-in-water emollient with glycerine. ** $P < 0.01$, *** $P < 0.001$, **** $P < 0.0001$ (Fisher's test following mixed-effects analysis).

species. Significant changes of twofold or greater were only observed after MVE+GL treatment (Figure 6). Apart from the NdS species, these ceramides were identified

in <50% samples at baseline but in far higher numbers of samples after treatment with MVE+GL. We looked for associations between lipid species abundance and

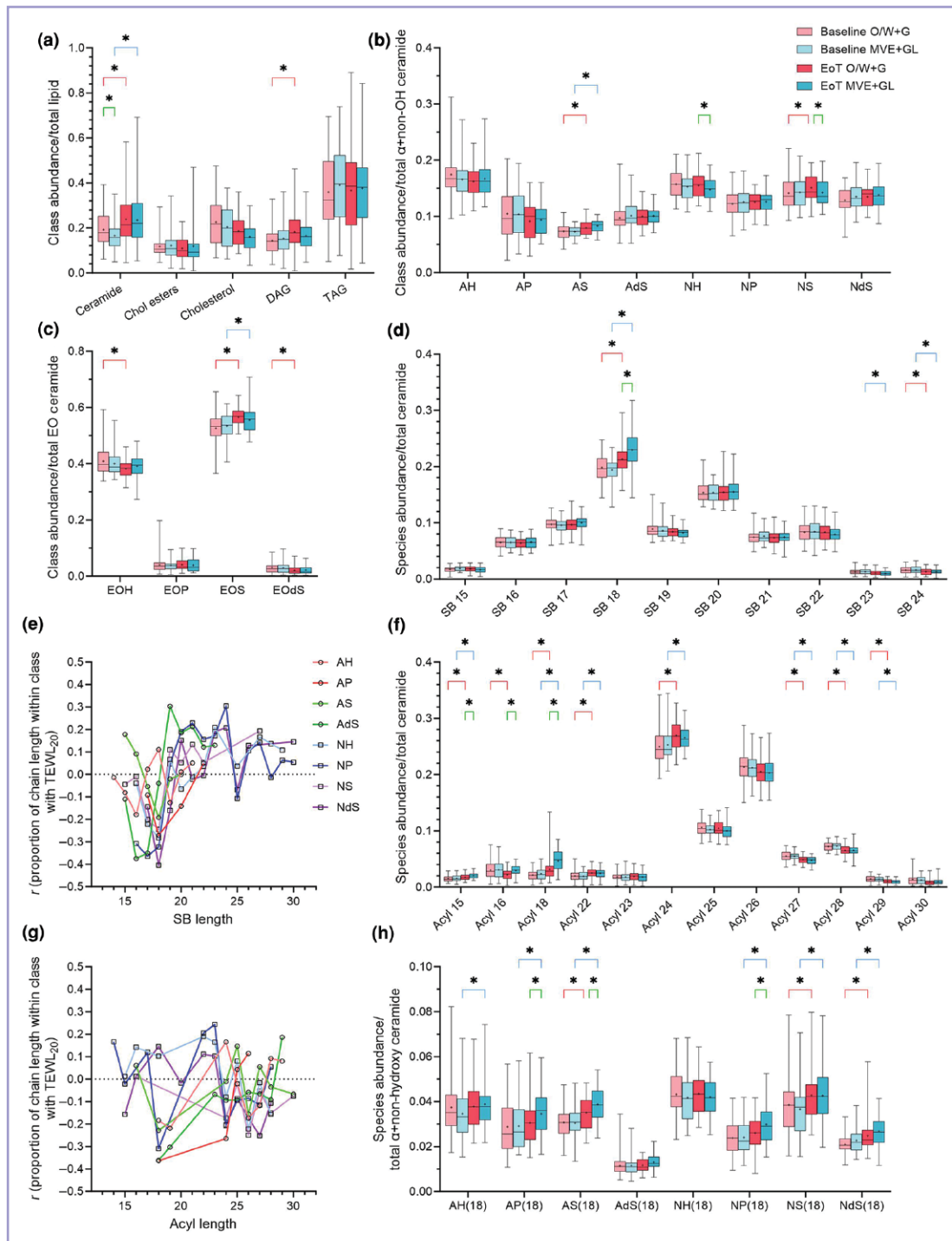


Figure 5 Change in the composition of stratum corneum (SC) lipids in response to emollient treatment: mass spectrometry analysis. Relative abundance of (a) major lipid groups, (b, c) ceramide subclasses and (h) α -hydroxy/non-hydroxy ceramides with an 18-carbon sphingoid base (SB) combined with any acyl chain (16–32 carbons). Distribution of α -hydroxy/non-hydroxy ceramides by (d) SB chain length, combined with any acyl chain (13–35 carbons), and (f) acyl chain length, combined with any SB (14–42 carbons). Only subclasses representing $>1\%$ of α -hydroxy/non-hydroxy ceramides are presented. (e, g) Correlation between the abundance of ceramide by chain length and barrier function. Mixed-effects analysis was used to assess differences between baseline and end of treatment (EoT) and between treatments, with control of the false discovery rate ($Q=0.01$); only comparisons where $q < Q$ are plotted (indicated by an asterisk). Green lines indicate treatment comparisons, red lines indicate timepoint comparisons for oil-in-water emollient with glycerine (O/W+G) and blue lines indicate timepoint comparisons for multivesicular emulsion with physiological lipids and glycerine (MVE+GL). Boxes represent the interquartile range, whiskers show the range, median is indicated as a horizontal line and '+' denotes the mean. AdS, α -hydroxy-dihydrosphingosine; AH, α -hydroxy-6-hydroxysphingosine; AP, α -hydroxy-phytosphingosine; AS, α -hydroxy-sphingosine; DAG, diacylglycerol; EO, ester-linked- ω -hydroxy ceramide; EOdS, ω -hydroxy-dihydrosphingosine; EOH, ω -hydroxy-6-hydroxysphingosine; EOP, ω -hydroxy-phytosphingosine; EOS, ω -hydroxy-sphingosine; NdS, non-hydroxy-dihydrosphingosine; NH, non-hydroxy-6-hydroxysphingosine; NP, non-hydroxy-phytosphingosine; NS, non-hydroxy-sphingosine; TAG, triacylglycerol; TEWL₂₀, transepidermal water loss after 20 tape strips.

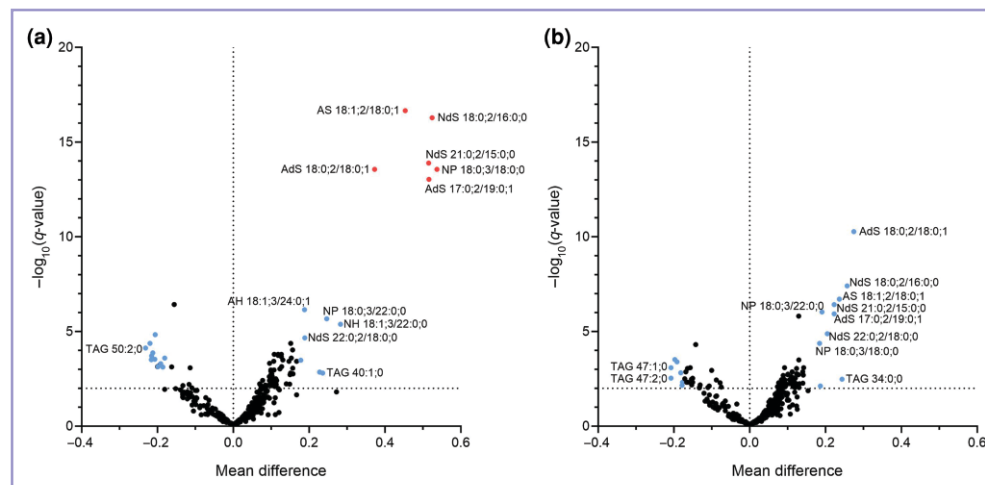


Figure 6 The abundance of lipid species within the stratum corneum changes in response to emollient treatment. Difference in molecular species abundance (\log_{10} pmol mg^{-1} protein) vs. baseline following treatment with (a) multivesicular emulsion with physiological lipids and glycerine (MVE+GL) and (b) oil-in-water emollient with glycerine (O/W+G). Species were included for analysis when detected in >70% of samples in at least one cohort. Differences were assessed using multiple *t*-tests, with control of the false discovery rate ($Q=0.01$). Red points represent species more than 2-fold abundant following treatment, blue points species with more than 1.5-fold change. AdS, α -hydroxy-dihydrospingosine; AH, α -hydroxy-6-hydroxysphingosine; AS, α -hydroxy-sphingosine; Nds, non-hydroxy-dihydrospingosine; NH, non-hydroxy-6-hydroxysphingosine; NP; non-hydroxy-phytosphingosine; TAG, triacylglycerol.

TEWL₂₀. Table 1 provides the top 10 associations (these were all negative; high species abundance associated with better barrier integrity) and change in species abundance with treatment. Notably, six of the species were NH ceramides. Four of the species were increased more than twofold in response to treatment with MVE+GL. As the relative balance of ceramide species appears to be an important determinant of barrier function,^{13,14} we determined the abundance of these four species relative to total NdS ceramides [an increase in NdS ceramides associated with worsening barrier integrity in this population (Williams *et al.*, submitted); Figure S4 a]. Increased abundance of these species relative to NdS was associated with improved barrier integrity, reduced skin sensitivity and greater SC hydration [$r=-0.447$, $r=-0.294$ and $r=0.437$ for NP18:0:3/18:0:0 relative to NdS, respectively (Figure S5; see Supporting Information)].

Discussion

In this study, the effects of two humectant-containing emollients, with and without physiological lipids, were compared in people with dry, eczema-prone skin. Over a 28-day treatment period, both emollients increased skin hydration and improved clinical signs of dryness. The onset of effects was more rapid with the multivesicular emulsion containing glycerine and physiological lipids (MVE+GL) and the water content of the SC was higher (25%). Only MVE+GL was associated with an improvement in the integrity of the skin barrier, and a reduction in sensitivity to the common household irritant SLS (TEWL reduced by 17%). Both treatments appeared to affect the composition of the SC lipidome. The proportion of AP(18) and NP(18) ceramides in the SC, both ingredients of MVE+GL, were increased by 19% and 24%, respectively, following MVE+GL treatment. The relative

Table 1 Changes in the abundance of lipid species correlated with differences in biophysical and functional properties of the skin

Species	Correlation (r), ^a TEWL ₂₀	O/W+G EoT – BL ^b	FC ^c	–Log ₁₀ q-value	Discovery (q < Q)	MVE+GL EoT – BL ^b	FC ^c	–Log ₁₀ q-value	Discovery (q < Q)	Species/NdS correlation (r) ^a TEWL ₂₀
AdS 18:0;2/18:0;1	–0.479	0.27	1.86	10.27	Yes	0.37	2.34	13.56	Yes	–0.510
NH 20:1;3/27:0;0	–0.410	0.035	1.08	0.44	No	0.11	1.29	2.11	Yes	–0.490
NH 18:1;3/27:0;0	–0.374	0.086	1.22	1.19	No	0.044	1.11	0.70	No	–0.496
NH 17:1;3/26:0;0	–0.364	–0.015	0.97	0.25	No	0.016	1.04	0.34	No	–0.516
NH 25:1;3/24:0;0	–0.354	–0.042	0.91	0.66	No	0.071	1.18	1.42	No	–0.500
AdS 17:0;2/19:0;1	–0.349	0.22	1.66	5.93	Yes	0.52	3.31	13.03	Yes	–0.435
NH 18:1;3/25:0;0	–0.342	–0.0014	1.00	0.11	No	0.022	1.05	0.37	No	–0.505
NP 18:0;3/18:0;0	–0.331	0.18	1.51	4.37	Yes	0.54	3.47	13.56	Yes	–0.447
NH 17:1;3/28:0;0	–0.325	–0.016	0.96	0.22	No	0.1	1.26	2.64	Yes	–0.567
AS 18:1;2/18:0;1	–0.313	0.24	1.74	6.71	Yes	0.45	2.82	16.66	Yes	–0.406

Bold text indicates a significant fold change. AdS, α -hydroxy dihydrospingosine; AS, α -hydroxy-sphingosine; BL, baseline; EoT, end of treatment; FC, fold change; MVE+GL, multivesicular emulsion with physiological lipids and glycerine; NdS, non-hydroxy ceramide dihydrospingosine; NH, non-hydroxy-6-hydroxysphingosine; NP, non-hydroxy-phytosphingosine; O/W+G, oil-in-water emollient with glycerine; TEWL₂₀, transepidermal water loss after 20 tape strips. ^aSpearman correlation; ^b \log_{10} pmol mg^{-1} ; ^c $10^{(\text{EoT} - \text{baseline})}$.

a paediatric cohort was not included. All participants had a propensity for eczema; there is no reference for 'healthy' skin due to a focus on an at-risk population in need of barrier restoration. The complexity of ceramide chemistry in the SC is emphasized. The effect of lipid-based treatments on the overall balance of ceramide species is important and so the findings of this work will not be generalizable to other ceramide-based therapies.

This study combined biophysical assessments of skin function with the identification of SC lipid changes at the species level, to evaluate the impact of emollient use on eczema-prone skin. Surprisingly, standard O/W emollient therapy (with glycerine) had no benefit to skin barrier function, despite indications that glycerine levels (and some ceramides) were increased in the SC. Marked changes to the SC lipidome associated with skin barrier strengthening, and a reduced response to irritant challenge, were observed using the MVE+GL. Maintaining a healthy skin barrier and avoiding exposure to irritants, are accepted strategies for managing AD.⁴⁶ In practice, avoidance of common irritants is challenging. Therefore, barrier-strengthening emollients with physiological lipids and glycerine offer benefits over and above emollients containing glycerine only.

Acknowledgements

We thank all of the volunteers who gave up their time to take part in this study. We would also like to acknowledge Lipotype GmbH (Dresden, Germany) for the lipidomic quantification and assistance in analysis. Graphical abstract created with BioRender.

Funding sources

This investigator-led study was supported by a grant from L'Oréal Dermatological Beauty, UK, CeraVe. The funder reviewed the manuscript and approved its submission for publication but was not involved in the study design, data collection or data analysis.

Conflicts of interests

S.G.D. has received fees for giving lectures and/or attending advisory boards and research funding from Almirall, Astellas Pharma, Bayer Dermatology, Hyphens, LEO Pharma, L'Oréal, MSD, Pfizer, Rohto Pharma, Sanofi and Stiefel-GSK. M.J.C. has been/is a clinical trial investigator for the following organizations: Atopix, Galapagos, Hyphens, Johnson & Johnson, Kymab, LEO Pharma, L'Oréal/La Roche Posay, Novartis, Pfizer, Regeneron and Sanofi-Genzyme. He is an advisory board member, consultant and/or invited lecturer for the following organizations: AbbVie, Amlar, Astellas, Atopix, Boots, Dermavant, Galapagos, Galderma, Hyphens, Johnson & Johnson, Kymab, LEO Pharma, L'Oréal/La Roche Posay, Menlo, Novartis, Oxagen, Pfizer, Procter & Gamble, Reckitt Benckiser, Regeneron and Sanofi-Genzyme. The other authors declare no conflicts of interest.

Data availability

The data underlying this article will be shared upon reasonable request to the corresponding author.

Ethics statement

The University of Sheffield Research Ethics Committee approved the study, under project reference 044518. The study was performed in accordance with the Declaration of Helsinki 1964 and its later amendments

Patient consent

Informed consent to participate in the study was obtained from all participants. Written patient consent for publication was obtained.

Supporting Information

Additional [Supporting Information](#) may be found in the online version of this article at the publisher's website.

References

- Menon GK, Cleary GW, Lane ME. The structure and function of the stratum corneum. *Int J Pharm* 2012; **435**:3–9.
- Ishikawa J, Narita H, Kondo N *et al.* Changes in the ceramide profile of atopic dermatitis patients. *J Invest Dermatol* 2010; **130**:2511–14.
- Kováčik A, Šilarová M, Pullmannová P *et al.* Effects of 6-hydroxy-ceramides on the thermotropic phase behavior and permeability of model skin lipid membranes. *Langmuir* 2017; **33**:2890–9.
- Mojumdar EH, Kariman Z, van Kerckhove L *et al.* The role of ceramide chain length distribution on the barrier properties of the skin lipid membranes. *Biochim Biophys Acta* 2014; **1838**:2473–83.
- Nädäban A, Rousel J, El Yachioui D *et al.* Effect of sphingosine and phytosphingosine ceramide ratio on lipid arrangement and barrier function in skin lipid models. *J Lipid Res* 2023; **64**:100400.
- Suzuki M, Ohno Y, Kihara A. Whole picture of human stratum corneum ceramides, including the chain-length diversity of long-chain bases. *J Lipid Res* 2022; **63**:100235.
- Wirén K, Nohlgård C, Nyberg F *et al.* Treatment with a barrier-strengthening moisturizing cream delays relapse of atopic dermatitis: a prospective and randomized controlled clinical trial. *J Eur Acad Dermatol Venereol* 2009; **23**:1267–72.
- Ng JP, Liew HM, Ang SB. Use of emollients in atopic dermatitis. *J Eur Acad Dermatol Venereol* 2015; **29**:854–7.
- Simpson E, Böhlting A, Bielfeldt S *et al.* Improvement of skin barrier function in atopic dermatitis patients with a new moisturizer containing a ceramide precursor. *J Dermatolog Treat* 2013; **24**:122–5.
- Elias PM, Wakefield JS, Man MQ. Moisturizers versus current and next-generation barrier repair therapy for the management of atopic dermatitis. *Skin Pharmacol Physiol* 2019; **32**:1–7.
- Boncheva M, Damien F, Normand V. Molecular organization of the lipid matrix in intact Stratum corneum using ATR-FTIR spectroscopy. *Biochim Biophys Acta* 2008; **1778**:1344–55.
- Danby SG, Andrew PV, Kay LJ *et al.* Enhancement of stratum corneum lipid structure improves skin barrier function and protects against irritation in adults with dry, eczema-prone skin. *Br J Dermatol* 2022; **186**:875–86.
- Rinnov MR, Halling AS, Gerner T *et al.* Skin biomarkers predict development of atopic dermatitis in infancy. *Allergy* 2023; **78**:791–802.
- Yokose U, Ishikawa J, Morokuma Y *et al.* The ceramide [NP]/[NS] ratio in the stratum corneum is a potential marker for skin

- properties and epidermal differentiation. *BMC Dermatol* 2020; **20**:6.
- 15 Danby SG, Chalmers J, Brown K *et al.* A functional mechanistic study of the effect of emollients on the structure and function of the skin barrier. *Br J Dermatol* 2016; **175**:2011–19.
 - 16 Danby SG, Andrew PV, Taylor RN *et al.* Different types of emollient cream exhibit diverse physiological effects on the skin barrier in adults with atopic dermatitis. *Clin Exp Dermatol* 2022; **47**:1154–64.
 - 17 Buraczewska I, Berne B, Lindberg M *et al.* Changes in skin barrier function following long-term treatment with moisturizers, a randomized controlled trial. *Br J Dermatol* 2007; **156**:492–8.
 - 18 Held E, Sveinsdóttir S, Agner T. Effect of long-term use of moisturizer on skin hydration, barrier function and susceptibility to irritants. *Acta Derm Venereol* 1999; **79**:49–51.
 - 19 Palmer CN, Irvine AD, Terron-Kwiatkowski A *et al.* Common loss-of-function variants of the epidermal barrier protein filaggrin are a major predisposing factor for atopic dermatitis. *Nat Genet* 2006; **38**:441–6.
 - 20 Elias PM, Hatano Y, Williams ML. Basis for the barrier abnormality in atopic dermatitis: outside-inside-outside pathogenic mechanisms. *J Allergy Clin Immunol* 2008; **121**:1337–43.
 - 21 Sugarman JL, Fluhr JW, Fowler AJ *et al.* The objective severity assessment of atopic dermatitis score: an objective measure using permeability barrier function and stratum corneum hydration with computer-assisted estimates for extent of disease. *Arch Dermatol* 2003; **139**:1417–22.
 - 22 Berdyshev E, Goleva E, Bissonnette R *et al.* Dupilumab significantly improves skin barrier function in patients with moderate-to-severe atopic dermatitis. *Allergy* 2022; **77**:3388–97.
 - 23 Berardesca E, Distanto F, Vignoli GP *et al.* Alpha hydroxyacids modulate stratum corneum barrier function. *Br J Dermatol* 1997; **137**:934–8.
 - 24 Chilcott RP, Dalton CH, Emmanuel AJ *et al.* Transepidermal water loss does not correlate with skin barrier function in vitro. *J Invest Dermatol* 2002; **118**:871–5.
 - 25 Ananthapadmanabhan KP, Moore DJ, Subramanyan K *et al.* Cleansing without compromise: the impact of cleansers on the skin barrier and the technology of mild cleansing. *Dermatol Ther* 2004; **17**(Suppl. 1):16–25.
 - 26 Egelrud T, Lundström A. The dependence of detergent-induced cell dissociation in non-palmo-plantar stratum corneum on endogenous proteolysis. *J Invest Dermatol* 1990; **95**:456–9.
 - 27 de Jongh CM, Jakasa I, Verberk MM, Kezic S. Variation in barrier impairment and inflammation of human skin as determined by sodium lauryl sulphate penetration rate. *Br J Dermatol* 2006; **154**:651–7.
 - 28 Coderch L, López O, de la Maza A, Parra JL. Ceramides and skin function. *Am J Clin Dermatol* 2003; **4**:107–29.
 - 29 Boer DEC, van Smeden J, Al-Khakany H *et al.* Skin of atopic dermatitis patients shows disturbed β -glucocerebrosidase and acid sphingomyelinase activity that relates to changes in stratum corneum lipid composition. *Biochim Biophys Acta Mol Cell Biol Lipids* 2020; **1865**:158673.
 - 30 Rawlings AV, Harding CR. Moisturization and skin barrier function. *Dermatol Ther* 2004; **17**(Suppl. 1):43–8.
 - 31 Uchida Y. Ceramide signaling in mammalian epidermis. *Biochim Biophys Acta* 2014; **1841**:453–62.
 - 32 Murakami I, Wakasa Y, Yamashita S *et al.* Phytoceramide and sphingoid bases derived from brewer's yeast *Saccharomyces pastorianus* activate peroxisome proliferator-activated receptors. *Lipids Health Dis* 2011; **10**:150.
 - 33 Jiang YJ, Uchida Y, Lu B *et al.* Ceramide stimulates ABCA12 expression via peroxisome proliferator-activated receptor (δ) in human keratinocytes. *J Biol Chem* 2009; **284**:18942–52.
 - 34 Bouwstra JA, Gooris GS, Dubbelaar FE, Ponc M. Phase behavior of stratum corneum lipid mixtures based on human ceramides: the role of natural and synthetic ceramide 1. *J Invest Dermatol* 2002; **118**:606–17.
 - 35 Bouwstra JA, Nädäban A, Bras W *et al.* The skin barrier: an extraordinary interface with an exceptional lipid organization. *Prog Lipid Res* 2023; **92**:101252.
 - 36 Danso M, Boiten W, van Drongelen V *et al.* Altered expression of epidermal lipid bio-synthesis enzymes in atopic dermatitis skin is accompanied by changes in stratum corneum lipid composition. *J Dermatol Sci* 2017; **88**:57–66.
 - 37 Janssens M, van Smeden J, Gooris GS *et al.* Increase in short-chain ceramides correlates with an altered lipid organization and decreased barrier function in atopic eczema patients. *J Lipid Res* 2012; **53**:2755–66.
 - 38 Di Nardo A, Wertz P, Giannetti A, Seidenari S. Ceramide and cholesterol composition of the skin of patients with atopic dermatitis. *Acta Derm Venereol* 1998; **78**:27–30.
 - 39 Jungersted JM, Scheer H, Mempel M *et al.* Stratum corneum lipids, skin barrier function and filaggrin mutations in patients with atopic eczema. *Allergy* 2010; **65**:911–18.
 - 40 Kim J, Kim BE, Goleva E *et al.* Alterations of epidermal lipid profiles and skin microbiome in children with atopic dermatitis. *Allergy Asthma Immunol Res* 2023; **15**:186–200.
 - 41 Janssens M, van Smeden J, Gooris GS *et al.* Lamellar lipid organization and ceramide composition in the stratum corneum of patients with atopic eczema. *J Invest Dermatol* 2011; **131**:2136–8.
 - 42 Berdyshev E, Kim J, Kim BE *et al.* Stratum corneum lipid and cytokine biomarkers at age 2 months predict the future onset of atopic dermatitis. *J Allergy Clin Immunol* 2023; **151**:1307–16.
 - 43 Sho Y, Sakai T, Sato T *et al.* Stratum corneum ceramide profiles provide reliable indicators of remission and potential flares in atopic dermatitis. *J Invest Dermatol* 2022; **142**:3184–91.
 - 44 van Smeden J, Janssens M, Kaye EC *et al.* The importance of free fatty acid chain length for the skin barrier function in atopic eczema patients. *Exp Dermatol* 2014; **23**:45–52.
 - 45 Stahlberg S, Školová B, Madhu PK *et al.* Probing the role of the ceramide acyl chain length and sphingosine unsaturation in model skin barrier lipid mixtures by $(2)H$ solid-state NMR spectroscopy. *Langmuir* 2015; **31**:4906–15.
 - 46 van Zuuren EJ, Fedorowicz Z, Christensen R *et al.* Emollients and moisturisers for eczema. *Cochrane Database Syst Rev* 2017; **2**:Cd012119.